Interactions between prefrontal cortex and cerebellum revealed by trace eyelid conditioning

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Eyelid conditioning has proven useful for analysis of learning and computation in the cerebellum. Two variants, delay and trace conditioning, differ only by the relative timing of the training stimuli. Despite the subtlety of this difference, trace eyelid conditioning is prevented by lesions of the cerebellum, hippocampus, or medial prefrontal cortex (mPFC), whereas delay eyelid conditioning is prevented by cerebellar lesions and is largely unaffected by forebrain lesions. Here we test whether these lesion results can be explained by two assertions: (1) Cerebellar learning requires temporal overlap between the mossy fiber inputs activated by the tone conditioned stimulus (CS) and the climbing fiber inputs activated by the reinforcing unconditioned stimulus (US), and therefore (2) trace conditioning requires activity that outlasts the presentation of the CS in a subset of mossy fibers separate from those activated directly by the CS. By use of electrical stimulation of mossy fibers as a CS, we show that cerebellar learning during trace eyelid conditioning requires an input that persists during the stimulus-free trace interval. By use of reversible inactivation experiments, we provide evidence that this input arises from the mPFC and arrives at the cerebellum via a previously unidentified site in the pontine nuclei. In light of previous PFC recordings in various species, we suggest that trace eyelid conditioning involves an interaction between the persistent activity of delay cells in mPFC—a putative mechanism of working memory—and motor learning in the cerebellum.

Results

Cerebellar learning requires that mossy fiber and climbing fiber inputs overlap in time

Recordings show that the activity of auditory-driven mossy fibers does not outlast the presentation of the tone—the majority of those recorded respond phasically to tone onset and a smaller
percentage show sustained responses that persist until tone offset (Boyd and Aitkin 1976; Aitkin and Boyd 1978; Freeman Jr. and Muckler 2003). Thus, during trace eyelid conditioning, mossy fiber activity directly driven by the tone does not overlap in time with the climbing fiber activity elicited by the US. To evaluate whether the cerebellum requires activity in mossy fibers other than those activated directly by the tone, we asked whether cerebellar learning fails when mossy fiber and climbing fiber activity do not overlap in time.

Identifying the temporal patterns of input necessary to engage cerebellar learning requires selective and precise temporal control over the activation of mossy fiber and climbing fiber inputs. Therefore, we used direct electrical stimulation of mossy fibers (see Fig. 3B) as a replacement for the tone (therefore precluding contributions from the forebrain) and periodical stimulation as the US since its activation of climbing fibers is relatively straightforward (Mauk et al. 1986; Sears and Steinmetz 1991; Hesslow 1994).

Trace eyelid conditioning using mossy fiber stimulation as the CS failed to promote learning, even though the same stimulation subsequently supported robust delay conditioning (Fig. 2). Nine rabbits were trained for 10 d using standard trace conditioning procedures (500 msec of mossy fiber stimulation and a 500-msec interval between stimulation offset and US). All subjects failed to learn (Fig. 2A,B, left) ($F_{9,72} = 0.78; P = 0.63$; main effect of trace conditioning session on response rate). The same subjects were then tested for their ability to learn eyelid responses with delay conditioning using the same stimulation parameters. For five subjects the interstimulus interval (ISI; the interval between CS and US onset) was 500 msec to match the stimulation duration used in trace conditioning. For the remaining four subjects, the CS was 1000 msec to match the ISI used in trace conditioning. All animals in both delay ISI 500 (Fig. 2A,B, middle) ($F_{5,20} = 21.93; P = 0.0001$; main effect of delay ISI 500 session on response rate) and delay ISI 1000 (Fig. 2A,B, right) ($F_{9,27} = 3.43; P = 0.006$; main effect of delay 1000 session on response rate) conditions showed robust learning.

Trace eyelid conditioning in rabbits is prevented by forebrain lesions when the trace interval (interval between CS offset and US onset) is 500 msec but not 300 msec (Moyer Jr. et al. 1990). To test whether this reflects the temporal limitations of cerebellar learning, we subsequently examined the ability of the cerebellum to learn at various trace intervals. We trained six additional rabbits using a 500-msec mossy fiber stimulation train as the CS and a trace interval of 200, 300, or 400 msec until a criterion level of responding (see Materials and Methods). If this criterion was achieved, the trace interval was increased by 100 msec and training continued. If not, the trace interval was decreased until learning was possible. This procedure permitted us to vary the trace interval several times in the same subject. We found that the capacity for learning decreased as the trace interval increased (Fig. 3A) ($F_{3,18} = 11.11; P = 0.0001$; main effect of trace interval on response rate). A 200-msec interval (Fig. 3C, left) supported robust learning in all animals, whereas learning was variable with trace intervals of 300 msec (Fig. 3C, middle) and 400 msec (Fig. 3C, right) and always failed with a 500-msec trace interval.

These data indicate that the cerebellum can learn when the climbing fiber input arrives no more than 200–400 msec after the offset of a mossy fiber input. This supports the hypothesis that the ability of forebrain lesions to impair trace conditioning with trace intervals longer than 300 msec (Moyer Jr. et al. 1990), and their inability to prevent delay conditioning (Solomon et al. 1986; Mauk and Thompson 1987; Kronforst-Collins and Disterhoft 1998; Weible et al. 2000; Powell et al. 2001; McLaughlin et al. 2002) relates to the temporal requirements of cerebellar learning. Combined with observations that the activity of auditory-driven mossy fibers does not outlast the presentation of a stimulus (Boyd and Aitkin 1976; Aitkin and Boyd 1978; Freeman Jr. and Muckler 2003), the data also support the assertion that during trace conditioning with relatively long trace intervals, the cerebellum requires mossy fiber activity during the trace interval that is distinct from activity driven directly by the tone. If this assertion is true and if the two sets of mossy fibers are anatomically segregated, it should be possible to abolish trace and not delay responding by inactivating the relevant mossy fiber inputs. We test this possibility in the next section.

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Figure 1. The procedures, neural pathways, and putative signals involved in delay and trace eyelid conditioning. (A) Stimulus timing for delay (left) and trace (right) training trials. For delay conditioning, the US overlaps in time with the tone CS. In this and subsequent figures, green is used to indicate the presentation of the CS for delay conditioning. For trace conditioning, the US is presented after CS offset, and “trace interval” refers to the period between CS offset and US onset. For convenience, we used red and maroon regions to represent the CS and trace interval, respectively. Sample conditioned eyelid responses are shown below, for which an upward deflection indicates closure of the eyelid. (B) Schematic representation of the pathways engaged by delay conditioning. The CS and US, respectively, engage mossy fibers and climbing fibers relatively directly, and forebrain input is not required for normal learning. (C) The signals hypothesized to engage the cerebellum during trace conditioning. The activity of mossy fibers directly activated by the tone CS does not significantly outlast the stimulus. Thus, a forebrain structure is thought to provide an input that overlaps in time with the US and is necessary to produce cerebellar learning.
Delay and trace conditioning engage cerebellar learning via separate sets of mossy fibers

One way to test this prediction is to train subjects on delay conditioning in the first phase and then trace conditioning in the second, while examining the effects of inactivating a specific set of mossy fibers after each training phase. However, any selective ablation of trace but not delay conditioned responses could be attributed to a number of differences during the separate test sessions (differences in drug concentration or diffusion, time-dependent effects, etc.) rather than successful inactivation of trace conditioning relevant mossy fibers. A more powerful way to control for such differences is to probe the effects of inactivation on delay and trace eyelid conditioning within the same animal and test session. We therefore developed a novel dual delay/trace conditioning procedure in which delay and trace conditioning trials were intermixed within the same training session. The CSs were two different tones (1 kHz and 9.5 kHz), with one used as the delay CS and the other as the trace CS (counterbalanced across subjects). We found that animals readily learned both delay and trace conditioned responses using this procedure, with delay responses learned in significantly fewer trials than trace responses (Fig. 4) \( (n = 11; F_{10,96} = 3.69; P = 0.004; \text{delay versus trace trials to criterion}). \)

To ensure that learning resulting from this procedure is comparable to that in which animals are trained with only trace or only delay conditioning, we examined the effects of inactivating the caudal mPFC and anterior interpositus nucleus (AIN) on learned responses. If learning in the dual training paradigm is comparable to learning established by training only one condition at a time, then inactivating the caudal mPFC should impair only trace conditioning (Mauk and Thompson 1987; Kronforst-Collins and Disterhoft 1998; Weible et al. 2000; Powell et al. 2001), while inactivating the AIN should abolish delay and trace conditioning (McComrick and Thompson 1984; Woodruff-Pak et al. 1985).

We trained animals using the dual delay/trace procedure and then tested the effects of reversibly inactivating the caudal mPFC or the AIN with the GABA\(_A\) agonist muscimol on the expression of learned responses. Consistent with previous observations, we found that infusing muscimol into the AIN abolished delay \( (\text{Fig. 5, filled symbols}) (F_{11,44} = 11.12; P < 0.0001; \text{block by session type [muscimol vs. last day of training interaction]}) \) and trace conditioning \( (n = 5; F_{11,44} = 16.13; P < 0.0001; \text{block by session type [muscimol vs. last day of training interaction]}) \). Also consistent with previous observations, muscimol infusions abolished trace \( (F_{11,55} = 5.03; P < 0.0001; \text{block by session type [muscimol vs. last day of training interaction]}) \) but not delay conditioning \( (n = 6; F_{11,55} = 1.09; P = 0.39; \text{block by session type [muscimol vs. last day of training interaction]}) \) in subjects with cannula placements in or near the anterior cingulate or prelimbic areas of caudal mPFC \( (\text{Fig. 6A, filled symbols, B, D, top row}) \). Artificial cerebrospinal fluid (ACSF) infusions had no effect on trace responses in these subjects \( (\text{Fig. 6A, open symbols}) (F_{11,55} = 1.15; P = 0.34; \text{block by session type [ACSF vs. last day of training interaction]}) \). Furthermore, muscimol infusions failed to abolish either delay \( (n = 7; F_{11,64} = 0.85; P = 0.59; \text{block by session type [muscimol vs. last day of training interaction] or trace conditioning}) \) or trace conditioning \( (F_{11,64} = 0.69; P = 0.75; \text{block by session type [muscimol vs. last day of training interaction]}) \) in subjects with cannula placements dorsal or caudal to effective cannula placements \( (\text{Fig. 6C,D, bottom row}) \).

These data demonstrate that the effects of lesions of the forebrain and cerebellum on responses using the dual delay/trace procedure parallel those seen previously for animals trained using delay and trace conditioning procedures in separate experiments. Inactivation of the AIN abolished delay and trace responses, while inactivation of mPFC abolished only trace responses. Because effective mPFC infusion sites \( (\text{Fig. 6D, top row}) \) tended to be more ventral and rostral than ineffective ones \( (\text{Fig. 6D}) \), our data suggest that the anterior cingulate and/or prelimbic areas of caudal mPFC are important for trace eyelid conditioning. Moreover, consistent with the relatively direct connections between the caudal mPFC and cerebellum in rabbits \( (\text{Weible et al. 2007}) \), the data indicate that the mPFC is a candidate source of the forebrain signal that engages cerebellar learning during trace eyelid conditioning.

The above data demonstrate that the dual delay/trace conditioning procedure is an effective assay for identifying sites that are
selectively important for trace eyelid conditioning. Therefore in another set of subjects, we used the procedure to identify mossy fibers important for trace conditioning. Muscimol infusions in eight well-trained animals with cannula placements in the lateral pontine nuclei (LPN), a source of mossy fibers that receives direct connections from areas of mPFC (Buchanan et al. 1994; Weible et al. 2007) close to our effective cannula placements (Fig. 6D, top), abolished trace ($F_{(11,77)} = 4.31; P < 0.0001$; block by session type [muscimol vs. last day of training] interaction) but not delay responses ($F_{(11,77)} = 0.98; P = 0.47$; block by session type [muscimol vs. last day of training] interaction) (Fig. 7A, filled symbols, B, D, top row). ACSF infusions in these animals had no effect on trace responses ($n = 5; F_{(11,44)} = 1.32; P = 0.25$; block by session type [ACSF vs. last day of training] interaction) (Fig. 7A, open symbols). Furthermore, muscimol infusions failed to abolish either delay ($F_{(11,66)} = 1.32; P = 0.24$; block by session type [muscimol vs. last day of training] interaction) or trace conditioning ($n = 7; F_{(11,66)} = 0.62; P = 0.80$; block by session type [muscimol vs. last day of training] interaction) in subjects with cannula placements near but outside of the LPN (by as little as < 500 μm in some cases) (Fig. 7C,D, bottom row).

These data show that trace conditioning using a tone CS and a 500-msec trace interval requires activation of mossy fiber inputs that are distinct from those that are driven directly by auditory input and that are important for delay conditioning. The data also suggest that these mossy fibers originate in the LPN, as effective infusions sites were always located within the LPN. Furthermore, given that the caudal mPFC projects directly to the LPN (Buchanan et al. 1994; Weible et al. 2007), the data also indicate that the caudal mPFC provides the cerebellum with the mossy fiber activity necessary to engage learning during trace eyelid conditioning with a relatively long trace interval.

An examination of the response rates during muscimol infusions into the AIN, mPFC, and LPN (Figs. 5–7, respectively) reveals what appears to be a delayed effect. During each type of infusion, responses decrease over the first few post-infusion blocks. For the LPN and AIN infusions, this is an averaging effect—some subjects’ responses abolished quickly, while others’ abolished more slowly. Thus, the apparent delayed effect of LPN and AIN infusions is likely due to individual differences in cannula placement and/or drug diffusion. For the mPFC infusions, we believe that the actual effective infusion site may be a small distance from our cannulae. We have noticed a strong tendency for ventral and anterior infusion sites to produce more robust and rapid effects on trace conditioning. Defining the precise region of mPFC necessary for the expression of trace responses will require systematically placing cannulae over a wide range of mPFC. It is also possible that our mPFC infusions caused extinction, rather than abolition of responses. We are currently investigating these possibilities.

**LPN involvement is specific to trace conditioning and is not related to the ISI**

Trace conditioning differs from delay conditioning not only because it has a stimulus-free trace interval but also because it usually involves a longer ISI. Because of this, the effects of forebrain lesions on trace eyelid conditioning have sometimes been attributed to the difficulty of learning eyelid conditioning procedures with long ISIs rather than the presence of the trace interval (Beylin et al. 2001). Thus, it is not clear whether the necessity of the forebrain during trace eyelid conditioning is attributable to the presence of the trace interval or to the long ISI.

![Figure 3](image-url) The ability of subjects to learn trace conditioning with a mossy fiber stimulation CS decreases as the trace interval increases. (A) Average response rates, normalized to delay conditioning trials, from the session after criterion was reached ($n = 6$). (B) Coronal section through the cerebellum shows a representative stimulation site in the middle cerebellar peduncle for the studies in this figure and in Figure 2. All stimulation sites were in the middle cerebellar peduncle. (C) Representative traces from the session after criterion was reached for trace intervals of 200, 300, and 400 msec.

![Figure 4](image-url) Learning during dual delay/trace conditioning. (Left) The acquisition of delay responses (green squares) was faster than the acquisition of trace responses (red circles) during dual delay/trace training. Note that both delay and trace responses decrease during each session. We also observe this phenomenon in animals trained with only trace or only delay conditioning. This effect increases with the ISI and will be presented in a forthcoming paper along with a more detailed description and mechanistic implications. (Right) The faster acquisition of delay conditioning can also be seen in the trials needed to reach a criterion level of responding (see Materials and Methods). **$P < 0.01$.**
We trained three subjects using the dual delay/trace conditioning procedure in which the delay conditioning ISI was longer than the trace conditioning ISI. The trace conditioning tone was 100 msec and was followed by a 500-msec trace interval for a total ISI of 600 msec, while the delay conditioning ISI was 1000 msec. Post-training infusions of muscimol into the LPN abolished trace responses (F(1,1) = 3.58; P = 0.005; block by session type muscinol vs. last day of training interaction) but not delay responses (F(1,1) = 1.44; P = 0.22; block by session type muscinol vs. last day of training interaction) (Fig. 8A, filled symbols, B). These data indicate that the LPN-and by inference the mPFC-is necessary for trace conditioning because of the presence of the trace interval and not because trace conditioning often has a longer ISI than delay conditioning (Beylin et al. 2001).

Discussion
We have tested two component assertions of the hypothesis that trace eyelid conditioning involves cerebellar learning in response to an input from the forebrain. First, we showed that the cerebellum cannot support learning when climbing fiber inputs arrive more than a few hundred milliseconds after the end of a mossy fiber input, as occurs for tone-activated mossy fibers during trace conditioning with trace intervals longer than 200–400 msec (Figs. 2, 3). Thus, our data indicate that under these conditions, mossy fiber inputs other than those activated by the auditory system must provide activity during the trace interval for learning to occur.

Second, we provided direct evidence that cerebellar learning during delay and trace conditioning using an auditory CS occurs to different mossy fiber inputs. We showed that cerebellar learning during trace eyelid conditioning occurs in response to activity in a set of mossy fibers that originate in the LPN and that are distinct from tone-CS activated mossy fibers (Fig. 7). Given that the mPFC projects to the LPN and that inactivating the caudal mPFC selectively abolishes trace conditioning (Fig. 6), we suggest that the mossy fibers necessary for trace conditioning convey input to the cerebellum from the mPFC. Together with previous observations (Kronforst-Collins and Disterhoft 1998; Weible et al. 2000; Simon et al. 2005), our data also indicate that this mPFC-driven input is necessary for both the acquisition and expression of trace conditioned responses.

Together, these observations provide a specific explanation for why trace conditioning is sensitive to lesions of both the cerebellum and forebrain structures when the trace interval is longer than ~400 msec. Given that mossy fiber inputs driven directly by a tone CS terminate shortly after CS offset (Boyd and Aitkin 1976; Aitkin and Boyd 1978; Freeman Jr. and Muckler 2003), the pattern of mossy fiber and climbing fiber inputs driven directly by the CS and US during trace conditioning is not appropriately timed to engage cerebellar learning. Thus, a second mossy fiber input is required that arises from the LPN, is driven by input from the mPFC, and presumably provides activity that overlaps in time with the US. Indirect evidence suggests that this input is learned-neurons in the hippocampus (Moyer Jr. et al. 1996; McEchron and Disterhoft 1997; McEchron et al. 2001) and mPFC (Weible et al. 2003) show learning-related changes during trace conditioning, and the acquisition of trace conditioned responses requires more training trials than is required for delay conditioning (Fig. 4; Beylin et al. 2001). We therefore suggest that trace conditioning with a trace interval longer than ~400 msec is mediated by a multistage process in which forebrain learning precedes, and is required for, subsequent cerebellar learning.

Implications for delay eyelid conditioning
It is interesting that inactivating the LPN did not affect the expression of delay conditioned responses, given that the LPN have previously been implicated as a locus of the auditory-driven mossy fibers that are necessary for delay eyelid conditioning to a tone (Steinmetz et al. 1987). Because the pontine nuclei are a large structure that span several millimeters in the anterior-posterior axis, our effective infusions may have been in an area of the LPN that lacks auditory-driven mossy fibers. This interpretation is supported by observations that short-latency auditory evoked activity is restricted to caudal aspects of the LPN and that lesions of the caudal, but not rostral, LPN affect delay conditioning to a tone (Steinmetz et al. 1987). Given that our effective infusions tended to be in rostral regions of the LPN (Fig. 7), mPFC-driven and auditory-driven mossy fibers may be segregated along the anterior-posterior axis of the LPN. A systematic mapping of single-unit, auditory, and mPFC-driven activity along the anterior-posterior axis of the LPN is needed to test this hypothesis.

The majority of auditory-driven mossy fibers that have been recorded do not show sustained activity in response to a tone (Boyd and Aitkin 1976; Aitkin and Boyd 1978; Freeman Jr. and Muckler 2003), suggesting the potential need for a mechanism to sustain mossy fiber activity during delay eyelid conditioning as well. Although the potential for selection bias with in vivo recordings requires that such conclusions remain tentative, the apparent majority of phasically responding auditory-driven mossy fibers would be unable to engage cerebellar learning mechanisms during delay conditioning because the offset of their activity would occur more than a few hundred milliseconds before a climbing fiber input. Thus, the cerebellum may require a mechanism...
that sustains mossy fiber activity to overlap with the US during delay conditioning. Our data indicate that this sustained input would not require the mPFC, as inactivating the mPFC did not abolish delay responses. There are a variety of alternatives.

Sustained mossy fiber responses to tones may be driven by the forebrain-auditory system (e.g., the inferior colliculus or auditory thalamus) (Halverson and Freeman 2006; Freeman et al. 2007) or by feedback from the cerebellum or red nucleus (Cartford et al.

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**Figure 6.** The mPFC is necessary for the expression of trace, but not delay, conditioning. (A) Infusing muscimol bilaterally into the mPFC after the third block of dual delay/trace conditioning (break in abscissa) abolished trace (filled red circles) but not delay responses (filled green squares), while infusing ACSF had no affect (open squares indicate delay; open circles, trace; n = 6). (B) Representative traces taken from an effective muscimol infusion session (same subject as shown in D, top left). (C) Representative traces taken from an ineffective muscimol infusion session (same subject as in D, bottom right). Note that in this example, neither trace nor delay responses were affected by the infusion. (D) Histological verification of cannula placements revealed that all cannula placements were in the vicinity of the caudal mPFC. Closer examination revealed that effective infusion sites tended to be more rostral and ventral than ineffective sites, suggesting that the necessary site(s) for trace eyelid conditioning is in the anterior cingulate and/or prelimbic cortices.

**Figure 7.** Mossy fibers originating in the LPN are necessary for the expression of trace, but not delay, conditioning. (A) Infusing muscimol into the LPN after the fourth block of dual delay/trace conditioning (break in abscissa) abolished trace (filled red circles) but not delay responses (filled green squares), while infusing ACSF had no affect (open squares indicate delay; open circles, trace; n = 8 for muscimol and 5 for ACSF). (B) Representative traces taken from an effective muscimol infusion session (same subjects as shown in D, top middle). (C) Representative traces taken from an ineffective muscimol infusion session (same subjects as in D, bottom left). (D) Histological verification of cannula placements revealed that effective infusion sites were located in the LPN, while ineffective sites were located near (<1mm), but outside of, the LPN.
Forebrain-cerebellum interactions during trace eyelid conditioning

Our findings establish trace eyelid conditioning as a useful tool to study cerebellar computation during interactions with the forebrain. Delay conditioning has proven particularly useful in characterizing the input/output transformations of the cerebellum (Mauk and Donegan 1997; Medina and Mauk 2000; Medina et al. 2000, 2002; Ohyama et al. 2003), due to the relatively direct ways that it engages cerebellar inputs (Mauk et al. 1986; Steinmetz et al. 1987, 1989; Sears and Steinmetz 1991; Hesslow 1994; Hesslow et al. 1999) and output (McCormick and Thompson 1984) and to its insensitivity to forebrain lesions (Mauk and Thompson 1987; Moyer Jr. et al. 1990; Kronforst-Collins and Disterhoft 1998; Weible et al. 2000; Powell et al. 2001). Despite its utility in this regard, delay eyelid conditioning represents a somewhat atypical situation where cerebellar afferents are activated relatively directly by somatosensory and auditory stimuli. Large numbers of mossy and climbing fiber inputs to the cerebellum are, in contrast, driven by descending forebrain projections (Ito 1984; Schmahmann and Pandya 1995; 1997; Middleton and Strick 2000; Weible et al. 2007). Thus, the typical mode of cerebellar information processing in the mammalian brain likely involves interactions with the forebrain. Our results therefore highlight the potential for using trace conditioning to study cerebellar information processing in a more typical context involving interactions with the forebrain.

More generally, our findings highlight the potential importance of learned (or potentially sensory-driven) cortical inputs driving the mossy fiber system for cerebellar learning. Given the massive amount of cortico-pontine input, it may be that the patterns of mossy fiber input driven relatively directly by the outside world are normally too brief to engage cerebellar learning mechanisms. Forebrain learning may therefore serve to expand the repertoire of mossy fiber inputs available for cerebellar learning by internally generating mossy fiber inputs that overlap with climbing fiber input (Clark et al. 2002).

Why does cerebellar learning fail when mossy fiber and climbing fiber activity do not overlap in time? Long-term depression (LTD) at the granule cell-Purkinje cell synapse and an increase in the efficacy of the mossy fiber to deep cerebellar nucleus cell synapse are two types of plasticity presumed to underlie cerebellar learning (Mauk and Donegan 1997; Clark et al. 1997; Bao et al. 2000). Alternatively, it may simply be the case that the number of mossy fibers that naturally show sustained responses to a tone is sufficient to support cerebellar learning during delay conditioning.

Finally, we showed that the necessity of the forebrain is related specifically to the trace interval and not the by-product of the long ISIs often used during trace conditioning. By use of combined delay/trace conditioning where the ISI for trace conditioning was shorter than the ISI for delay (600 msec for trace vs. 1000 msec for delay), we demonstrated that inactivating the LPN abolishes trace responses but not delay responses (Fig. 8). These data reinforce the interpretation that the forebrain activity relayed to the cerebellum via the LPN is required to span the stimulus-free trace interval and is not required because the ISI is long. This interpretation is also supported by the observation that subjects acquired delay conditioning with 1000-msec ISI, but not trace conditioning with 1000-msec ISI with mossy fiber stimulation as the CS (Fig. 2).

Figure 8. The necessity of the LPN in trace conditioning is due to the trace interval and not a longer ISI. For these experiments, the trace conditioning tone was 100 msec, the trace interval was 500 msec, and the delay conditioning ISI was 1000 msec. (A) Infusing muscimol into the LPN after the fourth block of dual delay/trace conditioning (break in abscissa) abolished trace (filled red circles) but not delay responses (filled green squares), while infusing ACSF had no affect (open squares indicate delay; open circles, trace; n = 3). (B) Representative traces taken from an effective muscimol infusion session.
overlaps in time with the US. From the cerebellum's point of view, this overlap, in affect, turns trace conditioning into delay conditioning with a relatively long ISI.

Future studies could test this forebrain-cerebellum hypothesis rather directly, given that the level of acetylcholine and certain monoamines in the PFC affect working memory and modulate its neural correlates (Williams and Goldman-Rakic 1995; Chudasama et al. 2004; Vijayraghavan et al. 2007; Wang et al. 2007). For example, relatively small doses of D1 dopamine receptor antagonists improve working memory and potentiate the task-related activity of PFC neurons (Williams and Goldman-Rakic 1995; Vijayraghavan et al. 2007). The working memory hypothesis predicts that trace eyelid conditioning should be similarly affected by these compounds. Another prediction is that recording from neurons in mPFC or LPN should reveal cells that fire persistently during the trace interval. This activity should gradually emerge early in training before overt, behavioral learning occurs and continue for as long as eyelid responses persist. Confirming this prediction would make trace eyelid conditioning a useful model system to study how delay cell activity in the PFC is learned.

Finally, the concrete framework provided by our hypothesis could help clarify the elusive role of the hippocampus in trace eyelid conditioning. Given the direct connections between the hippocampus and mPFC (Swanson 1981; Ferino et al. 1987; Jay and Witter 1991) and the time-dependent effects of hippocampal lesions on trace conditioning (Kim et al. 1995; Takehara et al. 2003), neural correlates of trace conditioning in the hippocampus (McEchron and Disterhoft 1997; McEchron et al. 2001) may play a crucial role in the establishment, but not the maintenance, of sustained activity in the PFC (Gilmartin and McEchron 2005; Hasselmo and Stern 2006).

Materials and Methods

Subjects

We obtained data from 57 male New Zealand albino rabbits (Oryctolagus cuniculus) (Myrtle's Rabbitry, TN). The animals weighed between 2.5 and 3 kg, were individually housed, and had free access to food and water. Treatment of animals and surgical procedures were in accordance with National Institutes of Health Guidelines and an institutionally approved animal welfare protocol.

Surgery

All animals were prepared with a head bolt cemented to the skull and a 26-gauge stainless steel guide cannula (Plastics One) or a tungsten stimulating electrode (A-M Systems; tip exposed to constant current pulses by stimulus isolators (World Precision Instruments, model A360) connected to the electrodes implanted in the middle cerebellar peduncle. As in previous studies, (Medina et al. 2000), an infrared emitter/detector was attached directly to the head stage of each subject to record movements of the left external eyelid by detecting changes in the amount of reflected light.

All daily conditioning procedures consisted of 12 blocks of nine trials (one CS-alone trial and eight paired trials/block). Trials were separated by a random intertrial interval in the range of 25 to 35 sec. The CS was either a 1-kHz, 85-dB tone; a 9.5-kHz, 85-dB tone; or cathodal stimulation (100 Hz, 40-μs pulse width, 100 μA) of the middle cerebellar peduncle. The US was a 50-msec train of constant current pulses (100 Hz, 1-msec pulse width, 2–3 mA) delivered through the periorbital electrodes. During all paired trace conditioning trials, the CS was presented for either 500 or 100 msec, and the US was presented after the offset of the CS by 100–500 msec, depending on the experiment. During paired delay conditioning trials, the CS was presented for either 550 or 1050 msec and coterminated with the US. For dual delay/trace conditioning procedures, a delay trial followed either one or two trace trials. Stimulus presentation was controlled by custom-designed software.

Infusions

After subjects were well trained using the dual delay/trace conditioning paradigm (10–15 sessions), we infused 1 mM of the GABAₐ agonist muscimol (Tocris), or ACSF through a 33-gauge internal cannula that extended 1.2 mm beyond the guide cannula during test sessions. Muscimol was dissolved in ACSF consisting of (in mM): NaCl 119, KCl 2.5, NaH₂PO₄ 1.2, MgCl₂ 2, CaCl₂ 2, NaHCO₃ 26, D-glucose 10, and HEPES 20 (pH 7.35–7.4). The internal cannula was coupled to a 50-μL Hamilton syringe that was mounted on an automated injector system (Bioanalytical Systems; model MD-1001) and driven by an electronic pump (model MD-1020). Infusions into cannula aimed at the LPN and AIN began after the fourth block and continued at a rate of 0.1 μL/min until training resumed 20 min later. Infusions into cannula aimed at the mPFC began after the third block and continued at a rate of 0.1 μL/min until training resumed 20 min later. Infusions into a given brain region were conducted within subjects. The order of infusion type (muscimol vs. ACSF) was counterbalanced across subjects and at least one day of retraining was given between infusion sessions to ensure there were no long-term effects.

Data analysis

We analyzed digitized sweeps of eyelid movements made 200 msec before and 2300 msec after CS onset with custom software. A conditioned response was defined as an eyelid movement of at least 0.3 mm within the ISI. Trials in which eyelid movements greater than 0.3 mm were made within 200 msec before CS onset were excluded from analysis. ANOVA was used to test for within- and between-subject differences. The significance level for all tests was 0.05. To determine the effects of muscimol or ACSF infusions on responding, we compared infusion data with those taken from the last day of training. We defined response rates as affected if there was a significant block by training session (muscimol, ACSF, or last day of training) interaction. We made this comparison...
because we observed a significant and reliable decrease in responding
that occurred naturally within sessions during both delay and
trace conditioning (a phenomenon we will describe in more detail
in a forthcoming paper), rendering a pre- versus post-infusion
comparison inappropriate. We used the Bonferroni method to
correct for type I errors associated with multiple comparisons.

Because the experiments of Figures 3 and 4 were designed for
different purposes, we defined two different criteria for learning.
In the experiments of Figure 3, where the objective was to detect
the first signs of learning regardless of its robustness, the criterion
for learning was defined as the session in which subjects made
three responses in each of two consecutive blocks. In our hands,
this criterion predicted well that a subject would reach asymptotic
responding the day after criterion was reached (unpublished
observations) and thus allowed us to vary the trace interval
multiple times in the same subject. Three subjects were given
the 200-msec trace interval first, one subject the 300-msec interval
first, and two subjects the 400-msec interval first. When a subject
reached criterion for a given trace interval, they were given one
more session with that interval before switching to a session with
a new interval. The response rate during the day after criterion was
reached was used in analysis. Because not all subjects received the
same trace procedures or order of training, we used a between-
subjects ANOVA to analyze these data. In the experiments of
Figure 4, where the objective was to detect asymptotic learning,
the criterion for learning was defined as five responses made in
each of two consecutive blocks. We used this criterion rather than
the classic criterion of eight responses in any nine consecutive
trials to accommodate the lower asymptote of trace conditioning
relative to delay conditioning.

Histology
Infusion and electrode sites were marked by passing 200 μA,
anodal current for 15 sec through a wire (cut to the length of
the internal cannula and threaded through the guide cannula) or
stimulating electrode, respectively. Animals were killed with an
overdose of sodium pentobarbital and were perfused transcardially
with 0.9% saline. Brains were imbedded in gelatin and sectioned
at 80 μm using a microtome. Sections were mounted and stained
with cresyl violet.

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References
Atkin, L.M. and Boyd, J. 1978. Acoustic input to the lateral pontine nuclei.
Hear. Res. 1: 67–77.
Bao, S., Chen, L., and Thompson, R.F. 2000. Learning and cerebellum-
dependent neuronal activity in the lateral pontine nucleus. Behav.
Neurosci. 114: 254–261.
Bevlin, A.V., Gandhi, C.C., Wood, G.E., Talk, A.C., Matzel, L.D., and
Shors, T.J. 2001. The role of the hippocampus in trace conditioning: Temporal
discontinuity or task difficulty? Neurobiol. Learn. Mem. 76: 447–461.
Bodner, M., Kroger, J., and Fuster, J.M. 1999. Auditory memory cells in
the rat. Exp. Brain Res. 100: 469–483.
Cardoff, M.C., Bodner, M., and Lavond, D.G. 1997. The role of awareness.
Science 280: 77–81.
Clark, R.E., Gohi, E.B., and Lavond, D.G. 1997. The learning-related activity
that develops in the pontine nuclei during classical eyeblink conditioning
is dependent on the interpositus nucleus. Learn. Mem. 3: 532–544.
Clark, R.E., Mandès, J.R., and Squire, L.R. 2002. Classical conditioning,
awareness, and brain systems. Trends Cogn. Sci. 6: 528–531.
Ferino, F., Thierry, A.M., and Giovisi, J. 1987. Anatomical and
electrophysiological evidence for a direct projection from Ammon's horn
to the medial prefrontal cortex in the rat. Exp. Brain Res. 65: 421–426.
Fisman Jr., J.H. and Mucke, L. 2003. Developmental changes in
eyeblink conditioned and neuronal activity in the pontine nuclei.
Learn. Mem. 10: 337–345.
Fisman Jr., J.H., Halverson, H.E., and Hubbard, E.M. 2007. Inferior colliculus
lesions impair eyelash conditioning in rats. Learn. Mem. 14: 842–846.
Fusinato, S., Bruce, C.J., and Goldman-Rakic, P.S. 1989. Mnemonic coding
of visual space in the monkey's dorsolateral prefrontal cortex. J.
Neuropsychol. 61: 331–349.
Fuster, J.M., Bodner, M., and Kroger, J.K. 2000. Cross-modal and cross-
temporal association in neurons of frontal cortex. Nature 405: 347–351.
Gilmartin, M.R. and McEchron, M.D. 2005. Single neurons in the medial
prefrontal cortex of the rat exhibit tonic and phasic coding during trace
ear conditioning. Behav. Neurosci. 119: 1496–1510.
Halverson, H.E. and Freeman, J.H. 2006. Medial auditory thalamic nuclei
are necessary for eyelash conditioning. Behav. Neurosci. 120: 880–887.
Hansel, C., Linden, D.J., and D'Angelo, E. 2001. Beyond parallel fiber LTD:
The diversity of synaptic and non-synaptic plasticity in the cerebellum.
Nat. Rev. Neurosci. 4: 467–475.
Hasselmo, M.E. and Stern, C.E. 2006. Mechanisms underlying working
memory for novel information. Trends Cogn. Sci. 10: 487–493.
Hesslow, G. 1994. Correspondence between climbing fibre input and motor
output in eyelash-related areas in cat cerebellar cortex. J. Physiol. 476:
229–244.
Hesslow, G., Svensson, P., and Iverson, M. 1999. Learned movements
elicited by direct stimulation of cerebellar mossy fiber afferents. Neuron
24: 179–185.
Ito, M. 1984. The cerebellum and neural control. Raven Press, New York.
Jay, T.M. and Witter, M.P. 1991. Distribution of hippocampal CA1 and
subicular efferents in the prefrontal cortex of the rat studied by means of
anterograde transport of Phaseolus vulgaris-leucoagglutinin. J. Comp.
Neurol. 313: 574–586.
Kim, J.J., Clark, R.E., and Thompson, R.F. 1995. Hippocampectomy impairs
the memory of recently, but not remotely, acquired trace eyelash
conditioned responses. Behav. Neurosci. 109: 195–203.
Kronforst-Collins, M.A. and Disterhoft, J.F. 1998. Lesions of the caudal area
of rabbit medial prefrontal cortex impair trace eyelash conditioning.
Neurobiol. Learn. Mem. 69: 147–162.
Mauk, M.D. and Donegan, N.H. 1997. A model of Pavlovian eyelid
conditioning based on the synaptic organization of the cerebellum.
Learn. Mem. 4: 130–158.
Mauk, M.D. and Thompson, R.F. 1987. Retention of classically conditioned
eyelash responses following acute decerebration. Brain Res. 403: 89–95.
Mauk, M.D., Steinmetz, J.E., and Thompson, R.F. 1986. Classical
conditioning using stimulation of the inferior olive as the
unconditioned stimulus. Proc. Natl. Acad. Sci. 83: 5349–5353.
McCormick, D.A. and Thompson, R.F. 1984. Cerebellum: Essential
involvement in the classically conditioned eyelid response. Science
223: 296–299.
McEchron, M.D. and Disterhoft, J.F. 1997. Sequence of single neuron
changes in CA1 hippocampus of rabbits during acquisition of trace
eyelash conditioned responses. J. Neurophysiol. 78: 1030–1044.
McEchron, M.D., Weible, A.P., and Disterhoft, J.F. 2001. Aging and learning-
specific changes in single-neuron activity in CA1 hippocampus during
rabbit trace eyelash conditioning. J. Neurophysiol. 86: 1839–1857.
McLaughlin, J., Skaggs, H., Churchwell, J., and Powell, D.A. 2002. Medial
prefrontal cortex and pavlovian conditioning: Trace versus delay
conditioning. Behav. Neurosci. 116: 37–47.
Medina, J.F. and Mauk, M.D. 2000. Computer simulation of cerebellar
information processing. Brain Res. 885: 1205–1211.
Medina, J.F., Garcia, K.S., Nores, W.L., Taylor, N.M., and Mauk, M.D.
2000. Timing mechanisms in the cerebellum: Testing predictions of a
large-scale computer simulation. J. Neurosci. 20: 5516–5525.
Medina, J.F., Nores, W.L., and Mauk, M.D. 2002. Inhibition of climbing
fibres is a signal for the extinction of conditioned eyelash responses.
Nature 416: 330–333.
Mediwake, N.A. and Strick, P.L. 2000. Basal ganglia and cerebellar loops:
Motor and cognitive circuits. Brain Res. 885: 1236–250.
Moyer Jr., J.R., Deyo, R.A., and Disterhoft, J.F. 1990. Hippocampus
mediates trace eye-blink conditioning in rabbits. Behav. Neurosci. 104:
243–252.
Moyer Jr., J.R., Thompson, L.T., and Disterhoft, J.F. 1996. Trace eyelash
conditioning increases CA1 excitability in a transient and learning-
specific manner. J. Neurosci. 16: 5536–5546.
Prefrontal cortex-cerebellum interactions

Narayan, N.S. and Laubach, M. 2006. Top-down control of motor cortex ensembles by dorsomedial prefrontal cortex. *Neuron* 52: 921–931.

Ohyama, T., Notes, W.L., Murphy, M., and Mauk, M.D. 2003. What the cerebellum computes. *Trends Neurosci.* 26: 222–227.

Ohyama, T., Notes, W.L., Medina, J.E., Rüsech, F.A., and Mauk, M.D. 2006. Learning-induced plasticity in deep cerebellar nucleus. *J. Neurosci.* 26: 12656–12663.

Powell, D.A. and Churchwell, J. 2002. Mediodorsal thalamic lesions impair trace eyeblink conditioning in the rabbit. *Learn. Mem.* 9: 10–17.

Powell, D.A., Skaggs, H., Churchwell, J., and McLaughlin, J. 2001. Posttraining lesions of the medial prefrontal cortex impair performance of Pavlovian eyeblink conditioning but have no effect on concomitant heart rate changes in rabbits (*Oryctolagus cuniculus*). *Behav. Neurosci.* 115: 1029–1038.

Pugh, J.R. and Raman, I.M. 2006. Potentiation of mossy fiber EPSCs in the cerebellar nuclei by NMDA receptor activation followed by postinhibitory rebound current. *Neuron* 51: 113–123.

Schmahmann, J.D. and Pandya, D.N. 1995. Prefrontal cortex projections to the basilar pons in rhesus monkey: Implications for the cerebellar contribution to higher function. *Neurosci. Lett.* 199: 175–178.

Schmahmann, J.D. and Pandya, D.N. 1997. Anatomic organization of the basilar pontine projections from prefrontal cortices in rhesus monkey. *J. Neurosci.* 17: 438–458.

Sears, L.L. and Steinmetz, J.E. 1991. Dorsal accessory inferior olive activity diminishes during acquisition of the rabbit classically conditioned eyelid response. *Brain Res.* 545: 114–122.

Simon, B., Knuckley, B., Churchwell, J., and Powell, D.A. 2005. Posttraining lesions of the medial prefrontal cortex interfere with subsequent performance of trace eyeblink conditioning. *J. Neurosci.* 25: 10740–10746.

Solomon, P.R., Vander Schaaf, E.R., Thompson, R.F., and Weisz, D.J. 1986. Hippocampus and trace conditioning of the rabbit's classically conditioned nictitating membrane response. *Behav. Neurosci.* 100: 729–744.

Steinmetz, J.E., Logan, C.G., Rosen, D.J., Thompson, J.K., Lavond, D.G., and Thompson, R.F. 1987. Initial localization of the acoustic conditioned stimulus projection system to the cerebellum essential for classical eyelid conditioning. *Proc. Natl. Acad. Sci.* 84: 3531–3535.

Steinmetz, J.E., Lavond, D.G., and Thompson, R.F. 1989. Classical conditioning in rabbits using pontine nucleus stimulation as a conditioned stimulus and inferior olive stimulation as an unconditioned stimulus. *Synapse* 3: 225–233.

Swanson, L.W. 1981. A direct projection from Ammon’s horn to prefrontal cortex in the rat. *Brain Res.* 217: 150–154.

Takehara, K., Kawahara, S., and Kurino, Y. 2003. Time-dependent reorganization of the brain components underlying memory retention in trace eyeblink conditioning. *J. Neurosci.* 23: 9897–9905.

Thompson, R.F. 1986. The neurobiology of learning and memory. *Science* 233: 941–947.

Vijayraghavan, S., Wang, M., Birnbaum, S.G., Williams, G.V., and Arnsten, A.F. 2007. Inverted-U dopamine D1 receptor actions on prefrontal neurons engaged in working memory. *Nat. Neurosci.* 10: 576–584.

Wang, S.S., Denk, W., and Hauser, M. 2000. Coincidence detection in single dendritic spines mediated by calcium release. *Nat. Neurosci.* 3: 1266–1273.

Wang, M., Ramos, B.P., Paspalas, C.D., Shu, Y., Simen, A., Duque, A., Vijayraghavan, S., Brennan, A., Dudley, A., Nou, E., et al. 2007. α2A-adrenoceptors strengthen working memory networks by inhibiting cAMP-HCN channel signaling in prefrontal cortex. *Cell* 129: 397–410.

Weible, A.P., McEchron, M.D., and Disterhoft, J.F. 2000. Cortical involvement in acquisition and extinction of trace eyeblink conditioning. *Behav. Neurosci.* 114: 1058–1067.

Weible, A.P., Weiss, C., and Disterhoft, J.F. 2003. Activity profiles of single neurons in caudal anterior cingulate cortex during trace eyeblink conditioning in the rabbit. *J. Neurophysiol.* 90: 699–612.

Weible, A.P., Weiss, C., and Disterhoft, J.F. 2007. Connections of the caudal anterior cingulate cortex in rabbit: Neural circuitry participating in the acquisition of trace eyelid conditioning. *Neuroscience* 148: 288–302.

Weiss, C. and Disterhoft, J.F. 1996. Eyeblink conditioning, motor control, and the analysis of limbic-cerebellar interactions. *Behav. Brain Sci.* 19: 479–504.

Williams, G.V. and Goldman-Rakic, P.S. 1995. Modulation of memory fields by dopamine D1 receptors in prefrontal cortex. *Nature* 376: 572–575.

Woodruff-Pak, D.S., Lavond, D.G., and Thompson, R.F. 1985. Trace conditioning: abolished by cerebellar nuclear lesions but not lateral cerebellar cortex aspirations. *Brain Res.* 348: 249–260.

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