Review

Contributions of ERK signaling in the striatum to instrumental learning and performance

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\section*{A B S T R A C T}

The striatum is critical for learning and decision making; however, the molecular mechanisms that govern striatum function are not fully understood. The extracellular signal regulated kinase (ERK) cascade is an important signaling pathway that underlies synaptic plasticity, cellular excitability, learning and arousal. This review focuses on the role of ERK signaling in striatum function. ERK is activated in the striatum by coordinated dopamine and glutamate receptor signaling, where it underlies corticostriatal synaptic plasticity and influences striatal cell excitability. ERK activation in the dorsal striatum is necessary for action-outcome learning and performance of goal-directed actions. In the ventral striatum, ERK is necessary for the motivating effects of reward-associated stimuli on instrumental performance. Dysregulation of ERK signaling in the striatum by repeated drug exposure contributes to the development of addictive behavior. These results highlight the importance of ERK signaling in the striatum as a critical substrate for learning and as a regulator of ongoing behavior. Furthermore, they suggest that ERK may be a suitable target for therapeutics to treat disorders of learning and decision making that arise from compromised striatum function.

\section*{1. Introduction}

The striatum, the largest of the basal ganglia nuclei, is a critical substrate for learning and decision making. Impairments in learning and decision making accompany a range of disorders that affect the striatum, including substance abuse disorder, obsessive-compulsive disorder, and Parkinson's disease\cite{1–3}. Furthermore, the application of sophisticated procedures derived from instrumental conditioning in animals has identified parallel corticostriatal circuits that mediate distinct learning and action control processes\cite{4–8}. Nevertheless, the molecular mechanisms that underlie learning and decision making in the striatum are not fully understood. One important regulator of neuronal function is the extracellular signal regulated kinase (ERK) pathway. ERK is a member of mitogen activated protein kinase (MAPK) family, and is critical for nervous system development and plasticity in the adult.

Abbreviations: ERK, extracellular signal regulated kinase; LTP, long-term potentiation; LTD, long-term depression; PIT, Pavlovian-instrumental transfer; pDMS, posterior dorsomedial striatum; DLS, dorsolateral striatum; NAc, nucleus accumbens.

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nervous system [9,10]. ERK signaling in the nervous system has a critical role in memory formation, affect and arousal. Furthermore, the efficacy of a number of psychoactive substances, such as mood stabilizers and addictive drugs, depend, in part, on their ability to activate ERK in the nervous system.

In the following review, we highlight the role of ERK signaling in the striatum in learning and decision making. We begin with a brief overview of ERK signaling in the nervous system and its role in learning and behavior regulation. This is followed by a discussion of the role of ERK signaling in striatal-based learning and decision-making tasks. We conclude with a description of the role of ERK signaling in substance abuse, in which we discuss how alterations in ERK signaling by repeated drug exposure may compromise striatum function and produce features of addictive behavior.

2. The ERK signaling pathway

Like other MAPK signaling pathways, the ERK cascade consists of three kinases: ERK, of which there are two isoforms, ERK1 and ERK2, and their upstream kinases MEK and raf (see [11,12] for review). ERK activation occurs in response to a variety of extracellular stimuli and forms an essential pathway for cells to generate adaptive responses to changing environments. These cellular responses include regulation of gene expression and synthesis of new proteins, alteration in cellular structure or metabolism, cellular growth, differentiation and apoptosis. ERK is activated when dual-phosphorylated by MEK at its serine and threonine residues. Raf, which phosphorylates MEK, consists of a family of kinases, and is activated when bound by the GTP-binding protein ras. Ras-ERK activity increases in response to a range of stimuli, including activation of G protein coupled receptors, tyrosine kinase receptors, and Ca²⁺ influx [9,13]. Scaffolding proteins and phosphatases regulate ERK activity, thus providing an additional level of control over ERK function [11]. Upon activation, ERK has many cellular targets and influences a wide range of cellular functions [12]. ERK influences gene expression through its interaction with transcriptional regulators, such as ribosomal S6 kinase (RSK), mitogen- and stress-activated protein kinase-1 (MSK1) as well as the transcription factor elk-1. ERK also influences translation and new protein synthesis [14]. These and many other cellular operations under ERK’s regulation enable the cell to produce a coordinated response to extracellular stimuli.

In the adult nervous system, ERK is intimately involved in synaptic plasticity (see [15] for review). ERK inhibition prevents the induction of various forms of long-term potentiation (LTP) and long-term depression (LTD) in the hippocampus and amygdala [16–20]. ERK activation is necessary for a number of physiological processes that underlie synaptic plasticity, including AMPA receptor trafficking and spine restructuring [21–24]. Among the ways ERK influences synaptic plasticity is through its regulation of transcription and translation [14,25,26]. Finally, the effects of various neuromodulators on LTP and LTD induction and maintenance depend on ERK signaling [26–30]. Consistent with its role in synaptic plasticity, ERK has been shown to be necessary for long-term memory formation across a variety of tasks. An increase in ERK activation, measured as the ratio of phosphorylated to total (phosphorylated and non-phosphorylated) ERK, occurs in a number of brain regions during learning. Furthermore, treatments that interfere with ERK signaling such as the MEK/ERK inhibitors SL327, U0126 or PD98059, impair long-term memory retention. ERK inhibition prevents the formation of lasting memories of an event or association, including spatial memory, fear memory, and object recognition memory [31–39]. More recently, genetic approaches have been used to examine ERK function and have yielded similar effects on behavior. ERK2 conditional knockouts show impaired long-term retention in spatial memory and fear conditioning tasks [14,40]. Based on these data, ERK activation is necessary for the establishment of long-lasting changes in behavior, likely by mediating changes in synaptic strength.

In addition to its role in synaptic plasticity, ERK activation in the nervous system is important for plasticity of cellular intrinsic excitability. This form of plasticity involves changes in the electrical properties of the cell membrane that render neurons more or less responsive to synaptic inputs [41]. In particular, A-type K⁺ channels located in distal dendrites are an important determinant of cellular excitability by limiting dendritic action potential back-propagation and raising action potential thresholds (see [41,42] for review). Recently it has been shown that the inwardly rectifying K⁺ channel Kᵢ,4.2 is regulated by ERK [see [43] for review]. ERK phosphorylates the pore-forming unit of the Kᵢ,4.2 channel, which results in down regulation of this channel’s activity [44–46]. The net effect of this modulation is to increase cellular excitability and facilitate LTP induction in neurons with elevated ERK phosphorylation [46,47]. Thus, Kᵢ,4.2 channel phosphorylation and its effect on membrane electrical properties enable ERK activation to increase cellular excitability and potentially alter cellular information processing and ongoing behavior.

Alterations in cellular excitability have been proposed to have a role in behavior regulation [48,49]. In this context, ERK’s regulation of cellular excitability may underlie its increasingly recognized role in regulating emotional arousal and affective states. For example, inhibition of ERK activation with the systemic ERK inhibitor SL327 reduces the amount of time rats swim in the forced swim test, which suggests that ERK inhibition has a depressive effect on behavior [50,51]. Likewise, knockout of ERK1 increases basal ERK2 activation and increases measures of behavioral arousal, an effect that is reversed by ERK inhibition [52,53]. These studies suggest a link between altered levels of basal ERK activation and mood disorders, such as mania and depression. This is supported by findings that chronic exposure to the stress hormone corticosterone produces a depressive phenotype in rats, reduces basal levels of ERK activation in the hippocampus and reduces performance on measures of motivation and arousal. Exposure to anti-depressants reverses the behavioral effects of corticosterone exposure and restores ERK phosphorylation levels in the hippocampus [54,55]. One explanation for these behavioral effects is that ERK activation modulates affect and arousal through its effects on cellular excitability. This would explain why ERK inhibition acutely disrupts behavior on these tasks, since it is unlikely that such rapid effects of ERK inhibition could be mediated by alterations in gene expression. Recent evidence showing that olfactory discrimination learning relies on increased basal ERK phosphorylation and changes in cellular intrinsic excitability in the piriform cortex supports this conclusion [56]. Of course, it is unlikely that such a general behavioral measure as arousal is mediated solely by the excitability of neurons in a single brain region. Additionally, changes in basal ERK activation not only influence intrinsic excitability, but also cause structural changes in neurons that are responsible for phenotypes characteristic of mood disorders such as mania and depression [57]. Nevertheless, these results highlight the importance of basal ERK signaling in regulation of arousal and affective states and its impact on ongoing behavior.

2.1. Summary

ERK is one of the primary cellular signal transduction pathways in the nervous system, and confers adult neurons with plasticity that is crucial for learning and adaptive behavior. ERK enables synaptic plasticity through its regulation of receptor trafficking, transcription and translation, operations that are vital for the formation of lasting memories. ERK enables plasticity of cellular
Excitability through its phosphorylation of Kv4.2 channels, which, unlike its effects on gene expression, allows ERK activity to rapidly modulate neural activity and behavior. These two components of ERK signaling likely work in parallel to simultaneously confer long-lasting experience-dependent changes in behavior as well as adaptability to changing environmental demands. In the following section, these themes will reappear when considering the role of ERK signaling in the striatum.

3. ERK signaling in the striatum

3.1. ERK signaling influences striatal synaptic plasticity and excitability

The striatum is the largest of the basal ganglia nuclei and serves as the primary input structure for basal ganglia-thalamo-cortical circuitry. The striatum receives excitatory glutamatergic inputs from cortical, limbic and thalamic regions, as well as dopaminergic input from the midbrain [58-60]. Dopaminergic input to the striatum accounts for the majority of dopaminergic signaling in the brain. Glutamatergic and dopaminergic synapses form in close proximity on striatal medium spiny neurons (MSN’s) [59]. Dopamine (DA) and glutamate signaling interact in important ways to influence synaptic plasticity at corticostriatal synapses (see [60-62] for review). In particular, the induction of corticostriatal LTP depends on the combined activation of post-synaptic D1 and NMDA receptors [63,64].

One consequence of combined D1 and NMDA receptor activation in striatal MSN’s is activation of ERK, and indeed ERK activation is necessary for corticostriatal synaptic plasticity [65-67]. This pattern of activation is the result of ERK’s complex regulation by phosphatase cascades and the phosphoprotein DARPP-32 [65,68,69] (Fig. 1). In striatal neurons, DA D1 receptor activation increases cAMP-PKA activity and phosphorylation by PKA of DARPP-32 at its Thr–34 residue. Thr–34 DARPP-32 becomes a potent inhibitor of a phosphatase cascade that targets ERK, thus freeing ERK from inhibition. NMDA signaling also activates ERK in striatal neurons via calcium-dependent activation of ras [62,70–72]. However, glutamate negatively regulates ERK signaling through calcium-mediated phosphorylation of DARPP-32 at Thr–75 and dephosphorylation of DARPP-32 at Thr–34, which disinhibits phosphatases that dephosphorylate ERK [73]. Thus, ERK appears to be optimally activated in the healthy striatum under conditions of combined NMDA and DA receptor activation. ERK likely enables corticostriatal plasticity, in part, through regulation of transcription factors such as CREM response element binding (CREB) protein, as disruption of CREB signaling in the striatum prevents striatal LTP and LTD induction [74].

In addition to its role in corticostriatal synaptic plasticity, ERK appears to be involved in regulating striatal cell excitability. MSN’s exhibit a hyperpolarized resting membrane potential known as the down state, which is characterized by high input resistance (see [62,75] for review). Coordinated glutamatergic input shifts MSNs to a less hyperpolarized “up-state”. During the up-state, D1 receptor activation increases cell excitability and facilitates glutamatergic signaling [76–79]. Striatal neurons contain Kv4.2 channels, which play an important role during striatal up-states to limit MSN polarization [80,81]. ERK phosphorylates Kv4.2 in striatal neurons, which down-regulates these channels, and ultimately increases cell excitability during up-states and promotes corticostriatal synaptic plasticity [82]. ERK’s ability to modulate MSN excitability during up-states through Kv4.2 phosphorylation may enable it to rapidly modulate striatal activity and behavior.

3.2. Striatal ERK signaling and instrumental learning

One of the clearest examples of striatal involvement in learning and control of actions is found in instrumental learning. During instrumental learning animals learn, in part, to associate their actions with specific outcomes or consequences. If outcomes are rewarding (or involve the removal of an aversive stimulus) they serve as positive reinforcers, whereas aversive outcomes (or those that involve removing an appetitive stimulus) act as negative reinforcers. During instrumental learning, animals form associations between their actions (such as pressing a lever in an operant chamber) and the specific consequences of those actions (such as delivery of food pellets or termination of an aversive footshock). The striatum is critical for this form of learning, and models of reinforcement learning have emphasized the importance of glutamate–DA interactions in the striatum during instrumental action-outcome learning [83-85]. Specifically, DA release in the dorsal striatum is hypothesized to underlie the encoding of action-outcome associations by enabling potentiation of corticostriatal synapses during learning [86]. In support of this hypothesis, inhibition of glutamate and DA receptors in the dorsal striatum interferes with action-outcome encoding [87,88].

Given the importance of glutamate and DA signaling in the striatum during instrumental learning, ERK signaling may be particularly relevant for this process, given that its activation in striatum depends on combined DA and NMDA receptor stimulation, whereas when these receptors are stimulated separately they weakly activate ERK [65]. Indeed, the requirement for dual-activation of NMDA and D1 receptors in order to activate ERK has lead to the proposal that ERK functions to selectively induce plasticity at corticostriatal neurons during highly relevant learning situations when coordinated DA and NMDA receptor activation is likely to occur, such as during reward learning [65,89]. Additionally, this selectivity would prevent arbitrary associations from being formed between actions and irrelevant outcomes.

Recent evidence supports the conclusion that ERK is important for instrumental action-outcome learning in the striatum. In
order to demonstrate action-outcome (or goal-directed) learning in instrumental conditioning, an animal must be shown to possess some knowledge of the outcome resulting from its actions and use this information to select among potential actions ([4,90] for review). One method of demonstrating this knowledge is through an outcome devaluation paradigm. In this paradigm the goal or outcome of an action is manipulated, with the expectation that the animal's behavior will adjust to changes in goal value. Using these procedures in rodents, action-outcome learning has been shown to rely on a posterior portion of the dorsomedial striatum (pDMS) [91]. Although still capable of performing instrumental responses, inactivation of the pDMS prior to instrumental learning produces outcome-insensitive responding following devaluation in a subsequent test conducted in extinction, thereby demonstrating that the pDMS has a critical role in action-outcome encoding.

We recently demonstrated that ERK signaling in the pDMS plays an important role in action-outcome learning [92]. We found that when rats are trained to press a lever for a food outcome, this experience will increase ERK phosphorylation in the pDMS compared to yoked controls that experience the same pattern of outcomes, but whose actions have no casual relation to outcome delivery. This suggests that ERK activation in the pDMS is necessary for action-outcome learning. To test this suggestion, we trained rats in several sessions to perform actions (left and right lever press) for a common outcome (20% polycose solution). Subsequently, in a single session, rats learned that the left and right lever responses delivered unique outcomes (sucrose pellets and 20% sucrose solution). Prior to learning these new action-outcome associations, rats received intra-pDMS U0126 infusions to interfere with ERK activation. We then tested the rats’ knowledge of action-outcome associations, by devaluing one of the two outcomes prior to a test in which rats could choose between the lever that produced the now devalued outcome or the still valued outcome. Control animals showed a significant devaluation effect, making fewer responses on the action that previously produced the now devalued outcome. In contrast, rats treated with U0126 showed no devaluation effect, responding equally on the valued and devalued lever, indicating that intra-pDMS U0126 prevented new action-outcome learning.

In addition to learning about the consequences of their actions, animals performing an instrumental task may also develop habitual responding, and indeed, this form of learning likely occurs in parallel and largely independent from action-outcome learning. Habitual behavior typically develops more slowly than goal-directed learning and is the product of extensive training. Once established, habitual responses are automatically elicited and performed without consideration of the outcome or consequence of their execution. Habitual responding is thought to be supported by stimulus-response associations. It has shown that habit formation in rodents depends on the dorsolateral striatum (DLS), an area that is richly innervated by sensorimotor areas of the frontal cortex. Lesions of this structure prevent the development of habitual responding, and inactivation after training renders behavior to an outcome-sensitive state [93,94].

Acquisition of habitual behavior depends on DA innervation of the DLS, suggesting that ERK may be involved in this form of learning [95]. However, it is not known whether the acquisition of habitual responding depends on ERK activation. It has been shown that ERK signaling in the dorsal striatum is necessary for the consolidation of a complex motor skill [96]. Inhibition of ERK activation in the dorsal striatum prevented improvement on the accelerating rotarod task following initial acquisition, suggesting that ERK signaling in the striatum has a role in the consolidation of this form of motor learning. Whether a similar process occurs in the development of habitual responding for rewards is not known.

Based on the findings reviewed above, ERK signaling in the striatum is necessary for instrumental learning: in the pDMS, ERK is necessary for action-outcome encoding, whereas in the DLS it may be necessary for the development of habitual behavior (Table 1). ERK is likely involved in these forms of learning through its role in corticostriatal plasticity. We propose that, during learning, ERK is “transiently” activated in the striatum (i.e., ERK activation increases during learning and returns to baseline following a learning episode) and engages transcriptional and translational mechanisms to enable corticostriatal plasticity and memory formation. In the DMS, action-outcome learning likely involves LTP, which is the dominant form of plasticity in this region, and where ERK has a well-described role in LTP induction. The role of synaptic plasticity in the DLS during habit formation is less clear, but may involve corticostriatal LTD. Although a requirement for ERK in corticostriatal LTD has not yet been demonstrated, ERK is necessary for LTD in other brain areas [20].

### Table 1

<table>
<thead>
<tr>
<th>Process</th>
<th>Dorsomedial striatum</th>
<th>Dorsolateral striatum</th>
<th>Nucleus accumbens</th>
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<tbody>
<tr>
<td>Learning</td>
<td>A-O learning</td>
<td>Habit formation?</td>
<td>S-O learning?</td>
</tr>
<tr>
<td>Performance</td>
<td>Goal-directed action</td>
<td>Habit performance?</td>
<td>PIT</td>
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**Striatal ERK signaling and instrumental performance**

In addition to a role in instrumental learning, the striatum is necessary for the performance of instrumental actions. As is the case with learning, instrumental control relies on parallel circuits involving the pDMS and DLS, which mediate performance of goal-directed and habitual action, respectively. Inactivation of the pDMS produces outcome-insensitive (i.e., habitual) instrumental behavior in rats that ordinarily would display outcome-sensitive responding, whereas inactivation of the DLS produces outcome-sensitive (i.e., goal-directed) instrumental behavior in rats that ordinarily would display outcome-insensitive responding [91,94]. These findings support the notion that parallel corticostriatal circuits mediate goal-directed and habitual action.

We recently demonstrated that ERK activation in the pDMS is necessary for the performance of goal-directed actions. We trained rats over several sessions to perform instrumental actions for a rewarding sucrose solution. We found an increase in ERK activation in the pDMS and DLS following the final training session [92]. To test the involvement of ERK in goal-directed behavior performance, we trained a separate set of rats to associate left and right lever presses with unique outcomes, after which one outcome was devalued and performance on each lever was assessed in an extinction test. Prior to the extinction test, rats received U0126 infusions into the pDMS or DLS. Rats that received U0126 infusions into the pDMS showed no preference for the valued action compared to the devalued action in a choice test. ERK inhibition had no effect on rats’ ability to make instrumental responses generally, nor did it interfere with the palatability of the food outcomes. Rather, it appears that inhibition of ERK activation in the pDMS prevents the use of outcome value to guide action selection. Interestingly, ERK inactivation in the DLS also interfered with goal-directed responding, suggesting that the DLS may also play a role in the performance of goal-directed actions.

In addition to outcome value, instrumental actions are also sensitive to cues associated with rewarding outcomes. Animals learn through Pavlovian conditioning to associate stimuli with outcomes or events, and these associations can affect instrumental responding by influencing the choice and vigor of instrumental actions. This process is known as Pavlovian-instrumental transfer (PIT). During
a typical PIT paradigm, animals are first trained under Pavlovian and instrumental procedures to separately associate a conditioned stimulus (CS) and an action with outcome delivery. During the PIT test, the CS is presented while animals have access to the instrumental levers, and typically CS presentation will cause animals to increase their instrumental performance compared to a neutral stimulus or baseline. This effect is thought to reflect the motivating or arousing state that CS presentations elicit.

PIT depends on the nucleus accumbens (NAc), a region within the ventral striatum: lesions of this structure do not prevent the acquisition or performance of instrumental responding; however, the motivating effect of CS presentation on instrumental performance is abolished in animals with NAc lesions [97,98]. We found that ERK signaling in the NAc has an important role in reward-seeking behavior, where it appears to be critically involved in PIT [99]. We trained rats over several sessions to associate a 2-min tone stimulus with delivery of food outcomes. In a subsequent test in which the tone was presented in extinction, we found that the tone stimulus increased ERK phosphorylation in the NAc when compared to control animals that were presented with a neutral stimulus. CS presentation also increased CREB phosphorylation in the NAc, suggesting that Pavlovian conditioning produces synaptic plasticity in this structure [100]. To examine the role of CS-evoked ERK activation in the NAc, we trained rats under Pavlovian and instrumental conditioning and assessed their performance in a PIT task. We found that intra-NAc U0126 infusion prior to the PIT task prevented the CS from elevating instrumental responding. ERK inhibition had no effect on instrumental responding generally, which suggests that ERK inhibition has a specific role in enabling cues to motivate instrumental behavior.

Based on these studies, ERK signaling in the striatum appears to be necessary for goal-directed learning as well as the modulation of acquired instrumental actions by outcome value or by cues associated with a rewarding outcome. We propose that ERK activation in the striatum influences performance of instrumental actions by modulating the intrinsic excitability of striatal neurons. As described above, ERK phosphorylates Kv4.2 channels, which increases cellular excitability in striatal neurons. The net effect of this modulation may be to allow synaptic inputs to the striatum representing appetitive stimuli (such as from the BLA [101]) to increase their control of striatal cellular activity, and as a consequence engage reward-seeking behavior. It is not known whether ERK acts as a permissive or instructive signal for modulating instrumental performance. Some evidence suggests that the level of ERK activation evoked during task performance may determine the magnitude of the behavioral effects mediated by ERK [102]. The level of ERK activation in the striatum may therefore act as a modulator of decision processes, with greater ERK activation evoked by a stimulus or event facilitating action initiation [103].

4. ERK and addiction

Exposure to a variety of substances with abuse potential, including psychostimulants, opioids and nicotine, acutely activate ERK in the striatum and other brain areas [104,105]. Many enduring behavioral effects of acute drug exposure depend on ERK signaling. For example, pairing a context with drug exposure produces a conditioned place preference (CPP), and ERK inhibition prior to drug-context pairing prevents the formation of this preference [105,106]. Since ERK does not prevent the hyperlocomotion induced by drug exposure, it is likely that the effect of ERK inhibition on CPP is the result of impaired memory formation [107,108]. ERK is also required for the development of locomotor sensitization, a process that occurs when repeated drug exposure heightens locomotor responses to subsequent drug presentations. ERK inhibition prior to drug exposure prevents the development of locomotor sensitization [109]. Similarly, over-expression of ERK2 increases locomotor sensitization and CPP [67,110]. These findings suggest that ERK is required for the neuroadaptive changes resulting from repeated drug exposure, and which are responsible for heightened sensitivity to the drug’s acute locomotor effects. Interestingly, ERK is also necessary for the expression of CPP [108,109]. Furthermore, the ability of stimuli to elicit ERK activation may itself become sensitized by repeated drug exposure, and this may also contribute to heightened relapse to drug-associated cues. Supporting this notion are findings that ERK is involved in the incubation of craving, in which responses to drug-associated stimuli increases with the withdrawal period [111,112]. Recent studies have shown that drug-associated cues elicit ERK activation in the central nucleus of the amygdala, and after 30 d of withdrawal this cue-driven ERK response is greater than after 1 d of withdrawal, mirroring the effects of these cues on relapse. Similarly, chronic food restriction increases sensitivity to drugs of abuse, and this is associated with enhanced striatal activation of ERK by D1 and NMDA receptor stimulation [113]. Taken together, these results parallel findings from instrumental tasks that ERK is necessary for learning and may also directly influence the expression of reward-seeking behaviors.

Although a single drug exposure acutely affects behavior, repeated drug administration produces a variety of changes in behavior that characterize addiction. Among the many physiological changes produced by repeated drug exposure is a sustained increase in baseline ERK activation in the brain, including in the striatum [114–116]. This change in baseline ERK activation has been observed in the NAc and accompanies changes in NAc excitability that results from repeated drug exposure, a process that depends on AMPA receptor insertion [116,117]. These findings suggest that ERK is dynamically regulated following repeated drug exposure and withdrawal, that changes in ERK activation directly influence striatal cell excitability, and that these effects may be responsible for the expression of addictive behavior. Based on the role that we have described for ERK signaling in striatum function, sustained ERK activation may promote excessive action initiation, consequently giving rise to impairments in action control. Excessive excitability in the pDMS may give rise to compulsive behavior as actions are repeatedly performed, whereas excessive excitability in the DLS may give rise to impulsive behavior as stimuli repeatedly elicit actions (Fig. 2). Such phenotypes have been observed in rodents following repeated drug exposure [118–120]. Within the NAc, excessive excitability may enhance the arousing effects of reward-associated cues on reward-seeking behavior, an effect that has been documented following repeated amphetamine exposure.
in rodents [121]. It is not known to what extent these behavioral patterns are the result of drug-induced alterations in striatal ERK signaling.

5. Conclusions

The role of ERK in addictive behavior in the striatum is consistent with its role in action-outcome learning and control of behavior by reward-associated stimuli more generally. This is not surprising given that drug-taking is a form of appetitive behavior. What is different about abused substances is that they activate ERK more strongly than natural rewards and that with repeated exposure they increase basal levels of ERK activation. These properties may be responsible for some features of addictive behavior. Indeed, the effects of abused substances, like the effects of mood regulators on ERK, show how up or down-regulation of baseline ERK activation influences behavior. In the context of ERK’s role in striatum function, we propose that changes in baseline activity give rise to impulsive or cue-driven behavior. By understanding ERK function in the striatum during processing of natural rewards, we may begin to decipher how dysregulation of ERK signaling gives rise to disorders such as addiction.

It is clear that striatal ERK mediates learning and decision processes; however, ERK is not necessary for the generation of motor action or for the processing of hedonics. This specialization in learning and decision making may be the consequence of ERK’s unique regulation in striatum: ERK is maximally activated under conditions of coordinated D1 and NMDA receptor signaling. This functional specialization suggests that ERK could serve as a target for therapeutics to treat disorders of learning and decision making (e.g., to prevent relapse) and spare other striatal functions. There is still much to be known about ERK’s regulation in the nervous system and the mechanisms through which it influences the structure and physiology of striatal neurons. Nevertheless, these data suggest that ERK is a promising target for treating disorders of learning and decision making that arise from compromised striatal function.

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