OPTOGENETIC INVESTIGATION OF STRIATAL CELL TYPE DURING TIMING BEHAVIOR

SOFIA LIMA DA SILVA SOARES

MESTRADO EM BIOLOGIA HUMANA E AMBIENTE

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Abreviations

AAV – Adeno-associated virus
ATR – All-trans retinal
BeT – Behavioral theory of timing
BG – Basal ganglia
BP – Band pass
ChAT – Choline acetyltransferase
ChR2 – Channelrhodopsin-2
CPu – Caudate-putamen
Cre – Cyclization-recombination enzyme
CRF – Continuous reinforcement schedule
D<sub>1</sub> – Dopamine type 1 (receptor)
D<sub>2</sub> – Dopamine type 2 (receptor)
DA – Dopamine
DIC – Differential interference contrast
DIO – Double-floxed inverted open reading frame
DNA – Deoxyribonucleic acid
EF-1α – Human elongation factor-1α
EI – Experienced interval
FI – Fixed interval
FS – Fast spiking
GABA – γ-aminobutyric acid
GP – Globus pallidus
GPe – Globus pallidus external
GPi – Globus pallidus internal
G-proteins – Guanine nucleotide-binding-proteins
HD – Huntington’s disease
HFT – Main dichroic beam splitter
IRI – Inter response interval
ITR – Inverted terminal repeat
LeT – Learning-to-time
L-dopa – L-3,4-dihydroxyphenylalanine
LoxP – Locus of X-over P1
LP – Long pass
LTS – Low-threshold spiking
MSN – Medium spiny neurons
ORF – Open reading frame
PBS – Phosphate buffered saline
PD – Parkinson’s disease
PFA – Paraformaldehyde
PI – Peak interval
PINP – Photostimulation-assisted identification of neuronal populations
PRP – Post-reinforcement pause
PV – Parvalbumin
SBF – Striatal beat-frequency
SET – Scalar expectancy theory
SFI – Serial fixed interval
SN – Substantia nigra
SNC – Substantia nigra pars compacta
SNr – Substantia nigra pars reticulata
STN – Subthalamic nucleus
TANs – Tonically active neurons
TTL – Transistor–transistor logic
YFP – Yellow fluorescent protein
WPRE – Woodchuck hepatitis virus posttranscriptional response element
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Resumo

Para conseguirem sobreviver num ambiente em constante mudança, é essencial que os animais consigam prever o desfecho de um determinado evento e/ou desenvolver comportamentos antecipadamente, no caso de esse mesmo evento voltar a acontecer. A cronometragem é um dos processos implicados nestes e noutros contextos de aprendizagem, já que confere ao organismo a capacidade de medir a duração do tempo entre eventos numa escala de segundos a minutos a horas.

Embora muitos modelos teóricos tenham sido desenvolvidos na tentativa de explicar a forma como o nosso cérebro codifica informação temporal, poucos são aqueles que tentam explicar a forma como os estímulos exteriores são integrados, o papel de diferentes tipos celulares nessa integração e ainda a influência do estado das redes neuronais no momento de integração. Apesar da sua extrema importância, o nosso conhecimento sobre como o cérebro representa a passagem do tempo é ainda muito limitado, parcialmente devido à falta de paradigmas comportamentais que acomodem electrotisiologia e também devido à incapacidade de identificar tipos neuronais específicos recorrendo a técnicas clássicas de electrotisiologia extra-celular. Em relação à cronometragem, diversos estudos indicam que os gânglios da base desempenham um papel fundamental no processamento de informação temporal. Anomalias no circuito dos gânglios da base estão intimamente ligados a doenças severas como a doença de Parkinson, doença de Huntington ou esquizofrenia, cujos pacientes demonstram incapacidades não só a nível motor, mas também na performance de tarefas de cronometragem. No entanto, as áreas envolvidas nestes processos, como é o caso do corpo estriado, possuem uma grande heterogeneidade a nível neuronal, tornando ainda mais complexo o estudo do papel de cada tipo de neurónio na representação temporal.

Existem pelo menos dois tipos de neurónios de projecção (MSNs) e quatro tipos de interneurónios no corpo estriado, e cada um deste tipo de células poderá desempenhar papeis distintos durante o processamento de informação temporal. Sabe-se que cada tipo de MSNs expressam predominantemente receptores para a dopamina do tipo 1 (D₁) ou do tipo 2 (D₂), formando subpopulações de neurónios D₁MSN ou D₂MSN distribuída no estriado. Sabe-se que os receptores D₁ e D₂ produzem efeitos antagónicos em cascadas intra-celulares, quando ligados à dopamina. Existem ainda estudos que sugerem um papel importante do receptor D₂ na capacidade de estimar intervalos de tempo. O desenvolvimento de linhas de ratinhos transgénicas torna a utilização deste modelo animal muito apelativa neste prisma, já que pode facilitar o desenvolvimento de técnicas para identificação in vivo destes diferentes tipos de
neurônios. No entanto, a real aplicabilidade destes animais a paradigmas comportamentais dinâmicos e complexos é questionada.

Pretende-se com este trabalho treinar ratinhos numa tarefa de cronometragem que acomode electrofisiologia, mas que seja mais dinâmica do que as tarefas clássicas utilizadas nesta área actualmente. Paralelamente, esperamos conseguir isolar diferentes tipos neuronais no corpo estriado de forma a que, a longo prazo, seja possível distinguir o tipo de célula cuja informação está a ser gravada durante experiências de electrofisiologia.

Treinámos ratinhos numa nova tarefa de cronometragem altamente dinâmica chamada Serial Fixed Interval (SFI). Esta tarefa foi desenvolvida no nosso laboratório, baseada num paradigma clássico no estudo de cronometragem, denominado Fixed Interval (FI) schedule. O equipamento da tarefa SFI é bastante simples, consistindo apenas numa alavanca de metal e um orifício por onde a recompensa (uma gota de água com açúcar) é entregue. Durante o período de treino e teste na tarefa SFI, os ratinhos são privados de água e estimam diferentes intervalos de tempo, tendo apenas a última recompensa como referência para estimarem a entrega da próxima recompensa. Após a entrega de uma recompensa, os ratinhos passam um determinado intervalo fixo (FI) de tempo sem terem a possibilidade de receber qualquer gota de água com açúcar, mesmo que pressionem a alavanca de metal. Após este intervalo fixo terminar, os animais têm 15 segundos durante os quais a primeira resposta na alavanca produz uma recompensa e o FI de espera repete-se. Após pelo menos 25 repetições do mesmo FI, que representa um bloco de tentativas, seleciona-se aleatoriamente uma nova duração, e durante uma sessão de duas horas são estimados vários blocos com intervalos de tempo diferentes.

Durante o decorrer da tarefa, os ratinhos desenvolveram um padrão de resposta que indica que estes animais estão a estimar intervalos de tempo. Depois de receberem uma recompensa, os ratinhos esperam um determinado intervalo de tempo antes de voltarem a pressionar a alavanca para tentar receber uma nova recompensa, e a esta latência para responder novamente chamamos PRP. Este padrão de respostas dos ratinhos é sensível à passagem do tempo, já que observamos que os PRPs variam no mesmo sentido da variação do FI entre cada bloco. Mais especificamente, obtivemos correlações significativas entre as PRPs e o intervalo anterior experimentado pelo animal, quer em sessões individuais ($R^2 = 0.45809$, valor $P < 0.001$) quer na análise de toda a população ($R^2 = 0.7813$, valor $P < 0.001$). Adicionalmente, a frequência com que os animais pressionam a alavanca desenvolve-se progressivamente mais lentamente quanto maior for o FI a ser estimado. Este perfil de resposta
foi acompanhado por rápidas curvas de aprendizagem, quer se tratasse da aprendizagem entre a passagem de um intervalo curto para um longo ou o inverso.

Paralelamente, começámos por tentar identificar dois tipos de neurónios do corpo estriado: os neurónios de projecção D₂-MSNs e um tipo de interneurónios que expressam parvalbumina (PV). Para isso, usamos duas linhas de ratinhos transgénicos: uma que expressa a enzima Cre recombinase (Cre) sob o controlo do promotor para o receptor D₂ (linha D₂-Cre); e outra que expressa Cre sob o controlo do promotor para a PV (linha PV-Cre). Estas linhas de ratinhos transgénicos foram sujeitas a micro-injecções de um virus adeno-associado (AAV) contendo um gene de fusão que codifica um canal iónico activado por luz, a channelrhodopsin-2 (ChR2), ligado a uma yellow fluorescent protein (YFP). Como a expressão de ChR2-YFP é dependente de dupla recombinação pela enzima Cre e as duas linhas de ratinhos transgénicos expressam Cre sob o controlo de dois promotores distintos, a acção do sistema cre-lox permite-nos expressar ChR2-YFP nestes dois tipos específicos de neurónios do corpo estriado. Através de técnicas de histoquímica, marcamos ainda os núcleos (no caso da linha D₂-Cre) e corpos celulares (no caso da linha PV-Cre) destes neurónios para uma melhor análise anatómica. Observamos que em cada linha transgénica estudada (D₂-Cre e PV-Cre), os tipos de neurónios identificados possuem as características morfológicas de neurónios D₂-MSNs ou de interneurónios PV, respectivamente. Observamos ainda a existência de pequenas inclusões fluorescentes presentes praticamente em todas as áreas onde existem células infectadas por virus. Algumas alternativas foram já pensadas de forma a evitar esta observação que parece ser um sinal de expressão elevada da proteína de fusão. Serão necessárias futuras experiências para avaliar até que ponto estas inclusões alteram as propriedades electrofisiológicas das células, assim como o efeito a expressão de ChR2-YFP por si só.

Com este trabalho, demos um passo importante para conseguirmos no futuro isolar subpopulações de neurónios e identificá-los durante gravações electrofisiológicas. Neurónios que expressem ChR2 serão identificados electrofisiologicamente recorrendo à técnica PINP (Photostimulation-assisted Identification of Neuronal Populations): uma vez estimulados com luz azul, a activação ChR2 causará uma despolarização da membrana do neurónio, e este produzirá um potencial de acção. Este efeito tornará possível identificar estes neurónios por técnicas extra-celulares de electrofisiologia enquanto os animais são testados na tarefa de SFI. Desta forma, poderemos estudar que tipo neuronal codifica que tipo de informação durante o processo de cronometragem. Esta informação poderá aprofundar o nosso conhecimento sobre os circuitos neuronais que estão na base deste processo fundamental. A compreensão da forma como integramos e processamos informação temporal poderá ainda ter enormes
vantagens no estudo de doenças neuro-degenerativas como as doenças de Parkinson e Huntington.

**Palavras-chave:** Cronometragem; interval fixo; gânglios da base; corpo estriado; ratinhos transgênicos; channelrhodopsin-2.
Abstract

The ability to time intervals in the range of seconds-to-minutes-to-hours, (interval timing) is a fundamental aspect of learning and behavior. Although many theoretical models have set out to address how the brain may process temporal information, little is known about the neural mechanisms that underlie this fundamental ability. Many studies indicate that the basal ganglia (BG) is the brain structure most involved in interval timing, but we still don’t know how the BG might process and encode duration information. Neurophysiological recording from single cells in a situation where a subject must access learned duration information provides a powerful tool to investigate interval timing mechanisms. However, information about neural cell type and connectivity within BG networks will ultimately be necessary to understand how timing information is computed, stored, and read out to guide behavior. We want to train transgenic mice in a dynamic paradigm and try to identify which signals are carried by which cell types during timing behavior. To achieve this, we trained mice in a new dynamic schedule, the Serial Fixed Interval (SFI) task. Animals showed a reliable response pattern that co-varied with the time duration being sampled and learned to adjust their response time rapidly in response to interval changes.

To ultimately identify recorded cell type during the SFI task, we used channelrhodopsin-2 to label two subpopulations of striatal neurons. We used adeno-associated virus to deliver channelrhodopsin-2 in to the striatum of two transgenic mouse lines in which the expression of the enzyme Cre was driven either by the parvalbumin promoter or by the dopamine type-2 receptor promoter.

Together, these results suggest that transgenic mice can be used in a highly dynamic timing paradigm, and that we may identify recorded cell types during such behavior, providing a powerful opportunity to study the neuronal circuit mechanisms of interval timing.

Keywords: Interval timing; serial fixed interval; basal ganglia; striatum; transgenic Cre mice; channelrhodopsin-2.
Introduction

Time and timing scales

Time is essential for innumerous daily life activities, like walking, speaking, playing music and learning. Such activities are possible because during our evolutionary history, we, like many organisms, have developed systems to deal with an environment where somewhat consistent variation of events across time is present, and producing behavior in such a manner is adaptive. This variation can occur at different scales. For example, the circadian rhythms are an adaptation to the regularity of days, interval timing for events within those days, and millisecond timing for proper perception and motor production. The mechanisms underlying circadian rhythms are perhaps the best understood of these three classes of timing. They operate roughly over the range of the 24-h light–dark cycle controlling the sleep-awake cycle, as well as metabolic and reproductive fitness. Millisecond timing is required for proper motor control, for generating and recognizing speech, or even for activities like dancing and playing music. Interval timing operates on a scale from seconds-to-minutes-to-hours and is involved in foraging, decision making and learning. We are surrounded by an environment in constant change, where important events can happen in regular intervals or at a predictable delay from other events. Having the ability to learn this regularity presents a great advantage and allows us to predict certain outcomes given a specific situation and develop anticipatory behaviors to prepare ourselves for the next time that a certain event will happen. For example, an animal in the jungle may have to wait still in a hidden place, for a certain amount of time, until predators are likely to have left an area before seeking out a food or water source. Therefore, interval timing allows a wide range of species to estimate short intervals of time which are essential for their survival.

Interval timing

Interval timing, defined as the perception, estimation and discrimination of durations in the range of seconds to minutes to hours, has been verified in organisms as diverse as insects, birds, fish, rodents, primates, human infant and adults. Traditionally, three types of behavioral protocols have been applied in the field of interval timing. These can be based on the estimation, production or reproduction of intervals of time. Both estimation and production protocols rely on verbal instructions, requiring the subject to convert the verbal representation of time durations into a performance of that representation. Because these protocols can lead to
confounds, using a reproduction procedure is a more reliable approach, and it can be applied to a wide variety of animal species, from humans to rodents\textsuperscript{14,15}. In a reproduction type behavioral protocol, the subject is presented with a given criterion duration and is then required to reproduce this duration. Usually, animals are deprived of either food or water, they are given a lever or a button to press and they are reinforced for that response (i.e. they receive a small amount of either food or liquid).

One of the most common schedules used to study interval timing is the Fixed Interval (FI) Schedule of Reinforcement. In this schedule, the first response of the subject after some FI of time since the last reinforcement has elapsed is reinforced. During that FI, no response is reinforced. Ferster and Skinner\textsuperscript{16} described that animals performing the FI schedule behaved in a characteristic sequential manner. First, just after receiving the reinforcer, animals stop responding and start engaging in self grooming or exploratory behaviors. Second, the animal's position and behavior are gradually oriented towards the response site. Finally, as the FI elapses, the animal's rate of response increases gradually and reaches the maximum near the moment when the reinforcer will be available again. Additionally, the three steps described above have different durations that are proportional to the duration of the FI.

Another classical schedule used to study interval timing is the Peak procedure\textsuperscript{17}, also known as the Peak Interval (PI) task. It is a modification of the FI schedule, where trials identical to those of the FI schedule are shuffled with a small percentage of so called probe trials. In these probe trials, no reward is presented to the animals, no matter when they respond. In the PI task, only probe trials are selected for data analysis. The subject's response profile in probe trials follows a normal distribution around the estimated FI and the width of this response distribution is proportional to the FI. Also, the manner in which both the mean and standard deviation of the response distribution co-vary is usually referred to as the scalar property, because the variability of timing performance scales proportionally to the interval being timed\textsuperscript{14}. It implies that, for example, one can estimate more precisely a 4 second interval than a 14 second interval. This property resembles Weber's Law, which is obeyed by many sensory dimensions\textsuperscript{18}. Weber's Law states that the relative increase in a stimulus needed to produce a noticeable difference is constant.

This scalar property of interval timing affects the system in such a way that it becomes increasingly less precise as the interval being timed lengthens and it applies not only to behavioral responses but also to neural activation\textsuperscript{19}.

Several models have been developed over the years, aiming to provide a possible explanation for the mechanisms by which the brain processes timed information. One of the
most popular views on interval timing in animals is the pacemaker-accumulator model, also called Scalar expectancy theory (SET)\textsuperscript{18}. This information-processing model is based on an internal clock model\textsuperscript{20} and proposes the existence of a Poisson-variable pacemaker that regularly emits pulses, which are temporarily stored in an accumulator. By the time of reward, the number of pulses present in the accumulator is stored in reference memory. The behavioral response results from the comparison of the ratio between the current time estimation, which is stored in the accumulator, and a sample selected from the distribution of previously estimated durations, which are represented as the number of pulses from previous reinforced readings stored in reference memory. This model presents an explanation for the scalar property of interval timing, since it suggests that the error generated during the accumulation of pulses is proportional to the criterion duration.

A second model, the Behavioral theory of timing (BeT), was developed by Killeen and Fetterman\textsuperscript{21}, based on the idea that reinforcement generates collateral behaviors that can be interpreted as a sequence of behaviors. BeT is a behavioral model that retains some aspects of SET, like the existence of a Poisson-variable pacemaker, but it proposes that behavior itself may serve as a signal for the passage of time, possibly mediating time discrimination. BeT suggests that each behavior is associated with a particular underlying state and that the transitions between one state to the next occur thanks to pulses that a Poisson-variable pacemaker produces. The speed of this pacemaker depends on the rate of reinforcement in a given experimental context, so that increases in reinforcement rate lead to an increase of the speed of the pacemaker.

Using BeT as a basis, Machado\textsuperscript{22} developed another behavioral model which he called Learning-to-time (LeT). His model consists of three major components: a sequential chain of behavioral states (similar to BeT), a vector of associative links connecting the behavioral states to the operant responses, and the actual operant responses. LeT hypothesizes that when the criterion duration starts, only the first behavioral state is active, but with the passage of time, the activation shifts from the previous to the next behavioral state in the sequence. Additionally, LeT proposes that each behavioral state is paired with the operant responses, and that the level of pairing varies during training, decreasing during extinction and increasing during reinforcement. Therefore, the strength of an operant response at a given moment is a result of the combination between which state is predominantly active at that moment and how strong is the association between that state and the response.

More recently, another model was developed, proposing that activity from loops involving the thalamus, cortex and striatum could be involved in timing. This new model was developed
taking the Striatal beat-frequency (SBF) computational model23 as a reference, but it was slightly modified and proposes specific roles for certain brain areas, trying to build a more biologically plausible model. This SBF based model suggests that some oscillatory cortical neurons can cause simultaneous activation of striatal projection neurons in the basal ganglia24. During a timing task, these striatal projection neurons would constantly compare the current pattern of activation of cortical neurons with the pattern present when reward is collected. The SBF based model suggests that dopamine plays a role in synchronizing these oscillatory cortical neurons at the onset of a trial as well as maintaining them at a fixed frequency of oscillation throughout the criterion interval. This hypothesis relies on biological observations such as that dopaminergic projections to the striatum are active in different ways during a timing task. At trial onset, they discharge a burst response, show a sustained activity during the criterion interval and display another burst response at the time when reward was expected25. Interestingly, the scalar property of interval timing was demonstrated in simulations using the modified SBF model24.

The models previously mentioned focus on greater extent on the integration of external stimuli. But, within complex neuronal networks, the response of a population of neurons may be determined both by the characteristics of the input stimuli and by the internal state of the network26. For example, a neuron can respond to a tone as a result of its specific frequency but also because of the strength of each of its synapses at that moment (which vary on a rapid timescale) or also because that neuron may be receiving additional internally generated excitatory and inhibitory inputs at the same time. Attempting to account for a role of the characteristics of external stimuli and also of the internal states during information processing within neural circuits, a new model has been developed. The State-dependent model was developed based mainly on cortical networks and accounts for both theoretical and experimental studies27,28. It suggests that both spatial and temporal information are processed taking into account the features of the incoming stimuli and the internal dynamic state of the neural network. This internal dynamic state can be divided in two components. The first component, called “active state”, comprises the ongoing pattern of activity in the network. The second component, known as the “hidden state”, includes the collection of time-dependent properties that are affected by prior activity and that can have an effect on whether, after receiving some input, a neuron fires or not (e.g. short- and long-term synaptic plasticity29,30). Therefore, the real internal state is created by combining the active and the hidden states, since both can influence the response of a local cortical network to a stimulus.
Regarding timing, state-dependent models predict that, for example, when two short tones are presented with a certain interval between each presentation, this inter tone interval can be encoded in the response of the network to the second tone presentation\textsuperscript{27}. A few studies in cortical areas, cerebellum as well as in the hippocampus have tested predictions that arise from State-dependent models. For example, in the cerebellum, the pattern of response of granule cell populations to a constant stimulus may depend on the stimulus features but also on the current state of the connections within that cerebellar network\textsuperscript{31,32}.

Data from our lab (personal communication) indicates that, during a Serial Fixed Interval (SFI) task, striatal cells are active at different times over tens of seconds, when their response profiles are aligned on reward delivery. Such a response profile could be helpful to encode time information, and is consistent with the predictions of State-dependent or SBF models, but not with the predictions of a pacemaker-accumulator model.

**Neurobiological systems involved in timing**

Many efforts have been made in the last decades in order to expand our understanding of the neurobiological systems that are involved in timing, and many brain areas have been implicated in these distinct timing systems. Regarding interval timing, researchers have been studying the basal ganglia (BG) and cerebellum, because there is evidence that these brain areas are involved in the integration of timed information. BG have been associated with interval timing\textsuperscript{24}, and the cerebellum has been charged with millisecond timing\textsuperscript{33,34}. This general attribution rises from studies that suggest that interval timing depends on the intact striatum\textsuperscript{35-37}, combined with studies where cerebellar lesions failed to affect performance on interval timing tasks\textsuperscript{38}. For example, a 1996 study showed that cerebellar lesions in rats led to an impaired performance during a duration discrimination task, when the stimulus range was centered at 500 ms, but did not affect performance when the range was centered at 30 sec\textsuperscript{39}. Also, because cerebellar injury fails to affect the scalar property of interval timing, it is presumed that the cerebellum is not fundamental for appropriate interval timing\textsuperscript{40}. Imaging studies also point to the BG as one of the players in timing behavior, together mainly with cortical areas. Jueptner and colleagues\textsuperscript{41} reported regional cerebral blood flow increases in the BG during a timing task, using positron emission tomography. Additionally, studies using ensemble recording techniques in animals or functional magnetic resonance imaging in humans, have also suggested a role for the BG during timing behavior\textsuperscript{42-44}.

Furthermore, patients with disorders that involve meso-striatal dopaminergic pathways, such as Parkinson's disease (PD)\textsuperscript{35,36} Huntington’s disease (HD)\textsuperscript{45} and schizophrenia\textsuperscript{46-48},
display impaired performances during an interval timing task. Patients’ performance is improved by L-dopa (L-3,4-dihydroxyphenylalanine) medication or by stimulation of the subthalamic nucleus, two standard treatments to alleviate PD symptoms. In PD, degeneration of the nigrostriatal dopaminergic projections occurs. This leads to low levels of dopamine (DA) in the striatum and the patient’s interval timing abilities are impaired in a variety of ways\textsuperscript{36,38}. Patients with PD, when performing a task where they had to time two different durations without medication during the training process, tend to overestimate both intervals. On the contrary, when the testing phase occurs without medication, the two remembered time intervals are estimated in such a way that overestimation of the shorter and underestimation of the longer one occurs. Also, the patient’s distributions of timed responses display the scalar property when medicated with L-dopa, but not when tested without medication.

DA is a major neuromodulator the activity of the projection neurons in the striatum\textsuperscript{49,50,51}. A specific type of DA receptor expressed in striatal neurons, the D\textsubscript{2} receptor, has been the target of many studies regarding its role on striatal modulation during interval timing. Meck\textsuperscript{52} assessed the effects of several D\textsubscript{2} receptor antagonists on rats performing an interval timing task and found that the dose of the drug needed to impair interval timing ability was negatively correlated with the drugs affinity for the D\textsubscript{2} receptor. Additionally, studies using a transgenic mouse line that selectively overexpresses the D\textsubscript{2} receptor in striatal neurons suggest that striatal D\textsubscript{2} overexpression impairs interval timing mainly by decreasing the animal’s motivation\textsuperscript{7,8} and by its impact on working memory and/or sustained attention. Together, these results suggest an important role for striatal D\textsubscript{2} receptor signaling in interval timing.

Although BG and the dopaminergic system seem to be major players in interval timing behavior, we still don’t know how the signals that are carried by neurons in the BG network processes temporal information to support the broad range of animals’ timing behavior. We propose to add a new layer of information on this subject, which is largely unavailable to neurophysiologists until now. Specifically, we aim to answer the question: what signals are encoded by which cell type during timing behavior?

**Basal ganglia organization**

The BG are a group of nuclei that are profoundly important clinically. Lesions of different cellular components of the BG lead to devastating neurological disorders including PD and HD\textsuperscript{55}, and BG dysfunction is associated with a range of neuropsychiatric disorders such as schizophrenia\textsuperscript{22} or obsessive-compulsive disorder\textsuperscript{23}. 
The position of the BG in the circuity of the brain gives a clue to their numerous functions. The BG occupy the center of the forebrain and it is composed by evolutionarily conserved motor nuclei that form loops with areas such as the cerebral cortex, thalamus and brainstem. Therefore, they are in a prime position to influence motor planning and even cognitive behaviors. A general canonical organization of the BG consists of four structures: the striatum, the globus pallidus (GP), the subthalamic nucleus (STN) and the substantia nigra (SN)\textsuperscript{58}.

The striatum is the main input structure of the BG (Fig1). In primates, the striatum is divided in caudate nucleus, putamen and ventral striatum. However, in rodents, the caudate nucleus and the putamen are fused in one structure often denominated CPU (caudate-putamen) and the ventral striatum is denoted as the nucleus accumbens.

The GP (divided in an external and an internal segment, GPe and GPI, respectively) is the site of therapeutic lesions (pallidotomy) and deep-brain stimulation procedures that were historically used to alleviate PD symptoms.

The SN is subdivided in two regions (pars compacta and pars reticulata, SNC and SNR, respectively) and it is a critical nucleus to BG function. DA-synthesizing neurons in the SNC degenerate in PD, and SNR, together with the GPI, represent the major output nuclei of the BG (Fig1). Finally, the STN is a key structure controlling GP function, and is now a favored site for deep-brain stimulation in the treatment of PD.

Glutamatergic layer 5 neurons from nearly all areas of the cortex project to the BG, which in turn provide output to brain systems that are involved in the generation of behaviors (e.g. output nuclei in the basal ganglia project to thalamic nuclei that in turn send inputs to frontal cortical areas involved in the planning and execution of movement; midbrain areas like the superior colliculus, which is involved in the generation of eye movements and orienting responses). It is hypothesized that BG receives inputs from these areas and acts as a filter, selecting activity and sending the result of its selection back to other brains systems. It is further thought that
reinforcement based plasticity occurs in the BG, and that this plasticity may somehow influence the input selection process based on previous experience\textsuperscript{58}.

In a classic view, information from the striatum can travel through the nuclei of the BG by two different parallel circuits, the direct and the indirect pathways (Fig2). Neurons within the striatum can project directly to the output nuclei of the BG (direct pathway), or instead, they can project to other nuclei in the BG that thereafter indirectly connect the striatum and the output nuclei of the BG (indirect pathway)\textsuperscript{55}.

These signaling pathways are regulated by DA in the striatum, and they have been the subject of intense study since it was discovered that the loss of DA in the BG leads to the psychomotor symptoms of PD\textsuperscript{59}.

**Fig2 – Direct and indirect striatal projection pathways in sagital schemes of rodents. A – Direct and indirect pathway rise from different poplations of striatal neurons. Direct projecting neurons provide an axon with collateral to the globus pallidus external (GPe) globus pallidus internal (GPI) and substantia nigra pars reticulata (SNr). Indirect projection neurons project to the GPe and are indirectly connected to the GPI and SNr through connections involving GPe and subthalamic nucleus (STN). Figure adapted from\textsuperscript{58}.**

**Anatomy and histochemistry of striatal neurons**

The primary input nucleus of the BG is the striatum. Although generally implicated in control of motor actions, recent studies account for a more numerous subdivision of functions within striatal areas. The dorsal striatum has been implicated in essential aspects of decision-making, sensorimotor, cognitive, and motivational information\textsuperscript{60-62}. Interestingly for interval timing, studies also suggest that neurons within dorsal striatum respond to sensory events related to the task in question, becoming active before task-related motor behaviors and, in addition, they display tonic activity until the time when reward was expected to be delivered\textsuperscript{63,64}.

The striatum is the largest nucleus within BG and it completely lacks glutamatergic neuronal cell bodies. Instead, most neurons with cell bodies in the striatum release γ-aminobutyric acid (GABA). Neurons within striatum have been characterized anatomically, histochemically, and physiologically\textsuperscript{65,66}. Regarding their anatomic characteristics, striatal neurons can be either medium spiny projection neurons (MSN) or aspiny interneurons. MSNs are the principal cell type of the striatum, since they account for more than 95% of striatal
neurons. These GABAergic neurons are the major receiver of extrinsic input and they are the only output neurons of the striatum. Glutamatergic inputs from cortex and thalamus to the MSNs terminate predominantly on dendritic spines. In addition, the MSNs are a main target of dopaminergic neuron axons from the ventral tegmental area and from the SNc, that form synapses on MSNs dendrites and spine necks.

The MSNs can be generally divided into two subpopulations. This division is based on their axonal projection targets, the expression of genes for certain peptides, and the expression of DA receptors. These two subpopulations of MSNs give rise to the two parallel BG circuits, the direct and indirect pathways. About half of MSNs send projections directly to the output neurons of the BG in the GPi and SNr. For that reason, they are called striatonigral MSNs and give rise to the so called direct pathway of the BG circuitry. The other half of MSNs, also called striatopallidal MSNs, belong to the indirect pathway because they don’t send direct projections to the BG output nuclei. Instead, MSNs from the indirect pathway send their projections to the GPe neurons, which then project to the STN, and then to SNr and GPi. The direct and indirect pathway model of the BG has been the target of many studies, more specifically studies regarding its role on motor control. It has long been thought that excitation of the direct pathway would facilitate movement and activation of the indirect pathway would inhibit movement.

Although the validity of this hypothesis has been challenged over the years, a recent study brought strong evidence in favor of this classical model. In this work, Kravitz and colleagues performed bilateral excitation of striatopallidal MSNs in transgenic mice using optogenetic methods and observed that this protocol induced a parkinsonian state, identified by increased freezing, decreased locomotor initiations and bradykinesia. Conversely, activation of striatonigral MSNs caused a decrease in freezing and an increase in locomotion. Additionally, in a mouse model of PD, activation of the direct pathway rescued the impairments displayed in freezing, bradykinesia and locomotor initiation.

MSNs belonging to either the direct or indirect pathway show selective expression of certain peptides and receptors for DA. Studies using in situ hybridization histochemistry combined with retrograde labeling of striatonigral neurons suggest that striatonigral MSNs express substance P, dynorphin and the DA type 1 (D₁) receptor, and therefore are also referred to as D₁ MSNs. The striatopallidal neurons express enkephalin and the D₂ receptor, and for that reason are also known as D₂ MSNs. Although initial studies using RT-PCR claimed that there was considerable co-localization of these DA receptor subtypes, improvements of this technique confirmed this partial division of MSNs. Within striatonigral and striatopallidal MSNs, there was found a 10%–20% co-localization of D₁ and D₂ receptor transcripts. Nonetheless,
immunohistochemical studies indicate that the co-localization between D1 and D2 receptors is only about 1%\textsuperscript{75}. This suggests the existence of major posttranscriptional control of the expression of DA receptors in MSNs.

Considerable progress has been made in characterizing the DA receptors in the striatum and the brain in general. Molecular cloning studies have helped to discover at least five DA receptor genes (D\textsubscript{1}, D\textsubscript{2}, D\textsubscript{3}, D\textsubscript{4}, D\textsubscript{5})\textsuperscript{76-80}. These receptors can be further grouped on the basis of their affinities for classical dopaminergic ligands and their coupling to intracellular signaling cascades into D\textsubscript{1} (D\textsubscript{1}, D\textsubscript{5}) and D\textsubscript{2} (D\textsubscript{2}, D\textsubscript{3}, D\textsubscript{4}). Both D\textsubscript{1} and D\textsubscript{2} receptors couple to adenylate cyclase via G (guanine nucleotide-binding)-proteins. D\textsubscript{1} receptors stimulate adenylate cyclase via stimulatory G\textsubscript{s}-proteins, whereas D\textsubscript{2} receptors inhibit adenylate cyclase via inhibitory G\textsubscript{i}-proteins\textsuperscript{81}. Although different cell types express distinct DA receptor subtypes, every cell type within striatum expresses DA receptors.

Anatomically, besides MSNs, there is another category of striatal neurons, the interneurons. They represent about 3-5% of striatal neurons and include cholinergic interneurons and several types of GABA-releasing interneurons\textsuperscript{82,83}. Within striatal neurons, only about 1-2% are cholinergic interneurons\textsuperscript{84}. These neurons are generally characterized by a large soma, often >50µm long and they stain positively for choline acetyltransferase (ChAT). Immunohistochemical analysis suggest that cholinergic interneurons express both D\textsubscript{2} and D\textsubscript{5} receptors\textsuperscript{85,86}. The GABAergic interneurons seem to express mainly D\textsubscript{5} receptors\textsuperscript{85} and can be divided into at least three groups based on their distinct histochemical and physiological properties\textsuperscript{82}. Histochemically, striatal GABAergic interneurons can be subdivided into: (a) parvalbumin (PV)-positive; (b) somatostatin-, neuropeptide Y-, and nitric oxide synthase-positive; and (c) calretinin-positive\textsuperscript{83}.

Physiologically, these three groups of striatal GABAergic interneurons can be further separated in at least two different types based on the firing patterns that they exhibit\textsuperscript{82}. While PV-positive neurons display rapid and continued firing rates post current injection, somatostatin-positive interneurons display lower firing rates and plateau potentials. Thus, PV-positive neurons are alternatively known as fast spiking (FS) interneurons and somatostatin-positive interneurons are known as low-threshold spiking (LTS) interneurons. Calretinin-positive interneurons appear to share some characteristics of LTS interneurons, but further physiological classification is required to confirm this suggestion\textsuperscript{87}. Although few in number, striatal GABAergic interneurons play a major role in regulating spike timing in the MSNs, mainly through feedforward inhibition\textsuperscript{87}. Like MSNs, striatal interneurons receive glutamatergic input from cortex and thalamus. Conversely, their output is directed primarily to MSNs and other
interneurons, thus forming microcircuits capable of modulating striatal output. Therefore, striatal interneurons may produce disproportionately strong effects despite being numerically in minority. As for the physiology of the cholinergic interneurons, they display spontaneous activity under physiological conditions, and therefore are also referred to as tonically active neurons (TANs). They exhibit significant hyperpolarization-activated currents, but they display a pause in their tonic firing in the presence of salient cues, including reward\textsuperscript{88,89}.

Though the basic physiology of striatal cell types has been characterized, a more detailed description of these neurons has been difficult. To distinguish striatonigral and striatopallidal MSNs \textit{in vitro} without recurring to post hoc analysis has been an unsuccessful attempt, since they share most anatomical and electrophysiological features. In contrast, striatal interneurons are more easily identifiable during electrophysiological recordings\textsuperscript{82}. Despite this, to target interneurons for electrophysiology is also challenging, since they represent a very small portion of the total number of striatal neurons and they are similar to MSNs under the light microscope.

Although researchers have developed new methods to visualize distinct neuron types, within striatum, for cellular and synaptic electrophysiology\textsuperscript{90}, these provide advances mainly for isolation of these different neuronal types \textit{in vitro}. Because we are interested in understanding which signals are carried by each striatal neuronal type during interval timing behavior, we want to be able to distinguish these different subpopulations for electrophysiological recordings \textit{in vivo}, while mice perform an interval timing task.

**Optogenetics**

Using classical extracellular recording techniques, it appears possible to distinguish just two general populations of striatal neurons. Berke and colleagues\textsuperscript{91} recorded from striatal neurons in freely moving rats and found that, although the majority of units recorded were probably MSNs based on their firing pattern, a second neuronal population displayed characteristic that were attributed to fast-spiking GABAergic interneurons. These include tonic activity, brief waveforms, and high-frequency bursts. Despite these advances, we cannot yet electrophysiologically distinguish between the two types of MSNs and between all the interneuron types in the striatum. These different neuronal subpopulations may play different roles during timing behavior. Isolating each neuronal type within striatum in behaving animals would lead to a deeper understanding of the BG circuitry and also of the signals that are carried by components of that circuitry. But because of the heterogeneity within striatal neurons (and within whole brain), resolving the role of particular neuron types in information processing poses fundamental challenges. Therefore, a noninvasive genetically based method, with a high
temporal resolution, to control neural activity may elucidate the spatial and temporal of activity encoded by particular neuronal populations that drive circuit dynamics and behavior.

Optogenetics may be the way to achieve the selectivity that classical extracellular recording techniques fail to provide. Optogenetics is a recent term that is used to describe an approach of using new optical methods, which operate at a high temporal resolution, for probing and controlling neurons that are genetically targeted within intact neural circuits. The introduction of optogenetics allows researchers to control not only electrical properties within targeted cell types, but also biochemical events, and all of this can be done in living, freely moving animals. Several of the methods that have been developed allow control of neuronal activity over timescales of seconds to minutes. Although this presents an advantage, a method with a higher temporal resolution would be required in order to enable control of individual spikes. Such a strategy was developed in 2005 and it uses a light-activated cation channel called channelrhodopsin-2 (ChR2) derived from the alga Chlamydomonas reinhardtii. ChR2 is a protein constituted by seven-transmembrane domains and an all-trans retinal (ATR) molecule located at the core, functioning as a photosensor. When illuminated with ~470 nm blue light, the ATR molecule undergoes isomerization and this event triggers a conformational change that ultimately leads to the opening of the channel’s pore. Following just 50 µs of illumination, ChR2 allows a depolarizing current to be evoked, and therefore neurons can be activated at the temporal precision of a single action potential. Moreover, engineered ChR2 fusions with several classes of fluorescent proteins have been expressed in both isolated neurons as well as in neural circuits.

While optogenetics has been widely used as a way to control neuronal activity, investigators have also been trying to employ it in the identification of different neuronal populations in vivo. In a recent study, Lima and colleagues developed a method of labeling neurons to be further identified during in vivo electrophysiological recording, called PINP (Photostimulation-assisted Identification of Neuronal Populations). Following expression of ChR2 in restricted neuronal subpopulations, neurons labeled with ChR2 can then be detected electrophysiological in vivo, because when illuminating a certain brain region with a short flash of blue light, neurons expressing ChR2 at their membrane will fire a short latency reliable action potential (Fig3B). Consequently, that specific group of neurons is distinguishable for electrophysiological recordings, and one can then study which signals are carried by that ChR2-tagged neuron. PINP was applied in mouse cortical FS interneurons and in excitatory neurons in the rat auditory cortex. ChR2 delivery was achieved by using, respectively, adeno-associated virus (AAV) and herpes simplex virus 1 for mouse and rat neurons.
Our ultimate objective is to train mice on a dynamic timing task and simultaneously record electrophysiologically from striatal neurons. Moreover, we aim to functionally tag striatal neuronal types, in order to identify from which neuron we are performing the recordings (Fig3). This will enable us to analyze which timing signals are carried by each neuronal type. To achieve this, we will selectively express a ChR2-YFP (yellow fluorescent protein) fusion in 4 different types of striatal neurons: D1-MSNs, D2-MSNs, PV+ interneurons and ChAT+ interneurons. Expression of the fusion gene will be Cre (cyclization-recombination enzyme) dependent and delivered by viral vectors in 4 different transgenic mouse lines, in which Cre expression is driven by a given promoter that provides cellular specificity, therefore restricting ChR2-YFP expression to one of the 4 neuronal types mentioned. We will then use PINP to identify each neuronal type in each mouse line, while they perform the interval timing task (Fig3).

**Fig3 – Diagram of an example of the method for applying PINP to identify neurons tagged with ChR-2 in a PV-Cre mouse line during timing behavior.**

A - In a PV-Cre transgenic mouse implanted with an electrode (grey thin line) and cannula (blue thin line), different units display different response patterns during the timing task. We don't know which cell types are responsible for each pattern. B – With the introduction of an optic fiber through the cannula (blue thin line), when flashes of blue light (blue thick bars) are sent through the optic fiber, a reliable response from the ChR2-tagged neurons will be recorded. Since ChR2-YFP expression is Cre dependent, and this mouse line only expresses Cre in PV interneurons, we can now assume that a PV interneuron is responsible for that specific pattern of response. C – Now, we identified the single unit tagged with ChR-2 as a PV interneuron and recordings from that single unit can be performed on mice while they perform the SFI task.
We predict that in general, different types of striatal neurons will display significant differences in their response profile during a timing task. We think that, because of DA’s antagonist effect on D₁MSNs and D₂MSNs, the response profile of D₁MSNs may tend to increase around the time of reward, and that D₂MSNs response profile may tend to decrease to reward. Also, due to the fact that the TANs cholinergic interneurons display a reduction in their tonic activity in response to salient cues like a reward, we think this class of interneurons will display a similar response profile during the SFI task.
Materials and Methods

Subjects

All procedures were carried out in accordance with European Union Directive 86/609/EEC and National Institute of Health guidelines on animal care and experimentation. Eight C57BL/6 male mice (Jackson Laboratory), aged 8-10 weeks were housed 2 per cage and kept under a 12 h light/dark cycle, at 22°C, relative humidity of 50%. Two lines of transgenic animals (kind gift from Rui Costa) expressing Cre under the promoter for D₂ receptor (D₂-Cre) or the promoter for PV (PV-Cre) were treated as above, but housed one per cage. Experimental procedures were performed during the light phase of their cycle. Each animal was identified by a system of ear marks and named with different numbers for selection, behavior and data analysis purposes.

Apparatus

Behavior training and testing procedures were performed in a 15 cm*30 cm*36 cm polypropene chamber with a covered ceiling to minimize distraction of the animal inside the chamber.

A lever (MedAssociates, Inc.) and a nose poke (Island Motion, Corp.) were placed in one of the widest walls of the chamber, as illustrated in Fig4. Because the lever was placed above the nose poke, the animal could not press the lever (Fig4A) and introduce its snout in the nose poke (Fig4B) at the same time. This design (and the small dimensions of the chamber) also
reduced the occurrence of other behaviors.

The nose poke was equipped with a small tube (from where the reinforcer was delivered), an infra-red light and a sensor placed just before the hole, to detect headentries (Fig 5). Rubber tubings connected a 1mL receptacle to a valve (Island Motion, Corp.) and the valve to a metal tube that was placed in the nose poke hole.

**Fig5 – Top view of the nose poke components.** It consists of an infra-red light and sensor, placed just before a small tube from where water is delivered. **A** – View of the nose poke without interruption. **B** – View of the nose poke with interruption, when the mouse approximates to collect the reinforcement.

The majority of the behavioral procedures were automated and did not require an active participation of the experimenter. The Bcontrol Behavioral Control System, developed by BrodyLab at Howard Hughes Medical Institute (Princeton University), was used to control the inputs (lever presses and headentries) and the output (sucrose 10% (m/v)). This system consists of two computers, one running Lynux Fedora OS with Real Time Linux Machine application and the other running Windows XP OS and MatLab 2007b (MathWorks, Inc.) application with a BControl library. Within the BControl system, each state had a specific relationship between inputs from and outputs to the behavioral chamber.

**Behavioral Procedures**

Mice were weighed for five consecutive days, while given *ad libitum* food and water, and their weights were averaged to calculate the baseline body weight. After those five days, water was removed from their home cages and they were given ~1mL of sucrose solution (10% (m/v)) every 24 h until they reached ~87% of their baseline weight. Mice were given 1mL of sucrose solution per day to maintain ~87% of their total body weight, the percentage at which they were maintained for behavioral training and testing in order to promote motivated behavior. The
sucrose solution at 10% (m/v) was the standard reinforcer for all the experiments and, since it is isotonic, there was no need to provide another source of water.

The training period was divided in 3 steps. First, mice were placed in the behavioral chamber and submitted to a continuous reinforcement schedule (CRF). In the CRF, every time the animal emitted a response, this is, pressed the lever, 3.5µL of sucrose solution were delivered in the nose poke. In this first step, animals were trained for 2 sessions every day. Each session lasted 30 min or until the animal received 100 rewards. Naïve mice learned to press the lever by repeatedly reinforcing approximations to the lever, until the actual lever press was achieved. Sometimes, treats were placed over the lever to lure the animal to emit a behavior closer the lever, in order to speed the learning process. Only this first step required an active participation of the experimenter and then automation began. When animals reached 200 rewards in one day, they were moved to the second training step, where the SFI training began. Only few aspects of the SFI changed in between training steps. In all training steps, animals were placed in the behavior chamber and the first lever press was rewarded and initiated the actual task. This means that the animal controlled when the task began. The lever presses that occurred before some FI of time after the last reward were not reinforced (Fig6A). After this FI elapsed, animals had limited time to respond and get a reward, or the trial restarted and a new FI during which there was no possibility to acquire the reward would be presented (Fig6A). This time to respond after the end of the FI was different for each training step. In all SFI sessions, trials were grouped in blocks of at least 25 trials (Fig6B). Each block had a different FI, which was randomly selected from a sequence of 5 intervals (12, 24, 36, 48 and 60 sec). In every training step, after at least 25 trials in a given block elapsed, the script checked if mice completed 80% of the last 15 trials. If mice achieved this precision criterion, the FI changed and a different block started. If not, mice were kept on the same block until they reached the precision criterion. The sessions ended when mice performed 5 block switches or 2 h have elapsed since the beginning of the session.

During the second step of the training period, mice had unlimited time to respond after the FI elapsed.

The third training step was the same as the second one, but now the animal had only 25 sec to respond after the FI elapsed in order to get the reward or the trial was restarted. When the animals performed at least 5 block switches in two consecutive training sessions, the training step moved forward, from second to third step and from third step to the data collection sessions.
**Fig6 – Illustration of the SFl task structure.** **A** – Diagram of the structure of a single trial. In blue is represented the reward of the last trial that is used as cue in the actual trial. The FI, represented in white, lasts from 12 to 68 seconds. During this FI, no lever press produces a reward. After the FI elapses, mice have 15 seconds to press the lever in order to get the reward. Time to respond is represented in green. **B** – Diagram of a session. First trial, represented in green, does not have a time limit. All the other colors represent the same parameters as described in **A**. The first lever press in the first trial will produce a reward and random selection of the first FI block of trials takes place. If there is no response during the time to respond, the trial is restarted. The FI changes when the animal reaches certain criteria.
During the data collection sessions, animals had 15 sec to respond after the FI elapsed in order to get a reward. Also, animals still had to perform at least 25 trials per block and get at least 80% of the last 15 trials correct to switch blocks. Sessions lasted 2 h and only those in which the animals performed at least 5 block switches were used for data analysis.

Data analysis

Sessions in which animals performed at least 5 block switches were selected for data analysis. In this set of data, lever presses were represented by the moment of occurrence in time (milliseconds) since the start of the session. We wanted to perform a trial by trial analysis, and therefore we transformed the absolute times into relative times, by subtracting those lever press times by the trial start time, also in milliseconds. Because mice needed a cue to measure the FI, and this cue was the reward from the previous trial, only data from reinforced trials that were also preceded by a reward were selected for data analysis. In addition, only trials with more than two lever presses were selected for analysis.

We also used a heuristic in order to further select responses from the relative times data set. A heuristic is an experience based method of solving problems, to determine something that cannot be formally proven. In our analysis, we developed a heuristic that comprised two points. First, we ignored lever presses that occurred on the first 2 sec after the reward was available because the immediate responses in the beginning of each trial were, most likely, an effect of high frequency responses from the previous trial. Also, mice always spent more than 2 sec collecting the reward. Second, we observed that mice occasionally pressed the lever in a manner that didn’t seem to reflect the true response. These occasional presses often occurred isolated in time, maybe reflecting an exploratory behavior inside each trial. In order to avoid these isolated lever presses, we measured the inter response interval (IRI), this is, the time between each lever press. This was done by subtracting the time of every lever press from the time of the next one and then removing from the analysis all the lever presses in which the IRI was over the percentile 85.

From the data set filtered with the heuristic, we measured the post reinforcement pause (PRP). The PRP is the time since the start of the trial until the first response of the data set filtered by the heuristic.

For single session analysis, we grouped blocks that had the same FI and calculated the average frequency rate of response within the groups of trials with the same FI. The rate was generated by taking the mean of responses in every 0.5 ms bin and then convolving this set of data with an 11 kernel size Gaussian. This step smoothed the average rates curves but,
because artifacts at the edges of the smoothed curves were generated by the convolution, we padded the beginning and the end of our data with vectors with the same size as the kernel. All the elements of the padding vectors had the same value (the first value of the data set for the beginning or the last value of the data set for the end of the curve).

Timing learning curves were generated by calculating Z-scores. These were calculated by taking the PRPs data from the first 15 trials after block switch in each block. PRPs mean and standard deviation were calculated using the distributions of the PRPs in each block, after excluding the first five trials post block switch. We then subtracted this mean from each of the first 15 trials PRPs and divided the result by the standard deviation within each block. We made 3 independent analysis: for all block switches; for shorter FI to longer FI block switches and finally for longer FI to shorter FI block switches.

Correlations between FIs or EIs (experienced interval, meaning, real interval between rewards experienced by the animal) and the PRPs were achieved by performing linear regression analysis. In the case of the single session analysis, we performed linear regressions using either the PRPs and the FI or the PRPs and the EIs.

For the population analysis, data from 91 individual single sessions was filtered as described above and combined in sets of blocks with the same FI size, each belonging to a single session. Then we calculated the PRPs mean for each block belonging to each individual session. We also generated average frequency rate of response in each individual session as described above, but additionally, we also calculated the average frequency rate of response for each block size, in the population. This was done simply by taking the average frequency rate of response from all the single sessions and averaging the data belonging to the same block size.

We also performed linear regression analysis for the population data set in order to look for correlations only between PRPs and EIs. We only used this combination because the EIs were a more realistic measure of the time between rewards experienced by the animal than the FI, and because the difference between the two analysis performed in the single session case was not significant.

Learning curves were generated in a similar way as in the single session analysis, but using a larger data set and averaging the results from the multiples sessions, in the trial by trial calculations of the Z-scores. We also calculated the standard error of the PRPs mean by taking the standard deviation of the mean distribution of PRPs in each session and dividing it by the square root of the sample size.

All data analysis was performed using the MatLab2009b software (MathWorks, Inc).
Viral construction and production

The AAV construct was a kind gift from Karl Deisseroth. It consisted of a ChR2-YFP coding region that was originally subcloned from a lentiviral vector into loxP/lox2722-flanked (locus of X-over P1), inverted ORF (open reading frame) AAV plasmid (Fig7). The ChR2-YFP coding region was located downstream of the elongation factor-1α (EF1α) promoter and upstream of the woodchuck hepatitis post-transcriptional regulatory element (WPRE), to enhance expression. Plasmid was verified by sequencing. High-titer stock of AAV-EF1α-DIO (double-floxed inverted open reading frame)-ChR2-YFP-WPRE expressing virus (8x10^{12} pfu/mL in PBS - phosphate buffered saline) was produced at the Penn Vector Core (University of Pennsylvania).

Viral injection

For the AAV injections, one PV-Cre mouse and one D2-Cre mouse, both two months old, underwent the same surgery protocol. They were anesthetized with isoflurane in a chamber and then positioned in a stereotaxic apparatus, where they continued under anesthesia through a gas anesthesia mask connected to the head holder section of the stereotaxic equipment (923-B, Kopf). The mouse’s upper front teeth were placed in the head holder and zygoma ear cups (921, Kopf) were positioned and tightened over the animal’s temporal lobes. This was done to make sure that the animal’s head was not moving through the surgery and therefore the injection coordinates could be measured correctly.

The scalp was rubbed with alcohol and then swabbed with iodine/betadine to sterilize. Then, a central incision was made, the skull was mapped stereotaxically for bregma, and two craniotomies approximately 2mm in diameter were drilled bilaterally (0.5 mm anterior-posterior, ± 2.25 mm medial-lateral; all coordinates relative to bregma). The virus (0.5, 0.75 and 1 µL into distinct injection sites) was delivered with a glass micropipette connected to a Nanoject automatic injector (Drummond, Sci. Co.). A pulse generator was connected to the Nanoject and sent one TTL (transistor-transistor logic) pulse every 5 sec, with 1 ms width. Viral injection to the dorsal striatum was achieved by lowering down the micropipette 2.25mm from pial surface.
Then, the virus was injected at 9.2 nL/sec. Different volumes were injected in each animal. The PV-Cre animal was injected with 1 µl in each hemisphere. In the case of the D₂-Cre animal, 0.5 µL were injected in the right hemisphere and 0.75 µL were injected in the left hemisphere. Since a TTL pulse was sent every 5 sec, it took 4 min and 30 sec, 6 min 45 sec and 9 min to inject 0.5, 0.75 and 1 µL, respectively.

In order to prevent the virus from being carried out along with the micropipette, we waited at least 10 min before removing the micropipette. After injection, the craniotomies were covered with silicone sealant, the skin was repositioned with tissue adhesive VetBond (3M, Co.) and the animals were returned to their home cages after regaining movement. During all surgical procedures, animals were placed on a heating pad and their eyes protected with Frakidex (Bausch & Lomb, Inc.).

**Histology**

Two weeks post surgery, both animals that undergo viral injection were deeply anesthetized intraperitoneally with Ketamine/Xilazine (120 mg/mL Ketamine, 1.8 mg/mL Xilazine). Next, their hearts were exposed, a needle connected to a pump (101U/R, Watson-Marlow) was introduced in the left ventricle and the right atrium was snipped. The animals were then slowly perfused with about 40-50 mL of cold 0.9% (m/v) solution of NaCl until the fluid coming from the right atrium was clear. Then, the NaCl solution was changed to cold 4% (m/v) paraformaldehyde (PFA) and the animal was again slowly perfused with about 50 mL of PFA. The brains were removed and fixated in PFA overnight at 4°C, and then stored in PBS 1x at 4°C. Then, 50 µm thick coronal sections were prepared with a VT100 vibratome (Leica Microsystems).

Free floating sections from PV-Cre injected mice were placed 4 per well, washed 3x5 min in PBS 1X and then incubated overnight at room temperature with a mouse monoclonal anti-PV antibody (Sigma) diluted 1:10000 in PBS 1x with 0.4% Triton X-100 (v/v). After washing 5x5 min in PBS 1x, the sections were incubated for 2 h at room temperature with the Alexa594-conjugated goat IgG fluorescent secondary antibody (Invitrogen), diluted 1:1000 in PBS 1x with 0.4% Triton X-100 (v/v). The slices were then rinsed in PBS 1x, mounted on glass slides and coverslipped with mowiol mounting media.

Free floating sections from D₂-Cre injected mice were placed 4 per well, washed 3x5 min in PBS 1X and then stained for 30 min at room temperature with TO-PRO-3 (Invitrogen) in PBS 1x with 0.4% Triton X-100 (v/v). After washing 5x5 min in PBS 1x, the sections were mounted on glass slides and coverslipped with mowiol mounting media.
In order to analyze the overall expression of ChR2-YFP in full slices of both mouse lines, we used a SteREO Lumar V.12 to acquire images in a low magnification with a ApoLumar 1,2 lens (all from Zeiss). For YFP detection, we used band pass (BP) 500/25nm excitation filters and BP 535/30nm emission filters. Identification of PV+ interneurons expressing ChR2-YFP was achieved by using AxioImager Z1 microscope, equipped with an Axiocam MR ver.3.0 camera and the Axiovision 4.7 software (all from Zeiss). Images were acquired with a 40x objective (ECplan Neofluar 40x, 0.75 numerical aperture), with BP 500/25nm excitation filters and BP 535/30nm emission filters for YFP and we used BP 546/12nm excitation filters and 590nm LP (long pass) emission filters for Alexa-594.

In order to assess labeling specificity, we used a confocal microscope (LSM-510/META, Zeiss) under a 63x objective (oil plan-Apochromat 63x, 1.4 numerical aperture) to acquire images and the labeled neurons were analyzed morphologically. Excitation was performed using 488nm and 543nm lasers for YFP and Alexa-594 respectively, and the same main dichroic beam splitter configurations (HFT 488/543/633nm) for both cases. The two emission records were captured using the META detector in the ranges of 500–550nm for YFP and 565–615nm for Alexa-594.

Identification of D2MSN expressing ChR2-YFP was achieved by using the same confocal microscope and objectives as in the PV+ interneurons imaging. Also, excitation of D2MSN samples was performed in a similar way as in the PV+ interneurons imaging, with the difference that the samples were excited using 488nm and 633nm lasers for YFP and TO-PRO-3, respectively. Again, two emission records were captured using the META detector in the ranges of 500–550nm for YFP and 650–710 nm for TO-PRO-3.
Results

Behavior during the SFI task

We first trained mice in the SFI schedule and analyzed the raw data in single sessions. We noticed that mice developed a reliable pattern of behavior that consisted in pressing the lever once and “checking” rapidly if the reward had been delivered. This was true even for times when the mice were pressing the lever at a very high frequency, and faithfully resemble the pattern of behavior displayed by rats performing the same task (personal communication).

In Fig8 we have an example of the lever press distribution in a single session, without any data filtering. Every trial and block is in the same sequence as they were during the session and all the single session plots here shown use data from this session. The gray dots in the very beginning of some trials represent a lasting effect of a high rate response from the previous trial. Therefore, for every trial, we selected the first response after reward using a heuristic and not the actual first response in the data set (see Methods). The distribution of these first responses after reward suggests that the PRP is sensitive to the block size, since the first lever press after reward appears later in trials with a long FI when compared to trials with shorter FI.

![Fig8 - Trial by trial distribution of all lever presses as a function of time, during a SFI single session. Each gray dot represents one lever press. The first response selected by the heuristic for data analysis is highlighted in red.](image)
Because we wanted to have a better visualization of the PRPs changes during the course of a single session, we plotted in Fig9 the running mean of the PRPs from the 3 previous trials, together with the mice PRPs and the FI sizes in each trial. In this and in all subsequent figures, data has been filtered and only trials that were preceded by a reward were used. Here we can see more clearly how the increase in FI size is followed by an increase in the running mean of the PRPs.

![Fig9 - Running mean of the PRPs during a SFI single session.](image)

For the population data set (n=91 sessions), we plotted the mean PRP in each block (Fig10) and, again, we observed an increase in the mean PRPs for longer FI durations.

![Fig10 - Distribution of the average rate of response and mean PRP as a function of block size.](image)
The EI, this is, the real time between rewards experienced by the animal, was slightly different on a trial by trial basis. This happened because, after the FI elapsed, mice had 15 sec to press the lever and receive the reward. Therefore, the EI varied from FI sec to FI+15 sec. To check whether the EI could have an effect on the PRPs, we calculated the EI and plotted it together with the running mean of the PRPs and the median PRP within each block (Fig1). Here we can see that the EI is very close to the FI within each FI size, because mice show a relatively high rate of lever presses right before the end of the FI, as it can also be seen in Fig10.

We then performed linear regression analysis to analyze possible correlations between the FIs and the PRPs (Fig12A) \( \left( R^2 = 0.43959, P\text{-value} < 0.001 \right) \), as well as between the PRPs and the EIs of the previous trial (Fig12B) \( \left( R^2 = 0.45809, P\text{-value} < 0.001 \right) \) and found a significant linear correlation in both cases. During the SFI task, animals can directly evaluate the EIs but not the actual FIs. Therefore, and because there was not a significant difference between both correlations, we only searched for correlations between PRPs and EIs in the population data set. We also found a significant linear correlation between PRPs and EIs when we analyzed the

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**Fig11** – Distribution of the median PRP and EI during a single session. Solid black line represents the FI (fixed interval) duration in each block. Solid gray line represents the interval between rewards delivery, this is, the actual interval experienced by the animal (EI). The running mean of PRPs (post reinforcement pauses) is represented in dashed dark gray line and the solid red line represents the median PRPs in each block. Solid blue line represents constant amount of reward delivered thorough the session.
population data set (Fig13) ($R^2 = 0.7813$, P-value < 0.001), confirming that PRPs distribution is time sensitive.

We found that, along with the PRPs, the response rate profile also showed this time sensitivity. In Fig14A we can see that, although the maximum average response frequency rate for each block of FIs is similar, its development is progressively slower for longer FIs. This result was also seen in the population analysis (Fig14B), both in the average response frequency rates in several sessions and in the averaged response frequency rates of the whole population. Thus, manipulating the FI during a SFI operant lever pressing task can be used to systematically vary the report of time until reward in mice. To check if mice could rapidly adapt their timing behavior to FI changes, we calculated the Z-Score for the early trials in each block of both single session and population data, as well as its standard error of the mean (SEM). In Fig15 we can see that mice change their PRPs to match the new FI size, and they do it considerably fast (~6 trials), even though this result is not seen in the individual data analysis (Fig16).
**Fig13** – Correlation between the latency to respond and the experienced intervals, in the population data sets. Correlation between the PRPs and the El between rewards in the population data set. Single trial PRPs are represented in red. Black dots represent the mean PRP per block of trials with the same FI within each session. Blue bars represent the standard deviation of the PRPs.

**Fig14** – Distribution of the average response rate in each block of trials with the fixed interval (FI). **A** – Each curve represents the single session average frequency rate of response in a given block size. Block sizes are color coded so that the greener, the longer the FI size. **B** – Each thin curve represents a single session average frequency rate of response in a given block size and many sessions are plotted in this figure that is representative of the population. Color code is identical to panel **A**. The four thicker curves represent the population average rate of response in each block of trials with the same FI. Again, color code is identical to panel **A**.
We then asked if mice could learn faster how to go from a long FI to a short FI than the other way around, so we analyzed separately block switches that were preceded by a smaller

Fig15 – Distribution of the population average learning curves. A – Distribution of the Z-scores extracted from the post reinforcement pauses (PRPs) in the first 15 trials after a switch from a long FI to a short FI in all the sessions. B – Distribution of the Z-scores extracted from the PRPs in the first 15 trials after a switch from a short FI to a long FI in all the sessions. C – Combined distribution of Z-scores extracted from the PRPs in the first 15 trials after a switch in FI duration.
The thin solid blue line represents the standard error of the mean (SEM) in all panels.

Fig16 – Distribution of the learning curves for block transitions in a single session. A – Distribution of Z-scores extracted from the post reinforcement pauses (PRPs) in the first 15 trials after a switch from a long FI to a short FI. B – Distribution of Z-scores extracted from the PRPs in the first 15 trials after a switch from a short FI to a long FI. C – Combined distribution of Z-scores extracted from the PRPs in the first 15 trials after a switch in FI duration.

We then asked if mice could learn faster how to go from a long FI to a short FI than the other way around, so we analyzed separately block switches that were preceded by a smaller
block (Fig15A) and block switches that were preceded by longer blocks (Fig15B). As we can see in these figures, mice performed very well in both scenarios.

**Preliminary results on ChR2 expression**

We looked at the expression of ChR2-YFP in the dorsal striatum of two mouse lines: D2-Cre and PV-Cre. The inverted ChR2-YFP coding region of the AAV was flanked by two loxP sites (one regular and one mutated, lox2722) on each side of the coding region, 4 in total (Fig17). Because the loxP sites upstream the coding region were on the opposite orientation of the ones downstream, double recombination between these sites leads to an inversion of the cassette. Selectivity in the expression of ChR2-YFP is achieved because the recombination only occurs in the presence of Cre.

![Diagram of recombination sequence](image)

**Fig17 – Scheme of the recombination sequence for inversion of ChR2-YFP coding region.** loxP sites don’t recombine with lox2772 sites, only with its equivalent pair. Since these are oppositely orientated, the inversion process occurs in two steps. First, the inversion occurs, independently of which pair of lox sites suffers recombination (A). This first step is followed by the excision of one lox site from each pair (B).

In Fig18A and Fig19A, fluorescence images of a D2-Cre mouse brain slice expressing YFP are shown in comparison to a schematic of a coronal section of the mouse brain, at the respective approximate anterior-posterior coordinates. These illustrate the areas of ChR2-YFP expression. The injection of the virus shown in Fig18B targeted the dorsolateral striatum. In figure Fig19B we see that, there is ChR2-YFP expression in GPe. Since the expression of D2-Cre in the transgenic mouse line we used is limited to a subset of adult D2 expressing neurons (primarily striatopallidal neurons), and no other area has shown ChR2-YFP expression, we can
Fig 18 - Selective viral-mediated ChR2 expression in striatal D2MSNs. A - A fluorescent image from a coronal section of a D2-Cre mouse brain (left) compared to a coronal mouse brain schematic (right). Anterior-posterior coordinates are ~+0.5mm. CPu – Caudate-putamen (striatum). B - Coronal section showing striatal D2MSNs expressing ChR2–YFP following injection of Cre-dependent AAV into D2-Cre BAC transgenic mice. Scale bar – 2mm. C – Confocal image of ChR2–YFP-expressing neurons in the striatum. Arrows indicate points of accumulation of ChR2-YFP. D – Confocal image of To-Pro-3 staining in the same region as in panel C. E – Panels C and D merged. Example of To-Pro-3 stained nucleus and ChR2–YFP expression in the same neuron (arrow). The white box indicates the region shown in panel F. F – Example of a spiny dendrite from a D2MSN expressing ChR2-YFP. Scale bars in C, D, E represent 15µm, in panel F it represents 5µm.

Fig 19 - Selective viral-mediated ChR2 expression in striatal D2MSNs projections to the GPe. A - A fluorescent image from a coronal section of a D2-Cre mouse brain (left) compared to a coronal mouse brain schematic (right). Anterior-posterior coordinates are ~0.46mm. CPu – Caudate-putamen (striatum); GPe – globus pallidus external. B - Coronal section showing striatal D2MSNs projections expressing ChR2–YFP in the GPe, following injection of Cre-dependent AAV into D2-Cre BAC transgenic mice (same animal as in Fig 15). Scale bar – 2mm. C – Confocal image of a region with processes expressing ChR2–YFP in the GPe. D – Confocal image of To-Pro-3 staining in the same region as in panel C. E – Panels C and D merged. Example of To-Pro-3 stained nuclei and ChR2–YFP expression in processes that do not belong to the same neuron. Scale bars in C, D and E represent 15µm.
deduce that the processes expressing ChR2-YFP that we see in the GPe are in fact projections from the striatal D₂ MSNs. Using confocal microscopy, we can identify neurons expressing ChR2-YFP (Fig18C). We also stained these slices with a deoxyribonucleic acid (DNA) binding fluorescent marker, TO-PRO-3 (Fig18D, Fig19D) in order to visualize cell nuclei. Combining both panels C and D in Fig18E, we see that there is co-localization of cells expressing ChR2-YFP at their membrane and nuclei stained with TO-PRO-3, that take over most of the cell body, a feature present in MSNs. Also, the labeled cells have spiny dendrites (Fig18F), another characteristic feature of MSNs. In (Fig19C), we see the MSNs projections to the GPe area, where no cell bodies express ChR2-YFP. Also, Fig19D shows stained nuclei that don’t have the same morphology as the ones in Fig18D. Analyzing both panels together (Fig19E), we find that the processes expressing ChR2-YFP don’t seem to belong to the stained nuclei, suggesting that there are no cells expressing ChR2-YFP in the GPe. We also observed accumulations of ChR2-YFP that are present across all slices that show ChR2-YFP expression, forming small spherical inclusions (Fig18C and Fig19C).

Additionally, we targeted PV+ interneurons by injecting the AAV in a PV-Cre mouse line. In Fig20A we have a fluorescence image of a brain slice from the PV-Cre line compared to a schematic of a mouse brain, illustrating the areas of ChR2-YFP expression. In Fig20B we have the complete fluorescent image: we can scarcely see any fluorescence in
the striatum and cortical neurons are labeled (arrows). When we look in a higher magnification to the striatal area, we can identify neurons expressing ChR2-YFP (Fig 20C). The labeled neurons possess the anatomical features of a PV+ interneuron. We immunostained the slices for PV and analyzed areas where there was expression of ChR2-YFP and in parallel the immunostaining result for the same area (Fig 20D). Merging the two panels, we see colocalization of ChR2-YFP cells and PV+ cells (Fig 20E).

Our results from the group of mice trained in the SFI task show that these animals display time sensitive behavior. In parallel, we achieved ChR2-YFP expression in two types of striatal cells (D2 MSNs and PV interneurons). We confirmed this neuron type selectivity by analyzing the cells projection sites and by using histochemical techniques.
Discussion

We trained mice in the SFI, a task developed in our lab that combines the required features for a behavioral paradigm to the electrophysiological research of timing. Although at first this task appears to be somewhat difficult, mice display a very reliable time sensitive behavior, since its PRPs significantly co-varied with the FI inside each block. This stable and reliable response pattern was accompanied by fast learning curves for short to long and long to short interval changes in the population, but not on a single session analysis. Also, animals do not appear to possess a directional facilitation for learning a new FI when trained in the SFI task.

We successfully expressed ChR2 in two transgenic mouse lines, D2-Cre and PV-Cre. Tagged neuronal population in each line display characteristic morphological features of D2MSNs and PV+ interneurons, respectively. Additionally, we immunostained slices with an antibody against PV in the PV-Cre line, and used the DNA binding marker TO-PRO-3 to identify nuclei within D2-Cre mice infected with the DIO AAV. PV immunostaining confirmed the identity of the tagged neurons, and TO-PRO-3 staining suggests that infected cells are MSNs, and based on their projection pattern (to the GPe area), they are most likely D2MSNs. ChR2-YFP expression was followed by the occurrence of inclusions of the fusion protein, which were more noticeable in the D2-Cre mice.

Regarding the behavioral results, in a single session analysis, mice did not achieve learning curves nearly as good as in the population set. The SFI task is more dynamic, in terms of the range of time intervals that the animal has to time during one single session, than other classical timing tasks. The SFI stands on a simple paradigm where every trial can provide valuable information, unlike other tasks where only a fraction of the total number of trials is used for statistical analysis. The use of several FIs in one single session with big time ranges (from 12s to 60s) allows the search for correlations between the animal’s behavior in specific time intervals and their electrophysiological activity during those intervals. Among rodents, rats have been the main animal model used for timing research. To train transgenic mice in such a task may lead to deeper understanding of the response profile of each cell type during timing behavior.

The SFI task was developed and implemented for the first time in our lab, using rats as a model (personal communication). The results here shown suggest a very similar pattern of behaviors between both species of rodents, making it more plausible to try and create
hypothesis for our future studies with mice, taking the electrophysiological data from the rats as a basis.

Regarding the preliminary results on the expression of ChR2-YFP, an intriguing observation was that expression of ChR2-YFP in the cell membrane was accompanied by the presence of inclusions of ChR2-YFP. These could be a result of high viral expression, maybe caused because of a too high viral titer, leading to accumulations of fusion protein within cells. If so, the physiological properties of the cell could be altered or it could even be causing cell death. We stained slices from infected D2-Cre mice with the DNA binding marker TO-PRO-3 to identify nuclei and try to detect some abnormality but stained nuclei looked normal. To exclude the hypothesis that these inclusions may be causing cell death, an apoptotic marker should be used in these slices in the future.

Although previous work suggests that ChR2–YFP expression alone does not alter the electrophysiological properties of MSNs, these used either different titers of this DIO viral construct or different viral constructs\cite{72,95,98,99}. Therefore, further testings in vitro are required in order to understand the impact of ChR2-YFP alone, as well as the impact of the inclusions in the electrophysiological properties of the cell types in question. One way to try and overcome the high levels of expression would be to reduce the viral titter. Interestingly, a recent study shows images of ChR2-YFP MSNs and inclusions seem to be present there as well (though they are not mentioned). This same study claims that electrophysiological properties of MSNs were not altered by ChR2-YFP expression alone\cite{72}.

In the PV-Cre line, in a low magnification, we could not detect the same levels of fluorescence as in the D2-Cre animals. This is probably due to the fact that this type of neurons represent only about 0.7% of all the striatal neurons\cite{100}. Another interesting observation was that, in the case of PV-Cre animals, because this transgenic line also expresses Cre in cortical interneurons, a population of cortical cells also expresses ChR2-YFP. Although the viral microinjection was performed in the striatum, the labeled cells in the cortex seem to reveal the “path” of the micropipette meaning that, most likely, viral particles traveled with the micropipette when the last was being removed from the animal's brain. Although we waited for more than 10 min post injection before removing the micropipette slowly, this step probably needs to be performed even more slowly. Moreover, another approach to try and avoid the problems related to the virus would be to perform multiple injections of about 100nL each. With these problems overcome, we believe that during image acquisition, it would be easier to acquire confocal images that more distinctly show dendrites from MSNs.
Future experiments, including slice physiology studies, will test what has been shown in other studies, that ChR2 expression allows for reliable, millisecond timescale control of neuronal spiking, as well as control of excitatory and inhibitory synaptic transmission\textsuperscript{72,95,97,98,99}. Further work is required to fully understand the extent of ChR2-YFP expression in cells other than D\textsubscript{2}MSNs and PV+ interneurons (false positives) and the existence of false negatives, although this would be potentially less problematic. Specifically, stereological cell counting should be performed to evaluate at which extent PV+ interneurons are co-localized with ChR2-YFP expression, and staining against several types of interneurons should be done in D\textsubscript{2}-Cre slices, because some interneuron classes also express D\textsubscript{2} receptors. Also, Cre mediated recombination could generate false positives but this should be of less great extent, since we used a double recombination scheme that is more reliable that using virus with a classic lox-stop-lox cassette.

Additional experiments need to be performed in order to express ChR2-YFP in other cell types, specifically D\textsubscript{1}MSNs and cholinergic interneurons. Although there are at least two more subtypes of interneurons, they either are much poorly characterized (calretinin-positive interneurons), or they present themselves very difficult to target (somatostatin-, neuropeptide Y-, and nitric oxide synthase-positive interneurons). As for the D\textsubscript{1}MSNs and cholinergic interneurons, there are already transgenic mouse lines expressing Cre in each neuronal type, allowing for the same approach to be applied.

We now have been able to train mice in a very reliable and highly dynamic timing paradigm. In parallel, we have a way to target two specific neuronal types within striatum. Now, animals can be trained in the SFI and then submitted to viral microinjection as well as cannula and electrode implantation. We will then record from specific neuronal types in each transgenic line and compare their behavioral responses with the electrophysiological activity of the neurons being recorded. This approach promises to add crucial information about how different cell types process timed information. Ultimately, this information will be essential for building accurate circuit models that faithfully replicate the computations normally carried out by brain circuits in behaving animals. In addition to a contribution in the development of new circuit models, the information concerning which signals are carried by which cell types during timing behavior can also be helpful to characterize brain circuits implicated in motor and cognitive dysfunctions in diseases such as PD and HD.
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