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Dopaminergic Modulation in Medial Prefrontal Cortex and its Dysfunction in Psychiatric Disease

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Dopaminergic Modulation in Medial Prefrontal Cortex and its Dysfunction in Psychiatric Disease

by

Sarah Elizabeth Robinson Schwartz

DISSERTATION

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DOCTOR OF PHILOSOPHY

in

Neuroscience

in the

GRADUATE DIVISION

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Author Contributions

The text of Chapter II of this dissertation is a reprint of the material as it originally appeared in The Journal of Neuroscience (Robinson et al., 2017). Sarah Robinson performed all experiments and analysis. Vikaas Sohal directed and supervised the research. Sarah Robinson and Vikaas Sohal co-wrote the paper.

The text of Chapter IV of this dissertation contains excerpts from two manuscripts in preparation. Sarah E. Robinson Schwartz performed all electrophysiology and behavior experiments as well as viral injections for the histology experiments. Bin Chen collaborated on generating the Tbr1 conditional mutant mouse. Siavash Fazel Darbandi generated the Tbr1 layer 5 and layer 6 mutant mice, performed ChIP-Seq, FACS RNA-Seq and in situ hybridization transcriptome analyses, as well as synaptic staining and all histology experiments. Jeremy Willsey, Jeff Mandell, and Qihao Qi performed bioinformatics analysis of the Tbr1 RNA-Seq data. Alex Nord and Rinaldo Catta-Preta conducted bioinformatics analysis of Tbr1 ChIP-Seq data. Vikaas Sohal, John Rubenstein, and Matthew State directed and supervised the research. S.E.R.S, S.F.D., V.S.S., J.L.R., and M.W.S. contributed to the text.
Abstract

The prefrontal cortex (PFC) mediates many executive functions including working memory, behavioral flexibility, decision making, and social cognition. The normal functioning of the PFC depends strongly on regulation by neuromodulators. Dysfunction of the PFC leads to deficits in these abilities and is a major factor in many neuropsychiatric disorders including schizophrenia and autism. Layer 5 (L5) of the PFC, a major site of dopaminergic modulation, is thought to play an important role in regulating these higher order processes. Indeed, many neurological disorders such as schizophrenia have been associated with imbalances in dopamine neurotransmission.

Prefrontal dopamine D2 receptors (D2Rs), the primary target for first generation antipsychotic drugs, specifically play a major role in tasks that are disrupted in schizophrenia. One major hypothesis is that excessive D2R activation within the PFC contributes to many of the cognitive impairments associated with schizophrenia. This dissertation investigates the mechanisms of D2R-mediated neuromodulation within the PFC and abnormalities within PFC circuitry and neuromodulation in mouse models of schizophrenia and autism.

We have previously described a phenomenon whereby D2R activation elicits afterdepolarization (ADPs) in subcortically-projecting (SC) pyramidal neurons within L5 of the PFC. Results presented in Chapter II of this dissertation show that this unusual physiological phenomenon, in which D2Rs enhance cellular excitability dependent on synaptic input, is mediated at the cellular level through the recruitment of signaling pathways associated with Gs, rather than Gi-associated mechanisms that have classically been ascribed to D2Rs.
In Chapter III, I discuss differences in D2R neuromodulation in mice with a dominant-negative mutation in disrupted in schizophrenia 1 (DISC1). DISC1 dysfunction has been associated with schizophrenia and causes deficits in working memory. Current studies, it has been shown that there is an upregulation of D2R expression and/or activity in DISC1 mouse models. Interestingly, DISC1 is also implicated in the regulation of cAMP. Here, I will show that these mice lack the quinpirole-induced $G_i$-independent ADP.

The medial prefrontal cortex (mPFC), particularly deep layer projection neurons, has also been implicated as a potential locus for Autism Spectrum Disorder (ASD) pathology. Previous work from our lab has shown that social exploration preferentially recruits mPFC D2R positive pyramidal neurons and that this recruitment is attenuated in three etiologically distinct mouse models of autism. In Chapter IV of this dissertation, we investigate the affects of $T$-brain-1 ($Tbr1$) selectively within layer 5 and layer 6 cortical neurons. We found that $Tbr1$ function is required to maintain many aspects of layer 6 identity, $Tbr1^{layer6}$ mutant neurons transform towards the identity of L5 neurons, including their transcriptome, dendritic pattern, and physiological properties. $Tbr1^{layer5}$ mutant neurons become a more homogeneous population transforming towards the identity of D2R expressing pyramidal neurons. Both $Tbr1^{layer6}$ and $Tbr1^{layer5}$ mutants have reduced excitatory and inhibitory synaptic density as well as reduced spontaneous EPSCs and IPSCs. We also present data suggesting that loss of $Tbr1$ function in layer 6 leads to increased anxiety and aggressive behavior whereas loss of $Tbr1$ function in layer 5 leads to decreased social behaviors, phenotypes consistent with different aspects of observed ASD patient behavioral characteristics.
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CHAPTER I

General Introduction
Normal and pathological roles of the PFC

The ability to appropriately make decisions which guide our subsequent goal-directed behaviors requires integration of previously learned action-outcome relationships and accurate environmental assessment. This ability is thought to be primarily mediated by the prefrontal cortex (PFC), which also plays a critical role in high-order executive tasks such as attention, behavioral flexibility, working memory, and social cognition (Druzin et al., 2000; Dalley et al., 2004; Floresco et al., 2006). Neuromodulation via dopamine plays a critical role in mediating these prefrontal functions. Dopamine dysfunction is strongly implicated in neuropsychiatric diseases (Roberts and Bruton, 1990; Kalivas and Volkow, 2005; Durstewitz and Seamans, 2008). This dissertation focuses on understanding the mechanisms of D2R-mediated neuromodulation within the PFC and abnormalities within the PFC circuitry in mouse models of schizophrenia and autism. By understanding these mechanisms, we hope to advance our understanding of how these mechanisms contribute to prefrontal dysfunction in psychiatric disease.

The effects of dopamine on the PFC

In order for the PFC to provide flexible top-down control of complex cognitive and behavioral processes it is heavily connected, sending and receiving inputs to and from numerous brain regions including sensory cortices, contralateral PFC, and subcortical regions (Wernicke, 1906; Carr and Sesack, 2000; Hoover and Vertes, 2007). In addition to these long-range projections, pyramidal neurons within the PFC also form highly recurrent connections with other PFC pyramidal neurons (Morishima and Kawaguchi, 2006). Importantly, the PFC receives strong mesocortical dopaminergic inputs from the ventral tegmental area (VTA) (Björklund and Dunnett, 2007). Traditionally, activity of the dopamine neurons within the VTA is believed to encode reward-prediction error (RPE), driving
learning by indicating a discrepancy between expectation and outcome (Schultz, 2007). However, the VTA contains anatomically and functionally heterogeneous dopamine neuron subpopulations with distinct axonal projections and input sources (Lammel et al., 2014). Recently, dopamine neuron activity has been associated with a variety of additional brain functions including aversion, salience, uncertainty, and novelty (Schultz, 2007; Bromberg-Martin et al., 2010; Ungless et al., 2010). For example, dopaminergic cells projecting to the nucleus accumbens respond to rewarding stimuli whereas dopaminergic cells projecting to the medial prefrontal cortex (mPFC) respond to aversive stimuli (Lammel et al., 2012). However, optogenetic stimulation of VTA terminals within the VTA rather than being positively or negatively reinforcing, instead shifts an animal between flexible or preservative behavior modalities (Ellwood et al., 2017). Neuromodulation by prefrontal dopamine is a topic of extensive research due to its critical role in higher cognitive processes and neuropsychiatric pathology (Seamans and Yang, 2004).

**Different effects of dopamine by D1R and D2R receptors in the PFC**

Dopamine modulates neuronal functioning by acting at five distinct G protein-coupled receptors (GPCRs). These dopamine receptors are canonically subdivided into two families: “D1-like” (D1 and D5) which tend to couple to the stimulatory G proteins $G_s$ and $G_q$, and “D2-like” (D2, D3, and D4) which tend to couple to the inhibitory proteins $G_{i/o}$ (Bonci and Hopf, 2005). Interestingly, the dopamine D1 receptor (D1R) family and the dopamine D2 receptors (D2R) family are thought to have opposing effects on PFC-dependent cognitive function (Sawaguchi and Goldman-Rakic, 1994; Seamans et al., 2001; Wang and Goldman-rakic, 2004; Tseng and O'Donnell, 2005). The “Dual-State Theory” of prefrontal function proposes that D1Rs within the PFC stabilize network activity, decreasing the
influence of weaker inputs thereby maintaining patterns of activity supporting previously learned behavioral strategies (Durstewitz and Seamans, 2008). Contrastingly, D2Rs are thought to destabilize the prefrontal network, shifting it toward promoting a more flexible behavioral state. Successful behavioral adaptation in the face of uncertainty therefore depends on a balance between these “exploit” and “explore” modalities (Daw et al., 2006).

D1Rs and D2Rs within the basal ganglia are expressed on non-overlapping populations of medium spiny neurons (MSNs) with distinct projection targets. “Direct pathway” MSNs in the striatum express D1Rs and project to the internal segment globus pallidus/sustantia nigra pars reticulate. Dopamine acting on these MSNs via D1Rs, acts through canonical GPCR signaling pathways, increasing intercellular levels of cAMP and cellular excitability (Surmeier et al., 2007). Activity in this pathway has been shown to facilitate movement whereas activity in the “indirect pathway” is thought to inhibit movement (Lalonde and Botez-Marquard, 2010). Indirect pathways MSNs project to the external segment of the globus pallidus and express D2Rs coupled to G_i/o, inhibiting cAMP production in response to dopamine. While this model is likely an oversimplification, it does provide a useful framework for testing the role of specific circuits and cell populations in basal ganglia function (DeLong and Wichmann, 2009; Kravitz et al., 2012). A similar detailed framework for dopaminergic modulation within the PFC is largely lacking.

The mechanisms by which D1Rs modulate excitability in the PFC have been extensively studied whereas the mechanisms through which D2Rs exert their influence remain mostly unclear. Neuromodulation by prefrontal dopamine D2 receptors (D2Rs) is of particular interest because of their implications in many neuropsychiatric disease pathologies. The infusion of D2R agonists and antagonists into the PFC modulates working memory and set-shifting in rodents (Druzin et al., 2000; Floresco et al., 2006; St Onge, Abhari and Floresco, 2011a). Selective photostimulation of mPFC pyramidal neurons
expressing D2Rs disrupts normal social behavior in mice whereas inhibition of these neurons improves social interaction in mouse models of autism (Brumback et al., 2017). Prefrontal D2Rs have also been shown to be necessary for neural activity associated with memory-guided saccades in non-human primates (Wang and Goldman-Rakic, 2004). In humans, genetic variation in D2Rs modulates prefrontal activity and working memory (Zhang et al., 2007). Interestingly, all known antipsychotics block prefrontal D2Rs supporting a major hypothesis that aberrant D2R activation contributes to prefrontal dysfunction in schizophrenia (Winterer and Weinberger, 2004; Durstewitz and Seamans, 2008). Thus, D2Rs play a major role in both normal and pathological prefrontal function. It is therefore crucial to reveal the underlying mechanisms of dopaminergic modulation in the prefrontal cortex.

Recent studies have classified separate populations of PFC pyramidal neurons with distinct morphology, physiological properties and dopamine receptor expression (Gee et al., 2012). D2Rs are selectively expressed in a subpopulation of layer 5 pyramidal subcortical-projection (SC) neurons, which have thick apical tufts and prominent h-current. A recent study by Clarkson et al. confirmed that D2R expressing pyramidal neurons within the mPFC exhibit prominent voltage sag during hyperpolarizing current steps and have a voltage rebound that depolarizes past resting membrane potential after the current step (Clarkson et al., 2017). This electrophysiological phenotype alone was enough to distinguish D2R expressing neurons from those lacking D2Rs with approximately 75% accuracy.

Within this D2R expressing SC-projecting population we found pharmacological activation of D2Rs by the agonist results in a pronounced afterdepolarization (ADP) (Gee et al., 2012). This ADP was not observed in quiescent slices and only occurred when NMDA receptors were activated, via either optogenetic stimulation of excitatory synapses or bath application of low dose NMDA. D2Rs are classically assumed to couple to inhibitory G
proteins, reducing neuronal excitability by activating $G_{\alpha i/o}$ and inhibiting adenylate cyclase (Bonci and Hopf, 2005). The $G_{\beta\gamma}$ subunits also directly interact with several different types of ion channels and can facilitate release of $Ca^{2+}$ from internal calcium stores. Prolonged activation of D2Rs leads to binding of $\beta$-arrestin which results in desensitization and internalization of the receptors. Recently it has been discovered that the D2R-$\beta$-arrestin complex can also initiate signal cascades which are temporally and spatially distinct from the D2R-G-protein signaling pathway (Beaulieu et al., 2009; Beaulieu and Gainetdinov, 2011). In Chapter II, we explore two different aspects of the intercellular signaling pathways through which activation of D2Rs elicits ADPs in prefrontal SC-projecting neurons using a combination of electrophysiological, transgenic, pharmacological, and chemogenetic approaches. We find that the prolonged ability of synaptic stimulation to facilitate D2R-dependent ADPs depends on voltage-dependent (likely $Ca^{2+}$) currents. Furthermore, we show that the D2R-dependent ADP persists following disruptions to $G_i$ or $\beta$-arrestin signaling. The ADP can also be mimicked by chemogenetic activation of $G_s$ signaling suggesting that this phenomenon reflects novel intracellular signaling downstream of D2Rs via stimulatory G proteins.

The role of dopamine in disease

Many neurological disorders have been associated with imbalances in dopamine neurotransmission. Prefrontal D2Rs have also been postulated to contribute to schizophrenia (Winterer and Weinberger, 2004; Durstewitz and Seamans, 2008), Tourette syndrome (Simonic et al., 1998; Minzer et al., 2004; Yoon et al., 2007; Steeves et al., 2010), and bipolar disorder (Minton et al., 2009). Thus, D2Rs play a major role in both normal and pathological prefrontal function. One major hypothesis regarding the role of D2Rs in
psychiatric disease postulates that D2Rs increase the variability of PFC, under normal conditions facilitating adaptation to a changing environment (Durstewitz et al., 2010;). However, excessive or imbalanced D2R activation could produce pathological variability that contributes to “prefrontal noise” and cognitive dysfunction in schizophrenia (Winterer and Weinberger, 2004; Durstewitz and Seamans, 2008).

Schizophrenia is a devastating psychiatric disorder affecting approximately 1% of the population (Cloutier et al., 2016). The majority of patients suffer lifelong symptoms as the age of onset is in late adolescence to young adulthood. While symptoms vary from patient to patient, typical symptoms can be grouped into three main categories: ‘positive’ symptoms include hallucinations and delusions, ‘negative’ symptoms include emotional blunting, reduced motivation, and a lack of social interactions, and cognitive impairments including deficits in executive function, attention, and working memory (Meltzer, 1997; Evans et al., 2004). D2Rs are the main target of antipsychotic medications and the clinical potency of antipsychotic drugs is correlated with D2R binding affinity (Seeman et al., 1976). However, current antipsychotics are ineffective at controlling symptoms in many patients and cause serious side effect. Thus understanding the specific mechanisms of D2R signaling and its dysfunction in schizophrenia is critical for developing novel more effective therapies.

Disrupted in schizophrenia 1 (DISC1) dysfunction is associated with an increased risk for schizophrenia and other psychiatric disorders (Pletnikov et al., 2008). The DISC1 gene was originally discovered in a Scottish family with high rates of schizophrenia (Jacobs et al., 1970). DISC1 is a scaffold protein which interacts with several other proteins in the dopamine system, including serine/threonine protein kinase Akt and glycogen synthase kinase-3 (GSK-3). It plays an important role in regulating the cAMP pathway and mGluR activation (El-Hassar et al., 2014). In DISC1 mouse models there is an upregulation of D2R expression in the mPFC and striatum (Niwa et al., 2013). DISC1 mutant mice exhibit
spontaneous hyperactivity in the open field, alterations in social interaction, and deficiency in spatial memory which a behavioral deficits similar to some features of schizophrenia.

In Chapter III of this dissertation, we investigate abnormalities in D2R-mediated modulation of PFC pyramidal cell excitability in transgenic mice with inducible expression of mutant human DISC1 (hDISC1) via the Tet-off system under regulation of the CAMKII promoter (Pletnikov et al., 2008). I will present results showing that SC-projecting pyramidal neurons within the mPFC lack the quinpirole-induced afterdepolarization.

Throughout my dissertation we focus on layer 5 pyramidal neurons because these neurons contain most prefrontal D2Rs (Lidow et al., 1998; Santana, Mengod and Artigas, 2009; Clarkson et al., 2017). Specific mechanisms for D2R-modulation of L5 pyramidal neurons in PFC remains elusive with several contradictory studies describing ways that D2Rs enhance (Wang and Goldman-rakic, 2004) and suppress (Gulledge and Jaffe, 1998; Tseng and O'Donnell, 2004) excitability in these neurons. Understanding how mechanisms of dopaminergic modulation are perturbed in various psychiatric disorders is critical for the development of safer and more efficient drug therapies.

Autism spectrum disorder (ASD) is a neurodevelopmental disorder clinically characterized by social communication and interaction impairments which causes lifelong disability in millions of people (Baio, 2012). Individuals with ASD have impaired social behaviors often accompanied by abnormalities in language development. Many genes have been implicated in ASDs but specific cellular abnormalities that link genes with behavior remain elusive. Several of the genes strongly linked to autism are convergently co-expressed within deep layer (L5 & L6) projection neurons in the PFC (Willsey et al., 2013). In humans, functional imaging studies reveal abnormally decreased activation of the mPFC during social tasks (Pierce et al., 2004). The mPFC has been specifically implicated in deficits associated with ASD: lesions cause impairments in emotional learning and loss of
social skills (Bachevalier et al., 1986; Morgan et al., 1993). Optogenetic manipulation of the excitatory/inhibitory balance within rodent mPFC disrupts normal social exploration (Yizhar et al., 2011). Another recent study from our lab found that social exploration preferentially recruits mPFC D2R+ neurons and is disrupted in valproic acid (VPA) in utero exposure, a mouse model of autism (Brumback et al., 2017). Additionally, optogenetic stimulation of D2Rs+ mPFC neurons disrupts normal social behavior in mice whereas inhibition of these neurons improves social interaction in three etiologically distinct mouse models of autism (Brumback et al., 2017).

T-brain-1 (Tbr1), a T-box transcription factor strongly expressed in deep cortical layers, has been identified as a high-confidence ASD gene. Together with the John Rubensteiin Lab, we focused on Tbr1 function in deep neocortical neurons using Cre-lox conditional mutagenesis. Siavash Fazel Darbandi created both Tbr1layer6 and Tbr1layer5 mutant mice. Results presented in Chapter IV of this dissertation show that late gestation/neonatal Tbr1 function is required to maintain many aspects layer 6 (L6) identity. Tbr1layer6 mutant neurons transformed towards identity of L5 neurons, including their transcriptome, their dendritic pattern and electrophysiological properties (h-current). Similarly, Tbr1layer5 mutant neurons have enhanced h-current typical of D2R expressing SC-projection pyramidal neurons. The Tbr1layer6 and Tbr1layer5 mutant neurons also have reduced excitatory and inhibitory synaptic density. We found that this reduction in excitatory and inhibitory synapse density functionally translates into reduced spontaneous EPSCs and IPSCs in mouse brain slices. We have also discovered that loss of Tbr1 function in layer 6 leads to increased anxiety and aggressive behavior, a phenotype that is also observed in ASD patients. Contrastingly we show that loss of Tbr1 function in layer 5 leads to a pronounced decrease in social behaviors, reminiscent of the main phenotype associated with Autism Spectrum Disorders.
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CHAPTER II

Dopamine D2 Receptors Modulate Pyramidal Neurons in Mouse Medial Prefrontal Cortex through a Stimulatory G-Protein Pathway

ABSTRACT

Dopaminergic modulation of prefrontal cortex is believed to play key roles in many cognitive functions and to be disrupted in pathological conditions such as schizophrenia. We have previously described a phenomenon whereby dopamine D2 receptor (D2R) activation elicits afterdepolarizations (ADPs) in subcortically-projecting (SC) pyramidal neurons within L5 of the prefrontal cortex. These D2R-induced ADPs only occur following synaptic input which activates NMDA receptors (NMDARs) even when the delay between the synaptic input and afterdepolarizations is relatively long, e.g., several hundred milliseconds. Here we use a combination of electrophysiological, optogenetic, pharmacological, transgenic, and chemogenetic approaches to elucidate cellular mechanisms underlying this phenomenon in male and female mice. We find that knocking out D2Rs eliminates the ADP in a cell autonomous fashion, confirming that this ADP depends on D2Rs. Hyperpolarizing current injection, but not AMPA receptor blockade, prevents synaptic stimulation from facilitating D2R-induced ADPs, suggesting that this phenomenon depends on the recruitment of voltage dependent currents, e.g., NMDAR-mediated Ca\(^{2+}\) influx, by synaptic input. Finally, the D2R-induced ADP is blocked by inhibitors of cAMP / PKA signaling, insensitive to pertussis toxin or β-arrestin knockout, and mimicked by G\(_s\)-DREADD stimulation, suggesting that D2R activation elicits the ADP by stimulating cAMP / PKA signaling. These results show that this unusual physiological phenomenon, in which D2Rs enhance cellular excitability in a manner that depends on synaptic input, is mediated at the cellular level through the recruitment of signaling pathways associated with G\(_s\), rather than the G\(_i/o\)-associated mechanisms that have classically been ascribed to D2Rs.
SIGNIFICANCE STATEMENT

Dopamine D2 receptors in the prefrontal cortex (PFC) are believed to play important roles in behaviors including working memory and cognitive flexibility. Variation in D2Rs has also been implicated in schizophrenia, Tourette syndrome, and bipolar disorder. Recently, we described a new mechanism through which D2R activation can enhance the excitability of pyramidal neurons in the PFC. Here we explore the underlying cellular mechanisms. Surprisingly, although D2Rs are classically assumed to signal through $G_{i/o}$-coupled $G$-proteins and/or scaffolding proteins such as $\beta$-arrestin, we find that the effects of D2Rs on prefrontal pyramidal neurons are actually mediated by pathways associated with $G_s$-mediated signaling. Furthermore, we show how, via this D2R-dependent phenomenon, synaptic input can enhance the excitability of prefrontal neurons over timescales on the order of seconds. These results elucidate cellular mechanisms underlying a novel signaling pathway downstream of D2Rs that may contribute to prefrontal function under normal and pathological conditions.
**Introduction**

Dopaminergic modulation plays a key role in the prefrontal cortex (PFC). Prefrontal neurons receive dopaminergic input from a specific subpopulation of ventral tegmental area neurons which are strongly activated by aversive stimuli and have unique physiological properties (Lammel et al., 2011; Lammel et al., 2014). The infusion of dopamine D2 receptor agonists and antagonists into the PFC modulates working memory and set-shifting in rodents (Druzin et al., 2000; Floresco et al., 2006; St Onge et al., 2011). In nonhuman primates, prefrontal D2Rs are specifically necessary for neural activity associated with memory-guided saccades (Wang et al., 2004). Consistent with these animal studies, genetic variation in D2Rs modulates prefrontal activity and working memory in humans (Zhang et al., 2007). Prefrontal D2Rs have also been hypothesized to contribute to schizophrenia (Winterer and Weinberger, 2004; Durstewitz and Seamans, 2008), Tourette syndrome (Simonic et al., 1998; Minzer et al., 2004; Yoon et al., 2007; Steeves et al., 2010), and bipolar disorder (Minton et al., 2009). Thus, D2Rs play a major role in both normal and pathological prefrontal function.

Recently, our laboratory and others have shown that in the mouse medial prefrontal cortex (mPFC), three major subtypes of dopamine receptors (D1Rs, D2Rs, and D3Rs) are differentially expressed by distinct subtypes of layer 5 (L5) pyramidal neurons (Gee et al., 2012; Seong and Carter, 2012; Clarkson et al., 2017). Specifically, we found that D2Rs are selectively expressed within thick-tufted, subcortically-projecting (SC) L5 pyramidal neurons which exhibit prominent hyperpolarization-activated cationic current ($I_h$) (Gee et al., 2012). We found that in this population, pharmacological activation of D2Rs by the agonist quinpirole elicits a profound afterdepolarization (ADP) following depolarizing current injection. This ADP was also associated with a progressive membrane depolarization and
reduction in action potential amplitude during the depolarizing current pulse. The ADP was not observed under baseline conditions, and only occurred when NMDA receptors were activated, via either optogenetic stimulation of excitatory synapses or bath application of low dose (4 μM) NMDA. Similar to plateau potentials previously described in L5 pyramidal neurons (Milojkovic et al., 2005; Major et al., 2008), this ADP seems to rely on the intracellular accumulation of Ca\(^{2+}\) following influx through a combination of NMDA receptors (NMDARs) and L-type Ca\(^{2+}\) channels (LTCCs). In particular, it could be blocked by the NMDAR antagonist AP5, the selective LTCC antagonist nimodipine, and the intracellular Ca\(^{2+}\) chelator BAPTA; however, additional mechanistic details of intracellular signaling pathways related to this phenomenon are not known.

D2Rs are classically assumed to couple to inhibitory G proteins, reducing neuronal excitability by activating G\(_{\alpha i/o}\) and inhibiting adenylate cyclase (Bonci and Hopf, 2005). D2Rs are also known to signal through at least two other pathways: via the G\(_{\beta\gamma}\) subunits, and ß-arrestin. Here we explore two aspects of the intracellular signaling pathways through which the activation of D2Rs elicits ADPs in prefrontal SC-projecting neurons using a combination of electrophysiological, transgenic, pharmacological, and chemogenetic approaches. We find that the ability of synaptic stimulation to facilitate D2R-dependent ADPs several hundred milliseconds later depends on voltage-dependent (likely Ca\(^{2+}\)) currents. Furthermore, we show that the D2R-dependent ADP persists following disruptions to G\(_i\) or ß-arrestin signaling, and can be mimicked by chemogenetic activation of G\(_s\) signaling, suggesting that this phenomenon reflects novel intracellular signaling downstream of D2Rs.
Materials and Methods

Electrophysiology

Coronal brain slices (250 µm) including medial prefrontal cortex were made from adult mice of either sex that were at least 8 weeks old. We used the following transgenic mouse lines: wild-type C57BL/6J mice (www.jax.org/strain/000664), Drd1-Cre^{+/-} (line EY262; www.gensat.org), Drd2-Cre^{+/-} (line ER44; www.gensat.org), Drd2^{fl/fl} (https://www.jax.org/strain/020631), Rosa26^{PTX} (from Shaun Coughlin, University of California, San Francisco), and βarr2-KO (https://www.jax.org/strain/011130). Slicing solution was chilled to 4°C and contained (in mM): 234 sucrose, 26 NaHCO₃, 11 glucose, 10 MgSO₄, 2.5 KCl, 1.25 NaH₂PO₄, 0.5 CaCl₂, bubbled with 5% CO₂/95% O₂. Slices were incubated in artificial cerebrospinal fluid (aCSF) at 32°C for 30 minutes and then at room temperature until recording. aCSF contained (in mM): 123 NaCl, 26 NaHCO₃, 11 glucose, 3 KCl, 2 CaCl₂, 1.25 NaH₂PO₄, 1 MgCl₂, also bubbled with 5% CO₂/95% O₂.

Neurons were visualized using differential interference contrast or DODT contrast microscopy on an upright microscope (Olympus). Recordings were made using a Multiclamp 700B (Molecular Devices) amplifier and acquired with pClamp. Patch pipettes (2-5 MΩ tip resistance) were filled with the following (in mM): 130 KGluconate, 10 KCl, 10 HEPES, 10 EGTA, 2 MgCl₂, 2 MgATP, 0.3 Na₃GTP. In some experiments, the internal solution contained 1mM GDP-βS or 100μM Rp-cAMPs. All recordings were made at 32-34°C. Series resistance was compensated in all current clamp experiments and monitored throughout recordings. Recordings were discarded if Rs changed by >25%.

D2R expressing, subcortically-projecting neurons were often identified by fluorescent visualization of retrograde tracer Alexafluor-tagged cholera toxin subunit B (CTb) injected
into the MD thalamus. In some experiments, pyramidal neuron subtypes were identified based on characteristic firing patterns, specifically h-current-induced “sag” greater than 3mV in response to hyperpolarizing current pulses.

All bath-applied drugs were dissolved in water (4 µM NMDA, 10 µM (-)quinpirole, 10 µM CNQX, 10 µM H89, 1 µM CNO), DMSO (5 µM sulpiride, 5 Aripiprazole), or 1M HCl (200 µM baclofen) before being diluted in aCSF.

**Viral injection for expression of ChR2, fluorescent reporter, or DREADDs.**

Viral injections were performed using standard mouse stereotactical methods. Mice were anesthetized for the duration of the surgery using isofluorane gas. After cleaning, an incision was made in the scalp, the skull was leveled, and small burr holes were drilled over the brain region of interest using a dental drill. Virus was injected through the burr holes using a microinjector at a speed of 150 nL/minute and the scalp was closed using tissue adhesive (Vetbond).

For expression of ChR2 in CC neurons, we injected a Cre-dependent ChR2 virus (AAV5-Ef1-DIO-ChR2-eYFP, 0.5 µL) into the mPFC of Drd1-Cre<sup>+/−</sup> mice (>p28) and waited 3-4 weeks for trafficking of ChR2 to the axon terminals in mPFC.

For fluorescent reporters, we injected a Cre-dependent eYFP virus (AAV5-Ef1-DIO-eYFP, 1 µL) into mPFC of Drd2-Cre<sup>+/−</sup>, Drd2<sup>fl/fl</sup>, or Rosa26<sup>PTX</sup> mice (>p28) and waited 3-4 weeks. For retrogradely labelling SC-projecting neurons, Alexafluor-tagged cholera toxin subunit B (CTb; ThermoFisher) was first dissolved in PBS to make a 1.0mg/mL solution. 350nL of CTb were then injected into the MD thalamus at a rate of 100nL/min and waited 3-5 days before electrophysiological experiments.
To selectively knockout D2Rs in SC-projecting cells, we injected canine adenovirus-2 encoding Cre virus (Cav2-Cre, 0.5 μL; Institut de Génétique Moléculaire de Montpellier) into the MD thalamus as well as 1.5 μL AAV5-Ef1-DIO-eYFP into mPFC of Drd2^fl/fl mice (>p28) and patched from both fluorescent and non-fluorescent cells after waited 4-5 weeks.

For DREADD experiments in D2R-expressing pyramidal neurons, we injected a Cre-dependent virus expressing the Gq-DREADD (AAV5-hSyn-DIO-hM3D(Gq)-mCherry, 750 nL), Gs-DREADD (AAV5-hSyn-DIO-rM3D(Gs)-mCherry, 750 nL), or Gi-DREADD (AAV5-hSyn-DIO-hM4D(Gi)-mCherry, 750 nL) into the mPFC of Drd2-Cre/+ mice and patched from fluorescent cells after waiting 5 weeks for expression.

MD thalamus injection coordinates were A/P = -1.7, M/L = +/- 0.3, D/V = -3.45. mPFC injection coordinates were A/P = 1.7, M/L = +/- 0.3, D/V = -2.75

ChR2 stimulation

We stimulated ChR2 in projections from the contralateral PFC using 5ms flashes of light generated by a Lambda DG-4 (Sutter Instruments) high-speed optical switch with a 300 W Xenon lamp delivered through a 470 nm excitation filter. For stimulation of fibers, 5 different trains of random light flashes (intensity: ~2mW, total train duration: 2.5 sec, ~60 flashes/train, flash duration: 2 msec) were delivered at through a 40x objective every 13 seconds.

Experimental Design and Statistical Analysis

All data are shown as mean +/- 1 SEM. Statistical significance was accepted at the level p < 0.05. All statistical computations were performed using GraphPad Prism 7.0 software. We used student’s t-test to compare pairs of groups if data were normally
distributed (verified using Lillie test). If more than two groups were compared, we used ANOVA with post-hoc tests between groups corrected for multiple comparisons (Holm-Sidak or Bonferroni). The specific post-hoc test as well as exact F and corrected p values can be found in the text.
Results

The quinpirole-induced ADP depends on D2Rs

First, to confirm the basic properties of the D2R-mediated ADP we had described previously (Gee et al., 2012), we labeled SC-projecting neurons by injecting a retrogradely transported fluorescent tracer (Alexafluor-tagged cholera toxin subunit B; CTb) into the mediodorsal (MD) thalamus (Figure 1A). Then, we patched from CTb+ L5 pyramidal neurons in mPFC, and measured their voltage responses to depolarizing current pulses (250 msec duration) (Figure 1B). We did not observe an ADP following depolarizing current pulses in the presence of just NMDA (4 μM), however, following the addition of the D2R agonist (−)quinpirole (10 μM) to the bath, we observed a robust ADP for ~100-300 msec after the end of each depolarizing current pulse (Figure 1C: n = 11/9/11/4, baseline/NMDA/QPL/SUL; One-way ANOVA, $F_{(3,31)} = 16.36, p < 0.0001$; t-test, Bonferroni correction, baseline vs NMDA: $t_{(18)} = 0.1408$, baseline vs QPL: $t_{(20)} = 6.178, p < 0.0001$, NMDA vs QPL: $t_{(18)} = 5.721, p < 0.0001$). Subsequent wash-in of the selective D2/D3 antagonist sulpiride (5 μM) eliminated this quinpirole-induced ADP (t-test, Bonferroni correction, QPL vs SUL: $t_{(13)} = 3.723, p = 0.0031$).

To further confirm that this ADP was mediated by D2Rs, we injected canine adenovirus-2 encoding Cre (Cav2-Cre) (Hnasko et al., 2006) into MD thalamus to drive Cre expression within a subset of SC-projecting neurons in D2R conditional KO mice (Drd2<sup>fl/fl</sup>, Jackson Labs) (Bello et al., 2011) (Figure 1D). We then injected AAV to drive Cre-dependent expression of EYFP into the mPFC to identify SC-projecting neurons which expressed Cre, and from which D2Rs should have been knocked out. Finally, we made simultaneous recordings from pairs of EYFP+ and EYFP- neurons in L5 of the mPFC. We identified EYFP-negative SC-projecting neurons based on the presence of prominent
voltage sag and rebound afterdepolarization during responses to hyperpolarizing current pulses (Gee et al., 2012). We confirmed that the quinpirole-induced ADP still occurred in EYFP-negative SC-projecting neurons, but was completely absent from simultaneously recorded EYFP+ SC-projecting neurons (Figure 1E-F: n = 8 cell pairs; Mixed-model two-way ANOVA, $F_{(1,14)} = 20.15$, $p = 0.0005$; t-test, Holm-Sidak correction, eYFP+ baseline vs QPL: $t_{(7)} = 6.461$, $p < 0.0001$, eYFP- baseline vs QPL: $t_{(7)} = 0.1137$). This confirms that the quinpirole-induced ADP can be eliminated in a cell autonomous fashion by knocking out D2Rs.

**Hyperpolarizing current suppresses the ability of synaptic input to facilitate the ADP**

Next, we investigated mechanisms through which optogenetic stimulation of callosal synapses can unmask the quinpirole induced ADP (Gee et al., 2012). Again, bath application of quinpirole alone does not elicit an ADP. The ADP is only observed when quinpirole application is combined with the optogenetic stimulation of synaptic inputs, e.g., from the contralateral mPFC, or by bath application of low-dose NMDA. Synaptic stimulation can facilitate a subsequent ADP even when this stimulation is relatively weak, i.e., insufficient to elicit spiking in the postsynaptic neuron, and occurs several hundred milliseconds before the current pulse. Synaptic stimulation appears to act by recruiting NMDARs, since the ADP is blocked in the presence of AP5. We hypothesized that there could be at least two distinct mechanisms through which synaptic stimulation activates NMDARs and facilitates the ADP. First, synaptic stimulation may recruit significant NMDAR-mediated currents; in particular, even though the resulting EPSPs may be relatively modest when measured at the soma (e.g., ~1 mV in amplitude), they may be substantially larger in the dendrites leading to significant NMDAR-mediated Ca$^{2+}$ influx, which persists for several
hundred milliseconds contributing to the subsequent ADP. Alternatively, synaptic stimulation may trigger ongoing network activity which continues for several hundred milliseconds.

To distinguish between these two possibilities, we compared the magnitude of the quinpirole-induced ADP under various conditions (Figure 2B). In each case we elicited the ADP as follows: we injected AAV5-Ef1α-DIO-ChR2-EYFP into the mPFC in one hemisphere of Drd1-Cre (D1-Cre) mice (to label callosally-projecting D1+ neurons), then after waiting 3 to 4 weeks for ChR2 expression, we made recordings from L5 SC-projecting neurons in the contralateral hemisphere (Figure 2A). After delivering a train of irregularly occurring light flashes (total train duration: 2.5 sec, ~60 flashes/train, flash duration: 2 msec) to stimulate callosal terminals, we waited for 500 msec, then injected a depolarizing current pulse (+400 pA, 500 msec duration) to elicit the ADP. Each experiment consisted of four phases. First, we simply recorded the ADP following synaptic stimulation (control; Figure 2C). Second, we injected hyperpolarizing current (-200 pA) while delivering the train of light flashes in order to suppress voltage-dependent currents (e.g., NMDAR-mediated Ca2+ influx) associated with synaptic stimulation (“Exp 1”; Figure 2C). Third, we washed on CNQX (10 μM) to block AMPA receptors and eliminate recurrent network activity elicited by synaptic stimulation (“Exp 2”; Figure 2D). Fourth, we delivered hyperpolarizing current during the light trains while CNQX remained in the bath (“Exp 1 + 2”; Figure 2D). To quantify the ADP, we measured the amount of time required the membrane potential to return to baseline following the depolarizing current pulse (more precisely, the time for 90% decay).

The duration of the ADP was significantly reduced when we injected hyperpolarizing current during the delivery of light flashes (Figure 2E: n = 12; 46 ± 8% reduction in ADP duration; repeated-measures ANOVA, F = 8.659, p = 0.0024; t-test, Holm-Sidak correction, for ACSF vs. QPL without hyperpolarizing current: t(22) = 5.067, p = 0.0036; for QPL with vs.
without hyperpolarizing current: $t_{22} = 3.544, p = 0.0318$). By contrast, there was relatively little effect of CNQX application alone (t-test, QPL with vs. without CNQX, no hyperpolarizing current: $t_{22} = 0.051$). Notably, the combined effect of CNQX application and hyperpolarizing current injection was significantly larger than that of hyperpolarizing current injection alone – in fact the ADP was almost eliminated in this case (70 ± 5% reduction in ADP duration; t-test, QPL with vs. without CNQX and hyperpolarizing current: $t_{22} = 4.988, p = 0.0037$), suggesting that dendritic space clamp may be significantly improved in the presence of CNQX. Note: hyperpolarizing current delivered during the period of synaptic stimulation also seemed to attenuate other phenomena elicited by combined D2R activation + synaptic stimulation, e.g., the progressive membrane depolarization and reduction in spike amplitude during a current pulse.

Finally, we explored the timescales over which synaptic input could contribute to the quinpirole-induced ADP. For these experiments, we simply varied the delay between the trains of light flashes and the subsequent depolarizing current pulses. Remarkably, we could still observe an ADP lasting ~100 msec even when current injection occurred up to 10 seconds following synaptic stimulation (Figure 2F; n = 5; Repeated measures 2-way ANOVA: $F_{(10,40)} = 4.8, p = 0.0002$; t-test, Holm-Sidak correction, for individual time delays were as follows: p = 0.0071 (0.5s), 0.0007 (1s), < 0.0001 (1.5s), < 0.0001 (2.0s), 0.1190 (2.5s), 0.0338 (5.0s), 0.0003 (10s), 0.6671 (30s), 0.4942 (60s), 0.6671 (90s), 0.1248 (120s)).

Inhibiting adenylate cyclase / PKA-dependent signaling inhibits the quinpirole-induced ADP
Next, we turned our attention to the intracellular signaling pathways through which D2R activation recruits the ADP. First, to verify that the quinpirole-induced ADP depends on G-protein signaling, we performed experiments with 1mM GDP-βS in our intracellular pipette. GDP-βS is a non-hydrolyzable form of GDP that binds to G-proteins, and prevents the conformational changes which normally follow receptor ligand binding, depend on the hydrolysis of GTP to GDP, and trigger downstream signaling (Seong and Carter, 2012). Indeed, whereas in control experiments (without GDP-βS in the intracellular pipette), we observed robust ADPs following application of (-)quinpirole (10 μM) + NMDA (4 μM), we observed no ADPs when 1mM GDP-βS was included in the intracellular pipette (Figure 3A-B; n=7/6, K-gluconate/GDP-βs; two-way ANOVA: F_{(2,27)} = 6.406, p = 0.0053; t-test, Holm-Sidak correction, quinpirole+NMDA condition for K-gluconate vs GDP-βs: t_{(11)} = 4.241, p = 0.0007).

Next, since D2Rs canonically signal through inhibitory G-proteins, we decided to test how inhibiting adenylate cyclase / PKA signaling would affect the ADP. Since inhibitory G-proteins normally inhibit these pathways, we expected PKA inhibitors such as Rp-cAMPs or H89 to mimic or enhance the ADP. In fact we found that when we included Rp-cAMPs (100 μM) in the intracellular pipette, we could initially elicit a quinpirole-induced ADP in a perforated patch recording configuration, but that when we subsequently broke in, shifted to a whole cell recording, and dialyzed the cell with our intracellular solution, the quinpirole-induced ADP was abolished (Figure 3C & E; n = 4; ANOVA, F_{(2,9)} = 7.154, p = 0.0138; t-test, Bonferroni correction; ACSF vs QPL: t_{(6)} = 3.271, p = 0.029, QPL vs Rp-cAMPs: t_{(6)} = 3.281, p = 0.0285). This demonstrates that Rp-cAMPs is actually sufficient to abolish the quinpirole-induced ADP. Similarly, the presence of H89 (10 μM) in the bath prevented us from inducing an ADP with bath application of quinpirole and NMDA (Figure 3D-E; n = 6;
ANOVA, $F_{(1,17)} = 5.841$, $p = 0.0272$, t-test, Holm-Sidak correction, Control ACSF vs QPL: $t_{(10)} = 3.524$, $p = 0.0052$, H89 control vs H89 + QPL: $t_{(6)} = 0.2426$. These results suggest that inhibition of adenylate cyclase / PKA signaling is sufficient to block the quinpirole-induced ADP in a cell-autonomous fashion.

The quinpirole-induced ADP is independent of β-arrestin

Activation of D2Rs with relatively high doses of quinpirole (10 mM) and for somewhat prolonged periods of time (e.g., 15-20 min) may induce β-arrestin-mediated desensitization and internalization. Thus, the preceding observations could be explained by a model in which at baseline, D2Rs inhibit adenylate cyclase / PKA signaling, suppressing the ADP, but that quinpirole induces receptor internalization, terminating this suppression, and leading to the appearance of the ADP. To explore whether the ADP might depend on β-arrestin signaling via such a mechanism, we measured the ADP after disrupting β-arrestin signaling in two ways. First, we studied β-arrestin2 KO mice (βarr2-KO) (Bohn et al., 1999), and found that quinpirole and NMDA continued to elicit ADPs (Figure 4A-B; $n = 18/11$, β-arrestin2 KO/C57BL/6; two-way ANOVA, $F_{\text{genotype}(1,75)} = 0.517$, $F_{\text{drug}(3,75)} = 16.56$, $p < 0.0001$; t-test, Holm-Sidak correction, β-arrestin2 KO ACSF vs QPL: $t_{(34)} = 5.759$, $p < 0.0001$).

Moreover, in βarr2 KO mice, this quinpirole-induced ADP was still eliminated by the subsequent wash-in of the selective D2R antagonist sulpiride (5 μM), confirming that it still depends on D2Rs (Figure 4A-B; t-test, Holm-Sidak correction, β-arrestin2 KO QPL vs SUL: $t_{(20)} = 2.595$, $p = 0.0448$). Second, we found that quinpirole and NMDA could still elicit an ADP even in the presence of Aripiprazole (ARP; 5 μM; Figure 4 C-D; $n = 6$; ANOVA, $F_{(1,52)} = 0.005$, $F_{\text{drug}(2,52)} = 16.34$, $p < 0.0001$, t-test, Holm-Sidak correction, ARP ACSF vs ARP QPL: $t_{(10)} = 3.329$, $p < 0.0416$). ARP is a functionally selective D2R ligand that is a complete
antagonist with respect to β-arrestin signaling, but a partial agonist with respect to canonical G-protein signaling (Burris et al., 2002; Madhavan et al., 2013). Together, these two experiments show that D2Rs can elicit the ADP in the absence of β-arrestin.

**Pertussis toxin does not block the quinpirole-induced ADP**

Next, we wanted to examine whether the quinpirole-induced ADP depends on G\textsubscript{i} signaling at all. For this, we expressed pertussis toxin, which selectively blocks all G\textsubscript{i}-mediated signaling. We crossed Drd2-Cre (D2-Cre) mice to a transgenic line expressing the Cre-dependent S1 catalytic subunit of pertussis toxin (PTX S1) inserted into the Rosa26 (Rosa26-PTX) (Regard et al., 2007) such that PTX S1 would be expressed in D2R+ cells which normally exhibit the D2R-mediated ADP (Figure 5A). We verified that pertussis toxin blocked G\textsubscript{i}-mediated signaling in D2R+ neurons by comparing baclofen-induced currents in D2R+ (SC-projecting) neurons and D2R-negative (IT-projecting) neurons (distinguished based on the Cre-dependent expression of EYFP) (Figure 5B). Baclofen is a GABABR agonist which normally recruits GIRK currents via G\textsubscript{i}signaling. We found that in voltage clamp recordings in wild-type mice, bath application of baclofen (200 μM), elicits large outward currents in both D2R+ and D2R-negative neurons (n = 4 of each; Figure 5C). In D2-Cre / Rosa26-PTX mice, baclofen continued to elicit GIRK currents normally in D2R-negative neurons; however, baclofen-induced currents were completely abolished in D2R+ neurons (Figure 5C; n = 4/4, D2R+ and D2R- from each genotype; two-way ANOVA, F\textsubscript{genotype(1,12)} = 620, p < 0.001, t-test, Holm-Sidak correction, t\textsubscript{(6)} = 31.48, p < 0.0001). We found that the quinpirole-induced ADP was intact in D2R+ neurons from D2-Cre / Rosa26-PTX mice (Figure 5D-E; n = 6/5, Drd2Cre::Rosa26-PTX/C57BL6; repeated-measures two-way ANOVA, F\textsubscript{genotype(1,9)} = 0.0002, F\textsubscript{drug(2,18)} = 45.4, p < 0.0001; t-test, Holm-Sidak correction,
C57BL/6 ACSF vs QPL: \( t_{(8)} = 5.581, p < 0.0001 \), Drd2Cre::Rosa26-PTX ACSF vs QPL: \( t_{(10)} = 6.047, p < 0.0001 \). These experiments show that D2Rs elicit the ADP even in the absence of G\(_i\)-signaling.

**Gs but not Gi or Gq coupled DREADDs elicit a quinpirole-like ADP**

The preceding results strongly suggest that D2Rs do not act through G\(_i\) or β-arrestin to elicit the ADP. Furthermore, the fact that the PKA inhibitors Rp-cAMPs and H89 can inhibit the quinpirole-induced ADP raise the possibility that signaling through G\(_s\) or a G\(_s\)-like pathway may actually elicit this phenomenon. To test this directly, we injected virus to drive Cre-dependent expression of G\(_s\), G\(_i\), or G\(_q\)-coupled DREADDs into the mPFC of D2-Cre mice. We found that the presence of low dose (4 μM) NMDA together with clozapine-N-oxide (1 μM; CNO), which activates each DREADD, could elicit an ADP identical to the sort normally induced by quinpirole in SC-projecting cells expressing G\(_i\) DREADDs (Figure 6A-B; \( n = 8 \) for each DREADD; Mixed-model two-way ANOVA, \( F_{(2,21)} = 15.94, p < 0.0001 \); t-test, Holm-Sidak correction, G\(_s\)-DREADD ACSF vs CNO+NMDA: \( t_{(14)} = 7.01, p <0.0001 \)). Activation of the G\(_s\)-DREADD increases the number of action potential during and immediately following the current pulse (i.e. including the ADP). By contrast, we observed no effect of CNO in mice expressing the G\(_q\) DREADDs. Furthermore, in mice expressing the G\(_i\) DREADDs, bath application of 1 μM CNO decreased action potential firing in response to a depolarizing current step (Figure 6C; \( n = 8 \) for each DREADD ; Mixed-model two-way ANOVA, \( F_{(2,21)} = 20.79, p < 0.0001 \); t-test, Holm-Sidak correction, G\(_i\)-DREADD ACSF vs CNO: \( t_{(14)} = 3.319, p = 0.0065 \), G\(_s\)-DREADD ACSF vs CNO: \( t_{(14)} = 5.621, p < 0.0001 \), confirming functional Gi DREADD expression in these neurons, and consistent with a G\(_i\)-induced reduction of neuronal excitability (Bonci and Hopf, 2005).
Discussion

Here we follow up on our earlier work (Gee et al., 2012), which described a new mechanism through which D2R activation could elicit an increase in excitability, specifically an ADP, within subcortically-projecting L5 pyramidal neurons in the mPFC. Several aspects of this phenomenon were unusual. First, we observed the ADP using relatively high doses of quinpirole (5-20 μM). This was in contrast to other effects of quinpirole can sometimes be observed using doses ~1 μM (Tseng et al., 2007). Second, the ADP represents an increase in excitability, whereas D2R activation has classically (in the striatum) been assumed to inhibit neuronal excitability. Third, this phenomenon was not readily observed under baseline conditions in brain slices, but required synaptic stimulation. All of these relatively unique features raised several questions, e.g., is this phenomenon, which is elicited by quinpirole and blocked by sulpiride, really mediated by D2Rs? If so, is this phenomenon mediated by canonical D2R signaling, the desensitization or internalization of D2Rs, or another pathway (e.g. β-arrestin-mediated signaling) altogether? And finally what is the mechanism through which synaptic input regulates the occurrence of an ADP several hundred milliseconds later?

This study answers those questions, at least in part. Knocking out D2Rs abolishes the quinpirole-induced ADP in a cell-autonomous manner, confirming that it is D2R-mediated. Hyperpolarizing current injection, but not CNQX, prevents synaptic stimulation from facilitating the quinpirole-induced ADP, showing that this effect depends on the ability of synaptic stimulation to recruit voltage-dependent currents, likely NMDAR-mediated Ca\(^{2+}\) influx. Finally, the quinpirole-induced depolarization is blocked by inhibitors of PKA signaling, insensitive to pertussis toxin, and mimicked by Gs-DREADD stimulation. Taken together, our observations, following numerous manipulations targeting multiple aspects of
G-protein coupled signaling, suggest that D2R activation elicits the ADP by stimulating cAMP / PKA signaling that is downstream of G\textsubscript{s} (Figure 7).

Importantly, neither quinpirole nor sulpiride distinguishes between D2Rs and D3Rs. Thus, pharmacology alone could not rule out the possibility that D3Rs, along with D2Rs, contribute to the quinpirole-induced ADP. However, another recent study shows that D3Rs are expressed by a distinct population of prefrontal neurons, and not by the prefrontal SC-projecting neurons which are the subject of this study (Clarkson et al., 2017).

**Limitations**

Of course, our experiments have some important limitations. For example, H89 inhibits other kinases besides PKA, although Rp-cAMPs (which we delivered intracellularly) is more selective. Our experiments have not directly tested whether G\textsubscript{i}-associated βγ subunits might contribute to the effects we have observed by stimulating PKA signaling, but we consider this possibility unlikely, because the D2R-induced afterdepolarization is not blocked by pertussis toxin. By preventing dissociation of the heterotrimeric G-protein complex, pertussis toxin should block signaling by both α and βγ subunits associated with G\textsubscript{i}. Similarly, although our results do suggest that D2Rs elicit an afterdepolarization via signaling pathways that are classically downstream of G\textsubscript{s}, we have not determined to what extent G\textsubscript{s}-associated βγ subunits might contribute to or modulate these effects (Figure 7). Finally, although our experiments with pertussis toxin strongly argue against a role for Gi-mediated signaling in the D2R-induced ADP, and our DREADD experiments show that Gs-mediated signaling is sufficient to elicit this phenomenon, we cannot completely rule out the possibility that native G\textsubscript{q}-signaling might contribute to this phenomenon in ways that are not captured by simply stimulating Gq DREADDs.
Relationship to other modes of D2R signaling

D2R mediated signaling is diverse. D2Rs have classically been assumed to inhibit neuronal excitability via G\(_i\) signaling, e.g., in indirect pathway medium spiny neurons and dopaminergic terminals (Bonci and Hopf, 2005). However other studies have described novel mechanisms whereby D2Rs can signal through G\(_{\beta\gamma}\) subunits or interactions with β-arrestin (Beaulieu et al., 2005) including in prefrontal interneurons (Urs et al., 2016). These results enlarge the family of possible signaling pathways downstream of D2R activation. At the same time, this study, which has focused on cellular physiology, does not elucidate the molecular or biochemical details of this pathway.

Relationship to D2R heterogeneity and heteroreceptors

D2Rs comprise 6 introns and are known to undergo alternative splicing giving rise to at least 2 major isoforms (Usiello et al., 2000; De Mei et al., 2009). There has also been controversy about whether D2Rs might form heteroreceptors with D1Rs in the cortex (Beaulieu and Gainetdinov, 2011; Zhang et al., 2011). Findings like these suggest that D2Rs exist in heterogeneous forms, which may explain how D2Rs can couple to various intracellular signaling pathways, giving rise to distinct effects on neuronal excitability, in different neuronal populations. It will be interesting for future studies to characterize the specific forms of D2Rs that are expressed in prefrontal SC-projecting neurons. In particular, there may be specific domains present in D2R isoforms expressed in prefrontal SC-projecting neurons which enable coupling to the G\(_s\) pathways, D1Rs, or other effectors.
Functional implications

The phenomenon studied here is one whereby synaptic activity, even when subthreshold with respect to action potential generation in the post-synaptic neuron, can enhance neuronal excitability several hundreds of milliseconds or even up to 10 seconds later. This could enable post-synaptic neurons to integrate their inputs over relative long durations. An alternative interpretation of this mechanism is that one set of inputs could switch neurons into a “high gain” mode whereby responses to subsequent inputs are potentiated, relative to the neuron’s baseline state. These effects could help to generate patterns of prefrontal activity e.g., the sequential patterns observed during the delay period of working memory or cued rule switching tasks (Bolkan et al., 2017; Schmitt et al., 2017). In these tasks, inhibiting prefrontal activity during the early portion of a delay disrupts subsequent sequential activity – early activity, if paired with D2R activation, may serve to “prime” the responses of neurons that fire later during the delay via this mechanism.

In this and previous studies (Gee et al., 2012) we have shown that optogenetically stimulating callosal inputs is sufficient to facilitate the D2R-dependent ADP. It will be interesting to explore whether other sources of inputs to mPFC can generate similar effects. We also have used trains of light flashes to stimulate synapses over periods of 2.5 seconds – it will be interesting to determine how this phenomenon depends on the duration of synaptic input. Finally, we have generally used patterns of synaptic stimulation that elicit EPSPs which are subthreshold for action potential generation in the postsynaptic neuron. An outstanding question is whether stronger patterns of stimulation are more or less effective for eliciting this phenomenon.
Clinical implications

Dopaminergic innervation of the prefrontal cortex is disrupted in schizophrenia (Slifstein et al., 2015), mutations in D2Rs are associated with schizophrenia (Schizophrenia Working Group of the Psychiatric Genomics Consortium, 2014), and all known clinically effective antipsychotics block D2Rs (Seeman, 2002). Together these observations raise the possibility that prefrontal D2R function may be abnormal in schizophrenia, although the precise nature of such dysfunction and its relationship to symptoms, remain largely unknown. As better models for schizophrenia are developed, e.g., based on neurons derived from patient iPS cells or mice with mutations in genetic loci strongly implicated in schizophrenia, it will be interesting to study whether this unusual form of prefrontal D2R signaling is altered.

Conclusion

Dopamine D2 receptors can enhance the excitability of subcortically-projecting pyramidal neurons in layer 5 of medial prefrontal cortex through a pathway that depends on increased cAMP / PKA signaling. This increase in neuronal excitability also appears to require the activation of synaptic NMDARs and voltage-dependent Ca\textsuperscript{2+} influx during the preceding several seconds. Important next steps will be understanding how this phenomenon contributes to normal brain function and is disrupted in the setting of diseases such as schizophrenia.
Figure 1. The quinpirole-induced afterdepolarization depends on D2Rs.

A) Schematic of coronal brain slice: we recorded from subcortically-projecting (SC) L5 pyramidal neurons labeled with CTB488 in the mouse mPFC (top). Images of a recorded neuron in DIC and exhibiting 488 fluorescence (bottom). B) Example pyramidal cell responses to hyperpolarizing and depolarizing current steps at baseline (black) and after application of drug (4 µM NMDA, purple; 4 µM NMDA + 10 µM Quinpirole, blue; 4 µM NMDA + 10 µM Quinpirole + 5µ Sulpiride, green). Arrow indicates the quinpirole-induced afterdepolarization (ADP). C) Changes in the ADP, quantified by the amount of time required for the membrane potential to return to baseline following the depolarizing current pulse (more precisely, the time for 90% decay), after pharmacological manipulations listed above. n = 11/9/11/4, baseline/NMDA/QPL/SUL. ***p < 0.0001. D) Experimental design: Drd2*fl/fl mice were injected with Cav2-cre virus in the MD thalamus and DIO-EYFP virus in the mPFC 4-5 weeks prior to slice experiments. E) Example pyramidal cell responses to hyperpolarizing and depolarizing current steps in 4 µM NMDA + 10 µM Quinpirole for EYFP negative (control) and EFYP positive (Drd2 knockout) cells (top). Images of a recorded neuron in DIC showing EYFP fluorescence (bottom). F) The quinpirole-induced ADP (arrows) is absent in cells lacking Drd2. n = 8 cell pairs. ***p < 0.0001.
Figure 2. Hyperpolarizing current suppresses the ability of synaptic input to facilitate the quinpirole-induced ADP.
A) Experimental design: We recorded from ChR2-negative layer 5 neurons while stimulating ChR2-expressing axon terminals from the contralateral mPFC with trains of light flashes (470 nm, 2.5 msec, ~2 mW). B) For each cell with bath application of 10 μM quinpirole, we recorded the neuronal response under four different stimulation paradigms: synaptic stimulation (control, black); hyperpolarizing current (-200 pA) during the train of light flashes (“exp 1”, blue); synaptic stimulation in 10 μM CNQX (“exp 2”, red); hyperpolarizing current during light flashes while in 10 μM CNQX (purple). C-D) Responses of a layer 5 SC-projecting pyramidal neuron to current injection before (left panel) and immediately following (middle and right panels) optogenetic stimulation of synaptic inputs. Blue bars indicate the times of light flashes. E) Quinpirole-induced ADP is reduced by injection of hyperpolarizing current. n = 12. *p = 0.0318, **p = 0.0037. F) The quinpirole-induced ADP persists up to 10 seconds after the synaptic stimulation. n = 5. *p < 0.0001-0.0338.
Figure 3. Inhibiting adenylate cyclase / PKA-dependent signaling inhibits the quinpirole-induced ADP
A) Sample traces in ACSF (black) and 4 μM NMDA + 10 μM quinpirole (blue) in control K-glucocate internal as well as internal including 1 mM GDP-βS, a G-protein inhibitor. B) GDP-βS blocks the quinpirole-induced ADP in a cell-autonomous fashion. n=7/6, K-glucocate/GDP-βs. ***p = 0.0007. C) Sample traces in ACSF (black) and 4 μM NMDA + 10 μM quinpirole (blue, left) in perforated patch and in whole-cell patch clamp dialyzed with internal solution containing 100 μM Rp-cAMPs (blue, right). D) Sample traces with bath application of 10 μM H89 in ACSF (black) and 4 μM NMDA + 10 μM quinpirole (blue). E) Inhibition of PKA suppresses the quinpirole induced ADP. n = 4 for perforated patch and 6 for H89. *p = 0.0285. **p = 0.0052.
Figure 4. The quinpirole-induced ADP is independent of β-arrestin.

A) Example recordings from β-arrestin2 KO mice in ACSF (black) and after application of drug (4 µM NMDA, purple; 4 µM NMDA + 10 µM Quinpirole, blue; 4 µM NMDA + 10 µM Quinpirole + 5µM Sulpiride, green). B) The quinpirole-induced ADP persists in β-arrestin2 KO mice. n = 18/11, β-arrestin2 KO / C57BL/6. ***p < 0.0001. C) Example traces in 5 M aripiprazole (ARP) at baseline (black) and after application of additional drugs (4 µM NMDA, purple; 4 µM NMDA + 10 µM Quinpirole, blue). D) The quinpirole-induced ADP persists in the presence of ARP, a biased D2R ligand that selectively antagonizes β-arrestin signaling. n = 6. ***p < 0.0001, * p=0.0416)
Figure 5. Pertussis toxin blocks G\textsubscript{i}-signaling but does not block the quinpirole-induced ADP

A) Experimental design: We crossed D2-Cre mice to Rosa26-PTX mice and injected the offspring in the mPFC with a Cre-dependent virus to drive EYFP expression. B) We then recorded from D2+ pyramidal neurons (i.e., EYFP+) and D2-negative pyramidal neurons (i.e., EYFP-negative) in L5 of mPFC. C) Pertussis toxin selectively suppresses baclofen-induced GIRK currents in D2+ neurons of Drd2Cre::Rosa26-PTX mice. n = 4. ***p < 0.0001. D) Sample traces from Drd2Cre::Rosa26-PTX mice in ACSF (black) and after application of drug (4 µM NMDA, purple; 4 µM NMDA + 10 µM Quinpirole, blue). E) The quinpirole-induced ADP is not blocked by pertussis toxin. n = 6/5, Drd2Cre::Rosa26-PTX/C57BL/6. ***p < 0.0001.
Figure 6. Signaling through Gs but not Gi or Gq coupled DREADDs elicit a quinpirole-like ADP

A) Example traces from L5 mPFC pyramidal neurons from D2-Cre mice injected with virus to derive Cre-dependent expression of either Gi (left), Gs (middle) or Gq (right) coupled DREADDs, at baseline (top, black) or in 4 μM NMDA + 1 μM CNO (bottom, blue). B) Activation of the Gs-DREADD with 1 μM CNO together with low dose NMDA elicits an ADP. n = 8. ***p < 0.0001. C) Activation of the Gi-DREADD with 1 μM CNO reduces the number of action potentials fired when +400pA depolarizing current is injected for 250msec. By contrast, activation of the Gs-DREADD increases the number of action potential during and immediately following the current pulse. n = 8. **p = 0.0065. ***p < 0.0001.
Figure 7. Schematic models for D2R signaling in mPFC SC-projecting projecting L5 pyramidal neurons

A) Schematic of the canonical D2R signaling cascade acting via Gi to inhibit cAMP/PKA activity. B) Schematic of the new model showing how D2R signaling might act via Gs or a Gs-like pathway to stimulate cAMP/PKA and thereby elicit an ADP. C) Summary of how various experimental manipulations we tested affected the quinpirole-induced ADP.
References


CHAPTER III

Inducible expression of mutant human DISC1 in mice abolishes quinpirole-induced afterdepolarization in mPFC
Introduction

Schizophrenia is a debilitating psychiatric disorder affecting millions of individuals (Cloutier et al., 2016). Patients with schizophrenia often experience a combination of ‘positive’ symptoms, including delusions and hallucination, ‘negative’ symptoms, including affective flattening and avolition, as well as cognitive impairments including deficits in working memory, cognitive flexibility, and executive functions (Meltzer, 1997; Evans et al., 2004). The prefrontal cortex (PFC) has been shown to play a critical role in many of these cognitive functions impaired in schizophrenia (Druzin et al., 2000; Floresco et al., 2006; Durstewitz and Seamans, 2008). Many neurological disorders including schizophrenia have been associated with imbalances in dopamine neurotransmission (Winterer and Weinberger, 2004). It has been suggested that balance between dopamine D1 receptor (D1R) and dopamine D2 receptor (D2R) activation is crucial for optimizing the signal-to-noise ratio of local microcircuits and thereby normal PFC function (Winterer and Weinberger, 2004; Winterer et al., 2004; Durstewitz and Seamans, 2008; Durstewitz et al., 2010).

The dopamine hypothesis of schizophrenia theorizes that excessive D2R activation within the PFC contributes to many of the cognitive impairments associated with the disorder (Coyle et al., 2012; Urs et al., 2017). The two early cornerstones of this hypothesis were the findings that antipsychotic drugs block D2Rs and increase presynaptic dopamine metabolism and that dopamine-mimetics are psychotogenic (Seeman et al., 1976; Beaulieu, Gainetdinov and Caron, 2009). Further studies have shown that infusion of D2R agonists and antagonists into the rodent PFC modulates cognitive functions disrupted in schizophrenia, including working memory and set-shifting (Druzin et al., 2000; Floresco et al., 2006; St Onge et al., 2011). Current antipsychotics, however, are not effective at reversing cognitive deficits associated with schizophrenia. Improving our understanding of
how normal dopaminergic modulation is perturbed in schizophrenia, and other neuropsychiatric diseases, can aid in the development of safer and more effective drug therapies.

Transgenic animal models of psychiatric disease can be useful tools for dissecting the function of individual components of neurotransmitter systems and their contribution to aberrant behaviors (Gainetdinov et al., 2001; Marcotte et al., 2001). Disrupted-in-Schizophrenia 1 (DISC1) is a gene known as a risk factor for mental illness and has been associated with impairments in dopamine neuromodulation (Jacobs et al., 1970). Several different DISC1 mutant animal models exhibit behavioral as well as neurochemical feature relevant to human schizophrenia pathologies (Su et al., 2014; Dahoun et al., 2017).

DISC1 is a scaffold protein which interacts with several other proteins in the dopamine system, including serine/threonine protein kinase B (Akt) and glycogen synthase kinase-3 (GSK-3). In DISC1 mouse models there is an upregulation of D2R expression in the mPFC and striatum altering typical D2R neuromodulation (Niwa et al., 2013). Dopamine receptors are G protein-coupled receptors (GPCRs) and D2Rs are canonically thought to act via coupling to inhibitory G-proteins (G_i) and thereby acting to inhibit adenylate cyclase, reducing intercellular levels of cAMP and protein kinase A (PKA) activation (Bonci and Hopf, 2005). However, β-arrestin, normally involved in receptor internalization after prolonged activation, can also form a complex with D2R and initiate signaling cascades which are independent from the D2R-G-protein signaling pathway (Lefkowitz, 2013). Activation of this β-arrestin pathway by dopamine inactivates Akt by dephosphorylation resulting in activation of its negatively regulated substrate (Beaulieu et al., 2005, 2007). Aripiprazole, a new generation antipsychotic acts as a biased agonist of the D2R pathway, partially agonizing G-protein signaling and antagonizing β-arrestin, suggesting that this non-canonical signaling pathway has a role in its antipsychotic effects (Urs et al., 2016, 2017). Interestingly
the D2R-DISC1 complex has also been shown to contribute to the antipsychotic effects of D2R antagonists (Su et al., 2014). DISC1 has also been shown to play important roles in regulating the cAMP pathway as well as Akt / GSK3 (El-Hassar et al., 2014; Su et al., 2014).

It is still not fully understood how D2Rs, G-proteins, β-arrestin, and DISC1 interact to modulate neuronal activity in response to dopamine or how these signaling complexes are impaired in schizophrenia. As discussed in chapter II, we have identified a new mechanism through which D2R activation can enhance excitability of pyramidal neurons in the PFC via pathways associated with Gs, rather and Gi, coupled proteins (Robinson and Sohal, 2017). We found that in layer 5 (L5) subcortical (SC) projection neurons, pharmacological activation of D2Rs by the agonist quinpirole elicits a profound afterdepolarization (ADP). Given that DISC1 is a susceptibility gene for schizophrenia and other psychiatric disorders and that the DISC1-D2R complex has been shown to regulate other non-canonical D2R signaling pathways including cAMP and Akt / GSK3 pathways, we investigated the role of DISC1 in the quinpirole-induced afterdepolarization. In this chapter, I will present results that subcortically (SC) projecting D2R positive layer 5 (L5) pyramidal neurons in mice with a dominant-negative mutation in disrupted in schizophrenia 1 (DISC1) lack the Gi and β-arrestin independent quinpirole-induced afterdepolarization (ADP).
Materials and Methods

Animals

Control experiments were conducted using wild-type C57BL/6J mice (www.jax.org/strain/000664). DISC1 experimental animals were breed by crossing Tg(tetO-DISC1*)1001Plet/J transgenic mice (https://www.jax.org/strain/008790, Pletnikov et al., 2008), which express a myc peptide-tagged, mutant (truncated) human disrupted-in-schizophrenia 1 (DISC1) gene under control of the tetracycline operator (tetO), with CaMKII-tTA transgenic mice (https://www.jax.org/strain/003010) expressing tetracycline-controlled transactivator protein (tTA) under regulatory control of the forebrain-specific calcium-calmodulin-dependent kinase II (CamkIIa) promoter thereby creating a bitransgenic animal expressing mutant DISC1 in CamkIIa expressing neurons (Fig. 1A).

Electrophysiology

Electrophysiological recordings were performed as described in Chapter II, Materials and Methods, “Electrophysiology.” In brief, coronal mPFC slices were made from adult mice of either sex at post-natal day (P)56-70. Following 30 minutes incubation at 32°C in ACSF recording solution, slices were moved to room temperature. Whole-cell current clamp recordings were then performed in mPFC L5, using a K+-gluc-based internal solution as reported in Chapter II. Pyramidal neuron subtypes were identified based on characteristic firing patterns, specifically h-current-induced “sag” greater than 3mV in response to hyperpolarizing current pulses (Gee et al., 2012). In some experiments, D2R expressing subcortically-projecting neurons were often identified by fluorescent visualization of retrograde tracer Alexafluor-tagged cholera toxin subunit B (Ctb) injected into the MD thalamus 3-5 days prior to electrophysiological experiments. All bath-applied drugs were dissolved in water (4 µM NMDA and 10 µM (-)-quinpirole) before being diluted in ACSF.
**CTb injection for retrograde tracing.**

Injections were performed using standard mouse stereotactical methods. Mice were anesthetized for the duration of the surgery using isoflurane gas. After cleaning, an incision was made in the scalp, the skull was leveled, and small burr holes were drilled over the brain region of interest using a dental drill. Alexa Fluor 488 conjugated to cholera toxin subunit B (CTb-488) was injected through the burr holes into the mediodorsal (MD) thalamus using a microinjector at a speed of 150 nL/minute and the scalp was closed using tissue adhesive (Vetbond). MD thalamus injection coordinates were A/P = -1.7, M/L = +/- 0.3, D/V = -3.45. (Fig. 1B)

**Experimental Design and Statistical Analysis**

All data are shown as mean +/- 1 SEM. Statistical significance was accepted at the level p < 0.05. All statistical computations were performed using GraphPad Prism 7.0 software. We used student’s t-test to compare pairs of groups if data were normally distributed (verified using Lillie test). If more than two groups were compared, we used ANOVA with post-hoc tests between groups corrected for multiple comparisons (Holm-Sidak or Bonferroni). The specific post-hoc test as well as exact F and corrected p values can be found in the text.
Results

**DISC1 mutants have increased input resistance in layer 5 pyramidal neurons.**

To investigate the function of *disrupted in schizophrenia 1 (DISC1)*, we generated a bitransgenic mouse expressing a dominant-negative mutant human DISC1 gene under control of the CaMKII promoter by utilizing the Tet-off expression system and a crossing a transgenic animal with inducible expression of the mutant human DISC1 gene (Pletnikov et al., 2008) and a transgenic animal expressing tTA under regulatory control of CaMKII (Fig. 1A).

Since the DISC1-D2R complex has been strongly implicated in schizophrenia, we wanted to investigate the role of DISC1 in dopamine D2 receptor (D2R) expressing neurons within the mPFC. We took advantage of the fact that in deep layers of the mPFC, D2Rs are mainly expressed by subcortical (SC) projection neurons, and injected a retrogradely transported fluorescent tracer (Alexa Fluor 488 conjugated cholera toxin subunit B; CTb-488) into the mediodorsal (MD) thalamus at P52-67 to label SC projection neurons and therefore putative mPFC D2R+ neurons. Then, 3 to 5 days after injection, we performed whole-cell current-clamp recordings, measuring their voltage responses to a series of hyperpolarizing and depolarizing pulses (250 msec duration) from L5 mPFC SC-projecting neurons identified by Ctb-488 fluorescence (Fig. 1B).

D2R+ pyramidal neurons can also be identified based on their prominent hyperpolarization-activated cation currents (Ih), which are mediated by hyperpolarization-activated cyclic nucleotide-gated (HCN) channels (Gee et al., 2012; Clarkson et al., 2017). In D2R+ neurons, Ih gives rise to a characteristic “sag” and rebound during and following hyperpolarizing current injection. We first estimated Ih by measuring the membrane potential sag and rebound elicited by a –250 pA current step (Fig. 1C) confirming this characteristic sag and rebound property in SC projection neurons labeled by Ctb-488. For
wildtype C57BL/6 and CaMKII::DISC1 mice, there was no difference in this estimate for Ih in SC projection neurons (Fig. 1G: n = 9/10, C56BL6/CaMKII::DISC1; unpaired t-test: t(17) = 0.04, n.s.). We also characterized other intrinsic electrophysiological properties in Ctb-488+ L5 pyramidal neurons within mPFC. We found no difference in resting membrane potential between neurons from wildtype and CaMKII::DISC1 mice (Fig. 1D: n = 9/10, C56BL/6/CaMKII::DISC1; unpaired t-test: t(17) = 0.06, n.s.) but did observe an increase in input resistance when subjected to a −50 pA current pulse in L5 SC projection neurons from CaMKII::DISC1 mice (Fig. 1E: n = 9/10, C56BL6/CaMKII::DISC1; unpaired t-test: t(17) = 3.01, p = 0.0079).

**DISC1 mutants lack the quinpirole-induced ADP.**

As previously shown in Chapter II of this dissertation, D2R activation elicits afterdepolarizations (ADPs) in this population of SC-projecting pyramidal neurons within L5 of the mPFC. These D2R-induced ADPs only occur following synaptic input which activates NMDA receptors (NMDARs) (Gee et al., 2012; Robinson and Sohal, 2017). To investigate whether mutant DISC1 disrupts this D2R-dependent ADP, we analyzed the neurons voltage responses to depolarizing current pulses (+400pA, 250 msec duration). To quantify the ADP, we measured the amount of time required for the membrane potential to return to baseline following the depolarizing current pulse (more precisely, the time for 90% decay). In C57BL/6 wildtype mice we confirmed that ADPs are absent following depolarizing current pulses in ACSF or the presence of just NMDA (4 μM), however, following the addition of the D2R agonist (-)quinpirole (10 μM) to the bath, we observed a robust ADP for ~100-300 msec after the end of each depolarizing current pulse (Fig. 1C(top), F: C57BL/6 n = 8, baseline / NMDA / QPL; Oneway ANOVA, F(3,21) =45.3, p < 0.0001; t-test, Tukey correction, baseline vs QPL: q(14) = 11.83, p < 0.0001, NMDA vs QPL: t(14) = 11.47, p < 0.0001). The
time for 90% decay back to baseline following the depolarizing current pulse is not statistically different between C57BL/6 wildtype and CaMKII::DISC1 mice in either control ACSF (Fig. 1F: n = 8/10, C56BL6/CaMKII::DISC1; unpaired t-test: t_{(16)} = 1.773, n.s.) or NMDA (Fig. 1F: n = 8/6, C56BL6/CaMKII::DISC1; unpaired t-test: t_{(12)} = 1.909, n.s.).

However, in CaMKII::DISC1 mice, the quinpiroline-induced ADP is completely abolished (Fig. 1F: CaMKII::DISC1 n = 7, baseline / NMDA / QPL; One-way ANOVA, F_{(3,21)} = 2.397, n.s.; n = 8/7, C56BL6/CaMKII::DISC1; unpaired t-test: t_{(13)} = 6.32, p < 0.0001), suggesting that the normal DISC1 function is necessary for the quinpiroline-induced ADP.
Discussion

Here we follow up our earlier work (Gee et al., 2012; Robinson and Sohal, 2017), which described a new mechanism through which dopamine D2 receptor (D2R) activation could elicit an increase in excitability, specifically an ADP, within subcortically projecting L5 pyramidal neurons in the mPFC by stimulating cAMP / PKA signaling that is downstream of Gs (Chapter II, Figure 7).

As discussed in Chapter II of this dissertation, D2R mediated signaling is diverse and the results presented in Chapter II only serve to enlarge the family of possible signaling pathways downstream of D2R activation. The work presented in both this chapter and Chapter II, focus on cellular physiology and do not elucidate the molecular or biochemical details of this pathway. Many recent studies suggest that D2Rs exist in heterogeneous forms, which may explain how D2Rs can couple to various intercellular signaling pathways, giving rise to distinct effect on neuronal excitability, in different neuronal populations. Disrupted-in-schizophrenia 1 (DISC1) is a scaffolding protein known to complex with D2R. Recent studies suggest important roles for this D2R-DISC1 complex in regulation of β-arrestin binding and internalization of the D2R as well as regulation of downstream signaling pathway including cAMP and Akt / GSK3. It will be interesting for future studies to characterize the D2R-DISC1 complex and the molecular mechanisms of its regulatory role on D2R intracellular signaling pathways.

Clinical Implications

Dysfunction of the dopamine system is heavily implicated in schizophrenia. Although the precise nature of such dysfunction remains unclear, many studies raise the possibility that prefrontal D2R function specifically may be abnormal in schizophrenia: mutations in D2Rs are associated with schizophrenia (Schizophrenia Working Group of the Psychiatric
Genomics Consortium, 2014), all known clinically effective antipsychotics block D2Rs (Seeman, 2002), and D2R agonists and antagonists modulate several cognitive functions disrupted in schizophrenia (Druzin et al., 2000; Floresco et al., 2006a; St Onge, Abhari and Floresco, 2011).

An interesting line of future investigation involves studying the impact of this novel mechanism for D2R-dependent enhancement of layer 5 mPFC excitability on performance during PFC-dependent behavioral tasks. In vivo cannula infusion of CNO into the mPFC of awake behaving mice injected with Gs-DREADD (AAV5-hSyn-DIO-rM3D(Gs)-mCherry) could mimic the quinpirole-induced ADP providing a mechanism for investigating the impact of this novel mechanism. While recent concerns have been raised regarding CNO being metabolized to clozapine, this is unlikely to occur over the time course of slice experiments and can be accounted for in behavior experiments with a CNO control group of uninjected animals (Smith et al., 2016; Gomez et al., 2017). It would be interesting to determine whether CNO paired with synaptic stimulation can elicit an ADP in CamKII::DISC1 mice injected with Gs-DREADD indicating whether DISC1 acts up or downstream of Gs. DISC1 mice have already been shown to have deficits in social interaction and behavioral flexibility. An interesting set of experiments would be to attempt to rescue performance during PFC-dependent behavior tasks by induction of the D2R/Gs-mediated ADP in CaMKII::DISC1 mice.

Improving our understanding of the D2R signaling mechanisms which are altered in schizophrenia will provide great insight aiding in the development of new and improved therapies to treat the symptoms of cognitive dysfunction with fewer adverse side effects.
Conclusion

Dopamine D2 receptors can enhance the excitability of subcortically-projecting pyramidal neurons in layer 5 of medial prefrontal cortex through a pathway that depends on

These results show that this unusual physiological phenomenon, in which D2Rs enhance cellular excitability in a manner that depends on synaptic input and is mediated through Gs associated signaling pathways is disrupted in at least one DISC1 mouse model of schizophrenia. As better models for schizophrenia are developed, it will be interesting to study whether this unusual form of prefrontal D2R signaling is also altered. Important next steps will be understanding how this phenomena contributes to normal brain function and is disrupted in other models of schizophrenia or psychiatric disease.
Figure 1. L5 mPFC neurons from \textit{DISC1} mutants have increased input resistance and lack the quinpirole-induced ADP.
A) Experimental design: We crossed Tg(tetO-DISC1) mice to CaMKII-tTA mice to drive expression of inducible mutant human DISC1 under control of the CaMKII promoter. B) We injected offspring in the mediodorsal thalamus with Ctb-488 to identify SC-projecting pyramidal neurons within the mPFC. 3 to 5 days later we recorded from Ctb-488+ pyramidal neurons in L5 of mPFC. C) Example pyramidal cell responses to hyperpolarizing (-250pA) and depolarizing (+400pA) current steps at baseline (black) and after application of drug (4 µM NMDA, purple; 4 µM NMDA + 10 µM Quinpirole, blue) in wildtype C57BL/6 mice (top) and CaMKII::DISC1 mice (bottom). Blue arrow indicates the quinpirole-induced afterdepolarization (ADP). D-G) Intrinsic electrophysiological properties determined by whole-cell patch clamp recordings from SC-projecting layer 5 mPFC at P56-70 show that expression of dominant-negative mutant human DISC1 results in an increased input resistance (E), but does not alter resting membrane potential (D), or current through HCN channels as indicated by sag and rebound (G). This characteristic "sag" and "rebound" during and following hyperpolarizing current (-250pA) injection is mediated by hyperpolarization-activated cyclic nucleotide-gated (HCN) channels. F) CaMKII::DISC1 mice lack the quinpirole-induced ADP characteristic of normal D2R+ L5 mPFC pyramidal neurons, quantified by the amount of time required for the membrane potential to return to baseline following the depolarizing current pulse (more precisely, the time for 90% decay), after pharmacological manipulations listed above. **p = 0.0079. ***p < 0.0001.
References


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CHAPTER IV

Functional importance of Tbr1 during later stages of cortical development
Introduction

The cortical circuits have central roles in brain functions (Rakic, 2009). While cortical evolution has endowed the human brain with robust computing abilities, these advances in cortical structure and its circuitry may have increased susceptibility to neuropsychiatric disorders (Johnson et al., 2009). Thus, elucidating the genetic underpinnings of cortical development and function are essential in understanding human disorders such as Autism Spectrum Disorders (ASDs).

Autism Spectrum Disorders (ASDs) are defined by impairments in reciprocal social interaction, often accompanied by abnormalities in language development as well as repetitive behaviors and/or restricted interests. Currently, several hundred genes are implicated in ASD risk, based on de novo copy-number variations or de novo point mutations (De Rubeis et al., 2014; Iossifov et al., 2014; Willsey et al., 2013). Thus, emerging biological insights from genetic and imaging studies have provided evidence that abnormal cortical development underlies at least some forms of ASD (Minshew & Keller, 2010; Willsey et al., 2013).

Cortical development requires the coordinated execution of a series of developmental processes, including regional and areal specification (Rash & Grove, 2006), the specification of neuronal and glia identity (Leone et al., 2008; Molyneaux et al., 2007), cell migration (Kwan et al., 2012), and wiring of neural circuits (De la Rossa et al., 2013). System analysis of co-expression networks of high-confidence ASD genes (hcASD) provided evidence that several of these genes are expressed in immature layer 5 and 6 cortical projection neurons (Willsey et al., 2013). This has provided a strong rationale for studying the function of these ASD genes in cortical development. Among the list of hcASD genes whose expression converges during development of deep cortical layers, several of the genes are already known to regulate mouse cortical development, including Tbr1,
Foxp1, and Bcl11a (Cánovas et al., 2015; De Rubeis et al., 2014; Dias et al., 2016; Dwyer & O’Leary, 2001; Han et al., 2011; Hisaoka et al., 2010; Huang et al., 2014; Iossifov et al., 2014; Sanders et al., 2012; Willsey et al., 2013). Here we have focused our work on defining the function of Tbr1 on the earliest-born cortical projection neurons of layer 6 and the subplate using conditional mutagenesis.

*T-brain-1 (Tbr1)*, a T-box transcription factor (TF), was the first gene shown to regulate development of early-born pallial projection neurons, including Cajal Retzius cells, subplate cells and layer 6 projection neurons (Bedogni et al., 2010; Bulfone et al., 1995; Hevner et al., 2001). In the telencephalon Tbr1 is expressed (beginning ~E10.5) only in postmitotic differentiating cortical projection neurons, and turns on following Tbr2 expression in the subventricular zone (Bulfone et al., 1995; Díaz-Alonso et al., 2015; Hevner et al., 2002; Hevner et al., 2003; Hevner et al., 2001; Vasistha et al., 2015). Tbr1 is expressed in the excitatory neurons of the neocortex (subplate, layer 6, rostral layer 5, layers 2/3), hippocampus, entorhinal cortex, pallial amygdala, piriform cortex, olfactory bulb and Cajal Retzius neurons (Hevner et al., 2003; Hevner et al., 2001). Analysis of Tbr1 constitutive null (*Tbr1*constitutive null) mice showed that it regulates layer 6 identity and the growth of their corticothalamic axons to the thalamus (Bedogni et al., 2010; Bulfone et al., 1998; Hevner et al., 2002). Tbr1 promotes the identity of layer 6 neurons by repressing the Fezf2 and Bcl11b TFs that promote layer 5 fate with subcerebral projections (McKenna et al., 2011). *Tbr1*constitutive null also have defects in cortical arealization (Bedogni et al., 2010; Hevner et al., 2001). Thus, early Tbr1 expression regulates neural fate in which the mutant has defects in molecular specification, neural lamination, and cortical connectivity with the thalamus (Alfano et al., 2014; Hevner et al., 2002; Hevner et al., 2001; McKenna et al., 2011; Mihalas & Hevner, 2017; Remedios et al., 2007). Most *Tbr1*constitutive null mice die at birth. TBR1 molecular functions are beginning to be elucidated. For instance TBR1 protein
interacts with the CASK (calcium/calmodulin dependent serine protein kinase) (Wang et al., 2004). The CASK-TBR1 complex promotes Grin2b transcription by directly binding to its promoter (Chuang, Huang, & Hsueh, 2014). Tbr1 promotes expression of many other genes, including those that encode regulators of cell adhesion and synaptogenesis including Cdh8, Cntn2, Ntng1 and Wnt7b (Huang et al., 2014).

While neocortical development appears normal in Tbr1^{constitutive heterozygous} mice, parts of their pallial amygdala are abnormal. They have abnormal inter- and intra-amygdalar axonal projections, reduced NMDAR expression and the central nucleus of amygdala is particularly susceptible to Tbr1 haploinsufficiency (Chuang, Huang, & Hsueh, 2015; Huang et al., 2014). The reduced NMDAR may be due to Tbr1 driving expression of the NMDAR subunit Grin2b (Chuang et al., 2014). These studies provide evidence that reduced Tbr1 dosage lead to abnormal brain wiring and excitation/inhibition imbalance (NMDAR hypoactivity), two prominent models for ASD etiology (Canitano & Pallagrosi, 2017; Chuang et al., 2014; Huang et al., 2014; Rubenstein, 2010, 2011; Rubenstein & Merzenich, 2003). This is particularly important, as ASD patients with mutations in hcASD genes are typically heterozygous.

Recurrent de novo disruptive mutations in the TBR1 gene have been found in patients with ASDs (De Rubeis et al., 2014; Sanders et al., 2012; Willsey et al., 2013). De novo truncating and missense mutations impairs TBR1’s subcellular localization and transcriptional repression (Deriziotis et al., 2014). Furthermore, TBR1 interacts with FOXP2 (Deriziotis et al., 2014), a transcription factor that is implicated in cortical development (Tsui et al., 2013) and speech/language disorders (Huang & Hsueh, 2017). ChIP-Seq, transcriptomic and computational analyses of TBR1 provided strong evidence that TBR1 regulates expression of multiple ASD risk genes (Notwell et al., 2016).
Given the complex spatio-temporal pattern of Tbr1 expression, we focused on its function in deep neocortical neurons using Cre-lox conditional mutagenesis by creating a Tbr1\textsuperscript{layer6} and a Tbr1\textsuperscript{layer5} mutant. Furthermore, we focused on its function beginning around the day of birth, nearly 10 days after the onset of its expression. We did not eliminate its expression in Cajal Retzius and thereby did not cause a radial migration/lamination defect. Both the Tbr1\textsuperscript{layer5} and Tbr1\textsuperscript{layer6} mutants are viable enabling the analysis of cortical physiology and behavior. We found that late gestation/neonatal Tbr1 function is required to maintain many aspects layer 6 identity.

The Tbr1\textsuperscript{layer6} mutant neurons transformed towards identity of layer 5 neurons, which include their transcriptome, their dendritic pattern and physiology. Previous studies have shown that different prefrontal pyramidal cell classes express varying degrees of hyperpolarizing-activated cyclic nucleotide-gated (HCN) channels (Dembrow \textit{et al.}, 2010; Gee \textit{et al.}, 2012; Seong and Carter, 2012). Quantification of the current through HCN channels, h-current (I\textsubscript{h}), can be used to distinguish dopamine D2 receptor (D2R) expressing neuron within the pre-frontal cortex (Clarkson \textit{et al.}, 2017). While layer 5 pyramidal neurons are of heterogeneous populations with various degrees of h-current, layer 6 pyramidal neurons classically lack h-current. The Tbr1\textsuperscript{layer6} mutant neurons had reduced excitatory and inhibitory synaptic density. Importantly, the physiological and synaptic phenotypes were also seen in Tbr1\textsuperscript{layer6} heterozygous. We found that this reduction in excitatory and inhibitory synapse density functionally translates into reduced spontaneous EPSCs and IPSCs in mouse brain slices.

RNA-seq of FACS purified layer 5 neurons from the Tbr1\textsuperscript{layer5} mutants did not show Tbr1 as a differentially regulated gene (data not shown). This is likely due to the heterogeneous nature of L5 cortical neurons (Clarkson \textit{et al.}, 2017; Dembrow \textit{et al.}, 2010). A more precise analysis of differential gene regulation in Tbr1\textsuperscript{layer5} mutant neurons could be
obtained by performing single-cell RNA-seq. Interestingly, electrophysiological properties of the \( Tbr1^{\text{layer5}} \) mutant neurons suggest a homogenization of the L5 population towards prominent h-current suggestive of D2Rpositive neuron identity (Clarkson et al., 2017). The \( Tbr1^{\text{layer5}} \) mutant neurons also had reduced excitatory and inhibitory synaptic density. We found that this reduction in excitatory and inhibitory synapse density functionally translates into reduced spontaneous EPSCs and IPSCs in mouse brain slices.

In the present study, we have demonstrated the neurobiology behind \( Tbr1 \) loss of function in layer 5 and layer 6 glutamatergic neurons as a part of a concerted effort to understand the neurobiology of Autism Spectrum Disorder (ASD) mutations. ASD is characterized by severe and sustained impairment in social communication and interactions. Humans with ASD also suffer from a range of additional behavior problems, including anxiety, compulsive behaviors, and aggression (Gadow et al., 2004; McClintock et al., 2003). We have also discovered that loss of \( Tbr1 \) function in layer 5 leads to a pronounced decrease in social behaviors consistent with ASD associated behaviors. Interestingly, loss of function of \( Tbr1 \) in layer 6 does not reduce social interaction but instead leads to increased anxiety and aggressive behavior. Aggressive behaviors are commonly observed in individuals diagnosed with autism spectrum disorder (ASD) and may be a phenotypic indicator of different subtypes within ASD. Interestingly, loss of function mutations of \( Tbr1 \) in layer 5 versus layer 6 have differential impacts on behavioral outcomes.
Materials and Methods

Animals

The Tbr1\textsuperscript{flox} allele was generated by inGenious Targeting Laboratory (Ronkonkoma, NY). LoxP sites were inserted into introns 1 and 3, flanking Tbr1 exons 2 and 3 (Fig. 1A). To enable selection of homologous recombinants, the LoxP site in intron 3 was embedded in a neo cassette that was flanked by Flp sites. The neo cassette was removed by mating to a Flp-expressing mouse (Rodriguez et al., 2000) to generate the Tbr1\textsuperscript{flox} allele. Cre excision removes exons 2 and 3, including the T-box DNA binding region, similar to the constitutive null allele (Bulfone et al., 1998). Ntsr1-cre mice were used to delete Tbr1 in layer 6 projection neurons. tdTomato\textsuperscript{fl/+} (Ai14) mice were crossed with Tbr1\textsuperscript{f/+} mice and used as an endogenous reporter. Tbr1 layer 6 knockout mice (Tbr1\textsuperscript{layer6} mutant) were generated by crossing Tbr1\textsuperscript{f/+}::tdTomato\textsuperscript{fl/+} mice with Tbr1\textsuperscript{f/+}::Ntsr1-cre\textsuperscript{+}. Rbp4-cre mice were used to delete Tbr1 in layer 5 projection neurons. tdTomato\textsuperscript{fl/+} (Ai14) mice were crossed with Tbr1\textsuperscript{f/+} mice and used as an endogenous reporter. Tbr1 layer 5 knockout mice (Tbr1\textsuperscript{layer5} mutant) were generated by crossing Tbr1\textsuperscript{f/+}::tdTomato\textsuperscript{fl/+} mice with Tbr1\textsuperscript{f/+}::Rbp4-cre\textsuperscript{+}. All strains were maintained on a C57Bl/6 background. Animals were housed in a vivarium with a 12hr light, 12hr dark cycle. Postnatally, experimental animals were kept with their littermates.

For timed pregnancies, noon on the day of the vaginal plug was counted as embryonic day 0.5. Animal care and procedures were performed according to the University of California San Francisco Laboratory Animal Research Center (LARC) guidelines.

Genomic DNA extraction and genotyping

Tissue samples were digested in a solution containing 1 mg/mL of proteinase K, 50 mM Tris-HCl pH 8.0, 100 mM EDTA, 100 mM NaCl and 1% SDS. Genomic DNA was extracted using a standard ethanol precipitation protocol. Genotyping was performed with PCR-based
assays using purified genomic DNA, and various primer-pair combinations flanking the deleted region and detecting Cre and tdTomato alleles.

**RNA extraction and cDNA synthesis**

Total RNA was extracted from the cortices of wildtype and Tbr1 constitutive null mice at E15.5 and P0 using RNeasy Plus® Mini Kit (QIAGEN) following the manufacturer’s protocol. First strand cDNA was synthesized using Superscript reverse transcriptase II following manufacturer’s protocol (Thermofisher).

**Quantitative real time PCR (qRT-PCR)**

Quantitative RT-PCR was performed to measure RNA levels using SYBR Green (Bio-Rad) and 7900HT Fast Real-Time PCR System. Gene-specific primers for Tbr1 exons 1, 2 and 4, Bcl11a, Grin2b and Hcn1 as well as ef1α housekeeping genes (HKG) were designed using the Primer 3 program (Rozen & Skaletsky, 1999). The expression levels of the genes in both wildtype and Tbr1 mutant mice were normalized to the expression levels of ef1α. Subsequently, the gene expression levels in Tbr1 mutant mice were measured relative to the wildtype littermates (Darbandi & Franck, 2009; Pfaffl, 2001).

**Western blot**

Cortices of 2 Tbr1 constitutive null and 2 wildtype brains were dissected at E15.5 and P0 in ice-cold PBS. Tissues were homogenized in 300 µL ice-cold RIPA lysis buffer. Following an incubation at 4°C for 2 hrs with agitation, the samples were centrifuged at 13,500 rpm for 20 min at 4°C. 20-30 µg total protein was combined with Laemmli buffer supplemented with 1:20 β-mercaptoethanol and was heat to 95°C for 5 min. The protein
lysate was electrophoresed using Mini-PROTEIN® TGX 4-20% precasted gels (Bio-Rad) and ran for 1-2 hrs at 100V. The fractionated proteins were transferred to a nitrocellulose membrane (GE Amersham Protran). The membrane was blocked with 7.5% nonfat dried milk, washed 3X with 1X PBS with 0.1% Tween-20, and then was incubated for 12 hrs with the primary antibody at 4°C. The following day, the membrane was washed 3X with 1X PBS with 0.1% Tween-20, incubated with the Goat Anti-Rabbit-HRP secondary for 1 hr. Signals were detected using a DAB system (Vector Laboratories) following manufacturer’s protocol.

RNA-seq on FAC-Sorted Cells

Layer specific transcriptome profiling was conducted by using RNA-seq on FAC-Sorted cells from somatosensory cortex of Tbr1\(^{\text{wildtype}}\), Tbr1\(^{\text{layer5}}\), and Tbr1\(^{\text{layer6}}\) mutants. The somatosensory cortex was dissected in HBSS from P5 mice (Thermofisher). Cortices were dissociated using a Papain Dissociation System (Worthington Biochemical Corporation) following manufacturer’s protocol. tdTomato\(^+\) cells were sorted using BD FACS Aria II Cell Sorter at Center for Advanced Technology (UCSF). Approximately 25,000 cells were collected from each sample and immediately proceeded with RNA extraction using RNeasy\(^\circledR\) Plus Micro Kit (QIAGEN) following manufacturer’s protocol. RNA quality was assessed using Agilent RNA 6000 Nano Kit (Agilent Technologies) and ran on Bioanalyzer 2100 (Agilent Technologies). RNA-seq libraries were generated by TruSeq\(^\circledR\) Stranded Total RNA Library Prep Kit with Ribo-Zero Gold Set A (Illumina). The quality of the libraries was assessed using Bioanalyzer 2100 (Agilent Technologies) and Agilent High Sensitivity DNA Kit (Agilent Technologies) following manufacturer’s protocol. Libraries were validated using qPCR and sequenced on Hiseq 4000 at Center for Advanced Technology (UCSF).
Bioinformatics analysis of FAC-Sorted layer 6 RNA-Seq

Collectively, we analyzed 8 RNA-Seq libraries, which comprised of 4 $Tbr1^{\text{wildtype}}$ and 4 $Tbr1^{\text{layer6}}$ mutant RNA-Seq libraries. Sequencing was conducted on HiSeq 4000 using Paired-End 100 (PE100) with the Library fragment size of approximately 300 bp.

RNA-Seq alignment, and quality control

The RNA-Seq reads were aligned to the mm9 mouse genome reference using STAR in gene annotation mode. Picard was utilized to generate alignment quality control (QC) metrics for every RNA-Seq samples. Principle component analysis (PCA) of the quality control matrices was employed to determine the presence of RNA-Seq sample outliers (The outliner is defined as a sample whose QC metrics are at least three standard deviations away from the mean in any of the first three principal components). The analysis did not indicate any outliers in these datasets.

Gene expression estimation and normalization

Gene expression was quantified with HTSeq in intersection-strict mode. Out of 24,015, 16,805 genes have at least one read in more than 50% of the samples. These 16,805 genes were normalized for gene length, GC content, and sample library size using CQN R-package. Gene length is obtained directly from the gene annotation file (.GTF) of mouse mm9 genome build reference. Bedtools is used to compute the gene GC content.

After normalization, the genes whose expression value doesn’t change across all samples are removed. PCA is applied to identify any sample outliners with those filtered and normalized gene expression. The expression values were scaled and centered before PCA. PCA over gene expression shows that there are not outliners in this dataset.
Differential gene expression analysis (DEX analysis):

To identify differentially expressed genes (DEX genes), we identified all possible confounding variables including ribosomal bases in the mapped reads, percentage of bases in intronic region, RIN, Sex and RNA concentration to produce a reliable conclusion. Thousands of negative binomial regression models are built to model expressions of each gene. The best model is formed using Bayesian information criterion (BIC) and forward stepwise algorithm. The DEX analysis was performed with edgeR (McCarthy, 2012; Robinson, 2010). Genes that pass the 0.05 significance threshold are considered as significantly differentially expressed genes.

Gene Co-expression network analysis with gene expression residual:

To avoid effect of confounding variables in gene co-expression network analysis, we mathematically removed the effects of variables, Sex and percentage of UTR region bases by computing the estimated gene expression with linear model coefficients and over-dispersion parameters output by edgeR. We use the estimated gene expression to construct a signed gene co-expression network. Hierarchical clustering with the average linkage method is employed to create modules of genes. Genes in one module have similar expression patterns in the data. The expression of genes belonging to a module is summarized using the eigengene expression of the module. We evaluated the degree of association between ‘Tbr1\textsuperscript{layer5} mutant vs. Control’ as well as ‘Tbr1\textsuperscript{layer6} mutant vs. Control’ status and module eigengene expression using Pearson correlation. Gene module which are statistically significant (adjusted p-value \(\leq 0.05\)) correlated (\(\geq 0.9\) or \(\leq -0.9\)) with status are reported. We conducted the GO pathway analysis using ToppFun.
online tool to understand the biological meaning of those gene modules.

**TBR1 Chromatin immunoprecipitation (ChIP)**

Transcription factor ChIP was performed as previously published with a few modifications (McKenna et al., 2011; Sandberg et al., 2016). P2 somatosensory cortices were dissected and dissociated by pipetting in cold PBS. Dissociated cells were fixed in 1% formaldehyde for 10 min at RT and neutralized with 1 mL 2.5M glycine. Fixed chromatin was lysed and sheared into 200 - 1,000 bp fragments using a Covaris S2 (13 cycles of duty cycle = 5%, intensity = 3 and cycles per burst = 200). Immunoprecipitation (IP) reactions of two biological replicates at P2 were performed using 5 μg TBR1 polyclonal antibody (Santa Cruz Biotech, SC48816 X (M-200)). 20X molar excess TBR1 blocking peptide was used as negative control. Protein/antibody complexes were collected using Dynabeads (20 mL protein A + 20 mL protein G) and processed as previously described (Vokes et al., 2007). ChIP-seq libraries were generated using Ovation Ultralow System V2 (NuGEN) following manufacturer’s protocol. Generated libraries were size selected (180–350 bp) and sequenced at the Center for Advanced Technology at UCSF (http://cat.ucsf.edu/) and the Genomics Core Facility (http://humangenetics.ucsf.edu/genomics-services/).

**ChIP-Seq Computational Analysis**

Clustering, base calling, and quality metrics were performed using standard Illumina software. Sequenced libraries were analyzed for overall quality and were filtered, and reads were mapped to the mouse genome (mm9) as previously described (Sandberg et al., 2016).
Primary Cell Culture and Luciferase assay

**Plasmids:** To generate luciferase constructs candidate regulatory elements of mouse *Tbr1* (hs416, chr2: 61494203-61494886, 683bp), *Foxp2* (chr6: 15097241-15098146, 905 bp), *Grin2b* (chr6: 135813640-135814770, 1,130 bp), *Bcl11a* (chr11: 24270818-24271924, 1,383 bp), *Drd1* (chr13: 54074453-54075131, 678 bp), *Hcn1* (chr13: 118669041-118670541, 1,500 bp), *Fezf2* (hs434, chr14: 13170235-13171693, 1,458bp), *Foxp1* (chr6: 99325484-99327361, 1,877 bp), and *Dlx5/6i* enhancer (chr6: 6819420-6819819, 400 bp) were amplified by PCR, and cloned into the pGL4.23 vector (Promega). The vectors were transformed with DH5α E. coli cells at 42°C.

**Luciferase assay:** Primary cortical neurons were harvested from P0 wildtype cortex and transfected using Lipofectamine 2000 (Invitrogen) and one of the regulatory element luciferase vectors that were generated as described above. To test whether TBR1 modified the regulatory elements activity, *pCAG-Tbr1-IRES-eGFP* was co-transfected together with one of the aforementioned regulatory element vectors. A renilla luciferase plasmid (pRL, Promega) was co-transfected to control for transfection efficiency. The luciferase assay was performed 48hrs after transfection using the dual-luciferase kit (Promega) according to manufacturer’s instructions. Reporter activity was measured using Veritas™ Microplate Luminometer (Turner BioSystems, Model# 9100-001).

**Histology**

At the time of experiment, for P0 and P3 experiments, animals were anesthetized on ice while postnatal (P21 and P56) animals were anesthetized with intraperitoneal avertin (0.015 ml/g of a 2.5% solution) injection. Animals were perfused transcardially with cold
PBS and then with 4% PFA in PBS, followed by brain isolation, 1-2 hr post-fixation, cryoprotected in 30% sucrose in PBS, and cut frozen (coronally or sagittally) on a sliding microtome at 40µm for immunohistochemistry or in situ hybridization. All primary and secondary antibodies were diluted in PBS containing 10% Normal Serum, 0.25% Triton X-100 and 2% BSA. The following primary antibodies were used: Chicken anti-GFP (1:2000, Aves), mouse anti-Vglut1 (1:200, Synaptic Systems), rabbit anti-Vgat (1:500, Synaptic Systems), rabbit anti-PSD95 (1:200, Cell Signaling), mouse anti-gephyrin (1:200, Synaptic Systems). The secondary antibodies for immunofluorescence were Alexa Fluor-conjugated and purchased from Thermofisher. For synapse immunohistochemistry, sections were pretreated with pepsin to enhance the staining as described before (Corteen, Cole, Sarna, Sieghart, & Swinny, 2011). Immunofluorescence specimens were counterstained with 1% DAPI to assist the delineation of cortical layers. For in situ hybridization a rostro-caudal coronal series of at least ten sections were examined. Anti-sense riboprobes for Tbr1, Foxp2, Wnt7b, Tle4, Nr4a2, Bcl11b, Fezf2, Foxp1, Sst, Ntng1, Cntn2, and Drd1 were prepared as previously described (Cobos et al., 2005; Long, Garel, Depew, Tobet, & Rubenstein, 2003). ISH was performed using digoxigenin-labeled riboprobes as previously described (Stanco et al., 2014).

**Image Acquisition and Analysis**

Fluorescent and brightfield images were taken using a Coolsnap camera (Photometrics) mounted on a Nikon Eclipse 80i microscope using NIS Elements acquisition software (Nikon). Confocal images were taken using Zeiss LSM 880 Confocal Microscope with a 63X objective at 1,024×1,024 pixels resolution using ZEN 2.0 software at Molecular Imaging Center (UC Berkeley). Brightness and contrast were adjusted and images merged.
using Photoshop or ImageJ software. ImageJ software was used for image processing. For synapse counting (presynaptic and postsynaptic boutons), confocal image stacks (0.4µm step size) were processed with ImageJ software. In brief, background subtraction and smooth filter were applied to each stack. Using a threshold function, each stack was converted into a ‘masks’ image. Furthermore, the channels were co-localized with the Image Calculator plugging. Lastly, the number of co-localizations were counted and the length of each dendrite was measured in each of the focal plane. Staining for control and mutant were done in parallel as well as the image capturing.

Cell counting

For assessing cell density in the neocortex on fluorescence sections 10X images at were taken of the somatosensory cortex, encompassing all neocortical layers, from both hemispheres for each replicate. Images were opened with ImageJ to delimit and measure the region of interest (ROI).

Electrophysiology

Coronal brain slices (250 µm) including medial prefrontal cortex were made from mice age p21-p28 and p56-p80. Slicing solution was chilled to 4°C and contained (in mM): 234 sucrose, 26 NaHCO₃, 11 glucose, 10 MgSO₄, 2.5 KCl, 1.25 NaH₂PO₄, 0.5 CaCl₂, bubbled with 5% CO₂/ 95% O₂. Slices were incubated in artificial cerebrospinal fluid (aCSF) at 32°C for 30 minutes and then at room temperature until recording. aCSF contained (in mM): 123 NaCl, 26 NaHCO₃, 11 glucose, 3 KCl, 2 CaCl₂, 1.25 NaH₂PO₄, 1 MgCl₂, also bubbled with 5% CO₂/ 95% O₂.

Neurons were visualized using differential interference contrast or DODT contrast microscopy on an upright microscope (Olympus). Ntsr1cre positive neurons were identified
by fluorescent visualization of cre-dependent tdTomato. We obtained somatic whole-cell patch clamp recordings using a Multiclamp 700B (Molecular Devices) amplifier and acquired with pClamp. Patch pipettes (2-5 MΩ tip resistance) were filled with the following (in mM): 130 K gluconate, 10 KCl, 10 HEPES, 10 EGTA, 2 MgCl₂, 2 MgATP, 0.3 Na₃GTP. All recordings were made at 32-34°C. Series resistance was compensated in all current clamp experiments and monitored throughout recordings. Recordings were discarded if Rs changed by >25%. For spontaneous EPSC and IPSC recordings cells were held in voltage clamp at -70 mV and +10 mV, respectively. In both cases patch pipettes were filled with the following (in mM): 135 Cesium Methanesulfonate, 8 NaCl, 10 HEPES, 0.3 EGTA, 5 QX314, 4 MgATP, 0.3 Na₃GTP.

**Viral injection for expression of fluorescent reporters**

Viral injections were performed using standard mouse stereotactical methods. Mice were anesthetized for the duration of the surgery using isoflurane gas. After cleaning, an incision was made in the scalp, the skull was leveled, and small burr holes were drilled over the brain region of interest using a dental drill. Virus was injected through the burr holes using a microinjector at a speed of 150 nL/minute and the scalp was closed using tissue adhesive (Vetbond).

For expression anterograde fluorescent reporters, we injected a Cre-dependent ChR2-eYFP virus (AAV5-Ef1-DIO-Chr2-eYFP, 0.5 µL). In $Tbr^{layer5}$ mutant and $Tbr^{control}$ mice, these viral injections targeted mPFC and medial dorsal thalamus (mdT) whereas in $Tbr^{layer6}$ mutant and $Tbr^{control}$ mice we targeted S1 and ventromedial (VM) thalamus of each mouse line (p21-p28) and waited 3-4 weeks for trafficking of ChR2 to the axon terminals in mPFC. For retrograde labelling, Alexafluor-tagged cholera toxin subunit B 488 (CTb488; ThermoFisher) was first dissolved in PBS to make a 1.0mg/mL solution. 350nL of
CTb488 were then injected into the MD thalamus at a rate of 100nL/min and waited 3-5 days before histology experiments. md thalamus injection coordinates were A/P = -1.7, M/L = +/- 0.3, D/V = -3.45. mPFC injection coordinates were A/P = +1.7, M/L = +/- 0.3, D/V = -2.75. VM thalamus injection coordinates were A/P = -1.7, M/L = +/- 0.7, D/V = -4.1. S1 injection coordinates were A/P = -0.75, M/L = +/- 2.5, D/V = -1.75

**Behavior Assays**

Mice are transferred to experimental room and allowed to habituate for at least 45 minutes each day. All behavior assays were performed on mice age P56 to P80. We were blind to genotype during scoring of videos.

**Open-field test:** An individual mouse was placed near the wall-side of 50 x 50 cm open-field arena, and the movement of the mouse was recorded by a video camera for 10 min. The recorded video file was later analyzed with Any-Maze software (San Diego Instruments). Time in the center of the field (a 25 x 25 cm square) was measured. The open field arena was cleaned with 70% ethanol and wiped with paper towels between each trial.

**Elevated plus maze test:** An individual mouse was placed at the junction of the open and closed arms, facing the arm opposite to the experimenter, of an apparatus with two open arms without walls (30 x 5 x 0.5 cm) across from each other and perpendicular to two closed arms with walls (30 x 5 x 15 cm) with a center platform (5 x 5 cm), and at a height of 40 cm above the floor. The movement of the mouse was recorded by a video camera for 10 min. The recorded video file was later analyzed and time in the open arms of the apparatus
was measured. The arms of the elevated plus maze apparatus was cleaned with 70% ethanol and wiped with paper towels between each trial.

**Rotarod test:** The assay consisted of four trials per day over the course of 2 days with the rotarod set to accelerate from 4rpm to 45rpm over 5 minutes. The trial started once five mice were placed on the rotarod rotating at 4rpm in separate partitioned compartments. Each trial ended when a mouse fell off, made three complete revolutions while hanging on, or reached 300 s. Digital videos of the mice on the rotarod were recorded from behind.

**Social interaction and novel object task:** An individual mouse was allowed to habituate for 5 minutes in their home cage prior to starting the trial. A juvenile (3-4 weeks old) mouse of the same strain and sex was introduced to the home cage. After 5 minutes, the juvenile was removed from the home cage. After a 5 minute break a novel object (typically a plastic test tube cap) was introduced into the home cage for five minutes. We scored videos offline, blind to genotype. We measured the number of seconds the mouse spent with its nose in direct contact with the novel object or engaged in social interaction with the juvenile (defined as sniffing, close following, or allo-grooming) in the 300 seconds following the time the juvenile or object was introduced into the cage. In addition, we noted any aggressive-appearing behaviors toward the juvenile, freezing, and grooming behaviors.

**Statistical Analysis**

All statistical analyses were done on GraphPad Prism 7.0 software. All data are shown as mean ± SEM. Statistical significance was accepted at the level p < 0.05. We used student’s t-test to compare pairs of groups if data were normally distributed (verified using
Lillie test). If more than two groups were compared, we used one-way ANOVA with post-hoc tests between groups corrected for multiple comparisons (Holm-Sidak or Tukey). The specific post-hoc test as well as exact F and corrected p values can be found in the text and corresponding figure legends.
Results
Characterization of the Tbr1 conditional mutant allele.

To investigate the function(s) of Tbr1 in specific subtypes of cortical neurons at later stages of development, we generated a Tbr1 conditional mutant (Tbr1\textsuperscript{flox}) allele. We constructed a targeting vector that inserts LoxP sites into introns 1 and 3 (Fig. 1A). Therefore, upon Cre-recombination, exons 2 and 3 (including the T-box DNA binding region) will be deleted; similar to the constitutive null allele previously published (Bulfone et al., 1998).

We validated recombination of Tbr1\textsuperscript{flox/flox} alleles with the \(\beta\)-actin-cre allele, to examine whether the conditional allele would result in Tbr1 loss of function upon Cre recombination. We measured Tbr1 RNA levels from the cortex (E15.5 and P0) via RT-qPCR using primers to exons 1 and 2, and to exons 1 and 4. Only the exon 1-2 reaction showed no signal in the Tbr1 mutant consistent with the location of the Cre-mediated deletion (Fig. 1C). We next investigated Tbr1 transcripts using in situ hybridization (ISH) at P0, using either a Tbr1 full-length (FL) probe, or a probe that only hybridizes to exons 2 and 3 (E2-3; the part deleted by Cre recombination). We did not observe any signal with the Tbr1 E2-3 probe; whereas we observed strong signal from the FL Tbr1 probe (Fig. 1D), suggesting that a stable truncated RNA is still produced in the conditional mutant.

We performed a western blot (WB) on cortical proteins from Tbr1 null and wildtype littermates at E15.5 and P0 to determine whether the Tbr1 truncated RNA generates a stable protein (Fig. 1E). The TBR1 antigen is encoded by exon 1 and maps to the N-terminus of TBR1 (red; Fig. 1A, 1B). The TBR1 protein is \(~75\) kDa in the wildtype E15.5 and P0 cortex; however, no signal was detected in the Tbr1 null cortex at E15.5 and P0 (Fig. 1E). Thus, it appears the Tbr1\textsuperscript{flox} allele generates a conditional Tbr1 allele that does not generate a stable TBR1 protein, and thus is likely to function as a null allele.
**Tbr1 maintains layer 6 identity in postnatal cortex.**

We next studied the effect of deleting Tbr1 in layer 6 and subplate approximately 5-6 days after Tbr1 expression begins in these neurons using *Ntsr1*-cre (Madisen et al., 2010). We refer to these as *Tbr1*\(^{layer6}\) mutants. *Ntsr1*-cre expression, detected using the Ai14 tdTomato Cre-dependent reporter, begins in layer 6 and subplate neurons around E16.5-E17.5 (data not shown). To identify dysregulated RNAs in layer 6 and subplate of *Tbr1*\(^{layer6}\) mutants we performed RNA sequencing (RNA-Seq) on FAC-sorted cells from the *Tbr1*\(^{wildtype}\) and *Tbr1*\(^{layer6}\) mutant somatosensory cortex at P5. We assessed a total of 8 samples, including 4 *Tbr1*\(^{layer6}\) mutants and 4 *Tbr1*\(^{wildtype}\) controls. After alignment, quality control, and normalization, we identified differentially expressed (DEX) genes in the *Tbr1*\(^{layer6}\) mutant versus the *Tbr1*\(^{wildtype}\) control with EdgeR (McCarthy, 2012; Robinson, 2010). We detected 178 genes DEX after correction for multiple comparisons (false discovery rate ≤ 0.05), including *Tbr1* (fold change [FC] = -1.93, p = 3.83 \times 10^{-7}). These 178 genes are enriched for gene ontology (GO) terms related to nervous system development, neurogenesis, and neuron development (Fig. 2), consistent with the known functions of *Tbr1* (Bedogni et al., 2010; Bulfone et al., 1995; Bulfone et al., 1998; Hevner et al., 2001).

The observed transcriptome changes provide evidence that *Tbr1* is required to maintain the identity of layer 6 neurons. Several key regulators of layer 6 identity are downregulated in the *Tbr1*\(^{layer6}\) mutants, including *Tle4* (FC = -2.11, p = 2.99 \times 10^{-8}) and *Wnt7b* (FC = -2.08, p = 1.83 \times 10^{-6}). Furthermore, we observed ectopic expression of regulators of layer 5 identity and properties, including *Foxp1* (FC = 1.41, p = 0.029), *Fezf2* (FC = 1.68, p = 0.000176), *Grin3a* (FC = 2.36, p = 4.65 \times 10^{-7}) and *Epha7* (FC = 1.95, p = 1.21 \times 10^{-6}; Fig. 2). We conducted ISH on a subset of these genes to verify the changes observed in the RNA-seq data. Fig. 4 shows the results in the somatosensory cortex at P3. We confirm that *Tbr1* and *Wnt7b* expression is reduced in layer 6 of *Tbr1*\(^{layer6}\) mutants (Fig.
4H-H', J-J'). This decrease appears to be accompanied by an increase in *Tbr1* and *Wnt7b* in the superficial layers of the somatosensory cortex (Fig. 4H-H', J-J'). We also observe that *Nr4a2* expression is reduced in the subplate (Fig. 4I-I'). We also confirmed the previously mentioned ectopic expression of layer 5 regulators in layer 6; for example, *Bcl11b* (Fig. 4K-K'), *Fezf2* (Fig. 4L-L') and *Foxp1* (Fig. 4M-M').

Overall, the RNA expression data (RNA-Seq and ISH; Figs 2 and 4) provide evidence that *Tbr1* is required not only to induce layer 6 identity (Hevner, Miyashita-Lin, & Rubenstein, 2002; Hevner et al., 2001), but is also essential approximately 5-6 days after *Tbr1* expression is established to maintain layer 6 identity.

*Tbr1* directly regulates the transcription of genes that control layer 6 identity.

To determine whether the changes in genes expression in *Tbr1* mutants is due to direct regulation by TBR1, we performed chromatin immunoprecipitation followed by high-throughput sequencing (ChIP-Seq) on P2 cortex and assessed whether TBR1 binds to the promoter and/or other candidate regulatory elements (REs) of key DEX genes. We also performed ChIP-Seq against H3K27Ac, a histone variant associated with active enhancers (Creyghton et al., 2010).

In P2 cortex, TBR1 binds to 27,228 regions genome-wide. To stratify these binding sites, we defined promoters as 1 kb upstream of the 5' UTR, all other regions as putative intragenic or distal regulatory regions. Approximately 40% of peaks overlap promoters and approximately 60% overlap intragenic or distal regulatory regions. To narrow to putative functional binding sites, we intersected these peaks with H3K27Ac peaks generated from the same tissue. To gain further evidence that these peaks represent candidate active regulatory elements influenced by TBR1, we assessed the number of genes with putative
TBR1 regulatory loci that are dysregulated in the $Tbr^\text{layer6}$ mutants. TBR1 binds to the 89% of the promoter regions and 77% of the candidate REs near the genes dysregulated in the $Tbr^\text{layer6}$ mutants (Fig. 5). Given the putative functional connection between $Tbr$ dysregulation and these regulatory elements, we next directly tested the activity of several REs, and their relationship to TBR1.

Thus, we cloned 9 REs into pGL4.23 luciferase expression vector upstream of a minimal promoter. We then transfected these into primary cultures made from P0 cortex, and assayed luciferase activity 3 days later. In parallel experiments, we co-transfected the RE vectors with a $Tbr$ expression vector. We tested 3 classes of candidate REs that were candidates for regulating genes that were (1) downregulated ($Tbr$, $Foxp2$, $Grin2b$, $Bcl11a$ and $Drd1$), (2) upregulated ($Hcn1$, $Fezf2$ and $Foxp1$) and (3) unchanged ($DlxI5/6i$) in the $Tbr^\text{layer6}$ mutants (Fig. 5). Luciferase activity was driven by all of the REs (Fig. 3), except the negative control $DlxI5/6i$ interneuron specific enhancer (Zerucha et al., 2000). On the other hand, $Tbr$ co-transfection only activated the $Tbr$, $Foxp2$, $Grin2b$, $Bcl11a$ and $Drd1$ candidate REs, consistent with down-regulation of these cognate genes in the $Tbr^\text{layer6}$ mutants (Fig. 2). In addition, $Tbr$ co-transfection reduced luciferase expression with the $Hcn1$, $Fezf2$ and $Foxp1$ REs, which corresponded to genes whose expression increased in the $Tbr^\text{layer6}$ mutants (Fig. 2).

Thus, using TBR1 ChIP-Seq and a RE functional assay, we have identified RE and provided evidence that they function either as activators or repressors of expression in the presence of TBR1. This data solidifies our evidence that TBR1 directly controls the molecular properties of layer 6 pyramidal neurons, and that TBR1 functions as an activator or a repressor depending on the nature of the RE.
**Tbr1** specifies a program that patterns apical dendritic lamination of layer 6 neurons.

A key feature of a neuron’s identity is its dendritic pattern (Lefebvre, Sanes, & Kay, 2015; Russ & Kaltschmidt, 2014). Normally, the apical dendrites of layer 6 pyramidal neurons grow into layer 4 where they elaborate their branches (Ledergerber & Larkum, 2010). We used tdTomato expression that was driven by *Ntsr1-cre* in layer 6 and the subplate, to compare the dendritic pattern of wild type and *Tbr1<sup>layer6</sup> mutants (Fig. 6). In the wildtype control, the apical dendrites of the layer 6 glutamatergic neurons extend to layer 4 at P3, P21 and P60 (Fig. 6A-C). On the other hand, in the *Tbr1<sup>layer6</sup>* mutants the apical dendrites extend to layer 1 at P3, P21 and P60 (white arrowheads, Fig. 6A’-C’). This change in the morphology of the dendrite is consistent with the hypothesis that the layer 6 neurons have changed fate to a layer 5 identity.

**Tbr1** is required after E17.5 for corticothalamic projections into the anteromedial thalamus.

Layer 6 and subplate projection neurons have a characteristic axon projection pattern that involves axons that extend through the basal ganglia to the thalamus, where they form a stereotypic topographic map between cortical areas and specific thalamic nuclei (Crandall, Cruikshank, & Connors, 2015). *Tbr1* constitutive null mutant axons fail to grow to the thalamus (Hevner et al., 2002; Hevner et al., 2001). Here, we investigated corticothalamic projections in *Tbr1<sup>layer6</sup>* mutants in two ways. First, we injected a Cre-dependent anterograde label (DIO-ChR2-EYFP) into the somatosensory cortex of wild type and *Tbr1<sup>layer6</sup>* mutants at P21. Histological analyses were performed at P60. Importantly, *Ntsr1-Cre* recombination of *Tbr1* does not extend into many cortical regions, including the olfactory bulb, medial prefrontal neocortex, dorsomedial neocortex (cingulate and
retrosplenial), hippocampus and parahippocampus, piriform cortex, and pallial amygdala; therefore we targeted the somatosensory cortex where there is significant overlapping expression of Ntsr1-cre and endogenous Tbr1 (data not shown). Despite the evidence that Tbr1\textsuperscript{layer6} mutant layer 6 pyramidal neurons have molecular and dendritic properties of layer 5 pyramidal neurons (Figs. 2, 3, 4, 6), the mutant layer 6 neurons, like in the wildtype, have corticothalamic projections that enter the thalamus (Fig. S1). However, the Tbr1\textsuperscript{layer6} mutants have reduced thalamic projections, which is most strongly seen in the anterior and anteromedial thalamus (seen in the rostral coronal sections). We confirmed this phenotype using a second approach using tdTomato expression that was driven in layer 6 neurons using Ntsr1-cre. This method also highlighted a reduction of mutant layer 6 cortical projections in the striatum.

To evaluate whether the phenotype was due to a failure to maintain the projections, or a failure to establish them, we studied neonatal Tbr1\textsuperscript{layer6} mutants at P3 (Fig. S2). The P3 and P21 phenotypes were very similar. This result supports a conclusion that Tbr1 expression in layer 6 and subplate neurons is required after ~E17.5 for the growth of corticothalamic projections in the anterior and anteromedial thalamus. Thus, while Tbr1 is required prior to E17.5 for corticothalamic projections to emerge from the subpallium and enter the diencephalon (Hevner et al., 2002; Hevner et al., 2001), these processes take place in the Tbr1\textsuperscript{layer6} mutants. However, the latter mutants show that after ~E17.5 Tbr1 is required for the maturation of corticothalamic connectivity preferentially in the anterior and anteromedial thalamus.

Excitatory synapse numbers are reduced in Tbr1\textsuperscript{layer6} and Tbr1\textsuperscript{layer5} mutants.

We next investigated excitatory synapses numbers of layer 6 neurons in the Tbr1\textsuperscript{layer6} mutants as well as layer 5 neurons in the Tbr1\textsuperscript{layer5} mutants. We used
immunofluorescence to label excitatory presynaptic terminals (VGlut1+) that are apposed to dendritic postsynaptic zones (PSD95+) in control, and Tbr1 heterozygous and homozygous mutants on the apical spines at P21 (layer 6, Fig. 7I; layer 5, Fig. 8I) and P56 (layer 6, Supplementary Fig 3I; layer 5, Data not shown). We continued to examine the loss of Tbr1 within layer 6, driven by Ntsr1-Cre recombination, in the somatosensory cortex. In the Tbr1layer5 mutants we targeted the medial prefrontal neocortex, a brain region strongly implicated in ASD dysfunction (Bachevalier and Mishkin, 1986; Morgan et al., 1993), because Rbp4-Cre recombination of Tbr1 occurred throughout many cortical regions including the mPFC (data not shown).

Tbr1 postnatally regulate the number of excitatory synapses at P21 and P56 in coronal sections from the somatosensory cortex (layer 6, Fig. 7A) and the medial prefrontal cortex (layer 5, Fig. 8A). We analyzed the number of excitatory terminals (VGlut1+ presynaptic structures) that are apposed to dendritic postsynaptic zones (PSD95+) of Tbr1wildtype, Tbr1 heterozygous mutants and Tbr1homozygous mutants at P21 (layer 6, Fig 7I; layer 5, Fig. 8I) and P56 (layer 6, Supplementary Fig 3I; layer 5, data not shown).

As depicted in Fig. 7I, brain sections from somatosensory cortex (Fig. 7A) of Tbr1wildtype (Ntsr1-cre::tdTomato/+ , Fig. 7D-D’), Tbr1layer6 heterozygous mutants (Tbr1flo/flo::Ntsr1-cre::tdTomato/+ , Fig. 7E-E’) and Tbr1layer6 homozygous mutants (Tbr1flo/flo::Ntsr1-cre::tdTomato/+ , Fig. 7F-F’) were tested. Confocal fluorescent microscopy analysis of the synapse numbers showed a ~35% reduction in excitatory synaptic punctae on Tbr1layer6 heterozygous (BD= 0.786, p<0.0001) and a ~75% decrease in Tbr1layer6 homozygous mutants (BD= 0.303, p<0.0001) at P21 (Fig. 7G). The decrease in excitatory synapse numbers is also present at P56 (Supplementary Fig. 3G), in which the excitatory synapses
have decreased ~35% in Tbr1\textsuperscript{layer6} heterozygous (BD=0.501, p<0.0001) and ~64% in Tbr1\textsuperscript{layer6} homozygous mutants (BD= 0.362, p<0.0001; Supplementary Fig. 3G).

Reduced excitatory synaptic density at P21 (Fig. 7D-G) and P56 (Supplementary Fig. 3D-G), led us to measure spontaneous EPSCs using whole-cell patch clamp at both P21 and P56. We recorded from neurons expressing Ntsr1-cre, identified using the fluorescent Ai14 tdTomato Cre-dependent reporter, in the somatosensory cortex (S1) from coronal slices (Fig. 7A) of Tbr1\textsuperscript{wildtype}, Tbr1\textsuperscript{layer6} heterozygous, and Tbr1\textsuperscript{layer6} homozygous mutant mice at P21 or P56. To examine the sEPSC, we made voltage clamp recordings at -70mV using a cesium-based intracellular solution. The frequency of sEPSC are reduced in Tbr1\textsuperscript{layer6} homozygous mutants as compared to cells from Tbr1\textsuperscript{wildtype} mice at P21 (Fig. 7H, 7I: n = 7/7/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, F\textsubscript{(2,18)} = 6.625, p = 0.007; t-test, Tukey correction, wildtype v. homozygous: q\textsubscript{(18)} = 5.123, p = 0.0053). The decrease in sEPSC frequency was also present at P56 (Supplementary Fig. 3H, 3I: n = 7/7/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, F\textsubscript{(2,18)} = 10.17, p = 0.0011; t-test, Tukey correction, wildtype v. homozygous: q\textsubscript{(18)} = 6.371, p = 0.0008).

As depicted in Fig. 8I, brain sections from mPFC (Fig. 8A) of Tbr1\textsuperscript{wildtype} (Rbp4-cre::tdTomato\textsuperscript{f/+}, Fig. 8D-D'), Tbr1\textsuperscript{layer5} heterozygous mutants ( Tbr1\textsuperscript{f/+}::Rbp4-cre::tdTomato\textsuperscript{f/+}, Fig. 8E-E') and Tbr1\textsuperscript{layer5} homozygous mutants (Tbr1\textsuperscript{f/f}::Rbp4-cre::tdTomato\textsuperscript{f/+}, Fig. 8F-F') were tested. Confocal fluorescent microscopy analysis of the synapse numbers showed a ~35% reduction in excitatory synaptic punctae on Tbr1\textsuperscript{layer5} heterozygous (BD= 0.786, p<0.0001) and a ~75% decrease in Tbr1\textsuperscript{layer5} homozygous mutants (BD= 0.303, p<0.0001) at P21 (Fig. 8D). The decrease in excitatory synapse numbers is also present at P56 (data not shown).
Reduced excitatory synaptic density at P21 (Fig. 8D-G) and P56 (data not shown), led us to measure spontaneous EPSCs using whole-cell patch clamp at both P21 and P56. We recorded from neurons expressing *Rbp4-cre*, identified using the fluorescent Ai14 tdTomato Cre-dependent reporter, in the mPFC from coronal slices (Fig. 8A) of *Tbr1*<sup>wildtype</sup>, *Tbr1<sup>layer5</sup*</sup> heterozygous, and *Tbr1<sup>layer5</sup*</sup> homozygous mutant mice at P21 or P56. To examine the sEPSC, we made voltage clamp recordings at -70mV. The frequency of sEPSC are reduced in *Tbr1<sup>layer5</sup*</sup> homozygous mutants as compared to cells from *Tbr1*<sup>wildtype</sup> mice at P21 (Fig. 8H, 8I: n = 6/6/6, wildtype/ heterozygous/ homozygous; One-way ANOVA, F<sub>(2,15)</sub> = 23.18, p < 0.001; t-test, Tukey correction, wildtype v. homozygous: q<sub>(15)</sub> = 9.416, p < 0.0001; heterozygous v. homozygous: q<sub>(15)</sub> = 6.455, p = 0.001). The decrease in sEPSC frequency was also present at P56 (Fig. 8H, 8I: n = 6/6/6, wildtype/ heterozygous/ homozygous; One-way ANOVA, F<sub>(2,15)</sub> = 19.76, p < 0.0001; t-test, Tukey correction, wildtype v. homozygous: q<sub>(15)</sub> = 8.582, p < 0.0001).

*Tbr1<sup>layer6</sup*</sup> and *Tbr1<sup>layer5</sup*</sup> mutants have altered cortical interneuron lamination and reduced inhibitory synaptic density at P21 and P56.

The pattern of SST<sup>+</sup> cortical interneuron (CIN) lamination is abnormal in P3 *Tbr1<sup>layer6</sup*</sup> mutants (Supplementary Fig. 4). In *Tbr1<sup>layer6</sup*</sup> mutants, the distribution of SST+ CINs are reduced in layer 6, unchanged in layer 5 and increased in layers 2-4 (Supplementary Fig. 4A-A”). These changes are consistent with previous studies that have demonstrated the role of cortical pyramidal neurons in regulating the distribution of cortical interneurons (Lodato et al., 2011)

This result, in conjunction with the reduction in excitatory synapses (layer 6, Fig. 7I; layer 5, Fig 8I) led us to measure inhibitory synapse numbers in the *Tbr1<sup>layer6</sup*</sup> and *Tbr1<sup>layer5</sup*</sup> mutants. From confocal images, we counted the numbers of inhibitory terminals (VGAT<sup>+</sup>
presynaptic structures) apposed to dendritic postsynaptic zones (Gephyrin⁺) of Tbr1wildtype (layer 6, Fig. 7J-J’, Supplementary Fig. 3J-J’; layer 5, Fig. 8. J-J’), Tbr1 heterozygous mutants (layer 6, Fig. 7K-K’, Supplementary Fig. 3K-K’; layer 5, Fig. 8 K-K’) and Tbr1layer6 homozygous mutants (layer 6, Fig. 7L-L’, Supplementary Fig. 3L-L’; layer 5, Fig. 8L-L’) onto the apical dendrites of layer 6 pyramidal neurons at P21 (Fig. 7II) and P56 (Supplementary Fig. 3II) as well as onto the apical dendrites of layer 5 pyramidal neurons at P21 (Fig. 8II) and P56 (data not shown).

Confocal fluorescent microscopy analysis of the synapse numbers demonstrated a ~37% reduction in inhibitory synaptic density in Tbr1layer6 heterozygous mutants (BD= 0.574, p<0.0001) and a ~72% decrease in Tbr1layer6 homozygous mutants (BD= 0.252, P<0.0001) at P21 (Fig. 7J). The decrease in inhibitory synapse numbers is also present at P56 (Supplementary Fig. 3M), in which the inhibitory synapses have decreased ~33% in Tbr1layer6 heterozygous mutants (BD= 0.673, p<0.0001) and ~66% in Tbr1layer6 homozygous mutants (BD= 0.346, p<0.0001; Fig. 6M).

To measure the inhibitory synaptic input onto the layer 6 and layer 5 pyramidal neurons in vivo, we measured spontaneous IPSCs (sIPSC) using whole-cell patch clamp at both P21 and P56. We made voltage clamp recordings at +10mV using a cesium-based intracellular solution. In slices from P21 Tbr1layer6 homozygous mutant mice, we observed reduced sIPSC frequency (Fig. 7N, 7O: n = 7/6/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, F(2,17) = 4.738, p = 0.023; t-test, Tukey correction, wildtype v. homozygous: q(17) = 3.847, p = 0.037, heterozygous v. homozygous: q(17) = 3.635, p = 0.0495). sIPSC frequency remained reduced at P56 (Supplementary Fig. 3N, 3O: n = 8/8/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, F(2,20) = 12.44, p = 0.0003; t-test,
Tukey correction, wildtype v. homozygous: \( q_{20} = 6.907, p = 0.0003 \), heterozygous v. homozygous: \( q_{(20)} = 4.901, p = 0.0066 \).

In slices from P21 \( Tbr1^{layer5} \) homozygous mutant mice, we observed reduced sIPSC frequency (Fig. 8N, 8O: \( n = 7/7/7 \), wildtype/ heterozygous/ homozygous; One-way ANOVA, \( F_{(2,18)} = 5.159, p = 0.0169 \); t-test, Tukey correction, wildtype v. homozygous: \( q_{(18)} = 4.534, p = 0.0129 \)). sIPSC frequency remained reduced at P56 (Fig. 8N, 8O: \( n = 7/7/7 \), wildtype/ heterozygous/ homozygous; One-way ANOVA, \( F_{(2,18)} = 6.694, p = 0.0067 \); t-test, Tukey correction, wildtype v. homozygous: \( q_{(18)} = 5.15, p = 0.005 \)).

**\( Tbr1 \) mutants have an increase in hyperpolarization-activated cation currents (\( I_h \)).**

Given that L6 pyramidal neurons in \( Tbr1^{layer6} \) mutants exhibit anatomical changes suggestive of L5 phenotypes (Fig. 4, 6), we wondered whether similar changes might be present in the intrinsic physiology of L6 pyramidal neurons in \( Tbr1^{layer6} \) mutants. We used whole-cell patch clamp to measure intrinsic physiological properties of labelled \( Ntsr1\)-cre expressing pyramidal neurons within L6 of S1 (Fig. 9a & b). At P56 and P21, resting membrane potential (Fig. 9a B, Supplementary Fig. 5B) and input resistance (Fig. 9a C, Supplementary Fig. 5C) were not different between \( Tbr1^{wildtype} \), \( Tbr1^{layer6} \) heterozygous, and \( Tbr1^{layer6} \) homozygous mutant cells (\( n=8 \) of each genotype). A prominent feature of many L5 pyramidal neurons that is largely absent from L6, is hyperpolarization-activated cation current (\( I_h \)) mediated by HCN channels (Gee et al., 2012). \( I_h \) causes a characteristic "sag" and "rebound" in current clamp recordings of responses to steps of hyperpolarizing current. We examined responses to a \(-200\) pA step, and found that S1 L6 pyramidal neurons from P56 \( Tbr1^{layer6} \) heterozygous and homozygous mutant mice exhibited significantly increased sag + rebound compared to \( Tbr1^{wildtype} \) controls, suggesting increased h-current, while other intrinsic electrophysiological properties were largely unaltered (Fig. 9a D: \( n = 7/6/7 \),
wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,17)} = 13.18$, $p = 0.0003$; t-test, Tukey correction, wildtype v. heterozygous: $q_{(17)} = 3.693$, $p = 0.0457$; wildtype v. homozygous: $q_{(17)} = 7.258$, $p = 0.0002$).

The neurons from $Tbr1^{layer6}$ homozygous mutants at P21 also exhibited an increased sag and rebound compared to $Tbr1^{wildtype}$ controls, suggesting increased h-current (Supplementary Fig. 5D: $n = 8/8/8$, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,21)} = 17.68$, $p < 0.001$; t-test, Tukey correction, wildtype v. homozygous: $q_{(21)} = 8.331$, $p < 0.0001$; heterozygous v. homozygous: $q_{(21)} = 5.16$, $p = 0.0041$). Other intrinsic electrophysiological properties were largely unaltered at P21 (Supplementary Fig. 5B, 5C).

In deep layer neocortical pyramidal neurons, the presence of $I_h$ shifts the resonant frequency towards higher frequencies (Dembrow, Chitwood, & Johnston, 2010). Therefore, to further characterize potential increases in $I_h$, we estimated the resonant frequency. For this, we injected constant current to hold $Ntsr1^{cre}$ positive neurons in current clamp near -70mV, then introduced a sinusoidal current stimulus with constant amplitude (100 pA peak-to-peak) and a frequency that increased linearly from 0 to 20 Hz over 20 seconds (Fig. 9a E). We used the ratio of the fast Fourier transform of the voltage response (Fig. 9a E top) to the fast Fourier transform of the sinusoidal current stimulus (Fig. 9a E middle) to calculate the impedance amplitude profile (Fig. 9a E bottom). We defined the resonant frequency as the frequency at which the impedance profile reached its peak. At P56-80, both $Tbr1^{layer6}$ heterozygous and $Tbr1^{layer6}$ homozygous mutants exhibit an increase in their resonant frequency compared to $Tbr1^{wildtype}$ controls (Fig. 9b G: $n = 7/8/8$, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,20)} = 16.24$, $p < 0.0001$; t-test, Tukey correction, wildtype v. heterozygous: $q_{(20)} = 7.075$, $p = 0.0002$; wildtype v. homozygous: $q_{(20)} = 7.038$, $p = 0.0002$).
Finally, we blocked $I_h$ by bath applying the specific HCN channel antagonist ZD7288 (25 μM; Fig. 9b F). The resonant frequency was reduced by over 50% in the $Tbr1^{\text{layer}6}$ heterozygous (Fig. 9b G: n = 8; paired T-test, t(7) = 7.723, p < 0.0001) and $Tbr1^{\text{layer}6}$ homozygous mutants (Fig. 9b G: n = 8; paired T-test, t(7) = 8.194, p < 0.0001). However, the resonant frequency was not significantly altered by ZD7288 in the neurons from the $Tbr1^{\text{wildtype}}$ mice, indicating that $I_h$ significantly shifts the resonant frequency of mutant, but not wild-type, L6 pyramidal neurons.

To compare electrophysiological abnormalities linked to loss of $Tbr1$ layer 5 cortical neurons we used whole-cell patch clamp to measure intrinsic physiological properties of labelled $Rbp4$-cre expressing pyramidal neurons within L5 of mPFC. At P21-28, resting membrane potential (Fig 10B) and input resistance (Fig 10C), were not different between $Tbr1^{\text{wildtype}}$, $Tbr1^{\text{layer}5}$ heterozygous, and $Tbr1^{\text{layer}5}$ homozygous mutant cells (n = 7 of each genotype).

A prominent feature of a subpopulation of dopamine D2 receptor (D2R) expressing L5 pyramidal neurons is hyperpolarization-activated cation current ($I_h$) mediated by HCN channels (Gee et al. 2012; Clarkson et al., 2017). We found that L5 mPFC neurons from P21-28 $Tbr1^{\text{layer}5}$ homozygous mutants have increased sag and rebound compared to $Tbr1^{\text{wildtype}}$ controls, suggesting increased $h$-current (Fig 10D: n = 7/7/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,18)} = 8.823, p = 0.0021$; t-test, Tukey correction, wildtype v. heterozygous: $q_{(18)} = 4.27, p = 0.0192$; wildtype v. homozygous: $q_{(18)} = 5.712, p = 0.0021$).

We also measured the intrinsic properties of mPFC L5 $Rbp4$-cre positive neurons from slices from P56-80 mice. Again, neurons from both $Tbr1$ heterozygous and homozygous mutant mice have significantly increased sag + rebound compared to controls,
while other intrinsic electrophysiological properties were largely unaltered (Fig 10H: n = 7/7/7, wildtype/ heterozygous/ homozygous; One-way ANOVA, \( F_{(2,18)} = 10.11, p = 0.0011; \) t-test, Tukey correction, wildtype v. heterozygous: \( q_{(18)} = 4.565, p = 0.0123; \) wildtype v. homozygous: \( q_{(18)} = 6.115, p = 0.0011 \)).

Additionally, to further characterize potential increases in \( I_{\text{h}} \), we estimated the resonant frequency \( Tbr1^{\text{layer5}} \) mutant neurons as described for \( Tbr1^{\text{layer6}} \) mutants in Figure 9. At P56-80, both \( Tbr1^{\text{layer5}} \) heterozygous and \( Tbr1^{\text{layer5}} \) homozygous mutants exhibit an increase in their resonant frequency over \( Tbr1^{\text{wildtype}} \) (Fig 11C: n = 8/6/8, wildtype/heterozygous/homozygous; One-way ANOVA, \( F_{(2,19)} = 23.93, p < 0.0001; \) t-test, Tukey correction, wildtype v. heterozygous: \( q_{(19)} = 6.229, p = 0.008; \) wildtype v. homozygous: \( q_{(19)} = 9.57, p < 0.0001 \)). Finally, we blocked \( I_{\text{h}} \) by bath applying the specific HCN channel antagonist ZD7288 (25 μM; Fig 11B). The resonant frequency was reduced by over 50% in the \( Tbr1^{\text{wildtype}} \) mice (Fig 11C: n = 8; paired T-test, \( t(7) = 4.646, p = 0.0024 \)), \( Tbr1^{\text{layer5}} \) heterozygous (Fig 11C: n = 6; paired T-test, \( t(5) = 8.099, p = 0.0005 \)) as well as in the \( Tbr1^{\text{layer5}} \) homozygous mutants (Fig 10C: n = 8; paired T-test, \( t(7) = 8.576, p < 0.0001 \)).

\( Tbr1^{\text{layer6}} \) mutants exhibit increased aggressive behaviors.

To identify behavioral abnormalities linked to loss of \( Tbr1 \) function in L6 neurons, we performed multiple behavioral assays in littermate cohorts of \( Tbr1^{\text{wildtype}}, Tbr1^{\text{layer6}} \) heterozygous, and \( Tbr1^{\text{layer6}} \) homozygous mutant male and female mice between P56-80. Because \( Tbr1 \) is a high confidence ASD gene, we explored the impact of \( Tbr1 \) loss of function mutation in layer 6 on ASD associated behaviors such as motor ability, anxiety, and social interaction. \( Tbr1^{\text{layer6}} \) heterozygous and \( Tbr1^{\text{layer6}} \) homozygous mutants did not show impairments in locomotion as measured by speed in the open field (Fig. 12A) or in motor coordination as measured by performance on a rotarod (Fig. 12 C) compared to wildtype.
controls. $Tbr1^\text{layer6}$ mutants did not show significant difference in anxiety, measured by time spent in the center of a novel open field (Fig. 12 B). However, in the elevated plus maze $Tbr1^\text{layer6}$ heterozygous, and $Tbr1^\text{layer6}$ homozygous mutants did spend an increased time in the closed arms (Fig. 12D: $n = 11/8/9$, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,25)} = 4.155$, $p = 0.028$; t-test, Tukey correction, wildtype v. heterozygous: $q_{(25)} = 4.065$, $p = 0.022$).

To assay social behavior, we measured the time each experimental mouse spent exploring a novel juvenile wildtype mouse of the same sex introduced to its home cage. Subsequently, we measured the amount of time the subject mouse spent exploring a novel object introduced to its home cage. We did not find differences in the time $Tbr1^\text{layer6}$ mutants spent exploring a novel object (Fig. 12E). However, in the $Tbr1^\text{layer6}$ homozygous mutants, we observed a significant increase in the total amount of time the adult spent with the novel juvenile mouse (Fig. 12F: $n = 11/8/9$, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,25)} = 4.534$, $p = 0.021$; t-test, Tukey correction, wildtype v. homozygous: $q_{(25)} = 3.731$, $p = 0.0364$). To further assess the type of interaction, we divided this time into “social interaction,” defined as time spent in sniffing, close following (not followed by periods of aggression), or allo-grooming, vs. “aggressive” behavior, defined as biting, fighting, or close following associated with periods of active fighting. $Tbr1^{\text{wildtype}}$, $Tbr1^\text{layer6}$ heterozygous, and $Tbr1^\text{layer6}$ homozygous mutants spent similar time in social interaction, particularly during the first 2 minutes of the 5 minutes social interaction task (Fig. 12F). $Tbr1^\text{layer6}$ homozygous mutants, however, also exhibited prolonged periods of aggressive interaction with the juvenile (Fig. 12F: $n = 11/8/9$, wildtype/ heterozygous/ homozygous; One-way ANOVA, $F_{(2,25)} = 10.97$, $p = 0.0004$; t-test, Tukey correction, wildtype v. homozygous: $q_{(25)} = 6.237$, $p = 0.0005$).
*Tbr1*<sub>layer5</sub> mutants exhibit increased open arm exploration and decreased social interaction.

We assessed ASD-related behaviors, specifically motor ability, anxiety, and social interaction, on littermate cohorts of *Tbr1*<sub>wildtype</sub>, *Tbr1*<sub>layer5</sub> heterozygous, and *Tbr1*<sub>layer5</sub> homozygous mutant male and female mice between P56-80. *Tbr1*<sub>layer5</sub> heterozygous and *Tbr1*<sub>layer5</sub> homozygous mutants did not show impairments in locomotion as measured by speed in the open field (Fig 13A) or in motor coordination as measured by performance on a rotarod (Fig 13C) compared to wildtype controls. *Tbr1*<sub>layer5</sub> mutants did not show significant difference in anxiety, measured by time spent in the center of a novel open field (Fig 13B). However, in the elevated plus maze *Tbr1*<sub>layer5</sub> homozygous mutants did spend an increased time in the open arms (Fig 13D: n = 11/8/9, wildtype/heterozygous/homozygous; t-test, wildtype v. heterozygous: t<sub>19</sub> = 2.355, p = 0.0294) suggesting decreased anxiety or increased exploratory drive in the *Tbr1*<sub>layer5</sub> homozygous mutant mice.

To specifically assay social behavior, we measured the time the experimental mouse spent exploring a novel juvenile wildtype mouse of the same sex introduced to its home cage. Subsequently, we measured the amount of time the subject mouse spent exploring a novel object introduced to its home cage. We did not find differences in the time *Tbr1*<sub>layer5</sub> mutants spent exploring a novel object (Fig 13E). However, in the *Tbr1*<sub>layer5</sub> homozygous mutants, we observed a significant decrease in the time spent interacting with the novel juvenile mouse (Fig 13F: n = 11/5/12, wildtype/heterozygous/homozygous; One-way ANOVA, F<sub>(2,25)</sub> = 16.51, p < 0.0001; t-test, Tukey correction, wildtype v. homozygous: q<sub>(25)</sub> = 7.124, p < 0.0001; heterozygous v. homozygous: q<sub>(25)</sub> = 6.295, p = 0.0004 ). This pronounced reduction of social exploration in the *Tbr1*<sub>layer5</sub> homozygous mutants is suggestive of core ASD behavioral characteristics.
Discussion

Persistent Tbr1 function drives a partially irreversible sequence of developmental processes: Neonatal Tbr1 specifies properties of a sublamina of neocortical layer 6 (corticothalamic) and represses layer 5 (corticofugal) identity.

Tbr1 is expressed in the excitatory neurons of the neocortex (subplate, layer 6, rostral layer 5, layers 2/3), hippocampus, entorhinal cortex, pallial amygdala, piriform cortex, olfactory bulb and Cajal Retzius neurons (Hevner, Neogi, Englund, Daza, & Fink, 2003; Hevner et al., 2001). Analysis of Tbr1\textsuperscript{constitutive} null mice demonstrated its function in the differentiation of the first waves of pallial glutamatergic neurons including Cajal Retzius cells, olfactory bulb mitral cells, subplate cells and layer 6 cells (Bedogni et al., 2010; Bulfone et al., 1998; Hevner, Miyashita-Lin, & Rubenstein, 2002). Further analyses of Tbr1\textsuperscript{constitutive} null mice revealed that Tbr1 promotes the identity of layer 6 neurons by repressing layer 5 molecular properties in layer 6 pyramidal neurons (McKenna et al., 2011).

In the present study, using conditional mutagenesis, we have demonstrated that Tbr1 is required in maintaining subplate and layer 6 identity. The impaired differentiation of subplate and layer 6 neurons is indicated both molecularly (Figs. 2, 3, 4) and by dendritic defects (Fig. 6). Molecularly, Tbr1\textsuperscript{layer6} mutant neurons have reduced expression of layer 6 markers including Wnt7b, Foxp2 and Tle4 (Figs. 2, 4) and possess an increased expression of genes controlling layer 5 molecular properties including Bcl11b, Fez1, Foxp1, Epha7 and Grin3a (Figs. 2, 4 and Supplementary Fig. 4). Strikingly, the expression of Tle4 and Foxp2 is mainly reduced in the deep layer 6 neurons (layer 6b), even though Tbr1 expression is uniformly reduced in the layer 6 of Tbr1\textsuperscript{layer6} mutant (Fig. 4). This phenomenon may suggest that Tbr1 may have preferential function within layer 6. The ectopic expression of layer 5 markers in the Tbr1\textsuperscript{layer6} mutants suggests that layer 6 neurons have changed fate to a layer
5 identity (Fig. 2, 4). The change in the dendritic patterning of the $Tbr1^{\text{layer6}}$ mutants further strengthens the hypothesis that the layer 6 neurons have changed fate to a layer 5 identity, in which the $Tbr1^{\text{layer6}}$ mutants extends their apical dendrites superficially into the marginal zone (Fig. 6) (Lefebvre, Sanes, & Kay, 2015).

Preliminary investigation into gene regulation by $Tbr1$ in layer 5 neurons did not draw any conclusive results. Batch-FACS RNA-seq on $Tbr1^{\text{layer5}}$ mutant cells did not find any significant variations in RNA expression suggesting differential gene regulation in $Tbr1^{\text{wildtype}}$ versus $Tbr1^{\text{layer5}}$ mutants. Previous studies have described layer 5 pyramidal neurons throughout somatosensory, visual, and frontal cortices as heterogeneous populations with different projection targets, morphology, electrophysiological properties, and receptor expression (Morishima and Kawaguchi, 2006; Dembrow et al., 2010, Gee et al., 2012; Harris and Shepherd, 2015; Clarkson et al., 2017). Batch RNA-seq likely failed to turn up significant results because of differential gene regulation by $Tbr1$ within the different subpopulations of pyramidal neurons within layer 5. In order to address this issue, we are planning to perform single-cell RNA-seq on $Tbr1^{\text{layer5}}$ mutant cells.

Lastly, both $Tbr1^{\text{layer6}}$ and $Tbr1^{\text{layer5}}$ mutant neurons have increased levels of h-current (layer 6, Fig. 9a & b and Supplementary Fig. 5; layer 5, Fig. 10), which is one of the physiological properties of D2R-expressing layer 5 pyramidal neurons (Clarkson et al., 2017; Gee et al., 2012; Lee et al., 2014). Recent studies have demonstrated that there are at least two subclasses of pyramidal neurons within layer 5 (Gee et al., 2012; Seong & Carter, 2012). These pyramidal subtypes are part of distinct circuits. ‘Type B’ pyramidal cells are thin-tufted pyramidal neurons that express $Drd1$ gene and are intratelencephalic (IT), with projections to contralateral cortex and lack h-current. ‘Type A’ cells are thick-tufted pyramidal neurons that express $Drd2$ gene, with projections to thalamus and pons but not to contralateral cortex and possess h-current (Dembrow, Chitwood, & Johnston, 2010; Gee et
al., 2012). In the $Tbr1^{layer6}$ mutants, the decreased expression of $Drd1$ (Fig. 2) and increased levels of HCN1 protein (Supplementary Fig. 5) suggests that during the layer 6 to layer 5 fate change, there is a bias towards generating more type A layer 5 pyramidal neurons. Furthermore, the elevated h-current resembles the physiological properties of type A pyramidal neurons that are usually positioned in layer 5b (Clarkson et al., 2017; Dembrow et al., 2010; Gee et al., 2012; Lee et al., 2014). Thus, $Tbr1$ persistent function is required to initiate, orchestrate and maintain layer 6 specific program of molecular, dendritic and physiological properties. Although we currently lack conclusive RNA-seq data from $Tbr1^{layer5}$ mutants, the electrophysiological characteristic of exaggerated h-current (Fig. 10 and Fig. 11) suggests that loss of $Tbr1$ function within layer 5 also drives generation of type A, D2R expressing, pyramidal neurons.

Contrary to the $Tbr1^{constitutive}$ null mice where corticothalamic axons fail to grow and enter the thalamus (Hevner et al., 2002; Hevner et al., 2001), $Tbr1^{layer6}$ mutants have corticothalamic projections that innervate the thalamus (Supplementary Fig. 1 and 2). However, their ramification within the thalamus is abnormal with decreased thalamic innervation in the anterior and anteromedial structure of thalamus (Supplementary Fig. 1 and 2). This suggests that even though $Tbr1$ expression is required to initiate the corticothalamic pathway, it is not required to maintain these axons through P56 in the $Tbr1^{layer6}$ mutant. Furthermore, ectopic subcortical projections or corpus callosum projections are not generated by the $Tbr1^{layer6}$ mutant. Thus, despite taking on many layer 5 properties, the mutant layer 6 neurons do not grow layer 5-like axonal projections. This suggests that once the layer 6 program regulating axonal pathway choice is established by $Tbr1$, it is irreversibly maintained after approximately E18, once $Tbr1$ is eliminated in the layer 6 neurons. On the other hand, $Tbr1$ dependent programs for promoting layer 6 gene expression, repression of layer 5 gene expression, layer 6-specific dendritic patterning and
H-current physiological properties remain plastic and are dependent upon Tbr1 function during later stages of development and adulthood (Fig. 2, 3, 4, and Supplementary Fig. 3 and 4).

**Tbr1 directly regulates the transcription of genes that control layer 6 identity**

Towards elucidating TBR1-regulated transcriptional pathways that control layer 6 properties, we combined transcriptomic analysis of FACS purified neonatal wildtype and Tbr1layer6 mutant (Fig. 2), with whole genome neonatal TBR1 ChIP-Seq (Fig. 5). This provides evidence to identify candidate regulatory elements (RE) that are bound by TBR1.

Transcription assays in neonatal neocortical primary cultures demonstrated that TBR1 functions as an activator or repressor of REs adjacent to genes whose expression changes in Tbr1layer6 mutant neurons (Fig. 3). Importantly, TBR1 activated REs near to genes whose expression was reduced in layer 6, and repressed REs near to genes whose expression was increased in layer 5. Future studies are needed to identify the nuclear cofactors that determine whether TBR1 acts as a transcriptional activator or repressor, although it is conceivable that the DNA sequence of the REs modifies TBR1’s confirmation to control its activity. The discovery of these TBR1 regulated REs opens the possibility that these elements will show in vivo layer-specific activity which could be elucidated using transgenic experiments. These REs also serve as essential nodes for establishing the transcriptional circuits that drive TBR1 mediated gene expression.

**Tbr1 may be involved in the processes that control excitatory and inhibitory synaptic development and maintenance.**

A reduced density of excitatory and inhibitory dendritic synapses is a central phenotype of both the Tbr1layer6 and the Tbr1layer5 heterozygous and homozygous mutants
as seen in tissue sections from P21 and P56 (layer 6, Fig. 7 and Supplementary Fig. 3; layer 5; Fig. 8). A similar reduction was also seen in primary cultures grown from P2 cortex (data not shown). These findings were substantiated using slice physiology, where the $Tbr1^{layer6}$ and $Tbr1^{layer5}$ mutant neurons exhibit reduced sEPSCs and sIPSCs at P21 and P56 (layer 6, Fig. 7 and Supplementary Fig. 3; layer 5, Fig. 8).

We took advantage of transcriptome analysis of FAC-sorted RNA-Seq from neonatal layer 6 neurons of wildtype and $Tbr1^{layer6}$ mutants to further elucidate and understand the $Tbr1$-regulated pathways that could contribute to the reduced synaptic phenotype (Fig. 2). $Tbr1$ regulates the transcription of a large cohort of genes that could be involved in synaptic development and maintenance through regulating processes that may involve retinoic acid levels, G-protein coupled receptors, Wnt canonical signaling, cell adhesion molecules, cell-cell signaling, axonal growth, neurotransmission, etc. (Krishnan & Schöth, 2015; Leung & Wong, 2017; Lim et al., 2009; Muram, Rowe, & Hirasawa, 2016; Yee & Chen, 2016). For instance, Wnt proteins comprise a large class of signaling molecules that regulate a variety of developmental processes, including synapse formation (Davis, Zou, & Ghosh, 2008; Liebl, McKeown, Yao, & Hing, 2010). Furthermore, it has been demonstrated that $Wnt7b$ activates canonical wnt signaling (Davis et al., 2008). The $Wnt7b$ activation of the canonical pathway has been linked to increased presynaptic inputs in the developing hippocampus (Davis et al., 2008).

Dysregulation of approximately 33% (66/178) $Tbr1$ targets in $Tbr1^{layer6}$ mutants of genes that are linked to biological processes such as regulating retinoic acid levels, G-protein coupled receptors, wnt canonical signaling and cell adhesion may contribute to the decreased synaptic density phenotype in the layer 6 mutant neurons (Fig. 7, Supplementary Fig. 3). The reduction in the synaptic transmission may lead to abnormal excitation and
inhibition levels in the cortical circuitry of the $Tbr1^{layer6}$ mutants. Although the etiology of ASDs is heterogeneous, the two most prominent models for autism pathogenesis are abnormal brain wiring and an imbalance of excitation/inhibition (Bernardinelli, Nikonenko, & Muller, 2014; Rubenstein & Merzenich, 2003; Walsh, Morrow, & Rubenstein, 2008). These two deficits lead to abnormal information processing and may result in autism-like behaviors. The absence of correct excitatory or inhibitory inputs could alter the activity of the target neurons. These inappropriate levels of neuronal activity may result in abnormal information processing, which may lead to aberrant behaviors.

$Tbr1^{layer6}$ mutants exhibit Autism-like behaviors with increased anxiety and aggressive behaviors.

$Tbr1^{layer6}$ mutants are viable allowing us to interrogate their behavior, which was remarkably normal in many assays including assays of their motor functions, interest in novel objects, an assay of social interaction (Fig. 12). On the other hand, a measure of exploration, (time spent in the closed arms of an elevated plus maze), showed that both heterozygote and homozygotes $Tbr1^{layer6}$ mutants spent more time in the closed arms (Fig. 12D), suggesting that they may have increased anxiety in open spaces. Furthermore, homozygous $Tbr1^{layer6}$ mutants exhibited prolonged periods of aggressive behaviors with the juvenile mice (Fig. 12F). Increased anxiety and aggressive behaviors have been observed in some human patients with ASD (Gadow et al., 2004; McClintock et al., 2003). That loss of function of $Tbr1$ within layer 6 impacts aggressive behaviors without deficits in normal social interactions suggests that $Tbr1$ play diverse roles in layer 6 and layer 5 which differentially impact behavior.

$Ntsr1-Cre$ recombination of $Tbr1$ does not extend into many cortical regions, including the olfactory bulb, medial prefrontal neocortex, dorsomedial neocortex (cingulate
and retrosplenial), hippocampus and parahippocampus, piriform cortex, and pallial amygdala, ruling out the possibility that defects in $Tbr1^+$ neurons in these structures contributes to the behavior phenotypes. This is pertinent, as $Tbr1^{\text{constitutive}}$ heterozygotes have abnormal amygdala connectivity that has been associated with deficits in social interaction, cognitive flexibility and associative memory (Huang et al., 2014). Thus, the highly specific molecular and physiological defects in the early born pyramidal neurons of the neocortical subplate and layer 6 can be implicated in the mutant’s increased aggression and anxiety.

$Tbr1^{\text{layer5}}$ mutants exhibit Autism-like behaviors with decreased social behaviors.

Unlike $Tbr1^{\text{constitutive null}}$ mice which typically die at birth, $Tbr1^{\text{layer5}}$ mutants are viable allowing us to study the role of $Tbr1$ in mature neurons. The $Tbr1^{\text{layer5}}$ mutants performance was notably normal in several behavior tasks including assays of their motor functions and exploration of novel objects (Fig. 13). On the other hand in an elevated plus maze, a measure of anxiety and exploration, showed that homozygotes $Tbr1^{\text{layer5}}$ mutants spent more time in the open arms (Fig. 13D), suggesting that they may have either decreased anxiety or an increased drive for exploration. Furthermore, homozygous $Tbr1^{\text{layer5}}$ mutants exhibited a pronounced reduction in novel juvenile social interactions (Fig. 13F). Interestingly, loss of $Tbr1$ function selectively throughout layer 5 and not layer 6 cortical neurons results in this severe social phenotype, reminiscent of a core ASD behavioral phenotypes in humans.
**Tbr1 loss-of-function mutations are associated with Autism Spectrum Disorders (ASDs).**

Genetic analyses of patients with ASD have identified TBR1 as a high confidence risk factor for ASDs (De Rubeis et al., 2014; Iossifov et al., 2014; Sanders et al., 2012; Willsey et al., 2013). Transcriptome analysis of FACS layer 6 neurons of neonatal wildtype and Tbr1\textsuperscript{layer6} mutant revealed that Tbr1 regulates a small group of ASD genes including Scn2a1, Foxp1, Wnt7b, Nuak1 (De Rubeis et al., 2014; Sanders et al., 2015) and Foxp2 (Gong et al., 2004; Li, Yamagata, Mori, & Momoi, 2005). Microarray analysis of wildtype and Tbr1 mutant (data not shown) as well as in situ hybridization experiments (Fig. 4H-H'), revealed that Bcl11a expression, a probable ASD gene (De Rubeis et al., 2014), is also reduced in Tbr1 mutants.

*De novo* and inherited mutations in TBR1 have been found in patients with ASDs (De Rubeis et al., 2014; Iossifov et al., 2014; Sanders et al., 2012; Willsey et al., 2013). Two of the TBR1 *de novo* mutations result in early termination and generate truncated proteins that lack a functional DNA-binding T-box domain (De Rubeis et al., 2014). It has been demonstrated that the two truncated TBR1 mutants can no longer function in transcription or in interactions with CASK (Huang & Hsueh, 2017; Wang et al., 2004) and FOXP2 (Deriziotis et al., 2014). Furthermore, it has been shown that *de novo* TBR1 mutations change the subcellular distribution of TBR1 (Deriziotis et al., 2014). All together, these studies demonstrated that *de novo* TBR1 mutations in ASD patients impair TBR1 ability to interact with FOXP2 (Deriziotis et al., 2014). Furthermore, FOXP2 and TBR1 are only co-expressed in layer 6 of the cerebral cortex and not in other layers of the cerebral cortex and amygdala. Thus, the interaction with FOXP2 can only partly account for the function of TBR1 during the development of pyramidal neurons of deep cortical layer 6.
Figure 1. Organization of the Tbr1 wildtype and conditional mutant (Tbr1f) alleles.

(A) Schematic representations of the Tbr1 wildtype allele, Tbr1 targeting vector, and Tbr1 conditional mutant allele (Tbr1f). The wildtype Tbr1 allele has six known exons (numbered black boxes, 1-6); the initiation codon is in exon 1, and the termination codon is in exon 6. The white boxes indicate the 5' and 3' UTRs. Red arrowheads correspond to the location of LoxP sites; the black boxes with an F inside are FRT sites. Flipase removes the Neomycin expression cassette (grey box with Neo inside). Upon Cre recombination, exons 2 and 3 are deleted to generate Tbr1 mutant allele. The positions of the qPCR primers used for genotyping are indicated with blue arrowheads under exons 1, 2 and 4. The location of TBR1 antigen that was detected with antibodies for western blotting (panel E) and for immunohistochemistry (Figure 5) is indicated by the red line under part of exon 1. (B) Amino acid alignment of full-length TBR1 protein (black) with TBR1 antigen (red). Black arrowheads represent exonic boundaries. Region highlighted in yellow indicates the amino acid sequence corresponding to the deleted region in the Tbr1 mutant allele. (C) qRT-PCR results corresponding to relative expression levels of Tbr1 exons 1 and 2 (E1+E2) as well as exons 1 and 4 (E1+E4) transcripts in the cortex of Tbr1 constitutive null and wildtype littermates at E15.5 and P0. Gene expression levels were analyzed using two biological replicates, each assayed in experimental triplicates. The error bars represent the standard error of the mean of all replicates; gene expression is normalized relative to a housekeeping gene (Ef1α). Relative expression levels in wildtype mice were defined as 1.0 (*P<0.05) (**P<0.01) (***)P<0.001. (D) In situ hybridization (ISH) on coronal sections of P0 forebrain from
wildtype and Tbr1 constitutive null. Tbr1 full-length (FL) probe and Tbr1 probe corresponding to exons 2 and 3 (E2-3) were used during the hybridization step. Layer 6 is labeled as VI. Scale bar = 50 µm. (E) Western blot (WB) to detect TBR1 protein isolated from E15.5 and P0 cortex of a Tbr1 constitutive null and wildtype littermate. Black arrowhead indicates the TBR1 protein (~75 kDa) detected in wildtype at E15.5 and P0.
Figure 2. RNA-seq analysis of FACS purified layer 6 neurons shows differentially regulated RNAs between wildtype and $Tbr1^{layer6}$ mutants.

Volcano plot of differentially expressed RNAs of $Tbr1^{wildtype}$ and $Tbr1^{layer6}$ mutant from FAC-sorted cells at P5. Up-regulated and downregulated genes in $Tbr1^{layer6}$ mutant are in represented in red and blue respectively. Black dots represent a group of genes that did not reach statistical significance ($\alpha=0.05$). The size of each point represents the difference in the median gene expression between $Tbr1^{wildtype}$ and $Tbr1^{layer6}$ mutant samples.
Figure 3. TBR1 transcriptional regulation of candidate enhancer regions in loci adjacent to Tbr1-regulated genes.

Luciferase transcription assay was utilized to measure activity of Tbr1, Foxp2, Grin2b, Foxp1, Fezf2, Drd1, Hcn1 candidate enhancers in primary cultures made from P0 cortex (transfected on 1 DIV; assay on 3 DIV). The reporter activity was measured under enhancer activity alone (red) and enhancer co-transfected with TBR1 (grey). TBR1 activates Tbr1 (FC= 2.3, p= 0.0007), Foxp2 (FC= 2.17, p= 0.0023), Grin2b (FC= 4.11, p= 0.0015), Bcl11a (FC= 3.46, p= 0.0002) and Drd1(FC= 3.57, p<0.0001), whereas TBR1 represses Foxp1 (FC= -2.52, p= 0.0087), Fezf2 (FC= -2.55, p= 0.0015) and Hcn1 (FC= -2.9, p=0.0248). I56i enhancer and pGL4.23 empty vector were used as negative controls. The error bars represent the standard error of the mean of all replicates for each enhancer relative to I56i enhancer and the empty pGL4.23 vector (in which the enhancers were inserted). T-test with Welch’s correction was used for the statistical analysis of the enhancer alone (red) and enhancer co-transfected with TBR1 (grey). (*p<0.05) (**p< 0.01) (***p<0.001).
Figure 4. *Tbr1* is required to maintain layer 6 identity in postnatal cortex.
In situ hybridization on coronal sections of P3 somatosensory cortex (S1) of Tbr1^{wildtype} (A-L) and Tbr1^{layer6} homozygous mutants (Tbr1^{f/f::Ntrs1-cre}) (A'-L'). In Tbr1^{layer6} mutants, Tbr1 (A-A'), Foxp2 (C-C'), Tle4 (D-D') and Wnt7b (E-E'), Ctnn2 (F-F'), Ptpk (G-G') and Bcl11a (H-H') expressions are reduced in layer 6. This decrease is accompanied by an increase in the expression of Tbr1 and Wnt7b in the superficial layers (A-A', E-E'). Nr4a2 expression decreases in the subplate of Tbr1^{layer6} mutants (B-B'). Tbr1^{layer6} mutants have ectopic layer 6 expression of Bcl11b (I-I'), Fez12 (J-J') and Foxp1 (K-K'). In Tbr1^{layer6} mutant, Sst+ CINs are reduced in layer 6, unchanged in layer 5, and increased in layers 2-4 (L-L'). II-IV = layers 2-4, V = layer 5, VI = layer 6. SP: Subplate. Scale bar = 50μm.
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Figure 5. TBR1 binds to the promoter and candidate enhancer regions of genes whose expression changes in Tbr1 mutant cortex.
TBR1 ChIP-Seq on wildtype whole cortex at P2 (red tracks) demonstrates TBR1 binding to either promoter or candidate enhancer regions. Red boxes below the ChIP-Seq track indicate that TBR1 binding reached statistical significance. Genes (exons and RNA transcripts) are shown in blue. Green boxes highlight the candidate enhancer regions that were tested in the luciferase transcription assay. Black boxes indicate genomic regions used in the luciferase assay, that have proven enhancer activity in the E11.5 cortex corresponding hs416 (Tbr1 locus), hs434 (Fezf2 locus) and hs399 (Bcl11a locus). Black arrow indicates the direction of transcription. Genomic scale (in kb) are shown for each locus.
Figure 6. Ectopic growth of layer 6 apical dendrites into superficial layer 1 in Tbr1$^{layer 6}$ mutants.

The endogenous tdTomato fluorescence (red) in the somatosensory cortex of (A) Tbr1$^{wildtype}$ and (B) Tbr1$^{layer 6}$ mutant. These lines had the Ntsr1-cre::tdTomato$^{+/+}$ alleles to label the layer 6 cell bodies and their dendrites. White arrowheads in panel A’- C’ indicate some of the apical dendrites extending through layers 2/3 to layer 1 in Tbr1$^{layer 6}$ mutant. Cortical layers are labelled. Scale bar: 50μm.
FIGURE 7. Loss of Tbr1 in layer 6 reduces excitatory (I) and inhibitory (II) synaptic input onto the layer 6 neurons in the somatosensory cortex at P21.
(A) Schematic representation of coronal slice of somatosensory cortex (S1, blue boxes). The blue box represents the region of somatosensory cortex (S1) utilized for imaging and whole-cell patch clamp experiments. (B, C) Schema of layer 6 projection neuron (red) in somatosensory cortex of $Tbr1^{\text{wildtype}}$ (B) and $Tbr1^{\text{layer6}}$ mutant (C). The rectangles indicate the zone where synapses were analyzed (B, C) at P21. Pipette tip indicates that the soma was patched during the electrophysiology recordings (B, C).

(I) Excitatory synaptic input was analyzed via synaptic bouton staining onto apical dendrites of layer 6 neurons (D–F) and spontaneous EPSC (sEPSC) recordings from the soma of the layer 6 pyramidal neurons (H) of $Tbr1^{\text{wildtype}}$ (D) $Tbr1^{\text{layer6}}$ heterozygous mutants, and (E) $Tbr1^{\text{layer6}}$ homozygous mutants (F) at P21 (D, E, F). These lines had the $Ntsr1^{-}\text{cre::tdTomato}^{f/+}$ alleles to label the layer 6 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (D–F) for quantification (D’–F’). (G) Quantification of VGlut1$^+$ boutons and PSD95$^+$ clusters co-localizing onto the apical dendrites of layer 6 neurons at P21. (H) Sample traces of sEPSC recordings in voltage clamp at -70mV in somatosensory cortex slices from 4 weeks old $Tbr1^{\text{wildtype}}$ (red), $Tbr1^{\text{layer6}}$ heterozygous mutants (green), and $Tbr1^{\text{layer6}}$ homozygous mutants (blue) at P21. (I) Quantification of the sEPSC frequency in layer 6 neurons at P21.

(II) Inhibitory synaptic input was examined through synaptic bouton staining onto apical dendrites of layer 6 neurons (J–L) and spontaneous IPSC (sIPSC) recordings from the soma of the layer 6 neurons (N) of $Tbr1^{\text{wildtype}}$ (J), $Tbr1^{\text{layer6}}$ heterozygote mutants (K), and $Tbr1^{\text{layer6}}$ homozygous mutants (L) at P21 (J, K, L). These lines had the $Ntsr1^{-}\text{cre::tdTomato}^{f/+}$ alleles to label the layer 6 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (J–L) for quantification (J’–L’). (M) Quantification of VGAT$^+$ boutons and gephyrin$^+$ clusters co-localizing onto the apical dendrites of layer 6 neurons at P21. (N) Sample traces of sIPSC recordings in voltage clamp at +10mV in somatosensory cortex slices from 4 weeks old $Tbr1^{\text{wildtype}}$ (red), $Tbr1^{\text{layer6}}$ heterozygous mutants (green), and $Tbr1^{\text{layer6}}$ homozygous mutants (blue) at P21. (O) Quantification of the sIPSC frequency in layer 6 neurons at P21. Two-way ANOVA was used for the statistical analysis of the control, heterozygous and null. (*p<0.05) (**p< 0.01) (***p<0.001) (****p<0.0001). Scale bar (in F and L) = 5µm.
Figure 8. Loss of Tbr1 in layer 5 reduces excitatory (I) and inhibitory (II) synaptic input onto the layer 6 neurons in the somatosensory cortex.
(A) Schematic representation of coronal slice of medial prefrontal cortex (mPFC, blue boxes). The blue box represents the region of mPFC utilized for imaging and whole-cell patch clamp experiments. (B, C) Schema of layer 5 projection neuron (red) in somatosensory cortex of Tbr1<sup>wildtype</sup> (B) and Tbr1<sup>layer5</sup> mutant (C). The rectangles indicate the zone where synapses were analyzed (B, C) at P21. Pipette tip indicates that the soma was patched during the electrophysiology recordings (B, C).

(I) Excitatory synaptic input was analyzed via synaptic bouton staining onto apical dendrites of layer 5 neurons (D-F) and spontaneous EPSC (sEPSC) recordings from the soma of the layer 5 pyramidal neurons (H) of Tbr1<sup>wildtype</sup>, Tbr1<sup>layer5</sup> heterozygous mutants, and Tbr1<sup>layer5</sup> homozygous mutants at P21 and P56. These lines had the Rbp4-cre::tdTomato<sup>f/+</sup> alleles to label the layer 5 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (D-F) for quantification (D’ – F’).

(G) Quantification of VGlut1<sup>+</sup> boutons and PSD95<sup>+</sup> clusters co-localizing onto the apical dendrites of layer 5 neurons at P21. (H) Sample traces of sEPSC recordings in voltage clamp at -70mV in somatosensory cortex slices from 4 and 8 weeks old Tbr1<sup>wildtype</sup> (red), Tbr1<sup>layer5</sup> heterozygous mutants (green), and Tbr1<sup>layer5</sup> homozygous mutants (blue) at P21 and P56.

(I) Quantification of the sEPSC frequency in layer 6 neurons at P21 and P56.

(II) Inhibitory synaptic input was examined through synaptic bouton staining onto apical dendrites of layer 5 neurons (J-L) and spontaneous IPSC (sIPSC) recordings from the soma of the layer 5 neurons (N) of Tbr1<sup>wildtype</sup>, Tbr1<sup>layer6</sup> heterozygote mutants, and Tbr1<sup>layer6</sup> homozygous mutants at P21 and P56. These lines had the Ntsr1-cre::tdTomato<sup>f/+</sup> alleles to label the layer 5 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (J – L) for quantification (J’ – L’).

(M) Quantification of VGat<sup>+</sup> boutons and Gephyrin<sup>+</sup> clusters co-localizing onto the apical dendrites of layer 5 neurons at P21. (N) Sample traces of sIPSC recordings in voltage clamp at +10mV in somatosensory cortex slices from 4 and 8 weeks old Tbr1<sup>wildtype</sup> (red), Tbr1<sup>layer5</sup> heterozygous mutants (green), and Tbr1<sup>layer5</sup> homozygous mutants (blue) at P21 and P56.

(O) Quantification of the sIPSC frequency in layer 5 neurons at P21 and P56. Two-way ANOVA was used for the statistical analysis of the control, heterozygous and null. (*)<i>p</i><0.05) (**<i>p</i><0.01) (**<i>p</i><0.001) (**<i>p</i><0.0001). Scale bar (in F and L) = 5µm.
Figure 9a. Loss of *Tbr1* in layer 6 somatosensory cortex results in an increase in hyperpolarization-activated cation currents (*Ih*).
Whole-cell patch clamp recordings from layer 6 S1 at P56 (A-D) show that many intrinsic electrophysiological properties were unaffected by loss of Tbr1, including resting membrane potential (B), input resistance (C), and action potential halfwidth (data not shown). We estimated Ih by measuring the membrane potential sag and rebound elicited by a –200 pA current step. This characteristic “sag” and “rebound” during and following hyperpolarizing current injection is mediated by hyperpolarization-activated cyclic nucleotide-gated (HCN) channels. Sag and rebound were increased in Tbr1layer6 heterozygous and Tbr1layer6 homozygous mutants at P56.

(E) Neurons were held in current clamp at -70mV. We provided a sinusoidal current of constant amplitude (100 pA peak-to-peak) with its frequency linearly increasing from 0 to 20 Hz over 20 seconds. We used the ratio of the fast Fourier transform of the voltage response (top) to the fast Fourier transform of the sinusoidal current stimulus (middle) to calculate the impedance amplitude profile (bottom). We estimated the resonant frequency to be the frequency at which the impedance profile reached its peak (arrows). Scale bar = 5 mV, 5 s.
Figure 9a. Increased hyperpolarization-activated cation currents (Ih) in Tbr1<sub>layer6</sub> mutants are blocked by HCN channel blocker ZD7288.

(F) Addition of ZD7288, an HCN channel blocker, did not significantly decrease the resonant frequency of Tbr1<sup>wildtype</sup> (red) but did decrease it by over 50% in Tbr1<sup>layer6</sup> heterozygote mutants (green), and Tbr1<sup>layer6</sup> homozygous mutants (blue). (G) Quantification of changes in resonant frequency of Tbr1<sup>wildtype</sup> (red), Tbr1<sup>layer6</sup> heterozygote mutants (green) and Tbr1<sup>layer6</sup> homozygous mutants (blue) after ZD7288 treatment. (**p< 0.01) (***p<0.001).
Figure 10. Loss of Tbr1 in layer 5 mPFC results in an increase in hyperpolarization-activated cation currents (Ih).
Whole-cell patch clamp recordings from layer 5 mPFC at p21-p28 (A-D) and p56-p80 (E-H) show that many intrinsic electrophysiological properties were unaffected by loss of *Tbr1*, including resting membrane potential (B, F), input resistance (C, G), and action potential halfwidth (data not shown). We estimated Ih by measuring the membrane potential sag and rebound elicited by a –200 pA current step. This characteristic “sag” and “rebound” during and following hyperpolarizing current injection is mediated by hyperpolarization-activated cyclic nucleotide-gated (HCN) channels. At both ages p21-p28 (D) and p56-p80 (H), sag and rebound were increased. (*p<0.05) (**p< 0.01).
Figure 11. Loss of Tbr1 increases layer 5 Rbp4-cre positive pyramidal neurons’ resonant frequency within somatosensory cortex.
(A) Neurons were held in current clamp at -70mV. We provided a sinusoidal current of constant amplitude (100 pA peak-to-peak) with its frequency linearly increasing from 0 to 20 Hz over 20 seconds. We used the ratio of the fast Fourier transform of the voltage response (top) to the fast Fourier transform of the sinusoidal current stimulus (middle) to calculate the impedance amplitude profile (bottom). We estimated the resonant frequency to be the frequency at which the impedance profile reached its peak (arrows). Scale bar = 5 mV, 5 s. 

(B) Addition of ZD7288, an HCN channel blocker, did not significantly decrease the resonant frequency of Tbr1\textsuperscript{wildtype} (red) but did decrease it by over 50% in Tbr1\textsuperscript{layer5} heterozygote mutants (green), and Tbr1\textsuperscript{layer5} homozygous mutants (blue). 

(C) Loss of Tbr1 increased resonant frequency in both Tbr1\textsuperscript{layer5} heterozygote mutants (green), and Tbr1\textsuperscript{layer5} homozygous mutants (blue) which was blocked by ZD7288. (**p<0.001).
Figure 12. Loss of Tbr1 in layer 6 Ntsr1cre positive neurons is associated with an increase in aggressive behaviors

All behavioral assays were performed on adult mice age p56-p80 with Tbr1\textsuperscript{wildtype} shown in red, Tbr1\textsuperscript{layer6} heterozygous mutants in green, and Tbr1\textsuperscript{layer6} homozygous mutants in blue. Loss of Tbr1 did not impair movement (A) or motor coordination (C) nor did it affect the time spent exploring the center of an open field arena (B). (D) Tbr1\textsuperscript{layer6} heterozygous mutants (green) and Tbr1\textsuperscript{layer6} homozygous mutants (blue) spent more time in the closed arms of an elevated plus maze than their Tbr1\textsuperscript{wildtype} littermates. Interestingly, loss of Tbr1 did not affect the time spent engaged in social interaction (F) or novel object exploration (E). Tbr1\textsuperscript{layer6} homozygous mutants (blue) of both genders displayed an increase in aggressive behaviors when interacting with a novel juvenile mouse (F). (*p<0.05) (**p<0.001).
Figure 13. Loss of Tbr1 in layer 5 Rbp4-cre positive neurons is associated with an increase in open arm exploration and decreased social interaction

All behavioral assays were performed on adult mice age p56-p80 with Tbr1\textsuperscript{wildtype} shown in red, Tbr1\textsuperscript{layer5} heterozygous mutants in green, and Tbr1\textsuperscript{layer5} homozygous mutants in blue. Loss of Tbr1 did not impair movement (A) or motor coordination (C) nor did it affect the time spent exploring the center of an open field arena (A). (D) Tbr1\textsuperscript{layer5} heterozygous mutants (green) and Tbr1\textsuperscript{layer5} homozygous mutants (blue) spent more time in the open arms of an elevated plus maze than their Tbr1\textsuperscript{wildtype} littermates. Loss of Tbr1 did not affect the time spent engaged novel object exploration (E). Tbr1\textsuperscript{layer6} heterozygous (green) and Tbr1\textsuperscript{layer6} homozygous mutants (blue) of both genders displayed an decrease in social interaction (F). (*p<0.05) (***p<0.001) (****p<0.0001).
A Summary Timeline of Tbr1 Function

Figure 14. Summary timeline for Tbr1 function.
Schematic representation of a timeline of Tbr1 function prenatally into adulthood in mouse. The red arrowhead at E17.5 corresponds to the timing of knocking out Tbr1 in layer 6 conditional mutant mice.
Supplementary Figure 1. Reduced corticothalamic innervation of the thalamus at P60 in Tbr1 layer 6 CKO.

(A) Injection sites of the anterograde tracer DIO-ChR2-EYFP (green ovals) in the somatosensory cortex of Tbr1\textsuperscript{wildtype} and Tbr1\textsuperscript{layer6} mutants. These lines had the Nttsr1-\textsuperscript{cre::tdTomato\textsuperscript{f/+}} alleles to label the layer 6 neurons and their axons by tdTomato's endogenous fluorescence (red). Numbered lines correspond to the location of each section plane presented in (B). The approximate location of the thalamus is written in orange letters.

(B) Rostro-caudal coronal section series shows of anterograde tracer (green) and tdTomato (from recombinated Ai14 allele; red) in Tbr1\textsuperscript{wildtype} and Tbr1\textsuperscript{layer6} mutants. In both genotypes, tdTomato\textsuperscript{+} corticothalamic axons (white arrowheads) innervate thalamus, although the cKO shows reduced corticothalamic innervation in the medial structures of the Tbr1\textsuperscript{layer6} mutant thalamus. Scale: 100 \(\mu\text{m}\).
Supplementary Figure 2. tdTomato expression in layer 6 neurons and axons shows reduced corticothalamic innervation in $Tbr1^{layer6}$ mutants at P3 and P21.

(A, B) Schematic representation of rostro-caudal coronal section planes (numbered lines) of $Tbr1^{wildtype}$ and $Tbr1^{layer6}$ mutants that were used to investigate the thalamic innervation of layer 6 axons. Prefrontal and somatosensory cortex are indicated in red and blue respectively. The approximate location of the thalamus is written in orange letters. (C, D) Rostro-caudal coronal section series shows tdTomato in $Tbr1^{wildtype}$ and $Tbr1^{layer6}$ mutants at P3 (C) and P21 (D). These lines had the Ntsr1-cre::tdTomato$^{+/+}$ alleles to label the layer 6 neurons and their axons by tdTomato’s endogenous fluorescence (red). The overlap between DAPI (blue) and tdTomato is shown. White arrowheads in panel 5 (at P3 and P21) correspond to the medial structures in the $Tbr1^{layer6}$ mutant thalamus that show reduced corticothalamic innervation. Scale: 100 $\mu$m in (C) and 50 $\mu$m in (D).
Supplementary Figure 3. Loss of Tbr1 in layer 6 reduces excitatory(I) and inhibitory (II) synaptic input onto layer 6 pyramidal neurons in somatosensory cortex at P56.
(A) Schematic representation of somatosensory cortex (S1, blue boxes) that was used during the imaging and patching experiments. (B, C) Schema of layer 6 projection neuron (red) in somatosensory cortex of Tbr1\textsuperscript{wildtype} (B) and Tbr1\textsuperscript{layer6} mutant (C). The rectangles indicate the zone where synapses were analyzed (B, C) at P56. Pipette tip indicates that the soma was patched during the electrophysiology recordings (B, C).

(I) Excitatory synaptic input was analyzed via synaptic bouton staining onto apical dendrites of layer 6 neurons (D-F) and spontaneous EPSC (sEPSC) recordings from the soma of the layer 6 pyramidal neurons (H) of Tbr1\textsuperscript{wildtype} (D) Tbr1\textsuperscript{layer6} heterozygous mutants, and (E) Tbr1\textsuperscript{layer6} homozygous mutants (F) at P56 (D, E, F). These lines had the Ntsr1-cre::tdTomato\textsuperscript{f/+} alleles to label the layer 6 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (D-F) for quantification (D’–F’). (G) Quantification of VGlut1\textsuperscript{+} boutons and PSD95\textsuperscript{+} clusters co-localizing onto the apical dendrites of layer 6 neurons at P56. (H) sEPSC recordings from the soma of layer 6 pyramidal neurons of Tbr1\textsuperscript{wildtype}, Tbr1\textsuperscript{layer6} heterozygous mutants, and Tbr1\textsuperscript{layer6} homozygous mutants at P56. (I) Quantification of the sEPSC occurrence in layer 6 neurons at P56.

(II) Inhibitory synaptic input was examined through synaptic bouton staining onto apical dendrites of layer 6 neurons (J–L) and spontaneous IPSC (sIPSC) recordings from the soma of the layer 6 neurons (N) of Tbr1\textsuperscript{wildtype} (J), Tbr1\textsuperscript{layer6} heterozygote mutants (K), and Tbr1\textsuperscript{layer6} homozygous mutants (L) at P56 (J, K, L). These lines had the Ntsr1-cre::tdTomato\textsuperscript{f/+} alleles to label the layer 6 neurons by tdTomato’s endogenous fluorescence (red). ImageJ software was used to process confocal images (J–L) for quantification (J’–L’). (M) Quantification of VGat\textsuperscript{+} boutons and Gephyrin\textsuperscript{+} clusters co-localizing onto the apical dendrites of layer 6 neurons at P56. (N) sIPSC recordings from the soma of layer 6 pyramidal neurons of Tbr1\textsuperscript{wildtype}, Tbr1\textsuperscript{layer6} heterozygous mutants, and Tbr1\textsuperscript{layer6} homozygous mutants at P56. (O) Quantification of the sIPSC occurrence in layer 6 neurons at P56. Two-way ANOVA was used for the statistical analysis of the control, heterozygous and null. (*p<0.05) (**p<0.01) (***p<0.001) (****p<0.0001). Scale bar (in F and L) = 5µm.
Supplementary Figure 4. *Tbr1* maintains cortical layer 6 identity through regulating the expression of layer 5 and layer 6 markers.

*In situ* hybridization of the latero-medial expression profile of a subset of layer 5 genes (green) and layer 6 genes (red) in the *Tbr1*mutants. The subset of genes was identified by utilizing FACS purified neurons of wildtype and *Tbr1*mutants at P5. Layer 5 markers such as *Cdh8, EphA7, Grin3a, Lypd1, Ntn1, Nrgn, Runx1II*, *Thrb* are upregulated (green), whereas, layer 6 markers including *Cntn2, Nfe2I3* and *Mc4r* are downregulated (red) in the *Tbr1*mutants. Image credit: Allen Institute. Cortical layers are labeled. I = layer 1, II/III = layers 2/3, IV = layer 4, V = layer 5 and VI = layer 6.
Supplementary Figure 5. Intrinsic properties of the pyramidal neurons of $Tbr1^{layer6}$ mutants revealed an increase in HCN channels at P21.
(A) The layer 6 pyramidal neurons were patched at P21 and intrinsic properties were collected from whole-cell patch clamp recordings from layer 6 somatosensory cortex. Recordings from $Tbr1^{\text{wildtype}}$ (red), $Tbr1^{\text{layer6}}$ heterozygous mutant (green) and $Tbr1^{\text{layer6}}$ homozygous mutant (blue) show that many intrinsic electrophysiological properties were unaffected by loss of $Tbr1$, including resting membrane potential (B), input resistance (C), and action potential halfwidth (data not shown). We estimated Ih by measuring the membrane potential sag and rebound elicited by a $-200$ pA current step. (D) sag and rebound is increased in $Tbr1^{\text{layer6}}$ mutants. (**p< 0.01) (***p<0.001).
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CHAPTER V

Concluding Remarks and Remaining Questions
The studies presented here reveal a novel mechanism of dopamine D2 receptor (D2R) signaling unique to a subpopulation of pyramidal neurons within layer 5 (L5) of the prefrontal cortex (PFC). I find this novel phenomenon, specifically an afterdepolarization (ADP), whereby D2R activation enhances cellular excitability in a manner dependent on synaptic inputs, to be mediated by the recruitment of Gs-associated signaling pathways rather than the classically described Gi/o-coupled mechanisms. The phenomenon studied here is one whereby even subthreshold synaptic activity can enhance neuronal excitability for a period of several hundreds of milliseconds or even up to 10 seconds later. Furthermore, I find that this phenomenon is absent in disrupted-in-schizophrenia 1 (DISC1) mouse model of schizophrenia, suggesting behavioral and clinical importance for this novel D2R signaling pathway. Additionally, and for the first time we investigated the role of Tbr1, an Autism Spectrum Disorder (ASD) risk gene, in these deep layer cortical neurons within adult mice. We found distinct molecular, physiological, and behavioral deficits specific to loss of Tbr1 function in layer 5 versus layer 6 suggesting that Tbr1 has diverse functions within different subpopulation of cortical circuits.

Many of the implications and caveats of the results presented here have been discussed individually in Chapters II, III, and IV. Therefore, in this chapter, I will discuss remaining questions, with a focus on larger implications and future directions.

Layer 5 neurons exhibiting this novel D2R-dependent afterdepolarization are well poised to affect the cognitive domains disrupted in schizophrenia and other psychiatric disorders. It is hypothesized that dopamine modulates PFC-dependent cognitive function by transitioning the cortical circuits between a state of stable active neural representations which facilitates maintenance of previously learned behavioral strategies and a destabilized state which promotes flexible behavioral responses important for successful adaptation.
(Durstewitz and Seamans, 2008). The D2R-mediated phenomenon studied here could alter neuronal integration of their inputs over relatively long durations or switch neurons into a “high gain” mode whereby responses to subsequent inputs are potentiated and thereby promote flexibility within the network.

As presented in Chapter III, a dominant negative mutation in human disrupted-in-schizophrenia 1 (DISC1) completely abolished this D2R-mediated phenomena. Should future studies confirm that the D2R-mediated ADP is important for dopaminergic modulation of neuronal modulation in this subpopulation of deep cortical neurons, my findings suggest that the disruption of the D2R-Gs signaling mechanism may disrupt network ability to switch into a flexible state and thereby play a critical role in cognitive deficits associated with schizophrenia.

Many questions do remain regarding the specific molecular mechanisms underlying this phenomenon and its role in normal cognitive function and implications in dysfunction associated with psychiatric disorders. How is the dopamine D2 receptor (D2R) coupled to Gs? Is DISC1, a scaffolding protein, necessary to facilitate this D2R/Gs-associated signaling pathway? D2Rs have been shown to be able to form heteroreceptors, might D2R act via Gs by forming heteroreceptor with another Gs-coupled GPCR in this subpopulation of L5 pyramidal neurons? Current studies have not conclusively ruled out coexpression of dopamine D1 receptors (D1Rs) and D2Rs, nor have the expression patterns of dopamine D4 receptors (D4Rs) and dopamine D5 receptors (D5Rs) been closely examined (Clarkson et al., 2017). There is controversy over whether D2Rs and D1Rs form heteroreceptors in the cortex (Beaulieu and Gainetdinov, 2011; Zhang et al., 2011). It will be interesting for future studies to investigate whether knocking out D1Rs in D2R+ neurons with L5 mPFC would abolish this D2R/Gs mediated phenomenon.
Work from our lab has also shown that the quinpirole-induced ADP is absent in mice younger than 8 weeks. As discussed in Chapter II, activity at the NMDARs is necessary to unmask the quinpirole-induced ADP. It is also known that NMDA receptor (NMDAR) subunit expression shifts over the course of development, at time points similar with the appearance of the quinpirole-induced ADP (Pian et al., 2010; Paoletti et al., 2013). How does the NMDAR facilitate this D2R/Gs-mediated phenomenon? Are these NMDAR subunits important for the unmasking of this D2R-mediated phenomenon? Interestingly, this developmental time point in mice corresponds to the onset of adolescence/young adulthood, similar to the age of onset in schizophrenia (Häfner et al., 1993).

The study presented in Chapter IV, shows that layer specific loss of function of T-brain-1 (Tbr1) leads to upregulation of both D2Rs and hyperpolarization-activated cyclic nucleotide-gated (HCN) channels also resulting in increased in hyperpolarization-activated cation current ($I_h$) mediated by these HCN channels in deep layer cortical neurons. $I_h$ can strongly modulate a neurons spike firing and synaptic potential integration (Shah, 2014). Future behavior experiments investigating the effect of loss of Tbr1 function within these specific cortical layer populations in prefrontal-dependent task such as set-shifting could further elucidate the roles and function of Tbr1 in the adult mouse cortex. Furthermore, it would be interesting to study the activity of these separate neuronal subpopulations during these PFC-dependent tasks.

Interestingly, both of the psychiatric disease models studied in this dissertation, a dominant negative mutation in human disrupted-in-schizophrenia 1 (DISC1) and layer specific loss of function of T-brain-1 (Tbr1), exhibited physiological perturbations impacting neuronal integration. The discovery of this D2R/Gs mediated mechanism provides a new framework in which to investigate the molecular functions of currently-prescribed
antipsychotics, as well as to explore the impact of perturbations in D2R-mediated neuromodulation on prefrontal cells and circuits. An improved understanding of how mechanisms of dopaminergic modulation are perturbed in psychiatric disease may thus aid in the development of safer and more efficient drug therapies.

Final thoughts

This work establishes a new understanding of D2R signaling within the prefrontal cortex, complicating but potentially resolving years of discrepancy in the field. It will be important to identify specific details of the molecular pathway underlying this effect and its impact on cortical circuit function and behavior. Elucidating these mechanisms has potential implications for psychiatric disease and could illuminate novel drug therapies or circuit-based approaches for understanding and treating prefrontal-related cognitive dysfunctions in psychiatric disease.
References


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