

Rapid Plasticity of Higher-Order Thalamocortical Inputs during Sensory Learning

Highlights

- Mice learn a multi-whisker sensory association during automated home-cage training
- Training, but not sensory stimulation alone, causes rapid synaptic plasticity in S1
- Synaptic plasticity observed only in higher-order thalamocortical pathways
- POm thalamic inputs are strengthened sequentially, first on L5 and then on L2 Pyr

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In Brief

Audette et al. use automated training and *in vitro* electrophysiology to define cortical circuit changes during sensory-association learning. Pathway-specific analysis identifies higher-order thalamic inputs to sensory cortex as a site of synaptic potentiation during the earliest stages of learning.



Rapid Plasticity of Higher-Order Thalamocortical Inputs during Sensory Learning

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<https://doi.org/10.1016/j.neuron.2019.04.037>

SUMMARY

Neocortical circuits are sensitive to experience, showing both anatomical and electrophysiological changes in response to altered sensory input. We examined input- and cell-type-specific changes in thalamo- and intracortical pathways during learning using an automated, home-cage sensory association training (SAT) paradigm coupling multi-whisker stimulation to a water reward. We found that the posterior medial nucleus (POm) but not the ventral posterior medial (VPM) nucleus of the thalamus drives increased cortical activity after 24 h of SAT, when behavioral evidence of learning first emerges. Synaptic strengthening within the POm thalamocortical pathway was first observed at thalamic inputs to L5 and was not generated by sensory stimulation alone. Synaptic changes in L2 were delayed relative to L5, requiring 48 h of SAT to drive synaptic plasticity at thalamic and intracortical inputs onto L2 Pyr neurons. These data identify the POm thalamocortical circuit as a site of rapid synaptic plasticity during learning and suggest a temporal sequence to learning-evoked synaptic changes in the sensory cortex.

INTRODUCTION

Experience-dependent plasticity is a cardinal feature of the neocortex. Abundant evidence indicates that motor or perceptual learning drives changes in neocortical circuits, with changes observed in fMRI signals (Shibata et al., 2016; Summerfield et al., 2006), altered topographic organization (retino-, tonotopy, or somatotopy; Harris et al., 2001; Kilgard and Merzenich, 1998; Schwartz et al., 2002), enhanced feature-selective responses and increased spike output to previously undetectable stimuli (Glazewski and Barth, 2015; Karni and Sagi, 1991), and increased synaptic strength (Cheetham et al., 2008; Clem et al., 2008; Rioult-Pedotti et al., 2000).

How is learning associated with plasticity in specific neocortical circuits? In rodent somatosensory cortex, experience-dependent changes are concentrated in infra- and supragranular layers rather than layer 4 (L4) (Chandrasekaran et al., 2015;

Diamond et al., 1994; Glazewski and Barth, 2015; Glazewski and Fox, 1996; Jacob et al., 2012; Oberlaender et al., 2012). This dissociation could be a product of postsynaptic differences but also aligns with the laminar targets of the two thalamic input streams that drive the barrel cortex. The primary, ventral posterior-medial (VPM) thalamic nucleus provides the dominant glutamatergic input to L4 (Feldmeyer et al., 2013), and neurons in L2 and L5 receive strong glutamatergic input from the higher-order posterior-medial (POm) thalamic nucleus (Audette et al., 2018; Bureau et al., 2006; Petreanu et al., 2009), raising the possibility that the two thalamocortical circuits might be related to differences in plasticity induction across different cortical layers.

VPM neurons receive ascending sensory information directly from the trigeminal brain stem nucleus and respond robustly to the deflection of a single whisker (Feldmeyer et al., 2013). VPM faithfully relays these signals to L4 of the cortex, where excitatory neurons fire short-latency action potentials time locked by fast, feedforward inhibition (Cruikshank et al., 2010; Feldmeyer et al., 2013). Although VPM input plasticity has been observed in early development, electrophysiological and anatomical changes in L4 of the adult mouse are typically only detectable after prolonged periods of drastically altered sensory input levels (Crair and Malenka, 1995; Diamond et al., 1994; Fox, 1992; Glazewski and Fox, 1996; Oberlaender et al., 2012).

The higher-order thalamic nucleus POm also receives sensory signals from the brainstem but integrates this information with strong cortical feedback from reciprocal connections to S1, S2, and M1 (Alloway et al., 2003; Groh et al., 2014; Urbain and Deschênes, 2007). Like other higher-order thalamic nuclei, such as the pulvinar, POm neurons are well positioned to provide contextual information to the cortex and are strongly modulated by arousal and cholinergic activity (Masri et al., 2006; Purushothaman et al., 2012; Roth et al., 2016; Sobolewski et al., 2015). POm axon terminals, concentrated in L1 and L5, provide direct synaptic input to excitatory neurons in L2 and L5 (Audette et al., 2018; Bureau et al., 2006), and POm activation can prolong and enhance sensory responses in S1 (Mease et al., 2016). Plasticity of POm afferents onto L2 excitatory neurons can be elicited over short timescales by artificial whisker stimulation in anesthetized mice, suggesting that these synapses possess the machinery for long-term potentiation (Gambino et al., 2014). Indeed, new findings indicate that POm activity in acute brain slices may facilitate intracortical plasticity through disinhibition (Williams and Holtmaat, 2019).



Because POM inputs can elicit activity-dependent synaptic strengthening, convey contextual and brain state information to the cortex, and drive activity in highly plastic cortical neuron populations, we hypothesized that POM-related pathways would undergo experience-dependent modifications during whisker-dependent learning. To test this, we developed a high-throughput, home-cage system for automated sensory association training that couples a multi-whisker stimulus to a water reward in freely moving mice. Animals exhibit behavioral evidence of learning within the first 24 h of training, and performance increases with longer training intervals.

Using acute brain slices for pathway-specific activation and precise targeting of postsynaptic neurons across different cortical layers, we identified changes in excitatory synapses during early learning and then later, as behavioral performance improved. Neocortical responses to optogenetic activation of thalamic inputs were changed at the earliest stages of learning, where POM, but not VPM, drove significantly greater evoked firing in neocortical neurons in both deep and superficial layers after just 24 h of training. Synaptic strengthening of direct POM inputs to L5 neurons was linked to increased spiking during early training but was not manifested in L2 neurons until 48 h of training. Evidence of synaptic strengthening in intracortical pathways, primarily at L2-L2 excitatory inputs, was delayed with respect to plasticity at POM synapses in L5. Importantly, these synaptic changes were not observed with unrewarded sensory stimulation. Together, our results indicate that plasticity at thalamic inputs from POM initiate cortical rewiring during sensory learning, revealing a temporal sequence of synaptic changes that begin in L5 and then progress to L2.

RESULTS

Automated, Home-Cage Sensory Association Training

Time-intensive animal training paradigms are not well-suited to a comprehensive electrophysiological analysis of cell-type- and input-specific excitatory synaptic changes during learning. Thus, we designed an automated home-cage training system for freely moving animals, where a multi-whisker stimulus was predictively coupled to the cage water source (Figures 1A and 1B). We developed a mouse-initiated training paradigm adapted from classical trace conditioning, where a conditioned sensory stimulus (CS) was followed by an unconditioned stimulus (US) at a fixed delay (Galvez et al., 2006). A gentle air puff was used as the CS, because it is a naturalistic stimulus and can activate multiple whiskers without precise animal positioning, well-suited for training freely moving mice. On training trials, snout entry into the water port triggered a short random delay (200–800 ms) followed by a gentle air puff (CS; 4–6 psi; 500 ms), a fixed delay (500 ms), and then water delivery (US; Figures 1B and 1D).

Blank trials, where no CS or US was delivered, were randomly interleaved on 20% of trials (Figures 1C and 1D). Prior to training, mice received 1 day of acclimation to the home-cage drinking setup, where they experienced an identical trial structure but without presentation of the CS (Figure 1C). Mice readily learned to drink from the lick port, and behavior data, including nose-poke times and licking, were recorded throughout the acclimation and training period.

In trace conditioning, the CS consistently predicts the US and becomes sufficient to evoke an unconditioned behavioral response, in this case, licking (Cohen et al., 2012; Galvez et al., 2006). CS-US association was monitored by comparing the lick frequency prior to the time of potential water delivery, i.e., anticipatory licking (referred to as “licking”), between training and blank trials. Before sensory association training (SAT), licking was identical for both trial types (Figure 1E), as expected because no cue differentiated water reward versus blank trials. A transient suppression of lick rates at the onset of SAT recovered rapidly, and the total trials/day were comparable between acclimation and training days (Figure 1G). By the end of the first day of training (24 h), mice increased their lick rate following the air puff, but not on blank trials, evidence of a predictive CS-US association (Figures 1F–1H; Cohen et al., 2012).

Behavioral evidence of learning after 24 h of training was monitored by assessing the difference in licking for air puff-water paired (L_{water} ; L_w) versus blank trials (L_{blank} ; L_b) over time to calculate a performance metric ($L_w - L_b$; Figure 1H). An increase in lick frequency for stimulus-paired trials was visible between 12 and 24 h and became significantly different from blank trials by 24 h. More than 75% of animals showed increased licking by the end of the first training day (Figure 1I). These data show that home-cage trace conditioning can drive the acquisition of a multi-whisker sensory association that is observable after just 24 h, which enabled electrophysiological investigation of circuitry changes at the earliest stages of sensory learning.

Increase in POM-Evoked Cortical Activity after 24 h of SAT

To systematically evaluate the location of SAT-dependent modifications across the earliest stages of cortical processing in primary somatosensory cortex, we measured the response properties of cortical neurons following channelrhodopsin-assisted stimulation of the primary sensory thalamic nucleus, VPM, or the higher order nucleus POM in acute brain slices. We examined how POM-evoked firing was changed by SAT by recording from pyramidal neurons (Pyr) in the major POM-recipient layers, L2 and L5 (Audette et al., 2018; Bureau et al., 2006; Viaene et al., 2011), after activation of thalamic axons. To isolate cortical responses specific to this pathway, ChR2 was expressed in POM (Figure S1). Although virally expressed ChR2 expression can vary across animals, strict criteria were used to ensure a minimum level of ChR2 viral transduction monitored by fluorescence signal in cortical POM afferents. Animals were assigned to control or SAT groups without prior knowledge of ChR2 expression levels, and average fluorescent labeling was comparable across groups.

In tissue from control animals (acclimation but no whisker stimuli), optogenetic stimulation of POM axons (5 pulses; 80 ms interstimulus interval [ISI]) drove short latency spiking in L5 Pyr (14.6 ± 3.4 ms; Figure S1) and subthreshold excitatory postsynaptic potentials (EPSPs) in L2 Pyr, consistent with prior work (Figures 2C and 2H; Audette et al., 2018). POM stimulation occasionally caused prolonged, recurrent sub- and suprathreshold activity in the post-stimulus period that lasted for several seconds, although action potentials were only rarely observed in L2. Both spike probability and EPSP amplitude depressed

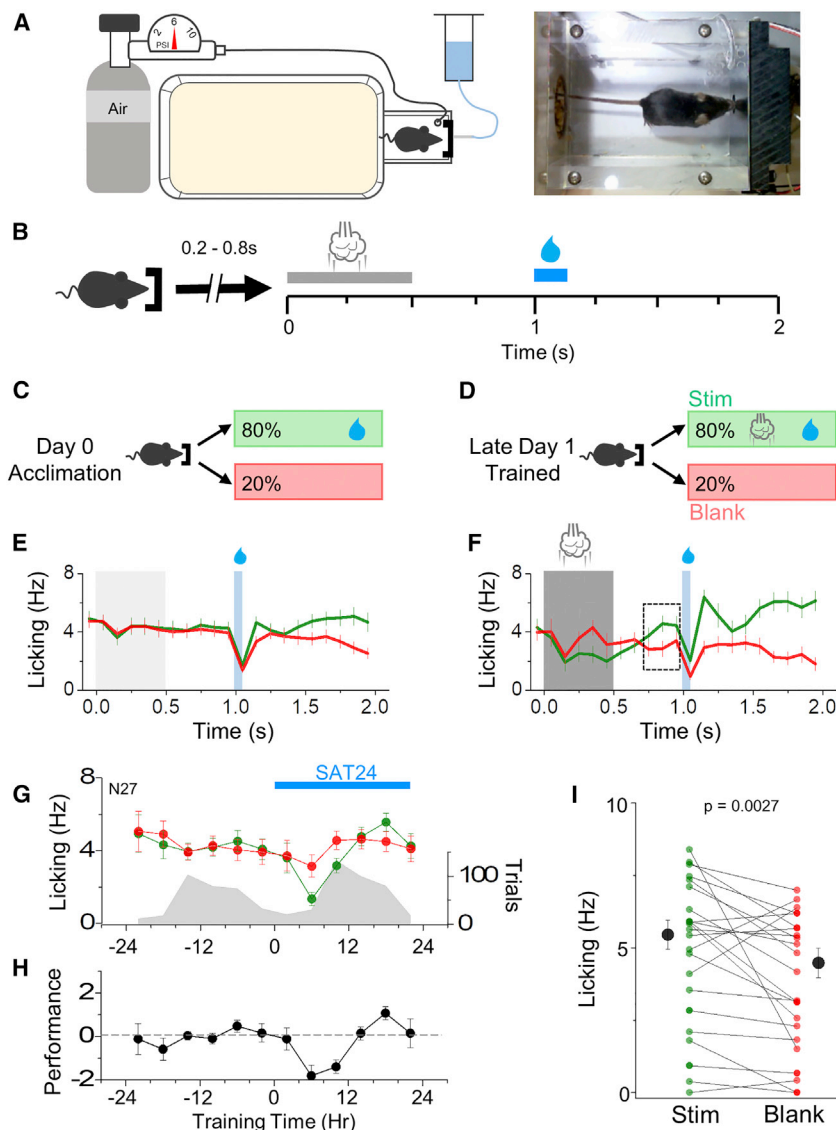


Figure 1. Automated Home-Cage Training Enables Rapid Acquisition of Multi-whisker Sensory Association

(A) Schematic of home-cage sensory association training cage (left) and image of mouse initiating a training trial (right).

(B) Sensory association training paradigm. Upon infrared (IR) beam-break measured nose poke, a random delay (200–800 ms) occurs prior to trial initiation. Air puff delivery period (CS; 500 ms duration; 4–6 psi) occurs at $t = 0$ following random delay with water delivery (US; 75 ms; $\sim 15 \mu\text{L}$) occurring at $t = 1$ s, leaving a 500 ms delay in between the CS and US. A new trial could not be initiated until $t = 2$ s.

(C and D) Identical trial structure during acclimation and SAT, with 80% of initiated trials providing water and air puff (no air puff during acclimation) and 20% of trials delivering neither air puff nor water.

(E and F) Average global lick rates (10 ms bins) of mice during training trials on either acclimation day (E) or the last 8 h (hour 16–24) of training day 1 (F). Grey and white shading represents stimulus and water delivery; black box shows 300 ms anticipatory lick window. $N = 11$ animals for (E) and (F).

(G) Time course of anticipatory lick rates (left axis; 300 ms prior to water delivery; 4 h bins) throughout training for blank (red) and stim+water trials (green) averaged across all animals. Trial initiation counts (right axis) are shown in gray for the same time bins.

(H) Performance defined as the difference in anticipatory lick rates between stim+water trials and blank trials during learning.

(I) Individual animal paired comparisons (Wilcoxon rank sum test) of lick rate for stim+water trials and blank trials on the last 20% of trials on day 1. Averages represented as mean \pm SEM.

with subsequent stimuli in L5 Pyr (Figures 2H, 2K, and S1; Audette et al., 2018).

After 24 h of SAT, POM-evoked cortical firing was dramatically increased (Figures 2D and 2I). The fraction of neurons spiking during the stimulus and/or the post-stimulus period increased from 60% to 92% in L5 and 14% to 63% in L2. In L5, the SAT-associated change in POM spiking was most notable in response to the first light pulse, where mean firing frequency in the first 10 ms after POM stimulation increased 4-fold (control 1.65 ± 1.0 Hz; 24 h SAT 4.25 ± 1.3 Hz) and the latency to spike was slightly reduced (control 14.6 ± 3.4 ms; 24 h SAT 13 ± 3.2 ms; Figure S1). L2 Pyr spiking during the 500 ms stimulus window increased nearly 20-fold (control 0.02 ± 0.02 Hz; 24 h SAT 0.33 ± 0.2 Hz), with the majority of spike output occurring later in the stimulus train. Spikes in L2 Pyr neurons typically occurred 10–40 ms after an individual light pulse stimulus and showed high trial-to-trial variability (Figure S1). Where whole-

cell recordings were performed, we calculated the subthreshold EPSP amplitude evoked by stimulation of POM inputs to the barrel cortex. Overall, EPSPs evoked by the first light pulse in the stimulus train were similar for control and SAT24 animals in L2 (control 4.24 ± 0.87 mV, $n = 11$ cells; 24 h SAT 4.36 ± 0.71 mV, $n = 18$ cells), although EPSPs increased modestly in L5 after training (control 7.14 ± 1.6 mV, $n = 6$ cells; 24 h SAT 8.27 ± 1.4 mV, $n = 4$ cells). However, these values are difficult to interpret, as the EPSP peak amplitude under these recording conditions likely represents a composite of thalamic and local polysynaptic input driven by strong ChR2 activation. Additionally, cells that consistently fired POM-evoked action potentials had to be excluded from PSP measurements because the presence of a spike made accurate detection of PSP amplitude impossible. Because L5 neurons were more likely to spike with POM activation after SAT, this dataset was particularly affected (8/12 cells fired with POM stimulation and were excluded from analysis).

In trained mice, POM stimulation evoked markedly increased levels of prolonged depolarization during the post-stimulus period for neurons in both L2 and L5 apparent in both firing

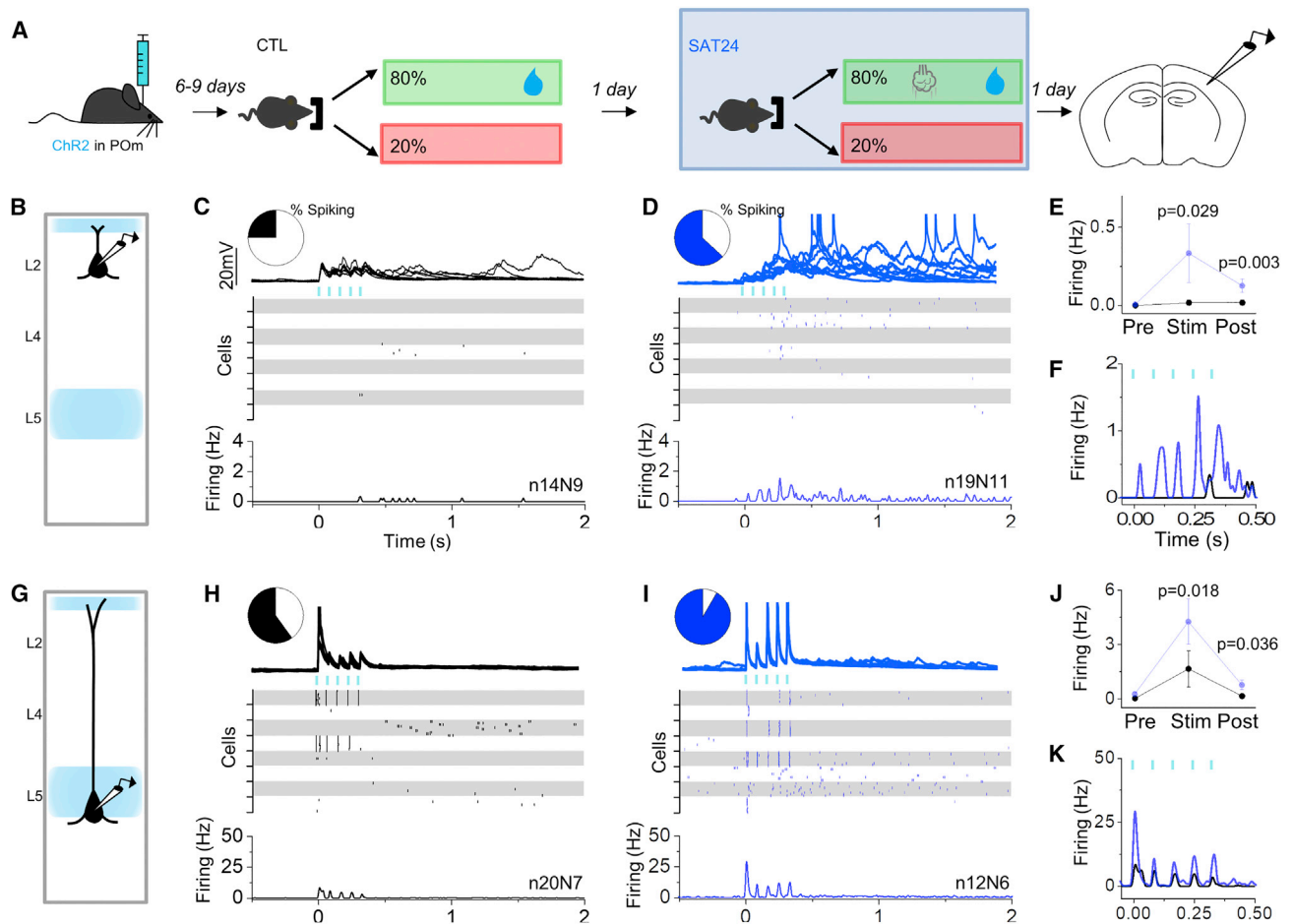


Figure 2. Increase in POM-Evoked Cortical Activity after 24 h of SAT

(A) Schematic of experiment with recordings performed in ChR2-injected mice after 24 h of acclimation and 24 h of SAT.

(B) Schematic of POM axonal labeling and laminar pyramidal neuron recording site in L2.

(C and D) POM-evoked activity (blue bars; 5 pulses; 5 ms; 80 ms ISI) in L2 Pyr neurons of control animals that received 24 h of acclimation (left, black) and 24 h SAT (blue, right). Pie chart shows fraction of neurons that generated any action potentials following stimulation. Example cell response (top) shows 10 consecutive trials for an individual neuron. Raster (middle) shows spiking activity on 10 consecutive trials for 8 example cells. Global peri-stimulus time histogram (PSTH) (bottom, 10 ms bins) shows average firing frequency across all cells in group.

(E) Average firing frequency across all cells during the 500 ms preceding POM stimulation (pre), during stimulation (stim), and directly following stimulation (post).

(F) Overlay of POM-evoked spiking activity (10 ms bins) for CTL (black) and SAT24 (blue) animals.

(G–K) Same as (C)–(F), but for L5 Pyr neurons.

Averages represented as mean \pm SEM. See also Figure S1.

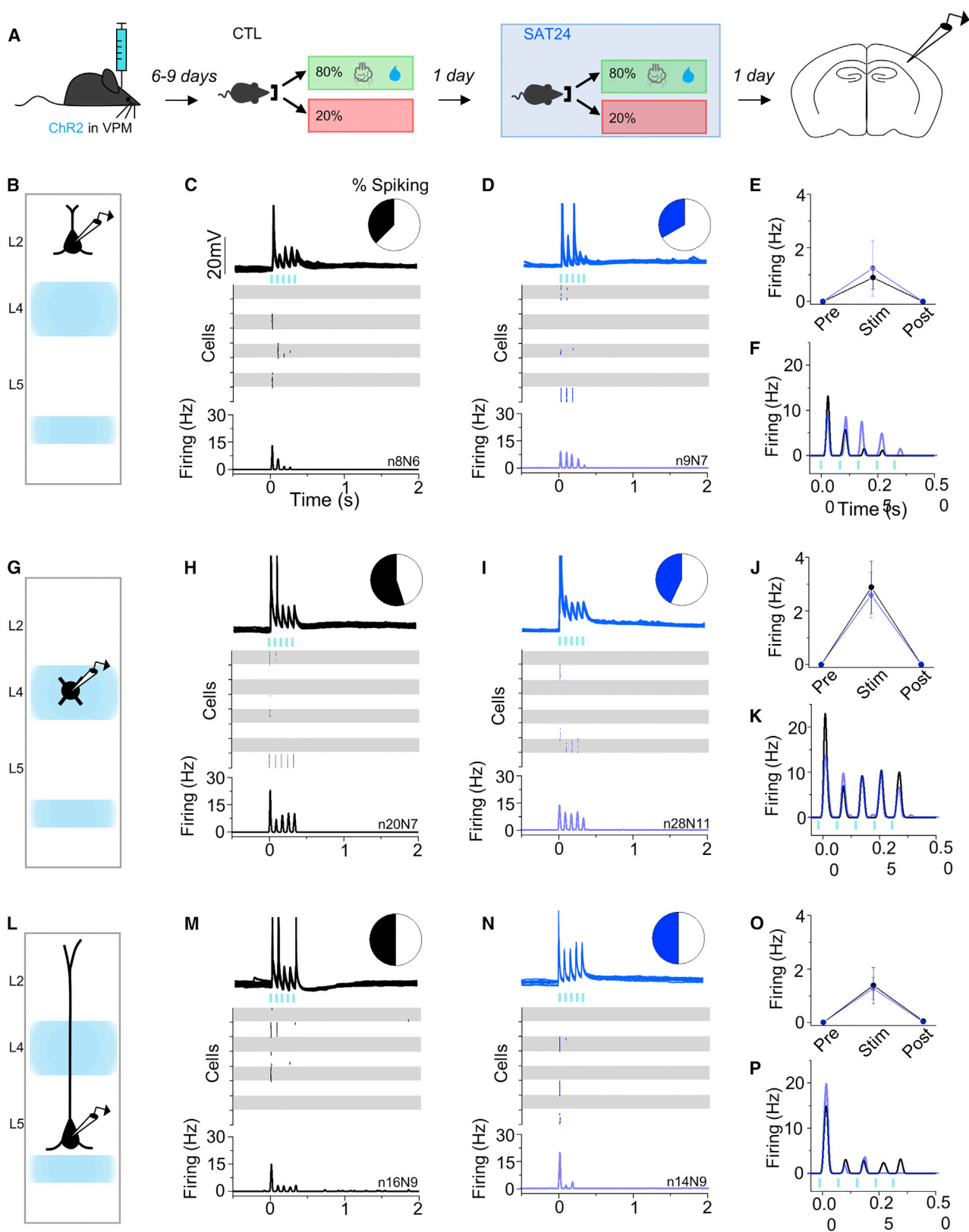
rate (L2 control 0.02 ± 0.02 Hz; L2 24 h SAT 0.13 ± 0.04 Hz; L5 control 0.15 ± 0.11 Hz; L5 24 h SAT 0.76 ± 0.26 Hz; Figures 2E and 2J) and peak subthreshold activity in the 1 s period following the stimulus train (L2 control 2.39 ± 0.48 mV, $n = 11$ cells; L2 24 h SAT 5.20 ± 0.68 mV, $n = 18$ cells; L5 control 1.86 ± 0.72 mV, $n = 6$ cells; L5 24 h SAT 2.79 ± 0.65 mV, $n = 6$ cells). These results indicate that 24 h of SAT drives a marked increase in Pyr responses to POM stimulation that are manifested across the cortical column.

VPM-Evoked Responses Are Unchanged

Although plasticity at VPM inputs is generally restricted to early development in L4, VPM axon remodeling in L4 has been observed in adult animals (Oberlaender et al., 2012), and VPM-

related plasticity in other cortical layers has not been well-investigated. We used optogenetic activation of ChR2-expressing VPM afferents to screen for changes in evoked firing of regular spiking (L4) or Pyr neurons in L2 and L5, which receive both direct and indirect VPM input.

We first investigated SAT-induced changes in L4, the main VPM-recipient layer (Bureau et al., 2006; Cruikshank et al., 2010; Meyer et al., 2010). In control animals, optogenetic stimulation of VPM axons drove precisely timed, short-latency (10.1 ± 1.5 ms) action potentials in the majority (55%) of L4 neurons that were restricted to the stimulus window (Figures 3H and 3I; Cruikshank et al., 2010). After SAT, a slightly smaller fraction of L4 neurons fired with VPM stimulation (43%) and spike latencies were unchanged (Figures S2G and S2H). The mean firing



(legend on next page)

frequency across the entire stimulus period was indistinguishable between control and SAT (control 2.89 ± 0.98 Hz; 24 h SAT 2.58 ± 0.86 Hz; [Figure 3J](#)), and subthreshold EPSP amplitude remained unchanged (control 10.18 ± 1.48 mV, $n = 12$ cells; 24 h SAT 9.59 ± 1.18 mV, $n = 9$ cells). Unlike POM-evoked activity in L2 and L5, VPM stimulation never drove recurrent firing in the post-stimulus period, consistent with strong feedforward inhibition in this layer ([Cruikshank et al., 2010](#); [Porter et al., 2001](#)).

VPM provides direct synaptic input to L5 ([Bureau et al., 2006](#); [Meyer et al., 2010](#)), and SAT could conceivably drive changes at this connection. Indeed, experience-dependent changes in L5 firing have been observed in some studies ([Diamond et al., 1994](#); [Jacob et al., 2012](#); [Ward et al., 2012](#)), and the change in POM-evoked firing in L5 we observed might reflect a special capacity for plasticity of neurons in this layer. However, optogenetic activation of VPM afferents revealed that SAT did not change the fraction of L5 spiking neurons (50% for both conditions) or their mean evoked firing frequency during or after the stimulus window ([Figures 3L–3P](#)). Thus, the pathway-specific activation of thalamic afferents in acute brain slices reveals an input-dependent, SAT-induced change in L5 response properties, something that would not have been easy to decipher from sensory stimulation *in vivo*.

L2 Pyr neurons receive minimal direct input from VPM but do receive strong ascending input from L4. Consistent with this circuitry, optogenetic activation of VPM afferents in control samples was sufficient to drive firing in a fraction of L2 Pyr neurons (38%) that was delayed compared to spikes generated in deeper layers (L2 24.3 ± 0.5 ms; L4 10.1 ± 1.5 ms; L5 14.0 ± 1.7 ms; [Figure S2](#)). SAT did not change the fraction of spiking neurons (33%) or the mean evoked firing response during the stimulus period. As in L4, VPM stimulation did not elicit prolonged depolarization in the post-stimulus window for either L2 or L5 ([Figures 3E and 3O](#)).

Lack of changes in VPM-evoked firing could result from a homeostatic reduction in intrinsic excitability ([Lambo and Turriano, 2013](#); [Mrsic-Flogel et al., 2007](#)). However, input-output curves and resting membrane potential for L2, L4, and L5 excitatory neurons showed no significant increase in either property between control and SAT neurons ([Figure 4](#)).

Our assay indicates that VPM-associated cortical pathways do not become potentiated during learning. Although L4 to L2/3 synapses can undergo spike-timing-dependent plasticity (STDP) *in vitro* ([Banerjee et al., 2009](#)), the conditions engaged by SAT *in vivo* may not be sufficient to activate these well-described mechanisms. These results point to a special role

for POM during learning-related changes in cortical response properties.

Target-Specific Potentiation of POM Inputs

The increase in POM-evoked firing in both supra- and infragranular layers following 24 h of SAT, despite unchanged intrinsic firing properties, suggested that POM inputs to Pyr neurons might be strengthened after SAT. To compare the amplitude of quantal excitatory postsynaptic currents (qEPSCs) directly evoked from POM, we carried out voltage-clamp recordings from Pyr neurons during ChR2 activation of POM afferents in the presence of Sr^{2+} to desynchronize neurotransmitter release. This method enables detection of EPSCs from individual synapses and can provide evidence for postsynaptic plasticity ([Clem and Barth, 2006](#)).

The short latency of POM-evoked spikes in L5 Pyr, both in control and SAT neurons, suggested that these L5 neurons were firing as a direct result of POM input. Thus, we hypothesized that POM inputs to L5 Pyr might be potentiated. After 24 h of SAT, the mean amplitude of POM-mediated qEPSCs was significantly increased (control 15.7 ± 0.5 pA; 24 h SAT 20.7 ± 0.6 pA; [Figures 5F–5H](#)) and the cumulative distribution of qEPSC amplitudes showed a rightward shift ([Figure 5H](#)). Consistent with other studies where experience has driven pathway-specific excitatory strengthening ([Biane et al., 2016](#); [Clem and Barth, 2006](#)), we also observed a significant increase in the ratio of ChR2-evoked AMPA receptor: NMDA-receptor-mediated currents (A:N ratio; control 1.89 ± 0.13 ; 24 h SAT 2.84 ± 0.29 ; $p = 0.027$; [Figures S3A–S3C](#)). Although changes in presynaptic release properties are difficult to accurately assess with ChR2-evoked release, we examined our data for training-induced changes in the paired-pulse ratio (PPR) and found it unchanged at 24 h of SAT (control 0.39 ± 0.04 ; 24 h SAT 0.41 ± 0.06 ; [Figures S3D–S3F](#)).

Although POM-recipient L2 Pyr neurons also exhibited an increase in evoked firing after 24 h SAT, mean POM-qEPSC amplitudes were unchanged between control and SAT neurons (control 17.8 ± 0.8 pA; 24 h SAT 18.3 ± 0.8 pA; [Figures 5B–5D](#)). Significant differences were not observed even in the cumulative distribution of qEPSCs, suggesting that this pathway is unaltered at this time point in learning ([Figure 5D](#)).

Synaptic strengthening at POM to L5 synapses may be well positioned to facilitate further changes in neocortical circuitry, because these neurons show short spike latencies in response to POM stimulation and increase their POM-evoked firing after SAT ([Figures 2 and S1](#)). The lack of even modest potentiation

Figure 3. No Change in VPM-Evoked Cortical Activity after 24 h of SAT

(A) Schematic of experiment with recordings performed in ChR2-injected mice after 24 h of acclimation and 24 h of SAT.
 (B) Schematic of VPM axonal labeling and laminar pyramidal neuron recording site in L2.
 (C and D) VPM-evoked activity (blue bars; 5 pulses; 5 ms; 80 ms ISI) in L2 Pyr neurons of control animals that received 24 h of acclimation (left, black) and 24 h SAT (blue, right). Pie chart shows fraction of neurons that generated any action potentials following stimulation. Example cell response (top) shows 10 consecutive trials for an individual neuron. Raster (middle) shows spiking activity on 10 consecutive trials for 8 example cells. Global PSTH (bottom, 10 ms bins) shows average firing frequency across all cells in a population.
 (E) Average firing frequency across all cells during the 500 ms preceding VPM stimulation (pre), during stimulation (stim), and directly following stimulation (post).
 (F) Overlay of VPM-evoked spiking activity for CTL (black) and SAT24 (blue) animals.
 (G–K) Same as (C)–(F) but for L4 excitatory neurons.
 (L–P) Same as (C)–(F) but for L5 Pyr neurons.
 Averages represented as mean \pm SEM. See also [Figure S2](#).

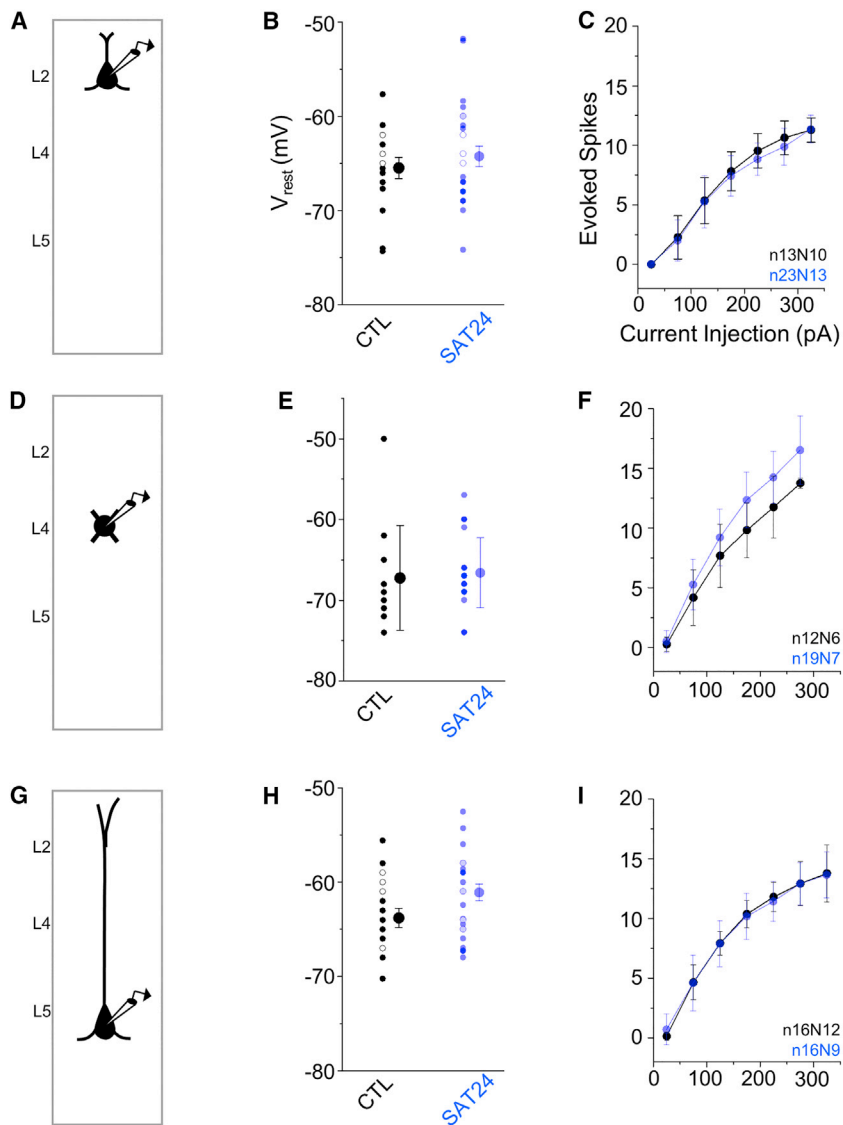


Figure 4. No Change in Intrinsic Properties of Cortical Excitatory Neurons after SAT

(A) Schematic of experimental setup. (B) Average resting membrane potential of L2 Pyr neurons from CTL (black) and SAT24 (blue) animals. Open circles indicate cells recorded in the presence of the synaptic blockers picrotoxin (50 μ M), APV (50 μ M), and NBQX (25 μ M) and are included in the displayed average. (C) Average spike count during 500 ms current injections (25 pA steps) for L2 Pyr neurons from CTL (black) and SAT24 (blue) animals. Displayed values include a subset of cells recorded in the presence of synaptic blockers described in (B). (D–F) Same as (A)–(C) but for L4 excitatory neurons. (G–I) Same as (A)–(C) but for L5 Pyr neurons. Averages represented as mean \pm SEM.

Thus, we hypothesized that POM-initiated, delayed synaptic input from deeper layers could contribute to the polysynaptic EPSPs and spiking activity observed in L2 Pyr. To test this, we compared the POM-evoked response properties of L2 Pyr neurons, before and after mechanical separation of supra- and infragranular layers, in acute brain slices from mice trained for 24 h (Figures 6A–6D). Neurons were paired from the same region of the same slice before and after transection to ensure the initial presence of POM-evoked activity. The incision through L4 should not affect light-evoked neurotransmitter release, because channelrhodopsin can drive vesicle release directly by illumination of the synaptic terminal.

Slice transection completely abolished POM-evoked action potentials in L2 Pyr neurons (Figures 6D and 6G), without changing resting membrane potential (control -66.7 ± 1.9 mV; 24 h SAT -69.0 ± 1.9 mV). EPSP onset latency and slope were not altered (onset latency: control 4.6 ± 0.3 ms; 24 h SAT 5.0 ± 0.6 ms; slope: control 0.37 ± 0.06 ; 24 h SAT 0.35 ± 0.09), but EPSP peak latency was significantly shorter (control 38.3 ± 3.8 ms; 24 h SAT 26.9 ± 1.5 ms), consistent with the loss of polysynaptic input originating in infragranular layers. After slice transection, POM stimulation no longer initiated prolonged depolarization in L2, indicating that POM-driven activity from deeper layers is also important in initiating or maintaining activity in the post-stimulus window. These data indicate that the increase in POM-evoked L2 firing relies upon convergent excitation from both POM and infragranular layers.

Intracortical Changes Are Not Present at 24 h of Training

The increase in L2 POM-evoked firing after 24 h of SAT could result solely from the increase in synaptic drive from POM to L5 that is inherited by L2 Pyr or could occur concurrently with excitatory synaptic potentiation within L5 or at L5 inputs to L2

at POM to L2 synapses at this time point suggests that the conditions for the induction of synaptic plasticity are different between L2 and L5 Pyr neurons in the behaving animal.

Elevated POM-Evoked Activity Is Driven by Ascending Input from Infragranular Layers

SAT increases POM-evoked firing of L2 Pyr neurons both during and after the optogenetic stimulus window. If POM inputs to L2 are not potentiated after 24 h of SAT, we hypothesized that this increased activity may be inherited from spiking in other cortical layers. For example, POM-evoked spikes in L5 occur within 15 ms of the optogenetic pulse, and ascending input from L5 Pyr to L2 could summate with direct POM EPSPs in L2 neurons to drive firing in these neurons. Indeed, POM-evoked EPSPs in L2 Pyr neurons were often complex, where individual trials showed multiple inflection points during the EPSP rise time that correspond to asynchronous synaptic inputs (Figure 6).

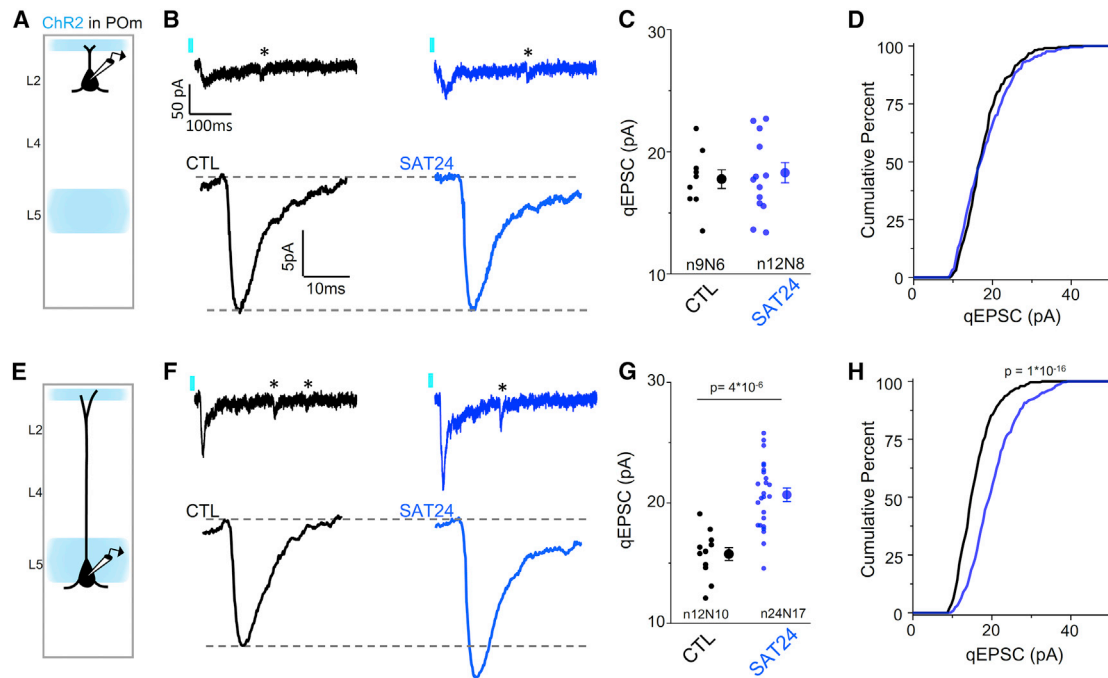


Figure 5. Twenty-four Hours of SAT Strengthens POM Synaptic Inputs onto L5 Pyr Neurons

(A) Schematic of experimental setup in L2 Pyr neurons.

(B) (Top) Example single trial showing Sr^{2+} -desynchronized POM-evoked response in a L2 Pyr neuron where individual, isolated quantal events (*) follow an initial multiquantal response. (Bottom) Global average qEPSCs from control animals (black, left) or in animals that received 24 h of SAT (blue, right) is shown. All well-isolated light-evoked qEPSCs in a cell (≥ 25 for inclusion) were aligned to rise time and averaged to generate an average cellular POM qEPSC. Cell averages were aligned to rise and averaged to generate global average qEPSC for each condition.

(C) Quantification of mean qEPSC amplitude for each cell, measured as the mean of individual qEPSC peak amplitudes within a cell.

(D) Cumulative distribution histogram of POM qEPSC amplitudes for CTL (black) and SAT24 (blue) animals. Distributions comprise 25 randomly selected events from each cell, compared using a Kolmogorov-Smirnov (K-S) test.

(E–H) Same as (A)–(D), but for L5 Pyr neurons.

Averages represented as mean \pm SEM. See also Figure S3.

Pyr. We examined both pathways using Sr^{2+} -replaced artificial cerebrospinal fluid (ACSF) to isolate pathway-specific quantal EPSCs.

To examine SAT-associated changes within the L5 local excitatory circuit, we used an extracellular stimulating electrode placed in L5 and recorded quantal EPSCs in L5 Pyr neurons (Figure S4). Although this method cannot isolate inputs from specific types of presynaptic neurons, the dense local connectivity within a cortical layer (Lefort et al., 2009) suggests that the majority of excitatory inputs will be from nearby neurons. Under these conditions, evoked L5 qEPSCs showed no difference between control and trained animals (control 19.6 ± 0.4 pA; 24 h SAT 19.85 ± 0.5 pA; Figures S4A–S4D).

To test whether L5 to L2 qEPSCs were increased after 24 h SAT, we expressed ChR2 specifically in L5 neurons using the *Etv1-Cre* driver line for ChR2 expression. A comparison of optogenetically evoked L5 qEPSC amplitudes in L2 Pyr neurons showed no difference between control and trained animals (control 20.1 ± 1.1 pA; 24 h SAT 19.6 ± 1.4 pA; Figures S4E–S4H). qEPSCs in L2 Pyr neurons evoked using an extracellular stimulating electrode in L2 also did not show any increase in mean amplitude after 24 h of training (Figures S4I–S4K). These results suggest that the training-induced increase in POM-initiated

spiking activity in L2 Pyr cannot easily be attributed to synaptic strengthening of intralaminar L5–L5 or translaminar L5 to L2 excitatory inputs and that this activity is driven by increased L5 Pyr firing associated with potentiation of POM to L5 excitatory inputs.

Sensory Stimulation Alone Does Not Drive POM Plasticity

To determine whether POM to L5 synaptic changes were linked to learning or more generally to stimulus exposure during the training period, we created a pseudotraining paradigm with identical trial and stimulus structure but with altered reward contingency such that the whisker stimulus no longer predicted reward (Figure 7A). Similar to SAT, we observed a transient dip in anticipatory lick rates following stimulation presentation early in training that rapidly recovered, indicating that the animals perceived the stimulus and rapidly habituated to it.

After 24 h of pseudotraining, animals displayed no difference in anticipatory licking behavior between stimulus trials and non-stimulus trials, as both trial types had a 50% chance of water delivery (Figures 7B and 7C). Importantly, the number of air-puff-exposed trials for pseudotrained animals was greater than the mean trial number for SAT animals (Figure 7C), likely

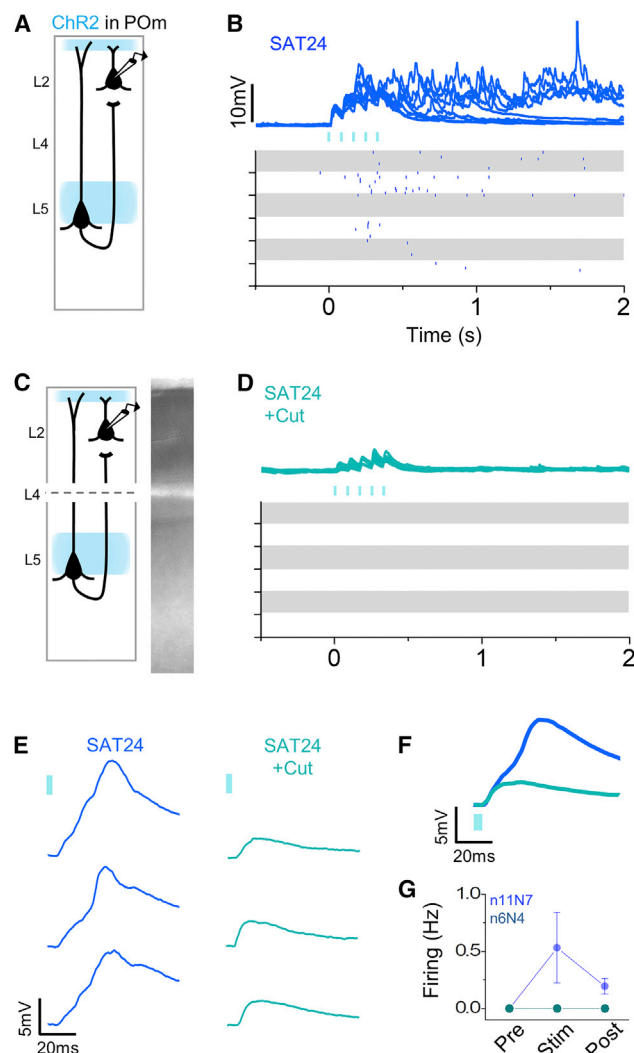


Figure 6. Elevated POM-Evoked Activity in Trained Animals Is Driven by Ascending Input from Infragranular Layers

(A) Schematic of experimental setup for ChR2-evoked firing of pre-cut L2 Pyr neurons in SAT24 animals.

(B) Light-evoked activity (blue bars; 5 pulses; 5 ms; 80 ms ISI) on 10 consecutive trials in an example L2 Pyr cell (top) and for 10 example cells (bottom) in SAT24 animals.

(C and D) Same as (A) and (B) but for L2 Pyr neurons after a mechanical incision through cortical L4. Each collected post-cut cell had at least one recorded L2 Pyr recording in the same slice prior to cut, and example cells in (B) and (D) were recorded in the same slice.

(E) Comparison of subthreshold responses on 3 consecutive sweeps following the first light pulse for example cells in (B) and (D).

(F) Average response (10 consecutive sweeps) to first light pulse for SAT24 (dark blue) and SAT24 + cut (light blue) example cells in (B) and (D).

(G) Average firing frequency across all cells during the 500 ms preceding VPM stimulation (pre), during stimulation (stim), and directly following stimulation (post).

Averages represented as mean \pm SEM. See also Figure S4.

because water was only provided on 50% of trials versus 80% during SAT, requiring more trials to receive the same amount of water.

To determine whether synaptic strengthening of POM inputs in L5 Pyr neurons was specific to sensory learning or could occur with passive sensory stimulation, POM qEPSC input strength was recorded in pseudotrained animals. After 24 h, mean POM synaptic strength onto L5 neurons was unchanged from control values and significantly smaller than POM qEPSCs in animals that had undergone SAT (control 15.8 ± 0.53 pA; pseudotrained 15.8 ± 0.59 pA; Figures 7E–7G). Thus, sensory stimulation decoupled from reward is not sufficient to drive POM thalamocortical plasticity at L5 Pyr neurons.

To determine whether POM input plasticity was specifically related to the rewarded whisker-deflection stimulus or might reflect the activation of non-specific sensory cues related to training, we replaced the air puff with a different sensory stimulus, a 500 ms-long light flash followed by a 500 ms delay to water delivery (Figure S5A). The light flash was delivered unilaterally from a triggered light-emitting diode (LED) at the same location as the air puff in other trials and generated a transient suppression in licking frequency at the onset of training, indicating animal detection. However, association of the light flash with the water reward proved more difficult to acquire, as animals did not display a behavioral change in anticipatory licking, even after 48 h of light-flash association training (Figure S5B). POM to L5 Pyr qEPSC amplitudes did not change with light-association training (Figures S5D and S5E). These data indicate that POM input plasticity in S1 occurs only when the stimulus is both whisker dependent and directly coupled to the water reward and that non-tactile arousal or attentional cues are unlikely to explain the pathway-specific changes observed.

Pathway-Specific Changes in L2 Pyr Neurons after SAT

The absence of plasticity at POM inputs to L2 Pyr neurons after 24 h SAT might suggest that the conditions required for synaptic potentiation have not yet been met or it could mean that L2 neurons do not possess the machinery for learning-dependent synaptic plasticity. To test whether longer periods of SAT might be sufficient to change POM input strength in L2 Pyr neurons, we examined ChR2-evoked POM-mediated qEPSC amplitudes at a later time point. After 48 h of SAT, task performance was further enhanced, driven primarily by an increase in stimulus-associated licking and not a depression of licking in blank trials (Figures 8B and 8C).

After 48 h SAT, qEPSC amplitude of POM to L2 Pyr neurons was significantly increased (control 17.8 ± 0.77 pA; 48 h SAT 22.2 ± 0.49 pA; Figures 8D–8G). The increase in POM input to L2 Pyr was not matched by a further enhancement of POM input strength onto L5 Pyr (Figures 8H–8K). Did this increase in POM response amplitude in L2 require 2 days of SAT or might it be a time-dependent process initiated during the first 24 h after training? To test this possibility, animals were trained for 24 h and then allowed to drink ad lib from the lick port without air puff stimulation for an additional 24 h. In this group, POM to L2 qEPSC amplitude was unchanged from control (control 17.8 ± 0.77 pA; 2 h SAT plus 24 h no stimulation 16.8 ± 0.43 pA; $n = 10$ cells; $N = 4$ animals).

Taken together, these data indicate that the absence of POM thalamocortical plasticity in L2 Pyr at 24 h SAT is not dependent

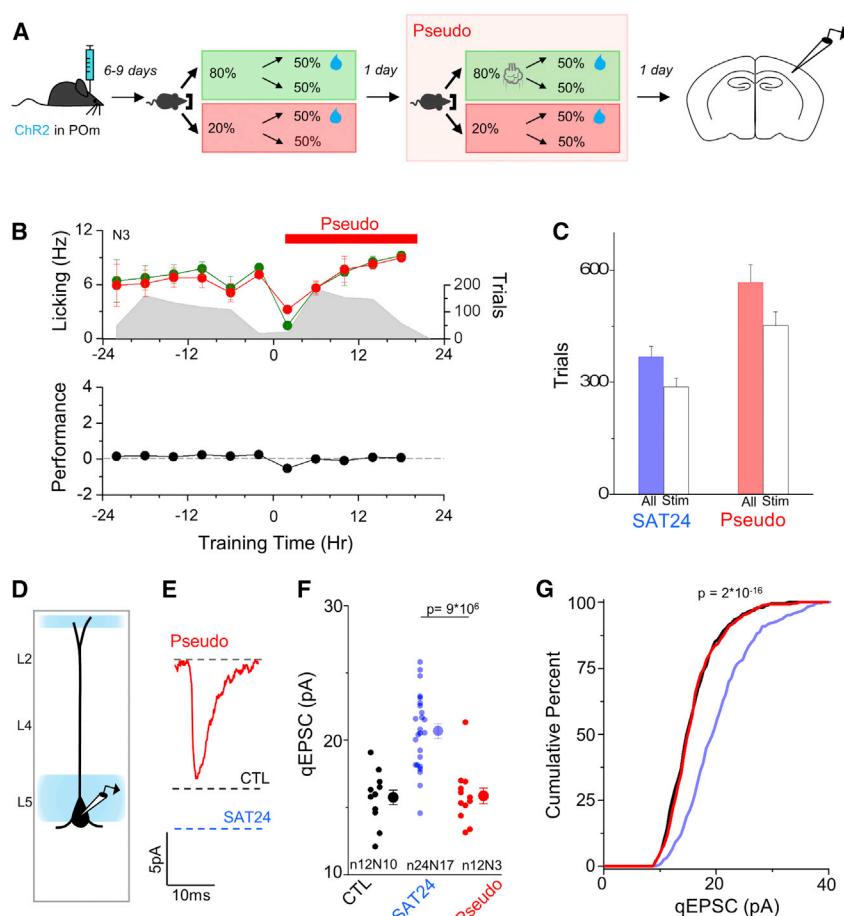


Figure 7. Sensory Stimulation Alone Does Not Drive POM Plasticity

(A) Schematic of experimental setup and pseudo-training behavioral paradigm. Cages, training structure, stimulus, and timing were identical to SAT, but water delivery (US) was uncoupled from (CS) and randomly delivered on 50% of trials, regardless of CS presentation.

(B) Time course of anticipatory lick rates (left axis; 300 ms prior to water delivery; 4 h bins) over the course of training for blank (20%, red) and stim trials (80%, green) averaged across all animals. Trial initiation counts (right axis) are shown in gray for the same time bins; red bar denotes pseudotraining period.

(C) Comparison of animal-initiated trial counts during 24 h of SAT (blue) and pseudotrained (red). Solid bars indicate total trials received, and white bars show the number of stimulus (CS) trials received.

(D) Schematic of experimental setup.

(E) Global average qEPSC in pseudotrained animals. All well-isolated, light-evoked qEPSCs in a cell (≥ 25) were aligned to rise time and averaged to generate an average cellular POM qEPSC. Cell averages were aligned to rise and averaged to generate global average qEPSC. Global average POM qEPSC amplitudes in control (black) and SAT24 (blue) animals for comparison are shown.

(F) Quantification of average qEPSC amplitude for each cell, measured as the average of individual qEPSC peak amplitudes within a cell. CTL and SAT24 replotted from Figure 4.

(G) Cumulative distribution histogram of POM qEPSC amplitudes for CTL (black) and SAT24 (blue) animals and pseudotrained animals (red). Distributions comprise 25 randomly selected events from each cell, and stats are for a K-S test between 24 h and pseudo.

Averages represented as mean \pm SEM. See also Figure S5.

on time since training initiation. Rather, we find that continuous training is important and 48 h of sensory association is sufficient to increase POM input strength onto L2 Pyr neurons.

Synaptic plasticity between L2 Pyr neurons in barrel cortex has been well-documented, both in acute brain slices and after *in vivo* sensory experience (Albieri et al., 2015; Cheetham et al., 2008; Rodríguez-Moreno and Paulsen, 2008; Wen and Barth, 2011); however, rewiring of this local network has not been linked to sensory learning. To determine whether POM thalamocortical plasticity during SAT occurred concurrent with changes in the L2 excitatory network, we examined changes in qEPSC amplitudes after 48 h SAT measured from local afferent stimulation, using an extracellular electrode placed in L2. At this stage, qEPSC amplitudes were significantly increased in L2 Pyr neurons compared to control and 24 h values (Figures S4I–S4L).

We also examined whether L5 to L2 or L5 to L5 qEPSC input strength might also increase after 48 h of SAT but did not detect a change in either pathway (Figures S4A–S4H). Overall, these data show that SAT-initiated increases in L2 excitatory strength are delayed with respect to L5 and that L2 synaptic potentiation may be pathway specific, detectable at thalamocortical, but not

within the aggregate, excitatory input to L2 and L5a Pyr neurons after 48 h of SAT.

We also examined the relationship between mean qEPSC amplitude for a given animal and metrics for trial completion and task performance (anticipatory licking) during SAT. No statistically significant, positive relationship between increased qEPSC amplitude and number of trials or animal performance was observed at either 24 or 48 h SAT for L2 and L5 Pyr neurons (Figure S6). However, qEPSC values can differ widely across cells, even within an animal, and in some cases, we had only a single qEPSC measurement from a given subject. Interestingly, POM input strength at L2 synapses was negatively correlated with animal trial number at 48 h of SAT ($p = 0.002$), suggesting a possible activity-dependent depotentiation (Clem et al., 2008).

DISCUSSION

We aimed to develop a comprehensive account of how a learned sensory-association task alters synaptic function in neocortical circuits, with isolation of specific input pathways and targeted recordings from specific cell types across different layers of the cortex. High-throughput, home-cage sensory-association

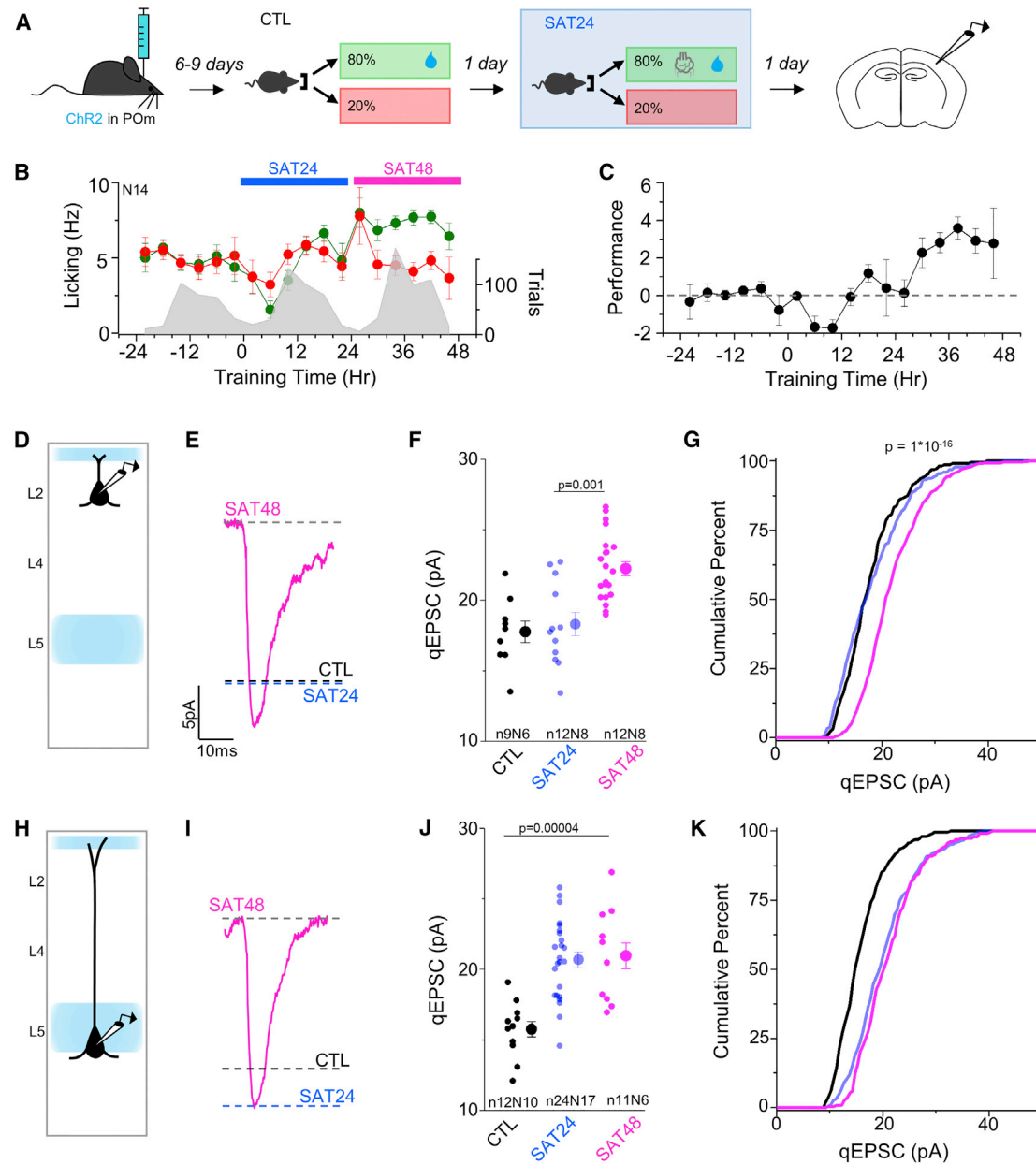


Figure 8. SAT Drives Sequential Thalamocortical Plasticity in L5 and then L2 Pyramidal Neurons

(A) Schematic of experiment with recordings performed in ChR2-injected mice after 24 h of acclimation and 48 h of SAT.

(B) Time course of anticipatory lick frequency (left axis; 300 ms prior to water delivery; 4-h bins) over the course of acclimation and training for blank (red) and stim+water trials (green) averaged across all animals ($N = 14$ for all time points). Trial initiation counts (right axis) are shown in gray for the same time bins.

(C) Quantification of performance defined as the difference in anticipatory lick rates between stim+water trials and blank trials during learning for the 4-h bins shown in (B).

(D) Schematic of experimental setup for recording POM qEPSCs in L2 Pyr neurons.

(E) Global average qEPSC in SAT48 animals. All well-isolated, light-evoked qEPSCs in a cell (≥ 25) were aligned to rise time and averaged to generate an average cellular POM qEPSC. Cell averages were aligned to rise and averaged to generate global average qEPSC. Global average POM qEPSC amplitudes in control (black) and SAT24 (blue) animals for comparison are shown.

(F) Quantification of average qEPSC amplitude for each cell, measured as the average of individual qEPSC peak amplitudes within a cell for control (black), SAT24 (blue), and SAT48 (magenta) animals.

(G) Cumulative distribution histogram of POM qEPSC amplitudes for CTL (black) and SAT24 (blue) animals and SAT48 animals (magenta). Distributions comprise 25 randomly selected events from each cell; K-S test compares pseudo and 24 h.

(H-K) Same as (D)-(G) but for L5 Pyr neurons.

Averages represented as mean \pm SEM. See also Figure S6.

learning revealed that thalamocortical synapses are a site of early synaptic change in primary somatosensory cortex, with selectivity for POM-related, but not VPM-related, pathways. Synaptic potentiation at POM inputs onto L5 Pyr neurons was then followed by POM input potentiation to L2 Pyr neurons and within the L2 excitatory circuit over subsequent days of training. Importantly, changes were not observed with pseudoconditioning, indicating that they were not driven by sensory stimulation alone. These data show that sensory association training drives sequential changes in excitatory synaptic strength in an input- and layer-specific manner and lays a foundation for understanding how learning alters the flow of information across the cortical column.

Thalamocortical Plasticity during Learning

Thalamocortical connections, particularly from VPM to L4, have been considered the major source of cortical input to somatosensory cortex, although there is an increasing awareness that POM inputs also provide significant drive (Audette et al., 2018; Gambino et al., 2014; Jouhannau et al., 2014; Mease et al., 2016). Abundant experimental evidence indicates VPM inputs are resistant to experience-dependent alterations after the early postnatal period (Feldman and Brecht, 2005). However, prior studies have largely ignored the potential role of POM in neocortical response plasticity. We used pathway-specific activation of VPM and POM thalamic inputs and targeted recordings in the cortex as a screen for learning-related changes across the cortical column. VPM-evoked cortical firing was remarkably stable after early sensory association training in L4 and L5, as well as in the downstream target L2/3. In contrast, POM-evoked firing was dramatically increased. POM input plasticity may contribute to increased evoked firing in L5 Pyr neurons following sensory experience (Diamond et al., 1994; Jacob et al., 2012; Ward et al., 2012).

What might explain the difference in plasticity between VPM and POM pathways? L4 neurons may be particularly resistant to post-critical period plasticity (Crair and Malenka, 1995; Fox, 1992), but we did not observe enhanced firing elsewhere in the VPM pathway for Pyr neurons in L2/3 and L5. Although we cannot rule out target-specific changes in VPM input strength—for example, opposite changes in VPM drive to excitatory and inhibitory neurons that result in no net change in firing output—our assay clearly revealed an increase in response output in POM-recipient layers. Differences in plasticity induction could be attributed to the pattern or duration of sensory-evoked activity in VPM and POM during sensory stimulation (Sobolewski et al., 2015). Because POM is more strongly influenced by descending cortical drive than VPM, it is possible that recurrent corticothalamic circuitry may prime POM circuits for learning-dependent plasticity. Finally, our assay revealed differences between VPM and POM-evoked intracortical dynamics. Optogenetic activation of VPM inputs never evoked prolonged depolarization in any cortical layers, whereas POM activation frequently did. This sub- and suprathreshold activity may be permissive to plasticity induction that is specific to POM-recipient circuitry.

The capacity for learning-induced strengthening of thalamic inputs, although unexpected in the sensory cortex, has been

previously described in the motor cortex (Biane et al., 2016). Rapid learning-related thalamic input plasticity may be an important feature of analogous higher-order sensory thalamic nuclei, such as the pulvinar (Arcaro et al., 2015; Purushothaman et al., 2012; Roth et al., 2016).

Sequence for Cortical Rewiring during Sensory Learning

Behavioral evidence for learning emerged early during SAT, and performance increased with longer training periods. The slow trajectory of learning enabled identification of progressive synaptic changes in the neocortex. Increased POM-evoked firing in both deep and superficial layers occurred at the same time as synaptic strengthening of POM inputs to L5 Pyr neurons, after only 24 h of training. The short latency of evoked spikes and the absence of enhanced excitability in L5 Pyr suggest synaptic strengthening at POM inputs to L5 Pyr drove the increase in spiking. Although L2 Pyr neurons also showed an increase in POM-evoked firing after 24 h of SAT, synaptic changes at thalamic or intracortical inputs to L2 were absent, and slice transection data indicate that this increased drive was indirect, arising from ascending input from infragranular layers. However, 48 h of SAT was sufficient to drive both POM to L2 Pyr synaptic strengthening as well as intralaminar plasticity in local L2 excitatory circuits.

Why was synaptic plasticity in L2 delayed relative to POM plasticity in L5? One possibility is that activity levels in superficial layers of the sensory cortex are too low initially to engage activity-dependent plasticity mechanisms. It is therefore notable that the increase in POM-evoked activity in L2 is evident prior to any observable change in synaptic strength in these neurons. Early synaptic strengthening in the more robustly active L5 could play a role in enhancing stimulus-evoked activity in L2 and drive POM input potentiation. Our results identify POM thalamic inputs to L5 as the cortical “first responder” for training-evoked cortical plasticity. It will be of great interest to identify the dependent processes that underlie the sequential pattern of cortical changes revealed in this study.

Notably, we did not find evidence for synaptic potentiation at translaminar L5 to L2 synapses or within the L5 excitatory circuit, despite well-documented anatomical pathways and prior experimental evidence that these connections can be plastic under some circumstances (Sjöström and Häusser, 2006). Our assays may have been too coarse to detect subtle changes between specific subtypes of L5 Pyr neurons or opposing changes within the same pathway. Learning-dependent changes in inhibitory synapses will be an important direction of future investigations.

Learning-Related Changes in Neocortical Circuits

POM-related synaptic changes, despite proceeding at the same timescale as behavioral learning, could have been generated by repeated sensory stimulation rather than learning. Although the air puff stimulus is likely to be particularly salient and thus a good activator of POM circuits (Jouhannau et al., 2014; Sosnik et al., 2001), it is notable that pseudoconditioning did not generate any change in POM input strength in L5 Pyr despite delivering the same or even a greater number of air puff stimuli to animals, decoupled from a water reward. These findings

indicate that synaptic plasticity in primary sensory cortex is strongly influenced by contingent reward, not just stimulus exposure.

Do cortical synaptic changes relate to sensory perception and task performance? POM input strength and task performance were not significantly correlated, possibly due to high variability in qEPSC measurements, which were often averaged from a small number of cells per animal due to the difficulty of obtaining these data from a given animal. However, the weak relationship between qEPSC values and the number of training trials suggest a progressive link between stimulus-reward associations and the degree of synaptic strengthening at POM inputs.

Wide-ranging approaches suggest varied, and often contradictory, locations and mechanisms underlying perceptual and associative learning, but perceptual learning has been more closely linked to changes in primary sensory cortex (Caras and Sanes, 2017; Makino et al., 2016; McGann, 2015). POM and analogous higher order thalamic nuclei interface between primary sensory cortex, association cortex, motor cortex, and the striatum, making them well positioned to integrate and transmit diverse information that may be important both for learning and initiation of cortical plasticity.

High-Throughput Home-Cage Behavioral Training

Analysis of learning-dependent reorganization of excitatory circuits across the cortical column with layer- and input-specific resolution was facilitated by a high-throughput home-cage training system, optimized for training large numbers of animals with minimal handling. Training occurred during the animal's normal active period and did not involve animal handling, a significant source of stress that may impair learning (Francis and Kanold, 2017). Sensory association training using a gentle air puff stimulus drove progressive changes in behavior that occurred quickly enough to enable detailed electrophysiological investigation but were slow enough to capture sequential changes in synaptic function across the cortical column (Poddar et al., 2013).

The training setup we developed was easily adapted to alter the contingency between water reward and conditioned stimulus for pseudotraining and could be further modified to alter other task parameters, such as reward valence, frequency, or sensory stimuli delivered. Home-cage training paradigms will be particularly useful for integrating cutting-edge recording, imaging, and stimulating technologies in freely moving animals.

Conclusions

Our analysis of synaptic changes reveals the footprints of plasticity that are induced during an associative learning task that might be difficult to isolate during dynamic recording of neural activity. Importantly, our study did not address how SAT activates thalamic and neocortical neurons *in vivo* or how it alters thalamic response properties after training. Future experiments with fine-scale cell type and temporal resolution will illuminate how activity in VPM, POM, and specific layers of the neocortex are engaged by and changed during learning.

The progressive emergence of POM plasticity at infragranular synapses followed by POM input potentiation at L2 Pyr neurons and an increase in intralaminar L2-L2 excitatory synaptic

strength suggests that supragranular layers may have a higher threshold but a large capacity for experience-dependent plasticity. These findings will be essential to develop and test data-constrained models of synaptic change and neural spiking.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.neuron.2019.04.037>.

ACKNOWLEDGMENTS

Special thanks to Joanne Steinmiller for expert animal care; Rogan Grant, Zhuopin Sun, Alexander Hsu, and Megumi Matsushita for technical assistance; and members of the Barth Lab for helpful comments. Special thanks also to Joanna Urban-Ciecko for essential contributions to the revised manuscript. This work was supported by the Carnegie Mellon University Hillman Presidential Fellowship (N.J.A.), the IISC-CMU BrainHub postdoctoral fellowship (A.R.), and NIH R01NS088958 (to A.L.B.).

AUTHOR CONTRIBUTIONS

Conceptualization, N.J.A. and A.L.B.; Methodology, N.J.A., S.M.B., and A.L.B.; Software, S.M.B.; Formal Analysis, N.J.A.; Investigation, N.J.A., L.T.S., and A.R.; Writing – Original Draft, N.J.A. and A.L.B.; Writing – Reviewing & Editing, N.J.A., S.M.B., L.T.S., A.R., and A.L.B.; Visualization, N.J.A.; Supervision, A.L.B.; Funding Acquisition, N.J.A., A.R., and A.L.B.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: July 16, 2018
 Revised: February 11, 2019
 Accepted: April 25, 2019
 Published: May 28, 2019

REFERENCES

Albieri, G., Barnes, S.J., de Celis Alonso, B., Cheetham, C.E.J., Edwards, C.E., Lowe, A.S., Karunaratne, H., Dear, J.P., Lee, K.C., and Finnerty, G.T. (2015). Rapid bidirectional reorganization of cortical microcircuits. *Cereb. Cortex* 25, 3025–3035.

- Alloway, K.D., Hoffer, Z.S., and Hoover, J.E. (2003). Quantitative comparisons of corticothalamic topography within the ventrobasal complex and the posterior nucleus of the rodent thalamus. *Brain Res.* 968, 54–68.
- Arcaro, M.J., Pinsk, M.A., and Kastner, S. (2015). The anatomical and functional organization of the human visual pulvinar. *J. Neurosci.* 35, 9848–9871.
- Audette, N.J., Urban-Ciecko, J., Matsushita, M., and Barth, A.L. (2018). POM thalamocortical input drives layer-specific microcircuits in somatosensory cortex. *Cereb. Cortex* 28, 1312–1328.
- Banerjee, A., Meredith, R.M., Rodríguez-Moreno, A., Mierau, S.B., Auberson, Y.P., and Paulsen, O. (2009). Double dissociation of spike timing-dependent potentiation and depression by subunit-preferring NMDA receptor antagonists in mouse barrel cortex. *Cereb. Cortex* 19, 2959–2969.
- Biane, J.S., Takashima, Y., Scanziani, M., Conner, J.M., and Tuszynski, M.H. (2016). Thalamocortical projections onto behaviorally relevant neurons exhibit plasticity during adult motor learning. *Neuron* 89, 1173–1179.
- Bureau, I., von Saint Paul, F., and Svoboda, K. (2006). Interdigitated paralemniscal and lemniscal pathways in the mouse barrel cortex. *PLoS Biol.* 4, e382.
- Caras, M.L., and Sanes, D.H. (2017). Top-down modulation of sensory cortex gates perceptual learning. *Proc. Natl. Acad. Sci. USA* 114, 9972–9977.
- Chandrasekaran, S., Navlakha, S., Audette, N.J., McCreary, D.D., Suhan, J., Bar-Joseph, Z., and Barth, A.L. (2015). Unbiased, high-throughput electron microscopy analysis of experience-dependent synaptic changes in the neocortex. *J. Neurosci.* 35, 16450–16462.
- Cheetham, C.E.J., Hammond, M.S.L., McFarlane, R., and Finnerty, G.T. (2008). Altered sensory experience induces targeted rewiring of local excitatory connections in mature neocortex. *J. Neurosci.* 28, 9249–9260.
- Clem, R.L., and Barth, A. (2006). Pathway-specific trafficking of native AMPARs by in vivo experience. *Neuron* 49, 663–670.
- Clem, R.L., Celikel, T., and Barth, A.L. (2008). Ongoing in vivo experience triggers synaptic metaplasticity in the neocortex. *Science* 319, 101–104.
- Cohen, J.Y., Haesler, S., Vong, L., Lowell, B.B., and Uchida, N. (2012). Neuron-type-specific signals for reward and punishment in the ventral tegmental area. *Nature* 482, 85–88.
- Crair, M.C., and Malenka, R.C. (1995). A critical period for long-term potentiation at thalamocortical synapses. *Nature* 375, 325–328.
- Cruikshank, S.J., Urabe, H., Nurmikko, A.V., and Connors, B.W. (2010). Pathway-specific feedforward circuits between thalamus and neocortex revealed by selective optical stimulation of axons. *Neuron* 65, 230–245.
- Diamond, M.E., Huang, W., and Ebner, F.F. (1994). Laminar comparison of somatosensory cortical plasticity. *Science* 265, 1885–1888.
- Feldman, D.E., and Brecht, M. (2005). Map plasticity in somatosensory cortex. *Science* 310, 810–815.
- Feldmeyer, D., Brecht, M., Helmchen, F., Petersen, C.C.H., Poulet, J.F.A., Staiger, J.F., Luhmann, H.J., and Schwarz, C. (2013). Barrel cortex function. *Prog. Neurobiol.* 103, 3–27.
- Fox, K. (1992). A critical period for experience-dependent synaptic plasticity in rat barrel cortex. *J. Neurosci.* 12, 1826–1838.
- Francis, N.A., and Kanold, P.O. (2017). Automated operant conditioning in the mouse home cage. *Front. Neural Circuits* 11, 10.
- Galvez, R., Weiss, C., Weible, A.P., and Disterhoft, J.F. (2006). Vibrissa-signaled eyeblink conditioning induces somatosensory cortical plasticity. *J. Neurosci.* 26, 6062–6068.
- Gambino, F., Pagès, S., Kehayas, V., Baptista, D., Tatti, R., Carleton, A., and Holtmaat, A. (2014). Sensory-evoked LTP driven by dendritic plateau potentials in vivo. *Nature* 515, 116–119.
- Glazewski, S., and Barth, A.L. (2015). Stimulus intensity determines experience-dependent modifications in neocortical neuron firing rates. *Eur. J. Neurosci.* 41, 410–419.
- Glazewski, S., and Fox, K. (1996). Time course of experience-dependent synaptic potentiation and depression in barrel cortex of adolescent rats. *J. Neurophysiol.* 75, 1714–1729.
- Gong, S., Doughty, M., Harbaugh, C.R., Cummins, A., Hatten, M.E., Heintz, N., and Gerfen, C.R. (2007). Targeting Cre recombinase to specific neuron populations with bacterial artificial chromosome constructs. *J. Neurosci.* 27, 9817–9823.
- Groh, A., Bokor, H., Mease, R.A., Plattner, V.M., Hangya, B., Stroh, A., Deschenes, M., and Acsády, L. (2014). Convergence of cortical and sensory driver inputs on single thalamocortical cells. *Cereb. Cortex* 24, 3167–3179.
- Harris, J.A., Harris, I.M., and Diamond, M.E. (2001). The topography of tactile working memory. *J. Neurosci.* 21, 8262–8269.
- Jacob, V., Petreanu, L., Wright, N., Svoboda, K., and Fox, K. (2012). Regular spiking and intrinsic bursting pyramidal cells show orthogonal forms of experience-dependent plasticity in layer V of barrel cortex. *Neuron* 73, 391–404.
- Jouhanneau, J.S., Ferrarese, L., Estebanez, L., Audette, N.J., Brecht, M., Barth, A.L., and Poulet, J.F.A. (2014). Cortical fosGFP expression reveals broad receptive field excitatory neurons targeted by POM. *Neuron* 84, 1065–1078.
- Karni, A., and Sagi, D. (1991). Where practice makes perfect in texture discrimination: evidence for primary visual cortex plasticity. *Proc. Natl. Acad. Sci. USA* 88, 4966–4970.
- Kilgard, M.P., and Merzenich, M.M. (1998). Cortical map reorganization enabled by nucleus basalis activity. *Science* 279, 1714–1718.
- Lambo, M.E., and Turrigiano, G.G. (2013). Synaptic and intrinsic homeostatic mechanisms cooperate to increase L2/3 pyramidal neuron excitability during a late phase of critical period plasticity. *J. Neurosci.* 33, 8810–8819.
- Lefort, S., Tómm, C., Floyd Sarria, J.C., and Petersen, C.C.H. (2009). The excitatory neuronal network of the C2 barrel column in mouse primary somatosensory cortex. *Neuron* 61, 301–316.
- Lu, J., Tucciarone, J., Padilla-Coreano, N., He, M., Gordon, J.A., and Huang, Z.J. (2017). Selective inhibitory control of pyramidal neuron ensembles and cortical subnetworks by chandelier cells. *Nat. Neurosci.* 20, 1377–1383.
- Makino, H., Hwang, E.J., Hedrick, N.G., and Komiyama, T. (2016). Circuit mechanisms of sensorimotor learning. *Neuron* 92, 705–721.
- Masri, R., Trageser, J.C., Bezdudnaya, T., Li, Y., and Keller, A. (2006). Cholinergic regulation of the posterior medial thalamic nucleus. *J. Neurophysiol.* 96, 2265–2273.
- McGann, J.P. (2015). Associative learning and sensory neuroplasticity: how does it happen and what is it good for? *Learn. Mem.* 22, 567–576.
- Mease, R.A., Metz, M., and Groh, A. (2016). Cortical sensory responses are enhanced by the higher-order thalamus. *Cell Rep.* 14, 208–215.
- Meyer, H.S., Wimmer, V.C., Hemberger, M., Bruno, R.M., de Kock, C.P.J., Frick, A., Sakmann, B., and Helmstaedter, M. (2010). Cell type-specific thalamic innervation in a column of rat vibrissa cortex. *Cereb. Cortex* 20, 2287–2303.
- Mrsic-Flogel, T.D., Hofer, S.B., Ohki, K., Reid, R.C., Bonhoeffer, T., and Hübener, M. (2007). Homeostatic regulation of eye-specific responses in visual cortex during ocular dominance plasticity. *Neuron* 54, 961–972.
- Oberlaender, M., Ramirez, A., and Bruno, R.M. (2012). Sensory experience restructures thalamocortical axons during adulthood. *Neuron* 74, 648–655.
- Petreanu, L., Mao, T., Sternson, S.M., and Svoboda, K. (2009). The subcellular organization of neocortical excitatory connections. *Nature* 457, 1142–1145.
- Poddar, R., Kawai, R., and Ölveczky, B.P. (2013). A fully automated high-throughput training system for rodents. *PLoS ONE* 8, e83171.
- Porter, J.T., Johnson, C.K., and Agmon, A. (2001). Diverse types of interneurons generate thalamus-evoked feedforward inhibition in the mouse barrel cortex. *J. Neurosci.* 21, 2699–2710.
- Purushothaman, G., Marion, R., Li, K., and Casagrande, V.A. (2012). Gating and control of primary visual cortex by pulvinar. *Nat. Neurosci.* 15, 905–912.
- Riout-Pedotti, M.S., Friedman, D., and Donoghue, J.P. (2000). Learning-induced LTP in neocortex. *Science* 290, 533–536.
- Rodríguez-Moreno, A., and Paulsen, O. (2008). Spike timing-dependent long-term depression requires presynaptic NMDA receptors. *Nat. Neurosci.* 11, 744–745.

- Roth, M.M., Dahmen, J.C., Muir, D.R., Imhof, F., Martini, F.J., and Hofer, S.B. (2016). Thalamic nuclei convey diverse contextual information to layer 1 of visual cortex. *Nat. Neurosci.* **19**, 299–307.
- Schwartz, S., Maquet, P., and Frith, C. (2002). Neural correlates of perceptual learning: a functional MRI study of visual texture discrimination. *Proc. Natl. Acad. Sci. USA* **99**, 17137–17142.
- Shibata, K., Sasaki, Y., Kawato, M., and Watanabe, T. (2016). Neuroimaging evidence for 2 types of plasticity in association with visual perceptual learning. *Cereb. Cortex* **26**, 3681–3689.
- Sjöström, P.J., and Häusser, M. (2006). A cooperative switch determines the sign of synaptic plasticity in distal dendrites of neocortical pyramidal neurons. *Neuron* **51**, 227–238.
- Sobolewski, A., Kublik, E., Swiejkowski, D.A., Kamiński, J., and Wróbel, A. (2015). Alertness opens the effective flow of sensory information through rat thalamic posterior nucleus. *Eur. J. Neurosci.* **47**, 1321–1331.
- Sosnik, R., Haidarliu, S., and Ahissar, E. (2001). Temporal frequency of whisker movement. I. Representations in brain stem and thalamus. *J. Neurophysiol.* **86**, 339–353.
- Summerfield, C., Greene, M., Wager, T., Egner, T., Hirsch, J., and Mangels, J. (2006). Neocortical connectivity during episodic memory formation. *PLoS Biol.* **4**, e128.
- Taniguchi, H., He, M., Wu, P., Kim, S., Paik, R., Sugino, K., Kvitsiani, D., Fu, Y., Lu, J., Lin, Y., et al. (2011). A resource of Cre driver lines for genetic targeting of GABAergic neurons in cerebral cortex. *Neuron* **71**, 995–1013.
- Urbain, N., and Deschênes, M. (2007). A new thalamic pathway of vibrissal information modulated by the motor cortex. *J. Neurosci.* **27**, 12407–12412.
- Viaene, A.N., Petrof, I., and Sherman, S.M. (2011). Properties of the thalamic projection from the posterior medial nucleus to primary and secondary somatosensory cortices in the mouse. *Proc. Natl. Acad. Sci. USA* **108**, 18156–18161.
- Ward, R.L., Flores, L.C., and Disterhoft, J.F. (2012). Infragranular barrel cortex activity is enhanced with learning. *J. Neurophysiol.* **108**, 1278–1287.
- Wen, J.A., and Barth, A.L. (2011). Input-specific critical periods for experience-dependent plasticity in layer 2/3 pyramidal neurons. *J. Neurosci.* **31**, 4456–4465.
- Williams, L.E., and Holtmaat, A. (2019). Higher-order thalamocortical inputs gate synaptic long-term potentiation via disinhibition. *Neuron* **101**, 91–102.e4.

STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
AAV1.CAG.hChR2(H134R)-mCherry.WPRE.SV40	Addgene	Cat#100054-AAV1
AAV2-hSyn-hChR2(H134R)-EYFP	UNC Vector Core, Deisseroth Lab	
Chemicals, Peptides, and Recombinant Proteins		
Tamoxifen	Tocris	Cat# 6342
Ketoprofen	Sigma-Aldrich	Cat# K1751
QX-314 (N-ethyl bromide)	Tocris	Cat# 2313
AP5	Tocris	Cat# 0106
Picrotoxin	Tocris	Cat# 1128
Experimental Models: Organisms/Strains		
C57Bl6 Mice (Harlan)	Jackson Laboratory	Cat# 000664
Nelf1Cre Mice	MMRC	Cat# 037424-UCD
ETV1creER Mice	Jackson Laboratory	Cat# 012048
Ai32 Mice	Jackson Laboratory	Cat# 024109
Software and Algorithms		
Igor Pro 6.0	Wavemetrics	
Minianalysis	Synaptosoft	
Other		
Custom Injection Cannulas	Plastics One	
IR Beam Break Sensor	Adafruit	Cat#2167
Yun Shield v2.4	Dragino	
Leonardo	Arduino	Cat# A000057
WifiShield 2.4	Df Robot	Cat# TEL0047
Solenoid Valve	The Lee Company	Cat# LHDA1233115H
Capacitive touch sensor AT42QT1010	Adafruit	Cat#1374
White LED (Version)	Prizmatix	UHP-T-LED-White-High-CRI
Bipolar Stimulating Electrode	FHC	Cat# CBCRC75

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Alison L. Barth (albarth@andrew.cmu.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals

All experimental procedures were conducted in accordance with the NIH guidelines and were approved by the Institutional Animal Care and Use Committee at Carnegie Mellon University.

Experiments targeting excitatory neurons were performed on C57Bl6 mice (Harlan). In a small subset of experiments, ChR2 expression was driven transgenically by crossing Nelf1Cre ([Gong et al., 2007](#)) (MMRC Stock No: 037424-UCD) animals or ETV1creER ([Gong et al., 2007](#); [Taniguchi et al., 2011](#)) (Jackson Laboratory Stock No: 013048) animals with Ai32 (Jackson Laboratory Stock No. 024109, ChR2(H134R)EYFP) animals to generate offspring that express ChR2 specifically in VPM and L5 Pyr neurons respectively. ETV1-Cre expression was initiated by injection of 2 mg tamoxifen (100 μ L of 20 mg/ml, Tocris Cat No 6342, <http://www.jax.org/research-and-faculty/resources/cre-repository/tamoxifen>) 8-12 days before behavioral training. We observed broad

expression of ChR2 across the entire extent of L5 in these animals. Experiments were performed on animals of both sexes. Animals were stereotactically injected between postnatal day 12–18 (P12–18), began training at P19–28, and were sacrificed for recording at P20–P30.

METHOD DETAILS

Viral Injections

ChR2 tagged with m-cherry or YFP (300–500 nL; AAV1.CAG.hChR2(H134R)-mCherry.WPRE.SV40, Catalog No. 100054-AAV1, Addgene, Cambridge, MA; AAV2-hSyn-hChR2(H134R)-EYFP, Deisseroth Lab, UNC Vector Core, Chapel Hill, NC) was stereotactically injected into the VPM or POM thalamic nucleus following a small craniotomy (VPM: bregma -1.3 , lateral 1.8 , depth 3.4 , POM: bregma -1.7 , lateral 1.00 , depth 3.25 mm) of isoflurane-anesthetized mice using a Hamilton syringe (Hamilton; Reno, NV), Stoelting infusion pump (Stoelting; Wood Dale, IL, Model #53210), and custom injection cannulas (Plastics One; Phoenix, AZ). Mice were treated once with ketoprofen after injection (5 mg/kg, Sigma-Aldrich; St. Louis, MO) and additional doses were administered as necessary. Mice recovered in their home cage for 7–13 days prior to sensory association training (SAT).

Automated home-cage sensory association training

Animals were singly housed in a 7x12 inches standard mouse cage outfitted with a custom-designed chamber with an infrared beam-break in front of a recessed lickport with a capacitor to detect individual lick events. Animals were maintained on a 12 h light-dark schedule, with lights on at 7 am. The lickport was the sole source of water in the cage, and animals were not otherwise water restricted. Food was provided *ad libitum*. Animals were typically introduced to the training cage at noon and allowed one day to acclimate to the cage. They readily learned to drink at the lickport without intervention or shaping, where ~ 1 – 3 ml of water were dispensed each day. Water was provided on 80% of the beambreak-initiated trials, without any predictive cue on the acclimation day. At noon on the second day, a small nozzle for air delivery (inner diameter $1/16$ in) was inserted into the ceiling of the chamber ~ 4 cm above the average location of the right vibrissa during drinking. Mouse position for the airpuff was not stereotyped, and the number and amplitude of whisker movements evoked were not monitored.

We elected to use a gentle airpuff as the stimulus in our sensory association task for several reasons. First, airpuff stimuli can target multiple whiskers without whisker contact, well-suited for automated home-cage-training. Second, animals can be directly introduced to the training cage without whisker trimming or whisker prostheses that can be difficult to maintain over long training periods. Third, because multiple whiskers can be stimulated in a single trial, the cortical region for analysis encompasses a wide area of S1, facilitating fine-scale analysis in acute brain slices (in comparison to single-whisker stimulation paradigms).

For all trials, including during acclimation, IR beam break triggered a variable delay (200–800 ms) before trial initiation, after which the nozzle delivered a short (500 ms) pulse of compressed air at 4–6 psi (measured by a gas regulator). Water was delivered 500 ms after the airpuff offset (1 s after trial start), and approximately 15 μ l of water was dispensed for each trial. Airpuffs and water were delivered as described for 80% of the beambreak-initiated trials (a random number between 0 and 100 was generated and if it was less than 80, water was delivered). The remaining 20% of trials had no airpuff and no water delivery but otherwise had identical trial structure and incidental auditory cues, including variable pre-trial delay. After trial initiation, a new trial could not be triggered until 1 s after water delivery, and additional IR beam breaks during the trial were ignored. In a subset of experiments, mice received an identical training paradigm but using a visible light-flash as the conditioned stimulus. The light flash was broad spectrum and directed downward into the drinking area from above on the right side of the drinking chamber. In an additional subset of experiments, mice received a pseudo-training paradigm which was identical to the previously described trial structure except that water was delivered on 50% of trials randomly determined irrespective of stimulus delivery. All animals performed either ≤ 25 or ≥ 150 trials, so a small minority of animals that failed to perform 25 trials was excluded from analysis.

Data were analyzed in MATLAB and excel using custom scripts that align measured licks to individual trials and measure the delay from trial start for each lick. The presence of a lick was determined by a change in capacitance at the lickport, surveyed in 100 ms time bins surrounding water delivery. PSTHs of lick behavior were generated relative to trial start (following the initial delay) so that changes in licking behavior could be aligned to CS and US delivery. To quantitate learning-related behavioral changes, lick rates were compared between water or stimulus trials to lick rates for no-water “blank” trials on Day 0 and in the last 8 h of training day 1 or 2. We differentiated between consummatory licking (required for water consumption) and anticipatory licking prior to water delivery (evidence that an association between the sensory stimulus and the prediction of a future water “reward”) (Cohen et al., 2012). Anticipatory lick events (occurring 300 ms preceding water delivery, 700–1000 ms after $t = 0$) were measured throughout training in 4 h bins for water-delivery trials and for blank trials. Lick frequency calculated over 100 ms time bins within this window were then converted into Hz.

To quantitate learning-related behavioral changes, lick rates were compared between water or stimulus trials to lick rates for no-water “blank” trials (100ms bins) on Day 0 and in the last 8 h of training day 1 or 2. We differentiated between consummatory licking (required for water consumption) and anticipatory licking prior to water delivery (evidence that an association between the sensory stimulus and the prediction of a future water “reward”) (Cohen et al., 2012). Anticipatory lick events (3, 100 ms time bins sampled for 300 ms preceding water delivery, 700–1000 ms after $t = 0$) were measured throughout training in 4 h bins for water-delivery trials and for blank trials. Lick frequency calculated over 100 ms time bins within this window were then converted into Hz. Performance was

defined as the difference between lick rate for stimulus and blank trials, which was measured and plotted in 4 h time bins. Regression of electrophysiology and behavior utilized each animal's performance quantified for the last 20% of trials performed. All measurements of behavior were calculated for individual animals and then combined to generate aggregate values.

Slice Preparation and Injection Site Confirmation

Injected mice were sacrificed by brief isoflurane anesthesia and decapitation between 11 am and 3 pm. 350 μ m thick off-coronal slices (One cut, 45° rostro-lateral and 25° rostro-dorsal) designed to preserve columnar connections in the somatosensory cortex were prepared in ice-cold artificial cerebrospinal fluid (ACSF) composed of (in mM): 119 NaCl, 3.5 KCl, 1 NaH₂PO₄, 26.2 NaHCO₃, 11 glucose, 1.3 MgSO₄, and 2.5 CaCl₂ equilibrated with 95%/5% O₂/CO₂. Slices were allowed to recover at room temperature for 45 min in the dark before recording. The injection site was confirmed anatomically using the mCherry-tagged ChR2 fluorescence in cell bodies at the injection site and the characteristic pattern of fluorescent axonal labeling in the barrel cortex, concentrated in L4 and L6 for VPM and L1 and L5a for POM (Meyer et al., 2010). Slices that had fluorescently labeled axons outside of the target layers were discarded. Retrogradely-labeled, ChR2+ neurons were not observed in the somatosensory cortex.

Although ChR2 expression levels could differ between animals, experiments were repeated across many animals, mice were assigned to experimental groups without expression information, and all controllable experimental variables were kept consistent. Additionally, fluorescent ChR2 labeling in the cortex was monitored. A consistent minimum expression threshold for inclusion was applied, and no difference was observed in the mean or range of fluorescent intensity between control and trained animals.

General Electrophysiology

Cortical excitatory neurons were targeted for whole-cell recording in the posteromedial barrel subfield using an Olympus light microscope (BX51WI) using borosilicate glass electrodes resistance 4–8 M Ω . Electrode internal solution for evoked activity experiments, was composed of (in mM): 125 potassium gluconate, 10 HEPES, 2 KCl, 0.5 EGTA, 4 Mg-ATP, and 0.3 Na-GTP, pH 7.25–7.30, 280 mOsm. Internal solution for quantal EPSC experiments was composed of (in mM) 130 cesium gluconate, 10 HEPES, 0.5 EGTA, 8 NaCl, 10 Tetraethylammonium chloride (TEA-Cl), 4 Mg-ATP and 0.4 Na-GTP, pH 7.25–7.30, 280–290 mOsm and typically contained QX-314 (5 mM, lidocaine N-ethyl bromide, Tocris). For some cells trace amounts of AlexaFluor 594 were added to the internal solution to confirm cell targeting.

Electrophysiological data were acquired using a Multiclamp 700A amplifier (Axon Instruments, Foster City, CA) and a National Instruments acquisition interface (National Instruments; Austin, TX). Data were filtered at 3 kHz, digitized at 10 kHz and collected by Igor Pro 6.0 (Wavemetrics, Lake Oswego, Oregon). Cells were allowed to recover from break-in for 5 min before data collection. Presumptive excitatory neurons were targeted for whole-cell or juxtacellular recording based on Pyr soma morphology (or stellate nature morphology in L4), intermediate Ri, (354 ± 23 M Ω), and regular-spiking phenotypes in response to current injections. Input-output firing curves were generated from neurons recorded in a potassium-gluconate based internal solution by injecting increasing amplitudes of a square pulse of current to elicit spikes. To ensure that potential changes in spontaneous synaptic currents did not influence measured intrinsic property values, we also recorded resting membrane potential and input-output curves in the presence of synaptic blockers for a subset of neurons control and SAT24 animals. We observed no difference across recording condition and data were pooled for further analysis. Rs and Ri were monitored for the duration of experiments and cells with Ri below 100 M Ω , Rs greater than 40 M Ω , or where Rs changed by more than 30% over the course of data collection were excluded from further analysis.

Following recording, cells were imaged to determine their laminar location based on depth from pial surface and relevant cytoarchitectural features. L2 neurons were defined as neurons up to 100 μ m below the cell-sparse area of L1, typically 50–150 μ m below the pial surface. L4 neurons are defined as inside the upper and lower limit of the L4 barrel, but were selected from both “barrel” and “septal” regions, since segregated barrel and septal circuits in mouse L4 are unclear (Feldmeyer et al., 2013). L5 neurons were recorded from the area up to 150 μ m below L4 barrels, an area corresponding to L5a.

Evoked cortical activity

ChR2-expressing thalamic axons were stimulated by delivering trains of light pulses (5 pulses, 80ms ISI, 0.05Hz inter-trial interval) through a 40x water-immersion objective (Olympus) at the recording site using a white LED (Prizmatix, Israel) in combination with an excitation filter (40 nm bandwidth centered at 480nm; Chroma; Bellows Falls, VT). Max light intensity at 470 nm was measured at 2.13 mW distributed over a beam area \sim 1 mm diameter, and the timing of optogenetic stimulation was controlled by a Master-8 (A.M.P.I.; Jerusalem, Israel). Responses in excitatory cortical neurons were measured in either the whole cell or juxtacellular configuration in a modified ACSF solution identical to cutting solution but with (in mM) 2.5 KCl, 0.5 MgSO₄, and 1 CaCl₂ (Audette et al., 2018).

Spike data from at least 10 consecutive trials for each cell was binned at 10ms intervals and averaged across all cells of a given population to generate an average PSTH. The average firing rate was calculated for 500ms pre-stim, during stim, and post-stim. A cell was included in the fraction of spiking cells if any action potential(s) was observed in the stimulus or post-stimulus window. Evoked activity experiments were performed for control animals, which had undergone at least one day of cage acclimation but received no sensory stimuli, and for animals that had undergone 24 h of sensory association training.

In a subset of experiments, POM- evoked activity in L2 pyramidal neurons was recorded from 24 h SAT animals before and after making an incision through L4. Incisions of 1–2 mm were manually performed with a custom knife. For each post-incision L2 cell to be

included in analysis, an adjacent pre-cut cell was recorded to insure that the slice had sufficient ChR2 expression to drive cortical activity. Evoked activity was measured as previously describe. To insure that our incision did not fundamentally alter the direct POM input onto L2 Pyr neurons, the onset latency and slope of the earliest identifiable POM-evoked EPSP were measured. The slope of the initial rise was defined as the maximum slope before a second inflection point.

Synaptic analysis

Subthreshold excitatory post-synaptic potentials were calculated for Pom- and VPM-evoked current clamp recordings made from control and SAT24 animals. For each cell, trials that did not contain an action potential in response to the first light pulse were averaged together to create an average of sub-threshold responses. Peak EPSP amplitude within 20 ms of the first light stimulus was then measured for each cell average trace. In many cells, especially after training, optogenetic stimulation of POM inputs drove action potentials on every trial, so PSPs could not be isolated, particularly for cells that had the largest PSPs. Additionally, measured sub-threshold EPSPs likely represent a composite of thalamic and local inputs since optical activation of thalamic inputs can drive rapid action potential generation in some cortical populations that feedback upon the recorded neuron. Peak subthreshold amplitudes of cell average responses in the one second following stimulus train offset (500-1500 ms after stimulus onset) were also measured to assess prolonged depolarization.

In a subset of experiments, thalamically evoked cortical responses were recorded in voltage clamp to assess paired pulse ratio and AMPA:NMDA ratio. For paired pulse experiments, neural responses to trains of light pulses (5 pulses, 80 ms ISI, 0.05 Hz inter-trial interval) were recorded at -70 mV (E_{Cl^-}). Paired pulse ratio was measured as the peak amplitude of the cell average response within 20ms of the second light pulse divided by the peak amplitude of the cell average response within 20 ms of the first light pulse. For AMPAR:NMDAR current ratio experiments, stimulus intensity was reduced to avoid polysynaptic activity that would contaminate accurate measurements of the direct POM response. Because this measurement is a ratio, it is independent of stimulus intensity and could thus be used across different slice preparations. EPSCs to a single light pulse were recorded at -70 mV (AMPA) and $+40$ mV (NMDA) in the presence of picrotoxin to block disynaptic thalamically-evoked inhibition. Light pulse duration and intensity were titrated for each cell to produce a response with minimal polysynaptic activity. The amplitude of the AMPA component was measured as the peak amplitude of the cell average response within 20 ms of the light pulse. The NMDA response was measured as the amplitude of the cell average response 50ms after the light pulse to ensure that the AMPA component of the response had returned to baseline.

Input-specific quantal EPSC measurements

Quantal amplitude measurements were performed in the standard ACSF solution used during cutting but containing $SrCl_2$ instead of $CaCl_2$ and in the presence of the NMDA receptor antagonist AP5 (50 μ M, Tocris Catalog No. 0106, Minneapolis, MN). Well-isolated individual quantal EPSCs were recorded at -70 mV following input-specific optical stimulation (1 pulse, 5 ms, variable intensity) of ChR2-expressing axons (Biane et al., 2016; Wen and Barth, 2011). Quantal events were manually selected but blind to cell identity based on their short rise time, isolated baselines, and absence of multiple inflection points indicative of a compound event. Events occurring between 50 and 500 ms following stimulation (minimum of 25 per cell) were analyzed using MiniAnalysis software (Synaptosoft, Decatur, GA; Detection parameters: Threshold 9 pA; local maximum period 3.5 ms; baseline period 6 ms; decay time period 10 ms; decay time fraction 0.333; baseline average period 4 ms; area threshold 10 pA; peak average points 3) aligned to rise, and averaged to generate an average qEPSC trace for each cell. The average qEPSC trace for each cell in an experimental group was then aligned to rise to generate a global qEPSC amplitude. The amplitude of each event was averaged to determine a cell's average qEPSC amplitude and cumulative distribution histograms were generated from a pool of qEPSCs containing 25 randomly selected events from each cell in an experimental group. Since the amplitude of quantal events can be influenced by Rs, only cells below 25 M Ω were included. Experiments in ETV1Cre mice were performed in the presence of the GABAA receptor antagonist Picrotoxon (50 μ M, Tocris, Cat. No. 1128) due to a small number of inhibitory neurons in the ETV1-Cre expressing population (Lu et al., 2017). L2-L2 and L5-L5 connections were stimulated using a concentric bipolar stimulating electrode (FHC Catalog No. CBCRC75, Bowdoin, ME) placed in L2 or L5 100-400 μ M away from the recorded neuron. For qEPSC experiments, light or electrical stimulus intensity was low and calibrated individually for each cell to evoke an initial multiquantal EPSC between 50 and 150 pA.

QUANTIFICATION AND STATISTICAL ANALYSIS

Unless specifically noted in figure legend, calculations and statistics were performed on cells and all statistical tests are non-parametric Mann-Whitney (unpaired) or Wilcoxon (paired) rank sum tests, and significance values are reported in figure or when references in results. All average values are mean \pm SEM unless indicated. Cell (n) and animal values (N) are reported in each figure. Animals were randomly assigned to experimental group, and quantal amplitude experiments were analyzed blind to cell identity. Potential relationships between trial number, performance (L_w-L_b), and qEPSC amplitude were plotted and evaluated using a Spearman rank-order correlation test. Analysis of qEPSC amplitude versus performance included some animals that did not show an increase in L_w-L_b ; thus, the regression analysis included performance values that were < 0 . Because we did not have lick frequency data for a subset of animals that went through SAT, the number of points differs for the regression of qEPSC amplitude and trial number versus qEPSC value and performance.

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Supplemental Information

**Rapid Plasticity of Higher-Order Thalamocortical
Inputs during Sensory Learning**

Nicholas J. Audette, Sarah M. Bernhard, Ajit Ray, Luke T. Stewart, and Alison L. Barth

Supplemental Information

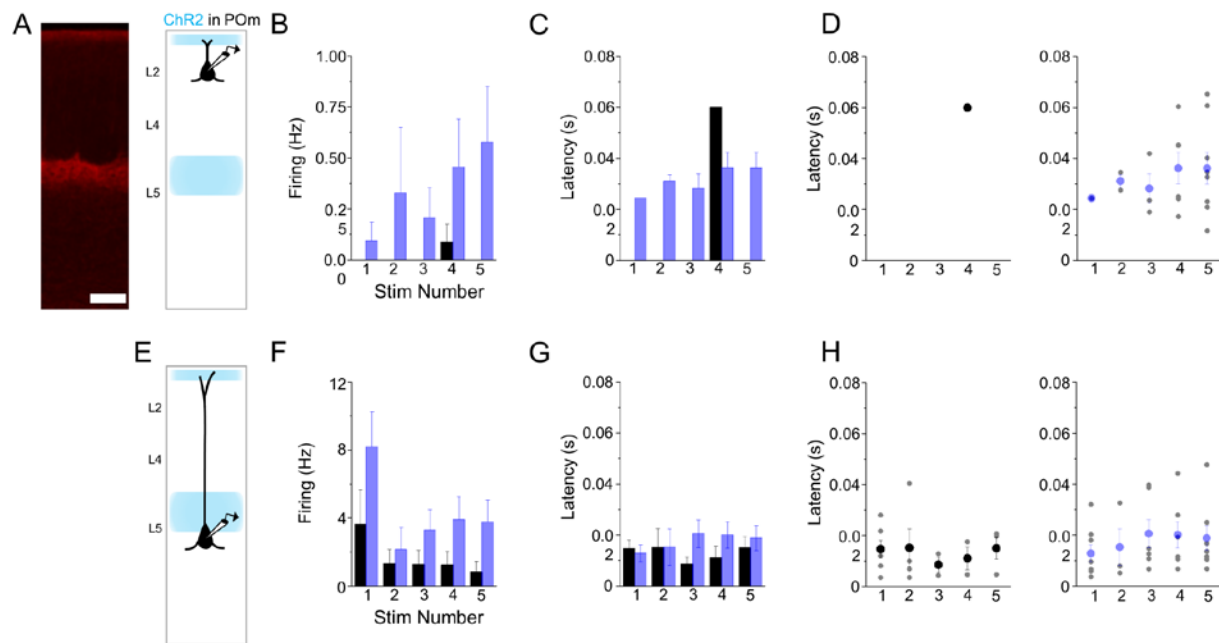


Figure S1. POM-evoked action potential latency changes after SAT, Related to Figure 2.

(A) Fluorescent image of POM ChR2-expressing axons in vS1, (Scale bar 250 μ m) and schematic of experimental setup. (B) Quantification of L2 Pyr spike rate in 80ms following each stim in a train for CTL (black) and SAT24 (blue) animals. (C) Latency of first spike occurring in the 80ms following each stim in a train for CTL (black) and SAT24 (blue) animals. Number of values for each stim is variable since not all cells fired action potentials. (D) First spike latency reported in (C) showing the average for each cell in CTL (black, left) and SAT24 (blue, right) animals. (E-H) Same as (A-D) but for L5 Pyr neurons.

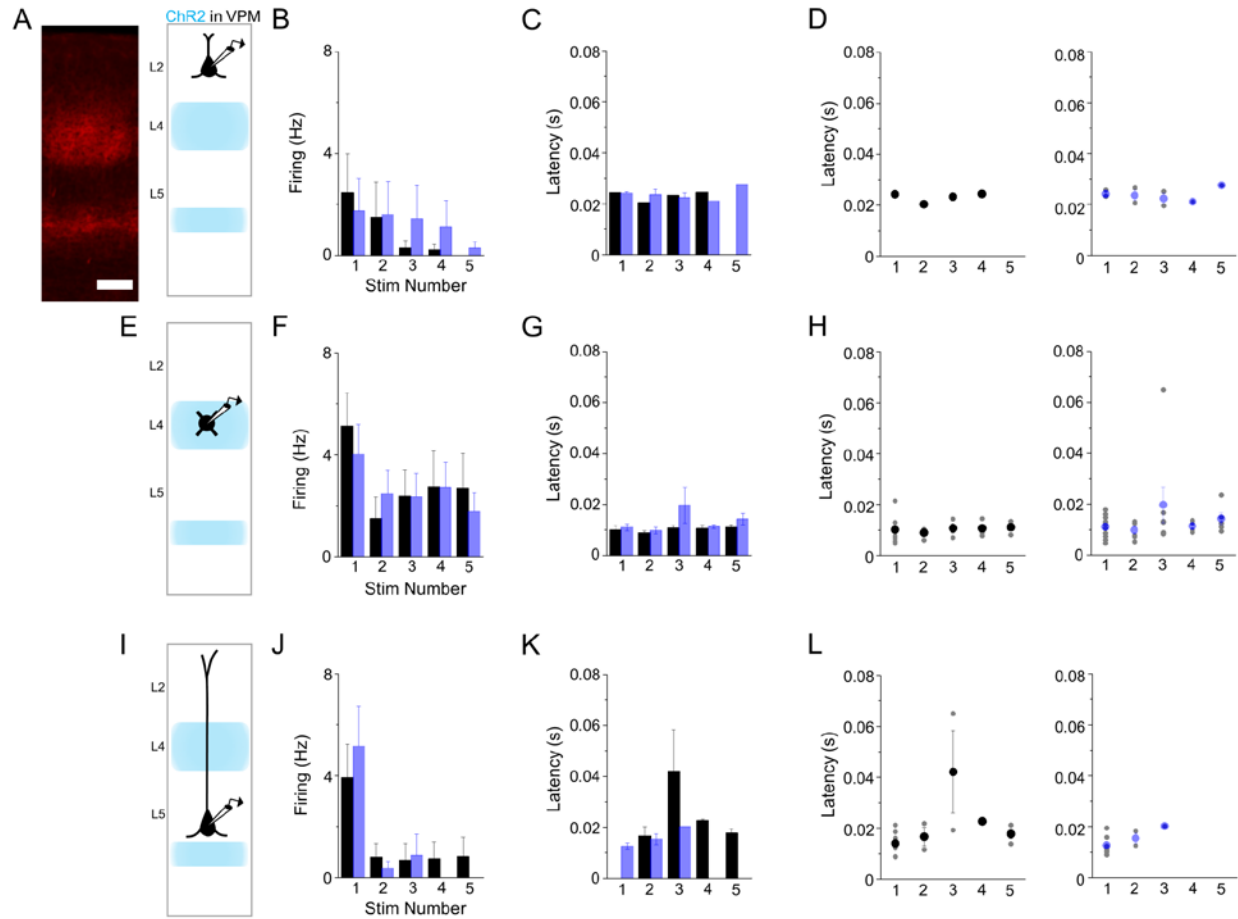


Figure S2. VPM-evoked action potential latency after SAT, Related to Figure 3. (A)

Fluorescent image of VPM ChR2-expressing axons in vS1, (Scale bar 200µm) and schematic of experimental setup. (B) Quantification of L2 Pyr spike rate in 80ms following each stim in a train for CTL (black) and SAT24 (blue) animals. (C) Latency of first spike occurring in the 80ms following each stim in a train for CTL (black) and SAT24 (blue) animals. Number of values for each stim is variable since not all cells fired action potentials. (D) First spike latency reported in (C) showing the average for each cell in CTL (black, left) and SAT24 (blue, right) animals. (E-H) Same as (A-D) but for L4 excitatory neurons. (I-L) Same as (A-D) but for L5 Pyr neurons.

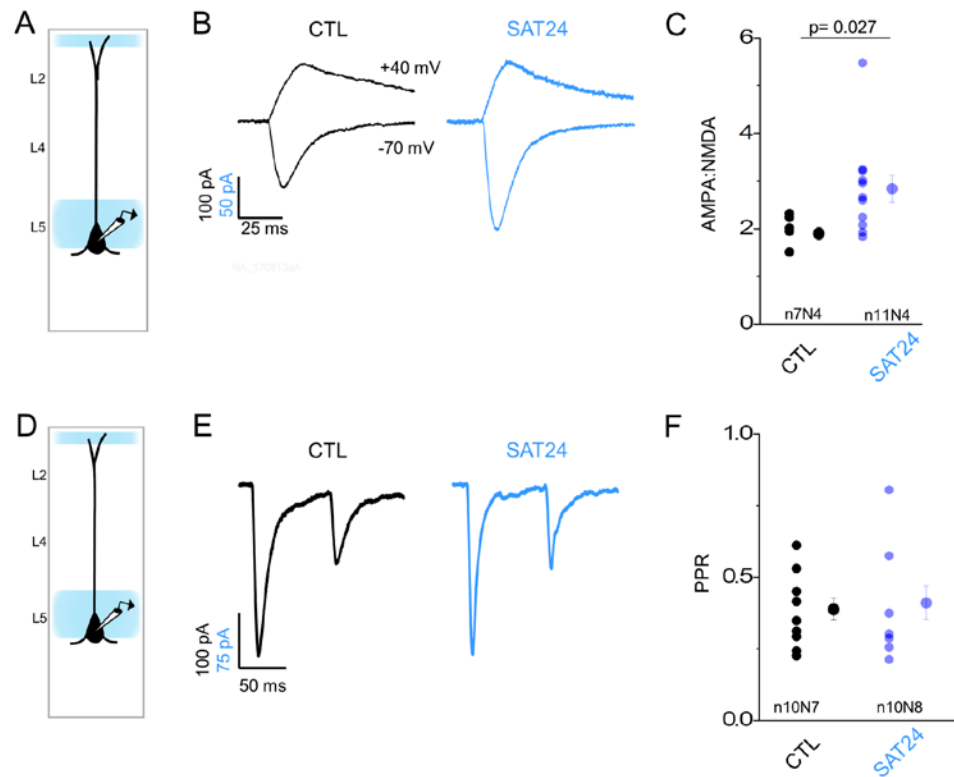


Figure S3. Postsynaptic plasticity at POM inputs following SAT, Related to Figure 5. (A,D)

Schematic of experimental setup for recording POM-evoked post-synaptic currents in L5 Pyr neurons. (B) Example traces (average of 10 consecutive sweeps in a single cell) showing AMPA and NMDA responses to POM optogenetic stimulation recorded in the presence of picrotoxin at -70mV and +40mV respectively for CTL (black) and SAT24 (blue) animals. (C) Quantification of cell average AMPA:NMDA ratio with NMDA amplitude measured 50ms after stimulation for all cells. (E) Example traces (average of 10 consecutive sweeps in a single cell) showing EPSCs evoked by the first two light pulses in a stimulation train (5ms, 80ms ISI) recorded at -70mV for control (black) and SAT24 (blue) animals. (F) Quantification of paired pulse ratio, measured as the average amplitude of EPSC evoked by the second light pulse divided by the average amplitude of EPSC evoked by the first light pulse for each cell.

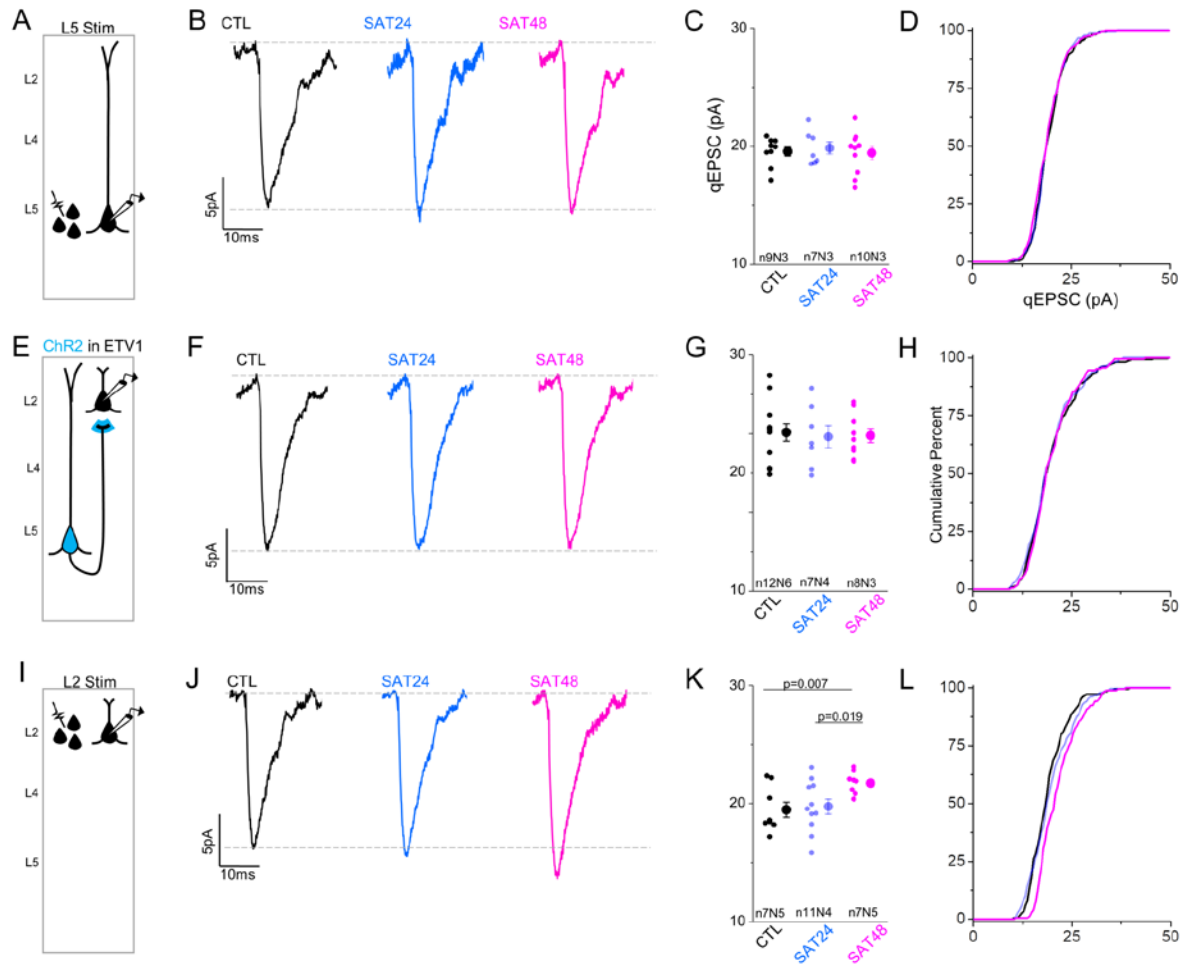


Figure S4. Average quantal amplitude of intracortical excitatory connections is unchanged following SAT, Related to Figure 6. (A) Schematic of experimental setup for assessing L5-L5 connection strength using local placement of an electrical stimulation electrode. (B) Global average qEPSC in CTL (black, left), SAT24 (blue, middle), or SAT48 (magenta, right) animals. All well-isolated light-evoked qEPSCs in a cell (≥ 25 for inclusion) were aligned to rise time and averaged to generate an average cellular qEPSC. Cell averages were aligned to rise and averaged to generate global average qEPSC for each condition. (C) Quantification of mean qEPSC amplitude for each cell, measured as the mean of individual qEPSC peak amplitudes within a cell. (D) Cumulative distribution histogram of qEPSC amplitudes for CTL (black), SAT24 (blue), and SAT48 (magenta) animals. Distributions

comprise 25 randomly selected events from each cell, compared using a K-S test. (E-H) Same as (A-D) but for L5 to L2 connections using transgenic expression of ChR2 in L5 under the control of the ETV1 promotor. (I-L) Same as (A-D) but for L5 to L5 connection strength using local placement of an electrical stimulation electrode. (L) K-S test: Control vs 48 hr $p=0.00045$, 24 hr vs 48 hr $p=0.0016$.

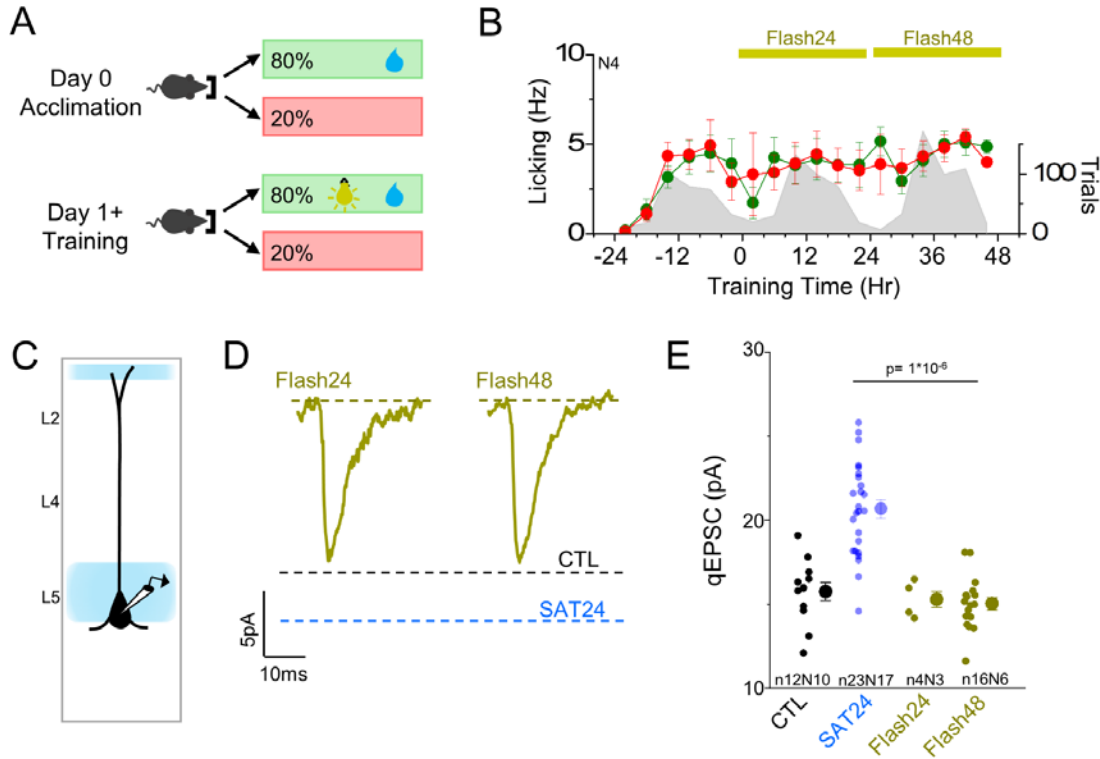


Figure S5. Light flash SAT does not alter POM input strength in barrel cortex, Related to

Figure 7. (A) Schematic of SAT paradigm identical to air puff training but with a light flash substituted as the CS. (B) Time course of anticipatory lick frequency (left axis, 300ms prior to water delivery, 4 hr bins) over the course of acclimation and training for blank (red) and stim/water trials (green) averaged across all animals (N=4 for all timepoints). (C) Schematic of experimental setup for recording POM-evoked qEPSCs in L5 Pyr neurons after light flash training. (D) Global average qEPSC in Flash24 and Flash48 trained animals. All well-isolated light-evoked qEPSCs in a cell (≥ 25 for inclusion) were aligned to rise time and averaged to generate an average cellular qEPSC. Cell averages were aligned to rise and averaged to generate global average qEPSC for each condition. Dotted lines represent the amplitude of global average qEPSCs for control (black) and SAT24 (blue) animals for comparison. (E) Quantification of mean qEPSC amplitude for each cell, measured as the mean of individual qEPSC peak amplitudes within a cell. Statistical comparison shown for SAT24 and Flash48 conditions.

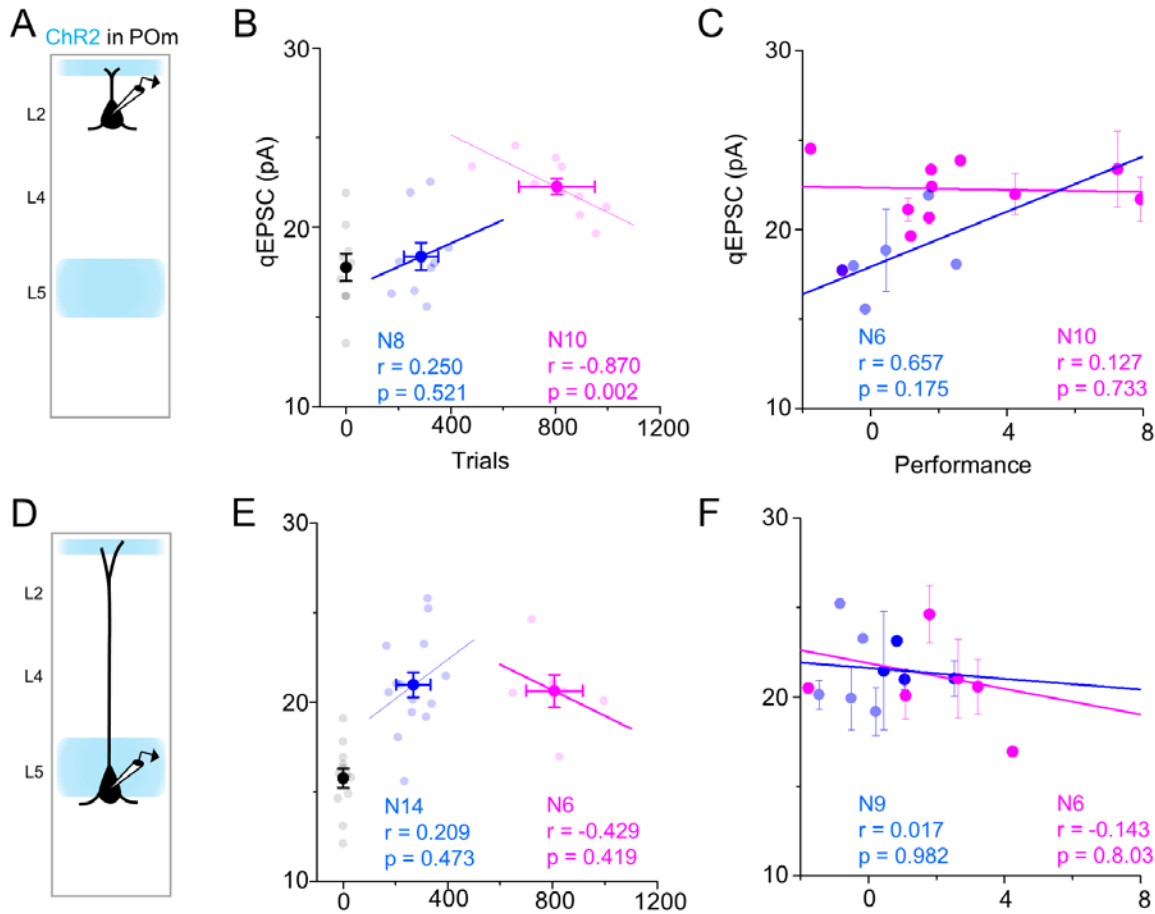


Figure S6. Relationship between synaptic strengthening and animal behavior during training, Related to Figure 8. (A) Schematic of experimental setup for recording POm-evoked qEPSCs in L2 Pyr neurons. (B) Relationship between total number of training trials (stimulus and blank) and average L2 POm-evoked qEPSC for control (gray), SAT24 (blue) and SAT48 (pink) trained animals. Each gray dot represents an animal average of qEPSC measurements. Regression analysis performed using spearman rank-order correlation, rho values and two-sided p value shown for each; N values indicate animals. (C) Relationship between performance on the final 20% of training trials and average L2 POm-evoked qEPSC amplitude in an animal for control (gray), SAT24 (blue) and SAT48 (pink) trained animals. Regression analysis is same as (B). (D-F) Same as (A-C) but for animal average L5 POm-evoked qEPSCs.