

Neurofunctional Reward Processing Changes in Cocaine Dependence During Recovery

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Although reward processing appears altered in addiction, few studies track neurofunctional changes following treatment or relate these to measures of reduced drug use. The current study examined neurofunctional alterations in reward processing in cocaine dependence (CD) pretreatment and posttreatment to determine whether these changes relate to clinically meaningful outcome indicators. Treatment-seeking CD outpatients ($N=29$) underwent functional magnetic resonance imaging while performing a monetary incentive delay task (MIDT) pretreatment and posttreatment. The MIDT parses anticipatory from outcome phases of reward/loss processing. Abstinence indicators (negative urines, days abstinent from cocaine during follow-up) were collected throughout treatment and up to 1 year later. Healthy control (HC) participants ($N=28$) were also scanned twice with the MIDT. Relative to pretreatment, at posttreatment CD participants demonstrated increased anticipatory reward activity in the midbrain, thalamus, and precuneus ($p_{FWE}<0.05$). Increased midbrain activity correlated with cocaine abstinence during the 1-year follow-up. Ventral striatal (VS) activity during loss anticipation correlated negatively with negative urine screens. HC group test–retest results showed decreased ventromedial prefrontal cortex activity during winning outcomes. CD–HC group-by-time differences revealed increased left inferior frontal gyrus activity in the CD group during anticipatory phases at posttreatment. In CD participants, increased posttreatment activity in dopamine-innervated regions suggests lowered thresholds in anticipatory signaling for non-drug rewards. Midbrain and VS responses may represent biomarkers associated with CD abstinence. Abstinence-related neurobiological changes occur in similar regions implicated during active use and may possibly be used to track progress during short- and long-term recovery.

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INTRODUCTION

Relapse rates for cocaine dependence (CD) remain among the highest of all illicit drugs (Heyman, 2013; Vocci, 2007). Although many neuroimaging studies examine chronic effects of cocaine on cognitive processes, relatively few examine brain changes occurring with discontinued drug use (Garavan *et al*, 2013; Hanlon *et al*, 2013). Investigating neurofunctional changes associated with abstinence may provide important insights regarding strategies for cessation and relapse prevention (Garavan *et al*, 2013; Hanlon *et al*, 2013).

To date, most neuroimaging studies exploring abstinence in CD have been cross-sectional, often with small samples and with varied testing times (eg, posttreatment, long-term abstinence). One of the few longitudinal neuroimaging CD

studies found that, at follow-up, CD individuals demonstrated an enhanced midbrain response during a cognitive-control task, which inversely related to simulated cocaine choice (Moeller *et al*, 2012). Although limited by a small sample and the absence of a control group, this study nonetheless provides some evidence of network recovery with sustained abstinence. Greater midbrain activation has also been observed in a reward processing task comparing former, relative to current, cocaine users (Patel *et al*, 2013); abstinence in the former users correlated with right ventral striatum (VS) activity during anticipatory processing (Patel *et al*, 2013). This study is limited in its cross-sectional design; current and former cocaine users differed in their self-reported cocaine use, with length of abstinence difficult to assess in the former cocaine-using group.

The current study aimed to provide insight into the neural basis of recovery; few studies track neural changes within individuals from baseline to treatment follow-up and relate these to abstinence indicators. This study employed a longitudinal design to follow a large group of CD individuals and examine brain changes in generalized reward processing

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Table 1 Participant's Demographic Information

	CD	HC
<i>n</i>	29	28
Male/female	21/8	14/14
Age (SD)*	41.34 (6.7)	31.25 (10.1)
White/Black/Hispanic/Asian/multiracial	12/13/2/0/2	18/9/1/1/0
Years of education*	12.52 (1.3)	15.36 (2.1)
Previous 28 days cocaine use	15.52 (6.9)	—
Previous 28 days cigarette use	21.07 (12.0)	—
Previous 28 days alcohol use	6.45 (7.7)	—
Previous 28 days marijuana use	1.21 (2.8)	—
Years cocaine use	9.41 (6.1)	—
Age at first cocaine use	22.28 (7.9)	—
Days in treatment	53.59 (31.9)	—
Sessions attended	6.21 (4.3)	—

* $p < 0.05$.

before and after treatment. Specifically, our main aims were to:

(1) Use a longitudinal, rather than cross-sectional, design to examine brain changes occurring from pretreatment to posttreatment in a large group of individuals participating in a randomized clinical trial of CD treatments. Few studies track neural changes within individuals from baseline to follow-up. Many addiction theories emphasize changes in the mesocorticolimbic dopamine system to motivationally salient stimuli across the addiction cycle; for example, increased release to drug-related cues (Goldstein *et al*, 2009) but diminished response to natural/generalized rewards (Volkow *et al*, 2007). If chronic drug use modifies dopamine signaling, then protracted abstinence may also alter activity in these regions. Consistent with previous CD studies (Moeller *et al*, 2012; Patel *et al*, 2013), we hypothesized that, at posttreatment, relative to pretreatment, CD individuals would demonstrate functional enhancement of dopaminergically innervated brain regions, including the midbrain, thalamus, and preceus.

(2) Examine generalized reward processing changes, particularly anticipatory processing. Many addiction treatments focus on reducing drug use by increasing the salience of non-drug-related stimuli (Prendergast *et al*, 2006; Vocci, 2007). Learning-based addiction theories underscore anticipatory processes as critical for associative learning mechanisms leading to habit formation (Everitt and Robbins, 2013). We hypothesized that functional enhancement of dopaminergically-innervated brain areas may occur specifically during anticipatory processing.

(3) Assess the degree to which functional changes in anticipatory processing relate to abstinence measures over time. Anticipatory processing recruits the VS, which effectively codes for reward-predicting cues, including drug cues (Knutson *et al*, 2001). Several cross-sectional studies link striatal reward processing with abstinence in CD, yet with somewhat differing results (Bustamante *et al*, 2013; Jia *et al*, 2011; Patel *et al*, 2013); this may relate to clinical differences, such as treatment-seeking status, length of abstinence, and recent cocaine use (Balodis and Potenza,

2015). Changes in the VS may act as a possible biomarker of relapse risk, although no study has directly examined this possibility. Given the importance of the VS in drug and reward processing, this area was selected as a region of interest (ROI) to examine relationships with abstinence indicators. Abstinence often occurs incrementally, rather than as a dichotomous process (Konova *et al*, 2013); negative urine screens are an objective measure of abstinence that has been identified as a clinically meaningful outcome indicator (Carroll *et al*, 2014). We hypothesized that pretreatment-posttreatment VS anticipatory changes would relate to negative urine screens.

(4) Examine practice effects on a reward processing task. In order to control for potential practice effects on the task, a control group was included who completed the task twice. Given the role of fronto-striatal circuitry in mediating reward, we hypothesized reduced activity in this network upon repeated testing.

MATERIALS AND METHODS

Participants

Participants consisted of 57 individuals providing written informed consent (see Table 1 and Supplementary Data for sample characteristics and diagnostic procedures). CD participants were scanned prior to treatment and at posttreatment. The average time between scanning sessions (ie, Scan1 and Scan2) for participants was 129 days (SD = 66.90) with no significant difference between CD and HC groups ($F(1,55) = 0.61$, $p > 0.05$). CD participants were treatment-seeking individuals recruited from a larger clinical research trial. CD participants received cognitive behavioral therapy with random assignment to 12 weeks of adjunct therapy and a 1-year follow-up; details of the RCT and outcomes are described in the main trial report (Carroll *et al*, 2016), here we focus on neural functioning and abstinence.

CD Group Assessments

Rather than one single measure, multiple outcome indicators have been identified as clinically meaningful (Carroll *et al*, 2014); here we present negative urine screens as our primary abstinence indicator. Negative urine screens comprise an objective measure of abstinence that has been identified in a recent review as a clinically meaningful indicator of treatment outcome (Carroll *et al*, 2014). Urine toxicology screens were obtained three times weekly during treatment and at each follow-up interview.

Reward Task

Participants completed two MIDT runs during each scanning session. Each trial includes anticipatory phases A1 (prospect of reward) and A2 (anticipation of reward receipt) and an outcome phase (OC). The task description can be found in Supplementary Data and previous publication (Patel *et al*, 2013).

Functional images were preprocessed using SPM5 (Wellcome Functional Imaging Laboratory, London, UK), normalized to the Montreal Neurological Institute template and smoothed with a 6-mm FWHM kernel. First-level modeling

used robust regression to reduce outlier influences (Wager *et al*, 2005). Motion and high-pass filter parameters comprised additional regressors of no interest. NeuroElf analysis package (www.neuroelf.net) was used for second-level random-effects analysis. Recommended cluster-extent-based thresholding analytic and reporting practices for fMRI were applied (Woo *et al*, 2014); correction for multiple comparisons was conducted using Monte-Carlo simulation (ie, AlphaSim), using a combined voxel-wise ($p < 0.001$) and cluster thresholds to result in a family-wise error (FWE) rate of $p_{FWE} < 0.05$. To investigate brain activation over time, we contrasted: (1) anticipation of monetary gain during Scan2 vs Scan1 for A1 and A2 phases (A1Win and A2Win, respectively); (2) anticipation of monetary loss during Scan2 vs anticipation of monetary loss during Scan1 for A1 and A2 phases (A1Loss and A2Loss, respectively); (3) outcome win processing during Scan2 from outcome win processing during Scan1; and (4) outcome loss processing during Scan2 from outcome loss processing during Scan1. Our analyses focused on Scan2 relative to Scan1 differences within the CD group, and further present test-retest differences in an HC group. Additionally, between-group comparisons among the treatment-seeking CD group and the HC group were examined.

ROI Analysis

Given the small VS volume, together with evidence implicating this area in anticipatory processing, the MIDT, and CD pathophysiology, the VS constituted an *a priori* ROI specifically during this reward phase (ie, A2). The ROI on the right side was defined and localized based on reward-processing findings in a previous cross-sectional MIDT study in former CD users (Patel *et al*, 2013), reporting a relationship between right anticipatory VS activity and abstinence duration. Additionally, this lateralized region was further identified in a meta-analysis of brain responses to cocaine cues, with the right VS also linked to drug craving (Kuhn and Gallinat, 2011). Activity from a spherical ROI (5 mm radius around 12, 12, -9; Figure 2) was extracted for each CD participant to examine the mean blood oxygen level-dependent percentage of signal change from baseline. Subsequently, Spearman's rho correlations in SPSS, version 17.0 (SPSS, Chicago, Illinois) tested the relationship between anticipatory VS changes and negative urine screens. Specifically, VS ROI activity during win and loss anticipation (Scan2 win cues > Scan1 win cues; Scan2 loss cues > Scan1 loss cues) during A2 was correlated with negative urine toxicology screens and Bonferroni-corrected for multiple comparisons.

RESULTS

Main Effect of Time/Treatment on Anticipatory Processing in the CD Group

No significant Scan2-Scan1 differences were observed during A1 phases, the A2Loss phase, or OC phases in CD participants.

During the A2Win phase (associated with the anticipation of reward), Scan2-Scan1 differences recruited bilateral thalamus extending to right caudate and lentiform nucleus (Table 2a; Figure 1b); precuneus; posterior cingulate

extending to culmen (Figure 1a and c); and right mid-brain/substantia nigra extending to lentiform nucleus (Figure 1d). A previous study in CD inversely linked enhanced midbrain response in an overlapping region with simulated cocaine choice (Moeller *et al*, 2012). In an effort to replicate and extend this aforementioned report, we conducted an exploratory analysis between the midbrain cluster and long-term abstinence; midbrain activity correlated with days of cocaine abstinence during follow-up ($r = 0.48$, $p < 0.01$; Figure 1e).

Changes in anticipatory VS activity and negative urines. Spearman's rho statistic between A2W posttreatment > pretreatment changes and negative urine screens was not significant. The total negative urines during treatment, however, correlated inversely with A2L posttreatment > pretreatment VS changes ($r_s = -0.42$, $p = 0.05$; Bonferroni-corrected; Figure 2).

HC Test-Retest Differences

Within this group, significant Scan2-Scan1 differences occurred only during the OCWin phase; showing a significant decrease in ventromedial prefrontal cortex (vmPFC) extending to anterior cingulate from Scan1 to Scan2 (Table 2b, Figure 3a and b).

Differences Between CD Posttreatment-Pretreatment and HC Test-Retest

There were no between-group differences during A1 or OC phases when comparing Scan2-Scan1 activity. During both A2Win and A2Loss phases, group differences appeared in left IFG driven by increased posttreatment-pretreatment activity in the CD group (Figure 3d). Groups also differed in the right superior temporal gyrus, where the CD group demonstrated Scan2-Scan1 increases (Table 2c; Figure 3c). Extracted signals from these regions covarying for age did not alter results.

DISCUSSION

Using a longitudinal design, the current study examined reward-processing changes in CD participants during treatment. Significant neurofunctional changes occurred predominantly during anticipatory processing with additional relationships to a clinically meaningful outcome indicator. Inclusion of HC subjects permitted characterizing test-retest effects and CD-HC group differences.

Within-Group CD Differences

Increased anticipatory reward processing. In line with our hypothesis, changes in anticipatory processing occurred from baseline to the end of treatment in the CD group. Specifically, CD participants showed increased anticipatory reward activity in the midbrain and thalamus areas extending to striatal regions. Our findings replicate and extend findings from two prior CD studies. Our longitudinal findings of increased midbrain activity from baseline to the end of treatment are consistent with prior cross-sectional CD data indicating greater midbrain activity in the former,

Table 2 Scan2–Scan1 Differences During the MIDT

MIDT phase	Structure	BA	Left/right	MNI coordinates				T-value
				x	y	z	k	
<i>(a) CD Group</i>								
A1Win	—	—	—	—	—	—	—	—
A1Loss	—	—	—	—	—	—	—	—
A2Win	Thalamus/caudate/lentiform nucleus	—	R	3	–12	12	118	4.11
	Midbrain/substantia nigra/lentiform nucleus	—	R	15	–21	–15	46	4.14
	Precuneus	7	L	–3	–60	57	33	4.02
	Posterior cingulate/culmen	29	R	3	–45	3	38	4.12
A2Loss	—	—	—	—	—	—	—	—
OCWin	—	—	—	—	—	—	—	—
OCLoss	—	—	—	—	—	—	—	—
<i>(b) HC Group</i>								
A1Win	—	—	—	—	—	—	—	—
A1Loss	—	—	—	—	—	—	—	—
A2Win	—	—	—	—	—	—	—	—
A2Loss	—	—	—	—	—	—	—	—
OCWin	vmPFC/anterior cingulate	10/24	R	6	45	–9	59	–4.22
OCLoss	—	—	—	—	—	—	—	—
<i>(c) CD vs HC Group</i>								
A1Win	—	—	—	—	—	—	—	—
A1Loss	—	—	—	—	—	—	—	—
A2Win	Inferior frontal gyrus	44	L	–53	10	18	23	3.95
A2Loss	Superior temporal gyrus	41	R	42	–33	12	17	3.85
	Inferior frontal gyrus	44	L	–51	15	15	26	3.77
OCWin	—	—	—	—	—	—	—	—
OCLoss	—	—	—	—	—	—	—	—

Abbreviations: A1Loss, prospect of loss phase; A2Loss, anticipation of loss; A1Win, prospect of reward phase; A2Win, anticipation of reward; BA, Brodman's area; k, cluster; MIDT, monetary incentive delay task; OCLoss, notification of loss; OCWin, notification of reward; vmPFC, ventromedial prefrontal cortex.

relative to current, cocaine users using the same reward-processing task (Patel *et al*, 2013). Additionally, our findings are consistent with other longitudinal data in CD participants performing a cognitive-control task with a pretreatment–posttreatment design (Moeller *et al*, 2012). Notably, the midbrain cluster coordinates reported here during anticipatory reward show significant overlap with those reported previously in which midbrain activity change correlated negatively with explicit cocaine selections on a neuropsychological drug-choice task (Moeller *et al*, 2012). Here, in an exploratory analysis, we extend this finding to link midbrain activity with real-world abstinence using 1-year follow-up data (Figure 1e). Notably, this correlation with the A2Win phase, one reflecting anticipatory activity with motor preparatory effects removed, suggests increased midbrain recruitment during passive anticipation of a rewarding non-drug cue. Nonetheless, the prior and present studies administered two different tasks (drug Stroop and MIDT, respectively) assessing distinct cognitive domains; enhanced midbrain response may therefore represent a biomarker for approach motivation relating to

choice behavior and longer-term cocaine use outcomes in CD. Altogether, these findings suggest the possibility that midbrain-related activation improvement with abstinence could relate to increases in attentional effort occurring with drug discontinuation (Sarter *et al*, 2006). Nevertheless, increased attention may itself represent a form of cognitive incentive, driven either through explicit and/or implicit motivational forces that might increase with recovery (Sarter *et al*, 2006). Future studies might disentangle motivational from attentional or working-memory processes occurring with recovery.

The midbrain contains dopaminergic cell bodies with ascending projections to striatal and thalamic regions responsive to reward-predicting cues (Schultz *et al*, 1997). Although BOLD fMRI cannot measure neurotransmitter activity directly, greater anticipatory activity in this network posttreatment may indicate a recovery of dopamine-related activity and endogenous response to non-drug rewards (Choi *et al*, 2006). Some evidence links dopamine D2 receptor availability with monetary expectation response in CD (Asensio *et al*, 2010), suggesting that resting availability of

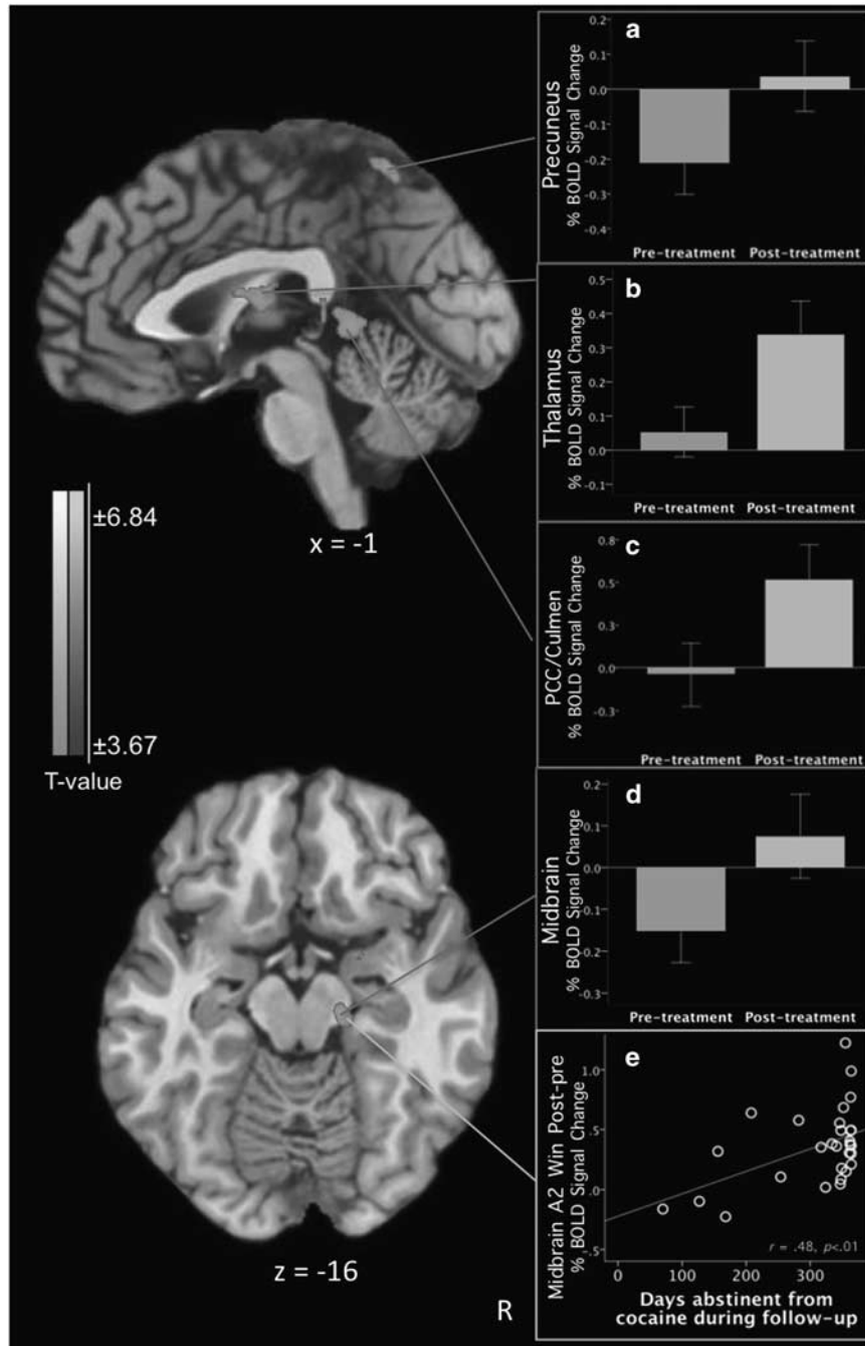


Figure 1 Posttreatment–pretreatment differences on the Monetary Incentive Delay Task in the cocaine-dependent (CD) group ($n = 29$) during the A2 winning phase (A2W, associated with the anticipation of potentially winning money). (a) The percentage of blood-oxygen-level-dependent (BOLD) signal change in the precuneus cluster during pretreatment and posttreatment; (b) BOLD signal change in the thalamus cluster during pretreatment and posttreatment; (c) BOLD signal change in the posterior cingulate (PCC)/culmen cluster during pretreatment and posttreatment; (d) BOLD signal change in the midbrain cluster during pretreatment and posttreatment; (e) scatterplot demonstrating a positive correlation between the midbrain A2W cluster difference and days abstinent from cocaine during follow-up ($r = 0.48$, $p < 0.01$). All contrast maps are thresholded at an uncorrected level of $p < 0.001$ two-tailed and family-wise-error-corrected at $p < 0.05$. Blue color demonstrates areas where subjects show relatively less activation at posttreatment vs pretreatment, and red color indicates where participants show relatively greater activation at posttreatment vs pretreatment. The right side of the brain is on the right. A full color version of this figure is available at the *Neuropsychopharmacology* journal online.

this receptor may predict functional responses to monetary reinforcers. Additionally, these findings provide some index of abstinence duration necessary to observe brain activation changes; increases in dopamine-innervated regions (eg,

midbrain, striatum) occurred following 4 months. Although the midbrain is susceptible to respiratory artifacts during imaging (Raj *et al*, 2001), the pattern of pretreatment–posttreatment activation increases through dopaminer-

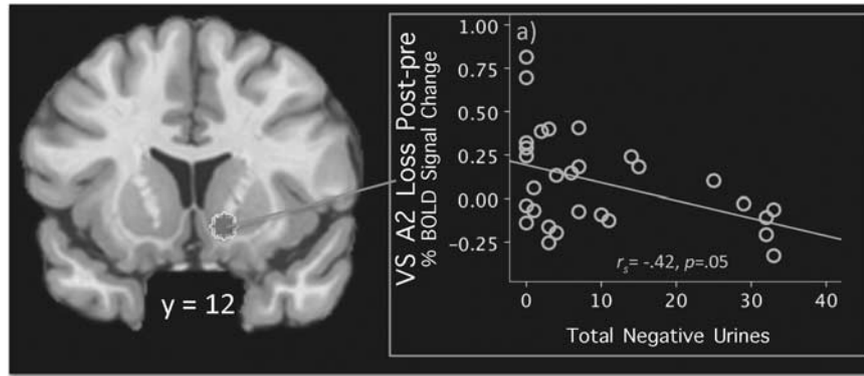


Figure 2 Coronal view of the ventral striatal Region of Interest (ROI) using coordinates reported by Patel *et al* (2013). A blue spot indicates a 5-mm sphere around the ventral striatum (VS) on the right (12, 12, -9) side. Scatterplots depict the percentage of blood-oxygen-level-dependent (BOLD) signal change extracted from the 5-mm ROI during the A2L Loss phase (posttreatment > pretreatment) correlated with (a) the total negative urines during treatment period ($r_s = -.42$, $p = 0.05$). A full color version of this figure is available at the *Neuropsychopharmacology* journal online.

gically-innervated regions, together with the striking overlap in cluster coordinates with a prior pretreatment–posttreatment fMRI study in CD, lend support to this as a biologically significant effect. Consistent with learning theories of addiction underscoring the importance of associative learning mechanisms, our findings highlight that recovery processes may recruit the same reward networks affected by chronic drug use. Furthermore, the plasticity of responsiveness in this circuitry support the goal of many therapeutic interventions in addiction aiming to increase the salience of non-drug-related cues. Future studies can focus on relating these recovery patterns with specific therapies and subjective effects in individuals.

Relationship between functional changes in VS anticipatory processing and abstinence. The VS ROI analyses link negative incentive signaling with individual variation in reduction of cocaine use within treatment as measured by negative urine screens. Our findings relating VS activity changes with abstinence are consistent with the right VS as a core region responding to cocaine cues and drug craving (Kuhn and Gallinat, 2011). Preclinical studies show craving-related VS dopamine signaling instigating drug-seeking behavior (Saunders *et al*, 2013); similarly, our findings link negative incentive VS signaling with cocaine. Notably, links with cocaine use measures occurred during loss anticipation, highlighting the possible role of negative incentive signaling in recovery.

Although increased VS signaling might be expected with reduction of cocaine use, the negative correlation between VS changes and negative urines is nonetheless consistent with findings from two cross-sectional CD MIDT studies linking reduced VS activity with greater abstinence (Bustamante *et al*, 2013; Patel *et al*, 2013). Our VS results extend these findings, demonstrating not only reduced VS activity, but lower VS fluctuations over time relate to stable remission. These findings provide insight into mixed findings in the literature, as cross-sectional designs may capture volatility in VS signaling in current users. These findings hint at dynamic VS dysregulation, particularly to anticipatory cues, as an

important clinical index in CD, potentially related to neurotransmitter bioavailability at different phases of the abuse–abstinence–relapse cycle. For example, actively using individuals may produce a greater dynamic response when challenged, through the presence of drug metabolites (ie, recent cocaine use) synergistically affecting the reward signal or through VS neuroadaptations from recent drug use. Moreover, imbalances in tonic–phasic signaling of cells from chronic drug exposure may dynamically alter the excitatory tone of cells across distinctive recovery phases (Phillips *et al*, 2003); similarly, alterations in VS dopamine receptor availability and affinity are observed in CD and relate to fMRI responses to non-drug rewards 3 years later (Asensio *et al*, 2010; Volkow *et al*, 2007). The data presented here suggest that abstinence-related neurobiological changes occur in similar regions showing neurofunctional alterations at pretreatment and that these might be used to track progress during protracted recovery. Additionally, correlational findings linking VS recruitment with changes in cocaine use provide support for this area as a possible predictor of better outcomes over time.

Test–Retest Differences in HCs

To date, few studies have examined test–retest effects in HCs on the MIDT; therefore, investigating these changes is important in characterizing neural responses associated with repeated task exposure. Differences emerged in the OCWin phase, with a significant reduction in vmPFC activity at retest. Given the ascribed role of the vmPFC in tracking monetary reward outcomes (Knutson *et al*, 2003), these findings suggest diminished outcome-associated reward responses and possibly reduced integration of incentive information with repeated MIDT exposure. These findings were observed in the absence of affective differences to winning outcomes across the two scans.

Differences Between CD and HC

Few prospective studies have examined brain-based mechanisms of change; to our knowledge, this is the first CD study

using fMRI at multiple time points to compare CD–HC differences in reward processing over time. Group differences emerged in the left IFG exclusively during the A2 phases,

whereby CD participants demonstrated significant increases relative to HC participants, who showed decreased activity here. The IFG is implicated in various functions, including

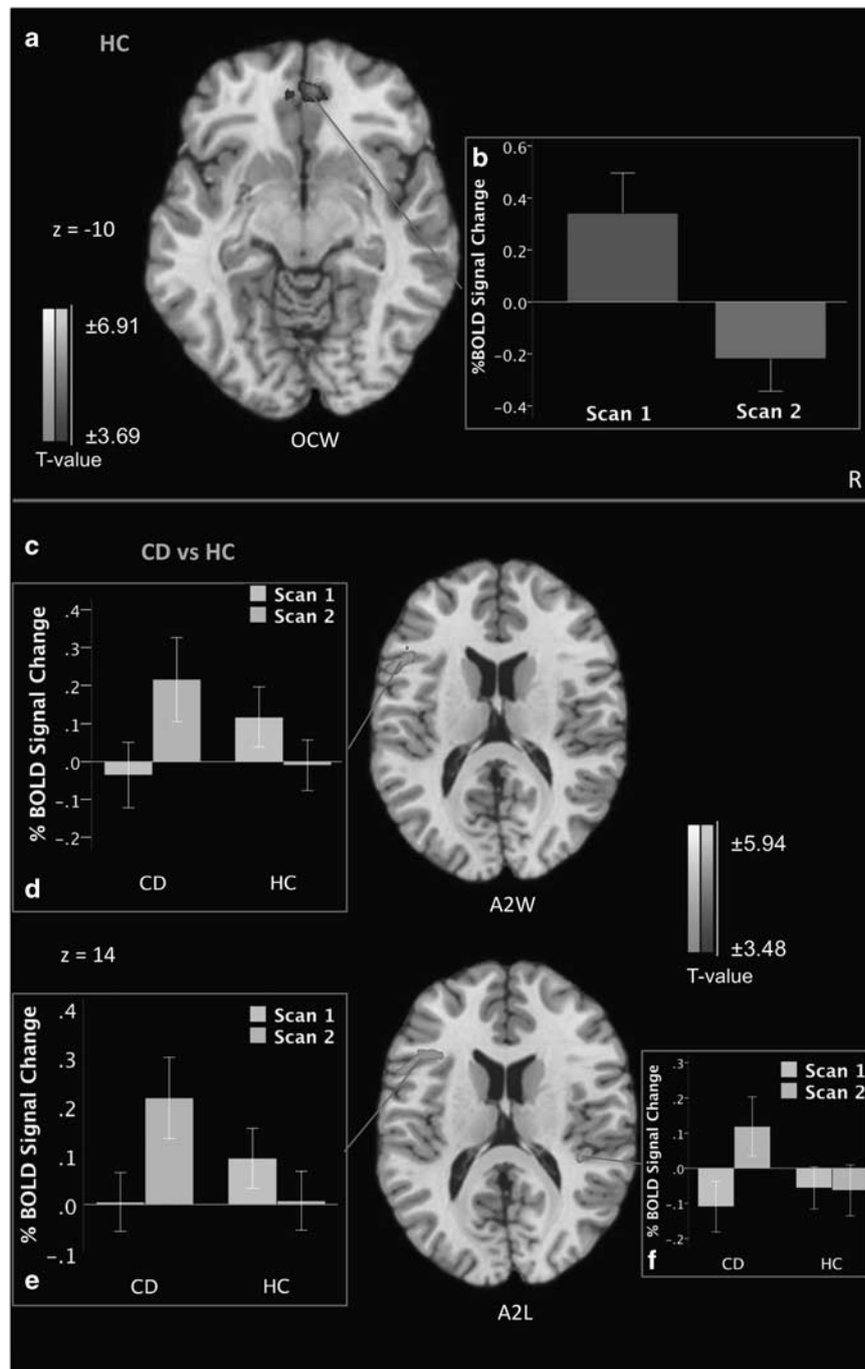


Figure 3 Scan2–Scan1 differences on the Monetary Incentive Delay Task in (a) the HC group ($n=28$) during the outcome winning phase (OCW, associated with the receipt of reward); (b) depicts the percentage of blood-oxygen-level-dependent (BOLD) signal change in the ventromedial prefrontal cortex (vmPFC; $z = -10$) in the HC group for their first and second scans. (c) Scan2–Scan1 contrast between the CD group ($n=29$) and the HC group ($n=28$); (d) depicts the percentage of blood-oxygen-level-dependent (BOLD) signal change in the left inferior frontal gyrus ($z = 14$) in the CD and HC groups for their first and second scans during the A2 winning phase (A2W, associated with the anticipation of potentially winning money); (e) depicts the percentage of blood-oxygen-level-dependent (BOLD) signal change in the left inferior frontal gyrus ($z = 14$) in the CD and HC groups for their first and second scans during the A2 losing phase (A2L, associated with the anticipation of potentially losing money); (f) depicts the percentage of blood-oxygen-level-dependent (BOLD) signal change in the right superior temporal gyrus ($z = 14$) in the CD and HC groups for their first and second scans during the A2 losing phase. All contrast maps are thresholded at an uncorrected level of $p < 0.001$ two-tailed and family-wise-error-corrected at $p < 0.05$. Blue color demonstrates areas where subjects show relatively less activation in the indicated contrast map, and red color indicates where participants show relatively greater activation. The right side of the brain is on the right. A full color version of this figure is available at the *Neuropsychopharmacology* journal online.

attention and behavior monitoring in goal-directed behavior (Corbetta and Shulman, 2002), and is described as a hub for limbic-executive functions (Bari and Robbins, 2013). Rather than subcortical areas, these findings characterize neurofunctional test–retest differences between CD and HC individuals as occurring during anticipatory processing in cortical areas involved in attention and executive control. Continued longitudinal fMRI investigations during recovery will establish whether normalization of functioning occurs between groups or whether dynamic processes persist with sustained abstinence. Comparisons with HC populations are critical for additionally tracking normative neuroadaptive changes occurring over time and further gauging neurobiological recovery profiles in addiction.

Strengths, Limitations, and Future Directions

The current study is novel in demonstrating changes in reward processing occurring in a CD population following interventions and linking these with clinically meaningful abstinence measures. This study also benefits from a comparatively large sample, a within-subject longitudinal design, rigorous fMRI analytical thresholds to reduce the rate of false positives, and further includes hypothesis-driven ROIs informed by the research literature. We report changes in similar/overlapping brain regions with comparable measures as reported by other groups (Bustamante *et al*, 2013; Moeller *et al*, 2012; Patel *et al*, 2013) and in populations with considerably more abstinence, thereby contributing to replication and broader generalizations beyond the current study. Therefore, results from the current study are not only consistent with previous CD study findings but also novel in presenting recovery-related findings; to our knowledge, this is the first study to longitudinally examine generalized reward processing changes in a CD population. Our prospective design using a well-validated reward-processing task contributes to the field's understanding of anticipatory processing fluctuations in addiction. These findings are particularly meaningful in the context of addiction literature, given the significant role of incentive signaling to reward-predicting cues thought to underlie addictive processes. Specifically, our findings of increased reward anticipation in mesocorticolimbic circuitry over time provide some evidence for recovery of generalized reward processing in this population. These findings are distinct from those of Moeller *et al* (2012), one of the few other studies also using a pretreatment–posttreatment design in a CD population. Moeller *et al* (2012) employed a drug Stroop task, which assesses cognitive domains of attention and inhibition specific to drug cue, rather than generalized reward processing. Our findings are also unique from Patel *et al* (2013), who employed a cross-sectional design and focused on loss outcome processing on the MIDT.

The inclusion of an HC group further permits examination of test–retest changes occurring with the MIDT and comparisons with the CD group. The current study examined outcomes across time in individuals participating in a range of treatments; it was not possible to evaluate treatment-specific effects or ascribe changes in neural activity to the effects of reduced cocaine use or specific effects of treatment. Additionally, a recent report demonstrates an inverse relationship between depressive symptoms

and right VS activity on the MIDT (Hagele *et al*, 2015). In the current study, depression scores were not collected in the CD group. Nonetheless, excluding the seven participants with a lifetime history of depression did not alter the correlations between A2L and outcome measures and, in fact, slightly strengthened them.

Gaining a better idea of neurofunctional changes during treatment represents an important first step in understanding the neurobiology of successful abstinence (Garavan *et al*, 2013; Hanlon *et al*, 2013). Additionally, our findings of increased precuneus/PCC activity during A2Win are consistent with a recent meta-analysis citing common effects of pharmacological and cognitive-based interventions in these areas (Konova *et al*, 2013). Finally, while the MIDT version administered dissociates specific phases of reward processing, BOLD fMRI cannot directly gauge neurotransmitter systems underlying changes in the motivational salience signals. Nonetheless, the results have implications for narrowing in on particular mechanisms and therapeutic targets. More research will be important in clarifying whether incentive motivational signals to non-drug cues directly reflect an adapted/recovered capacity to recruit dopamine synthesis. Further, studying VS signal volatility and adaptations over time merits additional investigations and could advance understanding of recovery processes.

CONCLUSIONS

This controlled, prospective study demonstrates a functional enhancement of dopaminergically innervated brain regions occurring specifically during anticipatory processing in a CD group following treatment. An exploratory analysis further linked midbrain activity with abstinence 1 year later. Additionally, functional changes in the VS relate to an objective measure of abstinence (urine screens). Altogether, this is one of the first longitudinal studies demonstrating how incentive signaling reflects recovery processes in CD. These findings suggest that both midbrain and VS responses during anticipatory processing may represent biomarkers for approach motivation relating to abstinence in CD. Although previous studies demonstrate striatal dopamine surges following drug-cue exposure in CD that positively correlate with craving measures (Boileau *et al*, 2007; Volkow *et al*, 2006; Wong *et al*, 2006), here we show that individual changes in non-drug anticipatory responsivity in the striatum relate to abstinence. The current study demonstrates abstinence-related neurofunctional changes during recovery, particularly relating to incentive salience signal readjustments linked to measures of reduced cocaine use over time. These findings have clinical relevance, as the effectiveness of many therapeutic interventions relies on the dynamic response to rewarding non-drug cues (Vocci, 2007). These findings support the idea that over time and with less expensive approaches to imaging, neural data outcome might be applied in the future to more specifically evaluate the clinical impact of particular therapies. Future studies longitudinally investigating individuals at multiple months and even years after treatment could clarify which changes are long-lasting and most predictive of sustained recovery.

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