Title:  Network failures: when incentives trigger impulsive responses

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**Abstract**

Adequate control of impulsive urges to act is demanded in everyday life but is impaired in neuropsychiatric conditions such as stimulant use disorder. Despite intensive research it remains unclear whether failures in impulse control are caused by impaired suppression of behavior or by the over invigoration of behavior by stimuli associated with salient incentives such as drugs, food and money. We investigated failures in impulse control using functional magnetic resonance imaging (fMRI) to map the neural correlates of premature (impulsive) responses during the anticipation phase of the Monetary Incentive Delay (MID) task in healthy controls (HC), stimulant dependent individuals (SDIs), and their unaffected first-degree siblings (SIB). We combined task-based fMRI analyses with dynamic causal modeling to show that failures of impulse control were associated with interactions between cingulo-opercular and dorsal striatal networks regardless of group status and incentive type. We further report that group-specific incentive salience plays a critical role in modulating impulsivity in SDIs since drug-related incentives specifically increased premature responding and shifted task modulation away from the dorsal striatal network to the cingulo-opercular network. Our findings thus indicate that impulsive actions are elicited by salient personally-relevant incentive stimuli and that such slips of action recruit a distinct fronto-striatal network.

**MAIN DOCUMENT**

**Introduction**

Behavior is often driven by the motivational properties of rewards, yet the interactions between motivation and cognitive control and their neural substrates are unclear [Botvinick and Braver, 2015; Hull, 1943]. In the Principles of Psychology, William James stated that motivated behaviors “are but results of the fact that certain things appeal to primitive and instinctive impulses of our nature, and that we follow their destinies with an excitement that owes nothing to a reflective source” [James, 1890] suggesting that motivated behavior can be accompanied by weakened “reflective” cognitive control over one’s impulses.

Impulsivity has been linked to numerous neuropsychiatric disorders and, as a multifaceted construct, incorporates a range of different traits and behaviors [Dalley et al., 2011; Evenden, 1999]. Key aspects of impulsivity include risky decision making and impulsive action [Dalley and Robbins, 2017]. Impulsive action is typically gauged by assessing the tendency to act prematurely without foresight or sufficient regard to negative consequences, or by assessing the ability to countermand or stop a prepotent response [Robbins et al., 2012]. Evidence from humans and rodents suggest the two complementary forms of impulsive action rely on overlapping, yet distinct, neural networks and neurochemical substrates [Dalley and Robbins, 2017]. While stopping prepotent responses is associated with dorsal striatal (dStriatum) along with ventrolateral and dorsomedial prefrontal (PFC) involvement [Swick et al., 2011], premature responding appears to involve the ventral striatum and ventromedial PFC [Dalley et al., 2011].

The inability to suppress inappropriate responses is believed to play a key role in stimulant drug addiction both as a vulnerability factor and as a consequence of chronic drug use [Goldstein and Volkow, 2011]. This can manifest in the day-to-day lives of drug users as difficulties in suppressing excessive approach behaviors and urges to act, particularly when exposed to incentivizing drug-related cues. Converging evidence shows that chronic stimulant drug use is associated with response inhibition impairments and top-down cognitive control abnormalities underpinned by aberrant fronto-striatal function [Morein-Zamir and Robbins, 2015]. Unaffected siblings of stimulant dependent individuals (SDI) also show response inhibition difficulties and associated structural brain abnormalities suggesting this may be a preexisting vulnerability factor [Ersche et al., 2013]. Whilst decisional impulsivity is often investigated in incentivized contexts [Kjome et al., 2010; Vigil-Colet, 2007], impulsive actions in humans have largely been investigated by assessing stopping or cancelling prepotent responses in non-incentivized contexts [Bari and Robbins, 2013].

Findings on the neural correlates of premature responding in incentivized contexts have largely stemmed from rodent research [Dalley et al., 2008; Eagle and Baunez, 2010]. Here, premature responses gauge difficulties in suppressing responses by capturing the inability to resist responding until a waiting interval has elapsed. Premature responding predicts the transition to compulsive cocaine-taking in rodents, suggesting it could also be a vulnerability factor [Belin et al., 2008]. Recent studies using a specialized paradigm adapted from the animal literature have begun to make inroads, demonstrating increased premature responding in abstinent SDIs [Voon, 2014]. Greater premature responding has also been reported for tobacco smokers, cannabis users and binge drinkers [Mechelmans et al., 2017; Morris et al., 2016; Sanchez-Roige et al., 2014; Voon et al., 2016] reinforcing the importance of this measure to addiction more broadly.

As noted above, impulsive actions are clearly intertwined with reward processing. Empirical research into mechanisms of reward has proceeded largely in parallel. This research has pointed to aberrant generalized reward processing in addiction, largely as a consequence of drug use [Balodis and Potenza, 2015; Cope et al., 2019; Koob and Moal, 2005]. For example, mesocortico-limbic abnormalities are believed to underscore exaggerated incentive salience to drug-associated stimuli [Berridge, 2007]. Thus, the process of ‘wanting’ triggered when faced with drug-related cues yields upregulation of reward-related regions [Berridge, 2007]. In humans, the widely-used monetary incentive delay (MID) task has been employed to assess reward related processing in addiction. Anticipating monetary rewards in this task has been associated with robust ventral striatum in addition to dorsal striatum and vlPFC activations [Oldham et al., 2018]. Contrary to expectations, drug users including SDIs do not appear to show consistent abnormalities in cross-sectional MID studies [Balodis and Potenza, 2015; Just et al., 2019]. One possible reason for this may be the use of monetary incentives in these studies. Addiction is associated with blunted brain response in a wide array of non-drug-related tasks, but with increased engagement of brain networks during exposure to drug cues or drug-related incentives [Zilverstand et al., 2018]. This is underscored in the theory of impaired response inhibition and salience attribution (iRISA) which points to the pivotal role of context and incentive type [Goldstein and Volkow, 2002; Goldstein and Volkow, 2011].

The current study investigates the neural correlates of premature responses in stimulant users and their unaffected siblings and healthy controls in different incentive conditions using functional Magnetic Resonance Imaging (fMRI). We focus on premature responding in the MID task [Peña-Oliver et al., 2016], introducing a novel analysis approach. Premature responses are akin to the everyday maladaptive behaviors exhibited by these individuals and to our knowledge their neural underpinnings using fMRI have yet been directly explored. By including two distinct contexts, one providing monetary cues as incentives and the other drug-related cues as incentives, we provide a direct empirical test of the iRISA model predictions. We predicted drug related cues to elicit increased impulsive behaviour and increased activation in fronto-striatal networks. Assessing the SIB allowed us to investigate whether excessive premature responding and corresponding brain activations constitutes a vulnerability factor preceding drug use [Just et al., 2019]. Given the importance of the striatum to top-down control and reward processing in addiction and its abnormal neuronal connectivity with the PFC [Ma et al., 2014; Ma et al., 2015], we also investigated the selective involvement of this region and its effective connectivity using dynamic causal modelling (DCM) within key nodes of the PFC.

**Methods**

**Participants.** Participants were recruited for this study by advertisements, by word of mouth, and from local treatment services (see Table 1 for demographic information). Recruitment and screening procedures have been described in detail elsewhere [Ersche et al., 2012; Just et al., 2019]. Briefly, three groups consisted of SDIs who met Diagnostic Statistical Manual (DSM-IV-TR) criteria for cocaine or amphetamine dependence, their biological siblings who had no history of substance dependence except nicotine, and healthy individuals without familial risk with no drug history. Urine screen results were positive for all but three SDI and negative for all other participants. The study was approved by the NHS Cambridge Research Ethics Committee (08/H0308/310) and all participants provided written informed consent. Data from these individuals as part of a larger sample have been published previously [Just et al., 2019]. This study introduces novel approach to evaluate premature responses in the MID task taking advantage of the incentivized setting. Additional inclusion criteria here were that participants exhibit at least one premature response in the MID task in each context.

**MID Task.** The task consisted of money and drug incentive blocks, with the two counterbalanced across subjects. While the stimuli displayed differed between the two contexts, timings and task structure were the same. Each incentive block consisted of 66 trials, of which 22 were neutral. Trials began with a cue (lasting 250 msec) signaling the reward. For money incentives, a circle with two, one or no horizonal lines indicated a possible win of 50 pence (large reward), 10 pence (small reward) or 0 pence (neutral), respectively. For drug incentives, images of cocaine, crack, IV or non-IV drugs (white powder) in commonly taken form, or a bottle of water as neutral, were used to signal the expected reward (see Figure 1A). Following cue presentation, participants awaited a target cue for an anticipation period lasting 3000-5000 msec. Subsequently, a white square target was presented, lasting 100-400 msec, with participants required to press a key while the target was still on screen. Target duration was titrated to maintain a 66% success rate. Target presentation was followed by a feedback message (1650 msec). For money incentives, successful responding was followed by a message “you’ve won 10p/50p” along with the respective coin. Neutral and unsuccessful trials were followed by a message “you’ve won 0 p” accompanied by a white circle. Participants received monetary rewards conditional on their performance in the monetary block at the end of the task. For drug incentives, successful responding was followed by an image of a person taking cocaine, crack, IV or non-IV drugs for the reward conditions, or a person drinking from a clear water bottle for the neutral or unsuccessful trials. An inter-trial interval lasted 2700-5700 msec, during which a fixation cross was displayed. Participants first completed 66 practice trials prior to scanning with both incentive types.

**Neuroimaging Acquisition.** Images were collected on a Siemens TIM Trio 3-Tesla scanner (Erlangen, Germany) using whole-brain echo planar images for functional data and T1 images for high resolution structural data. The following parameters were applied for the functional scan: repetition time =2000ms; echo time=30ms, flip angle=78°; 32 slices with a thickness of 3mm plus a 0.75mm gap; matrix=64x64 field of view=192 x 192 mm with an in-plane resolution of 3x3mm. T1 scans were acquired using: TR=2300ms; TE=2.98ms; TI=900ms; flip angle=9°; FOV=240x256mm, resulting in 176 slices of 1 mm thickness.

**Data Analysis.**

***Task performance.*** Analyses contrasted the number of premature responses between groups, for the money and drug contexts. As data were positively skewed and deviated significantly from the normal distribution (Kolmogorov-Smirnov tests, p<0.05), one-way nonparametric comparisons using Kruskal-Wallis tests were conducted in each incentive condition and Dunn’s post hoc tests were used for pairwise group comparisons.

***Imaging data***. Following the discarding of the first 5 volumes, 330 volumes were analyzed in each incentive context. First level analyses were carried out using FSL FEAT (FMRI Expert Analysis Tool, www.fmrib.ox.ac.uk/fsl) with standard settings in FSL 6.0. Image preprocessing included brain extraction using BET [Smith, 2002] nonlinear registration of T1 images to Montreal Neurological Institute (MNI) standard space with 10mm warps and boundary-based registration of the functional image to the corresponding T1 image. Isotropic smoothing kernel with Gaussian full width half measure of 5 mm was chosen. Second level analyses were in MNI standard space (resampled to 2x2x2 mm). Motion correction (mcflirt, Jenkinson *et al*, 2002) was used to linearly register all images in a 4D volume to an average image and to estimate head motion parameters (rotation, temporal derivatives). Temporal derivatives of explanatory variables (EVs) were included in the first level General Linear Model (GLM), in lieu of slice-timing correction.

  These analyses focused on trials where participants responded during the anticipation window, i.e. before target presentation. For the GLM fMRI analyses, premature responses were matched with corresponding correct trials where participants responded during target presentation. Specifically, a correct trial of the same type (e.g., neutral) appearing as close in presentation as possible to the premature trial was selected. Matching correct trials could occur both before and after the corresponding premature trial. This procedure ensured that contrasts included the same number of premature and correct events. For the purpose of the logistic regression fMRI analysis, all correct trials were used.

  The first level design matrix included six EVs and their temporal derivatives: 1) premature responses for money incentive, 2) corresponding correct responses for money incentive, 3) premature responses with drug cue incentives, 4) corresponding correct responses for drug incentives, 5) all remaining correct responses, and 6) all feedback events. EV1-EV5 were modelled with event onset and duration as the start and duration of the anticipation window. EV6 was modelled with onset times and durations for the feedback. Effects of motion were controlled by including 24 motion parameters in the design matrix. Parameter estimate contrasts were calculated for EV1-EV2 (premature > correctmoney) and for EV3-EV4 (premature > correctdrug).

  Whole brain maps from the first level analyses were passed to a second level design matrix that tested mean group activations and pairwise group differences using one-sample and independent sample t-tests, respectively. Second level analyses used FEAT GLM with FLAME1 and a cluster forming threshold of z=2.3 and voxel-wise threshold of p<0.05. Demeaned gender was included as a covariate in the GLM. The striatum region of interest (ROI) mask was created by combining bilateral masks for caudate, putamen, and nucleus accumbens from the Harvard-Oxford atlas [Desikan et al., 2006].

  Interaction effects between groups (HC, SIB and SDI) and the within-subject factor of incentive block (money vs drug) were tested in the striatum region of interest (ROI) as follows: for each participant, difference maps were calculated for contrast of parameter estimates between the premature>correctmoney and premature > correctdrugcontrasts. Using FSL randomise with n=5000 samples [Nichols and Holmes, 2003; Winkler et al., 2014], one sample t-tests revealed the areas in which the groups showed greater activation to the premature>correctmoney contrast than to the premature > correctdrugcontrast. An F-test comparing the three group means was used to detect any significant interaction between all three groups. This interaction was further investigated using independent sample t-tests in FSL randomize (n=5000 samples), comparing the contrast of parameter estimates difference maps between the groups (HCs vs SDIs, SIBs vs SDIs) to indicate regions with a significant interaction (Figure 2). The resulting familywise-error corrected p-value maps with threshold-free cluster enhancement [Smith and Nichols, 2009] from the striatum ROI were thresholded at p<0.05 and the ROI comprising voxels with a significant interaction between both HCs and SDIs and SIBs and SDIs was selected for extraction of % signal change using featquery (<http://mumford.fmripower.org/perchange_guide.pdf>; https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FEAT/UserGuide#Featquery\_-\_FEAT\_Results\_Interrogation).

  A potential limitation of the traditional mass univariate GLM in this instance is the low number of events (premature and corresponding correct responses, see Figure 1A) as explanatory variables of interest. To test whether the GLM findings were robust, we selected ROIs that were significantly active in group mean activation maps. We then extracted and preprocessed the timeseries of these ROIs, located the anticipation windows in which premature and correct responses were made and used the BOLD activation in each of these anticipation windows to predict whether this window included a premature or a correct response. If ROIs are more active during premature than during correct responses, their activation should be sufficient to classify response type. For this analysis, all correct responses were included. Inferior frontal cortex (IFC), anterior cingulate cortex (ACC), parietal operculum (pO) and striatal ROIs were chosen based on group mean activation. More details on timeseries extraction and preprocessing can be found in supplementary information

In HCs, the BOLD signal on each trial in the IFC, striatum, ACC and parietal operculum (pO) was used to predict trial type (premature vs correct) in money context, while the BOLD signal in the ACC was used to predict trial type in drug context.  In the SIBs, the BOLD signal on each trial in the IFC, striatum and ACC was used to predict trial type (premature vs correct) in money context only.  In the SDIs, BOLD signal on each trial in the IFC, ACC and the pO was used to predict trial type in the money context, and BOLD signal in the IFC, ACC, striatum and pO was used to predict trial type in drug context. These regions were significantly activated in the group mean maps in the GLM results and were thus selected to validate the group results. Hierarchical logistic regression with subject-level random effects in RStudio (glmer function in the lme4 package) tested whether the ROI timeseries significantly predicted trial type. Receiver Operating Curves and area under the curve (roc and auc functions in the caTools package, plot and lines functions in the ggplot package) provided an additional metric of model performance.

***Brain-Behavior Correlations.*** To assess possible brain-behavior relationships, logistic regression models were fitted for each participant individually (using glm, MATLAB). Beta regression weight values for the IFC, striatum and ACC were correlated with BIS11 [Patton et al., 1995] and a self-reported estimate of likelihood to pick up money on the floor (Value of Money). For SDIs, correlations were also assessed with the obsessive-compulsive drug use scale (OCDUS) scores. In the SDI group, ACC, striatum and IFC beta values from the drug incentive condition were analyzed, whereas in the HC and SIB groups, ACC, striatum and IFC values were used in the money incentive condition. Since the beta value distribution was non-normal, Spearman’s rank correlations with confidence intervals are reported, with p-values using Bonferroni-correction for multiple comparisons within each group.

***Dynamic Causal Modelling.*** To explore directional interactions (effective connectivity) between the regions identified by the mass univariate GLM analysis, a set of dynamic causal models were created, estimated and tested in SPM12 (v6906). Following the GLM activations, dynamic causal models (DCM) were tested with the money incentive condition for the HC group and with the drug incentives for the SDI group. In two participants in each group the DCM analyses for three or more models failed to converge, resulting in these participants being excluded for this analysis. During the DCM analyses for the SIB group, the Bayesian model selection (BMS) phase did not yield a single winning model limiting any interpretation of their results. Additional information is reported in the supplementary information. The network of interest included three ROIs: IFC (pars opercularis), Caudate and ACC, in keeping with the recommendation that only commonly activated regions in both groups be included as nodes (Seghier et al., 2010). Time series were extracted in similar way to the logistic regression analysis, i.e. based on the individual peak activations in response to the premature>correct contrast. Here, time series preprocessing only included despiking (>4SD from the mean) of the SDI group to address severe motion artifacts and were not mean centered (since the Eigenvariate extraction results in a mean of 0 and an SD of 1). Firstlevel GLM analyses were re-estimated in SPM12 for the money and drug incentive conditions in a separate GLM and combined with the preprocessed timeseries in the DCM. Second level group maps estimated in SPM12 were consistent with 2nd level group maps estimated in FSL.

 Model space definition aimed to address two issues: firstly, we wanted to confirm the interactive architecture of the ACC – Caudate – IFC network by comparing a fully interactive model family (family A, Figure 3a) with architectures where one of the connections is pruned (model families B-G). Secondly, we wanted to investigate at which node or connection in the network the modulation by premature>correct contrast occurs (models A1-A9). All principle experimental conditions (correct, premature, failure) were used as driving inputs to the ACC and IFC, similarly to previous DCM analyses of response inhibition [Rae et al., 2015; Rae et al., 2016]. Modulatory effects were placed at each possible node and connection. Random effects (RFX) Bayesian model selection was ran on the full model space including families A-G, and on family A only; with both yielding the same results. Since strong *a priori* evidence from functional and structural connectivity studies [Choi et al., 2017b; d’Acremont et al., 2013; Menon, 2015; Sadaghiani and D’Esposito, 2015; Uddin, 2016] and family comparison results supported the fully interactive model, the comparisons reported focus on models A1-A9. We computed exceedance probabilities (EPs) for each model and used those to determine the winning model (family) in a one-state, bilinear, deterministic DCM. Bayesian model averaging (BMA) provided estimates of the fixed and modulatory connections in each subject, weighted by the evidence of each model tested (A1-A9). Using individual parameter estimates, group mean activations and group comparisons were tested using one sample t-tests and independent sample t-tests (Figure 3).

**Results**

**SDIs exhibit greater premature responding to drug incentives.**Participants in all three groups occasionally made ‘premature’ responses in the anticipation phase while waiting for the target to appear unpredictably (Figure 1A). There were significant differences between HCs, SIBs and SDIs in the number of premature responses for drug-related incentives using the nonparametric Kruskal-Wallis test (χ2(2)=10.56, *p*=0.005, Figure 1B). The groups did not significantly differ for money incentives (χ2(2)=3.39, *p*=0.18). Additionally, there were no significant group differences in performance accuracy for either money (F2,129=2.83, *p*=0.06) or drug incentives (F2,129=0.63, *p*=0.53). Mean (SD) numbers of correct trials for HCs, SIBs and SDIs were 41(4), 41(4), and 39(5), respectively in the money context, and 41(5), 41(4), and 40(5), respectively in the drug context. The behavioral results indicate that while infrequent (9% of total responses), premature responding is ubiquitous, exhibited by the majority of participants. SDIs exhibited elevated levels, specifically with drug-related incentives, with no differences between HCs and SIBs. This is consistent with greater self-reported general impulsivity in SDI (Table 1), whilst capturing their behavioral difficulties in action-restraint when anticipating drug-related rewards.

**Brain activation associated with premature responding depends on group-specific incentives*.***We next investigated whole-brain group activations associated with premature responding. To this end, trials where a premature response was detected were contrasted with adjacent trials with a correct response. fMRI analyses of the premature>correct contrast revealed wide-spread incentive-specific neural correlates of premature responses in the three groups. Group level activations (Figure 1C) for the money incentive condition in HCs and SIBs and for the drug incentive condition in SDIs included the inferior frontal cortex *pars opercularis* (IFC), dorsal anterior cingulate cortex (dACC) and dorsal striatum (dStriatum encompassing the caudate and putamen). For money incentives, the HC and SDI groups showed activation in the parietal operculum and the thalamus in addition to the dACC and IFC. For drug incentives, SDIs also activated portions of the motor, temporal, parietal cortices and the thalamus (Figure 1C, one sample t-tests, cluster forming threshold z > 2.3, p<0.05).

These findings suggest a similar pattern for the three groups with money incentives, but with drug incentives only SDIs exhibited activations associated with restraint failures. To assess this interpretation, group differences were interrogated using a whole brain mask. This revealed no significant differences in the monetary incentive condition. In contrast, in the drug condition SDIs showed greater activation than SIBs and HCs in the IFC, striatum, primary motor cortex, PCC, parietal, temporal areas and the thalamus in addition to the ventromedial PFC (SDI>HC only) and amygdala (SDI>HC only) (independent samples t-tests, cluster forming threshold z > 2.3, voxelwise FWE corrected p<0.05, SI Appendix Table S1). The BOLD results thus converge with the behavioral findings, with SDI showing clear abnormalities with drug incentives, and with SIBs differing from SDIs and being no different from controls.

If the BOLD activity were dependent on personal relevance regardless of incentive, then considerable overlap would be expected between money incentives in the HC and SIBs, and drug incentives in the SDI group. Conjunction analyses (fsl easythresh\_conj) indeed confirmed large-scale overlap in the IFC, dACC, striatum and parietal operculum between the HCs and SIBs (money incentives) and SDIs (money and drug incentives). We can therefore conclude that across three separate groups, the findings specify a set of fronto-striatal regions that are involved in impulsive responding when anticipating a personally-desired outcome. That similar regions were activated in the drug users, regardless of incentive type, suggests a general system that is associated with failures in self-restraint that is also sensitive to motivational processes.

**BOLD activation in the IFC, dACC, dStriatum and parietal operculum predicts trial type, thus validating the GLM results*.***To ascertain whether the GLM group mean activation maps in the three groups were robust to the limited number of premature trials, we assessed whether the resulting BOLD activations could reliably predict trial type. Specifically, BOLD timeseries activation in the IFC (*pars opercularis*), dACC, dStriatum and the parietal operculum were used to predict whether trials were correct or premature (Figure 1D). Hierarchical logistic regression models with subject-level random intercepts performed well in differentiating premature from correct responses in the money context (AUCHC=0.77; AUCSIB=0.73; AUCSDI=0.75) and in the drug context (AUCSDI=0.71). The differential contributions of each region of interest (ROI) to the prediction analyses can be found in SI appendix Table S2. The results for financial incentives, found independently in the three groups, further lends support to the reliability of the involvement of these regions in premature responding. This, together with the considerable overlap between groups noted above supports and validates our fMRI findings.

**Differential dStriatum involvement in SDI compared to HC and SIB.**Previous work has established striatal involvement in addiction, task control and reward processing [Everitt and Robbins, 2016; Oldham et al., 2018]. We thus focused on a ROI encompassing bilaterally the caudate, putamen and nucleus accumbens. To assess the selective involvement of the striatum in failures of restraint specifically to drug cues in SDIs, interaction effects between group (SDI vs SIB vs HC) and incentive type (money vs drug) were examined within this striatal mask. Significant interaction effects in the caudate (MNI [-11 10 16], t=3.2, PFWE=0.05) and putamen (MNI [-34 -2 -4], t=3.9, PFWE<0.04) were driven by greater activation in money compared to drugs in the HCs versus the SDIs (SI Appendix Figure S2). Similarly, interactive effects in the caudate (MNI [-12 12 16], t=3.26, PFWE=0.04) and putamen (MNI [-34 -2 -4], t=4.1, PFWE=0.01) were driven by greater activation in money compared to drugs in the SIBs versus the SDIs. Figure 2B shows a subregion of the caudate, in which SDIs show a different pattern of activation from *both*SIBs and HCs (MNI [-12 12 16], HC vs SDI t=3.31, PFWE=0.05; SIB vs SDI t=3.26, PFWE=0.04; region extent 6 voxels). This more stringent and focused approach indicated that the BOLD response in the dStriatum was differentially sensitive to impulsive responding for money in non-dependent individuals and to impulsive responding to drug cues in SDIs, regardless of familial vulnerability (Fig 2C). Overall, there was a clear opposing pattern in the dStriatum with reduced activation for monetary incentives in SDIs compared to the other two groups, pointing to specific abnormalities in this region in the SDIs.

**Limited associations between traits, behavior and BOLD signal.**We next investigated the relationship between neural activation associated with premature responses and self-reported impulsivity (Barratt Impulsivity Scale; BIS), money valuation (Value of Money), and in SDIs also compulsive drug use (OCDUS). For dACC, IFC and striatal regions, we used corresponding beta values from individual logistic regression models as a measure of each region’s sensitivity to premature responses. In SDIs responding to drug incentives, caudate activation was positively associated with self-reported impulsivity (rs=0.36, p=0.02, 95%CI [0.06 0.6]) and with compulsive drug use (rs=0.40, p=0.011, 95%CI [0.1 0.63]). In this group IFC beta values in the drug context were also significantly associated with impulsivity (rs=0.36, p=0.02, 95%CI [0.06 0.6]), and with compulsive drug use (rs=0.42, p=0.008, 95%CI [0.14 0.64]). In HCs, there was a significant negative association between caudate beta values for monetary incentives and money valuation ratings (rs=-0.36, p=0.02, 95%CI [-0.1 -0.6]). No correlations survived Bonferroni correction, with nine comparisons in SDIs and six in the HCs, and hence are reported within an exploratory framework. Finally, no significant correlations were found in SIBs. Whilst only suggestive, these correlations appear consistent with the interpretation above of an abnormal involvement of the striatum in impulsive responding in SDIs.

**Group differences in effective connectivity in the same network.**Although HCs and SDIs showed activation in similar brain regions when failing to restrain responses to different incentives, it is possible that the underlying network architecture and their connectivity differ in some way. This would be consistent with evidence positing deficient communication between prefrontal and subcortical regions in addiction [Goldstein and Volkow, 2011; Ma et al., 2015]. Namely, the direction of effective connectivity between regions or the influence of response type (premature vs correct) on connectivity may differ between the two groups. To test these possibilities, we used DCMs based on neurobiologically plausible circuits [Choi et al., 2017b; Haber, 2016]. Such DCMs fit generative models to assess the directed influence of one region over another, allowing us to compare competing hypotheses about functional interactions in a set of ROIs.

First, we assessed the underlying functional architecture of the network including the IFC, dACC and striatum in the SDI (drug context) and HC (money context) groups. We tested models with either a pruned connection or the full model yielding seven model families. Within each family we varied all possible modulation locations, yielding 57 models in total (Figure 3A). These models were estimated and the evidence for each family was compared using family-wise Bayesian Model Selection (BMS). For both groups independently, the fully interactive model architecture, including bidirectional connections between all regions of interest, was confirmed by family-wise BMS (exceedance probabilities>0.99). This is consistent with the literature reporting the presence of structural and functional connections between these regions[Choi et al., 2017b; Haber, 2016].

Additionally, we interrogated which connections were modulated by trial type. Focusing on the fully-interacting model architecture of family A, BMS on models differing in the location of the modulation  (IFC, dACC, dorsal striatum nodes, or one of the six directed connections) revealed the same winning model for the HC group network active in money context as for the SDI group network active in drug context (Figure 3B). Therefore, both HC and SDI participants appeared to activate the same network, provided that they find themselves in the appropriate incentive condition (money or drug, respectively). At the same time, exceedance probabilities of the winning models (Figure 3B), were below 0.9 (protected exceedance p=0.14 and p=0.21 for HCs and SDIs, respectively), suggesting some heterogeneity in the location of the modulation in both HC and SDI groups.

Next, the strength of the modulatory effects of the task (premature>correct trial type) and the fixed connections between ROIs were explored within each group using one-sample t tests. We also questioned whether the coupling parameters of the network were different in the two groups, using independent sample t-tests. Bayesian Model Averaging (BMA) allowed us to compute means for each model parameter, weighted by the posterior probability of each model for each subject. A summary of all parameters, including fixed connections between ROIs and task modulatory effects can be seen in Figure 3C. In HCs, BMA revealed a negative modulation of striatal activity by trial type (t39=2.0, p=0.03), while in SDIs trial type showed negative modulation of dACC activity (t38=2.0, p=0.03) and positive modulation of the dACC to dStriatum projection (t38=2.1, p=0.02). A dStriatum negative (autoinhibitory) fixed connection was thus amplified on premature trials in the HC group, whereas a dACC autoinhibitory connection and an excitatory connection from the dACC to the dStriatum was amplified on premature trials in the SDI group.

Independent sample t-tests, comparing the groups in modulation strength, showed only a significant difference in modulation of the dACC to dStriatum projection by trial type, which was higher in SDIs than in HCs (t77=2.13, p=0.036, Figure 3D). Fixed connections between nodes did not differ between the two groups (all *p*s>0.05) except for ACC to ACC autoinhibitory connection, which was higher in HC than SDI (t77=2.36, p=0.021).

To summarize, Bayesian model selection revealed a fully-interactive model architecture given relevant incentives, with bilateral connections between the IFC, dACC and dStriatum. Among these interactive models, the dStriatum is critically involved as it is modulated by the premature vs correct trial condition. Bayesian model averaging suggested differences between HC and SDIs, with task-based modulation changing striatal activity directly in HC, while in SDIs striatal modulation was driven by the dACC.

**Discussion**

Our results integrate research on inhibitory response control and reward-related processing, offering insight into how these manifest jointly in the human brain and relate to impulsivity. The findings point to the intrinsic importance of context and the nature of the relevant incentives in modulating inhibitory control processes, particularly in relation to drug addiction. We show that failure of response control when faced with anticipating rewards is underpinned by fronto-striatal and cingulo-opercular network activations, wherein dorsal striatum regions play a key role. By focusing on failures of restraint in the MID task, the study demonstrates for the first time how impulsive actions can be triggered by drug incentive cues in individuals addicted to stimulant drugs.

**Shared networks for premature responding to personally salient cues.** This study identified key brain regions associated with failures of restraint in the presence of reward in SDI, their unaffected siblings and HC. Robust activation of the cingulo-opercular and fronto-striatal network regions comprising dACC, IFC (pars opercularis), inferior parietal cortex (parietal operculum), striatum and thalamus were present in healthy individuals and an attenuated version of this network minus thalamic and parietal activation in the unaffected siblings with monetary incentives. Activations were also seen in response to failed impulse control in SDI for money and drug incentives. That all three groups independently exhibited a similar pattern reinforces its robustness.

The regions identified here are known to be differentially activated to errors across a variety of cognitive tasks and populations [Neta et al., 2015; Norman et al., 2019]. This is in keeping with the notion that premature responses are brought about by failure of control processes. Analyses of failures when trying to stop a prepotent response reveal similar regions encompassing the IFC, ACC, dorsal caudate and inferior parietal cortex [Whelan et al., 2012]. At the same time, premature responding is of special interest as it is instigated in the presence of an incentive-driven impulse, pointing to general error processing mechanisms transcending different motivational contexts.

At the group level the neural correlates of action restraint were intimately linked to the personal relevance of the incentives. Thus, subjective value across all incentive types elicited activation in the regions described above [Bartra et al., 2013]. Subjective value here refers to individuals’ neural and behavioral response rather than self-reported valence or arousal ratings, which have previously been shown to diverge in studies of alcohol dependence [Nees et al., 2012]. Using multiple incentives revealed that healthy controls and the unaffected siblings exhibited activations when responding prematurely to monetary incentives but not to drug-related cues. SDI demonstrated similar activations but most robustly to drug-related cues where they further showed increased activation in brain areas that encode motivation and emotional salience (e.g., amygdala, OFC). Greater activation to drug incentives in these regions was accompanied by elevated premature responding in the active users, suggesting that incentive salience was driving this behavior, consistent with studies in experimental animals and in other addiction-related clinical populations [Dalley and Ersche, 2019; Voon, 2014].

**Neural correlates of impulsivity in drug addiction**. Increased impulsive behavior and corresponding increased activation in the presence of drug cues fit well within the iRISA framework [Goldstein and Volkow, 2002; Goldstein and Volkow, 2011; Zilverstand et al., 2018]. Drug cues such as those used in our version of the MID task are known to have abnormally high motivational significance to cocaine users [Goldstein et al., 2008] and elicit increased approach behaviors and upregulation across the brain including the cingulo-opercular network [Zilverstand et al., 2018]. These networks are generally underactive in active stimulant users during standard cognitive task performance including inhibitory control processing [Morein-Zamir and Robbins, 2015; Zilverstand et al., 2018]. While this is generally true of stimulant addiction, previous studies of reward anticipation in nicotine addiction [Bühler et al., 2010] suggest that dependent smokers can show similar levels of reactivity to cigarette compared to monetary rewards. Presently, whole brain analyses did not find significant hypoactivation in the SDI compared to control participants in whole-brain analyses with monetary incentives. This is in keeping with the limited differences in task performance and function between these same individuals in monetary reward processing [Just et al., 2019] and consistent with the broader literature on processing of non-drug rewards in addicted individuals [Zilverstand et al., 2018]. Whilst consistent with the notion that brain activity during inhibitory control is stimulus-dependent [Czapla et al., 2017], this limits proposals that drug use diminishes the perceived value of non-drug rewards to only specific brain regions [Goldstein and Volkow, 2011]. Taken together the results support an imbalance between incentive salience of drug and non-drug rewards in drug users [Bühler et al., 2010]. The iRISA model will likely benefit from further refinement by taking into account the nature of the reactivity measures, including behavioral, neural, physiological, or subjective responses [Nees et al., 2012].

Nevertheless, the importance of incentive type to impulsive responding in addiction is reflected across the brain and most clearly by the activation of caudate and putamen subregions of the dorsal striatum. Here, not only was there greater activation in SDI in the presence of drug incentives, but there was indeed a blunted response in the presence of monetary incentives compared to either of the two other groups. The temporal and correlational features of fMRI preclude us from determining the exact nature of dorsal striatal involvement in the processing of premature responding. This region could be directly implicated in triggering the impulsive responding or could be linked to monitoring processes. The dorsal striatum is involved in action control [Graybiel, 1995; Haber, 2016], but is also activated when participants anticipate rewards [Oldham et al., 2018], with neurons integrating reward information with movement processing [Schultz, 2016]. We therefore tentatively attribute the striatal involvement to the failure of control in the face of personally meaningful incentives. This explanation dovetails with the role of the dorsal striatum in cancelling planned motor responses [Bari and Robbins, 2013; Eagle and Baunez, 2010] and impulsive choice [Kim and Im, 2018], whilst providing evidence for its involvement in additional aspects of impulsivity in humans.

Abnormal striatal processing in addiction has been consistently linked to general aberrations in cortico-striatal circuitry that subserves motor, cognitive and motivational processes [Choi et al., 2017b; Choi et al., 2017a]. When considering this circuit, we show that SDIs and HCs share a strikingly similar network architecture, provided that a relative incentive is present. Nevertheless, whilst the striatum was consistently modulated by premature responding in the two groups, effective connectivity between the IFC, dACC and striatum pointed to the differential involvement of the latter two regions. The dACC sits at the connectional intersection of the brain’s reward and action networks [Haber, 2016], and is ideally situated to regulate impulsive behaviors. This region thus provides top-down inhibitory control over striatal representations of action and stimulus values and has been found to over-activate in SDIs with failures of response inhibition [Morein-Zamir et al., 2013]. Here the observed enhanced directed connectivity from dACC to dorsal striatum in SDIs relative to HCs suggests that interactions between the salience network and dorsal striatum via the ACC play a critical role in impulsive responding in drug abuse. Investigations using dynamic causal models in a response inhibition task also pointed to aberrant modulation of the dACC to dorsal striatum projections in cocaine dependent individuals [Ma et al., 2015]. Present results converge to support deficient communication between prefrontal and subcortical regions in addiction [Goldstein and Volkow, 2011].

**Additional implications of the present findings.** Interactions between the PFC and striatum are also found in non-human animal studies investigating PFC involvement in failures of restraint. The rodent prelimbic cortex, a homologue of human dACC in addition to rodent infralimbic cortex, a homologue of the human ventromedial PFC, has been implicated in premature responding [Dalley et al., 2011; Dalley and Robbins, 2017] and behavioural control [Gourley and Taylor, 2016]. We observed ventromedial PFC activations in SDIs consistent with previous human and animal literature [Dalley et al., 2011; Morris et al., 2016], again reinforcing the importance of incentive salience in this group. We note that the ventral striatum did not appear to be uniquely activated by failures of restraint, though it was robustly associated in all groups with general reward anticipation [Just et al., 2019]. Given its role in addiction, reward seeking, impulsivity and action initiation, striatal dopaminergic dysregulation may be a putative mechanism underlying the present findings in SDIs. Lower dopamine D2 receptor binding in the human striatum is associated with impulsivity [Buckholtz et al., 2010] and addiction [Trifilieff and Martinez, 2014]. In chronic cocaine users elevated dopamine neurotransmission in the dorsal striatum specifically was noted in the presence of cocaine-related cues [Volkow et al., 2006], dovetailing with its present involvement in incentive salience. Changes to dorsal striatum physiology involving D2 receptors have also been implicated in chronic exposure to cocaine in experimental animals [Porrino et al., 2004], consistent with its relationship to impaired impulsive control in the present study.

Assessing the unaffected siblings of SDI allowed us to test whether premature responding is a familial predisposition to stimulant drug addiction. Performance and neural correlates in siblings indicated this was not the case for both incentive types. As the connectivity analyses for the siblings did not result in a single winning model (see SI Appendix Figure S3) we could not extend this conclusion to network connectivity. Nevertheless, on balance we conclude that an impulsive endophenotype does not appear to extend to premature responding, at least under reward-based conditions. Indeed, it may be that increased restraint served as a protective factor for the siblings. Alternatively, it remains possible that the ability to restrain responding may be compromised in unaffected family members under more constrained conditions, such as those eliciting negative urgency [Um et al., 2019], with greater cognitive demands, or drug exposure [Sanchez-Roige et al., 2016].

The MID task has been favored in reward research partly because it requires participants to make simple decisions, minimizing cognitive confounds [Oldham et al., 2018]. This allowed us to attribute premature responses to failures of inhibitory control specifically. This approach offers a parsimonious measure of impulsive responding to diverse incentives in clinical and non-clinical populations. The presence of a relatively wide anticipation window likely increased premature responding frequency overall and allowed greater individual variability. Future research may further explore how varying specific task parameters in the MID task such as the nature and timing of reward delivery, can elicit or minimize premature responding, testing convergence with preclinical models. Additionally, performance and network changes in at-risk groups or following prolonged abstinence should be explored. This could better elucidate the exact role of various risk factors in addiction and point to potential mitigating factors.

In conclusion, our findings provide insight into the brain networks recruited during failures of restraint in a well-characterized sample of SDIs, their unaffected siblings and HC. Limited cognitive control over action was evident in the SDI group, particularly involving incentive-based situations in line with prominent theorizing. Whole brain analyses provide evidence for cortico-striatal network disruption in addiction involving top-down control by the PFC and its interactions with striatal structures with altered connectivity and abnormal striatal activation patterns. By capitalizing on the presence of premature responses in this version of the MID and introducing a novel analysis, our results indicate that different forms of impulsive behaviors are comprised of separable, though often overlapping neural networks and are selectively modulated by incentive motivational processes.

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**Tables**

**Table 1.** Demographic, personality and clinical measures for the three groups. Data are means ± standard deviation (SD). Significant differences (p<0.05) are highlighted in bold; Chi-square tests were used for categorical comparisons and one-way analyses of variance were used to test continuous outcomes. Gender (Chi2=36.48, p< 0.001), Impulsivity (Barratt Impulsivity questionnaire, BIS-11, F­2,123= 38.22, p<0.001) and Value Money (“How likely are you to pick up 10p/50p?”, F2,123=4.439, p=0.0138) were significantly different between the HC, SIB and SDI groups.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | **Healthy Controls** | | **At-risk Siblings** | | **Stimulant Dependent Individuals** | |
| **(n=42)** | | **(n=43)** | | **(n=41)** | |
| **Demographics** | **Mean** | **(SD)** | **Mean** | **(SD)** | **Mean** | **(SD)** |
| Age (years) | 32.6 | (8.8) | 32.3 | (8.4) | 34.7 | (7.4) |
| Gender (% male) | 64.3 |  | 47.8 |  | 88.1 |  |
| Verbal Intelligence (NART) | 112.0 | (8.4) |  |  | 110.6 | (7.4) |
| Monthly disposable income (£) | 695 | 1000 | 403 | 410 | 399 | 672 |
| Duration of stimulant use (years) |  |  |  |  | 16.1 | (6.5) |
| Compulsive Stimulant Use (OCDUS) |  |  |  |  | 23.6 | (9.3) |
| Impulsivity (BIS-11) | 59.7 | (7.9) | 67.3 | (10.5) | 77.3 | (9.3) |
| Value Money Ratings | 71.3 | (27.0) | 58.6 | (32.8) | 76.1 | (26.8) |

NART, National Adult Reading Test; OCDUS, Obsessive Compulsive Drug Use Scale; BIS-11, Barret Impulsivity Scale.

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**Figure Legends**



**Figure 1.**   Impulsive responding when anticipating an incentive. (A) Cues indicated the incentive available on each trial, and participants prepared during the anticipation phase to respond during the brief period when the target was presented. Incentives could be monetary (a small sum of money) or drug-related images, with 44 incentive and 22 neutral trials in each of the two contexts.  (B) The number of premature responses, where participants responded during the anticipation phase and before the target appeared, for each of the three groups and to the two incentives. Post-hoc tests with false discovery rate correction were used. (C) Axial brain slices (z=4, 21, 39, 56) demonstrating prefrontal and striatal activations when participants responded prematurely during the anticipation phase (whole brain cluster corrected z=2.3, p=0.05). (D) Receiver Operating Curves demonstrating good model performance of regions of interest timeseries in predicting whether participants responded prematurely or accurately to the target. Asterisks in figure denote level of significance (\*p<.05; \*\*p<.01)​

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**Figure 2**. Striatum ROI activation during premature responding. (A) HC and SIB exhibit striatal activation during premature responding to money incentives, while SDI exhibit a similar pattern but with drug incentives (MNI [y z]=[10 0]).  Activations in all groups span the caudate and putamen. (B) The striatum ROI (outlined in red), with the subregion in yellow in theanterior dorsal caudate (n=6 voxels) exhibiting an opposing pattern with increased activation in both HC and SIB and decreased activation in SDI when responding prematurely to money versus drug incentives. (C) Mean percent signal change (with 95% CI) of the difference between activation for money versus drug incentives in the anterior dorsal caudate. A significant interaction, F(2, 123) = 5.99, p=0.003, was found between group and incentive condition.  Specifically, Money-Drug difference was significantly greater in the HC than in SDI and in the SIB than in SDI (post-hoc tests with false discovery rate correction, p=0.0016 p=0.0074, respectively). Asterisks in figure denote level of significance (\*\*p<0.01)​

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**Figure 3.** DCM connectivity analyses. (A) Model space and structure of the DCMs compared. Top row shows the 7 families, testing different possible connections between the 3 regions. Black arrows denote driving inputs (all MID trials) and red arrows indicate the modulatory effect of premature responding (premature versus correct). Within the fully interactive family A, model A3 with modulation to the striatum is highlighted as having the greatest exceedance probabilities for both HC with money incentive and for SDI with drug incentives (see also SI Appendix S3). (B) Random effects Bayesian Model Selection within family A, demonstrating the evidence in support of model A3 for both groups, each with their relevant incentive. (C) Average connectivity estimates. All fixed connections survive Bonferroni corrections. Modulatory connections are significant at p<0.05, uncorrected. (D) Mean group modulation strength estimates (with 95% CI) from the BMA for HC and SDI (each with their respective incentive). Only the ACC to striatum modulation differed significantly between the groups. Asterisk in figure denotes level of significance (\* p<.05, uncorrected); ACC: anterior cingulate; STR: striatum incorporating caudate and putamen subregions that were active in the HC and SDI group, respectively; IFC: inferior frontal cortex including the frontal operculum only.