Temporal learning among prefrontal and striatal ensembles

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1 Abstract

2	Behavioral flexibility requires the prefrontal cortex and striatum. Here, we investigate neuronal
3	ensembles in the medial frontal cortex (MFC) and the dorsomedial striatum (DMS) during one
4	form of behavioral flexibility: learning a new temporal interval. We studied corticostriatal
5	neuronal activity as rodents trained to respond after a 12-second fixed interval (FI12) learned to
6	respond at a shorter 3-second fixed interval (FI3). On FI12 trials, we discovered time-related
7	ramping was reduced in the MFC but not in the DMS in two-interval vs. one-interval sessions.
8	We also found that more DMS neurons than MFC neurons exhibited differential interval-related
9	activity on the first day of two-interval performance. Finally, MFC and DMS ramping was
10	similar with successive days of two-interval performance but DMS temporal decoding increased
11	on FI3 trials. These data suggest that the MFC and DMS play distinct roles during temporal
12	learning and provide insight into corticostriatal circuits.

13 Introduction

14	Behavioral flexibility requires learning to adapt to uncertainty. Two forebrain structures
15	critical for flexibility are the prefrontal cortex and striatum (Fuster, 2008; Kehagia et al., 2010).
16	Prefrontal cortical neurons densely innervate the striatum (Gabbott et al., 2005; Wall et al., 2013)
17	and disruptions of either structure profoundly impact the learning of new goals, rules, and
18	strategies (Hart et al., 2018; Ragozzino, 2007). Dysfunctional corticostriatal circuits and
19	connectivity are implicated in a range of psychiatric and neurological disorders (Deutch, 1993;
20	Shepherd, 2013). However, the relative roles of prefrontal and striatal networks during
21	behavioral flexibility are unclear.
22	One task that provides an ideal window into behavioral flexibility is interval timing,
23	which requires participants to estimate an interval of several seconds via a motor response.
24	Across species, interval timing requires the prefrontal cortex and striatum (Coull et al., 2011;
25	Dallérac et al., 2017; Emmons et al., 2017, 2016; Matell and Meck, 2004; Merchant and de
26	Lafuente, 2014). Work from our group and others has shown that both prefrontal and striatal
27	neurons encode temporal information via 'time-related ramping' activity-or monotonic changes
28	in firing rate over a temporal interval (Bakhurin et al., 2017; Donnelly et al., 2015; Emmons et
29	al., 2017; Kim et al., 2018; Narayanan, 2016; Wang et al., 2018). Our past work suggested that
30	ramping activity in neurons of the medial frontal cortex (MFC) and the dorsomedial striatum
31	(DMS) is very similar, with ~40% of neurons in each area exhibiting such activity (Emmons et
32	al., 2017). We have also found that MFC inactivation attenuates DMS ramping (Emmons et al.,
33	2019, 2017) and that MFC stimulation is sufficient to increase DMS ramping (Emmons et al.,
34	2019). These data suggest that DMS ramping is closely linked to MFC ramping and suggest the
35	hypothesis that MFC and DMS ensembles respond similarly as animals learn new temporal

36	intervals. By contrast, recordings from primate lateral prefrontal cortex and caudate indicate that
37	striatal ensembles encode stimulus-response associations earlier than prefrontal ensembles,
38	leading to the hypothesis that prefrontal and striatal ensembles play differential roles during
39	learning (Antzoulatos and Miller, 2011; Histed et al., 2009; Pasupathy and Miller, 2005).
40	We tested these hypotheses by recording MFC and DMS activity in rodents as they
41	learned to respond to a new interval. Specifically, rodents previously trained to perform a 12-
42	second fixed-interval task learned a new version of the task that included two fixed-intervals—3
43	seconds and 12 seconds. We report three main results. First, time-related ramping activity in the
44	MFC decreased on 12-second interval trials during two-interval sessions compared to one-
45	interval sessions, whereas activity in the DMS on 12-second interval trials did not change.
46	Second, DMS neurons were more likely to have distinct firing patterns during the 12-second vs.
47	3-seconds interval than those of the MFC, but this interval-related activity normalized between
48	MFC and DMS over subsequent days of two-interval performance. Finally, MFC and DMS
49	ramping did not change consistently over subsequent two-interval sessions, but temporal
50	decoding by DMS ensembles improved for FI3 trials. These data suggest that the MFC and DMS
51	play distinct roles during temporal learning.

52 Methods

53 *Rodents*

All procedures were approved by the University of Iowa IACUC, and all methods were 54 performed in accordance with the relevant guidelines and regulations (protocol #7072039). 55 Seven male Long-Evans rats were trained on the 12-second fixed-interval timing task (FI12) 56 according to procedures described in detail previously (Emmons et al., 2017, 2016). In brief, the 57 rats were autoshaped to press a lever for water reward using a fixed-ratio task before being 58 59 trained on 12-second fixed-interval timing (FI12). Trials began with the presentation of a house light, and the first response made after 12 seconds resulted in the delivery of a water reward, a 60 61 concurrent click, and termination of the house light (Fig. 1A; video S1). Responses made before 62 the interval ended were unreinforced. Trials were separated by a randomly chosen 6-, 8-, 10-, or 12-second intertrial interval. After animals behaved consistently, the MFC and DMS were each 63 implanted with recording electrodes (Fig. 1B; see below). Animals were then acclimatized to the 64 recording procedures and recordings were made during behavior in the FI12 task (Day 0). The 65 following day, an additional 3-s interval (FI3) was added to the task and cued by a light distinct 66 from the one used to indicate FI12. Behavior and simultaneous neuronal activity in the MFC and 67 DMS were recorded over the following three days (Day 1, Day 2, and Day 3). Some data from 68 subsequent recording sessions in these rodents were included in prior manuscripts (Emmons et 69 70 al., 2017, 2016).

71

72 Surgical and histological procedures

73	Animals were anesthetized using ketamine (100 mg/kg) and xylazine (10 mg/kg), and a
74	surgical level of anesthesia was maintained using ketamine supplements (10 mg/kg).
75	Craniotomies were drilled above the left MFC and left DMS and four holes were drilled for skull
76	screws, which were connected to electrode recording arrays via a separate ground wire.
77	Microelectrode arrays were composed of 4x4 50-µm stainless steel wires (250 µm between wires
78	and rows; impedance measured <i>in vitro</i> at ~400 k Ω ; Plexon, Dallas, TX). These arrays were
79	positioned in the MFC (coordinates from bregma: AP +3.2, ML \pm 1.2, DV -3.6 @ 12° in the
80	lateral plane) and the DMS (coordinates from bregma: AP +0.0, ML \pm 4.2, DV -3.6 @ 12° in the
81	posterior or lateral plane) while recording neuronal activity to verify that implantation was in
82	correct brain area. The craniotomy was sealed with cyanoacrylate ('SloZap', Pacer Technologies,
83	Rancho Cucamonga, CA), and the reaction was accelerated by 'ZipKicker' (Pacer Technologies)
84	and methyl methacrylate (AM Systems, Port Angeles, WA). Rats recovered for one week before
85	being acclimatized to behavioral and recording procedures.
86	Following these experiments, the rats were anesthetized and sacrificed by injection with
87	100 mg/kg sodium pentobarbital and transcardially perfused with 4% formalin. Brains were post-
88	fixed in a solution of 4% formalin and 20% sucrose before being sectioned on a freezing
89	microtome. Brain slices were mounted on Superfrost Plus microscope slides and stained for cell
90	bodies using either DAPI or Cresyl violet. Histological reconstruction was completed using
91	postmortem analysis of electrode placement by slide-scanning fluorescent microscopy
92	(Olympus).

93

94 Neurophysiological recordings and neuronal analyses

95	Neuronal ensemble recordings were made using a multi-electrode recording system
96	(Plexon). In each animal, one electrode without single units was reserved for local referencing,
97	yielding 15 electrodes per animal. Offline Sorter (Plexon) was used to analyze the signals after
98	the experiments and to remove artifacts. Spike activity was analyzed for all cells that fired at
99	rates above 0.1 Hz. Principal component analysis (PCA) and waveform shape were used for
100	spike sorting. DMS neurons were classified as either medium spiny neurons (MSNs) or
101	interneurons based on peak-to-trough ratio and the spike half-peak width of spike waveforms
102	(Fig. 1C; Berke, 2011). Single units were defined as those 1) having a consistent waveform
103	shape, 2) being a separable cluster in PCA space, and 3) having a consistent refractory period of
104	at least 2 ms in interspike interval histograms.

105

106 *Statistics*

107 Basic analyses were performed via ANOVA, Wilcoxon rank-sum tests, and Cohen's D. As in our prior work, we quantified temporal control of action during fixed-interval performance 108 in two ways. We calculated the curvature of time-response histograms (Emmons et al., 2019; Fry 109 et al., 1960; Narayanan et al., 2012). Curvature values range between -1 and 1 and are calculated 110 111 from the cumulative response record by deviation from a straight line; 0 indicates a constant response rate throughout the interval. Curvature indices are resistant to differences in response 112 113 rate, smoothing, or binning. Second, we modeled each response using generalized linear mixedeffects models (GLMM; *fitglme.m* in MATLAB) where the outcome is response time, the 114 predictor variable was Day, and the random effect was animal. For two-interval trials, single-trial 115 116 analyses were used to find start times and coefficients of variation for FI3 and FI12 trials (Church et al., 1994). All data and statistical approaches were reviewed by the Biostatistics, 117

Epidemiology, and Research Design Core (BERD) at the Institute for Clinical and Translational
Sciences (ICTS) at the University of Iowa.

120 Analyses of neuronal activity and basic firing properties were carried out using 121 NeuroExplorer (Nex Technologies, Littleton, MA) and custom routines for MATLAB, as described in detail previously (Emmons et al., 2019, 2017; Parker et al., 2014). Peri-event rasters 122 123 and time-histograms were constructed around houselight and lever press. As in our past work, 124 neuronal modulations were quantified in two ways. First, we used principal component analysis 125 (PCA), a data-driven set of orthogonal basis vectors that captures patterns of activity in 126 multivariate neuronal ensembles. PCA was calculated from average peri-event time histograms computed from kernel-density estimates (ksdensity.m; bandwidth 0.5) and normalized using 127 128 zscore. As in our past work, we used absolute values of PC1 scores (indicated by |PC1|) compare ramping strength across areas and days (Emmons et al., 2017; Kim et al., 2017; Parker et al., 129 130 2017, 2014).

We also used *fitglme.m* to construct GLMMs to analyze neuronal modulations. For 131 neuron-by-neuron analysis, we used GLMMs where the outcome variable was the firing rate 132 (binned at 0.1 s) and the predictor was time in the interval (0 to 3 seconds on FI3, or 0 to 12 133 seconds on FI12), interval (FI3 or FI12), or responses. Neurons with a main effect of time were 134 considered 'ramping' neurons, neurons with a main effect of interval-type were considered 135 136 interval-modulated, and neurons with a main effect of response were considered responsemodulated. Neurons with time-related ramping were defined as those with a main-effect of time 137 138 in the interval via GLMMs where the outcome was firing rate binned at 0.1 s, and the predictor 139 was time in the interval. We also performed trial-by-trial GLMMs for all trials and neurons where the outcome was firing rate, and the predictors were area (MFC or DMS) or Day 0 or Day 140

1, and random effects were lever presses and neurons (Table 1). To examine interval-related 141 modulation for all trials and neurons, we used an equation where the outcome was firing rate, the 142 predictors were interval type (FI3 or FI12), area, or Day, and random effects were lever presses 143 and neurons (Table 2). For two-interval performance, we used GLMMs for FI3 trials (Table 3) 144 and FI12 trials (Table 4) on Days 1-3. Poisson distributions were used for all firing rate models. 145 146 We used a naïve Bayesian classifier to examine neuronal ensemble decoding, as we have 147 in our past work (Emmons et al., 2017; Kim et al., 2017). We calculated kernel density estimates (bandwidth: 1.2) of trial-by-trial firing rates from MFC and DMS neurons. To prevent edge 148 149 effects that might bias classifier performance, we included data from 6 seconds prior to trial start 150 and 6 seconds after interval end. We used leave-one-out cross-validation to predict objective time from firing rate within a trial. We evaluated classifier performance by computing the R^2 of 151 152 objective time and predicted time only for bins during the interval. With perfect classification,

the R^2 would approach 1. Classifier performance was compared to ensembles with time-shuffled

154 firing rates. For each area and interval, performance was compared via GLMMs of R^2 vs. each

155 day.

157 **Results**

158	We studied temporal learning in the MFC and DMS by introducing a new 3-s fixed
159	interval (FI3) to rats after they had been trained on a task with 12-second fixed-intervals (FI12;
160	Fig. 1A). We compared behavior on FI12 trials only in sessions with one interval ("Day 0") to
161	two-interval sessions in which FI12 trials were randomly intermixed with FI3 trials ("Day 1").
162	We quantified the temporal control of action via a 'curvature' index calculated from the
163	cumulative distribution of time-response histograms. We have used this index extensively in the
164	past (Emmons et al., 2019; Narayanan et al., 2012). On FI12 trials, the curvature index trended
165	towards being lower on Day 1 vs. Day 0 (Fig. 2A-B; Day 0 curvature: 0.29 ± 0.04 , mean \pm SEM;
166	Day 1 curvature: 0.17 ± 0.06 ; signrank $p = 0.08$; Cohen's d = 0.86). Response times were
167	significantly shorter on Day 1 vs. Day 0 (Fig. 2C-D; Day 0: 10.14 ± 0.09 vs. Day 1: 9.61 ± 0.10
168	seconds; main effect of Day: $F_{(1,3784)} = 12.28$, $p = 0.0005$; $R^2 = 0.003$). These results suggest that
169	the timing of FI12 responses was shifted earlier on FI12 trials on Day 1 when they were
170	intermixed with FI3 trials compared to Day 0, when FI12 trials were presented alone.
171	We recorded neuronal ensembles simultaneously in the MFC and DMS as animals well-
172	trained on FI12 trials on Day 0 learned to perform a two-interval task with FI3 and FI12 trials
173	randomly intermixed on Day 1. As in our past work, we found that neurons in both brain regions
174	exhibited time-dependent ramping, i.e. monotonic increases in firing across the interval
175	(Emmons et al., 2017; Fig. 3A-B). On FI12 trials 35 of 59 (59%) MFC neurons exhibited
176	ramping activity on Day 0, surprisingly, there were only 19 of 47 (40%) ramping neurons in
177	MFC on Day 1 (Fig. 3C; $X^2 = 3.74$, $p = 0.05$). In the DMS, 32 of 67 (48%) neurons ramped Day
178	0 and 29 of 58 (50%) ramped on Day 1, ($X^2 = 0.06$, $p = 0.80$).

179	To further compare ramping activity, we turned to principal component analysis (PCA)
180	as a data-driven approach to compare neuronal activity patterns (Chapin and Nicolelis, 1999;
181	Emmons et al., 2017; Narayanan and Laubach, 2009). Consistent with our prior work, we found
182	that principal component 1 (PC1) exhibited time-related ramping (Fig 3H; Emmons et al., 2017;
183	Kim et al., 2017; Parker et al., 2015, 2014; Zhang et al., 2019). For the MFC, the strength of
184	PC1 was lower on Day 1 than Day 0 (Fig. 3I; signrank $p = 0.03$; Cohen's d = 0.50), whereas
185	there was no consistent difference for the DMS (Fig 3I; signrank $p = 0.60$). Trial by-trial analysis
186	of firing rate on FI12 trials revealed that time-related ramping interacted with both the brain area
187	(MFC and DMS) as well one- vs. two-interval sessions (i.e., Day 0 vs. Day 1). There was a
188	three-way interaction between time-related ramping, brain area, and one vs. two intervals (Table
189	1). For FI12 trials, these data suggest that time-related ramping in the MFC was stronger on Day
190	0 compared to Day 1 when FI12 trials were intermixed with FI3 trials. Our results suggest that
191	neuronal ensembles in the MFC, but not in the DMS, are sensitive to the temporal context
192	(Jazayeri and Shadlen, 2010; Shi et al., 2013).

193 Next, we compared MFC and DMS activity on FI3 and FI12 trials. First, we analyzed 194 fixed-interval behavior using single-trial analysis, which was developed for peak-interval timing but can be useful to analyze start times during fixed-interval tasks (Church et al., 1994; Emmons 195 et al., 2019). On the first day of the shorter 3-second interval, single-trial analysis revealed that 196 animals had shorter start times on FI3 trials compared to FI12 trials (FI3: 3.18 ± 0.69 vs. FI12 197 7.06 ± 0.34 , signrank p = 0.03; Cohen's d = 2.9; single-trial analysis could not compute FI3 start 198 199 times from one animal, which was subsequently removed from this analysis). One indication that timing processes are scalar is that the coefficient of variation (CV-the ratio of standard 200 201 deviation of temporal estimates to the mean) is relatively constant at different intervals (Gibbon

et al., 1984; Rakitin et al., 1998). Accordingly, we found that during fixed-interval performance, 202 start time CVs were similar on FI3 and FI12 trials (FI3: 0.59+/-0.12 vs. FI12: 0.41+/-0.03 203 signrank p: 0.56). These data suggest that start times during fixed-interval timing exhibit scalar 204 properties (Gibbon et al., 1984). 205 Ramping neurons can have distinct slopes of firing rates vs. time on FI3 and FI12 trials 206 207 (Fig 4A). We ran GLMMs where firing-rate slope vs. time was the outcome variable, and FI3 vs. FI12 interval and Day were predictor variables; note that we were interested in the magnitude of 208 209 the slope and we focus on its absolute value, indicated by slope. Consistent with past work by 210 our group and others, ramping neuron |slopes| were consistently steeper on FI3 vs. FI12 trials for both the MFC (Fig. 4B; main effect of interval: $F_{(143)} = 4.52$, p = 0.04, $R^2 = 0.22$) and for the 211 DMS (main effect of interval: $F_{(214)} = 6.91$, p = 0.01; $R^2 = 0.40$ (Emmons et al., 2017; Mello et 212 al., 2015; Wang et al., 2018). There was no effect of Day or higher interactions for either MFC 213 or DMS. These data are consistent with influential drift-diffusion models of interval timing, 214 suggesting that drift rates increase with shorter intervals (Simen et al., 2011). 215 Next, we searched for neurons in which firing rates were a function of interval duration 216 (Fig 4C). Specifically, we used GLMMs to identify neurons with a main effect of interval on 217 firing rate (Fig. 4E; FI3 vs. FI12 trials). Interval-modulated neurons were more common in the 218 DMS than the MFC on Day 1 of two-interval performance (Fig 4D; MFC 10 of 47 vs. DMS: 28 219 of 58; $X^2 = 8.20$; p = 0.004). Interestingly, ~50% of interval-modulated neurons also had ramping 220 activity in MFC (5 of 10) and DMS (15 of 28). Of note, the number of interval-modulated 221 neurons was not different between MFC and DMS on Days 2 and 3 (Fig 4D). Consistent with 222 223 these analyses, GLMMs revealed a significant interaction between interval-modulation, brain area, and Days 1-3 (Table 2). 224

225	A comparison of behavior across the three days of two-interval performance revealed that
226	response times shortened for both FI3 and FI12 trials (Fig. 5A-B: GLMM FI3: $F_{(1741)} = 8.35$, $p =$
227	0.0002; $R^2 = 0.05$; FI12: $F_{(4796)} = 6.98$, $p = 0.0009$; $R^2 = 0.002$), but the curvature of time-
228	response histograms did not reliably change (FI3: $F_{(40)} = 0.03$, $p = 0.87$, $R^2 = 0.17$; FI12: $F_{(40)} = 0.03$
229	1.05, $p = 0.31$, $R^2 = 0.61$). As in prior work demonstrating temporal scaling in the MFC and
230	DMS, PCA revealed that the principal components for FI3 and FI12 trials were very similar (Fig.
231	5C-D; Pearson's <i>rho</i> correlation: PC1: = 0.99 <i>p</i> < 0.001; PC2: -0.95, <i>p</i> < 0.001; PC3: 0.95, <i>p</i> <
232	0.001; (Emmons et al., 2017; Mello et al., 2015; Wang et al., 2018). However, PC1 did not
233	change consistently over the three days of two-interval performance for MFC or DMS (Fig. 5E-
234	H; Table 3-4). Taken together, these data indicate that ramping-related patterns of activity of
235	corticostriatal ensembles did not consistently change as animals performed two-interval tasks on
236	Days 1-3.

We turned to decoding analyses based on machine learning to capture more complex 237 features of MFC and DMS ensembles (Fig 6; Emmons et al., 2017; Gouvea et al., 2015; Kim et 238 239 al., 2017). Specifically, we constructed neuron-dimensional arrays of smoothed trial-by-trial 240 firing rates over the interval binned at 0.1 seconds. We decoded time in the interval from ensemble firing rates using naïve Bayesian classifiers. Classifier performance was assessed by 241 computing the variance explained (R^2) of predicted vs. observed time. For all sessions, R^2 was 242 much less for time-shuffled ensembles—i.e., ensembles constructed from neuronal activity 243 shuffled in time (Fig. 6B-G; signrank $p = 4*10^{-47}$; Cohen's d = 1.57). We found that temporal 244 decoding had a main effect of Day only for the DMS Ensembles on FI3 trials (Fig. 6F&G; F₍₁₀₈₎ 245 = 6.07, p = 0.02, $R^2 = 0.06$). These results suggest that as the response times shortened with two-246 interval performance, temporal decoding improved only for DMS ensembles on FI3 trials. 247

248 Discussion

249	We found three key distinctions between MFC and DMS during temporal learning. First,
250	time-related ramping in the MFC decreased as animals that had been trained on a one-interval
251	task learned to respond to a second interval that was novel and shorter. Second, interval-
252	modulated neurons were more common in the DMS early in two-interval performance. Third,
253	time-related ramping in the MFC and DMS did not change with two-interval performance, but
254	temporal decoding improved only for the DMS on FI3 trials. Our data suggest that MFC
255	ensembles are sensitive to the context or 'rules' in the task $-i.e.$, FI12 vs. FI3/FI12, while the
256	DMS optimizes behavior particularly on FI3 trials. These data provide insight into the relative
257	roles of prefrontal and striatal networks during temporal learning.

258 These results contradicted our hypothesis that time-related ramping in the MFC and DMS would be similarly affected by the introduction of a new temporal interval. Our hypothesis was 259 260 based on five lines of evidence: 1) the existence of strong projections from the MFC to the DMS 261 (Gabbott et al., 2005; Han et al., 2017; Wall et al., 2013), 2) clear roles for both structures in interval timing (Coull et al., 2011; De Corte et al., 2019; Emmons et al., 2017; Meck, 2006), 3) 262 similarities in time-related ramping in the MFC and DMS (Emmons et al., 2017), 4) the necessity 263 of MFC activity for DMS ramping (Emmons et al., 2019, 2017), and 5) our recent demonstration 264 that the stimulation of axons that project from the MFC to the DMS is sufficient to increase time-265 266 related ramping in the DMS (Emmons et al., 2019). Given these data, it is notable that the MFC and DMS play distinct roles during temporal learning, although this observation is concordant 267 with the vastly different connectivity and synaptic organization of these two structures 268 269 (Shepherd, 2003). Nevertheless, decreases in MFC ramping after the introduction of a shorter interval suggest that MFC ramping is sensitive to the temporal context of the one-interval vs. 270

two-interval task, and they may reflect Bayesian priors of temporal probabilities (Jazayeri and
Shadlen, 2010; Shi et al., 2013).

273 Differences between the MFC and DMS were anticipated based on a recent comparison 274 of neuronal ensembles during a temporal categorization task that involved maze running (Kim et 275 al., 2018). This study indicated that ramping was more prevalent in the MFC than the dorsal 276 striatum. In this task as the intervals became longer, temporal decoding by the MFC was less 277 effective than that by the striatum. Although this task was more complex than ours and a number 278 of others (Bakhurin et al., 2017, 2016; Donnelly et al., 2015; Narayanan, 2016; Wang et al., 279 2018), the strong temporal encoding across corticostriatal ensembles is consistent with our 280 findings here.

281 We found that the DMS contained more neurons in which there was a main effect of interval compared to the MFC on the early days of two-interval performance. Notably, half of 282 283 interval-modulated neurons were not ramping. These data suggest that patterns beyond timerelated ramping encode information about temporal intervals. On progressive days of two-284 285 interval performance, interval-related activity between the MFC and DMS equalized. Because 286 our task design involved a second cue for FI3 intervals, we cannot distinguish whether this activity was related to working memory for temporal intervals, cue-related processing, or other 287 288 aspects of interval timing. Future work using more advanced learning paradigms may clarify 289 these patterns of activity.

Our findings are in line with drift-diffusion models of two-interval tasks, as we find that time-related ramping scales with the interval duration (Simen et al., 2011). We find that MFC ramping is sensitive to temporal context whereas DMS ramping is not, and that non-ramping interval-related modulations and temporal predictions in the DMS change with two-interval

performance. These results suggest that time-related ramping reflects distinct processes in MFC 294 and DMS. Given that MFC activity influences ramping in DMS (Emmons et al., 2019, 2017), 295 DMS ramping activity might integrate aspects of MFC ramping as well as non-ramping activity. 296 297 Because time-related ramping activity in MFC and DMS ensembles did not change during two-interval performance, ramping activity may be remarkably stable in both brain 298 299 regions when the temporal context does not change. It is unclear how ramping might change with extended periods of behavior over several days or weeks (Barnes et al., 2005; Graybiel, 2008; 300 301 Yin et al., 2005). However, we did find that on FI3 trials, temporal decoding in the DMS 302 improved even though DMS ramping was stable. In the DMS patterns beyond ramping activity might change during two-interval performance and contribute to improved temporal decoding 303 (Paton and Buonomano, 2018). The improvement in temporal prediction despite unchanged 304 ramping activity supports improved 'population clock'-based temporal predictions during FI3 305 trials (Karmarkar and Buonomano, 2007; Laje and Buonomano, 2013). 306

To our knowledge, our study is one of the first to record from corticostriatal ensembles 307 during temporal learning. The striatum has a well-established role in learning other contexts 308 including habit formation, reversal learning, and instrumental learning (Graybiel and Grafton, 309 2015; Kimchi and Laubach, 2009; Yin and Knowlton, 2006). However, direct comparisons of 310 cortical and striatal learning in rodents, in any context, are rare. One exception is a recent study 311 312 of prenatal alcohol exposure, which showed that the orbitofrontal cortex disengages and the dorsal striatum updates reward contingencies (Marquardt et al., 2020). These findings parallel 313 314 the changes in the MFC and DMS that we report here. The observation that prenatal exposure to 315 alcohol leads to changes in cortical activity underscores the clinical significance of this brain circuit. 316

During associative learning in primates, corticostriatal ensembles are highly sensitive to 317 learning, with striatal neurons rapidly encoding new associations and the prefrontal cortex 318 learning more slowly (Pasupathy and Miller, 2005). Primate striatal neurons rapidly encoded 319 stimulus-response associations, whereas primate prefrontal neurons encoded category abstraction 320 (Antzoulatos and Miller, 2011). In line with these results, we found that time-related ramping 321 322 decreased in two-interval vs. one-interval sessions, suggesting that prefrontal ensembles may be sensitive to temporal categories or context. It is important to note that these primate studies 323 recorded from lateral prefrontal areas, which lack a clear rodent analogue (Laubach et al., 2018), 324 325 and that they employed vastly different task conditions. Nevertheless, our work provides insight into the dynamics of rodent corticostriatal ensembles during an elementary temporal learning 326 paradigm. 327

Our study has several limitations. First, we used fixed-interval timing; peak-interval 328 timing tasks might enable more precise dissection of start and stop times (Rakitin et al., 1998). 329 Second, our techniques cannot identify the genetic or molecular identity of recorded neurons. 330 This detail would be of particular interest in the case of the DMS, which contains D1 and D2 331 MSNs (Kreitzer, 2009). Third, we are unsure if the MFC and DMS neurons we captured were 332 333 connected, because of the sparsity of cortical projections and constraints of our recording techniques (Wall et al., 2013). This limitation might be overcome in future work by exploiting 334 335 optogenetic tagging and retrograde viral tracing to isolate corticostriatal projections (Otis et al., 336 2017). Studying how MFC-DMS connectivity changes with learning might provide further insight in corticostriatal circuits. Fourth, we cannot reliably follow neurons over separate 337 338 sessions. Finally, we were unable to clearly identify clear correlates of temporal learning during the first two-interval session on Day 1. Corticostriatal ensembles may rapidly learn the new 339

interval. Such rapid learning has been observed in the striatum, but capturing it might require a
different task design to capture trial-by-trial neuronal dynamics during learning (Kimchi and
Laubach, 2009).

343 In summary, we investigated corticostriatal ensembles in rodents that had been welltrained to perform a single fixed-interval timing task while they learned to incorporate a new 344 345 interval. We discovered that time-related ramping activity in the MFC decreased following introduction of the shorter interval, whereas ramping activity in the DMS was unchanged. We 346 also found that more DMS neurons fired differentially on each interval compared to the MFC 347 348 early in two-interval performance. Finally, corticostriatal ramping activity did not change on the days following the initiation of two-interval performance, yet DMS temporal decoding 349 350 improved. Taken together, our data suggest that the MFC and DMS play distinct roles in 351 temporal learning.

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356

- 357 **Contributions:** EE, YK, and NN designed experiments; EE and GTC collected data, EE, YK,
- 358 KC, and NN analyzed data, AB independently checked the code and data, and EE, GTC, MW,
- 359 AB, YK, and NN wrote the manuscript.

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361 **Data and Code**: Available at narayanan.lab.uiowa.edu

362 **References**

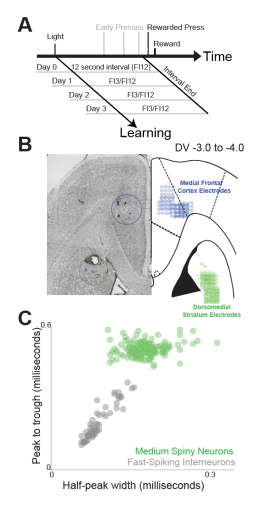
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512 Figures:



514 Figure 1: Fixed-interval timing tasks and recording locations: A) On Day 0, rodents performed fixed-interval timing tasks in which a reward was given for the first lever press after a 515 12-second interval (FI12). Interval start was cued by a house light, motivation was a liquid 516 reward, and presses before interval end were unreinforced. On Day 1, a second, shorter 3-second 517 518 interval (FI3) was introduced and cued by a distinct light. FI3 trials were randomly intermixed with FI12 trials. Recordings were performed for two days following the initial two-interval 519 520 performance (Day 2 and Day 3). B) Animals were implanted with neuronal ensemble arrays targeting the medial frontal cortex (MFC) and dorsomedial striatum (DMS). C) MSNs within the 521 DMS were identified based on waveform shape. 522

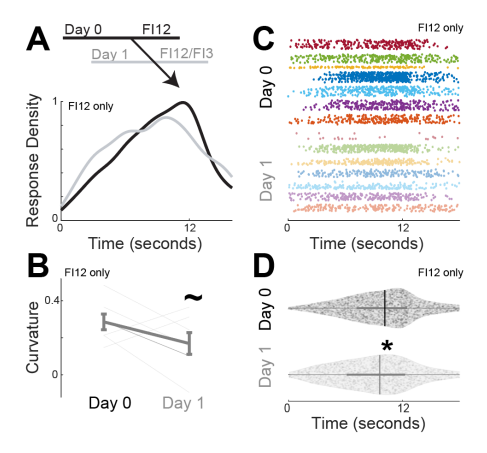
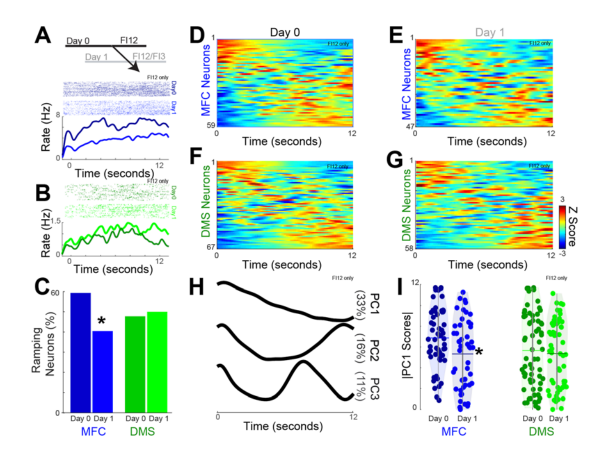


Figure 2: Response times reflect temporal context. A) Kernel density estimates of time-524 525 response histograms across animals on FI12 trials on Day 0 (black) when FI12 trials were presented alone to Day 1, with FI12 trials were presented alongside randomly intermixed FI3 526 trials. B) Curvature indices of time-response histograms from 7 animals on Day 0 vs. Day 1. C) 527 528 Compilation of every FI12 response from every animal on Day 0 (darker colors) vs. Day 1 (lighter colors); each animal is represented by dots of a different color. D) Violin plot of all 529 responses; vertical lines denote the mean and thicker horizontal gray lines span the interquartile 530 range. Data from FI12 trials from seven animals; ~ indicates a trend via Wilcoxon rank-sum; * 531 indicates p < 0.05 via GLMMs. 532

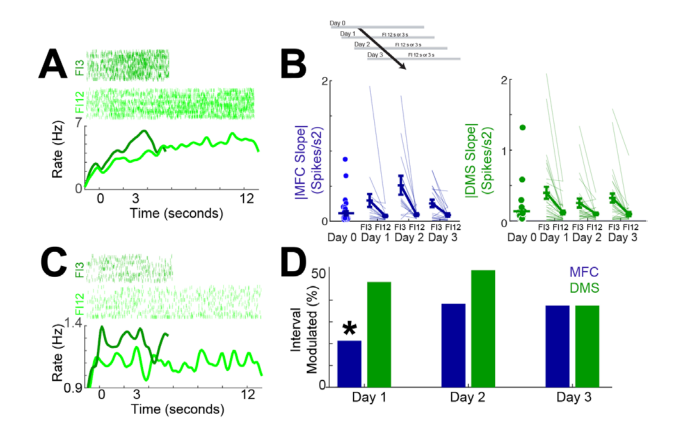
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Figure 3: MFC ramping reflects temporal context. Peri-event rasters for single neurons in the 535 A) MFC (blue) and B) DMS (green). Top panels: each row represents a trial; each tick is an 536 action potential; darker colors represent Day 0 (FI12) and lighter colors represent Day 1 537 (FI12/FI3); all data are from FI12 trials only. C) Quantification of neurons that underwent time-538 related ramping via GLMMs in the MFC and DMS for Day 0 and Day 1; * indicates p < 0.05 via 539 a chi-squared test. Peri-event histograms from all neurons in the MFC on Day 0 (D) and Day 1 540 541 (E) and all MSNs in the DMS on Day 0 (F) and Day 1(G). H) Principal component analyses revealed three main components; percentage of variance is indicated in parentheses. I) |PC1 542 scores for MFC and DMS ensembles on Day 1 vs. Day 0. Each circle represents the PC1 score 543 544 from a single neuron; horizontal lines denote the mean and thicker vertical lines span the

- interquartile range. * indicates p < 0.05 via Wilcoxon sign-rank. Data from MFC and DMS
- 546 recordings in 7 animals.



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Figure 4: MFC and DMS activity is distinct on FI3 vs. FI12 trials. A) An exemplar ramping neuron from the DMS; the slope of firing rate vs. time was steeper on FI3 vs. FI12 trials. B) |Slopes| for MFC (blue) and DMS (green) neurons on Day1, Day 2, and Day 3 for FI3 and FI12 trials; for both MFC and DMS |slopes| were higher on FI3 trials. C) An exemplar neuron from the DMS that fired differentially on FI3 vs. FI12 trials. D) The number of neurons with a main effect of firing rate vs. interval for the MFC and DMS. * indicates p < 0.05 via a chi-squared test; Data from MFC and DMS recordings in 7 animals.

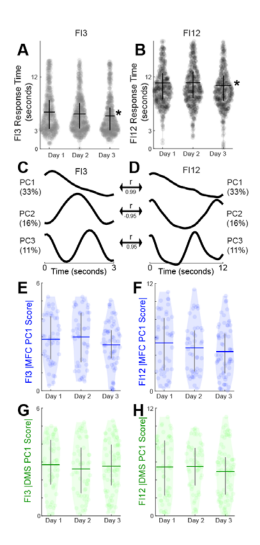


Figure 5: MFC and DMS ramping is stable with two-interval performance. A) Response

- times by day for FI3 and B) FI12 trials over the three days of two-interval performance. Principal
- components for C) FI3 and D) FI12 trials. Ramping activity as measured by |PC1| scores for E)
- 560 FI3 trials in the MFC (blue), F) FI12 trials in the MFC, G) FI3 trials in the DMS, and H) FI12 in
- the DMS. Horizontal lines denote the mean and thicker vertical gray lines span the interquartile
- range. * indicates p < 0.05 via GLMMs. Data were from MFC and DMS recordings in 7 animals.

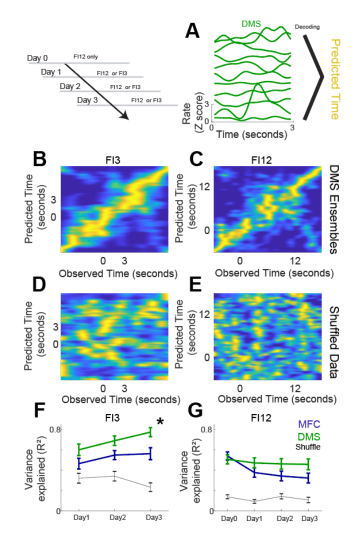


Figure 6: Decoding reveals that DMS improves temporal predictions with two-interval 564 performance. A) We trained decoders (naïve Bayesian classifiers) to predict time from firing 565 566 rate. B) Decoder performance for DMS ensembles for FI3 trials and C) for FI12 trials. Predicted time is on the y axis and observed time is on the x axis; decoded time is in yellow. D) Decoded 567 performance for the same DMS ensembles with time-shuffled data for FI3 trials and E) FI12 568 trials. F). We measured decoder performance by calculating the variance explained (R^2) of 569 predicted vs. observed time. For FI3 trials, DMS ensembles increased R² with two-interval 570 performance, while MFC and G) FI12 MFC and DMS decoding was unchanged. * indicates a 571 main effect of Day 1 \rightarrow Day 3 via GLMMs; data from MFC and DMS recordings in 7 animals. 572

573 **Tables** (significant values in **bold**)

Table 1: Day 0-Day 1 effects on firing rate in MFC and DMS

Model: FiringRate~Times*Area*Learning+(1|Response)+(1|Neurons) Obs 2294280 Model R² 0.19

Predictor	$oldsymbol{F}$	р
Times	111.69	4.00E-26
Area	6.06	0.014
Learning	1.77	0.183
Times:Area	686.31	3.00E-151
Times:Learning	96.96	7.00E-23
Area:Learning	1.39	0.238
Times:Area:Learning	97.5	5.00E-23

Table 2: Firing rate on FI3 vs. FI12 trials

Model: FiringRate~Area*Learning*Interval+(1|Response)+(1|Neurons) Obs 3018780 Model R² 0.15

Predictor	F	р
Area	0.48	0.49
Learning	0.58	0.45
Interval	3.89	0.05
Area:Learning	0.81	0.37
Area:Interval	0.32	0.57
Learning:Interval	13.16	0.0003
Area:Learning:Interval	6.35	0.01

Table 3: FI3 Day 1-Day 3 effects on firing rate

Model: FiringRate~Times*Area*Learning+(1|Response)+(1|Neurons) Obs 533580 R² 0.15

Predictor	$oldsymbol{F}$	р
Times	0	0.959
Area	0.5	0.481
Learning	0.56	0.456
Times:Area	5.41	0.02
Times:Learning	0.04	0.846
Area:Learning	0.69	0.407
Times:Area:Learning	1.2	0.274

Table 4: FI12 Day 1-Day 3 effects on firing rateModel: FiringRate~Times*Area*Learning+(1|Response)+(1|Neurons)Obs 2485200 R² 0.16

Predictor	F	р
Times	21.84	3.00E-06
Area	0.84	0.361
Learning	0.16	0.69
Times:Area	34.34	5.00E-09
Times:Learning	7.41	0.006
Area:Learning	0.74	0.388
Times:Area:Learning	7.08	0.008