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Nicholas A. Lusk, Warren H. Meck, Henry H. Yin

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Mediodorsal Thalamus Contributes to the Timing of Instrumental Actions

Nicholas Lusk,¹ Warren H. Meck,¹ and Henry H. Yin^{1,2}

¹Department of Psychology and Neuroscience, Duke University, Durham, North Carolina 27708, and ²Department of Neurobiology, Duke University School of Medicine, Durham, North Carolina 27708

The perception of time is critical to adaptive behavior. While prefrontal cortex and basal ganglia have been implicated in interval timing in the seconds to minutes range, little is known about the role of the mediodorsal thalamus (MD), which is a key component of the limbic cortico–basal ganglia–thalamocortical loop. In this study, we tested the role of the MD in timing, using an operant temporal production task in male mice. In this task, the expected timing of available rewards is indicated by lever pressing. Inactivation of the MD with muscimol produced rightward shifts in peak pressing on probe trials as well as increases in peak spread, thus significantly altering both temporal accuracy and precision. Optogenetic inhibition of glutamatergic projection neurons in the MD also resulted in similar changes in timing. The observed effects were found to be independent of significant changes in movement. Our findings suggest that the MD is a critical component of the neural circuit for interval timing, without playing a direct role in regulating ongoing performance.

Key words: instrumental action; interval timing; mediodorsal thalamus; operant conditioning; optogenetics; prefrontal cortex

Significance Statement

The mediodorsal nucleus (MD) of the thalamus is strongly connected with the prefrontal cortex and basal ganglia, areas which have been implicated in interval timing. Previous work has shown that the MD contributes to working memory and learning of action–outcome contingencies, but its role in behavioral timing is poorly understood. Using an operant temporal production task, we showed that inactivation of the MD significantly impaired timing behavior.

Introduction

The ability to track the passage of time is ubiquitous across the animal kingdom and vital for survival (Buhusi and Meck, 2005). Neural correlates of timing have been found across the cortex as well as the basal ganglia (Meck, 2006; Kim et al., 2013; Yin, 2014b; Finnerty et al., 2015; Gouvêa et al., 2015; Bakhurin et al., 2017; Tiganj et al., 2017). Recent work has focused on the role of the prefrontal cortex (PFC) in timing (Jones et al., 2004; Buhusi et al., 2018; Kim and Narayanan, 2019). In rodents, the PFC is defined largely by the existence of strong reciprocal connections with the mediodorsal nucleus (MD) of the thalamus (Heidbreder and Groenewegen, 2003; Jones, 2007). The MD not only receives strong projections from the basal ganglia, but also sends

projections that drive feedforward inhibition within the PFC (Delevich et al., 2015). As a key thalamic component in the limbic cortico–basal ganglia–thalamocortical loop, the MD is in a position to contribute to interval timing, yet its specific functional contribution remains unclear.

Because of its strong connections with the PFC, which is often associated with working memory, considerable research has been devoted to elucidating the role of MD in working memory (Isseroff et al., 1982; Stokes and Best, 1990; Hunt and Aggleton, 1991; Watanabe and Funahashi, 2012). Based on this work, there is consensus that selective lesions of the MD do not impair working memory in rodents (Neave et al., 1993; Hunt and Aggleton, 1998a), though recent optogenetic work has suggested that PFC–MD interactions are important for working memory (Bolkan et al., 2017; Ferguson and Gao, 2018).

Previous work has also implicated the MD in the acquisition and selection of goal-directed actions controlled by outcome value and action–outcome contingencies (Corbit et al., 2003; Ostlund and Balleine, 2008; Yu et al., 2012; Parnaudeau et al., 2015; Alcaraz et al., 2018). In addition, MD lesions impair acquisition and expression on a temporal differentiation task, in which animals learn to produce lever presses with specific durations to earn rewards (Yu et al., 2010). MD appears to be required when animals have to adjust the timing of their actions, consistent with

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Correspondence should be addressed to Henry H. Yin at hy43@duke.edu.

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the claim that the MD is critical for behavioral flexibility (Hunt and Aggleton, 1998b).

In this study, we use pharmacological and optogenetic techniques in mice to examine the role of the MD in interval timing. We first inhibited the MD during a temporal production task, using local infusions of muscimol, a GABA_A receptor agonist. MD inhibition reduced temporal precision, as indicated by peak responding on unrewarded probe trials. Optogenetic inhibition of glutamatergic projection neurons in the MD also produced similar effects on timing, resulting in rightward shifts in peak pressing when stimulation was delivered at trial onset, and lengthened a bout of lever pressing when stimulation was delivered during lever pressing.

Materials and Methods

Viral constructs

AAV5-EF1 α -DIO-eNpHR3.0-eYFP (halorhodopsin for optogenetic inhibition) and AAV5-EF1 α -DIO-eYFP (control viral vector with no opsin) were obtained from the Duke Viral Vector Core (Duke University, Durham, NC).

Animals

Data for muscimol experiments ($n = 10$) was collected from adult wild-type mice (C57BL/6). Optogenetic experiments used adult *Vglut2-IRES-Cre* mice ($n = 11$). Male mice were used in experiments. Mice were maintained on a 12 h light/dark cycle. During training of 30 s peak interval procedure, animals were food deprived, receiving 2.0–2.5 g of rodent chow in addition to food pellets during testing. During food deprivation, the weights of the animals were monitored daily to ensure they remained at >85% of their *ad libitum* weights. All experimental procedures were approved by the Duke University Institutional Animal Care and Use Committee.

Surgery

Mice were anesthetized with 1.0–2.0% isoflurane mixed with 1.0 L/min oxygen for surgical procedures and placed into a stereotaxic frame (David Kopf Instruments). For pharmacological experiments, adult C57BL/6 mice ($n = 10$) were implanted, bilaterally, with 7 mm, 26 gauge guide cannulas over the MD [measured from the skull surface at 20° from vertical: anteroposterior (AP), –1.35 mm; ML, \pm 1.48 mm; DV, –3.0 mm]. Guide cannula coordinates were chosen accounting for infusion cannulas extending out an extra 0.5 mm. Guide cannulas were secured in place with dental acrylic adhered to skull screws. For optogenetic experiments, adult *Vglut2-IRES-Cre* mice were randomly assigned to *Vglut2::eNpHR3.0* ($n = 6$) or *Vglut2::eYFP* ($n = 5$) groups. Virus was bilaterally microinjected into the MD (0.3 μ l in each hemisphere; measured from the skull surface at 20° from vertical: AP, –1.35 mm; ML, \pm 1.48 mm; DV, –3.3 mm). All measurements are relative to bregma. Mice were bilaterally implanted with custom-made fiber optics aimed directly above the MD (AP, –1.35 mm relative to bregma; ML, \pm 1.48 mm. DV: –3.1 mm from skull surface at 20° from vertical). Fibers were secured in place with dental acrylic adhered to skull screws. Mice were singly housed during recovery for at least 2 weeks before training began.

Operant training

Lever-press training consisted of the extension of a single lever, either to the right or left of a food port. A food pellet reward (20 mg Dustless Precision Pellet, Bio-Serv) was delivered for each lever press. Every fifth

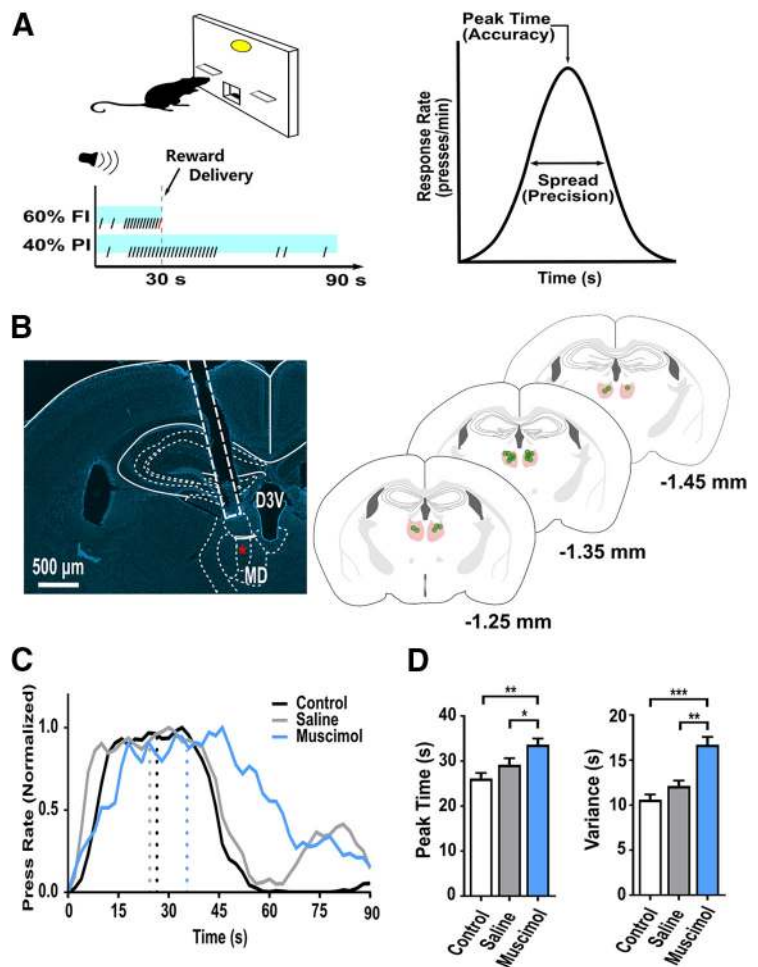


Figure 1. Pharmacological inhibition of MD disrupts interval timing behavior. **A**, Schematic of behavioral task and idealized behavioral output. **B**, Coronal section displaying cannula placement. Infusion cannula (red asterisk) was located 0.5 mm below the guide cannula (white box). **C**, Representative performance of a mouse under control (black), saline (gray), and muscimol (blue) conditions. Dotted lines indicate peak times defined by Gaussian + Ramp fit. **D**, Muscimol has effects on timing performance, from left to right: peak time, variance across, and elevation ratios. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$.

press led to an alternation of which lever was extended. Sessions ended after either successfully receiving 40 rewards or 30 min. Training was considered complete after an animal received 40 rewards before the 30 min time limit. The median number of training sessions it took for the animals to collect 40 rewards within 30 min was 5 (two to eight sessions).

Fixed interval (FI) training sessions began with the extension of a single lever to either the left or right of the food port, counterbalanced across mice, as well as the illumination of a house light. Individual trials were demarcated by the presence of an auditory cue (white noise; 68 dB). An animal was rewarded for the first lever press occurring after the cue had been on for 30 s. At this time, a food reward was delivered, the cue was turned off, and the session entered a variable intertrial interval ranging from 90 to 180 s randomly selected from a uniform distribution of values in 5 s increments. If there was no press within 8 s of the 30 s criterion, the trial would end as stated above, but without a reward. There was no penalty for presses occurring before 30 s. The development of a “scalped” profile for session-averaged lever pressing indicated the acquisition of timing behavior. Each session lasted 2 h. The median number of sessions it took to complete the FI training phase was 10 (7–27 sessions).

Peak interval (PI) training sessions consisted of the following two trial types: the aforementioned FI trials, and unrewarded probe trials. During probe trials, the house light was turned on for a minimum of 3× the longer target duration (90 s) plus an additional random amount of

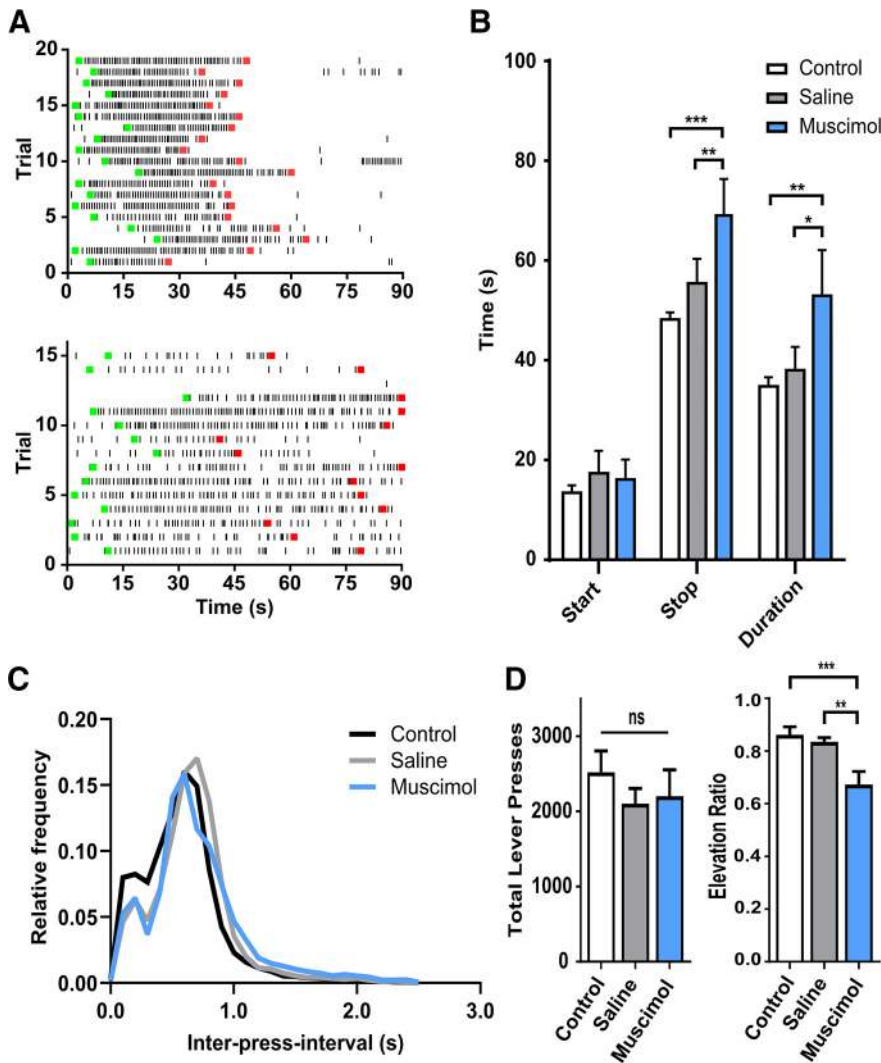


Figure 2. Single-trial analysis demonstrates differences in pressing profiles. **A**, Representative raster plot of individual trials during control (top) and muscimol (bottom) conditions from an individual mouse. Each black dash represents a single lever press. Green and red squares indicate the start and stop times, respectively (see Materials and Methods). **B**, Average start time, stop time, and duration of the pressing bout for mice across conditions. **C**, Histogram of IPIs across conditions. **D**, Total lever presses (left) and elevation ratio (right) during a session. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, ns, not significant.

time with a mean of 20 s and a Gaussian distribution. Each trial type was randomly selected, with a 40% chance of being a probe trial. Each session lasted 150 min. Animals vary in how long it took for a clear peak response function to emerge on the probe trials. The median number of sessions for the PI phase was 12 (6–21 sessions) before any neural manipulations were performed.

Local muscimol infusion

Mice received local bilateral infusions of either 0.9% saline or muscimol (0.01 mg/ml) solution. All infusions contained a total volume of 400 nl (i.e., 200 nl/hemisphere) infused at a rate of 0.05 ml/h. Infusion cannulas were left in place for 10 min postinfusion. After the removal of cannulas, animals were again given 10 min before starting any training. The condition order was counterbalanced across animals. Control sessions consisted of the insertion of injector cannula, but without infusing any solution. The injectors were left in place for 10 min, and animals were given 10 min after removal before starting any training.

Optogenetic stimulation

Optical stimulation began after an animal had progressed to the PI procedure and developed a distinctive peak in pressing activity centered at

~30 s. Mice were connected to a 589 nm DPSS laser (Shanghai Laser) via fiber optic cables (Doric; core = 200 μm ; numerical aperture = 0.22) and were placed inside the testing chamber (Rossi et al., 2012, 2015). A rotating optical commutator (Doric) divided the beam (50:50) permitting bilateral stimulation. Stimulation (5–8 mW; constant; 15, 10, or 5 s duration) was delivered at the following three timepoints: cue onset ($t = 0$); 15 s after cue onset ($t = 15$); or 60 s after cue onset ($t = 60$). The first two conditions occurred on both FI and probe trials, while the final condition only occurred on probe trials.

Histology

After testing was completed, mice were deeply anesthetized with isoflurane and perfused with 0.1 M PBS followed by 4% paraformaldehyde. Brains were postfixed in 4% paraformaldehyde for 24 h followed by 30% sucrose solution. After sinking in the sucrose solution, brains were sliced coronally at 60 μm using a Leica CM1850 Cryostat. For cannula localization, sections were mounted and immediately coverslipped with Fluoromount G with DAPI medium (catalog #17984–24, Electron Microscopy Sciences). To confirm proper fiber placement, coronal sections were compared with those from a mouse brain atlas (Paxinos and Franklin, 2003).

Experimental design and statistical analysis

Experimental procedures were controlled by a MED Associates interface using MED-PC software system. Lever presses were recorded in real time with 10 ms resolution. Session-averaged data were analyzed using Python scripts. All statistical analysis was performed in Prism 8.

Lever-pressing data for each trial was placed into 2 s bins and collapsed across all trials. The cumulative pressing data were smoothed using a Savitzky–Golay filter and then fit using a Gaussian curve with the addition of a linear ramp accommodating the right-tail skew, resulting from the temporal asymmetry of expected reward time relative to length of probe trials. Fits were used to obtain peak times (accuracy) and peak spread (precision), as previously described (Cheng and Meck, 2007). For optogenetic experiments, fits were calculated by combining data from two sessions to provide a sufficient number of trials for single-trial analysis, because only about half of the trials in each session were stimulation trials.

Single-trial analysis of lever-pressing data was performed by first putting the data in 1 s bins (Church et al., 1994). As the probe trials were of variable length with a minimum duration of 90 s, only the first 90 bins were used for analysis. The press rate data were padded on both sides and smoothed using a median filter, followed by a Savitzky–Golay filter. The smoothed data were log normalized, and then change points were detected using Python ruptures library with a Pelt search method and “L2” cost function. To account for differences in press rate between animals, the penalty and median filter length values were allowed to vary.

Results

Pharmacological inhibition of MD

Mice ($n = 10$ males) were implanted with guild cannulas before being trained on a 30 s peak interval procedure (Fig. 1A). Following experimentation, localization of cannula placement was verified for all animals (Fig. 1B). After progressing to the

final stage of training (see Materials and Methods), animals were assigned an infusion schedule, counterbalanced across subjects. Session-averaged response rates during nonrewarded probe trials were normalized in amplitude and fit using a Gaussian + Ramp function. Control animals exhibited peak times at 26.33 ± 1.32 s for the 30 s PI procedure, which was lower than the 30 s criteria, but in accord with previous work in mice using the same task (Buhusi et al., 2009). The peak functions revealed differences between muscimol and control conditions (Fig. 1C). A repeated-measures one-way ANOVA found significant effect of treatment on peak time ($F_{(1.95,17.58)} = 10.65$, $p < 0.001$) and variance ($F_{(1.88,16.90)} = 20.25$, $p < 0.001$). A Tukey's *post hoc* test revealed significant increases in the muscimol peak time (33.46 ± 1.18 s) and variance (15.86 ± 0.92 s) relative to control (26.33 ± 1.32 s, $p = 0.002$; 10.37 ± 0.34 s, $p < 0.001$) as well as saline (29.19 ± 1.39 s, $p = 0.05$; 12.07 ± 0.59 s, $p = 0.01$). No significant differences were found between control and saline conditions for either peak time ($p = 0.25$) or variance ($p = 0.14$).

To shed light on behavioral patterns that give rise to the session-averaged data shown in Figure 1, we conducted a single-trial analysis. For each trial, the start time, stop time, and duration of lever pressing were calculated (Fig. 2A). A one-way repeated-measures ANOVA found a significant treatment effect for the stop times ($F_{(1.78,16.02)} = 24.95$, $p < 0.001$) and duration of pressing ($F_{(1.93,17.41)} = 14.14$, $p < 0.001$), but not for the start times ($F_{(1.92,17.27)} = 1.80$, $p = 0.20$).

A Tukey's *post hoc* test revealed significant increases in the stop time ($p < 0.001$; $p = 0.006$) as well as duration of pressing ($p = 0.002$; $p = 0.022$) for the muscimol condition compared with control and saline conditions, respectively (Fig. 2B). The distribution of interpress intervals (IPIs) shows qualitatively similar profiles among muscimol (0.69 ± 0.004 s), saline (0.65 ± 0.003 s), and control (0.57 ± 0.003 s) conditions (Fig. 2C). Thus, the pattern of pressing is not affected by muscimol, suggesting that the observed effects are unlikely to be the result of changes in general motor activity.

In addition, no significant difference in total lever presses ($F_{(1.98,17.78)} = 0.77$, $p = 0.48$) or in press rate ($F_{(1.50,13.54)} = 3.31$, $p = 0.078$) was found, indicating that increases in response variance could not be solely explained by differences in the level of motor output in

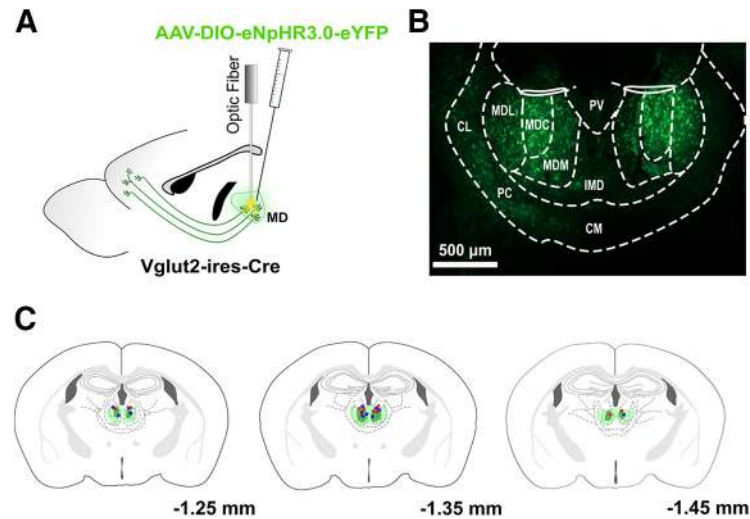


Figure 3. Optogenetic experimental design and histology. **A**, Schematic of viral infection and fiber optic implantation. **B**, Histology demonstrating viral expression and fiber placement in MD in coronal section. **C**, Extent of viral spread indicated by the maximum (light green) and minimum (dark green) expression volume, as well as fiber placement in experimental (blue dots) and control (red dots) mice.

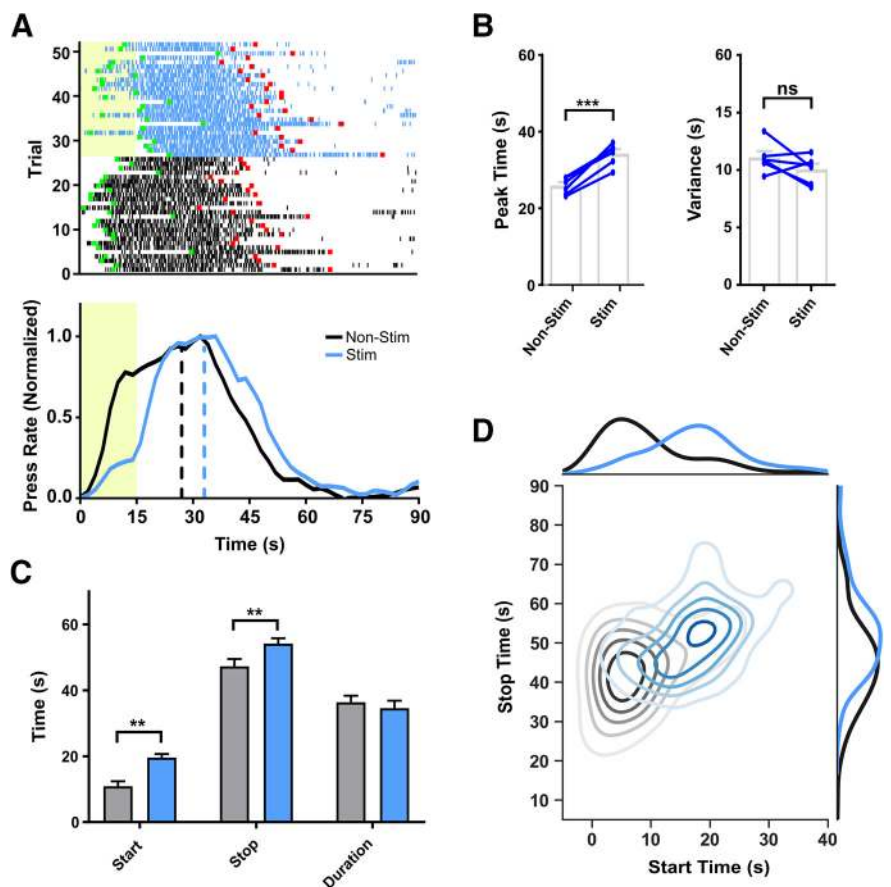


Figure 4. Optical inhibition of MD at cue onset delays the initiation of lever pressing. **A**, Example of single-trial (top) and averaged (bottom) pressing data over two sessions with 15 s of stimulation at the onset of the auditory cue (yellow rectangle). Each tick represents a single lever press. Green and red squares indicate the start and stop times, respectively. Dashed lines indicate the peak time of Gaussian + Ramp fit. **B**, Stimulation delays the peak time (left) but not the variance (right) of peak fits. Group average (gray) and individual (blue) data are shown. **C**, Stimulation significantly affects the average start time and stop time, but not press bout duration. **D**, Distributions of start and stop times for all trials. $**p < 0.01$, $***p < 0.001$, ns, not significant.

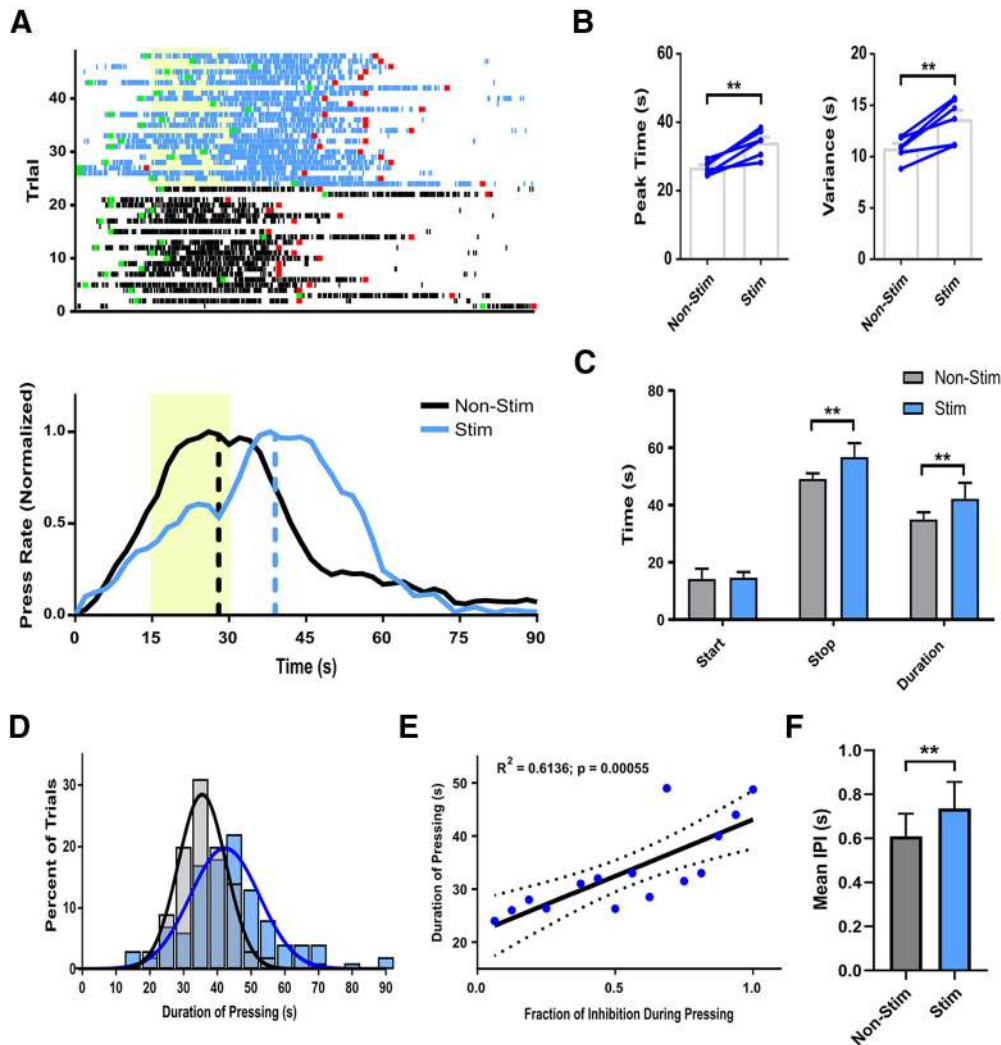


Figure 5. Optical inhibition of MD during pressing increases the length of the pressing bout. **A**, Example of single-trial (top) and averaged (bottom) pressing data over two sessions with 15 s of stimulation at the onset of auditory cue (yellow rectangle). Each tick represents a single lever press. Green and red squares indicate the start and stop times, respectively. Dashed lines indicate the peak time of Gaussian + Ramp fit. **B**, Stimulation delays the peak time (left) and variance (right) of peak fits. Group average (gray) and individual (blue) data are shown. **C**, Stimulation significantly affects the average start time and stop time, but not the press bout duration. **D**, Histogram of press durations for stimulation (Stim) and nonstimulation (Non-Stim) trials. **E**, Linear regression of press duration relative to the percentage of the total stimulation time received. **F**, Average interpress interval for each animal ($n = 6$) during the high state when the laser was on (Stim) and when the laser was off (Non-Stim). $**p < 0.01$.

muscimol animals (Fig. 2D). There was also a significant main effect of treatment on elevation ratio (Buhusi et al., 2009), which measures the number of presses during the trial relative to the total number of presses ($F_{(1.46, 13.12)} = 21.93$, $p < 0.001$) and muscimol condition (0.70 ± 0.035), which was significantly lower than both control (0.87 ± 0.022 , $p < 0.001$) and saline (0.83 ± 0.015 , $p = 0.007$) conditions (Fig. 2D). Therefore, while the muscimol doses given in this study did not significantly change overall lever pressing, it did alter sensitivity to the discriminative stimulus that indicates when presses can be rewarded.

Optical inhibition of MD

While muscimol inhibition of the MD demonstrated its importance to proper timing behavior, this manipulation lacks temporal precision. A given trial can be divided into the following three phases: initial “low-state” with little pressing followed a pressing “high-state” and concluded with another “low-state” (Church et al., 1994). To investigate the contributions of MD to behavior at different time points during the probe trial, we used optogenetics to inactivate the MD (Fig. 3). We used a Cre-dependent halorhodopsin

(DIO-eNpHR3.0) in vGlut2-Cre mice to selectively inhibit vGluT2⁺ projection neurons in the MD (Zhang et al., 2007). This manipulation provides more spatially and temporally selective inactivation of the MD. Although most MD cells are assumed to be glutamatergic, it is not possible to confirm that muscimol inactivation only affected glutamatergic neurons. By using the vGlut2-Cre driver line, optical inhibition can be limited to the glutamatergic projection neurons. To inactivate the MD at specific time points during the trial, we selected three timepoints for inhibition (0, 15, and 60 s). Each time point occurs primarily during one of the three phases. Inhibition was constant over 15 s, as a similar stimulation parameter was previously shown to effectively disrupt prefrontal-dependent processes such as working memory maintenance (Bolkan et al., 2017).

Performance on noninhibition trials was consistent with previous work. Peak time (25.53 ± 0.719 s) was significantly < 30 s, but was not significantly different from control data from muscimol experiments ($t_{(14)} = 0.44$, $p = 0.67$), nor was the peak variance ($t_{(14)} = 0.84$, $p = 0.41$). Therefore, despite having MD inhibition on 50% of the reward trials, there is likely no residual influence

on nonlaser peak trials. When optical inhibition was concurrent with the onset of the trial, mice showed a systematic delay in the onset of lever pressing, evidenced by a rightward shift in peak time (Fig. 4A). However, the rightward shift in peak times ($t_{(5)} = 9.05$, $p < 0.001$) was not accompanied by a significant difference in temporal variance ($t_{(5)} = 1.31$, $p = 0.25$; Fig. 4B). Single-trial analysis using a multiple-comparison t test with a Holm–Sidak correction demonstrated the rightward shift to be a product of delays in the start (planned comparison: $t_{(30)} = 3.14$, $p = 0.011$) and stop ($t_{(30)} = 2.69$, $p = 0.023$) of the pressing high state with no changes in the duration of pressing ($t_{(30)} = 0.45$, $p = 0.65$; Fig. 4C). A two-dimensional density plot shows qualitative similarities in the distribution of start and stop times (Fig. 4D). Despite having significantly different means, an F test for equality of variance found no significant differences between the two conditions for start times ($F_{(1,26)} = 2.69$, $p = 0.11$) or stop times ($F_{(1,32)} = 1.33$, $p = 0.26$).

The rightward shift in peak time during optical inhibition is seemingly at odds with the findings from the muscimol experiment, which found no changes in the start time for lever pressing (Fig. 2B). As mice under the muscimol condition pressed more during the intertrial interval (ITI), one possible explanation is that mice may already be engaged in pressing at the start of a trial. This is confirmed by analysis of lever pressing, including 60 s (the shortest ITI duration) before trial onset: under muscimol, more trials had start times that either preceded or coincided with the onset of the trial cue compared with the saline condition ($t_{(9)} = 2.32$, $p = 0.046$).

Next, we started optogenetic inhibition 15 s into the trial. Average start times during nonlaser trials for the $t = 0$ s condition (11.23 ± 1.001 s) indicated this to be a likely time for mice to be in the early stages of their pressing. As with optical inhibition at onset, animals produce rightward shifts in peak pressing ($t_{(5)} = 5.35$, $p = 0.0031$), but this was accompanied by a significant increase in peak variance ($t_{(5)} = 4.79$, $p = 0.0049$; Fig. 5A,B). Single-trial analysis was again performed to evaluate the contributing factors to differences found in session-averaged data between conditions (Fig. 5C). Contrary to the effects seen with inhibition at trial onset, inhibition at $t = 15$ s produced significant delays in pressing stop times ($t_{(30)} = 3.60$, $p = 0.003$) as well as increases in the duration of the pressing high state ($t_{(30)} = 3.38$, $p = 0.004$) with no significant differences in start times ($t_{(15)} = 0.21$, $p = 0.838$). In addition to the increases in the mean length of pressing, how inhibition of the MD affected the variability of press duration was investigated through the assessment of Gaussian fits of press duration histograms (Fig. 5D). An extra sum-of-squares F test found the variance of the best fits of the inhibition condition to be significantly greater than that of noninhibition trials ($F_{(1,32)} = 15.14$, $p = 0.0005$). Therefore, the inhibition of the MD does not only lead to increased duration of pressing, but also to increased variability in the length of a pressing bout.

As the start of the lever pressing bout can vary from trial to trial, a linear regression was performed between the bout duration and the total amount of inhibition as a fraction of the possible 15 s (Fig. 5E). When only part of the inhibition occurred

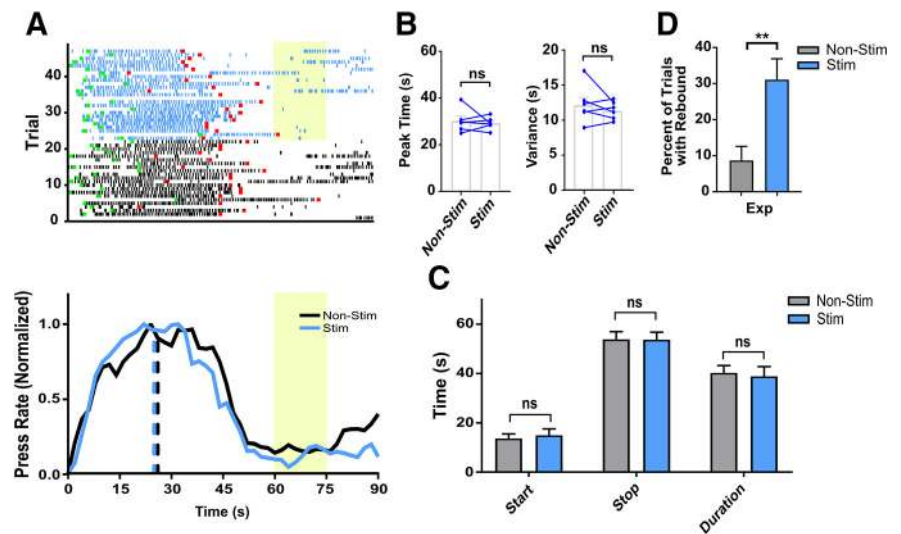


Figure 6. Optical inhibition following pressing increases the probability of reengagement. **A**, Example of single-trial (top) and averaged (bottom) pressing data over two sessions with 15 s stimulation at $t = 60$ s of auditory cue (yellow rectangle). Each tick represents a single lever press. Green and red squares indicate the start and stop times, respectively. Dashed lines indicate the peak time of Gaussian + Ramp fit. **B**, Stimulation has no effect on peak time (left) or variance (right) of peak fits. Group average (gray) and individual (blue) data shown. **C**, Stimulation has no significant effect on start time and stop time, or press bout duration. **D**, Percentage of trials with reengagement (rebound). ** $p < 0.01$, ns, not significant.

during pressing, no distinction was made between whether inhibition occurred at the beginning or the end of the pressing. The fraction of inhibition showed a strong linear correlation with the duration of the lever-pressing bout ($R^2 = 0.61$, $p = 0.00055$). To test whether optical inhibition reduces the rate of pressing, the IPI within the high state during the time of optical inhibition was compared with the IPI within the high state when optical inhibition was not occurring (Fig. 5F). Surprisingly, a paired t test found that during optical inhibition mice pressed more rapidly, evidenced by a significantly shorter IPI ($t_{(5)} = 4.05$, $p = 0.0098$). While this finding demonstrates a degree of hyperactivity, it rules out the possibility that a slowing of motor output could explain the timing effects.

The final time point for optical inhibition, $t = 60$ s, occurs after the animal has finished pressing. As expected, inhibition at this point has no significant effects on performance (session averaged, Fig. 6A,B; single trial, Fig. 6C). However, mice exhibited a significant increase in rebound pressing during inhibition trials (Fig. 6D; $t_{(5)} = 3.30$, $p = 0.03$). This is unlikely to be explained by confounding factors such as visually perceiving light from the laser as enhanced yellow fluorescent protein (eYFP)-control mice did not exhibit any changes in rebound pressing ($t_{(5)} = 0.03$, $p = 0.49$).

To address potential confounds related to light delivery, AAV5-EF1 α -DIO-eYFP controls ($n = 5$) were tested on the 30 s PI procedure using identical parameters. A paired t test on session-averaged data showed no difference in peak times ($p = 0.24$, $p = 0.20$, $p = 0.090$) or variance ($p = 0.43$, $p = 0.091$, $p = 0.84$) for control mice at stimulation times $t = 0$, $t = 15$, and $t = 60$ s, respectively (Fig. 7A–C). Additionally, single-trial analysis at all three stimulation timepoints was conducted to confirm that averaging effects did not mask differences at a lower level of analysis. There were no differences in start ($p = 0.74$, $p = 0.89$, $p = 0.57$), stop ($p = 0.94$, $p = 0.80$, $p = 0.64$), or duration ($p = 0.75$, $p = 0.72$, $p = 0.84$), respectively (Fig. 7D).

Last, we investigated how the length of optical inhibition modulates the timing behavior at all three time points across various lengths of inhibition (5, 10, and 15 s). For optical inhibition

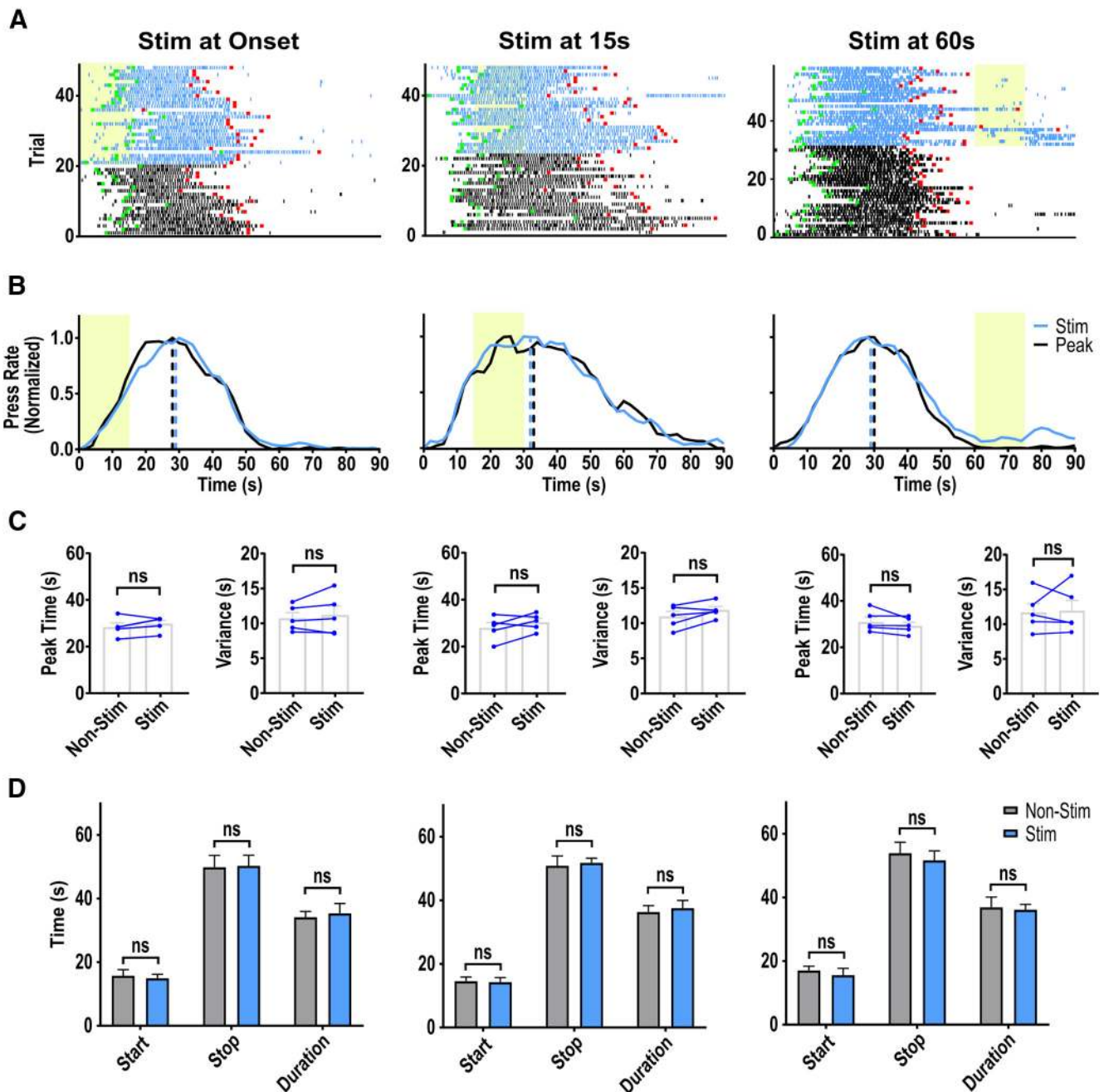


Figure 7. Photograph stimulation did not change behavior in eYFP controls. **A**, Raster plots for eYFP control mice at timepoints, from left to right, $t = 0$ s, $t = 15$ s, and $t = 60$ s. **B**, Session-averaged data for eYFP control animals for stimulation (Stim; blue) and nonstimulation (Non-Stim; black) trials at timepoints, from left to right, $t = 0$ s, $t = 15$ s, and $t = 60$ s (yellow). Peak times for Stim (blue dash) and Non-Stim (black dash) defined by Gaussian + Ramp fits. **C**, Stimulation does not modify the peak time (left) or variance (right) of peak fits. Group average (gray) and individual (blue) data are shown. **D**, Stimulation has no significant effect on the average start time and stop time, or the press bout duration. ns, not significant.

at trial onset ($t = 0$ s), a repeated-measures ANOVA found a significant main effect of stimulation length ($F_{(2,8)} = 9.85$, $p = 0.007$). *Post hoc* tests found a significant difference between 15 s of inhibition and 5 s ($t_{(8)} = 4.29$, $p = 0.0079$) as well as 10 s ($t_{(8)} = 3.13$, $p = 0.042$; Fig. 8A). Moreover, inhibition at $t = 15$ s demonstrated a significant treatment effect ($F_{(2,8)} = 30.21$, $p < 0.001$) with longer durations of MD inhibition leading to increases in variance. A *post hoc* test found 15 s of inhibition to produce significantly greater increases in variance than 5 s ($t_{(8)} = 7.39$, $p < 0.001$) or 10 s of inhibition ($t_{(8)} = 5.77$, $p = 0.001$; Fig. 8B). Moreover, this finding aligns with single-trial data demonstrating that increases in the length of the high-state correlate with the

length of optical inhibition that occurs over the pressing phase. A two-way repeated-measures ANOVA with stimulation duration (5, 10, 15 s) and stimulation condition (stimulation, nonstimulation) as factors revealed a significant main effect of duration on the percentage of rebound trials ($F_{(2,8)} = 5.20$, $p = 0.036$), but no main effect of stimulation condition or interaction between these factors. *Post hoc* tests found that inhibition lengths of 15 s ($t_{(8)} = 5.53$, $p = 0.002$) and 10 s ($t_{(8)} = 4.44$, $p = 0.007$) result in a higher proportion of rebound trials than noninhibition trials (Fig. 8C). Additionally, the 5 s inhibition condition did not significantly increase rebound pressing ($t_{(8)} = 1.56$, $p = 0.47$).

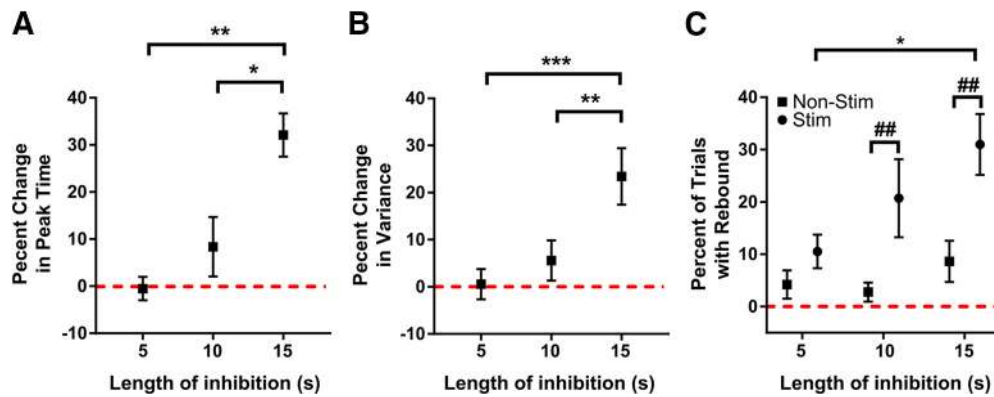


Figure 8. Effects of the optical inhibition scale with length of inhibition. **A**, Percentage increase in session peak time relative to the length of optical inhibition at $t = 0$ s. **B**, Percentage increase in session variance relative to the length of optical inhibition at $t = 15$ s. **C**, Percentage of trial with rebound pressing relative to the length of optical inhibition at $t = 60$ s. Within condition: * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$; across condition, ## $p < 0.01$.

Discussion

Using pharmacological and optogenetic manipulations, we provide evidence for the involvement of the MD in interval timing. We first trained mice on an FI task, in which rewards become available after 30 s. The internal estimate of time is indicated by the peak lever pressing on occasional probe trials with no reward. Here we show rightward shifts in the peak timing function in addition to increases in timing variance after MD inactivation. Such effects are not confounded by general changes in motor output, and therefore are likely due to changes in internal timing mechanisms.

Reversible pharmacological inhibition of the MD with muscimol modified timing performance, impairing both timing accuracy and precision, as indicated by peak time and spread of session-averaged performance on probe trials. In addition to timing behavior, mice exhibited a decrease in the ratio of presses during cued trials quantified through the elevation ratio (Figs. 1, 2). These findings resemble the effects of mPFC inactivation as muscimol inactivation of the prelimbic cortex also produced similar effects (Buhusi et al., 2018). Additionally, mixed effects on response rate were found depending on the dosage of muscimol, with low doses leading to increases in response rate. This is akin to the discrepancy found in the current work with the muscimol dosage used, producing no significant changes in response rate, but the optogenetic inhibition leading to increases in response rate. The strong reciprocal connectivity between the MD and prefrontal cortical areas and similarities in effects on timing from inactivation of these two regions suggests a critical role of cortico–thalamo–cortical interactions in interval timing.

To further elucidate MD contribution to various stages of timing behavior, we used optogenetics to inhibit MD output neurons at various time points. Inhibition of the MD was found to increase the duration of the lever pressing bout for the animal during a trial, resulting in delays in the onset of pressing (Fig. 4) and the offset of pressing (Fig. 5). The duration of inhibition during pressing strongly correlates with the increase in the length of the pressing bout. This suggests that the mouse may time how long to wait before starting to press and then time how long to press once initiated. That stimulation duration during pressing correlates with the length of the pressing bout suggests that each stage of the task is timed independently. That is, the animal will time when it should begin pressing and then when to stop, sequentially. This interpretation of timing was proposed previously (Killeen and Fettermann, 1988). However, this is in conflict with early analysis of individual trial studies that found support

for parallel processing theories, most notably the Scalar Expectancy Theory, according to which all behavioral output is generated in relation to a single timed duration (Gibbon, 1977). Previous work found that the covariance pattern between start and stop times of pressing bouts supported the use of a single temporal sample from memory with different decision thresholds for when to start and stop pressing (Church et al., 1994). However, the ability to maintain the necessary covariance pattern does not require parallel processing, as theories based on memory decay have demonstrated (Staddon and Higa, 1999). Our results favor a serial timing process in which animals time the length of each individual stage during a peak interval procedure.

Our findings also have implications for computational models of interval timing. Many timing models focus on short intervals in the subsecond range, but neural networks relying solely on cortical-based connectivity rules have trouble scaling to longer time intervals (>10 s; Laje and Buonomano, 2013). It has been suggested that shifts in the balance of excitation and inhibition (E/I) within thalamocortical networks allow for the emergence and stabilization of sequential activity patterns (Hardy and Buonomano, 2016). Recent work has suggested that the MD is important for maintaining E/I balance within the mPFC (Ferguson and Gao, 2018), by modulating feedforward inhibition via GABAergic interneurons in the cortex (Delevich et al., 2015). Adjusting E/I balance via cortico–thalamo–cortical interactions may improve the maintenance of working memory in these networks and allow the timing of longer durations.

Thus, we show for the first time the importance of the MD in timing long intervals. The durations used in the current work span 30 s, much longer than that used in previous work on the role of thalamus in timing (Wang et al., 2018). While shorter durations (i.e., <2 s) can be effectively timed in localized networks such as the cerebellum (Johansson et al., 2014), as the relevant time interval increases, the neural substrate of timing appears to require a larger network of brain regions, requiring synchronization, coordination, or integration of temporal information from distributed sources (Matell and Meck, 2004; Hass and Durstewitz, 2016; Petter et al., 2016). Substantial evidence supports the role of higher-order thalamic nuclei, such as the MD, in synchronizing neural activity in the cortex (Theyel et al., 2010; Saalmann, 2014). This role of the MD also explains why changes in timing behavior after MD inactivation resemble those found after inactivation of connected cortical regions (Buhusi et al., 2018).

In addition to interactions with the frontal cortex, the MD is also a major target of basal ganglia output nuclei, such as the substantia nigra pars reticulata (SNr; Gerfen et al., 1982; Haber et al., 1985; Kuroda and Price, 1991; Miyamoto and Jinnai, 1994). Extensive work has established the role of the basal ganglia circuit in interval timing (Coull et al., 2004; Matell and Meck, 2004; Gouvêa et al., 2015; Bakhurin et al., 2020). However, recent optogenetic work also has implicated the nigral output in both ongoing performance and the timing of future behavior. While the nigral output is known to play a key role in posture control and top-down regulation of movement parameters (Barter et al., 2014, 2015; Yin, 2017), activation of the GABAergic nigrothalamic projection not only reduces ongoing licking, but proportionally delays subsequently timed behavior (Toda et al., 2017). However, in the present study, MD inhibition does not appear to impair ongoing performance. It is possible that manipulations of the SNr can alter performance via direct projections to the tectum and brainstem pattern generators (Yin, 2014a; Rossi et al., 2016). However, collaterals of nigral efferent projections may send a copy of the ongoing motor commands to the MD. This is consistent with previous work in monkeys showing that the MD does not play a role in movement generation, but is critical for internal monitoring of movement sequences by signaling an efference copy or corollary discharge to frontal cortical regions (Sommer and Wurtz, 2002). Although projections from the SNr usually target more lateral regions of the MD than those targeted in the current work, it is possible that there is significant overlap in the target region and the region inactivated in our experiments, or that different regions in the MD may signal different types of efference copy signals corresponding to different types of actions. Future work targeting specific SNr–MD and MD–PFC pathways may shed light on how efference copy signals are computed and used in the timing of instrumental actions.

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