### The integration of Gaussian noise by long-range amygdala inputs in frontal circuit promotes fear learning

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#### 20 ABSTRACT

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Survival depends on the ability of animals to select the appropriate behavior in response to threat and safety sensory cues. However, the synaptic and circuit mechanisms by which the brain learns to encode accurate predictors of threat and safety remain largely unexplored. Here, we show that frontal association cortex (FrA) pyramidal neurons integrate auditory cues and basolateral amygdala (BLA) inputs non-linearly in a NMDAR-dependent manner. We found that the response of FrA pyramidal neurons was

more pronounced to Gaussian noise than to pure 8 kHz tone, and that the activation of BLA-to-FrA axons was the strongest during safe periods in between conditioning pairings. Blocking BLA-to-FrA signaling specifically at the time of presentation of Gaussian noise

30 (but not 8 kHz tone) as a safety signal impaired the formation of auditory fear memories. Taken together, our data reveal a circuit mechanism that facilitates the formation of fear traces in the FrA, thus providing a new framework for probing discriminative learning and related disorders.

#### **INTRODUCTION**

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Discriminative learning is an important survival strategy that depends on the repeated contingency and contiguity between sensory cues (conditioned stimuli, CS) and the events (e.g., danger, safety) that they must predict (unconditioned stimuli, US) (Hall, 2002). It has been classically studied by using differential fear conditioning paradigms where two different auditory CS are positively (CS+) and negatively (CS-) paired in time

- with an aversive US (e.g., foot shock). This learning protocol is taken to assign appropriate 40 emotional valence to the two incoming CSs (Hall, 2002; LeDoux, 2000; Likhtik and Paz, 2015), thereby providing an accurate representation of the environment by increasing discriminative skills between threat and safety signals. Whereas CS+ promotes conditioned fear responses (e.g., freezing behavior) when presented alone, CS- has been shown to serve as a learned safety predictor by reducing fear behavior and increasing 45 positive affective responses (Rogan et al., 2005). While previous work has thoroughly investigated how CS+ generate fear responses (Dejean et al., 2016; Karalis et al., 2016)., it remains unclear whether and how the brain learns to encode CS- and thus discriminates between threat and safety.
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The medial prefrontal cortex (mPFC) has appeared over the past decade as a critical region that shapes behaviors in response to both aversive and non-aversive environmental cues (Likhtik and Paz, 2015; Likhtik et al., 2014; Stujenske et al., 2014). These antagonistic effects of the mPFC possibly develop through specific interaction between its different subdivisions (*i.e.*, prelimbic (PL) and infralimbic (IL) cortices) and 55 the basolateral complex of the amygdala (BLA) (Senn et al., 2014; Sierra-Mercado et al., 2011; Vidal-Gonzalez et al., 2006). However, the mPFC does not receive direct sensory information neither from sensory cortical areas or from the thalamus (Hoover and Vertes, 2007), thereby supporting the idea that a higher-order neuronal network above the mPFC might encode specific memories that are later selected preferentially during recall together with its downstream cortical (e.g., PL or IL mPFC) or subcortical structures (e.g., 60 BLA). Specifically, the superficial frontal association cortex (FrA) has been shown to contribute to memory formation during associative learning (Lai et al., 2012; Nakayama

et al., 2015; Sacchetti et al., 2002). This region of the lateral part of the agranular cortex (AGI) (Paxinos and Watson, 2007; Uylings et al., 2003) receives inputs from the BLA (Lai et al., 2012; Mátyás et al., 2014; Nakayama et al., 2015) and sensory cortices (Hoover and 65 Vertes, 2007; Zhang et al., 2016), and is non-reciprocally connected to the PL/IL subdivisions of the mPFC (Zhang et al., 2016), raising the possibility that it may function as a relay station during learning from sensory cortical areas and the BLA to the mPFC. However, whether and how the FrA integrates the variety of sensory information required for discriminative learning is not understood.

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The involvement of the FrA in auditory fear conditioning has constantly been reported. For example, the pharmacological inactivation of FrA neurons alters both the expression and extinction of learned fear (Lai et al., 2012; Nakayama et al., 2015; Sacchetti et al., 2002). Recently, fear conditioning and extinction have been shown to induce dendritic spine elimination and formation in the FrA, respectively (Lai et al., 2012). Importantly, this phenomenon occurs within the same dendritic branch, supporting the idea that a unique FrA circuit could form memory traces with distinct emotional values. Nonetheless, no previous evidence has demonstrated the contribution of the FrA in the encoding of incoming sensory cues as threat or safety predictors and, if so, how such a process may be controlled by inputs from the BLA.

To address the possible role of the FrA in discriminative learning and the mechanisms behind it, we investigated auditory-evoked computations of layers II/III FrA pyramidal neurons, as well as the dynamics of long-range projections from the BLA during the acquisition and recall of fear memory traces. By using two-photon (2P) calcium imaging in head-restrained mice, *in vivo* whole cell recordings and optogenetics, we found that FrA pyramidal neurons process auditory tones based on their spectral properties. Unlike pure frequency tones, Gaussian noise produced somatic and dendritic depolarizations in FrA pyramidal neurons. The photo-stimulation of BLA-to-FrA neurons resulted in the supra-linear integration of auditory tones. During conditioning, the activity

- 90 of BLA-to-FrA axons was stronger between CS+/US pairings (that is, when CS- is presented) than during pairings. Inhibiting these axons during CS- impaired auditory fear learning but only when Gaussian noise is used as CS-. Taken together, our data support the idea that FrA and BLA-to-FrA neurons gate learning by promoting the integration of non-conditioned Gaussian noise (*i.e.*, not paired to the footshock). In conclusion, the study
- 95 reveals a potent dendritic mechanism for encoding predictors in the FrA, and thus extends the cortical framework for probing discriminative learning and related disorders.

#### RESULTS

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#### 100 Auditory tones recruit NMDARs conductances in FrA L2/3 pyramidal neurons

We performed somatic whole-cell recordings in anesthetized naive animals and characterized the activation of FrA L2 pyramidal cells to different sounds by presenting a pure tone (*8 kHz*) and a broad-band mixture of pure tones (white Gaussian noise, *WGN*) (**Fig. 1**), which have been widely used in previous fear conditioning studies (Dejean et al., 2016; Grosso et al., 2015; Karalis et al., 2016; LeDoux, 2000; Likhtik et al., 2014; Park et al., 2016; Senn et al., 2014; Stujenske et al., 2014). For each recorded cell, membrane potential (Vm) was monitored prior to, during and after the random presentation of both tones, each consisting of 27 pips (50 ms, 0.9 Hz for 30 s) (**Fig. 1B**-F). Membrane potential spontaneously fluctuated between up and down states (**Fig. 1B**). Therefore, to detect changes in Vm specifically induced by the auditory stimulation, we computed the cumulative depolarization over time (cVm), from which was then subtracted the linear regression calculated during the baseline period prior to auditory stimulation (cVm change) (**Fig. 1C, D; Fig. S1A, B**). This allowed us to minimize the variability related to spontaneous activity and thus compare evoked depolarizations under different conditions.

*WGN* evoked a subthreshold depolarization in naive animals (27.6 ± 4 mV, n=22) that lasted for at least 30 sec after the end of the stimulation (32.5± 5 mV, n=22) (Fig. 1C, D). This change in Vm was abolished by the application of the NMDAR antagonist D(-)-2-amino-5-phosphonovaleric acid (dAP5) or the presence of the NMDAR open-channel blocker MK-801 in the intracellular solution (end of the stimulation; control: 27.6 ± 4 mV, n=22; +dAP5: -13.4± 5 mV, n=14; +iMK801: -2.2 ± 2 mV, n=5; p<0.001, *anova*) (Fig. 1D; Fig. S1C-F). This suggests that *WGN* generates NMDAR-mediated depolarization that appears similar to somatic plateau potentials (Gambino et al., 2014).

- In contrast, pure auditory tone (*8 kHz*) did not seem to affect membrane potential either during (*8 kHz*: 1.2 ± 4 mV; *WGN*: 27.6 ± 4 mV; n=22, p<0.001, paired t-test) or after auditory stimulation (*8 kHz*: -3.8 ± 3 mV; *WGN*: 32.5± 5 mV; n=22, p<0.001, *paired t-test*). Nonetheless, we found that *8 kHz* tones hyperpolarized pyramidal neurons in the presence of dAP5 or iMK801 (end of stimulation; control: 1.16 ± 3.9 mV, n=22; +dAP5: -16.7 ± 4 mV, n=10; +iMK801: -21 ± 3.5 mV, n=5; p=0.004, *anova*), thus revealing an
- 130 NMDAR component that was masked under control conditions. (Fig. 1C-F). In fact, *WGN*

and 8 kHz tones hyperpolarized FrA neurons to a similar level under NMDAR blockade (*WGN*: -10.5 ± 3.6 mV, n=19; 8 kHz: -18.2 ± 2.9 mV, n=15; p=0.125, *t-test*; dAP5 and iMK801 conditions pooled together), suggesting that a fraction of the NMDAR conductances recruited by auditory stimulations is masked by non-specific inhibition under control conditions (Doron et al., 2017). However, on average, the NMDAR-mediated component of the evoked cVm change was much larger in response to WGN, indicating that WGN recruited more NMDAR conductances than pure auditory tones (*WGN*: 33.1 ± 4.2 mV; 8 kHz: 9.7 ± 3.4 mV; n=5; p=0.017, paired *t-test*) (**Fig. S1G-J**). Altogether, our data indicate that, during anesthesia, Gaussian noise activates enough NMDAR-mediated synaptic inputs to produce sustained depolarization of the cell body.

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### Auditory tones generate local calcium events in distal FrA dendrites

NMDARs conductances confer unique computational capabilities to pyramidal neurons by operating supra-linear signaling in dendrites (Antic et al., 2010a; Major et al., 2013). These NMDAR-mediated events are highly localized to a small dendritic segment,
but can spread towards the soma to produce plateau potentials (Gambino et al., 2014; Palmer et al., 2014). Thus, we investigated whether auditory tones generate local dendritic events. We infected mice with an AAV9-Syn-flex-GCaMP6s together with a 1:10000 dilution of AAV1-hSyn-cre (Fig. 2A) in order to obtain a sparse labeling with few neurons expressing GCaMP6s. The activity of non-overlapping distal dendritic branches
150 was imaged in superficial layer 1 through an implanted cranial window in anesthetized (139 dendrites from 5 mice; Fig. 2B-D) and awake mice (104 dendrites from 8 mice; Fig. 2B, E, F). We isolated calcium transients and segregated them based on their spatial spread along individual dendrites (Fig. 2C; Fig. S2A-C).

We detected calcium transients in apical dendritic tufts of mice that were
anesthetized with isoflurane (1.5%). These calcium events occurred both spontaneously
(*i.e.* during baseline prior to stimulation) and upon auditory stimulation. The presentation of *WGN* evoked more events (*WGN*: 409; 8 kHz: 139; 2 kHz: 102; p<0.001, *McNemar's χ<sup>2</sup>* test), with a significantly higher number of local dendritic events per dendrite as compared to pure frequency auditory tones (*WGN*: 3.28 ± 0.4; 8 kHz: 1.19 ± 0.3; 2 kHz:
1.26 ± 0.25; n=8; p<0.001, anova) (Fig. 2C, D). 8 kHz and 2 kHz tones did not evoke calcium</li>

transients more frequently than baseline (**Fig. 2D**). Although it remains possible that pure

tones activate basal dendrites, our data demonstrate that, during anesthesia, apical dendrites located in L1 are specifically activated by *WGN*.

- During anesthesia, auditory-evoked calcium events were mostly local, with a full width at half maximum (fwhm: 13.5 ± (s.d.) 13 µm, n=650 events) that fell into the spatial range of NMDAR-mediated spikes(Gambino et al., 2014; Palmer et al., 2014) (**Fig. 2G**). Global calcium transients with a longer spatial extent (fwhm  $\ge$  50µm; 81 ± (s.d.) 35 µm, n=313 events) were additionally observed in awake mice (**Fig. 2E, H**). These events, which could reflect backpropagating somatic action potentials, occurred independently of the nature of the auditory tone (**Fig. S2**). In contrast, most of the local calcium transients were concurrent with *WGN* (*WGN*: 339; *8 kHz*: 218; *2 kHz*: 253; p<0.001, *McNemar's*  $\chi^2$  *test*). Again, and as expected, *WGN* evoked significantly more local events per dendrite than pure tones (*WGN*: 3.37 ± 0.3; *8 kHz*: 2.49 ± 0.2; *2 kHz*: 2.01 ± 0.2; n=5; p<0.001, *anova*)
- (Fig. 2F), thus mirroring what we observed during anesthesia (Fig. 2I). Nevertheless, 8 *kHz* and 2 *kHz* tones generated more local events as compared to baseline (Fig. 2F), suggesting that anesthesia might reduce dendritic signaling of pure frequency auditory inputs (1.22 ± 0.3 *vs.* 2.24 ± 0.2; p=0.037) (Fig. 2I). However, the specific effect of *WGN* in dendrites was not attributable to noise-induced pain, since exploratory motor behaviors were not affected by *WGN* or pure tones (Fig. S3).
- Taken together, our results show that FrA pyramidal neurons process auditory tones differently according to their spectral properties in both anesthetized and awake mice. This occurs at the subthreshold level with Gaussian auditory tones being more efficient in producing somatic, plateau-like depolarizations (**Fig. 1**) and local dendritic events within the same tuft dendritic branch (**Fig. 2**) than pure tones.

#### 185 Non-conditioned Gaussian noise promotes fear memories in FrA circuit

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WGN and pure frequency tones produce distinct forms of synaptic plasticity during auditory fear conditioning (Park et al., 2016). This raises the possibility they might play a specific role during learning (Grosso et al., 2015; Park et al., 2016). To test this hypothesis, we injected mice bilaterally with an AAV expressing the light-activated proton pump *archaerhodopsin* (AAV9.CaMKII.ArchT.GFP, n=40; or AAV9.CamKII.eGFP for controls, n=35) into the FrA. We transiently inhibited the activity of the FrA during learning with light through implanted optical fibers (**Fig. 3A, B**). Auditory fear conditioning (FC) was induced by using a classical discriminative protocol, during which five auditory stimuli

(each consisting of 27 WGN or 8 kHz pips, 50 ms, 0.9 Hz for 30 s) were positively (CS+) or negatively (CS-) paired with the delivery of a mild electrical shock (0.6 mA) to the paws 195 in a pseudorandom order (Fig. S4). The auditory tones (8 kHz and WGN) used for CS+ and CS- during conditioning were counterbalanced across mice (protocol 1, CS+/CS-: 8 *kHz/WGN* respectively; protocol 2, CS+/CS-: *WGN/8 kHz* respectively) (Fig. S4A, B), and learning was tested 24h later during recall by measuring cue-induced freezing in a novel context (Fig. 3B). Mice were classified as learners (learning+) when the learning index was higher than 20% during recall (Fig. S4E, F).

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First, we suppressed the activity of FrA L2/3 neurons by delivering light each time a CS+ was presented during conditioning (Fig. 3C, D). Surprisingly, neither the fraction of mice that learned the cue-shock association (protocol 1, GFP: 75%, n=8, ArchT: 87%, n=8,

- p=0.522; protocol 2, GFP: 87%, n=8, ArchT: 65%, n=8, p=0.248, Pearson's  $\chi^2$  test), nor the 205 percentage of freezing induced by CS+ during recall (GFP: 48.5 ± 5%, n=16; ArchT: 38.5 ± 5%, n=16; p=0.2) were affected by the inhibition of FrA neurons (Fig 3D-F). In contrast, ArchT stimulation during CS- presentation (Fig. 3G, H) significantly decreased the fraction of mice that were conditioned. However, this occurred only when WGN was used
- as CS- (protocol 1, GFP: 75%, n=12, ArchT: 30%, n=13, p=0.02; protocol 2, GFP: 85%, n=7, 210 ArchT: 91%, n=11, p=0.7, *Pearson's*  $\chi^2$  *test*) (**Fig. 3I**). In those mice, the magnitude of CS+ evoked freezing responses was significantly lower during recall than in control mice (protocol 1, GFP: 51 ± 7%, n=12; ArchT: 24.5 ± 5%, n=13; p=0.002, post-hoc Holm-Sidak test) (Fig. 3J).

215 These data demonstrate that when *WGN* is paired negatively to footshock during conditioning, it promotes fear memory traces in the FrA circuit, thus confirming the specific nature of complex auditory cues during learning (Grosso et al., 2015; Park et al., 2016). To further clarify the synaptic mechanisms involved, we measured auditoryevoked dendritic and somatic responses following conditioning protocol 1. As compared

to naïve mice (Fig. 2), WGN (CS-) generated less local dendritic activation during recall 220 (protocol 1; *naive*: 3.37 ± 0.3, n=9; *FC*: 1.88 ± 0.3, n=5 mice; p=0.003) (**Fig. S5**). In contrast, local dendritic events evoked by a conditioned cue (8 kHz) or cues that were not presented during conditioning (2 kHz) were not altered, indicating that fear learning specifically affected WGN-mediated dendritic signaling. In addition, WGN-induced somatic plateau potentials were reduced in conditioned animals in a learning-dependent 225 manner, suggesting that NMDAR-dependent plasticity mechanisms in FrA neurons were engaged during fear learning (Basu et al., 2016; Lai et al., 2012; Rioult-Pedotti et al., 2000) (**Fig. S6**).

#### BLA-to-FrA axons are recruited during fear conditioning

The above mentioned results indicate that the FrA is required for fear learning and 230 guides behaviors by integrating non-conditioned complex auditory cues during conditioning. Given the well-established role of the BLA and its cortical projection during the acquisition and expression of auditory cue fear learning (LeDoux, 2000; Likhtik and Paz, 2015; Likhtik et al., 2014; Nakayama et al., 2015; Senn et al., 2014; Stujenske et al., 2014), we next investigated the information transmitted from the BLA to the FrA (Lai et 235 al., 2012; Mátyás et al., 2014; Nakayama et al., 2015) during conditioning. We injected a virus expressing the genetically encoded calcium indicator GCaMP6f into the right BLA and imaged axonal Ca<sup>2+</sup> responses in the superficial L1 of the right FrA of awake headrestrained mice during fear conditioning (Fig. 4A, B). Results confirmed that BLA neurons project to the superficial layer 1 of the ipsilateral FrA (<150µm) (Lai et al., 2012; Mátyás 240 et al., 2014; Nakayama et al., 2015) (Fig. 4B; see also Fig. S8). We conditioned awake mice (n=10) under the 2-photon microscope by using the same counterbalanced protocols as described previously (protocol 1, n=7; protocol 2, n=3) (Fig. S4). GCaMP6f calcium transients ( $\Delta F/F_0$ ) provided a direct measure of the activation of BLA neurons projecting to the FrA (Fig. 4B). Again, learning was tested 24h later and quantified by the percentage 245 of freezing (Fig. 4D). We then compared the activity of individual boutons between mice that learned (*learning+*, n=5) and those that failed to learn (*learning-*, n=5).

As a first metric to quantify the activity of BLA-to-FrA axons, we measured the number of calcium transients in individual boutons observed during baseline and in conditioning pairings. While the activity of individual BLA boutons in FrA was relatively low at rest, it increased significantly upon successive pairings (**Fig. 4D**). This occurred independently of the conditioning protocol and only in mice that learned (*learning+*, all CS+:  $1.73 \pm 0.3$ , all CS-:  $2.07 \pm 0.45$ , n=5; *learning-*, all CS+:  $0.62 \pm 0.08$ , all CS-:  $0.77 \pm 0.13$ , n=5; p=0.015) (**Fig. 4D-F**; **Fig. S7**). Interestingly, it also never occurred before the end of the first US presentation. Indeed, the number of transients observed during the first CS+ (*i.e.* before the delivery of the first US) was significantly lower than the other CS that were paired with the US (*learning+*, CS/No US:  $1.03 \pm 0.15$ , CS/US:  $1.97 \pm 0.4$ , n=5, p=0.011; *learning-*: CS/No US:  $0.74 \pm 0.05$ , CS/US:  $0.68 \pm 0.08$ , n=5, p=0.863) (**Fig. 4F**). As a

consequence, the activity of boutons measured during the first CS- was always higher than

- during the first CS+ (*learning*+, CS+1: 1.03 ± 0.15, CS-1: 2.03 ± 0.4, n=5, p=0.003; *learning*-260 , CS+1: 0.74 ± 0.05, CS-1: 0.88 ± 0.09, n=5, p=0.425) (arrow in **Fig. 4F**). These data suggest that neither the tone alone nor the foot shock influence the activity of BLA-to-FrA axons. This was corroborated by the observation that BLA-to-FrA axons were never activated by auditory stimulations in naive mice (*i.e.* before fear conditioning) (Fig. S7). Instead, our data support the idea that BLA axons projecting to the FrA convey information about 265 learning, *i.e.* the CS+/US association itself rather than about the nature of the auditory

tones (Nakayama et al., 2015).

Then, we summed the amplitude of all calcium transients detected during each CS presentation (cumulative  $\Delta F/F_0$  averaged across all CS+ or CS-). The averaged cumulative  $\Delta F/F_0$  measured during CS- was always higher than during CS+ (CS-: 0.27 ± 0.07  $\Delta F/F_0$ , 270 CS+: 0.21  $\pm$  0.06  $\Delta$ F/F<sub>0</sub>, n=10; p=0.012), revealing that the overall activity of BLA-to-FrA axons was stronger between conditioning trials, notably at the time when CS- occurred (Fig. 4G). Importantly, the difference between CS+ and CS- related axonal activity ( $\Delta$ cumulative:  $\Delta$ F/F<sub>0 CS-</sub> -  $\Delta$ F/F<sub>0 cs+</sub>) was significantly higher in mice that learned the association than in non-learners (*learning*+:  $0.1 \pm 0.02 \Delta$ cumulative, n=5; *learning*-: 0.01 275 ± 0.008 Δcumulative, n=5; p=0.013, *t-test*) (**Fig. 4G**). In fact, when plotted as a function of freezing percentage observed during recall,  $\Delta$ cumulative correlated positively with learning performance (r<sup>2</sup>=0.89, p<0.001) (**Fig. 4H**), suggesting that the level of activity of BLA-to-FrA axons during CS- is critical for the acquisition of fear memories.

#### Non-linear interaction in FrA L2/3 pyramidal neurons between segregated BLA and 280 auditory inputs

The activation of BLA neurons instructs prefrontal circuits during learning and memory recall (Klavir et al., 2017; Nakayama et al., 2015; Stujenske et al., 2014). However, the optical activation of the BLA alone is not sufficient to produce learned associations (Johansen et al., 2010). Therefore, we hypothesized that the activation of BLA axons 285 during CS- (Fig. 4), along with the synaptic non-linearities evoked by WGN (Fig. 1 and 2), could gate fear learning (Fig. 3) by controlling L2/3 FrA pyramidal neurons through their projections into L1.

To test this hypothesis, we first addressed the properties of BLA-to-FrA synapses in naive mice. We expressed the recombinant light-gated ion channel *channelrhodopsin*-290

2-YFP (ChR2; AAV9-CamKIIa-hChR2-eYFP) in the BLA and performed intracellular recordings in L2/3 FrA neurons from naive mice (Fig. 5). BLA neurons expressing ChR2 projected to the superficial layer 1 of the ipsilateral FrA (<150µm) (Fig. 5A; Fig. S8A-C), thereby most likely contacting dendritic tufts of L2/3 pyramidal neurons. Local photostimulation of ChR2-BLA axons in acute slices produced excitatory postsynaptic 295 current (EPSC) in FrA pyramidal neurons with short latencies  $(3.5 \pm 0.36 \text{ ms}, n=9)$  and low jitter (0.289  $\pm$  0.04 ms, n=9), suggesting that a fraction of BLA neurons are connected monosynaptically to L2/3 FrA pyramidal neurons (Fig. S8D, E) (Klavir et al., 2017). In vivo, the photostimulation of BLA neurons with an implanted optical fiber produced plateau-like depolarizations in all FrA neurons (averaged peak amplitude: 6.2 ± 1.2 300 mV\*sec, full-width at half-max (fwhm): 551 ± 80 ms; n=13). However, BLA-to-FrA inputs were mostly undetectable and became visible only when the stimulation was delivered during down states (Fig. S8F, G). When detected, the distribution of amplitudes across cells suggested that, on average, they were highly variable (range of amplitude: 0.4 mV-14.5 mV) (Fig. S8H). The rhythmic stimulation of ChR2-expressing BLA neurons at 0.9 Hz 305 for 30 sec (27 square pulses, 50ms) (Fig. 5B, C), a protocol that mimicked the pattern of auditory stimuli, generated modest cumulative depolarization (8 ± 4 mV; n=21). Taken together, our data indicate that BLA-to-FrA synapses are likely to be weak and unreliable.

We next investigated the effect of BLA activation during auditory stimulation on L2/3 FrA pyramidal neurons (Fig. 5D-I). We first verified that WGN alone was able to 310 activate FrA pyramidal neurons in mice chronically implanted with optical fibers. Similar to the effect of auditory stimulation in non-implanted mice (Fig. S9A), WGN but not 8 kHz evoked a long-lasting subthreshold depolarization (WGN: 18.9±7 mV, n=11; 8 kHz: -9.7 ± 3 mV, n=8; p=0.005; *t-test*). The coincident photo-activation of BLA neurons during the presentation of WGN resulted in somatic responses (observed cVm) that were 315 significantly higher than the arithmetic sum (expected cVm) of individual depolarizations evoked by the stimulation of BLA or auditory tones alone (observed cVm: 44 ±6.4 mV, expected cVm: 18 ± 8.8 mV, n=11; p=0.002, paired t-test) (Fig. 5D, E, H; Fig. S9B, C). We then plotted the observed vs. expected cVm change and found that the observed cVm exceeded the expected cVm, indicating that the interaction in FrA neurons between BLA 320 and WGN-related inputs was clearly supralinear (Spruston and Kath, 2004; Tran-Van-Minh et al., 2015) (Fig. 5H). Supra-linear operations have been shown to depend on active dendritic conductances (Spruston and Kath, 2004; Tran-Van-Minh et al., 2015).

Accordingly, we found that the application of dAP5 (1mM) to the cortical surface blocked the effect of BLA activation during *WGN* (BLA+*WGN*:  $39 \pm 6.2 \text{ mV}, \text{ n=13};$ 325 BLA+WGN/+dAP5:  $-26.6 \pm 6.7$  mV, n=7; p<0.001; *t-test*), indicating that NMDARs are involved in the supra-linear integration in FrA neurons (Fig. S9F). The photo-activation of BLA also significantly affected the cumulative potential evoked by 8 kHz (8 kHz: 9.7± 3.3 mV vs. 8 kHz+BLA: 16.6 ± 3.1 mV; n=8; p<0.001; paired t-test) (Fig. 5F, G; Fig. S9D, E). Although supra-linear (observed cVm:  $16.6 \pm 3.1$  mV, expected cVm:  $-2.4 \pm 3.3$  mV, n=8; 330 p<0.001, *paired t-test*), the BLA+8 *kHz* integration remained significantly lower than the supra-linearity generated by BLA+*WGN* (observed cVm; *WGN*: 44 ± 6.4 mV, n=11; 8 kHz: 16.6 ± 3.1 mV, n=8; p=0.003, *t-test*). This indicates that BLA axons are necessary to produce non-linear integration of auditory inputs in FrA neurons, which is stronger for Gaussian noise than for pure tones. 335

#### BLA-to-FrA non-linear integration of Gaussian noise gates fear learning

We next examined whether this BLA-mediated, non-linear integration of auditory inputs in FrA pyramidal neurons could play a role in the acquisition of fear memories. This question was addressed by silencing specifically the BLA-to-FrA axons during
conditioning with optogenetics (Fig. 6 and Fig. S10). Mice were injected bilaterally with a retrograde Cav-2-CMV-Cre (Hnasko et al., 2006) into the FrA together with either AAV9. CBA.Flex.ArchT.GFP (ArchT-expressing mice, n=24) or AAV9.CAG.Flex.eGFP (control GFP-expressing mice, n=25) into BLA bilaterally (Fig. 6A). This resulted in the restricted expression of the light-driven inhibitory proton pump ArchT (or GFP for controls) in a
target-specific fraction of BLA neurons that project to the FrA (Fig. 6B-D). Mice were then submitted to auditory fear conditioning, and we analyzed the impact of opto-stimulation on freezing behaviors for each counter-balanced protocol (Fig. S10).

We found that the time-locked suppression of BLA-to-FrA communication during negatively-paired *WGN* (CS-, protocol 1b) significantly decreased the fraction of ArchTmice that learned the association (GFP: 87.5%; ArchT: 22%, p=0.005, *Pearson's*  $\chi^2$  *test*) and freezing behaviors upon subsequent CS+ (8 kHz) presentation (GFP: 55.9 ± 8 %, n=8; ArchT: 34.9 ± 3 %, n=7; p=0.037, *t-test*) (**Fig. 6E, F; Fig. S10**). These results were similar to those obtained when FrA neurons were inhibited during conditioning (**Fig. 3; Fig. S10**), thus confirming that negatively paired *WGN* participates in the formation of fear traces. Surprisingly, however, ArchT-expressing mice showed increased fear responses in response to CS- during recall (GFP:  $5.3 \pm 1 \%$ , n=8; ArchT:  $13.7 \pm 4.5 \%$ , n=7; p=0.04, *t*-*test*) (**Fig. 6F**). In agreement with previous studies(Likhtik and Paz, 2015; Likhtik et al., 2014; Sangha et al., 2013; Senn et al., 2014; Stujenske et al., 2014), this suggests that CS-also acquires safety properties through the activation of a specific population of BLA neurons.

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In contrast, neither the fraction of mice that learned (GFP: 78%; ArchT: 44%, p=0.145, *Pearson's*  $\chi^2$  *test*) nor freezing responses during recall (CS+; GFP: 34.8 ± 3.9, n=9; ArchT: 29.3 ± 8, n=9; p=0.530, *t-test*) were affected by blocking the activity of BLA-to-FrA axons during negatively-paired *8 kHz* (CS-, protocol 2b) (**Fig. 6E, G**). Similar results were obtained when photo-stimulation was administered during CS+ (*WGN*, protocol 2a) (**Fig. 510**). Altogether, our data support the idea that if Gaussian noise is presented when the activity of BLA neurons is the strongest (*i.e.* between conditioning trials, **Fig. 4**), it facilitates discriminative learning.

#### 370 **DISCUSSION**

The present study investigates the role of the BLA-to-FrA circuit in the integration of auditory cues, and how this process participates in the acquisition of fear traces during conditioning. Taken together, our data demonstrate that rather than being an obstacle, Gaussian noise facilitates auditory fear learning when not paired to the foot shock (i.e. delivered between conditioning trials), thereby confirming the sophisticated nature of 375 differential conditioning protocols (Grosso et al., 2015; Hall, 2002; McDonnell and Abbott, 2009). This is likely due to: 1) the specific dendritic and somatic responses of FrA pyramidal neurons to WGN; 2) the activation of BLA-to-FrA axons between conditioning pairings, which might support 3) the non-linear integration of Gaussian noise in FrA 380 neurons. Unlike *WGN*, none of the optogenetic manipulations aiming at altering pure tone processing in FrA and BLA during conditioning had an effect on fear learning. Although our results might also depend on the specific association between pure and Gaussian auditory tones during conditioning, they bring new conceptual perspectives to central questions regarding how frontal circuits contribute to learning, thus going beyond the BLA-mPFC interactions classically described in fear learning studies (Likhtik and Paz, 385 2015).

Accumulating evidence from anatomical and functional studies has demontrated that despite its pivotal role in the acquisition and expression of associations between sensory stimuli and the emotional valence of these stimuli (Laviolette et al., 2005; Roesch et al., 2010), the mPFC is not directly involved in sensory processing (Hoover and Vertes, 390 2007; Martin-Cortecero and Nuñez, 2016; Zhang et al., 2016). In contrast, due to its anatomical connections with distributed cortical and subcortical regions (Hoover and Vertes, 2007; Zhang et al., 2016), the FrA might serve as a hub that coordinates incoming sensory information before reaching the mPFC. Here, we show that it is indeed required for fear learning in a rather unconventional way, and further clarify the underlying 395 synaptic mechanisms. To our knowledge, this is the first demonstration that auditory sensory stimulation produces NMDAR-dependent depolarization in FrA L2/3 pyramidal neurons and activates their dendrites during both anesthesia and wakefulness. These somatic and dendritic events were more pronounced with Gaussian noise than with pure frequency auditory stimulation. Although we cannot rule out that WGN tones are 400 structured and further abstracted throughout the entire auditory system (Deneux et al., 2016), the simplest explanation is that frequency-tuned spines are distributed widely and heterogeneously throughout the same FrA dendrite, as already reported in the auditory cortex of anesthetized mice (Chen et al., 2011). As a consequence, the multiple frequencies
composing *WGN* might promote the activation of a dense pattern of neighboring spines that in turn facilitate the generation and propagation of local non-linear events towards the soma(Antic et al., 2010b). Multiple calcium transients occurring simultaneously in multiple dendritic branches are necessary to affect somatic voltage (Palmer et al., 2014). Both those findings and ours suggest that WGN-induced depolarization is the consequence of multiple calcium events that occur in different dendritic branches of the same neuron. In contrast, pure-frequency tone appeared unable to activate enough branches simultaneously, thereby making the alteration of somatic voltage less probable.

The FrA and the BLA are anatomically interconnected (Lai et al., 2012; Mátyás et al., 2014; Nakayama et al., 2015). However, the functional properties of these connections remain unknown, notably during learning. Our results confirm that BLA neurons project 415 to the superficial layer of the FrA, thereby most likely contacting dendrites of L2/3 pyramidal neurons. The photo-stimulation of ChR2-expressing BLA neurons produced plateau-like depolarizations that were rather weak and unreliable. It thus seems unlikely that they can summate during rhythmic activation to create favorable conditions for the integration of coincident sensory-driven inputs (Antic et al., 2010b; Brandalise et al., 420 2016; Gambino et al., 2014; Xu et al., 2012). Alternatively, it is possible that the modest activation of BLA synapses in FrA apical dendrites facilitates or gates the propagation towards the soma of tone-evoked dendritic events (Jarsky et al., 2005). In agreement, we observed that the coincident activation of BLA-to-FrA inputs increased both WGN and 8 kHz-evoked depolarization non-linearly. Nevertheless, the BLA+WGN nonlinearities 425 appeared to be much stronger than those generated by BLA+8 kHz. Strikingly, however, only *WGN*, but not pure-frequency tone, potentiated FrA pyramidal neurons when combined with the photo-stimulation of BLA-to-FrA inputs. Although this effect was analyzed no longer than 30 sec after the end of auditory stimulation, it prompts the speculation that the additional non-linear dendritic depolarization gains control over 430 WGN-related synapses. Indeed, compelling experimental evidence has demonstrated that non-linear interactions between compartmentalized streams of neural activity induce long-lasting changes in synaptic strength and intrinsic excitability (Dudman et al., 2007; Gambino et al., 2014; Jarsky et al., 2005; Larkum, 2013; McGaugh, 2013; Xu et al., 2012) that could have permanently affected the dynamics of the FrA membrane potential. 435

The BLA presumably transfers information to the FrA that is relevant for fear learning (Lai et al., 2012; Nakayama et al., 2015). Surprisingly, inhibiting the activity of FrA neurons during CS-, or BLA neurons projecting to the FrA, attenuated freezing responses in response to CS+. However, this counterintuitive effect occurred only when *WGN*, but not 8 *kHz* tone, was used as CS- during conditioning (protocol 1), suggesting 440 that the BLA-mediated, non-linear integration of WGN in FrA neurons is critical for the acquisition of fear memory traces. This alteration of learning is unlikely to be the consequence of insufficient activation of the BLA. Instead, given the low number of BLA neurons expressing ArchT and their precise inhibition during CS-, an alternative explanation is that when *WGN* is combined with the activation of BLA projecting axons, 445 which is maximal between conditioning trials, it might promote the representation of sensory cues predicting threat (i.e. CS+, 8 kHz) within the FrA (Hall, 2002). This hypothesis is supported by the late modification of membrane potential fluctuations observed after the activation of BLA with WGN presentation, which occurred when an 8 *kHz* tone was presented during conditioning. In addition, we found that blocking the 450 activity of BLA-to-FrA neurons transiently during CS- also increased freezing behaviors in response to CS- during recall. While these data are consistent with the role of BLA axons in the encoding of CS- as a safety predictor (Likhtik and Paz, 2015; Likhtik et al., 2014; Rogan et al., 2005; Sangha et al., 2013; Senn et al., 2014; Stujenske et al., 2014), they contrast with the effect of FrA inhibition, suggesting that this effect is perhaps dependent 455 on the mPFC (Likhtik et al., 2014; Stujenske et al., 2014), possibly through specific individual axons targeting both the mPFC and the FrA (Mátyás et al., 2014).

BLA neurons send multiple projections to cortical and subcortical areas that have
been shown to project also to the FrA. It is thus possible that the information is
transmitted from the BLA to the FrA through an indirect pathway (Nakayama et al., 2015;
Price, 2003). Here, we demonstrate that the expression of the genetically encoded calcium
indicator GCaMP6 in BLA neurons makes it possible to monitor optically the activity of
target-specific BLA axons during learning. Using this strategy in awake mice, we
demonstrate for the first time that BLA-to-FrA axons are progressively recruited upon
successive conditioning trials, thereby ruling out the indirect activation of the FrA circuit.
First, we show that BLA-to-FrA axonal activity is never affected by the presentation of
auditory cues alone. Our data contrast with previous work showing an increase in local
field potential and unit activity in the BLA upon auditory stimulation (Collins and Paré,

2000), and suggest instead the existence of a subpopulation of BLA neurons projecting specifically to the FrA that might play a specific role during the learning of emotion. 470 Indeed, these axons are activated only after the first CS+/US pairing, and thus seem to transmit integrated information about the association itself (Nakayama et al., 2015). The level of activity of BLA-to-FrA axons was stronger between conditioning trials (*i.e.* during CS- presentation) and correlated with learning performance. This suggests a putative Hebbian-like frontal mechanism that could integrate any Gaussian noise that is contiguous 475 to the maximal activation of BLA-to-FrA axons (Johansen et al., 2010, 2014; Larkum, 2013; Nakayama et al., 2015). While this mechanism would occur only if WGN is presented between conditioning trials (and thus used as CS-), it might eventually facilitate the recruitment of neurons into specific cue memory traces. In agreement, we found that auditory fear conditioning significantly decreased the number of WGN, but not 8 kHz-480 evoked local dendritic transients. Dendritic plateau potentials have been shown to regulate synaptic strength and synaptic plasticity (Cichon and Gan, 2015; Du et al., 2017; Gambino et al., 2014; Humeau and Lüthi, 2007; Palmer et al., 2014), which might subsequently facilitate the stabilization or pruning of synaptic inputs during learning (Holtmaat and Caroni, 2016; Li et al., 2017). In support of the latter, the level of fear 485 learning has been shown to correlate with the percentage of spine elimination in FrA (Lai et al., 2012) which possibly explains the negative relation we observed during anesthesia

Collectively, our data reveal the specific properties of Gaussian noise in FrA during 490 fear conditioning. The question arises as to the function or benefit of Gaussian noise in the BLA-to-FrA circuit during learning. Previous studies highlighted the critical function of the BLA in attention for learning (Laviolette et al., 2005; Roesch et al., 2010). Here, we show that the activation of BLA-to-FrA axons is independent of the nature of the CS presented. It thus seems unlikely that BLA-to-FrA axons convey the emotional valence of this association. In addition, the activation of BLA alone, while necessary, is not sufficient 495 to trigger learning (Johansen et al., 2010, 2014). BLA neurons might signal to frontal circuits any new association independently of its valence, which might subsequently be assigned by the mPFC (Klavir et al., 2013; Likhtik and Paz, 2015). While acoustic noise is often viewed as a disturbing variable, it can enhance signal processing, facilitate sensory signaling and improve cognitive performance, notably in individuals with poor attention 500 (McDonnell and Abbott, 2009; McDonnell and Ward, 2011; Stein et al., 2005). Therefore,

between *WGN*/(CS-)-evoked subthreshold depolarizations and the strength of learning.

given that noise is abundant in the environment and communication of most mammals, it might facilitate learning in coordination with BLA-to-FrA inputs.

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#### **AUTHOR CONTRIBUTIONS**

MA, EA, VK, NC performed the experiments. MA and FG conceived the studies and analyzed the data with the help of NC and EA. FG supervised the research and wrote the manuscript with help from MA, EA and NC.

#### **DECLARATION OF INTERESTS**

The authors declare no competing financial interests.

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### **FIGURES AND LEGENDS**

Figure 1. Gaussian auditory stimulation depolarizes FrA pyramidal neurons

Figure 2. Gaussian auditory stimulation evokes more local dendritic events than pure frequency tones in anesthesia and awake

Figure 3. The FrA is engaged during fear learning when WGN is used as a CS-

Figure 4. The activation of BLA-to-FrA axons between conditioning trials predicts the level of auditory fear learning

Figure 5. The activation of BLA-to-FrA axons supports the non-linear integration of auditory tones

Figure 6. BLA-to-FrA projecting neurons are required for auditory fear learning when WGN is used as a CS-

Fig. S1. dAP5 application impacts spontaneous and auditory-evoked FrA membrane potential changes

705 **Fig. S2. Auditory-evoked calcium events in dendritic tufts are similar between brain** states

Fig. S3. Auditory tones do not affect locomotor activity.

Fig. S4. Effect of FrA opto-inhibition during conditioning on freezing responses and learning during recall

710 Fig. S5. Fear conditioning (protocol 1) specifically decreases WGN-induced local dendritic events

Fig. S6. Fear conditioning (protocol 1) occludes WGN-evoked somatic plateau potentials

Fig. S7. BLA-to-FrA axons do not encode the nature nor the valence of auditory tones

Fig. S8. BLA-mediated synaptic inputs onto FrA L2/3 pyramidal neurons

Fig. S9. The activation of BLA-to-FrA axons supports the non-linear integration of auditory tones

Fig. S10. Statistical comparisons between all behavioral optogenetic experiments

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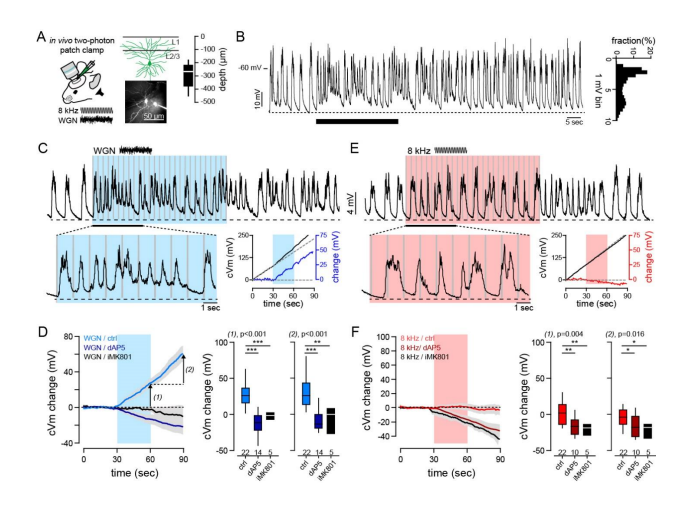


Figure 1. Gaussian auditory stimulation depolarizes FrA pyramidal neurons
 A) FrA pyramidal neuron membrane potential (Vm) was recorded from L2/3 FrA pyramidal under urethane anesthesia with 2P visual guidance. Depth of recorded cells is indicated.

**B)** Example of recorded FrA neuron showing typical spontaneous slow wave fluctuations.

## 730 Black bar indicates timing of Gaussian stimulation. *Right*, membrane potential probability histogram.

**C)** Example traces of postsynaptic membrane potential recorded from an individual FrA L2/3 pyramidal neuron upon white Gaussian noise (*WGN*) auditory stimulation (gray bars: 27 pips, 50 ms in duration, 0.9 Hz, 30 s). *Bottom right panels*, auditory-evoked cVm changes that were calculated by subtracting the sumulation Vm (aVm, calculated by subtracting the sum value) by subtracting the sum value (aVm, calculated by subtracting the sum

- changes that were calculated by subtracting the cumulative Vm (cVm, solid black line) by its linear regression during the baseline period (solid black line).
  D) *Left*, averaged cVm change (± sem) evoked by *WGN*, with or without the blockade of NMDARs (dAP5 or iMK801). Auditory stimulation is depicted by the blue bar; arrows, analysis time points 1 (end of stimulation) and 2 (30 sec later). *Right*, effect of dAP5 and
- iMK801 on cVm change at time points 1 (left, p<0.001, *one way anova*) and 2 (right, p<0.001, *one way anova*). Boxplots represent median and interquartile range (Ctrl, n=22, dAP5, n=14, iMK801, n=5; \*\*\*, p<0.001, \*\*, p<0.01, Holm-Sidak multiple comparisons).</li>
  E, F) Same representation as (C, D) but for pure 8 kHz-evoked cVm change (Ctrl, n=22, dAP5, n=10, iMK801, n=5; p=0.04 and 0.016, *one way anova*; \*\*, p<0.01, \*, p<0.05, Holm-</li>
- *Sidak comparisons*). The effect of both stimuli was tested on the same cell.

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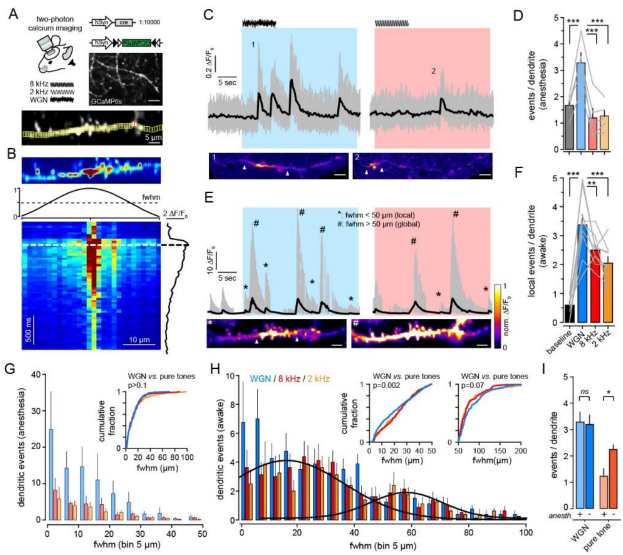


Figure 2. Gaussian auditory stimulation evokes more local dendritic events than pure frequency tones in anesthesia and awake

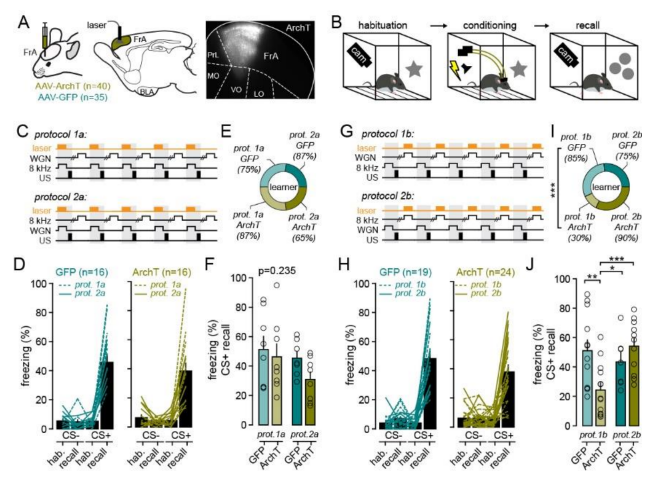
- A) *Top*, experimental strategy. Mice were double-infected with AAV9.Flex.GcaMP6s and 1/10000 diluted AAV1.hSyn.Cre viruses resulting in sparse expression of GCaMP6s. Scale bar, 10 μm. Ca<sup>2+</sup> events in individual dendrites were then imaged in L1 with 2P upon auditory stimulation (*WGN*, 8 kHz and 2 kHz) in awake or anesthetized mice. *Bottom*, example of GCaMP6s fluorescence standard deviation image with ROIs in yellow. Red ROIs overlapped with another dendrite, and were hence excluded from the analysis.
- **B)** The spread of  $Ca^{2+}$  events was quantified by calculating the full-width at half-max (fwhm, dashed line) of the normalized Gaussian fit at the time when the averaged  $\Delta F/F_0$  was maximal.
- **C)** *Top*, example of multiple Ca<sup>2+</sup> transients ( $\Delta F/F_0$ ) recorded in a single dendrite in anesthetized mouse, during baseline and upon *WGN* (blue) and *8kHz* (red). Grey lines,  $\Delta F/F_0$  measured from small ROIs (all ROIs superimposed). Black line, mean  $\Delta F/F_0$ averaged over all ROIs, respectively. *Bottom*, examples of local dendritic Ca<sup>2+</sup> events upon *WGN* (1) and *8 kHz* (2). Scale bar, 5µm.
- **D)** The number of all Ca<sup>2+</sup> events observed during baseline and upon auditory stimulations was averaged over 5 anesthetized mice (p<0.001, *one way repeated measures anova*; \*, p<0.05, \*\*\*, p<0.001, *Holm-Sidak multiple comparisons*). Grey lines represent individual mice.

**E)** *Top*, same representation as **(C)** but in awake mouse. *Bottom*, example of local (\*) and global (#) dendritic Ca<sup>2+</sup> events. Scale bar, 5µm.

**F)** Same presentation as in (**D**) but for local events from 9 awake mice (p<0.001, *one way repeated measures anova*; \*, p<0.05, \*\*, p<0.01, \*\*\*, p<0.001, *Holm-Sidak multiple comparisons*). Gray lines represent individual mice.

**G**, **H**) Distribution histogram of dendritic events fwhm upon *WGN* (blue), *8 kHz* (red) and *2 kHz* (orange) tone presentation, from anesthetized (**G**) and awake (**H**) mice. *Inset,* cumulative probability histograms of auditory-evoked events fwhm.

cumulative probability histograms of auditory-evoked events fwhm.
I) Dendritic events evoked by *WGN* and pure tones (8 *kHz* and 2 *kHz* tones pooled together) in anesthetized (+) and awake (-) mice.



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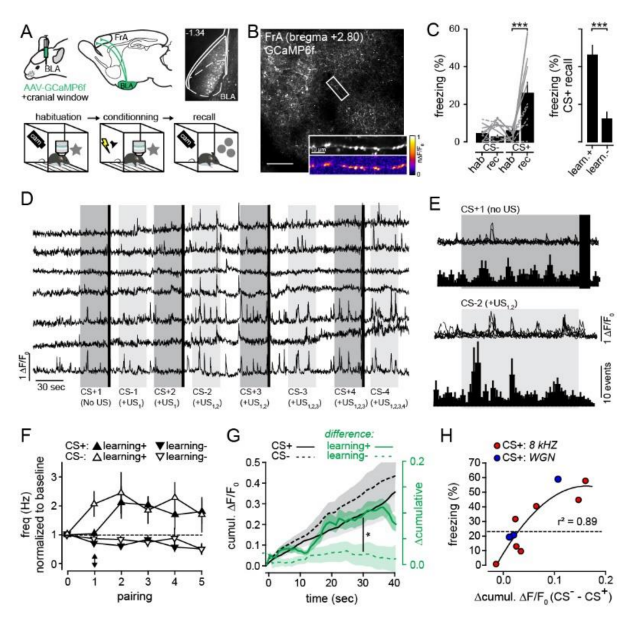
#### Figure 3. The FrA is engaged during fear learning when WGN is used as a CS-

**A)** *Left*, experimental strategy. Mice were infected with AAV9.CaMKII.ArchT (n=40) or AAV9.CaMKII.eGFP for controls (n=35) in FrA, and implanted bilaterally with optical fibers in FrA. Right, expression profiles of ArchT in the FrA (scale bar, 50 μm). PrL, *prelimbic cortex*; MO, VO, LO, *medial, ventral and lateral orbitofrotal cortex* respectively.

B) Timeline of auditory fear conditioning behavioral protocol, with habituation and fear conditioning done in one context and fear learning quantified 24 h later in a new context.
C) Experimental fear conditioning protocols. FrA neurons expressing ArchT (or GFP) were photo-stimulated during the presentation of CS+ (*8 kHz* for protocol 1a or *WGN* for protocol 1b). US, unconditional stimulus (footshock).

**D)** Effect of light during learning on freezing behaviors during recall as compared to habituation (hab.) in GFP-expressing mice (left; n=16) and ArchT-expressing mice (right, n=16). Turquoise and khaki lines represent individual mice.

Fraction of mice that learned the cue-shocked association for each protocol.
 F) Freezing behaviors quantified during CS+ in GFP- and ArchT-expressing mice (p=0.235, *anova*). Circles, individual mice.
 G-J) Same representation as (C-F) but for experiments with photo-stimulations delivered during CS-.



## 805 Figure 4. The activation of BLA-to-FrA axons between conditioning trials predicts the level of auditory fear learning

**A)** *Top left,* experimental protocol. BLA neurons were transfected with AAV1.Syn.GCaMP6f and their boutons were imaged in the superficial layer of the FrA. *Top Right,* expression profiles of GCaMP6f in the BLA. *Bottom,* GCaMP6f-expressing mice were fear conditioned under the 2P microscope (both protocols pooled together), and fear

fear conditioned under the 2P microscope (both protocols pooled together), and fear learning was quantified 24 h later in a new context.
B) GCaMP6f-expressing axons were imaged in the FrA of awake mice through a cranial window (scale bar, 200 μm). *Inset*, Example of Ca<sup>2+</sup> transients in individual BLA-to-FrA axon.

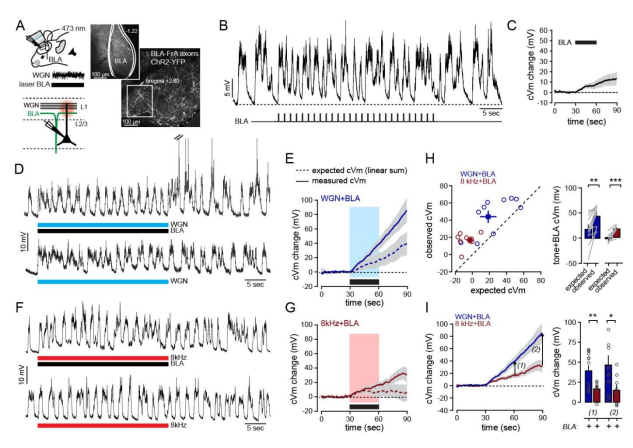
- **C)** *Left*, conditioning under the microscope induced robust fear behaviors (n=10; p=0.007, *two-way repeated measures anova*): in contrast to CS-, CS+ increased fear freezing responses during recall as compared to habituation (hab.) (\*\*\*, p<0.001, *Holm-Sidak multiple comparisons*). Gray solid lines, % of freezing of mice with learning index >20% (learning+, n=5); gray dashed lines, % of freezing of mice with learning index <20%
- 820 (learning-, n=5). *Right*, freezing responses induced by CS+ during recall for learners (learn. +) and non-learners (learn. -) mice (p<0.001, *t-test*).

**D)** Examples of Ca<sup>2+</sup> transients ( $\Delta F/F_0$ ) from individual boutons recorded from one mouse upon consecutive CS+ / US pairings. Dark grey bars, CS+; light grey bars, CS-; black bars, footshock (US).

**E)** Examples of axonal Ca<sup>2+</sup> transients (*top*) and averaged histograms of distribution (± sem) (*bottom*), for the first CS+ presentation (CS+1) and the second CS- presentation (CS-2).

**F)** Frequency of axonal Ca<sup>2+</sup> transients recorded during successive CS. For the first CS+, the activity of boutons was monitored in absence of footshock. Black arrow points to the difference between the first CS+ (before the first footshock) and the first CS- (after the

line, n=5) or not (learning-, dotted green line, n=5) the association. \*, p=0.013, *t-test*.
H) Relation between the Δcumulative during conditioning and the % of freezing during recall for protocol 1 (blue, CS+: 8 kHz) and protocol 2 (red, CS+: WGN). Circles, mice.



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# Figure 5. The activation of BLA-to-FrA axons supports the non-linear integration of auditory tones

**A)** *Left,* co-activation protocol. ChR2-expressing BLA neurons were photo-stimulated during auditory stimulation. Right, representative example of the ChR2-GFP expression profile in the mouse BLA and FrA (ChR2-expressing axons were imaged in superficial layer 1 with 2P microscopy before recording).

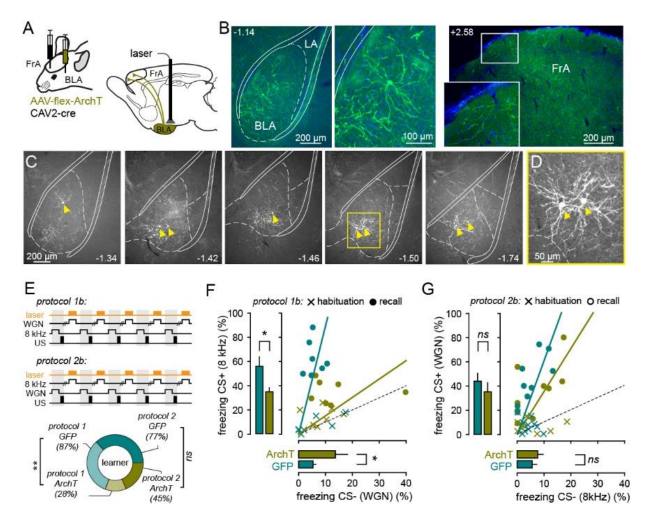
**B)** Single-cell example of depolarizations in FrA by the rhythmic photostimulation of BLA neurons (27 square stimulations, 50 ms, 0.9 Hz for 30 s).

**C)** Averaged cVm change (± sem) evoked by the photostimulation of BLA neurons. Photostimulation is depicted by the black bar.

**D)** Example of traces of postsynaptic membrane potential recorded from individual FrA L2/3 pyramidal neurons upon *WGN* paired (*top*) or not (*bottom*) with the photostimulation of ChR2-expressing BLA neurons. Black and blue bars below the traces indicate the duration of the stimulation.

**E)** Averaged cVm change (± sem) observed upon paired stimulation (solid line) or expected from the arithmetic sum of individual depolarizations evoked by the stimulation of BLA or auditory tones alone (dotted line). Blue and black bars, auditory and BLA stimulations, respectively.

- F, G) Same representation as in (D, E) but for 8 kHz.
  H) Left, relation between observed and expected cVm. Circles, individual cells, squares, mean ± sem. Right, Averaged cVm (± sem). \*\*, p<0.01; \*\*\*\*, p<0.001; paired t-test. Gray lines indicate paired experiments.</li>
- I) *Left*, averaged observed cVm change (± sem). Arrows, analysis time points 1 (end of stimulation) and 2 (30 sec later). *Right*, effect of photostimulation (+) on *WGN* (dark blue) and 8 kHz (black red)-evoked cVm at time points 1 and 2. \*\*, p=0.003; \*, p=0.035; *t-test*. Circles, individual cells.



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## Figure 6. BLA-to-FrA projecting neurons are required for auditory fear learning when WGN is used as a CS-

**A)** Experimental strategy. Mice were infected with AAV9.CBA.Flex.ArchT-GFP (or AAV9.CAG.Flex.eGFP for controls) in BLA and CAV2.CMV.CRE in FrA, and chronically implanted bilaterally with optical fibers in BLA.

**B)** Example of expression profile of ArchT in BLA (left) and FrA (right). BLA, basolateral amygdala; LA, lateral amygdala.

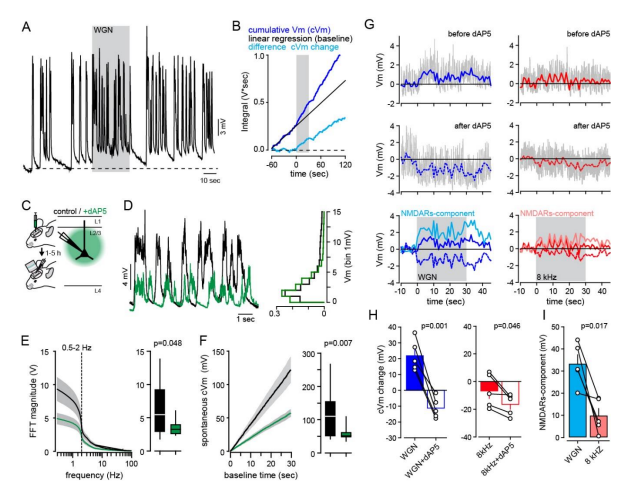
**C**, **D**) Few neurons expressing ArchT were found throughout the entire BLA, but not the LA.

**E)** *Top*, experimental fear conditioning protocols. FrA neurons expressing ArchT (or GFP) were photo-stimulated during the presentation of CS- (*WGN* for protocol 1b or *8 kHz* for protocol 2b). US, unconditional stimulus (footshock). *Bottom*, fraction of mice that learned the cue-shocked association for each protocol. \*\*, p<0.01; ns, non-significant; *Pearson*  $\chi^2$  *test*.

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**F)** Effect of light on freezing responses during recall upon 8 kHz (CS+ protocol 1b; GFP vs ArchT, p=0.037, *t-test*) and *WGN* (CS- protocol 1; GFP vs ArchT, p=0.04, *t-test*) presentation. The relation between 8 kHz and WGN-evoked freezing during habituation (crosses) and recall (dotes) is also indicated.

**G)** Same presentation as in (**F**) but for *WGN* (CS+, protocol 2b) and *8 kHz* (CS-, protocol 2b). ns, p>0.05, *t-test*.



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## Fig. S1. dAP5 application impacts spontaneous and auditory-evoked FrA membrane potential changes

A) Single-cell example of typical spontaneous slow wave fluctuations.

**B)** WGN-evoked cVm change (light blue) is calculated by subtracting the cumulative Vm (cPSP, blue) by its linear regression during the baseline period (black line).

**C)** Experimental protocol: dAP5 (1 $\mu$ m) was applied through the dura and L2/3 pyramidal neurons were patched 1 to 5 hours later.

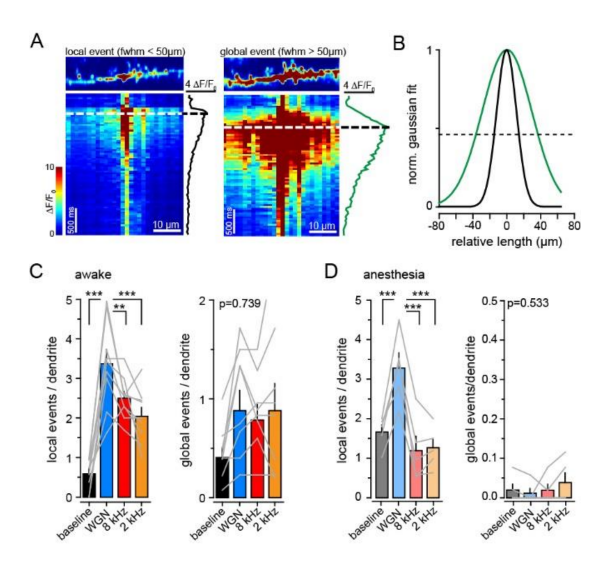
**D)** *Left*, examples of single-cell spontaneous membrane potential (Vm) fluctuations during anesthesia in controls (black) and upon dAP5 injection (green). *Right*, corresponding membrane potential probability histograms.

- 905 corresponding membrane potential probability histograms.
   E) *Left*, average (± sem) fast Fourier transform (FFT) of spontaneous membrane potentials in controls (black) and upon dAP5 injection (green). *Right*, blocking NMDARs significantly decreases the 0.5-2 Hz range (p=0.048, *Mann-Whitney rank sum test*). Box plot represents median and interquartile range (control, n=22; dAP5, n=19).
- **F)** *Left*, average (± sem) cumulative PSP (cPSP) of spontaneous membrane potentials in controls (black) and upon dAP5 injection (green). *Right*, blocking NMDARs significantly decreases cVm amplitude after 30 sec (p=0.007, *Mann-Whitney rank sum test*). Box plot represents median and interquartile range (control, n=22; dAP5, n=19).

**G)** Single cell example of Vm fluctuation upon WGN (left) and 8 kHz (right) before and

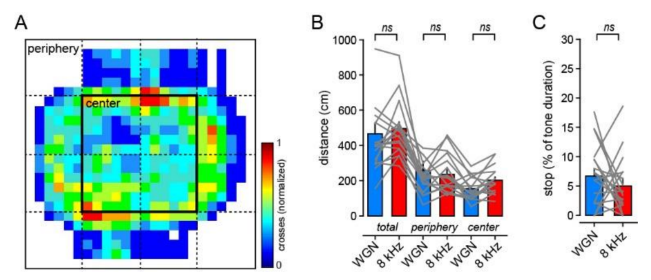
915 after dAP5 application. Colored bars represent averaged Vm over 1 sec time window. The NMDARs-mediated component is calculated by subtracting the averaged Vm before and after dAP5 application.

**H**, **I)** Average cVm (± sem) change before and after dAP5 application. Circles and lines, paired cells.



### Fig. S2. Auditory-evoked calcium events in dendritic tufts are similar between brain states

- **A)** *Top*, example of single-dendrite local (*left*) and global (*right*) Ca<sup>2+</sup> events. *Bottom*, raster plot of  $\Delta F/F_0$  for multiple ROIs along the dendrite represented above. *Right*, average  $\Delta F/F_0$ . Dashed line, time point of the maximum peak amplitude of averaged  $\Delta F/F_0$ . **B)** Gaussian fits (normalized to max) of the local (black) and global (green) Ca<sup>2</sup> events shown in (**A**).
- 930 **C, D)** number of local (left) and global (right) Ca<sup>2+</sup> events observed during baseline and upon auditory stimulations in awake (**E**) and anesthetized (**F**) mice (n=9 and 5, respectively). p<0.001, *one way repeated measures anova*; \*\*, p<0.01, \*\*\*, p<0.001, *Holm-Sidak multiple comparisons*). Grey lines represent individual mice.



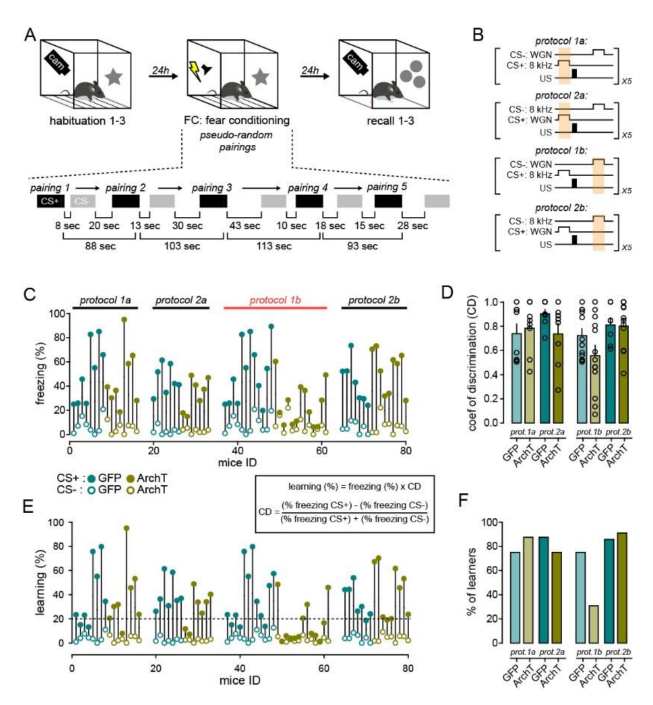
### Fig. S3. Auditory tones do not affect locomotor activity.

**A)** Heatmap representation of mouse activity over 12 minutes. Each auditory tone was presented 4 times for 30 sec, every 10 to 60 seconds.

**B)** Averaged distance travelled (total, in the periphery, and in the center of the open field) upon *WGN* and *8 kHz* tone presentation. Gray lines, individual mouse (n=17). ns, non-significant, *post-hoc comparison after repeated measures anova*.

**C)** Averaged stop duration upon tone presentation. Gray lines, individual mouse (n=17). ns, *paired t-test*.

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## Fig. S4. Effect of FrA opto-inhibition during conditioning on freezing responses and learning during recall

A) Timeline of behavioral experiments. Random delays (between 10 and 60 seconds) between CS were used and one example is indicated below.

**B)** Fear conditioning protocols. FrA neurons expressing ArchT (or GFP) were photostimulated during the presentation of CS+ (*8 kHz* for protocol 1a or *WGN* for protocol 2a) or CS- (*WGN* for protocol 1b or *8 kHz* for protocol 2b). US, footshock. Photo-simulations (orange box) were given during CS+ (protocols 1a and 2a) or CS- (protocols 1b and 2b).

(orange box) were given during CS+ (protocols 1a and 2a) or CS- (protocols 1b and 2b).
 C) Freezing responses during recall upon CS+ (filled circles) and CS- (open circles) in GFP (turquoise) and ArchT (khaki) expressing mice. Significant differences between GFP and ArchT-mice were seen only for protocol 1b (red).

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**D)** Coefficient of discrimination (CD) was calculated as follows: CD = (CS+ % freezing- CS- % freezing) / (CS+ % freezing+ CS- % freezing). Circles, individual mice.

**E)** Learning index was quantified for each CS by multiplying the % of freezing in each condition by the corresponding coefficient of discrimination. Mice with a learning index below 20 % (dotted line) were considered as non-learners.

**F)** fraction of mice that learned the cue-shock association (learning>20%).

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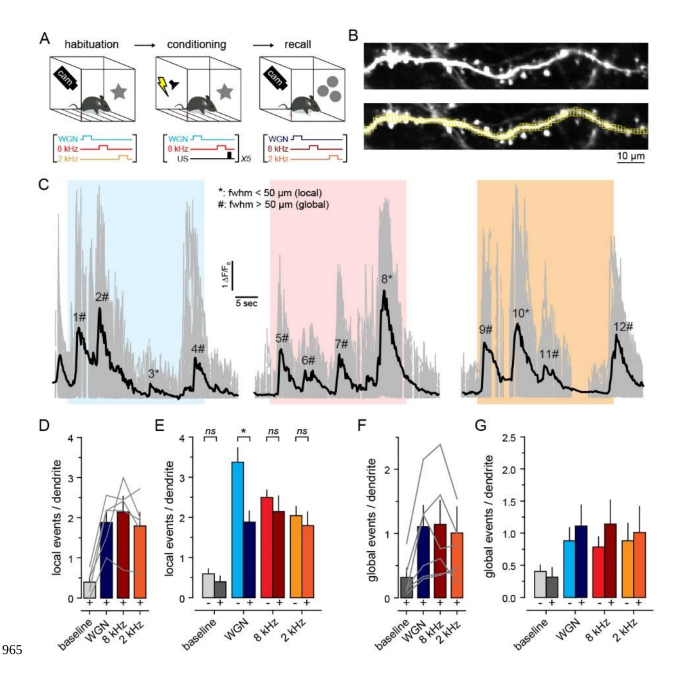


Fig. S5. Fear conditioning (protocol 1) specifically decreases WGN-induced local dendritic events

A) *Top*, Experimental protocol. GCAMP6s-expressing dendrites were imaged in awake
 mice before and after fear conditioning during presentation of *WGN*, *8 kHz* and *2 kHz* tones. *Bottom*, fear conditioning protocol. Only *WGN* and *8 kHz* were used during conditioning, while *2 kHz* served as a neutral tone that was only presented before and after conditioning.

**B)** Example of GCaMP6s fluorescence standard deviation projection (top) with ROIs in yellow (bottom).

**C)** Example of multiple Ca<sup>2+</sup> transients ( $\Delta F/F_0$ ) recorded in a single dendrite, upon *WGN* (blue), *8 kHz* (red) and *2 kHz* (orange). Grey lines,  $\Delta F/F_0$  measured from small ROIs (all ROIs superimposed). Black line, mean  $\Delta F/F_0$  averaged over all ROIs, respectively.

**D)** Examples of dendritic Ca<sup>2+</sup> events upon *WGN* (left, blue), *8 kHz* (middle, red), and *2 kHz* (right, orange). The spread of Ca<sup>2+</sup> events was quantified by calculating the full-width at

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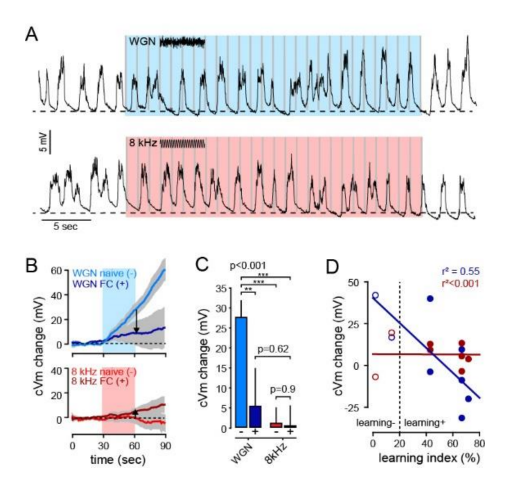
half-max (fwhm) of the normalized Gaussian fit (colored line) at the time when the averaged  $\Delta F/F_0$  was maximal.

**E**, **F**) Number of local Ca<sup>2+</sup> events per dendrite observed during baseline and upon auditory stimulations in awake mice before (-, n=9) and after (+, n=5) fear conditioning. The averaged number of local dendritic events per dendrite observed upon *WGN* presentation in awake mice was significantly lower after fear learning (*WGN*-: 3.37 ± 0.3, n=9 naive mice; *WGN*+: 1.88 ± 0.3, n=5 conditioned mice; p=0.025, *t-test*).

**G**, **H**) Same presentation as in (**E**, **F**) but for global events.

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# Fig. S6. Fear conditioning (protocol 1) occludes WGN-evoked somatic plateau potentials

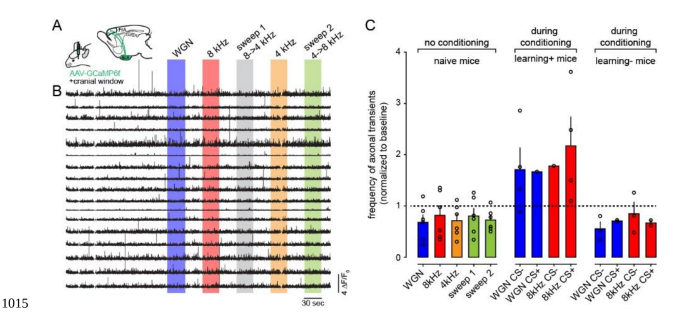
A) Example traces of postsynaptic membrane potential recorded from an individual FrA pyramidal neuron upon *WGN* (CS-, top) and *8kHz* (CS+, bottom) auditory stimulation after fear conditioning. The effect of both stimuli was tested on the same cell.
 B) Averaged cVm change (+ sem) evoked by *WGN* (top) and *8kHz* topes (bottom) before

**B)** Averaged cVm change (± sem) evoked by *WGN* (top) and *8kHz* tones (bottom) before (naïve (-)) and after fear conditioning (FC (+)).

- C) Effect of WGN and pure 8kHz tone on cVm change before (-, n=22) and after learning (+; n=8). As compared to naive mice, WGN failed to activate FrA pyramidal neurons in conditioned mice (WGN+: 5.4 ± 9 mV, n=8; WGN-: 27.6 ± 4 mV, n=22; p=0.008). Thus, the difference between WGN and 8 kHz-induced cVm change observed in naive mice (14 naive mice; WGN-: 27.6 ± 4 mV; 8kHz-: 1.1 ± 4 mV; n=22; p<0.001; paired t-test) disappeared</p>
- after fear conditioning (5 conditioned mice; *WGN+*: 5.4 ± 9 mV; 8 kHz+: 0.5 ± 5 mV; n=8; p=0.698; paired t-test). p<0.001, one-way anova; \*\*\*, p<0.001, \*\*, p<0.01, Holm-Sidak comparisons.

**D)** Relation between the learning index and WGN (blue)- and 8kHz (red)-induced cVm change after conditioning. As opposed to 8kHz-cVm change that was not different

1010 between behavioral performance ( $r^2 < 0.001$ ), *WGN* induced stronger cVm change in low freezing mice indicating that WGN- somatic plateaus are negatively correlated with behavioral performance (*WGN*:  $r^2 = 0.55$ ; *8 kHz*:  $r^2 < 0.001$ ). Open / filled circles, individual cells from non-learning / learning mice, respectively. bioRxiv preprint doi: https://doi.org/10.1101/569137; this version posted February 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

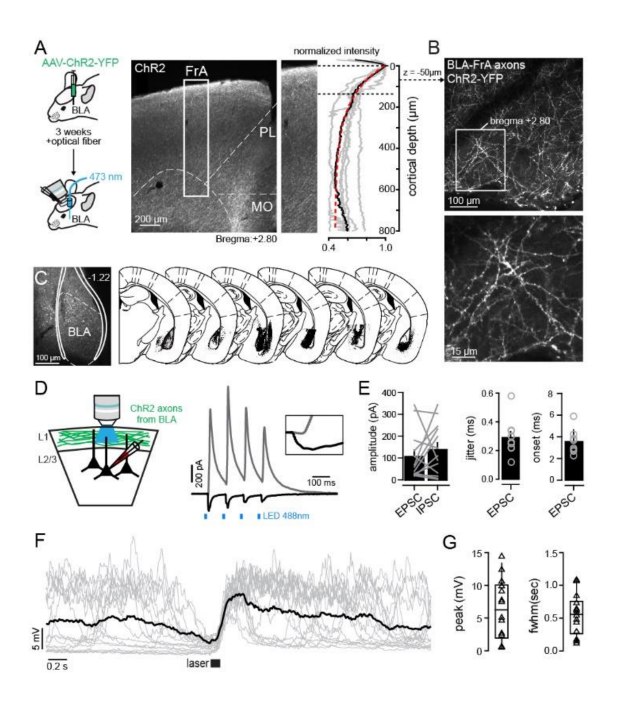


# Fig. S7. BLA-to-FrA axons do not encode the nature nor the valence of auditory tones

A) Experimental protocol. BLA neurons were transfected with AAV1.Syn.GCaMP6f.
 GCaMP6f-expressing axons were imaged in the FrA of awake mice through a cranial window.

**B)** Examples of Ca<sup>2+</sup> transients ( $\Delta F/F_0$ ) from individual boutons recorded from one mouse upon different auditory tones.

**C)** Frequency of axonal Ca<sup>2+</sup> transients (normalized to baseline) recorded in naïve mice (no conditioning, n=6 mice) and during auditory fear conditioning (n=10 mice) with protocol 1 (CS+: *8 kHz*, CS-: *WGN*, n=7) and protocol 2 (CS+: *WGN*, CS-: *8 kHz*, n=3).



# 1030 Fig. S8. BLA-mediated synaptic inputs onto FrA L2/3 pyramidal neurons

**A)** Experimental strategy. *Left*, BLA neurons were transfected with AAV9-CamKIIahChR2-eYFP, and ChR2-expressing BLA neurons were then photo-stimulated *in vivo* with a DPSSL ( $\lambda$  = 473 nm) through an implanted optical fiber. *Middle*, example of a cortical coronal slice with ChR2-YFP fluorescence in the FrA. *PL, prelimbic cortex; MO, medial orbital cortex. Right*, fluorescence intensity profiles measured in the FrA (white box) of 6 different animals in which injections targeted BLA.

**B)** Examples of ChR2-expressing axons imaged in superficial layer 1 with 2P microscopy. **C)** *Left*, representative example of the ChR2-YFP expression profile in the mouse BLA. *Right*, Coronal diagrams of the brains from 6 mice showing the expression profiles (in black) of ChR2-YFP. Diagrams were adapted from the Paxinos atlas.

**D)** Schematic of the slice experiment. BLA-to-FrA axons expressing ChR2 were photo-activated through the objective. *Right*, gray and black lines, representative examples of

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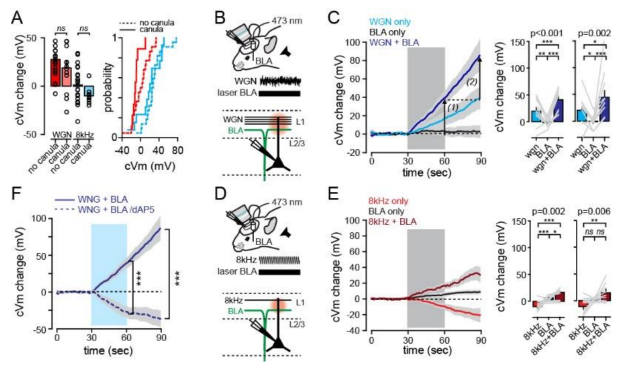
excitatory post-synaptic current (EPSCs) and feed-forward inhibitory post-synaptic currents (IPSCs) respectively evoked on L2/3 pyramidal neurons with an ultrahigh power 488 nm LED.

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**E)** Amplitude (*left*) and jitter (*right*) of light-evoked EPSCs. Gray lines indicate pairs. Open circles indicate individual cells.

**F)** Single-cell examples of depolarizations in FrA neuron evoked by the photostimulation of BLA neurons. Gray lines, single trials; black line, average of 20 trials.

**G)** Peak amplitude (*left*) and fwhm (*right*) of evoked PSP. Triangles, cells. 1050



# Fig. S9. The activation of BLA-to-FrA axons supports the non-linear integration of auditory tones

A) Optical fibers implantation (canula) do not affect auditory-induced cVm changes as 1055 compared to non-implanted mice (no canula).

**B)** Co-activation protocol. ChR2-expressing BLA neurons were photo-stimulated during WGN auditory stimulation.

**C)** Averaged cVm changes (± sem) in response to BLA photo-stimulation alone (black).

*WGN* presentation alone (light blue) and *WGN*+BLA (dark blue). Arrows, analysis time 1060 points 1 (end of stimulation) and 2 (30 sec later). \*, p<0.05; \*\*, p<0.01; \*\*\*, p<0.001; Holm-Sidak multiple comparisons after one-way anova (p<0.001 for time point 1; p=0.002 for time point 2).

**D**, **E**) Same presentation as in (**B**, **C**) but for 8 kHz.

**F)** Averaged cVm change (± sem) during *WGN*+BLA co-stimulation in controls (solid line) 1065 and after dAP5 application (dashed line). \*\*\*, p<0.001; *t-test*.

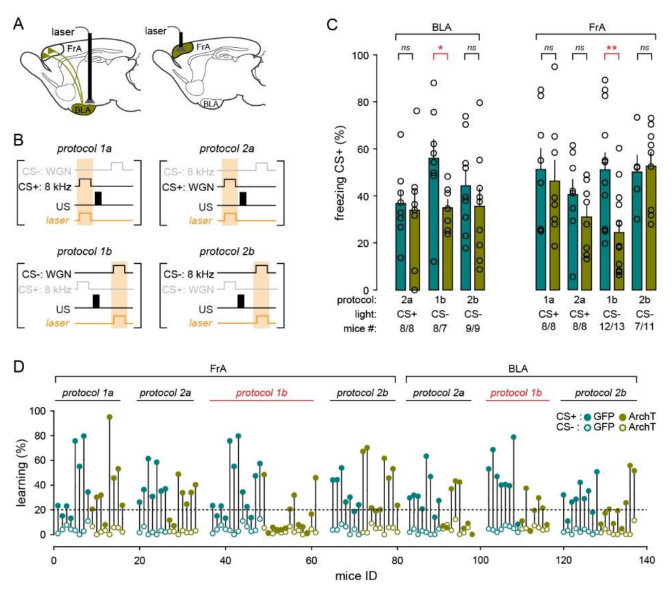


Fig. S10. Statistical comparisons between all behavioral optogenetic experiments

- 1070 **A)** Experimental strategies. *Left*, mice were infected with AAV9.CBA.Flex.ArchT (n=24) or AAV9.CAG.flex.eGFP (for controls, n=25) in BLA and CAV2.CMV.Cre in FrA, and chronically implanted with optical fibers bilaterally in BLA. *Right*, mice were infected with AAV9-CAG-ArchT (n=40) or AAV9-CaMKII-eGFP for controls (n=35) in FrA, and chronically implanted with optical fibers bilaterally in FrA.
- B) Fear conditioning protocols. FrA neurons expressing ArchT (or GFP) were photostimulated during the presentation of CS+ (*8 kHz* for protocol 1a or *WGN* for protocol 2a) or CS- (*WGN* for protocol 1b or *8 kHz* for protocol 2b). US, footshock. Photo-simulations (orange box) were given during CS+ (protocols 1a and 2a) or CS- (protocols 1b and 2b).
   C) Freezing responses during recall upon CS+ for all behavioral conditions.
- 1080 **C)** Learning index during recall upon CS+ (filled circles) and CS- (open circles) in GFP (turquoise) and ArchT (khaki) expressing mice. Significant differences between GFP and ArchT-mice were seen only for protocol 1b (red).

fig.	variable /units	group	Ν	mean	Std dev	median	25%	75%	test	p -value
	WGN cVm change (mV)	control	22	27.62	19.741				One way anova	p<0.001
	(time point1)	dAP5	14	-13.4	17.589				with multiple comparisons	p<0.001 (control vs dAP5)
		iMK801	5	-2.24	5.368				(Holm-Sidak method)	p=0.002 (control vs iMK801)
1D										
10	WGN cVm change (mV)	control	22			25.653	14.3	41.2	Kruskal-Wallis one way anova	p<0.001
	(time point 2)	dAP5	14			-13.74	-19	- 0.93	with multiple comparisons	p<0.05 (control vs dAP5)
		iMK801	5			-0.263	-16	4.81	(Dunn's method)	p<0.05 (control vs iMK801)
	8kHz cPSP change (mV)	control	22	1.16	18.145				One way anova	p=0.004
	(end of stimulation)	dAP5	10	-16.8	12.998				with multiple comparisons	p=0.006 (control vs dAP5)
		iMK801	5	-21	7.657				(Holm-Sidak method)	p=0.008 (control vs iMK801)
1F										
	8kHz cPSP change (mV)	control	22	-3.82	14.122				One way anova	p=0.016
	(time point 2)	dAP5	10	-15.4	15.447				with multiple comparisons	<i>p</i> =0.012 (control vs dAP5)
		iMK801	5	-22.7	12.508				(Holm-Sidak method)	p=0.041 (control vs iMK801)
	events per dendrites	baseline	139	1.664	0.394				One way anova	p<0.001
	during anesthesia	WGN	139	3.282	0.864				repeated measures	p<0.001 (WGN vs baseline)
	(number)	8kHz	139	1.19	0.813				with multiple comparisons	p<0.001 (WGN vs 8kHz)
2D		2kHz	139	1.261	0.52				(Holm-Sidak method)	p<0.001 (WGN vs 2kHz)
										p=0.056 (8kHz vs baseline)
										p=0.097 (2kHz vs baseline)
										<i>p</i> =0.789 (8 <i>k</i> Hz vs 2 <i>k</i> Hz)
	local events per dendrites	baseline	104	0.54	0.38				One way anova	p<0.001
	awake	WGN	104	3.193	1.005				repeated measures	p<0.001 (WGN vs baseline)
	(number)	8kHz	104	2.484	0.567				with multiple comparisons	p=0.031 (WGN vs 8kHz)
2F		2kHz	104	1.999	0.715				(Holm-Sidak method)	p<0.001 (WGN vs 2kHz)
										p<0.001 (8kHz vs baseline)
										p<0.001 (2kHz vs baseline)
										p=0.128 (8kHz vs 2kHz)
	events per dendrites	WGN+	139	3.282	0.864				Two way anova	p=0.026
	anesthesia(+) vs awake(-	WGN-	104	3.193	1.005				repeated measures	p=0.844 (WGN+ vs WGN-)
2I	) (number)	pure+	139	1.226	0.659				with multiple comparisons	p=0.037 (pure+ vs pure-)
	pure tones pooled	pure-	104	2.241	0.5117				(Holm-Sidak method)	
	Learning+ (total mice)	GFP+	6 (8)		-				pearson $\chi^2$ test ( $\chi^2$ =0.4103)	p=0.522
	protocol 1a	ArchT+	7 (8)							×
3E	Learning+ (total mice)	GFP+	7(8)			L			pearson $\chi^2$ test ( $\chi^2=1.3333$ )	p=0.248
	protocol 2a	ArchT+	5(8)							*
	freezing (%)	CS- hab	16	5.811	4.9544	5.236	1.58	8.58	Two way anova	p<0.001 (interaction)
	GFP (all protocols)	CS-rec	16	5.11	5.9977	2.833	1.04	7.83	repeated measures	p=0.8 (CS+hab vs CS-hab)
	hab=habituation	CS+ hab	16	5.33	3.9664	5.015	1.91		with multiple comparisons	p<0.001 (CS+ rec vs CS- rec)
3D	rec=recall	CS+ rec	16	48.48	19.608	45.5	30.5		(Holm-Sidak method)	p=0.99 (CS- hab vs CS- rec)
									,	<i>p</i> <0.001 ( <i>CS</i> + <i>hab vs CS</i> + <i>rec</i> )
			l			l				1

1	<b>•</b> • • • • • • • • • • • • • • • • • •						4.00	0.50	-	
	freezing (%)	CS- hab	16	5.669	6.7446	2.968	1.38	8.58	Two way anova	p<0.001 (interaction)
	ArchT (all protocols)	CS-rec	16	3.866	3.4275	3.59	1.69	6.29	repeated measures	p=0.67 (CS+hab vs CS-hab)
	hab=habituation	CS+ hab	16	4.064	4.3073	2	1.67	7.25	with multiple comparisons	p < 0.001 (CS + rec vs CS - rec)
	rec=recall	CS+ rec	16	40.22	20.911	37.195	27.7	48.2	(Holm-Sidak method)	p=0.72 (CS- hab vs CS- rec)
										<i>p</i> <0.001 ( <i>CS</i> + <i>hab vs CS</i> + <i>rec</i> )
	CS+ freezing (%)	GFP 1a	8	51.21	25.031	50.25	25.6	74.1	one way anova	p=0.235
3F	during recall	ArchT 1a	8	46.25	25.041	37.667	28.9	61.8		
51		GFP 2a	7	45.37	12.038	41.893	36.1	56.6		
		ArchT 2a	8	30.79	14.51	32.447	16.2	43.7		
	Learning+ (total mice)	GFP+	9(11)						pearson $\chi^2$ test ( $\chi^2$ =4.893)	p=0.0270
	protocol 1b	ArchT+	4(13)							
31	Learning+ (total mice)	GFP+	6(7)						pearson $\chi^2$ test ( $\chi^2=0.1169$ )	p=0.7324
	protocol 2b	ArchT+	10(11)							
	freezing (%)	CS- hab	19	4.069	5.007	2.567	1.11	5.92	Two way anova	p<0.001 (interaction)
	GFP	CS-rec	19	7.008	6.258	4.4	3.03	11.1	repeated measures	p=0.994 (CS+ hab vs CS- hab)
	hab=habituation	CS+ hab	19	4.044	3.976	2.5	1.09	6.79	with multiple comparisons	<i>p</i> <0.001 ( <i>CS</i> + <i>rec vs CS</i> - <i>rec</i> )
	rec=recall	CS+ rec	19	48.25	22.389	45.5	26.6	63	(Holm-Sidak method)	p=0.46 (CS- hab vs CS- rec)
										p < 0.001 (CS+ hab vs CS+ rec)
3H	freezing (%)	CS- hab	24	6.133	7.253	2.894	1.83	8.64	Two way anova	p<0.001 (interaction)
	ArchT	CS-rec	24	6.285	6.201	4.889	1.67	7.75	repeated measures	p=0.988 (CS+ hab vs CS- hab)
	hab=habituation	CS+ hab	24	6.079	4.925	6.167	1.75	8.87	with multiple comparisons	p<0.001 (CS+ rec vs CS- rec)
	rec=recall	CS+ rec	24	37.96	23.036	35	18.3	59.3	(Holm-Sidak method)	p=0.97 (CS- hab vs CS- rec)
	ice_recan	Corrice	24	57.70	23.030	55	10.5	57.5	(1101m-Staak method)	
		GFP 1b	10	51.00	25 147	40.012	25.0	741	T	p < 0.001 (CS + hab vs CS + rec)
	CS+ freezing (%)		12	51.09	25.147	49.912	25.6	74.1	Two way anova	p=0.003
3J	during recall	ArchT 1b	13	24.34	17.665	18.5	8	36.9	repeated measures	p=0.002 (GFP1b vs ArchT1b)
		GFP 2b	7	43.4	17.348	42.833	29.5	52.3	with multiple comparisons	p<0.001 (ArchT1b vs ArchT2b)
		ArchT 2b	11	54.07	17.867	58.333	35.8	69	(Holm-Sidak method)	<i>p</i> =0.049 ( <i>GFP2b</i> vs ArchT1b)
	freezing (%)	CS- hab	10	4.799	4.077	4.107	1		Two way anova	p=0.007 (interaction)
	hab=habituation	CS-rec	10	7.304	4.783	8.958	2.83		repeated measures	p=0.742 (CS+hab vs CS-hab)
	rec=recall	CS+ hab	10	3.199	3.131	3.153	0	5.17	with multiple comparisons	<i>p</i> <0.001 ( <i>CS</i> + <i>rec vs CS</i> - <i>rec</i> )
4C		CS+ rec	10	29.24	20.241	25.733	12.5	44.3	(Holm-Sidak method)	p=0.613 (CS- hab vs CS- rec)
										<i>p</i> <0.001 ( <i>CS</i> + <i>hab vs CS</i> + <i>rec</i> )
	freezing (%)	learn+	5	46.24	11.619	44.333	37.8	57.6	t-test	p<0.001
	CS+ during recall	learn-	5	12.25	8.028	12.5	6.98	19.2		
	learning+ vs learning-									
	frequency of events	CS+ learn+	5	1.723	0.733	1.76	1.22	2.05	Two way anova	p=0.015 (learn.+ vs learn)
	normalized to baseline	CS-learn+	5	2.071	0.999	1.667	1.4	2.76	repeated measures	p=0.081 (CS+ vs CS-)
		CS+ learn-		0.616	0.184	0.692	0.48	0.74	with multiple comparisons	
		CS-learn-		0.776	0.295	0.72	0.58	0.92	(Holm-Sidak method)	
4F	frequency of events	learn+ US-	5	1.036	0.339	0.933	0.75	1.33	Two way anova	p=0.042 (interaction)
4F		learn+ US+	5	1.97	0.948	1.835	1.27	2.55	repeated measures	p=0.01 (learn+, US+ vs US-)
4F	normalized to baseline								1	
4F	normalized to baseline w/ US(+) vs w/o US(-)	learn- US-	5	0.742	0.115	0.79	0.68	0.81	with multiple comparisons	p=0.845 (learn-, US+ vs US-)
4F			5 5	0.742 0.688	0.115 0.165	0.79 0.729	0.68 0.56	0.81 0.81	with multiple comparisons (Holm-Sidak method)	p=0.845 (learn-, US+ vs US-)

	c c .	<b>CC</b> 1	r.	1.026	0.220	0.022	0.75	1.22	<b>T</b>	
	frequency of events	CS+ learn+	5	1.036	0.339	0.933		1.33	Two way anova	p=0.036 (interaction)
	normalized to baseline	CS-learn+	5	2.035	0.954	1.58	1.41	2.51	repeated measures	<i>p</i> =0.003 ( <i>learn</i> +, <i>CS</i> +1 <i>vs CS</i> -1)
	CS+1 vs CS-1	CS+ learn-	5	0.678	0.175	0.781	0.53	0.8	with multiple comparisons	p=0.425 (learn-, CS+1 vs CS-1)
		CS- learn-	5	0.879	0.217	0.773	0.73	1.05	(Holm-Sidak method)	
	cumulative $\Delta F/F_0$	CS+	10	0.213	0.201	0.128	0.08	0.29	paired t-test	p=0.012
4G		CS-	10	0.273	0.244	0.204	0.12	0.3	rank sum test	p=0.006
	cumulative $\Delta F/F_0$	learn+	5	0.101	0.0571	0.108	0.05	0.15	paired t-test	p=0.013
	CS CS+	learn-	5	0.016	0.0186	0.0207	0.01	0.03	rank sum test	p=0.032
	cVm change (Vm)	obs.	11	43.92	21.295				paired t-test	p=0.002
	WGN	exp.	11	18.01	29.395					
5H										
	cVm change (Vm)	obs.	8	16.67	8.928				paired t-test	p<0.001
	8kHz	exp.	8	-2.84	9.271					
	observed cVm change	WGN	11	43.92	21.295	18.567	14.7	22.1	t-test	p=0.003
	(mV) time 1	8kHz	8	16.67	8.928	54.172	22.9	62.6	rank sum test	p=0.035
5I										
	observed cVm change	WGN	11	48.54	43.225	34.954	25.1	64.2	t-test	p=0.056
	(mV) time 2	8kHz	8	14.85	18.769	12.97	-2.9	28.5	rank sum test	p=0.035
	Learning+ (total mice)	GFP+	7 (8)						pearson $\chi^2$ test ( $\chi^2$ =7.77)	p=0.0053
Œ	protocol 1b	ArchT+	2 (7)							
6E	Learning+ (total mice)	GFP+	7(9)						pearson $\chi^2$ test ( $\chi^2=2.01$ )	p=0.1469
	protocol 2b	ArchT+	4(9)							
	CS+(8kHz) freezing	GFP	8	55.96	22.305	56.083	49	68.8	t-test	p=0.037
	(%) protocol 1b	ArchT	7	34.91	9.337	34.5	26.2	42.1	Mann-Whitney rank sum	p=0.014
6F	CS- (WGN) freezing		/			54.5			test Mann-Whitney rank sum	•
	(%)	GFP	8	5.354	3.0517	5	3.17	7.08	test	p=0.04
	protocol 1b	ArchT	7	13.69	12.02	10.167	6.71	14.1		
	CS+: (WGN) freezing (%)	GFP	9	44.23	19.692	39.402	28.8	57.9	t-test	p=0.396
	protocol 2b	ArchT	9	35.54	22.461	37.46	17.3	46.1	Mann-Whitney rank sum	p=0.427
6G	CS-: (8kHz) freezing								test	
	(%)	GFP	9	5.171	4.91	3.755	1.37	9.05	t-test	p=0.578
	protocol 2b	ArchT	9	6.847	7.351	5.785	0	13	Mann-Whitney rank sum test	p=0.928
a	FFT magnitude (V)	ctrl	22			5.452	2.97	9.07	Mann-Whitney rank sum test	p=0.048
S1E	0.5-2Hz	dAP5	19			3.26	2.65	3.92		
	spontaneaous 30s-	ctrl	22			110.39	52.3	148	Mann-Whitney rank sum	p=0.007
S1F	cumulative								test	p=0.007
	potential (mV)	dAP5	19	01.0	0.050	51.515	48.8	61.2		0.001
	cVm change (Vm)	ctrl	5	21.8	9.858				paired t-test	p=0.001
S1H	WGN	dAP5	5	-11.3	6.652					
	cVm change (Vm)	ctrl	5	-6.81	12.19				paired t-test	p=0.046
	WGN	dAP5	5	-16.5	7.741					
S1I	NMDARs component	WGN	5	33.12	9.404				paired t-test	p=0.017
	(mV)	dAP5	5	9.695	7.562					

	local events per dendrites	baseline	104	0.54						
			104	0.54	0.38				One way anova	p<0.001
. 1	awake	WGN	104	3.193	1.005				repeated measures	p<0.001 (WGN vs baseline)
	(number)	8kHz	104	2.484	0.567				with multiple comparisons	p=0.031 (WGN vs 8kHz)
		2kHz	104	1.999	0.715				(Holm-Sidak method)	p<0.001 (WGN vs 2kHz)
S2C										p<0.001 (8kHz vs baseline)
52C										p<0.001 (2kHz vs baseline)
										p=0.128 (8kHz vs 2kHz)
	global events per dendrites	WGN	104	0.881	0.61				One way anova	p=0.739
	awake	8kHz	104	0.786	0.469				repeated measures	
	(number)	2kHz	104	0.882	0.813					
	local events per dendrites	baseline	139	1.664	0.394				One way anova	p<0.001
	during anesthesia	WGN	139	3.282	0.864				repeated measures	p<0.001 (WGN vs baseline)
	(number)	8kHz	139	1.19	0.813				with multiple comparisons	p<0.001 (WGN vs 8kHz)
	()	2kHz	139	1.261	0.52				(Holm-Sidak method)	p<0.001 (WGN vs 2kHz)
									(	p=0.056 (8kHz vs baseline)
S2D										p=0.097 (2kHz vs baseline)
										p=0.789 (8kHz vs 2kHz)
	global events per	WGN	139	0.012	0.0258				One way anova	p=0.533
	dendrites								-	p=0.555
	awake	8kHz	139	0.019	0.0333				repeated measures	
	(number)	2kHz	139	0.039	0.0544	207	201	526		
	distance (cm)	total-WGN total-8kHz	17	424.2	178.65	397	301	526	one-way anova on ranks	p < 0.001
			17	487.6	156.48	454.5	382	518	repeated measures	p > 0.05 (WGN vs 8kHz total)
s3B		in-WGN	17	214	82.676	210	168	246	with multiple comparisons	p > 0.05 (WGN vs 8kHz outer)
		in-8kHz	17	240.1	111.59	215.5	159	324	(Tukey method)	p>0.05 (WGN vs 8kHz inner)
		out-WGN	17	153.1	65.353	136.5	96.5	220		
		out-8kHz	17	201.1	77.608	182.5	141	249		- 0.262
s3C	stop (%)	WGN	17	6.676	5.312				paired t-test	p=0.362
	% of tone duration	8kHz	17	4.975	4.813	0.447	0.10	0.05		0.407
	local events	Baseline-	8	0.54	0.38	0.647	0.19	0.85	t-test Mann-Whitney rank sum	p=0.496
	per dendrites	Baseline+	5	0.397	0.308	0.5	0.13	0.59	test	p=0.524
	awake	WGN-	8	3.193	1.005	3.298	2.58	3.61	t-test	p=0.025
	(number)	WGN+	5	1.882	0.623	2.176	1.38	2.27	Mann-Whitney rank sum test	p=0.045
s5E	naive(-) vs FC(+)	8 kHz-	8	2.497	0.532	2.556	2.21	2.7	t-test	p=0.360
		8 kHz+	5	2.145	0.866	2.412	1.81	2.58	Mann-Whitney rank sum test	p=0.548
		2 kHz-	8	2.043	0.682	2	1.55	2.46	t-test	p=0.544
		2 kHz+	5	1.796	0.762	1.935	1.43	2.18	Mann-Whitney rank sum	p=0.739
	Global events	Baseline-	8	0.377	0.283	0.4	0.13	0.48	test t-test	p=0.712
	per dendrites	Baseline+	5	0.317	0.326	0.167	0.09		Mann-Whitney rank sum	p=0.734
	•								test	
s5G	awake	WGN-	8	0.819	0.583	0.75	0.27	1.33	t-test Mann-Whitney rank sum	p=0.411
	(number)	WGN+	5	1.108	0.741	1.294		1.51	test	p=0.65
	naive(-) vs FC(+)	8 kHz-	8	0.729	0.461	0.786	0.32	0.94	t-test	p=0.113
• 1		8 kHz+	5	1.143	0.837	0.765	0.55	1.8	Mann-Whitney rank sum test	p=0.571

1		2 kHz-	8	0.826	0.75	0.833	0.24	1.07	t-test	p=0.968
		2 kHz+	5	0.811	0.507	0.824	0.41	1.14	Mann-Whitney rank sum	p=0.910
		2 KHZ+	5	0.011	0.507	0.024	0.41	1.14	test	p=0.910
	We share (mW)	WCN	22	27.62	10 741					0.001
s6C	cVm change (mV) time point 1	WGN- WGN+	22 8	27.62 1.16	19.741 18.145				one-way anova with multiple comparisons	p<0.001 p=0.008 (upWGN vs npWGN)
	naive(-) vs FC(+)	8kHz-	22	5.407	26.645				(Holm-Sidak method)	p=0.008 (up WGN vs up WGN) p<0.001 (up WGN vs up 8kHz)
	protocol 1	okfiz- 8kHz+	8	0.526	14.083				(Holm-Slaak melnoa)	p < 0.001 (up WGN vs up 8kHz) p < 0.001 (up WGN vs pp 8kHz)
		οκπ2+	0	0.520	14.065					p < 0.001 (up WGN vs pp8KHz) p = 0.601 (npWGN vs up8Khz)
										p=0.620 (npWGN vs up8Khz) p=0.620 (npWGN vs pp8Khz)
										p=0.938 (np8kHz vs pp8kHz)
	cVm change (mV)	WGN-	22	27.62	19.741	25.888	16.2	36.3	t-test	p=0.251
	-								Mann-Whitney rank sum	-
s9A	canula (+)	WGN+	13	18.84	24.246	19.955	5.79	42.6	test	p=0.322
5711	vs non caunla(-)	8kHz-	22	1.16	18.145	1.748	-13	11.7	t-test	p=0.091
		8kHz+	9	-10	8.845	-11.41	-15	- 8.17	Mann-Whitney rank sum test	p=0.086
	cVm change (mV)	WGN	11	18.9	23.505	19.955	11.5	37.3	One wayanova	p<0.001
	time point 1	BLA	11	-0.89	8.132	-0.189	-6.1	4.73	repeated measures	p=0.002 (WGN vs BLA)
		WGN+BLA	11	43.92	21.295	54.172	22.9	62.6	with multiple comparisons	<i>p</i> <0.001 (WGN vs WGN+BLA)
	-	-	_	_	_	_	_	_	(Holm-Sidak method)	<i>p</i> <0.001 (BLA vs WGN+BLA)
s9C	cVm change (mV)	WGN	11	22.43	27.812	27.535	5.23	35	One wayanova	p=0.002
	time point 2	BLA	11	-0.46	11.482	-0.125	-2.4	2.36	repeated measures	<i>p</i> =0.044 (WGN vs BLA)
		WGN+BLA	11	48.54	43.225	34.954	25.1	64.2	with multiple comparisons	<i>p</i> =0.037 ( <i>WGN vs WGN</i> + <i>BLA</i> )
									(Holm-Sidak method)	<i>p</i> <0.001 (BLA vs WGN+BLA)
	cVm change (mV)	8kHz	8	-9.7	9.409	-10.29	-16	- 7.55	One wayanova	p=0.002
	time point 1	BLA	8	6.866	4.765	7.155	2.56	11.2	repeated measures	p<0.001 (8kHz vs BLA)
	-	8kHz+BLA	8	16.67	8.928	18.567	14.7	22.1	with multiple comparisons	p<0.001 (8kHz vs 8kHz+BLA)
									(Holm-Sidak method)	p=0.023 (BLA vs 8kHz+BLA)
9E	cVm change (mV)	8kHz	8	-10.9	15.645	-11.36	-25	- 0.03	One wayanova	p=0.006
	time point 2	BLA	8	2.324	3.503	3.37	-0.5		repeated measures	p=0.064 (8kHz vs BLA)
	unie point 2	8kHz+BLA	8	14.85	18.769	12.97	-2.9		with multiple comparisons	p=0.002 (8kHz vs 8kHz+BLA)
		onini ( D Di i	U	1 1100	101107	1207	2.7	2010	(Holm-Sidak method)	p=0.078 (BLA vs $8kHz+BLA$ )
	cVm change (mV)	control	13	39.1	22.593	48.968	15.4	61	t-test	p<0.001
	WGN+BLA time point							-	Mann-Whitney rank sum	
9F	1	dAP5	7	-26.7	17.794	-29.68	-33	24.3	test	p<0.001
	cVm change (mV)	control	13	46.38	41.988	34.954	20.7	68.2	t-test	p=0.003
	WGN+BLA time point 2	dAP5	7	-9.44	15.34	-2.961	-21	2.25	Mann-Whitney rank sum test	p<0.001

# Supplementary Table 1. Data and Statistic

1085 All statistics were performed using Matlab (Mathworks) with an α significant level set at 0.05 (in red within the table). Normality of all value distributions and the equality of variance between different distributions were first assessed by the Shapiro-Wilk and Levene median tests, respectively. Standard parametric tests were used when data passed the normality and equal variance tests. Non-parametric tests were used otherwise. Only 1090 two-sided tests were used. When applicable, pair-wise multiple post-hoc comparisons were done by using the Holm-Sidak method.

#### **METHODS**

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All experiments were performed in accordance with the Guide for the Care and Use of Laboratory Animals (National Research Council Committee (2011): Guide for the Care and Use of Laboratory Animals, 8th ed. Washington, DC: The National Academic Press.) and the European Communities Council Directive of September 22th 2010 (2010/63/EU, 74). Experimental protocols were approved by the institutional ethical committee guidelines for animal research (N°50DIR\_15-A) and by the French Ministry of Research (N°02169.01). We used male C57Bl6/J 6-weeks old mice from Charles River that were housed with littermates (3-4 mice per cage) in a 12-h light-dark cycle. Cages were 1100 enriched and food and water were provided *ad libitum*.

#### Surgery and virus injection

Mice were anesthetized with an intraperitoneal (i.p.) injection of a mix containing medetomidine (sededorm, 0.27 mg kg<sup>-1</sup>), midazolam (5 mg kg<sup>-1</sup>) and fentanyl (0.05 mg kg<sup>-1</sup>) <sup>1</sup>) in sterile NaCl 0.9% (MMF-mix). Analgesia was achieved by local application of 100 μl 1105 of lidocaine (lurocaine, 1%) and subcutaneous (s.c.) injection of buprenorphine (buprécare, 0.05 mg kg<sup>-1</sup>). 40 µl of dexamethasone (dexadreson, 0.1mg ml<sup>-1</sup>) was administrated intramuscularly (i.m.) in the quadriceps to prevent inflammation potentially caused by the friction of the drilling. A heating-pad was positioned underneath 1110 the animal to keep the body temperature at 37°C. Eye dehydration was prevented by topical application of ophthalmic gel. The skin above the skull was disinfected with modified ethanol 70% and betadine before an incision was made. Stereotaxic injections were done as previously described (Gambino et al., 2014). Briefly, the bregma and lambda were aligned (x and z) and a hole for injection was made using a pneumatic dental drill (BienAir Medical Technologies, AP-S001). The injections were targeted either to the layer 1115 2/3 of the FrA (from bregma: AP, +2.8 mm; DV, -0.2-0.3 mm; ML ±1.0 mm) or to the BLA (from bregma: AP, -1.3 mm; DV, -4.5 to 4.8 mm; ML, ±2.9 mm), or to both at the same time. 200 nl of virus were injected at a maximum rate of 60 nl/min, using a glass pipette (Wiretrol, Drummond) attached to an oil hydraulic manipulator (MO-10, Narishige).

The following viruses were used depending on the experiments. AAV-ChR2 1120 (AAV9.CamKIIa.hChR2(H134R).eYFP.WPRW.SV40, Penn Vector Core) was unilaterally injected in the right BLA, whereas AAV-ArchT-Flex (AAV9.CBA.flex.Arch-GFP.WPRE.SV40,

Penn Vector Core) and CAV2-Cre (Cav2.CMV.Cre, IGMM BioCampus Montpellier) were bilaterally injected into the BLA and FrA, respectively. Control experiments were performed using an AAV containing the DNA construct for GFP 1125 (AAV9.CAG.flex.eGFP.WPRE.bGH). calcium For axonal imaging, AAV-GCaMP6f (AAV1.Syn.GCaMP6f.WPRE.SV40, Penn Vector Core) was injected to the right BLA. For dendritic calcium imaging, AAV-GCaMP6s (AAV9.Syn.Flex.GCaMP6s.WPRE.SV40, Penn Vector Core) and a 1:10000 dilution of AAV-Cre (AAV1.hSyn.Cre.WPRE.hGH, Penn Vector Core) were injected together into the right FrA. After injections, the viruses were allowed 1130 to diffuse for at least 10 min before the pipette was withdrawn. Mice were then either prepared for cranial window implantation or waked-up by a sub-cutaneous injection of a mixture containing atipamezole (revertor, 2.5 mg kg<sup>-1</sup>), flumazenil (0.5 mg kg<sup>-1</sup>), and buprenorphine (buprécare, 0.1 mg kg-1) in sterile NaCl 0.9% (AFB-mix).

- The cranial windows were made as previously described (Gambino et al., 2014). Briefly, after skull's exposure a ~5 mm plastic chamber was attached on the area of interest and a 3 mm craniotomy was made on the right hemisphere above FrA and M2, with a pneumatic dental drill, leaving the dura intact. The craniotomy was covered with sterile saline (0.9% NaCl) and sealed with a 3 mm glass cover slip after viral injection (for imaging experiments). The chamber, the cover slip and a custom-made stainless steel head stage were well attached to the skull using dental acrylic and dental cement (Jet Repair Acrylic, Lang Dental Manufacturing).
- To evaluate the viral expression profiles in BLA and FrA, fixed brain slices were imaged post-hoc using a wide-field epifluorescence microscope (Nikon, Eclipse N-iU). Illumination was set such that the full dynamic range of the 16-bit images was utilized. A two-dimensional graph of the intensities of pixel was plot using Fiji Software. 16-bit images' brightness was processed and masks were registered to the corresponding coronal plates (ranging from -1.94 to -2.70 mm) of the mouse brain atlas using Illustrator (Adobe), at various distances anterior (FrA) or posterior (BLA) to the bregma.

### 1150 Fear conditioning and quantification of learning

At least 5 days before starting behavioral experiments, mice went through handling with the same experimenter that performed the experiments in order to decrease stress. For consistency across experiments, mice were then habituated to

auditory tones during 3 successive days. During habituation, mice were placed on the conditioning compartment (context A, consisting of a squared box with a grid floor that 1155 allows the delivery of a foot shock and with home cage litter under; cleaned between individuals with 70% ethanol). Two conditional auditory stimuli (CS) (8 kHz pure tone; and white Gaussian noise (WGN); each composed of 27 pips, 50 ms in duration, 0.9 Hz for 30 s) were presented 4 times with a 80 dB sound pressure level and variable inter stimulus interval (ISI). The freezing time during each CS presentation was measured and 1160 the mice returned to their home cage. Mice were fear conditioned 24 hours after the last habituation phase by using a classical differential protocol. Briefly, mice were exposed to context A and 5 auditory tones (CS+) were paired with the unconditional stimulus (US, 1s foot-shock, 0.6 mA). The onset of US coincided with the CS+ offset. 5 CS- presentations were intermingled with CS+ presentations with a variable (10-60 s) ISI. CS were 1165 counterbalanced with WGN and 8 kHz pure tones being used as CS+ and CS-, respectively. Recall tests were carried out 24, 48 and 72 hours after the conditioning phase by measuring the freezing time during the presentation of 2 CS+ and 2 CS- in a new context (context B, consisting of a cylindrical white compartment with home cage litter on the 1170 floor; cleaned between individuals with septanios MD 2%).

For optogenetic experiments using *archeorhodopsin* (ArchT) or GFP controls, mice were subjected to the same behavioral protocol described above. Optogenetic inhibition of FrA neurons or BLA-to-FrA projections upon CS presentation was achieved during the conditioning phase by synchronizing each pip (50 ms) composing the CS+ or the CS- with a 50 ms-laser pulse. For the experiments in which the conditioning phase was taken place under the 2 photon microscope, the context consisted of the microscope shading box in which the mice were head-restrained in a custom tube containing a shocking grid at the bottom. CS and US presentations were triggered by a MATLAB routine, associated to a pulse-stimulator (Master-8, A.M.P.I) capable of triggering the foot shock. For somatic and dendritic calcium imaging experiments, behavior was assessed at least 6 hours after imaging sessions. For whole-cell recordings experiments, mice were anesthetized and prepare for patch recordings immediately after behavior.

For each behavioral session, the total time duration (sec) of freezing episodes upon CS+ and CS- presentation was quantified automatically using a fire-wire CCD-camera connected to an automated freezing detection software (AnyMaze, Ugo Basile, Italy), and expressed as % of freezing. Learning index was further quantified for each CS by multiplying the % of freezing in each condition by the corresponding index of discrimination by using the following equation:

$$learning index (\%) = freezing (\%) \times \frac{freezing CS^{+}(\%) - freezing CS^{-}(\%)}{freezing CS^{+}(\%) + freezing CS^{-}(\%)}$$

Learning index <20% during recall was considered as a failure of conditioning.

#### In vivo whole cell recordings

- Isoflurane (4% with  $\sim 0.5$  l min<sup>-1</sup> O<sub>2</sub>) combined with an i.p. injection of urethane (1.5 g kg<sup>-1</sup>, in lactated ringer solution containing in [mM] 102 NaCl, 28 Na L Lactate, 4 KCl, 1195 1.5 CaCl<sub>2</sub>) was used to induce anesthesia and prolonged by supplementary urethane (0.15 g kg<sup>-1</sup>) if necessary. To prevent risks of inflammation, brain swelling and salivary excretions, 40 µl of dexamethasone (dexadreson, 0.1 mg ml<sup>-1</sup>, i.m.) and glycopyrrolate (Robinul-V, 0.01 mg kg<sup>-1</sup>, s.c.) were injected before the surgery. Adequate anesthesia (absence of toe pinch and corneal reflexes, and vibrissae movements) was constantly 1200 checked and body temperature was maintained at 37°C using a heating-pad positioned underneath the animal. Ophthalmic gel was applied to prevent eye dehydration. Analgesia was provided as described for viral injection (with lidocaine and buprenorphine). After disinfection of the skin (with modified ethanol 70% and betadine), the skull was exposed and a  $\sim$ 3mm plastic chamber was attached to it above the prefrontal cortex using a 1205 combination of super glue (Loctite) and dental acrylic and dental cement (Jet Repair FrA (+2.8 mm from bregma, ±1.0 mm midline) was made using a pneumatic dental drill, leaving the dura intact.
- 1210 Whole-cell patch-clamp recordings of L2/3 pyramidal neurons were obtained as previously described (Gambino et al., 2014) Briefly, high-positive pressure (200–300 mbar) was applied to the pipette (5–8 M $\Omega$ ) to prevent tip occlusion, when passing the pia. Immediately after, the positive pressure was reduced to prevent cortical damage. The

pipette resistance was monitored in the conventional voltage clamp configuration during the descendent pathway through the cortex (until -200  $\mu$ m from the surface) of 1  $\mu$ m 1215 steps. When the pipette resistance abruptly increased, the 3–5 G $\Omega$  seal was obtained by decreasing the positive pressure. After break-in, Vm was measured, and dialysis was allowed to occur for at least 5 min before launching the recording protocols. Currentclamp recordings were made using a potassium-based internal solution (in mM: 135 potassium gluconate, 4 KCl, 10 HEPES, 10 Na2-phosphocreatine, 4 Mg-ATP, 0.3 Na-GTP, 1220 and 25 µM, pH adjusted to 7.25 with KOH, 285 mOsM), and acquired using a Multiclamp 700B Amplifier (Molecular Devices). Spontaneous activity was recorded prior, during and after the presentation of auditory stimulation. Spiking pattern of patched cells was analyzed to identify pyramidal neurons. dAP5 (1 mM, Tocris) was topically applied to the dura mater, before whole cell recordings. Offline analysis was performed using custom 1225 routines written in Sigmaplot (Systat), IGOR Pro (WaveMetrics) and Matlab (Mathworks).

#### In vivo optogenetics

After virus injection for ChR2 or ArchT expression, mice were subsequently implanted with fiber optic cannula for optogenetics (CFML22U, Thorlabs) in the BLA. The optic fibers were previously cleaved with a fiber optic scribe (S90R, Thorlabs) at 4.5mm 1230 for BLA. The cannula were guided and stereotaxically inserted inside the brain with the help of a cannula holder (XCL, Thorlabs) through the same burr hole used for the viral injections (BLA coordinates from bregma: AP, -1.3mm; DV, -4.5 mm; ML, ±2.9mm) and secured in place with a mix of super glue (Loctite) and dental acrylic and dental cement (Jet Repair Acrylic, Lang Dental Manufacturing). Anesthesia was reversed using AFB-mix 1235 for mice assigned to behavioral experiments. For *in vivo* photostimulation of ChR2expressing BLA neurons, the fiber optic cannula and the optogenetic patch cable (M83L01, Thorlabs) were connected through a ceramic split mating sleeve (ADAL1, Thorlabs). The patch cable was then coupled to a blue DPSS laser (SDL-473-050MFL, Shanghai Dream Lasers Technology) which was triggered by a pulse-stimulator (Master-1240 9, A.M.P.I), able to synchronize 50 ms laser pulses with 50 ms sound pips composing the CS. For inhibition of BLA-to-FrA projections during learning, in vivo bilateral optic stimulation of ArchT-expressing neurons was achieved by coupling the optic fibers implanted in BLA to a multimode fiber optic coupler (FCMH2-FCL, Thorlabs), with a

1245 ceramic split mating sleeve, and subsequently connected to a yellow DPSS laser (SDL-LH-1500, Shanghai Dream Lasers Technology).

## In vitro whole-cell recordings

Mice were anesthetized with a mixture of ketamine/xylazine (100mg/kg and 10mg/kg respectively) and cardiac-perfused with ice-cold, oxygenated (95% 02, 5% CO2) cutting solution (NMDG) containing (in mM): 93 NMDG, 93 HCl, 2.5 KCl, 1.2 1250 NaH2PO4, 30 NaHCO3, 25 Glucose, 10 MgSO4, 0.5 CaCl2, 5 Sodium Ascorbate, 3 Sodium Pyruvate, 2 Thiourea and 12mM N-Acetyl-L-cysteine (pH 7.3-7.4, with osmolarity of 300-310 mOsm). Brains were rapidly removed and placed in ice-cold and oxygenated NMDG cutting solution (described above). Coronal slices (300 µm) were prepared using a Vibratome (VT1200S, Leica Microsystems, USA) and transferred to an incubation 1255 chamber held at 32°C and containing the same NMDG cutting solution. After this incubation (9-11 min), the slices were maintained at room temperature in oxygenated modified ACSF containing (mM): 92 NaCl, 2.5 KCl, 1.2 NaH2PO4, 30 NaHCO3, 20 HEPES, 25 Glucose, 2 MgSO4, 2 CaCl2, 5 Sodium Ascorbate, 3 Sodium Pyruvate, 2 Thiourea and 12mM N-Acetyl-L-cysteine (pH 7.3-7.4, with osmolarity of 300-310 mOsm) until 1260 recording.

Whole-cell recordings of layer 2/3 FrA principal neurons were performed on coronal slices (from bregma: +2.58 mm to +3.08 mm) at 30-32°C in a superfusing chamber. Patch electrodes (3-5 MΩ) were pulled from borosilicate glass tubing and filled
with a K-gluconate-based intracellular solution (in mM: 140 K-gluconate, 5 QX314-Cl, 10 HEPES, 10 phosphocreatine, 4 Mg-ATP and 0.3 Na-GTP (pH adjusted to 7.25 with KOH, 295 mOsm). BLA-to-FrA monosynaptic EPSCs were elicited by 1-50 ms light stimulations delivered by an ultrahigh power 460 nm LED (Prizmatix Ltd, Israel). Data were recorded with a Multiclamp700B (Molecular Devices, USA), filtered at 2 kHz and digitized at 10 kHz. Data were acquired and analysed with pClamp10.2 (Molecular Devices).

### 2-photon laser-scanning microscope (2PSLM)-based calcium imaging

Head-fixed awake mice were placed and trained under the microscope every day for at least 7 days prior to the experiment, and then imaged 21 to 35 days after virus

injection using an in vivo non-descanned FemtoSmart 2PLSM (Femtonics, Budapest, 1275 Hungary) equipped with a ×16 objective (0.8 NA, Nikon). The MES Software (MES v.4.6; Femtonics, Budapest, Hungary) was used to control the microscope, the acquisition parameters, and the TTL-driven synchronization between the acquisition and auditory/footshock stimuli. The GCaMPs were excited using a Ti:sapphire laser operating 1280 at  $\lambda$ =910 nm (Mai Tai DeepSee, Spectra-Physics) with an average excitation power at the focal point lower than 50 mW. Time-series images were acquired within a field-of-view of 300 x 300 µm (256 lines, 1ms/line) for axons; for dendrite: 200 x 60 µm (64 lines, 0.5ms/line). Each imaging session consisted of 30 s of baseline recording followed by 8 gaussian and 8 pure (8kHz)-tone auditory stimuli delivered with pseudo-random delays. 1285 We imaged on average 3500 frames ( $\sim$ 900 s) per session, and no visible photo-bleaching was observed. Images were then analyzed as previously described (Gambino et al., 2014) using custom routines written in Fiji and Matlab (Mathworks). We registered images over time and corrected XY motion artifacts within a single imaging session by using crosscorrelation based on rigid body translation (Stack aligner, Image J, NIH, USA). Motion corrections were then assessed by computing pair-wise 2D correlation coefficient (Image 1290 correlation, Image J, NIH, USA), and frames were discarded from the analysis if lower than 0.7. Regions of interest (ROIs) for pyramidal neurons and putative axonal boutons were selected and drawn manually. All pixels within each ROI were first averaged providing a single time-series of raw fluorescence. To limit the effect of fluorescence drift over time, the baseline fluorescence ( $F_0$ ) was calculated as the mean of the lower 50% of previous 3 1295 s fluorescence values. Change in fluorescence ( $\Delta F_t/F_0$ ) was defined as ( $F_t-F_0$ )/ $F_0$ , were  $F_t$ is the fluorescence intensity at time t (time of the first pixel in each frame). Calcium events were then detected using a template-based method with a custom library of calcium transients. Templates were created by extracting and averaging segments of data that 1300 were visually identified as corresponding to a transient. Calcium transients whose peak amplitude reached a 3 X background standard deviation threshold were further considered for analysis. Each detected event was inspected visually and analysis was restricted to detected events rather than on raw fluorescence. For extracting spatial profiles of dendritic calcium events, small ROIs of 2 X 2 pixels are generated along the dendrite by using custom routine in Fiji. The spread of Ca<sup>2+</sup> events was then quantified by 1305 calculating the full-width at half-max (fwhm, expressed as % of total dendritic length) of the normalized Gaussian fit at the time when the averaged  $\Delta F/F_0$  was maximal.

# Data availability and Statistics

- All data generated or analyzed during this study are included in the manuscript, and provided in the supplementary statistical table. Experiments and analysis were conducted blind to the operator. Data are presented as the median  $\pm$  interquartile range or mean  $\pm$  sem (except where stated differently). All statistics were performed using Matlab (Mathworks) and Sigmaplot (Systat) with an  $\alpha$  significant level set at 0.05. Normality of all value distributions and the equality of variance between different distributions were first assessed by the Shapiro-Wilk and Levene median tests, respectively. Standard parametric tests were only used when data passed the normality and equal variance tests. Non-parametric tests were used otherwise. Only two-sided tests were used. When applicable, pair-wise multiple post-hoc comparisons were done by using the Holm-Sidak method. No statistical methods were used to estimate sample size,
- 1320 but  $\beta$ -power values were calculated for parametric tests.