Cross-Modal Reinstatement of Thalamocortical Plasticity Accelerates Ocular Dominance Plasticity in Adult Mice

Highlights

- Thalamocortical (TC) plasticity is limited in adults
- A short duration of deafening specifically recovers TC-LTP in adult V1
- Deafening increases NMDAR function and E/I balance at TC synapses
- Deafening accelerates ocular dominance plasticity in adult V1

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In Brief

Plasticity of thalamocortical (TC) synapses is limited in adults. Rodríguez et al. demonstrate that a brief period of deafening adults recovers LTP at TC synapses in visual cortex and accelerates ocular dominance plasticity. These results suggest that cross-modal sensory deprivation may be an effective way to promote adult cortical plasticity.
Cross-Modal Reinstatement of Thalamocortical Plasticity Accelerates Ocular Dominance Plasticity in Adult Mice

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SUMMARY

Plasticity of thalamocortical (TC) synapses is robust during early development and becomes limited in the adult brain. We previously reported that a short duration of deafening strengthens TC synapses in the primary visual cortex (V1) of adult mice. Here, we demonstrate that deafening restores NMDA receptor (NMDAR)-dependent long-term potentiation (LTP) of TC synapses onto principal neurons in V1 layer 4 (L4), which is accompanied by an increase in NMDAR function. In contrast, deafening did not recover long-term depression (LTD) at TC synapses. Potentiation of TC synapses by deafening is absent in parvalbumin-positive (PV+) interneurons, resulting in an increase in feedforward excitation to inhibition (E/I) ratio. Furthermore, we found that a brief duration of deafening adult mice recovers rapid ocular dominance plasticity (ODP) mainly by accelerating potentiation of the open-eye responses. Our results suggest that cross-modal sensory deprivation promotes adult cortical plasticity by specifically recovering TC-LTP and increasing the E/I ratio.

INTRODUCTION

Thalamocortical (TC) inputs convey sensory information to the cortex for further processing and are shaped by sensory experience during a defined early developmental period (Barkat et al., 2011; Crair and Malenka, 1995). Notably, across different sensory cortices, TC synapses in layer 4 (L4) exhibit the earliest and shortest critical period that precedes synaptic plasticity in other layers, which often persists through adulthood (Barkat et al., 2011; Barth and Malenka, 2001; Crair and Malenka, 1995; Desai et al., 2002; Goel and Lee, 2007; Jiang et al., 2007). Therefore, the loss of TC synaptic plasticity may be an essential factor contributing to the limited ability of the adult brain to undergo plasticity. In the mouse visual pathway, TC synapses undergo experience-dependent reorganization and refinement that end between the second and third postnatal week (Gu and Cang, 2016; Jiang et al., 2007). Ascending TC inputs onto L4 neurons express NMDA receptor (NMDAR)-dependent long-term potentiation (LTP) and long-term depression (LTD) during this limited time window, while L4 to L2/3 synapses remain plastic into adulthood in rodent primary visual cortex (V1) (Jiang et al., 2007). The same laminar progression of plasticity has been described in studies of experience-dependent homeostatic synaptic scaling in V1, where L4 plasticity ends earlier while that in L2/3 persists (Desai et al., 2002; Goel and Lee, 2007). Studies performed in vivo also showed that ocular dominance plasticity (ODP) is more robust in L4 early in development, while in adults, it is more evident in L2/3 (Pham et al., 2004). Taken together, these cases support the view that TC synapses onto V1 L4 undergo plasticity early on, and subsequent experience-dependent changes are mainly mediated by plasticity in the superficial layers.

Recent studies have challenged this notion by demonstrating TC plasticity in adults under particular manipulations, such as environmental enrichment (Mainardi et al., 2010), prolonged visual deprivation (Montey and Quinlan, 2011), or peripheral nerve transection (Chung et al., 2017; Yu et al., 2012). Furthermore, we previously reported that visual deprivation results in strengthening of TC synapses to auditory cortex (A1) while deafening leads to TC potentiation in V1 (Petrus et al., 2014). This indicates that cross-modal sensory deprivation in adults readily reactivates TC plasticity, which could represent the cellular basis for cross-modal plasticity that allows enhanced processing in spared cortices after sensory loss (Lee and Whitt, 2015). However, the synaptic mechanisms underlying this change remain unexplored. Here, we used targeted optogenetic activation of TC synapses to demonstrate the reemergence of NMDAR-dependent LTP at V1 TC synapses following a week of deafening adult mice (post-natal day 90 [P90] to P120), which occurred without recovery of TC-LTD. In addition, we found that deafening...
increases the excitation to inhibition (E/I) ratio of TC inputs onto L4 principal neurons. Such a shift in the E/I ratio has been linked to heightened cortical plasticity (Froemke, 2015; Froemke et al., 2007). Consistent with this notion, we report that restoration of V1 TC plasticity by cross-modal sensory deprivation accelerates ODP in adults. Our findings suggest cross-modal sensory manipulations as a potential method to facilitate plasticity in the adult brain.

RESULTS

Reemergence of LTP at TC Synapses in V1 L4 after Deafening

We previously showed that a brief period of sensory deprivation produces cross-modal strengthening of TC synapses in L4 of the spared sensory cortices of adult mice (Petrus et al., 2014). Importantly, this change in TC synapses was dependent on the spared cortex retaining its own sensory inputs. Hebbian forms of synaptic plasticity have been established as the mechanisms driving experience-dependent changes of TC inputs to L4 in primary sensory cortices during the critical period (Crair and Malenka, 1995; Dudek and Friedlander, 1996; Kirkwood et al., 1995).

Therefore, we tested whether the cross-modal potentiation of TC synapses by a period of deafening is due to reengagement of LTP mechanisms in adult V1. We induced hearing loss in adult animals (P90–P120) via ototoxic lesioning of hair cells by injection of kanamycin (175 mg/mL) into the inner ear coupled with tympanic rupture. We confirmed the efficiency of our approach by measuring auditory brainstem responses (ABRs) in a subset of mice, as well as by verifying cochlear damage in the deafened group by phalloidin staining of hair cells (Figure S1).

In order to selectively activate TC synapses, we expressed ChR2 in the visual thalamus (dLGN) via adeno-associated virus (AAV)-mediated transfection as in our published study (Petrus et al., 2014). We found that ChR2 can only follow stimulation faithfully up to ~40 Hz (Figure S2). Therefore, we induced LTP at TC synapses using a pairing protocol, which consists of pairing low-frequency activation (1 Hz, 200 pulses) of ChR2-expressing TC synapses using light emitting diode (LED) (455 nm) with postsynaptic depolarization (0 mV) of a L4 principal neuron (Figure 1A). This is a modification of a protocol used successfully in cortical slices from young (P8–P17) mice to study TC-LTP using electrical stimulation of the white matter (WM) (Jiang et al., 2007). The location of recorded neurons and expression of ChR2 were verified post hoc (Figure 1B). Consistent with previous studies showing
A week of prior deafening did not restore TC-LTD in adults (Figure 1D). Collectively, our results indicate that a brief cross-modal sensory deprivation specifically restores TC-LTP in adults. To do this, we repeated the same protocol to induce TC-LTP in V1 slices of normal-reared (NR) adults (Figure 1C). In contrast, V1 slices from adult mice that were deafened early critical period for WM to L4 LTP, our pairing protocol failed to induce TC-LTP in V1 slices of normal-reared (NR) adults (Figure 1C). In contrast, V1 slices from adult mice that were deafened (DF) a week (7 ± 1 days) prior to the experiment exhibited robust TC-LTP, which was blocked by bath application of 100 μM DL-2-aminoo-5 phosphonopentanoic acid (APV) (Figure 1C). These results indicate that TC-LTP recovered in adults by cross-modal sensory deprivation is dependent on NMDARs, which is similar to what has been described for TC-LTP in normal young mice (Crair and Malenka, 1999).

Next, we examined whether deafening will also restore LTD at TC synapses in adults. To do this, we repeated the same experimental design, except that the postsynaptic cells were held at −40 mV during pairing protocol (Figure 1A). We did not observe TC-LTD in normal adults, which is consistent with a loss of this form of LTD beyond the third postnatal week (Jiang et al., 2007), and a week of prior deafening did not restore TC-LTD in adults (Figure 1D). Collectively, our results indicate that a brief cross-modal sensory deprivation specifically restores TC-LTP in adults.

Cross-Modal Regulation of NMDARs at TC Synapses

Recovery of NMDAR-dependent TC-LTP in V1 of DF adult mice suggests that deafening may alter NMDAR function. To examine this possibility, we first determined the relative contribution of NMDARs to excitatory postsynaptic currents (EPSCs) evoked from thalamic terminals by measuring the NMDA/AMPA ratio. There was no significant change in the average NMDA/AMPA ratio between the NR and DF groups (Figure 2A). We previously reported that the same duration of DF increases AMPA receptor (AMPA)-mediated currents at TC synapses (Petrus et al., 2014), which suggests that there is concurrent potentiation of NMDAR-EPSCs to maintain the NMDA/AMPA ratio at TC synapses following DF.

A switch in NMDAR GluN2 subunit composition is known to coincide with decreased synaptic plasticity and the closure of the critical period for cortical plasticity (Barth and Malenka, 2001; Philpot et al., 2001; Quinlan et al., 1999). GluN2B-containing NMDARs predominate at synapses early on but undergo a developmental switch to incorporate GluN2A subunits, which have faster decay kinetics (Monyer et al., 1994; Sheng et al., 1994). Furthermore, changes in GluN2 subunit composition have been shown to alter the threshold for LTD induction (Philpot et al., 2001; 2003; Quinlan et al., 1999). In order to determine if changes in NMDAR subunit composition allowed for the recovery of TC-LTP with DF, we measured isolated NMDAR-evoked responses at TC synapses onto L4 in V1 and compared their weighted decay time constants between groups as reported previously (Philpot et al., 2001; Rumbaugh and Vicini, 1999). We observed no significant changes in the NMDAR decay kinetics at TC synapses between the NR and DF groups (Figure 2B). Our results suggest that cross-modal sensory deprivation in adults potentiates the function of synaptic NMDARs without detectable changes in NR2 subunit composition, which contrasts what is observed early in development.

Deafening Increases the E/I Ratio of TC Inputs to V1 L4 Principal Neurons

Thalamic inputs strongly drive feedforward inhibition onto cortical L4 neurons, which serves to regulate coincidence detection by improving the precision of postsynaptic spikes (Chittajallu and Isaac, 2010; Cruikshank et al., 2007; Daw et al., 2007; Gabernet et al., 2005; Kloc and Maffei, 2014). In V1, like other sensory cortices, TC inputs recruit feedforward inhibition mainly by activating parvalbumin-positive (PV+) interneurons (Cruikshank et al., 2007; Daw et al., 2007; Kloc and Maffei, 2014). Maturation of cortical inhibition mediated by PV+ neurons is also implicated in gating cortical plasticity (Jiang et al., 2005). Therefore, we examined if DF could lead to changes in thalamic engagement of cortical inhibition.

We targeted recordings to PV+ cells in L4 V1 by using mice expressing tdTomato in these neurons (PV-tdTomato) and expressed ChR2 in dLGN using targeted injection of AAV (Figures 3A and 3B). To compare the strength of individual TC synapses regardless of ChR2 expression or stimulation level, we recorded LED-evoked Sr2+ desynchronized (Lev-Sr2+) mEPSCs as described previously (Petrus et al., 2014, 2015) (see STAR Methods). In this scheme, LED-evoked responses are desynchronized such that recorded events represent single-vesicle release and thus, quantal synaptic response size can be determined (Gil et al., 1999). We found no significant differences between the average amplitude of Lev-Sr2+ miniature EPSCs (mEPSCs) recorded
from PV+ neurons in the NR and DF groups (Figure 3C). This result contrasts our previous finding that TC synapses on L4 principal neurons potentiate with DF (Petrus et al., 2014), which suggests that a postsynaptic target determines the plasticity of TC synapses with cross-modal deprivation. To test whether the selective potentiation of TC inputs to principal neurons alters the balance of recruiting feedforward excitation and inhibition, we measured the E/I ratio in V1 L4 principal neurons following TC input activation using ChR2 while holding the postsynaptic cell at reversal potential for glutamatergic (E$_\text{GABA}$) or GABAergic (E$_\text{GABA}$) synaptic transmission (Figure S3). We observed a significant increase in the average E/I ratio values after DF (open circles represent each cell; NR, 4 mice; DF, 4 mice; t test, *p = 0.0127).

Bar graphs: mean ± SEM.

Deafening Restores Rapid ODP in Adult V1
Based on our result that short-term deafening of adult mice restores TC-LTP in V1, we next tested whether deafening could restore experience-dependent V1 plasticity. ODP is a classic approach to studying experience-dependent synaptic plasticity in V1, and the mechanisms underlying it in both juveniles and adults have been well characterized (Frenkel and Bear, 2004; Gordon and Stryker, 1996; Sato and Stryker, 2008; Sawtell et al., 2003). It is well established that a short period (3–4 days) of monocular deprivation (MD) elicits quick and robust ocular dominance (OD) shift in V1 during the critical period, which in mice starts at P19 and extends until P35 (Gordon and Stryker, 1996) and mainly involves weakening of the closed-eye inputs (Sato and Stryker, 2008; Sawtell et al., 2003). In adults (P60–P90), however, ODP requires a longer period of MD (5–7 days) and is driven predominantly by a delayed potentiation of the open-eye inputs (Frenkel and Bear, 2004; Sato and Stryker, 2008; Sawtell et al., 2003). Since deafening promotes TC-LTP in adult V1, we reasoned that it may accelerate the open-eye potentiation in adults to allow ODP with a shorter duration of MD, which is normally ineffective.

We measured ODP using optical imaging of V1 intrinsic signals before and after periods of MD (Figures 4A and 4B) and calculated an ocular dominance index value (ODI) at each time point (Figure 4C). First, we verified that brief (3–4 days) MD shifts ODI in young mice within the established critical period (P25–P32) (Figure 4A), which in our hands was due to a combination of weakening of the closed contralateral eye inputs and potentiation of the open ipsilateral eye inputs (Figures 4D and 4E).
adult mice (P90–P110), that were deafened 1 week prior to initiating the brief (3–4 days) MD, we observed a significant shift in the ODI which was driven exclusively by potentiation of the open ipsilateral inputs (Figures 4D and 4E). Deafening by itself did not alter the ODI (Figure S4). Consistent with previous studies, the same brief (3–4 days) duration of MD failed to alter the ODI in normal hearing adults, and consequently there was no change in the strength of either the closed or open eye inputs to V1 (Figures 4D and 4E). Normal adults required a longer duration of MD (5–6 days) to produce a significant shift in the ODI,
which was driven by potentiation of the open ipsilateral eye inputs (Figures 4D and 4E) as reported in prior studies (Frenkel and Bear, 2004; Ranson et al., 2012; Sato and Stryker, 2008; Sawtell et al., 2003). Our data indicate that deafening accelerates the emergence of open-eye potentiation in the adults but does not engage the fast depression of the closed-eye inputs, as seen in juveniles.

DISCUSSION

In this study, we investigated the molecular mechanisms and functional consequences underlying the reactivation of TC plasticity by cross-modal sensory deprivation. In particular, we addressed the impact deafening has on TC plasticity in the spared V1. We observed a reemergence of NMDAR-dependent TC-LTP, which is driven by potentiation of NMDAR currents and an increased E/I ratio. However, deafening did not restore TC-LTD in adults, which is consistent with the expectation that increase in NMDAR function and E/I ratio will promote LTP rather than LTD by providing greater summation of responses and Ca$^{2+}$ influx. Moreover, we demonstrated that deafening accelerates ODP in adult mice through potentiation of open-eye inputs without any depression of the deprived eye inputs. Notably, the mechanisms described here allow for TC plasticity 10–12 weeks after the defined critical period for these synapses, suggesting that cross-modal sensory deprivation may be an effective means to recover cortical plasticity in adults that are specifically dependent on synaptic potentiation.

Recent studies have begun to highlight the potential for TC synapses to undergo plasticity in adulthood under certain conditions (Chung et al., 2017; Mainardi et al., 2010; Montey and Quinlan, 2011; Petrus et al., 2014; Yu et al., 2012). While our findings are similar to those describing TC-LTP in the spared whisker barrel of adult mice following infraorbital nerve (ION) lesions (Chung et al., 2017; Yu et al., 2012), we extend this by showing that cross-modal sensory deprivation, not just within-sensory modality deprivation, can recover TC-LTP (Figure 1). Furthermore, we show that cross-modal sensory deprivation specifically recovers TC-LTP, but not TC-LTD, in adults. In terms of molecular mechanisms, we found a requirement for NMDARs in adult TC-LTP induction and a potential potentiation of NMDAR function as a result of cross-modal sensory deprivation as deduced by a preserved NMDA/AMPA ratio despite previously reported potentiation of AMPAR current (Petrus et al., 2014) (Figure 2A). However, we did not find observable changes in the kinetics of NMDAR current as would occur if GluN2 subunit composition was altered (Figure 2B). This differs from the findings reported in the ION lesion model, which showed increased ifenprodil sensitivity at spared TC synapses indicative of a switch in NMDAR composition (Chung et al., 2017). Our results suggest that cross-modal NMDAR regulation is likely mediated by a change in the number of functional NMDARs. This is in line with a previous study describing activity-dependent concomitant regulation of AMPAR and NMDAR synaptic components, which is driven mainly by modulation in the number of open NMDAR channels (Watt et al., 2000).

We also found that while TC synapses to L4 principal neurons strengthen (Petrus et al., 2014), those to PV+ inhibitory cells remain unaltered after DF (Figure 3), which suggests that cross-modal reactivation of TC plasticity is specified by the postsynaptic cell type. One functional consequence is an increase in the balance of feedforward excitation and inhibition recruited at TC synapses on V1 L4 principal neurons. Increased E/I ratio is reminiscent of early developmental stages with high potential for plasticity, which becomes restricted as inhibition matures to decrease the E/I ratio (Zhang et al., 2011). Several studies suggested that E/I balance is relatively maintained in the mature cortex (Dehghani et al., 2016; House et al., 2011; Xue et al., 2014), yet our results show that DF causes a deviation from this equilibrium. Changes in E/I balance have been implicated in recruitment of TC plasticity after environmental enrichment (Mainardi et al., 2010) and for learning-induced cortical rearrangement (Fromemke, 2015; Fromemke et al., 2007). Our result similarly suggests that the E/I ratio is not fixed in adults but can be regulated based on sensory experience to reopen the window for plasticity to sculpt neural circuits in accordance with environmental changes and the need of the cortical network for sensory processing.

While the consequences of cross-modal plasticity on V1 function are still unclear, we demonstrate that a short-term deafening can accelerate ODP in adults. This effect was driven selectively by accelerated potentiation of open-eye inputs without major changes in the deprived-eye inputs. This is qualitatively different from other means of recovering ODP in the adult brain such as dark exposure, which has been shown to recover juvenile-like plasticity that involves weakening of the deprived eye inputs (He et al., 2006, 2007) and a change in NMDAR subunit composition to juvenile form (He et al., 2006). We found that DF specifically recovers TC-LTP, but not TC-LTD, by increasing NMDAR function without significant alterations in subunit composition. In addition, deafening increased the E/I ratio at TC synapses on L4 principal neurons. These cellular level changes correlate with acceleration of open-eye potentiation mechanisms and ODP in the adult V1. In sum, our results suggest that cross-modal sensory deprivation may be an effective way to promote experience-dependent cortical plasticity in adults that are specifically dependent on TC-LTP.  

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 includes four figures and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.08.072.

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AUTHOR CONTRIBUTIONS

Conceptualization, G.R. and H.-K.L.; Investigation and Formal Analysis, G.R. (electrophysiological recordings, optical intrinsic signal imaging), D.C. (LTP experiments), R.S. and I.U. (cochlear staining and imaging), K.M.S. and A.M.L. (ABR recording), and H.-K.L. (final analysis), Writing – Original Draft, G.R., A.M.L., and H.-K.L.; Visualization, G.R. and H.-K.L.; Supervision, H.-K.L.; Project Administration, H.-K.L.; Funding Acquisition, H.-K.L. and A.M.L.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Alexa Fluor 488-phalloidin conjugate | Molecular Probes | Cat# A-12379; RRID:AB_2315147 |
| Bacterial and Virus Strains |        |            |
| AAV5.hSyn.ChR2(H134R)-YFP.WPRE.hGH | Penn Vector Core | Addgene26973P |
| Chemicals, Peptides, and Recombinant Proteins |        |            |
| kanamycin           | Sigma-Aldrich | 60615; CAS: 70560-51-9 |
| DAPI                | Life technologies | D-1306 |
| Avidin-Texas red conjugate | Life technologies | A-820 |
| chlorprothixene     | Sigma-Aldrich | C1671; CAS: 6469-93-8 |
| formalin            | Sigma-Aldrich | HT5014; MDL: MFCD00003274 |
| biocytin            | Sigma-Aldrich | B4261 |
| Isoflurane          | Patterson Veterinary | 07-890-8115 |
| Tetrodotoxin citrate (TTX) | Abcam | ab120055 |
| DL-APV              | Sigma-Aldrich | A5282 |
| NBOX disodium salt hydrate | Sigma-Aldrich | N183 |
| glycine             | Fisher | BP381-1 |
| xylazine            | Patterson Veterinary | 07-808-1947 |
| ketamine            | Patterson Veterinary | 07-890-8598 |
| Bicuculline methiodide | Enzo | BML-EA149-0050 |
| picrotoxin          | Tocris | Cat#1128 |
| Deposited Data      |        |            |
| Dataset for each figure | Mendeley | https://doi.org/10.17632/drgg7v5zjx.1 |
| Experimental Models: Organisms/Strains |        |            |
| C57BL/6J           | Jackson Laboratories | RRID:IMSR_JAX:000664 |
| B6.129P2-Pvalbtm1(cre)Arbr/J | Jackson Laboratories | RRID:IMSR_JAX:017320 |
| B6.Cg-Gt(ROSA)26Sortm14(CAG-tdTomato)Hze/J | Jackson Laboratories | RRID:IMSR_JAX:007914 |
| Software and Algorithms |        |            |
| MATLAB              | Mathworks | https://www.mathworks.com/products/matlab.html; RRID:SCR_001622 |
| Prism 7.0           | GraphPad Software | https://www.graphpad.com/scientificsoftware/prism/; RRID:SCR_002798 |
| Igor Pro            | Wavemetrics | https://www.wavemetrics.com/products/igorpro/igorpro.htm; RRID:SCR_000325 |
| Mini-Analysis       | Synaptosoft | http://www.synaptosoft.com/MiniAnalysis/; RRID:SCR_002184 |
| Other               |        |            |
| gel foam            | Pfizer | 0315-08 |
| CCD camera          | Dalsa | DS-1A-01M30-12E |
| Axon patch-clamp amplifier | Molecular Device | Multiclamp 700B |
| processor           | Tucker-Davis Technologies | RX6 Multifunction Processor |
| attenuator          | Tucker-Davis Technologies | PA5 Programmable Attenuator |
| monitor             | ViewSonic | VX2268WM |
| 455 nm LED          | Thor Labs | M455L3-C5 |
| Vibratory tissue slicer | Ted Pella | PELCO easiSlicer product#: 11000 |
| ProLong Gold antifade | Life Technologies | P36930 |
Further information and requests for resources and reagents should be directed to the Lead Contact, Hey-Kyoungh Lee (heykyounghlee@jhu.edu).

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

B6 (C57BL/6J, Jackson Laboratories; RRID: IMSR_JAX:000664) and PV-tdT (offsprings of PV-Cre x Ai14; PV-Cre: Pvalb<sup>tm1(cre)Arbr</sup>/J, RRID:IMSR_JAX:017320; Ai14: Gt(Rosa)26Sor<sup>tm14(CAG-tdTomato)Hze</sup>/J, RRID:IMSR_JAX:007914) mice were reared in a 12 hours light/dark cycle with water and food pellets ad libitum. All experiments were performed on adult mice (P90-P120) of both sexes where littermates were randomly assigned to experimental groups. All protocols were approved by the Institutional Animal Care and Use Committee (IACUC) at Johns Hopkins University and followed the guidelines established by the Animal Care Act and National Institutes of Health (NIH).

**METHOD DETAILS**

**Thalamic Viral Transfection**

Mice (P40-60) of either sex were anesthetized and head fixed in a stereotaxic device (Kopf Instruments, California) under 1.5%–2% isoflurane/oxygen mix. The dorsal lateral geniculate nuclei (dLGN; −2.3 mm lateral: 2 mm from Bregma; 2.42 mm depth from pia) were bilaterally injected with an adeno-associated virus expressing channelrhodopsin (ChR2) under the control of the human synapsin (hSyn) promoter (AAV5.hSyn.ChR2(H134R)-YFP.WPRE.hGH, Penn Vector Core, University of Pennsylvania). Mice recovered on a heated pad and were returned to the animal colony, housed with 2-3 same sex mice until experimental procedures.

**Deafening**

At P90-110, one week (± 1 day) prior to experiments, male and female mice were deafened as in previous studies (Petrus et al., 2014; Petrus et al., 2015). In brief, animals were anesthetized by exposure to isoflurane vapors in an induction chamber. Following the absence of the toe pinch reflex, the animal was placed in a stereotaxic device (Kopf Instruments, California) under constant administration of 2% isoflurane/oxygen mix. The pinnae were cut and the ventral surface of the ear was slit to aid visualization of the inner ear. The tympanic membrane was punctured with a 30-gauge needle, the ear cavity was injected with 50 μL kanamycin (175 mg/mL), and stuffed with kanamycin soaked gel foam (Pfizer) (Hashimoto et al., 2007). The ventral incision and remainder of pinnae were sutured shut (PSDII; Ethicon) and animals were allowed to recover on a heating pad until movement and drinking behaviors were evident.

**Confirmation of deafening**

Auditory brainstem responses (ABRs) were measured for a subset of animals as described previously (McGuire et al., 2015). Mice were anesthetized with ketamine/xylazine (i.p., 100 mg/kg and 20 mg/kg respectively), then placed on a heating pad inside a sound-attenuating chamber (IAC) lined with Sonex Acoustic foam. Mice were placed facing a speaker (FT28D; Fostex, Tokyo, Japan) positioned 30 cm from the pinnae. Temperature was monitored via a rectal probe and maintained at 36°C ± 1°C. Subcutaneous platinum needle electrodes were placed over the left bulla and at the vertex of the skull, and a ground electrode was inserted into the leg muscle. The electrodes were attached to a preamplifier and amplifier (ISO-80; World Precision Instruments, Sarasota, FL). Stimulus generation, presentation and response acquisition were controlled using custom MATLAB-based software (Mathworks; RRID:SCR_001622), a Tucker Davis processor (RX6; Tucker-Davis Technologies, Alachua, FL) and programmable attenuator (PAS). Stimuli consisting of clicks and 5-ms tones (0.5 ms onset/offset) at frequencies of 8, 12, 16, 24 and 20 kHz were presented. Stimuli were generated with a sampling frequency of 195 kHz, and presented at a rate of 20/s. Responses were sampled at 9.5 kHz, bandpass filtered from 300–3000 kHz, and averaged over 300 stimulus repetitions. Clicks were tested first to verify electrode placement and the presence of a clearly observable response, and then tones were tested in random order of frequency. We presented a tone of a given frequency at a sound level of 85–105 dB SPL (depending on frequency), and then continued presenting the same tone at lower sound levels until a threshold was reached. Threshold was defined as the sound level at which the peak-to-peak (any peak) amplitude of the response was two standard deviations above the average baseline noise amplitude during a time window when no sound stimulus was present. Testing lasted approximately 40-60 minutes, and mice were returned to their home cages following testing and monitored until recovery.

We also confirmed the effectiveness of deafening by observation of hair cell loss using phalloidin staining. Whole cochleae were dissected from experimental animals and stored in 10% formalin solution (Sigma) at 4°C. Sectioning and staining of cochleae was done in a blinded manner. Before staining cochleae were washed in 0.1 M PB and decalcified in 3% EDTA for 48 hours. Apical, middle and basal turns were dissected. Each turn was permeabilized with 0.2% Triton-X for 1 hour and subsequently incubated with Alexa Fluor 488-phalloidin (Invitrogen; 1:200) and DAPI (1:5000) for 2 hours. Cochlear turns were whole mounted on glass slides using ProLong Gold antifade mounting medium (Life Technologies). Confocal images of NR and DF cochleae were taken using the same settings.
Cortical slice preparation

Each mouse was deeply anesthetized using isoflurane vapor until absence of corneal reflex and toe pinch response, then transcardially perfused with ice cold artificial cerebrospinal fluid (ACSF, in mM: 124 NaCl, 5 KCl, 1.25 NaH2PO4, H2O, 26 NaHCO3, 10 dextrose, 2.5 CaCl2. 1.5 MgCl2, bubbled with 95% O2/5% CO2) and immediately decapitated. The brain was removed and immersed in ice-cold dissection buffer (in mM: 212.7 sucrose, 10 dextrose, 3 MgCl2, 1 CaCl2, 2.6 KCl, 1.23 NaH2PO4, H2O, 26 NaHCO3, bubbled with 95% O2/5% CO2). Blocks containing V1 were isolated and sectioned coronally into 300-μm thick slices using a vibratome (PELCO easiSlicer, Ted Pella). Slices were incubated in a light-tight holding chamber filled with ACSF at 30°C for 30 minutes and then allowed to recover at room temperature for at least 30 minutes. The slices were then transferred to a submersion-type recording chamber mounted on an upright microscope (Nikon, E600FN) with oblique infrared illumination.

Electrophysiology

Whole-cell current clamp recordings for LTP and LTD

Visually identified neurons were targeted at a 40%–50% depth from the pia corresponding to L4. The location of the majority of recorded cells was confirmed post hoc by biocytin labeling. Recording pipettes (3-5 MΩ) were filled with an internal solution consisting of (in mM): 130 K-glucuronate, 8 NaCl, 0.2 EGTA, 10 HEPES, 3 ATP, 10 Na-phosphocreatine, and 0.5 GTP, pH 7.4, 275–285 mOsm. Excitatory postsynaptic potential (EPSPs) were evoked by shining blue light (Thorlabs 455-nm LED, 5-ms duration) through a 40x objective lens to activate thalamic terminals. Stimulus intensity was adjusted to the minimum required to reliably produce single-peaked, short-onset latency (< 4 ms) EPSPs. Recording configuration switched from current-clamp to voltage-clamp for the plasticity induction protocol. The paired stimulus protocol entailed 200 pulses of presynaptic stimulation at 1-Hz coupled with postsynaptic depolarization to 0 mV or −40 mV (Jiang et al., 2007) to induce LTP or LTD, respectively. An Axon patch-clamp amplifier 700B (Molecular Devices) was used for whole cell recordings and data were acquired through Igor Pro software (WaveMetrics; RRID: SCR_000325). Changes in synaptic strength were quantified as the initial slope of the EPSP normalized by the average baseline slope obtained during the first 5 minutes of stable recordings. Cells were discarded if Vm > −65 mV or if R changed more than 30% during the experiment.

NMDAR/AMPA ratio

Isolated monosynaptic glutamatergic currents were recorded in the presence of bicuculline (20 μM) in ACSF with 4 mM Ca2+ and 4 mM Mg2+ using internal solution consisted of (in mM): 102 Cs-glucuronate, 5 TEA-chloride, 3.7 NaCl, 20 HEPES, 0.3 Na-GTP, 4 MgATP, 0.2 EGTA, 10 BAPTA, and 5 QX-314 chloride, pH 7.2, 300 mOsm. The LED intensity was adjusted to twice the minimum value required to consistently produce single-peaked, short-onset latency (< 4 ms) AMPAR-EPSCs. NMDAR and AMPAR responses were distinguished based on their kinetics and voltage dependence. The NMDAR component was measured as the amplitude 70-ms after the response onset at +40 mV, whereas the AMPAR component was measured as the peak amplitude recorded at −80 mV.

NMDAR kinetics

NMDAR-EPSCs were pharmacologically isolated with 20 mM bicuculline and 100 μM DL-2-amino-5 phosphonopentanoic acid (APV) and (ii) isolated with the same drugs with 1 μM tetrodotoxin (TTX) to confirm monosynaptic responses resulting from ChR2 activation of presynaptic inputs. Recording pipettes (3-5 MΩ) were filled with internal solution containing (in mM): 130 Cs-glucuronate, 8 KCl, 1 EGTA, 4 ATP, and 5 QX-314, pH 7.4, 285–295 mOsm. Visually identified neurons were voltage clamped at −80mV. Only recordings with Rs < 25 MΩ and Ri > 150MΩ that changed less than 15% were included in the analysis. The responses were evoked using a 455-nm LED (Thor labs) illuminated through a 40x objective lens (Nikon). Stimulation intensity was set to the minimal light required to produce a reliable response with 5-ms stimulus duration delivered every 10 s. Spontaneous events were quantified during a 400-ms time window before LED illumination (preLED) and LED-evoked Sr2+ desynchronized (LEV-Sr) events were quantified in a 400-ms window that started 50-ms from LED onset (postLED). Events were selected using MiniAnalysis (Synaptosoft; RRID: SCR_002184) with threshold set to 3 times the root mean square (RMS) noise. Cells with an RMS noise > 2 were excluded. To calculate the mean amplitude of evoked desynchronized events without baseline spontaneous activity, we used the following equation: [postLED amp x postLED frq] - (preLED amp x preLED frq) / (postLED frq - preLED frq) where amp is amplitude and frq is frequency. Calculated LEV-Sr2+ mEPSC amplitudes between the two recording conditions (with and without TTX) showed no significant changes (NR = 2.1 ± 2.8 pA, NRTTX = 16.4 ± 1.4 pA, unpaired Student’s t test p = 0.2080; mean DF = 18.61 ± 2.4 pA, DF_TTX = 20.8 ± 2.3 pA, unpaired t test p = 0.5200). Therefore, the data were pooled for Figure 3.
**E/I ratio**

Cortical L4 principal neurons were patched in voltage-clamp configuration with internal solution consisting of (in mM): 130 Cs-gluconate, 8 KCl, 1 EGTA, 10 HEPES, 4 ATP, and 5 QX-314, pH 7.4, 285–295 mOsm. Only recordings with Rs < 25 MΩ and Ri > 150 MΩ were included in the analysis. Monosynaptic EPSCs and disynaptic inhibitory postsynaptic currents (IPSCs) were recorded from each cell at the reversal potential for inhibitory currents and excitatory currents, respectively. After compensation for the junction potential, $E_{GABA}$ was −52 mV and $E_{GLU}$ was 0 mV. Thalamic terminals were activated with light flashes (LED 455-nm, 5-ms duration) delivered at several intensities until responses maximized, but did not evoke polysynaptic events. E/I ratios were calculated at each stimulation intensity for each cell. We noticed a tendency for E/I values to increase along with stimulation power until it reached a plateau at higher intensities (Figure S3). Therefore, we averaged the first three values of E/I ratio at intensities that produced the plateau for each cell and report this as the E/I ratio.

**Biocytin labeling**

Biocytin was added to the internal solution during most experiments to allow confirmation of neuronal location and morphology. Cortical slices were fixed in 4% paraformaldehyde overnight at 4°C, rinsed 10 minutes in 0.1 M phosphate buffer (PB) at room temperature and permeabilized in 2% Triton X-100 for one hour. Slices were then incubated in avidin-Texas red conjugate (Fisher) diluted 1:2000 in 1% Triton X-100 overnight at 4°C. Slices were washed twice in 0.1 M PB and mounted on glass slides with Prolong Gold antifade (Invitrogen). Images were taken on a laser scanning confocal microscope (Zeiss LSM 700) to confirm location of recorded cells post hoc.

**Monocular deprivation**

Mice were deeply anesthetized with isoflurane gas (3%) in an induction chamber. After the disappearance of toe pinch response, animals were transferred to a stereotaxic apparatus where oxygen supply was supplemented with isoflurane (1%–2%). The upper and lower margins of one eyelid were trimmed and sutured (PSD II; Ethicon) shut. The eye contralateral to the hemisphere being imaged was always sutured. Animals of the same sex were housed together (2-3 mice/cage) and disqualified if sutures opened.

**Optical imaging of intrinsic signals**

Mice were placed in a stereotaxic apparatus under constant supply of an oxygen/isoﬂurane mixture (0.7%–1.5% isoflurane), supplemented by a single injection of chlorprothixene (0.2 μg/g, ip). Body temperature was maintained at 37°C and heart rate was monitored throughout the experiment. The skull over V1 on the left hemisphere (contralateral to lid suture) was exposed and washed with hydrogen peroxide. Low melting point agarose (3%) and a glass coverslip were placed over the exposed area and allowed to solidify. V1 responses were recorded using the method previously developed by Kalatsky and Stryker (2003) and optimized for ocular dominance (OD) measurements (Cang et al., 2005). Optical images of cortical intrinsic signals were acquired using a Dalsa CCD camera (Dalsa, Waterloo, Canada) controlled by custom software. The surface vasculature and intrinsic signals were visualized with illumination wavelengths of 555-nm and 610-nm, respectively. After focusing on prominent vasculature marks, the camera was focused 600-μm below the surface. A high refresh rate monitor (ViewSonic) was placed 25-cm in front of the animal for stimulus presentation. The visual stimulus presented was restricted to the binocular visual field (5° to +15° azimuth) and consisted of a horizontal bar ($x = 5^\circ$, $y = 0^\circ$, width = 20) continuously presented for 5 minutes in upward ($90^\circ$) and downward ($270^\circ$) directions to each eye separately. The cortical response at the stimulus frequency was extracted by Fourier analysis and used to calculate the ocular dominance index (ODI). Two maps were averaged per eye for each animal to compute the ODI following the formula: (C-I)/(C+I) where C and I are the response magnitudes of each pixel to visual stimulation to the contralateral (C) and ipsilateral (I) eye respectively. The binocular area was selected as a region of interest (ROI) in the ODI map and the ODI values within this region were averaged.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

All statistical analysis was performed using Prism 7.0 (GraphPad Software; RRID: SCR_002798) and data are presented as mean ± SEM. Sample number for each experiment can be found in the figure legends. Unpaired two-tailed Student’s t test was used to compare recordings between NR and DF groups. Paired two-tailed t tests were used to compare ODI and eye-specific cortical activation responses measured from the same animals before and after MD. In all cases, p < 0.05 was considered statistically significant.
Supplemental Information

Cross-Modal Reinstatement of Thalamocortical Plasticity Accelerates Ocular Dominance Plasticity in Adult Mice

Gabriela Rodríguez, Darpan Chakraborty, Katrina M. Schrode, Rinki Saha, Isabel Uribe, Amanda M. Lauer, and Hey-Kyoung Lee
Supplementary Figures

Figure S1, Related to Figure 1. Confirmation of deafening.

(A) Increased threshold to elicit ABRs in DF mice across all frequency of tones presented as well as for a click (n=3 animals each; unpaired Student’s t-test: 8 kHz *p= 4.90e-006, 12 kHz *p=0.0001, 16 kHz *p=0.0002, 24 kHz *p=0.0004, click *p=0.0023). Note that 2 DF mice did not exhibit measureable ABR to determine the threshold when presented with a 32-kHz tone even at the maximum sound intensity used. Filled symbols: mean±SEM. Open symbols: individual mouse.  (B) Confirmation of cochlear damage by phalloidin staining (green) of hair cells from NR (top) and DF (bottom) animals. Sections were counterstained with DAPI (blue). Images were taken using the same setting on a confocal microscope. Note reduced phalloidin stain intensity in hair cells from DF. IHC: Inner hair cells. OHC: Outer hair cells. Scale bar: 20 µm.
Figure S2, Related to Figure 1. Failure of ChR2 to follow higher frequency activation.

A ChR2 expressing cortical neuron was recorded under current clamp configuration and activated with blue LED (455-nm, 5 ms duration; blue upward arrows indicate when LED stimulation occurred) to elicit action potentials. The neuron reliably elicited action potentials when stimulated at 20-Hz (A), but at 40-Hz stimulation frequency (B) there was occasional failure of action potentials (noted by the red arrows).
Figure S3, Related to Figure 3. Measurement of E/I ratio of LGN inputs to V1 L4.

LGN axons expressing ChR2 were activated by blue LED (455-nm, 5-ms duration) at different intensities, and monosynaptic EPSC and disynaptic IPSC were recorded in a V1 L4 principal neuron. As noticed, lower stimulation intensities recruited larger IPSC, which resulted in a lower E/I ratio. E/I ratio values gradually increased to reach a plateau as stimulation light intensity increased. Maximum stimulation intensity was the highest intensity used without recruiting polysynaptic events. Average of the three E/I ratio values at the plateau range (marked with a blue oval) was used for comparing E/I ratio between NR and DF groups shown in Fig. 3. Top: Traces of monosynaptic EPSC (inward traces) and disynaptic IPSC (outward traces) corresponding to the lowest intensity stimulation (yellow) and the start of the plateau (orange).
Figure S4, Related to Figure 4. Deafening adult mice for 1 week does not significantly alter ocular dominance index (ODI).

(A) No significant change in the average ODI measured before and after the 1 week deafening (paired t-test: N.S. p=0.141). Bars show mean±SEM. Gray lines connect ODI value for each animal. (B) No significant change in the cortical activation signal for contralateral eye (squares connected with solid line, paired t-test: N.S. p=0.426) and ipsilateral eye (triangles connected with dashed line, paired t-test: N.S. p=0.206). Mean±SEM.