Acquisition of “Start” and “Stop” response thresholds in peak-interval timing is differentially sensitive to protein synthesis inhibition in the dorsal and ventral striatum

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INTRODUCTION

Throughout their daily routines, humans and other animals perceive events as a function of the flow of time and make decisions based upon the relative durations of those events by means of a process known as scalar timing (Gibbon et al., 1984; Meck, 2003; Buhusi and Meck, 2005; Buhusi et al., 2009). Therefore, it is perhaps not surprising that timing and time perception within the seconds-to-minutes range—i.e., interval timing—plays a fundamental role in adaptive behavior, including optimal foraging, temporal discounting, and other aspects of intertemporal preferences and neuroeconomics (Brunner et al., 1992; Bateson, 2003; Zauberman et al., 2009; Cui, 2011; Ray and Bossaerts, 2011). Indeed, interval timing contributes importantly to decision-making as well as learning predictive relationships among stimuli in both appetitive and aversive contexts (Gibbon et al., 1997; Gallistel and Gibbon, 2000; Harrington et al., 2004; Tanaka et al., 2004; Maimon and Assad, 2006; Droit-Volet and Meck, 2007; Meck and MacDonald, 2007; Forstmann et al., 2008; Jones et al., 2011). In fact, temporal discounting and the anticipation of a future event’s time of occurrence often emerge as critical variables that influence performance during tasks intended to characterize a variety of basic cognitive processes, including memory (Malapani et al., 1998; Lustig et al., 2005; Lustig and Meck, 2005, 2011), attention (Lustig and Meck, 2001; Buhusi and Meck, 2002, 2006, 2009a,b; Meck and Benson, 2002; Coull et al., 2004), choice (Fantino et al., 1979; McClure et al., 2004; Rudebeck et al., 2006), and reaction time (MacDonald and Meck, 2004, 2006).

Although dopamine-glutamate interactions within cortico-striatal circuits have been shown to support numerous aspects of interval timing (Meck, 1983, 1996, 2006a,b; Matell et al., 2004; Liao and Cheng, 2005; Cheng et al., 2006, 2007a,b,c; Drew et al., 2007; Pennartz et al., 2009; Meck et al., 2008, 2012; Williamson et al., 2008; Agostino et al., 2011; Coull et al., 2011; Gu et al., 2011; Hata, 2011; Höhn et al., 2011; Jones and Jahanshahi, 2011; Allman and Meck, 2012), it remains unclear what types of neural mechanisms are involved in the construction of decision thresholds for determining when to start and stop responding (see Meck, 2002, 2006c; Vink et al., 2005; Bromberg-Martin et al., 2010; van der Meer et al., 2010). In order to investigate this issue, we employed the peak-interval (PI) timing procedure which is a “classic” example of a duration reproduction task that has proven useful in the study of temporal cognition due to its applicability across a wide-variety of animal species—including fish, birds, rodents, and primates (Church et al., 1991, 1994; Rakitin et al., 1998; Paule et al., 1999; Buhusi and Meck, 2000; Buhusi et al., 2002, 2009; Gallistel et al., 2004; Hinton and Meck, 2004; Drew et al., 2005, 2007; Penney et al., 2008; Ward et al., 2009, 2012; Cheng et al., 2011) and stimulus modalities (Meck and Church, 1982; Meck, 1991; Penney et al., 2000; Buetsi, 2011; Lustig and Meck, 2011; Allman and Meck, 2012). In the PI timing procedure, the subject is initially exposed exclusively to fixed-interval (FI) trials throughout a session, during which the onset of a signal (e.g., houselight) reliably predicts the time at which feedback/reward will be provided—thus establishing a temporal criterion. After sufficient training with FI trials, unreinforced probe trials are randomly intermixed within a session in order to characterize the accuracy and precision of temporal memory as illustrated in Figure 1A. During a probe trial, the signal stays on long past the temporal criterion and no feedback is provided. Following sufficient training with probe trials, the mean response rate for an individual subject averaged across all probe trials within a session resembles a Gaussian-shaped distribution—the peak function—whose mode is centered on the temporal criterion (Meck and Church, 1984; Church et al., 1994; Rakitin et al., 1998). In such
FIGURE 1 | (A) An illustration depicting the two trial types randomly intermixed throughout a standard peak-interval (PI) training session. The houselight serves as the signal being timed in this example. On FI trials, the houselight’s onset predicts the availability of reinforcement at 20 s. Delivery of reward is contingent on a lever press made by the rat at or after this time, which also terminates the houselight and initiates an ITI. On probe trials, the houselight remains on for an extended period of time with a minimum of 60 s and reinforcement is never delivered. (B) A representative rat that is well-trained on the PI procedure. The raster plot (top) depicts each probe trial as a row and responses (red tics) are displayed across time during the probe trial. On each trial, a response sequence contains both a “Start” and “Stop” response (blue tics), which demarcates the beginning (“Start”) and end (“Stop”) of the “high state” (see Introduction and Methods). Probe trial responses are collapsed across trials within 1 s bins to create the peak function, which is normalized so that it expresses the percentage of maximum response rate as a function of signal duration (bottom). Because the signal extends well past the temporal criterion (e.g., 20 s), the accuracy and precision of the remembered time of reward is inferred by the rat’s response rate before and after 20 s. (C) Before probe trials are randomly intermixed within a session, rats are exposed only to FI trials for multiple sessions (see Methods). This panel depicts a normalized peak function (± SEM) averaged across all rats during the first eight sessions in which probe trials are randomly intermixed with FI trials. Note that the degree of response inhibition after the remembered time of reward increases across sessions as the rats obtain more experience with partial extinction during unreinforced probe trials.

With rats and pigeons, for whom an appetitive reward can readily serve as a time marker (Freestone and Church, 2010), performance during a single probe trial can be described by an abrupt transition both into and out of a “high state” of responding, during which the subject responds at a relatively rapid,

cases, a balance is often struck between starting the sequence of reinforced behavior in anticipation of the reward, but also giving up responding when the opportunity for reward is thought to have passed (Brunner et al., 1992, 1996, 1997; Bateson, 2003; Silva and Timberlake, 2005).
constant rate as illustrated in Figure 1B. Remarkably, the variability in the onset (“Start”) and offset (“Stop”) of the “high state,” as well as the duration of the “high state” itself, is proportional to the interval being timed (Church et al., 1994; Rakitin et al., 1998). This feature contributes to the scalar property, which describes the proportionality between the spread of the peak function and the value of the temporal criterion (i.e., Weber’s law—Gibbon et al., 1984; Cheng and Meck, 2007). Importantly, the “Start” and “Stop” times are considered independent of one another because they are not well correlated on a trial-by-trial basis (Church et al., 1994; Gallistel et al., 2004; Matell et al., 2006; Buhusi and Meck, 2009b). In effect, these “Start” and “Stop” times are independent decision thresholds demarcating an action sequence that the subject “attempts” to center on the expected time of reward. Moreover, the presentation of unreinforced probe trials subsequently leads to the acquisition of the “Stop” response threshold (see Balci et al., 2009). At steady-state levels of performance this action sequence may be conceptualized as a distinct behavioral unit whose organization depends on the memory of the target duration (see Gibbon et al., 1984; Church et al., 1994).

However, despite the evidence supporting a clear dissociation between “Start” and “Stop” times, there has been no anatomical localization of these response thresholds for interval-timing tasks (see Mansfield et al., 2011). In addition, there is little understanding concerning (1) how the “Start” and “Stop” thresholds are acquired and (2) the underlying neural substrates for temporal conditioning (Gallistel and Gibbon, 2000). Regarding the first point, in the PI timing procedure the subject must learn to inhibit responding after the target duration during probe trials, which ultimately leads to the gradual acquisition of the “Stop” response as illustrated by the PI data from Experiment 1 plotted in Figure 1C. This occurs when partial extinction/unrewarded probe trials are first introduced, typically long after the emergence of the “Start” response which defines the beginning of the “high state” of responding acquired during initial FI training. Regarding the second point, there is reason to believe that the striatum is an important neural substrate for interval timing in a wide variety of timing procedures (Gibbon et al., 1997; Matell and Meck, 2000, 2004; Buhusi and Meck, 2005; Meck, 2006b; Harrington et al., 2011a,b; Kotz and Schwartzte, 2011; Matell et al., 2011; Portugal et al., 2011; van Rijn et al., 2011). Indeed, the firing rate of neurons in the dorsal striatum (DS) is modulated with respect to the temporal criterion acquired in the PI timing procedure (Matell et al., 2003). Outside the context of interval timing, one proposed function of the DS is organizing the hierarchical structure of learned response sequences (Graybiel, 1998; Fujii and Graybiel, 2005; Jin et al., 2009; Jin and Costa, 2010), which is consistent with a direct relationship to interval timing. For example, single neurons within the rat’s DS also encode the hierarchal context of a specific response within a systematic, natural response sequence like grooming (Albridge and Berridge, 1998) and lesions of the DS can disrupt the expression of both natural and learned sequences (Cromwell and Berridge, 1996; Bailey and Mair, 2006). While the DS is believed to contribute to encoding of the temporal criterion, the ventral striatum (VS) is thought to play a modulatory role for interval timing, especially with respect to changes in the reward value associated with feedback at the temporal criterion (McClure et al., 2004; O’Doherty et al., 2004; MacDonald and Meck, 2005; Meck, 2006b). This connection is consistent with a broader framework for understanding VS function in which this region of the striatum appears to hold a closer relationship to affective processing than the DS and plays a role in “behavioral flexibility” by modulating ongoing natural and/or learned behaviors (Cardinal et al., 2002; Kelley, 2004; Pleil et al., 2011). For example, the VS contribute to inhibiting behavior that is made inappropriate so that novel behavioral strategies may be acquired (Tanaka et al., 2004, 2007; Floresco et al., 2006).

With these functional parallels in mind, we were interested in characterizing the role of the DS and VS in the PI timing procedure while rats acquired the “Start” and “Stop” responses. Learning entails enduring changes in the brain, which presumably requires a cascade of molecular events to take place beforehand (Abel and Lattal, 2001). Protein synthesis inhibitors, such as anisomycin (ANI), have proven to be useful tools for the characterization of this memory consolidation process in Pavlovian conditioning (Stafford and Lattal, 2009), spatial navigation (Morris et al., 2006), instrumental learning (Hernandez et al., 2002), and motor-skill acquisition (Wächter et al., 2010). In the following experiments, we usedANI microinjections administered prior to rats performing on variants of the PI timing procedure to conclude that the acquisition of the “Start” response depends on normal functioning (including protein synthesis) within the DS during training (see Castellucci et al., 1989; Okamoto et al., 2011). In contrast, disruption of normal functioning within the VS during training, but not the DS, impairs the acquisition of the “Stop” response.

MATERIALS AND METHODS

ANIMALS

Across both experiments, a total of 49 experimentally naive Sprague–Dawley rats (Charles-River Laboratories, Raleigh, NC) weighing between 250–350 g at the start of the experiment were used. Rats were housed in pairs in a 12:12 light:dark cycle with lights on from 7:00 AM to 7:00 PM. Rats were given continuous access to water and maintained at 85% free-feeding weight with a daily ration of rat chow (Rodent Diet 5001, PMI Nutrition International, Inc., Brentwood, MO) given shortly after the test session. All procedures were conducted in accordance with the policies of the Duke University Institutional Animal Care and Use Committee.

APPARATUS

The apparatus consisted of 10 standard lever boxes (Model ENV-007, MED Associates, Inc., Albans, VT) housed in light and sound attenuating cubicles (Model ENV-019, MED Associates, Inc., Albans, VT). Each lever box had inside dimensions of approximately 24 cm × 31 cm × 31 cm. The top, side walls, and door were constructed of clear acrylic plastic. The front and back walls were constructed of stainless steel, and the floor was comprised of 19 parallel stainless steel bars. Each lever box was equipped with two retractable response levers (Model ENV-112, MED Associates, Inc., Albans, VT) situated on opposite sides of the front wall of the lever box. Precision food pellets (45 mg—Research Diets, Inc., New Brunswick, NJ) could be delivered by a pellet dispenser (Model ENV-203, MED Associates, Inc., Albans, VT) to a food
cup on the front wall, 1 cm above the floor, and in between the two levers. A 28 V, 100 mA, 2500 lx house light was mounted at the center-top of the front wall and could be used to illuminate the box. A white noise amplifier/speaker system (Model ENV-225, MED Associates, Inc., Albans, VT) was mounted on the opposite wall from the levers but was not used. A 66 dB sound produced by a ventilation fan was present throughout all procedures.

**DRUGS AND INFUSIONS**

ANI (Sigma-Aldrich, St Louis, MO) was dissolved in equimolar HCl and diluted with CSF (Harvard Apparatus, Holliston, ME—Experiment 1) or 0.9% saline (Sigma-Aldrich, St Louis, MO—Experiment 2), and adjusted to pH 7 with NaOH for a final concentration of 62.5 μg/0.5 μl. The ANI dose was selected based on preliminary data from our laboratory and a previous study that confirmed it as being an effective dose for behavioral tasks of this sort (Hernandez and Kelley, 2004). In memory consolidation experiments, protein synthesis is often inhibited after training is completed. Depending on the experimental question, this strategy can be especially effective when using tasks that require a limited numbers of trials (e.g., contextual fear conditioning) to meet some learning criteria. However, learning the “Stop” response requires multiple daily sessions that each include 60 or more total trials/session (Figure 1C). Because of this we couldn’t be sure which specific trial or trials within the session constitute the “learning episode” as well as when the “Start” and “Stop” response thresholds are consolidated (or in fact whether these task components are reconsolidated at any point during the session). As a consequence, we chose to infuse ANI before the training session began in order to ensure that protein synthesis was maximally impaired throughout the session (see Wanisch and Wotjak, 2008 for data on the time course of protein synthesis inhibition).

All rats were given at least three days of mock infusions to habituate them to the injection procedure before testing. Bilateral infusions were given immediately before the sessions at a rate of 0.25 μl/min over the course of two minutes. Following the infusion, the injector cannulae were maintained in place for at least 1 min before being placed immediately in the lever box to begin the experiment.

**HISTOLOGY**

In order to confirm cannulae placement, following the experiment, all rats were perfused transcardially with 0.9% saline followed by 10% formalin. Brains were removed and stored in 30% sucrose solution before sectioning. The tips of the injector cannulae extended ~1.5 mm from the base of the guide cannulae. The cannulae were fixed using dental cement and three small, skull screws. All rats were given at least five days of recovery.

**SURGERY**

Rats were anesthetized with i.p. injections of both ketamine hydrochloride (100 mg/kg) and xylazine hydrochloride (20 mg/kg). Bilateral guide cannulae (Plastics One Inc., 26 gauge with stylets) were implanted into either the DS or VS cerebral nuclei using aseptic technique. All rats were given at least five days of recovery.

**EXPERIMENT 1: BEHAVIORAL TRAINING: EFFECTS OF ANISOMYCIN INFUSION ON “START” AND “STOP” RESPONSES DURING THE ACQUISITION OF A PEAK-INTERVAL (PI) 20 S RESPONSE SEQUENCE**

**Pre-training (Sessions 1–5)**
All rats received six sessions of combined magazine and lever training. During these sessions, a food pellet was delivered once a min for 60 min, and was signaled at the 58 s mark by a 2 s lever retraction/extension sequence. Throughout the session, a lever press also resulted in the delivery of a food pellet.

**Fixed-interval (FI) 20 s training (Sessions 6–55)**
During FI 20 s training, trials were cued by the onset of a houselight and extension of the response lever for the duration of the trial. Any lever press before 20 s had no consequence. However, any lever press after 20 s resulted in (1) delivery of a food pellet, (2) offset of the houselight, (3) retraction of the lever, and (4) the beginning of an intertrial interval (ITI). An ITI was always 60 s plus a duration randomly selected from one of seven values (min = 1.1 s, max = 70.4 s) that were geometrically distributed. A new trial began at the end of an ITI and sessions lasted for approximately 2 h.

**Post-surgical fixed-interval (FI) 20 s training (Sessions 56–65)**
Following surgery, rats were given 10 additional sessions of FI training to ensure stable pre-treatment performance for the groups.

**Peak-interval (PI) 20 s training (Sessions 66–80)**
PI 20 s training consisted of 50% FI trials (see above) and 50% unreinforced probe trials. On any given trial, the chosen trial type was random. Probe trials were like FI trials except that (1) lever presses after 20 s did not result in delivery of a food pellet and (2) the houselight remained on and the lever extended for 60 s plus a duration randomly selected from one of seven values (min = 1.1 s, max = 70.4 s) that were geometrically distributed. The end of a trial signaled the beginning of an ITI, whose parameters were the same as that described during FI training and sessions lasted approximately 2 h. Rats were microinjected with the appropriate solution (ANI or vehicle) 10 min prior to the beginning of the first 6 PI 20 s training sessions.

**EXPERIMENT 2: EFFECTS OF ANISOMYCIN INFUSION ON “START” RESPONSES DURING THE TRANSITION FROM 20 S TO 50 S TEMPORAL CRITERION**

**Pre-training (Sessions 1–5)**
All rats received six sessions of combined magazine and lever training. During these sessions, a food pellet was delivered once a min for 60 min, and was signaled at the 58 s mark by a 2 s lever retraction/extension sequence. Throughout the session, a lever press also resulted in the delivery of a food pellet.

**Fixed-interval (FI) 20 s training (Sessions 6–20)**
During FI training, trials were cued by the onset of a houselight and extension of the response lever. Any lever press before 20 s had no consequence. However, any lever press after 20 s resulted in (1) delivery of a food pellet, (2) offset of the houselight, (3) retraction of the lever, and (4) the beginning of an ITI. An ITI was always 60 s plus a duration randomly selected from one of seven values
A temporally controlled “Stop” response requires normal functioning (including protein synthesis) in the VS, but not the DS. Thirty rats were trained on a 20 s PI procedure. Depending on the group assignment, rats were microinjected with ANI (DS-ANI, n = 7; VS-ANI, n = 7) or vehicle solution (DS-CON, n = 8; DS-CON, n = 8) before the first six sessions during which probe trials were first introduced in a training session. We divided the data into two blocks and calculated the mean peak functions for treatment groups during these blocks.

Figure 2 illustrates the mean peak functions (left column) and results of the single-trials analysis (right column) for the CON treatment groups combined over the DS and VS conditions as well as the ANI treatment groups for the separate DS and VS conditions. In each case, the acquisition data are plotted in blocks of 3 sessions (rows), ANI Sessions 1–3 and ANI Sessions 4–6. The data from the CON rats for the DS and VS conditions were combined because there were no reliable differences between these two groups for the “Start” measures during sessions 1–3 ($t_{1.40} = 0.61, p > 0.05$) and sessions 4–6 ($t_{1.22} = 1.17, p > 0.05$), as well as for the “Stop” measures during sessions 1–3 ($t_{1.40} = 0.01, p > 0.05$) and sessions 4–6 ($t_{1.22} = 1.96, p > 0.05$). A One-Way ANOVA was conducted on the single-trial “Stop” times for ANI Sessions 1–3 and ANI Sessions 4–6. During ANI Sessions 1–3, there was no reliable effect of Treatment on the “Stop” times ($F_{2.87} = 0.76, p > 0.05$). However, during ANI Sessions 4–6, there was a significant effect of Treatment ($F_{2.69} = 11.32, p < 0.0001$). Post-hoc tests [Fischer’s Least Significant Difference (LSD), $\alpha = 0.05$] confirmed that the VS-ANI group differed significantly from both the CON and DS-ANI groups in terms of the time of the “Stop” response. No other significant differences were confirmed, i.e., there were no significant differences between the CON-DS and ANI-DS groups or the CON-VS and ANI-VS groups, $p's > 0.05$.

During acquisition of the “Stop” response, anisomycin infusion in the dorsal striatum resulted in a more precise “Start” response

A similar One-Way ANOVA was used to determine the effect of Treatment on the single-trial “Start” response times. During ANI Sessions 1–3, there was no reliable effect of Treatment on the “Start” times ($F_{2.87} = 1.12, p > 0.05$), however, during ANI Sessions 4–6, a significant effect of Treatment on “Start” times was observed ($F_{2.69} = 5.13, p < 0.01$). Post-hoc tests (Fischer’s LSD, $\alpha = 0.05$) confirmed that only the DS-ANI treatment group differed from both the CON and VS-ANI treatment groups. With respect to the expected time of reward, the DS-ANI treatment group withheld responding to a greater degree beginning from the onset of a probe trial. Therefore, rats in the DS-ANI treatment group entered the “high state” at a time that was significantly closer to the target duration, which suggests an increase in temporal precision (Church et al., 1991). The significant main effects observed during ANI Sessions 4–6 continued to be observed during post-ANI Sessions 1–3 during which no ANI treatments were given, $F_{3.69} = 3.92, p < 0.001$ and $F_{3.69} = 0.52, p < 0.0001$ for “Start” and “Stop” response times, respectively.

The differential effects of ANI infusion on the acquisition of “Stop” response were essentially non-existent following extended training without ANI treatment as demonstrated by the lack of
Figure 2 | Normalized mean (± SEM) peak functions (left column) for CON (blue), DS-ANI (red) and VS-ANI (green) treatment groups during four phases of the experiment: (A) ANI Sessions 1–3, (B) ANI Sessions 4–6, (C) post-ANI Sessions 1–3, and (D) post-ANI Sessions 12–14. Insets (right column) are included for each peak function to describe the “Start” and “Stop” times obtained from the signal-trials analysis conducted on the data obtained from that phase. Asterisks denote a significant difference between the treatment groups compared, which are bracketed by dark lines. ANI = anisomycin; CON = control; DS = dorsal striatum; VS = ventral striatum.

A One-Way ANOVA conducted on both the “Start” and “Stop” times during these post-ANI Sessions using Treatment condition as the main factor confirmed non-significant effects on both the “Start” and “Stop” measures, ($F_{2,69} = 0.37, p > 0.05$) and ($F_{2,69} = 1.87, p > 0.05$), respectively. These observations indicate that although the effects of
ANI lingered for a few sessions following its discontinuation (e.g., post-ANI Sessions 1–3), it did not damage the brain in a way that permanently disrupted the temporal control of behavior.

Anisomycin treatment lowered response rates to a comparable degree in both the dorsal striatum and ventral striatum treatment groups, but had no long-lasting effects on timing behavior

Mean (± SEM) response rate during probe trials for CON, DS-ANI, and VS-ANI treatment groups is plotted as a function of the mean (Anisomycin treatment lowered response rates to a comparable degree in both the dorsal striatum and ventral striatum treatment groups, but had no long-lasting effects on timing behavior. The baseline peak functions corresponding to probe trials obtained from the final two 20 s PI training sessions before ANI treatment are illustrated in Figure 4A. The peak functions are normalized with respect to the time-bin with the highest response rate (peak rate) in order to determine the shape of the temporal gradient and its centering around the target duration (peak time). Peak time and peak rate have been shown to be independent measures of performance with the former reflecting motivational factors and the later reflecting timing factors (Roberts, 1993; Cheng and Meck, 2007). While the response rate in both ANI treatment groups decreased by the same degree relative to the CON treatment groups, the effects of ANI administration on the “Start” and “Stop” times for the DS-ANI and VS-ANI treatment groups were significantly different from each other as reported above.

EXPERIMENT 2: ANISOMYCIN INFUSION INTO THE DORSAL STRIATUM IMPAIRS THE ACQUISITION OF THE “START” RESPONSE IN REFERENCE TO A SECOND TEMPORAL CRITERION

In the present experiment, normal functioning was disrupted by ANI infusion into the DS while a group of rats was transitioned from a previously acquired 20 s temporal criterion to a new 50 s criterion. The effect of ANI on the “Start” response was evaluated during the acquisition of a second temporal criterion rather than the initial temporal criterion in order to avoid non-specific disruption of behavior due to failure to consolidate the association of the lever press with food reinforcement as opposed to failure to establish a decision threshold associated with a specific temporal criterion. Investigation of the effects of striatal disruption (including inhibition of protein synthesis) on the acquisition of a second “Start” threshold provides the subject with the option of maintaining the original “Start” threshold in the absence of the ability to acquire a new “Start” threshold, thus providing a more specific test of decision thresholds as opposed to simple instrumental associations (see Meck et al., 1984b; Balci et al., 2009).

Nineteen rats were first trained on a 20 s PI procedure (see Methods) and were then divided into two groups, which differed with respect to whether ANI (DS-ANI, n = 10) or vehicle (DS-CON, n = 9) would be microinjected through bilateral cannulae placed into the DS. Within a session, 50% of the trials were FI trials. The remaining 50% of the trials were unreinforced probe trials during which the houselight stayed on for at least 60 s and no lever presses were reinforced (see Figure 1A and Methods section).

The baseline peak functions corresponding to probe trials obtained from the final two 20 s PI training sessions before ANI treatment are illustrated in Figure 4A. The peak functions are normalized with respect to the time-bin with the highest response rate (peak rate) in order to determine the shape of the temporal gradient and its centering around the target duration (peak time). Peak time and peak rate have been shown to be independent measures of performance with the former reflecting motivational factors and the later reflecting timing factors (Roberts, 1993; Cheng and Meck, 2007). “Start” times obtained from the single-trials analysis conducted across each of the PI 20 s baseline sessions are depicted in Figure 4D (left panel). A Two-Way ANOVA with Treatment and Session as factors indicated non-significant effects of Treatment ($F_{1,31} = 0.15, p > 0.05$) and Session ($F_{1,31} = 0.003, p > 0.05$), as well as a non-significant interaction between these two factors ($F_{1,31} = 0.02, p > 0.05$) during the initial training phase.

Each treatment group was then transitioned to a 50 s PI training protocol and rats were microinjected with ANI or CON prior to the first 10 test sessions. Figure 4B represents the mean peak functions obtained from Sessions 7–8 of ANI treatment while Figure 4D (center panel) illustrates the “Start” times obtained during each session of this transition phase as well as the PI 50 s baseline sessions (right panel). A Two-Way ANOVA conducted
on the 20 s → 50 s transition phase data revealed significant effects on “Start” times with respect to Treatment and Session, \((F_{1,17} = 4.47, p < 0.05)\) and \((F_{7,119} = 7.14, p < 0.0001)\), respectively. In addition, a significant Treatment × Session interaction was observed, \((F_{7,119} = 2.91, p < 0.01)\). The mean response rates on probe trials during the three specified phases of the experiment for the DS-CON and DS-ANI treatment groups are illustrated in Figure 4E. A Two-Way ANOVA was conducted on each of these phases (Baseline-20 s, Transition Phase, and Baseline-50 s). During the Transition Phase, a significant effect of Session was observed \((F_{7,119} = 5.46, p < 0.0001)\), but non-significant effects of Treatment \((F_{1,17} = 2.60, p > 0.05)\) as well as the Session × Treatment interaction \((F_{7,119} = 0.69, p > 0.05)\). In addition, there were no reliable effects of Session or Treatment on response rates during either of the 20 s or 50 s Baseline phases, \(p's > 0.05\).

Following the completion of ANI or vehicle administration (CON), each treatment group continued to be trained on the 50 s PI procedure, but with microinjections discontinued. While ANI administration significantly impaired the acquisition of the “Start” response across the 10 treatment sessions, there did not appear to be any long-lasting functional damage to the DS with respect to the temporal control of behavior. Figure 4C shows the mean peak functions for each treatment group obtained approximately three weeks (post-ANI Sessions 21–22) following microinjections while Figure 4D (right panel) depicts the “Start” times during this Baseline-50 s phase. A Two-Way ANOVA with Treatment and Session as factors indicated non-significant effects of Treatment \((F_{1,32} = 0.71, p > 0.05)\), Session \((F_{1,32} = 1.07, p > 0.05)\), as well as the Treatment × Session interaction \((F_{1,32} = 0.11, p > 0.05)\).

Histological examination for rats bearing bilateral, indwelling cannulae aimed at the DS and VS verified accurate placements for all rats in the ANI and CON treatment groups at both anatomical locations. A summary of these results for Experiment 1 and 2 is provided in Figure 5 (panels A–C).
Indeed, the VS inhibit habitual responding in situations during which there is no reward or low stimulus control (Reading and Frontiers in Integrative Neuroscience www.frontiersin.org March 2012 | Volume 6 | Article 10 | DS and VS.

In this series of experiments we provide the first demonstrations that differential patterns of activity in the dorsal and VS are required for the types of decision-making used in the temporal control of response sequences. In particular, normal functioning of the DS is necessary for the acquisition of a timed “Start” response and similar activation in the VS, but not the DS, is required for the acquisition of a timed “Stop” response. Interestingly, during acquisition of the “Stop” response, the “Start” responses exhibited by the rats in the DS-ANI treatment group were sharper relative to the timing performance of the rats in the other treatment groups. This observation supports our finding in Experiment 2 that the DS is connected with the “Start” response. Therefore, ANI infusion into the DS maintained and/or increased the precision of the temporal estimate for the “Start” response while rats were acquiring the “Stop” response. The basis for this double dissociation might be grounded in the behavioral context that underlies PI training and the acquisition of a “Start” response. Indeed, this finding suggests that a decrease in the probability of reward relating to a predictive cue (i.e., houselight), which until this point in training had reliably signaled the expected time of reward delivery, brings about activational changes (possibly involving protein synthesis) in both the DS and VS.

In concert with mesolimbic dopaminergic input, the VS is believed to contribute to “behavioral switching,” which reflects the reallocation of behavioral resources from current behavior(s) to subsequent one(s) in a flexible manner (Koob et al., 1978; Cools, 1980; Robbins and Brown, 1990; Redgrave et al., 1999). Indeed, the VS inhibit habitual responding in situations during which there is no reward or low stimulus control (Reading and Dunnett, 1991). Though interval timing is typically not studied within an associative framework (Kirkpatrick and Church, 1998), it is well-known that FI schedules can promote responding guided by “habitual” (S-R) associative content (Dickinson et al., 1983) and the transition to habitual responding can occasion changes within neurotransmitter systems that are directly related to interval timing (Choi et al., 2005; Faure et al., 2005; Yin and Knowlton, 2006; DeRusso et al., 2010). Moreover, it is often suggested that the VS contributes to goal-directed behavior by serving as an interface between limbic regions, which are thought to underlie affective (e.g., amygdalar) and contextual (e.g., hippocampal) processing, and provides a structural nexus whereby motivational information can gain access to predictive stimuli (Cardinal et al., 2002; Kelley, 2004; van der Meer et al., 2010). With this in mind, the VS might contribute to the acquisition of the “Stop” response by integrating updated, behavioral-relevant information with output from a neural timing mechanism, which provides a signal for when the expected time of reward has passed. Downstream motor structures could then be triggered to coordinate the termination of an ongoing response sequence (see Roesch et al., 2009). In this sense, the “Stop” response in the PI timing procedure corresponds to the “giving-up time” in foraging situations where a subject has to decide when to leave a patch. “Giving-up times” have been shown to be more sensitive to the distribution of reinforcement times and magnitudes than the decision of when to enter a patch—which have been related to the differential control of “Start” and “Stop” responses in the PI timing procedure (Bruner et al., 1992, 1996, 1997; Taylor et al., 2007).

Conversely, activational changes (possibly involving protein synthesis) in the DS appear to contribute to the shaping of the “Start” response, which determines when the onset of the “high state” occurs. Some researchers have proposed that the DS plays an important role in the initiation of response sequences (Carli et al., 1985; Robbins and Brown, 1990; Graybiel, 1998; Bailey and Mair, 2006), but precise role(s) of the basal ganglia in kinematics and habits remain controversial (Graybiel, 2008; Desmurget and Turner, 2010; Turner and Desmurget, 2010). In Experiment 2, ANI infusion into the DS impaired the rightward transition in the “Start” response that normally takes place after switching from a 20 s to a 50 s temporal criterion. However, from our results, it is unclear how the acquisition of a “Start” response is mediated by activational changes in the DS. Indeed, in both Experiment 1 and 2, the differences observed with respect to the “Start” response in the DS-ANI group emerged (1) after an initial “Start” response was well-learned (i.e., following extensive training with one target duration) and (2) under conditions where uncertainty with regard to the expected time of reward delivery is present. This suggests that activational changes (including protein synthesis) in the DS can come on-line during transitional conditions in which the temporal contingencies that guided the “Start” response over an extended period need to be updated. Similarly, for example, the DS neuronal population changes on many dimensions as an animal learns to respond optimally on a T-maze. In this case, neurons gradually transition to preferably encode the beginning and end of the acquired response sequence in the T-maze. However, the specificity of the response profile for the beginning and end abruptly decreases during transitional conditions such as those reported in Barnes et al. (2005). In our experiment, ANI infusion
the PI timing procedure (Meck et al., 1984a,b; Lejeune et al., 1997; Rodríguez-Gironés and Kacelnik, 1999). Consequently, abrupt transitions from a 20 s to a 50 s temporal criterion would be unimpaired by ANI administration, whereas the second transition to an “intermediate” state appears to depend upon whether the adjustment is from a lower to a higher target duration or vice versa, the ratio of the times of reinforcement, the ratio of FI trials to unreinforced probe trials, and the subject’s previous experience with such transitions (Meck et al., 1984a,b; Lejeune et al., 1997; Rodríguez-Gironés and Kacelnik, 1999; Simen et al., 2011). Whether or not these transitions initially involve an abrupt transition to an “intermediate” state appears to depend upon whether the adjustment is from a lower to a higher target duration or vice versa, the ratio of the times of reinforcement, the ratio of FI trials to unreinforced probe trials, and the subject’s previous experience with such transitions (Meck et al., 1984a,b; Lejeune et al., 1997). Under the conditions used in the current experiment, it is expected that when rats are transitioned from a 20 s PI procedure to a 50 s PI procedure using a ratio of 50% FI trials to 50% unreinforced probe trials, they will initially exhibit a relatively abrupt transition of their “Start” times from those appropriate to the 20 s target duration to an intermediate value near the geometric mean of the 20 s and 50 s target durations. This step is thought to result from the subject comparing the remembered rate of reinforcement under the 20 s condition to the experienced rate of reinforcement under the new 50 s condition and not necessarily the rapid acquisition of a new target duration—which would be expected to occur in a more gradual manner. Such continuous updating of the subjective probabilities of reinforcement operates in parallel to the adjustment of “Start” and “Stop” response thresholds centered around specific times of reinforcement—much like the processing of variable-interval vs. fixed-interval schedules of reinforcement (Hinton and Meck, 1997a,b; Rodríguez-Gironés and Kacelnik, 1999). Consequently, in the current experiment it might be expected that when rats are transitioned from a 20 s to a 50 s temporal criterion the initial adjustment of the “Start” response to an intermediate state would be unimpaired by ANI administration, whereas the second step involving the acquisition of a “Start” response appropriate to the 50 s target duration would be impaired as this requires the acquisition of a new temporal criterion and associated response thresholds as opposed to the monitoring of changes in the overall rates of reinforcement and the adjustment of previously acquired response thresholds (Lejeune et al., 1997). Moreover, the abrupt jump to an intermediate duration isn’t influenced by hippocampal lesions and may be indicative of subjects using their old “Stop” threshold as a temporary “Start” threshold during this transition period (Meck et al., 1984a,b; Meck, 1988; MacDonald et al., 2007, 2009, 2011; Yin and Troger, 2011).

Our results also show that VS can contribute importantly to interval timing during learning. On the surface, it seems plausible that the “Stop” response ought to be mediated by behavioral and neural processes similar to those that operate during extinction (Myers and Davis, 2002; Lattal et al., 2006). Indeed, like extinction the acquisition of the “Stop” response appears to reflect new learning that is independent of the “Start” response. Moreover, several experiments have shown that VS lesions can lead to persistent responding during extinction of other operant tasks (Annett et al., 1989; Reading and Dunnett, 1991; Reading et al., 1991). However, in the case of the PI timing procedure, the extinction is partial. Reward is obtained on a random proportion (e.g., 50%) of PI trials and “extinction” of responding during the unreinforced probe trials exhibits a temporal gradient indicating that it is strongly guided by the memory of the target duration (see Kaiser, 2008).

Regardless of its relationship to extinction, the emergence of the “Stop” response might reflect learning that there is little value to sustaining a high response rate once the target duration has passed. In this regard, the VS may be involved in the “invigoration” of behavior (Cardinal et al., 2002; Robbins and Everitt, 2007), and may do so by dynamically tracking the value of the predictive signal O’Doherty et al., 2004; Atallah et al., 2007; Niv, 2007. Therefore, in order to terminate the ongoing response sequence, activation changes (possibly including protein synthesis) in the VS are required for the integration of time-related information with the “Stop” response. This allows the animal to reallocate its behavioral resources from current behavior(s) to subsequent one(s) in a flexible manner (Koob et al., 1978; Cools, 1980; Robbins and Brown, 1990).

The basal ganglia are known to receive rich and diffuse input from many brain regions, especially the midbrain dopaminergic area, which is sensitive to both the expected time of reward and changes in reward probability—two conditions that characterize each of our experimental contexts (Hollerman and Schultz, 1998; Fiorillo et al., 2003). Our results suggest a regional dissociation between how these different types of information might be integrated to influence behavior. Moreover, they are supported by a study that confirmed a double dissociation with respect to the role of the DS and VS on reconsolidation during the instrumental learning of a lever press (Hernandez et al., 2002). In this case, ANI impeded the acquisition of lever pressing following post-session administration into the accumbens core of the VS, the brain area primarily targeted in our experiments. In contrast, post-session ANI injections into the DS facilitated the acquisition of an instrumental response, which we speculate might arise from a decrease...
in the effect of task-irrelevant stimuli impinging on the DS. An intriguing possibility is that these two striatal regions might function as a competitive network early in learning, which ultimately promotes optimal behavioral allocation (Hernandez et al., 2002; MacDonald and Meck, 2004, 2005; Kimchi and Laubach, 2009; Kimchi et al., 2009; Yin et al., 2009; Smith et al., 2010). In the present experiment, ANI was microinjected before the sessions began so we can’t distinguish whether the effects we observed are targeting the memory consolidation, or a reconsolidation process. However, our experiments support and extend earlier findings by implicating a dissociable role for the DS and VS regarding when a response sequence should be initiated and terminated on the basis of a real-time temporal expectation.

The striatal beat-frequency (SBF) model of interval timing ascribes a mechanism for detecting event durations to medium spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2004; Buhusi and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998). Briefly, the SBF model posits that medium spiny neurons in the DS become entrained to fire in response to oscillating, coincident cortical inputs that become active at a particular duration. This timing model is particularly useful insofar as the striatal spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998). Briefly, the SBF model posits that medium spiny neurons in the DS become entrained to fire in response to oscillating, coincident cortical inputs that become active at a particular duration. This timing model is particularly useful insofar as the striatal spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998). Briefly, the SBF model posits that medium spiny neurons in the DS become entrained to fire in response to oscillating, coincident cortical inputs that become active at a particular duration. This timing model is particularly useful insofar as the striatal spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998). Briefly, the SBF model posits that medium spiny neurons in the DS become entrained to fire in response to oscillating, coincident cortical inputs that become active at a particular duration. This timing model is particularly useful insofar as the striatal spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998). Briefly, the SBF model posits that medium spiny neurons in the DS become entrained to fire in response to oscillating, coincident cortical inputs that become active at a particular duration. This timing model is particularly useful insofar as the striatal spiny neurons within the DS (MacDonald and Meck, 2004; Matell and Meck, 2005; Lustig et al., 2005; Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). These striatal neurons have a set of functional properties that place them in an ideal position to detect behaviorally relevant patterns of afferent cortical input (Beiser and Houk, 1998).

SUMMARY

Despite numerous claims that the inhibition of protein synthesis following ANI administration directly interferes with memory formation (Abel and Lattal, 2001; Alberini, 2008; Kwapis et al., 2011; Okamoto et al., 2011), there are compelling reasons to be more cautious in this interpretation. Gold and colleagues, for example, have argued that ANI induces temporary amnesia rather than inhibition of memory formation per se (Canal and Gold, 2007; Canal et al., 2007; Gold, 2008; Qi and Gold, 2009; Sadowski et al., 2011). Such amnesia might result, in part, from the disruption of neurotransmitter systems, including the increased release of norepinephrine, dopamine, and serotonin (Canal et al., 2007). When these unintended “side effects” were blocked, leaving the inhibition of protein synthesis largely unchanged, no reliable memory impairment was observed (Canal et al., 2007; Qi and Gold, 2009; Sadowski et al., 2011). As a consequence, some investigators have been skeptical of the role of protein synthesis in memory formation (for review see Alberini, 2008; Gold, 2008; Rudy, 2008; Rudy and Sutherland, 2008). Moreover, even when ANI is administered post-session and, therefore, thought to only affect processes involved in memory consolidation and not behavioral performance during the task, there is the potential to impair instrumental learning by drug-induced devaluation of the reward (e.g., food or sucrose pellet) which can affect behavioral performance in subsequent sessions as demonstrated by Jonkman and Everett (2009).

In the current study, ANI infusions were given prior to the start of a training session in order to maximize the chances of affecting learning and memory consolidation that would be expected to occur within the 2 h session. As a consequence, all of the concerns expressed above concerning performance variables are valid. The dissociation of timing performance for “Start” and “Stop” response thresholds as a function of ANI infusion into either the dorsal or VS suggests selective effects depending upon the type of response and brain region, as well as the degree of behavioral training and whether short-term or long-term memory processes are involved. In Experiment 2 for example, during the transition from a 20 s to a 50 s target duration, rats initially used a criterion that was intermediate between the two times of reinforcement. The determination of this intermediate target duration (which was never reinforced) presumably requires some sort of short-term memory process to be engaged. The observation that these early transitional processes were unaffected by ANI infusion, but that the more gradual acquisition of the “Start” response for the new 50 s target duration was selectively impaired by ANI infusion into the DS is consistent with a “protein synthesis” argument, i.e., that only those mechanisms responsible for the formation of long-term memories/response thresholds associated with the 50 s target duration were impacted by ANI infusion.

Taken together, our results point to a regional dissociation between how temporal information is used to guide separate components of a response sequence. Both the “Start” and “Stop” response thresholds serve as temporal boundaries that define transitions between two measurably different behavioral “states” occur. Clearly, there are many more questions as to why these transitions are different in the first place and require different brain structures. For example, it might be the case that the response topography of the state from which the transition leaves, or to which the transition leads represents an important factor. For some types of responses (e.g., discrete or sustained), response duration and the probability of response initiation can be differentiated by drugs that target D1 and D2 receptors, respectively (Gooch et al., 2007; Choi et al., 2005). Whatever the case, an intriguing possibility is that the dorsal and ventral striatal regions might function as a competitive network designed to respond to temporal information during the early stages of learning. The tracking of temporal regularities in the environment ultimately enhances adaptive behavior and may eventually lead to habit formation (Hernandez et al., 2002; MacDonald and Meck, 2004, 2005; Cheng et al., 2007a,c; Kimchi and Laubach, 2009; Kimchi et al., 2009; Yin et al., 2009; Smith et al., 2010). As such, most neurophysiological models of interval timing assume that target durations are encoded into a region’s neuronal
network in part through the modification of synaptic weights (Coull et al., 2011; Oprisan and Buhusi, 2011; Allman and Meck, 2012). In this way, it is important to explicitly test how and where these long-term modifications might be carried out at the cellular level by making use of what we know in other learning settings. In this regard, while there remains much to explore, these experiments using ANI infusions are among the first to isolate the roles of the dorsal and VS in temporal processing (see Meck, 2006b).

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