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Proactive and reactive cognitive control and dorsolateral prefrontal cortex dysfunction in first episode schizophrenia

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\begin{abstract}
Cognitive control deficits have been consistently documented in patients with schizophrenia. Recent work in cognitive neuroscience has hypothesized a distinction between two theoretically separable modes of cognitive control—reactive and proactive. However, it remains unclear the extent to which these processes are uniquely associated with dysfunctional neural recruitment in individuals with schizophrenia. This functional magnetic resonance imaging (fMRI) study utilized the color word Stroop task and AX Continuous Performance Task (AX-CPT) to tap reactive and proactive control processes, respectively, in a sample of 54 healthy controls and 43 patients with first episode schizophrenia. Healthy controls demonstrated robust dorsolateral prefrontal, anterior cingulate, and parietal cortex activity on both tasks. In contrast, patients with schizophrenia did not show any significant activation during proactive control, while showing activation similar to controls during reactive control. Critically, an interaction analysis showed that the degree to which prefrontal activity was reduced in patients versus controls depended on the type of control process engaged. Controls showed increased dorsolateral prefrontal cortex (DLPFC) and parietal activity in the proactive compared to the reactive control task, whereas patients with schizophrenia did not demonstrate this increase. Additionally, patients’ DLPFC activity and performance during proactive control was associated with disorganization symptoms, while no reactive control measures showed this association. Proactive control processes and concomitant dysfunctional recruitment of DLPFC represent robust features of schizophrenia that are also directly associated with symptoms of disorganization.
\end{abstract}

1. Introduction

While decades of research illustrate that schizophrenia is associated with deficits across a wide variety of cognitive domains, including attention, memory, and language, recent theories propose that impaired cognitive control-related dysfunction of the prefrontal cortex may account for many of these findings (Barch and Ceaser, 2012; Lesh et al., 2011). Cognitive control involves online maintenance of goals and task sets to guide adaptive behavior (Miller and Cohen, 2001). An extensive literature suggests that patients with schizophrenia show cognitive control deficits (Cohen and Servan-Schreiber, 1992; Cohen et al., 1996, 1999; Servan-Schreiber et al., 1996) associated with impaired recruitment of the dorsolateral prefrontal cortex (DLPFC) and related circuitry, including the anterior cingulate cortex (ACC) and parietal cortex (Barch et al., 2001; MacDonald and Carter, 2003; MacDonald et al., 2000; Yoon et al., 2008).

Recent work highlights a distinction between proactive and reactive modes of cognitive control, which Braver and colleagues have termed the dual mechanisms of control (DMC) theory (Braver et al., 2007, 2009). Proactive control is conceptualized as maintenance of goal-relevant information to optimally bias attention, perception and response preparation ahead of a cognitively demanding event. In contrast, reactive control reflects transient, ‘on the fly’ engagement of control processes at the onset of challenging task demands. Due to its connectivity with sensorimotor regions, DLPFC plays a central role in the maintenance of goals and rules for action (Asaad et al., 2000; Watanabe, 1990, 1992), which should be reflected in increased DLPC activity during proactive control. Proactive control may depend more on the ability to mount a sustained pattern of neural activity, a well-characterized aspect of prefrontal cortex that is observed during both physiological recording in non-human primates (Goldman-Rakic, 1995) and fMRI in...
Humans (MacDonald et al., 2000). In contrast, reactive control may be associated with more transient neural engagement of not only DLPFC, but also ACC, which is associated with detecting conflict and recruiting DLPFC engagement on a trial to trial basis (Egner and Hirsch, 2005; Kerns et al., 2005; MacDonald et al., 2000). Edwards et al. (2010) investigated proactive and reactive control processes in patients with schizophrenia using a version of the AX Continuous Performance Task (AX-CPT), in which the cue (proactive control) and probe (reactive control) phases were examined separately. Patients with schizophrenia showed more probe-related activity while controls demonstrated greater cue-related PFC activity, suggesting that patients were relying on reactive control. Further, after training patients to recognize and attend to contextual information, they showed a significant shift in brain activation (i.e., increased cue-related activity and decreased probe-related activity) reflecting changing emphasis from reactive to proactive control.

While these data provide evidence for the dissociation between proactive and reactive control, examining these processes within the same task has some limitations. First, reactive control is necessarily linked to contextual processes engaged during the cue, thus the reactive control measure is inherently confounded by the degree to which proactive processes are engaged. This might attenuate reactive activity in controls due to their intact proactive processes. While this may be unavoidable for any task, as individuals often engage context maintenance to guide responding, this limitation may be partially mitigated by choosing tasks specifically biased towards one type of control process. The B-cue trials of the AX-CPT offer an excellent example of proactive control (see Fig. 1 for AX-CPT illustration). During the AX-CPT, subjects make a target response to the probe letter X, only when it follows the cue letter A. All other trials in which X is preceded by any letter other than A (collectively referred to as B-cue trials) require a non-target response. B-cue trials require preparation to inhibit a prepotent response due to the high proportion of AX (target) trials, as one must prepare to inhibit the incorrect, but more frequent, response in order to correctly respond to the subsequent probe letter. In contrast, the single trial Stroop task is uncued and can be biased to either cue-related PFC activity, which proactive processes are engaged. This might attenuate reactive activity in controls due to their intact proactive processes. While this may be unavoidable for any task, as individuals often engage context maintenance to guide responding, this limitation may be partially mitigated by choosing tasks specifically biased towards one type of control process. The B-cue trials of the AX-CPT offer an excellent example of proactive control (see Fig. 1 for AX-CPT illustration). During the AX-CPT, subjects make a target response to the probe letter X, only when it follows the cue letter A. All other trials in which X is preceded by any letter other than A (collectively referred to as B-cue trials) require a non-target response. B-cue trials require preparation to inhibit a prepotent response due to the high proportion of AX (target) trials, as one must prepare to inhibit the incorrect, but more frequent, response in order to correctly respond to the subsequent probe letter. In contrast, the single trial Stroop task is uncued and can be biased towards reactive control by decreasing the frequency of Incongruent trials, as one must prepare to inhibit the incorrect, but more frequent, response due to the high proportion of AX (target) trials, as one must prepare to inhibit the incorrect, but more frequent, response in order to correctly respond to the subsequent probe letter.

2. Material and methods

2.1. Participants

Forty-three first episode patients with schizophrenia spectrum disorders (see Table 1 for diagnoses and medication status at time of testing) were recruited along with 54 control subjects. Of the present sample, 16 schizophrenia and 19 control participants were included in a previous study focused only on the AX-CPT (Yoon et al., 2008). Schizophrenia participants were outpatients within one year of their first psychotic episode. All participants were assessed using the Structured Clinical Interview for the DSM-IV-TR [SCID-I/P; First et al., 2002]. Clinical interviews were conducted by clinicians with masters or doctoral degrees trained to high reliability (kappa > .70; range = .70–1.0). Schizophrenia participants followed longitudinal and diagnoses were confirmed 6 months after ascertainment. Clinical ratings were collected in the schizophrenia sample using the Scale for the Assessment of Negative Symptoms [SANS; Andreasen, 1983]. Scale for the Assessment of Positive Symptoms [SAPS; Andreasen, 1984], and Brief Psychiatric Rating Scale [BPRS; Lukoff et al., 1986]. Three schizophrenia participants had missing data on the SAPS and were not included in symptom analyses using those items (e.g., disorganization). Exclusion criteria for both groups

![Fig. 1](image-url). Task parameters for the a) AX-CPT and b) Stroop. Tasks were presented using E-prime software (http://www.pstnet.com/eprime.cfm) running on an IBM-compatible computer. Presentation of stimuli was pseudorandom, and the first two stimuli for each subject were target (AX) trials for the AX-CPT and Congruent trials for the Stroop.
Table 1
Demographic, clinical, and behavioral characteristics for patients with schizophrenia and healthy controls.

| Characteristic                  | Schizophrenia (n = 43) | Healthy control (n = 54) |
|--------------------------------|------------------------|-------------------------|
| Years of education: mean ± SD  | 12.67 ± 1.91 (9-17)    | 14.09 ± 2.02 (10-20)*   |
| (range)                        |                        |                         |
| Years of parental education:   | 14.87 ± 2.30 (10-19)   | 14.28 ± 2.44 (8-20)     |
| mean ± SD (range)              |                        |                         |
| Gender: n males (%)            | 34 (79%)               | 35 (65%)                |
| Ethnicity: n Caucasian (%)     | 22 (51%)               | 28 (52%)                |
| Handicapped: n left-handed (%) | 0 (0%)                 | 4 (7%)                  |
| WASI IQ: mean ± SD (range)     | 102.12 ± 13.10 (78-131)| 113.63 ± 10.67 (85-135)|
| Diagnosis: n (%)               | 41 (96%)               |                         |
| Schizoaffective                 | 1 (2%)                 |                         |
| Schizophreniform                | 1 (2%)                 |                         |
| Medication status: n (%)       | 15 (35%)               |                         |
| Unmedicated                    | 1 (2%)                 |                         |
| Atypical antipsychotic         | 27 (63%)               |                         |
| Typical and atypical           | 1 (2%)                 |                         |
| Antipsychotic                  |                        |                         |
| Symptom scores: mean ± SD      |                        |                         |
| (range)                        |                        |                         |
| Disorganization symptoms       | 7.03 ± 3.53 (3-16)     |                         |
| Reality distortion symptoms    | 16.07 ± 6.18 (4-29)    |                         |
| Poverty symptoms               | 13.49 ± 5.69 (3-26)    |                         |
| AX-CPT accuracy: mean ± SD     | .94 ± .08              | .98 ± .03               |
| BX reaction time: mean (ms) ± SD| 613 ± 184              | 545 ± 122               |
| AX                             | .92 ± .21              | .84 ± .19               |
| AX                             | .86 ± .14              | .94 ± .08               |
| AX                             | .98 ± .04              | .99 ± .03               |
| AX-CPT reaction time: mean     | .97 ± .03              | .99 ± .02               |
| AX                             | .92 ± .11              | .95 ± .06               |
| AX                             | 685 ± 131              | 622 ± 85                |
| AX                             | 785 ± 158              | 717 ± 120               |

SD, standard deviation.
* p < 0.05.

Included: Wechsler Abbreviated Scale of Intelligence (WASI) IQ score below 70, alcohol or drug dependence or abuse within 3 months before testing, positive urine toxicology screen for illicit drugs, prior head trauma worse than a Grade I concussion, or contraindication to MRI scanning. Healthy controls were excluded for the following additional criteria: any lifetime diagnosis of an Axis I or Axis II disorder or any first-degree relatives with a psychotic disorder. Before testing, a detailed description of the study was provided and written informed consent obtained. The study was approved by the University of California, Davis Institutional Review Board. Subjects completed the AX-CPT and Stroop task during one fMRI session, with the order counterbalanced across subjects. All subjects were paid for their participation.

2.2. Measures and data analysis

Task parameters and visual depictions of the AX-CPT and Stroop are presented in Fig. 1. Briefly, in the AX-CPT, subjects make a target response (index finger button press) to the probe letter X only if it was preceded by the cue letter A. All cues and non-target probes require non-target responses (middle finger button press). Target sequence trials are frequent and set up a prepotent tendency to make a target response when the probe letter X occurs. As a result, non-target sequence trials where any non-A cue (collectively called B-cues) is presented and followed by a probe letter X, require the most cognitive control. In the Stroop task, stimuli consisted of one of three color words (RED, GREEN, and BLUE) that were written in one of three color inks (red, green, and blue). Stimuli could be Congruent (word and ink match) or Incongruent (word and ink do NOT match). Subjects respond with a button press corresponding to the color of the ink of the word.

AX-CPT accuracy and reaction time were examined using ANOVA with task condition (AX, AY, BX, BY) as a within-subjects factor and diagnosis as a between-subjects factor. Accuracy and reaction times were based upon probe responses, which were only analyzed if the subject responded correctly to the cue. D-prime context (d’-context) (Cohen et al., 1999; Swets and Sewall, 1963) was computed from AX hits and BX false alarms and analyzed using an independent samples t-test. Stroop accuracy and reaction time were analyzed using ANOVA with task (Congruent, Incongruent) as a within-subjects factor and diagnosis as a between-subjects factor. To evaluate whether the degree of group differences depended upon the task, difference scores were calculated for each task [i.e., raw AX percent correct (hits) minus BX percent error (false alarms) and Congruent percent correct minus Incongruent percent error] and evaluated in a 2 × 2 ANOVA. Although d’-context is computed to provide continuity with previous studies of the AX-CPT, these raw difference scores provide a similarly derived measure of specific control processes that can be used to compare the AX-CPT to the Stroop. To address concerns that these raw score differences may not be directly comparable, z-transformed difference scores were created using the entire sample and also evaluated in a 2 × 2 ANOVA. Group comparisons on measures that violated sphericity assumptions were adjusted using Greenhouse-Geisser and tests in which equal variances were not assumed.

Notably, comparing two tasks and identifying potential differential deficits in a patient group may be confounded by differences in the measurement properties of the tasks (Chapman and Chapman, 1973, 1978). Therefore, estimates of true score variance for each task and trial type in the control group were computed and compared using the methods described by Chapman and Chapman (1978) in which true score variance is the product of the reliability of the test (coefficient alpha) and the variance of the observed scores.

2.3. Functional imaging parameters and data analysis

Imaging data were obtained using a 1.5 T General Electric MRI scanner. Coplanar T1-weighted and T2-weighted structural images were acquired prior to each fMRI sequence. For the AX-CPT, T2*-weighted echoplanar imaging (EPI) sessions were acquired with the following settings: TR = 2000-msec, echo time = 40-msec, flip angle = 90°, and field of view = 22 cm. Functional images consisted of 24 contiguous and interleaved 4.0-mm axial slices with a 3.4-mm2 in-plane resolution. For the Stroop Task, T2*-weighted EPI sessions were acquired with the following settings: TR = 1500-msec, echo time = 32-msec, flip angle = 90°, and field of view = 22 cm. Functional images consisted of 20 contiguous and interleaved 4.0-mm axial slices with a 3.4-mm2 in-plane resolution and extended 80 mm above to 16 mm below the anterior–posterior commissure line. Preprocessing was completed using Statistical Parametric Mapping-8 (SPM8, http://www.fil.ion.ucl.ac.uk/SPM8), including slice timing correction, spatial realignment, spatial normalization to the EPI Montreal Neurological Institute (MNI) template using a rigid-body transformation followed by non-linear warping, and spatial smoothing using a Gaussian 8-mm full-width half-maximum kernel. Individual fMRI runs were removed from the analysis if translational movement exceeded 4-mm or rotational movement exceeded 3° of within-run movement.

Functional imaging analysis was performed for both event-related tasks in SPM8 with multiple regression in the general linear model.
framework. All task regressors were modeled and only correct responses were included in the reported contrasts. AX-CPT regressors included the A cue, B cue, and each probe (i.e., AX, AY, BX, BY), while Stroop regressors included Congruent and Incongruent trials. Translational and rotational movement data were included as covariates. Group-level random-effect comparisons were performed between groups for the AX-CPT contrast subtracting the A cue from the B cue regressor (AX-CPT B–A contrast) and Stroop task contrast subtracting the Congruent from Incongruent regressor (Stroop I–C contrast). All between-group contrasts were thresholded at the voxel level with $p < 0.01$ and clusters were considered significant if they survived cluster level FWE correction of $p < 0.05$. To examine the interaction between task and group, a mixed-model ANOVA was implemented in SPM8 with task as the within-subject factor (AX-CPT B–A contrast and Stroop I–C contrast) and diagnosis as the between-subject factor.

In addition to whole-brain analyses, a priori hypotheses regarding the DLPC and ACC prompted interrogation of two regions of interest (ROI). The DLPC ROI consisted of a bilateral Brodmann Area 46 anatomical mask obtained from the Wake Forest University PickAtlas (Maldjian et al., 2003, 2004). The second ROI was obtained using bilateral anterior cingulate regions defined in the Automated Anatomical Labeling (AAL) atlas (Tzourio-Mazoyer et al., 2002), with pre-genual portions manually removed to isolate dorsal ACC. Beta weights for each ROI were extracted and analyzed from the AX-CPT B–A and Stroop I–C contrasts and evaluated in separate 2 × 2 ANOVAs with diagnosis and task as between- and within-subject factors, respectively. Given the a priori directional hypothesis that patients would show attenuated activation specifically on the Stroop task in the ACC, a one-tailed t-test was used to test for group differences on this measure. Additionally, in order to accurately interpret patterns of activity in the whole-brain group by task analysis, beta weights were extracted post-hoc from regions surviving whole-brain FWE correction.

### 2.4. Correlation analyses with clinical symptomatology

Planned bivariate correlations (two-tailed, alpha set at $p < 0.05$) were performed between composite symptom measures [disorganization, reality distortion, and poverty symptoms; see Barch et al., 2003] and cognitive/physiological measures (AX-CPT and Stroop accuracy difference scores, DLPCF and ACC beta weights for AX-CPT B–A and Stroop I–C contrasts). Steiger's Z tests (one-tailed) were performed to compare correlations between tasks. Steiger's Z is preferred over Fisher's Z-test as Steiger's Z takes into account the dependency of correlation coefficients that have an index in common (i.e., comparing correlation coefficients between disorganization and an AX-CPT variable to disorganization and a Stroop variable).

### 3. Results

#### 3.1. Demographic results

Participant demographic information is presented in Table 1. The groups did not differ significantly on age, gender, handedness, or parental education. Control subjects completed more years of education ($t = 3.521$, $p = 0.001$) and had higher estimated IQ ($t = 4.724$, $p < 0.001$) than schizophrenia participants.

#### 3.2. Behavioral results

See Table 1 for behavioral data. For AX-CPT accuracy, ANOVA revealed significant main effects of diagnosis, $F(1, 95) = 7.61$, $p = 0.007$; and condition, $F(3, 93) = 48.43$, $p < 0.001$; and a significant diagnosis by condition interaction, $F(3, 93) = 3.58$, $p = 0.017$. Bonferroni-corrected (adjusted alpha level of 0.0125 per test) post-hoc comparisons of individual trial types revealed the predicted pattern of performance, with patients showing worse performance on AX ($t = 2.77$, $p = 0.008$) and BX ($t = 3.29$, $p = 0.002$), but not on AY ($t = 0.569$, $p = 0.571$), or BY ($t = 1.61$, $p = 0.112$). Independent sample t-test of d’-context scores revealed significantly lower d’-context in patients with schizophrenia ($t = 4.22$, $p < 0.001$). ANOVA of reaction time data revealed a significant main effect of diagnosis $F(1, 95) = 6.14$, $p = 0.015$; a significant main effect of condition, $F(3, 93) = 170.18$, $p < 0.001$; and a trend level diagnosis by condition interaction, $F(3, 93) = 2.45$, $p = 0.068$. Bonferroni-corrected (adjusted alpha level of 0.0125 per test) post-hoc reaction time comparisons for each condition revealed a trend towards longer reaction times for schizophrenia patients on BX ($t = 2.48$, $p = 0.015$) and BY ($t = 2.55$, $p = 0.013$), with no significant differences on AX ($t = 2.10$, $p = 0.039$) or AY trials ($t = 1.84$, $p = 0.070$).

For Stroop accuracy, ANOVA revealed significant main effects of diagnosis, $F(1, 95) = 4.89$, $p = 0.029$; and condition, $F(1, 95) = 38.49$, $p < 0.001$; but no significant diagnosis by condition interaction, $F(1, 95) = 1.62$, $p = 0.206$. ANOVA of reaction time data revealed a significant main effect of diagnosis, $F(1, 95) = 7.24$, $p = 0.008$; a significant main effect of condition, $F(1, 95) = 211.02$, $p < 0.001$; but no significant diagnosis by condition interaction, $F(1, 95) = 0.13$, $p = 0.721$. In other words, patients demonstrated lower accuracy and longer reaction times overall compared to controls, but the extent of the performance decrement was not dependent upon the trial type.

ANOVA of accuracy difference scores for the AX-CPT (AX minus BX false alarms) and Stroop (Congruent correct minus Incongruent incorrect) revealed significant main effects of diagnosis, $F(1, 95) = 15.73$, $p < 0.001$; and task, $F(1, 95) = 16.56$, $p < 0.001$; and a significant diagnosis by task interaction, $F(1, 95) = 6.82$, $p = 0.010$ (Fig. 3a). Z-transformed difference scores were complicated by the presence of outliers. Consequently, a Winsorizing procedure (Dixon, 1960; Hastings et al., 1947) was applied to values exceeding the 99th percentile. ANOVA of these z-transformed difference scores revealed a significant main effect of diagnosis, $F(1, 95) = 14.38$, $p < 0.001$; no main effect of task, $F(1, 95) = 0.157$, $p = 0.692$; and a significant diagnosis by task interaction, $F(1, 95) = 3.97$, $p < 0.050$. While the z-transformed measure paralleled the raw difference score measure in terms of a main effect of group and group by task interaction, the main effect of task was not replicated. These results are encouraging and suggest that when z-transformed, the difference scores reflect comparable levels of cognitive control difficulty overall, but still reveal an interaction such that patients with schizophrenia show a decline in performance in the proactive compared to reactive condition. Given that the z-transformed and raw difference scores showed similar results in the interaction, raw difference scores were used for correlation analyses.

#### 3.3. True score variance measurement

Overall, across all trial types, AX-CPT true score variance ($\sigma^2_T = 64.30$) was markedly similar to Stroop true score variance ($\sigma^2_T = 69.98$). Examining individual trial types for the AX-CPT, AX trials showed the highest true score variance ($\sigma^2_T = 23.86$), followed by AY ($\sigma^2_T = 7.58$), BX ($\sigma^2_T = 2.74$), and BY ($\sigma^2_T = 0.25$) trials. For Stroop, Congruent trials ($\sigma^2_T = 22.81$) showed slightly higher true score variance than Incongruent trials ($\sigma^2_T = 15.76$). Given that our main comparisons of interest involve AX hits versus BX false positives in the AX-CPT and Congruent correct versus Incongruent errors, true score variance values for these trial types are the most critical. While AX and Congruent true score variance estimates are very similar, Incongruent true score variance is over five times larger than BX true score variance. Given that tasks with greater true score variance are more likely to (erroneously) show greater differential deficits, the fact that BX trials show lower true score variance argues against the possibility that AX-CPT specific deficits (and relatively intact Stroop performance) in patients with schizophrenia is due purely to differences in measurement properties of the tasks.
3.4. Functional MRI results

3.4.1. Within-group analyses

Table 2 summarizes data for regions with significant activation, while Fig. 2 illustrates uncorrected whole-brain results. In the AX-CPT B–A contrast, controls showed activation in the predicted network of frontal and parietal regions, including bilateral DLPCF, bilateral parietal cortex, and anterior cingulate/supplementary motor area. In contrast, patients did not demonstrate significant suprathreshold activation in this contrast, although they did show activation across this network at lower, uncorrected thresholds.

In the Stroop I–C contrast, controls showed robust activation in many regions typically activated during cognitive control, including bilateral DLPCF, anterior cingulate/supplementary motor area, and left parietal cortex. Patients showed a similar pattern of activation, including bilateral DLPCF, bilateral parietal cortex, and anterior cingulate/supplementary motor area.

To evaluate the potential impact of medication on brain activity, individuals who were currently taking antipsychotic medication were compared to unmedicated subjects at both a priori regions of interest. There was no significant effect of medication on DLPCF activity during the AX-CPT (t = 1.63, p = 0.120; Cue B minus Cue A), DLPCF activity during the Stroop (t = 0.71, p = 0.483; Incongruent minus Congruent), ACC activity during the AX-CPT (t = 1.58, p = 0.121), or ACC activity during the Stroop (t = 0.32, p = 0.753; Incongruent minus Congruent).

3.4.2. Between-group analyses

In the whole-brain AX-CPT B–A contrast, control subjects demonstrated significantly more activity in right DLPCF as well as right inferior parietal cortex compared to patients. Similarly, control subjects demonstrated significantly more activity in the a priori DLPCF ROI compared to patients with schizophrenia (t = 2.45, p = 0.016; Cue B minus Cue A). No significant differences were found in the a priori structurally defined dorsal ACC ROI for the AX-CPT B–A contrast (t = 1.28, p = 0.205; Cue B minus Cue A). No clusters reached significance in the whole-brain between-group comparison of the Stroop I–C contrast. However, the a priori structurally defined dorsal ACC ROI revealed a trend towards greater activity in controls compared to patients in the primary Stroop contrast (t = 1.42, p = 0.080; Incongruent minus Congruent). No significant difference was identified in the DLPCF ROI for the Stroop task (t = 0.47, p = 0.643; Incongruent minus Congruent).

3.4.3. Task by group analyses

The whole-brain task by group interaction analysis revealed significant clusters in bilateral DLPCF and right inferior parietal cortex, with control subjects showing greater increases in activation in the AX-CPT B–A contrast compared to the Stroop I–C contrast, while patients with schizophrenia did not (Fig. 4a). This pattern of results was corroborated in the a priori DLPCF ROI (Fig. 4c), which revealed a significant group by task interaction, F(1, 95) = 6.66, p = 0.011; a trend level main effect of group, F(1, 95) = 3.64, p = 0.059; and no significant main effect of task, F(1, 95) = 1.30, p = 0.257. The a priori ACC ROI revealed no group by task interaction, F(1, 95) = 0.44, p = 0.834; a trend level main effect of group, F(1, 95) = 3.88, p = 0.052; and no significant main effect of task, F(1, 95) = 1.80, p = 0.183. To accurately interpret activity in regions surviving FWE correction in the whole-brain task by group interaction, we also evaluated beta weights extracted from the local maximum coordinates in each significant region (left and right DLPCF and right parietal cortex; see Table 2 for coordinates and Fig. 4 for graphs). Left DLPCF revealed a significant group by task interaction, F(1, 95) = 15.91, p < 0.001; a trend level main effect of group, F(1, 95) = 2.93, p = 0.090; and no significant main effect of task, F(1, 95) = 1.48, p = 0.227. Right DLPCF revealed a significant group by task interaction, F(1, 95) = 10.72, p = 0.001; a main effect of group, F(1, 95) = 4.69, p = 0.033; and no significant main effect of task, F(1, 95) = 2.05, p = 0.155. The right inferior parietal ROI revealed a significant group by task interaction, F(1, 95) = 10.74, p = 0.001; a significant main effect of group, F(1, 95) = 9.05, p = 0.003; and no significant main effect of task, F(1, 95) = 1.89, p = 0.173. In contrast there were no regions in the whole-brain interaction analysis in which patients demonstrated greater increases than controls.

3.5. Associations with clinical symptomatology

Bivariate correlations in the patient group examined the relationship between performance (i.e., AX hits minus BX false alarms and Congruent hits minus Incongruent errors) and DLPCF activity (proactive and reactive). Better AX-CPT performance was significantly associated with higher proactive DLPCF activity (Cue B minus Cue A), r(43) = 0.343, p = 0.025, while Stroop performance was not significantly associated with reactive DLPCF activity (Incongruent minus Congruent), r(43) = -0.228, p = 0.141. No relationships were found with ACC activity. Lower disorganization symptomatology was associated with a higher AX-CPT accuracy difference score, r(40) = −0.56, p < 0.001, as well as higher BOLD recruitment in
DLPFC during the AX-CPT B–A contrast, \( r(40) = -0.33, p = 0.041 \) (Fig. 5). There was no association between disorganization and the Stroop accuracy difference score or BOLD in DLPFC or ACC during the Stroop (all \( p > 0.12 \); Incongruent minus Congruent).

Steiger’s Z transformations determined whether disorganization symptoms were more strongly associated with performance and DLPFC activity on the AX-CPT (Cue B minus Cue A) compared to the Stroop (Incongruent minus Congruent). Behavioral performance on the AX-CPT showed a significantly stronger relationship to disorganization, compared to the relationship between Stroop performance and disorganization \( (p = 0.03) \). There was not a significant difference in the strength of the relationship between disorganization and DLPFC activity during the AX-CPT compared to the Stroop task \( (p = 0.28) \). Notably, the relationship between disorganization and proactive control processes was specific in that correlations with reality distortion, \( r(42) = -0.108, p = 0.50 \), and poverty symptoms, \( r(43) = -0.131, p = 0.40 \), were low.

4. Discussion

We used two well-validated cognitive paradigms to probe the neural substrates supporting reactive and proactive control processes in a large sample of patients with first episode schizophrenia. Behaviorally, patients with schizophrenia generally demonstrated worse performance during the AX-CPT compared to the Stroop task.
on both tasks compared to controls. However, the extent of the performance decrement was much larger for the proactive control measure examining AX hits and BX false alarms compared to the reactive control measure examining Congruent correct and Incongruent error trials. Neuroimaging results revealed robust activation in both reactive and proactive tasks in healthy controls, represented by lateral prefrontal (BA9, BA46, ACC), and inferior parietal regions during the Stroop I–C contrast and prefrontal (BA9, BA46, ACC) and parietal cortex during the AX-CPT B–A contrast. While patients demonstrated activity comparable to controls in the Stroop, activity was markedly reduced during the AX-CPT, with no activity surviving the cluster-wise threshold. More importantly, the interaction analysis of group and task showed that controls increased activity in DLPFC as well as inferior parietal cortex during proactive compared to reactive control, while patients did not show this increase. Taken together these results suggest that schizophrenia individuals demonstrate relatively preserved engagement of the fronto-parietal network during reactive control, but show a reduced ability to increase recruitment in DLPFC and parietal cortex for proactive control.

Additionally, we identified significant relationships between disorganization and both performance and reduced DLPFC activity during proactive control (AX-CPT B–A contrast). These data corroborate the previous work identifying a relationship between disorganization and DLPFC activity (Edwards et al., 2010) and DLPFC connectivity (Yoon et al., 2008). In contrast, no significant relationships were found between disorganization and any metric of reactive control (e.g., Stroop performance, DLPFC I–C activity, or ACC I–C activity). Furthermore, the relationship between disorganization and AX-CPT performance was stronger than the relationship to Stroop performance, suggesting that disorganization is more strongly associated with proactive control processes. These data, taken in the context of the existing literature, suggest that while reactive control deficits have been identified on the Stroop [for review see Henik and Salo, 2004], proactive control processes may be a more robust link to disorganized clinical symptomatology and underlying neuropathophysiology.

These data shed additional light on the pathophysiology of impaired cognition in schizophrenia in several ways. First, they add to an already substantial literature identifying DLPFC impairment in schizophrenia [see Glahn et al., 2005; Minzenberg et al., 2009 for meta-analytic reviews]. Consistent with recent work (Edwards et al., 2010; Perlstein et al., 2003; Yoon et al., 2008), between-group comparisons revealed significantly reduced DLPFC recruitment during the AX-CPT in schizophrenia individuals. Our findings of adequate recruitment of DLPC and trend level hypoactivation of ACC during reactive control must be considered in the context of some inconsistency in the literature, with some studies of the Stroop revealing PFC hyperactivation in patients (Weiss et al., 2003) and others identifying PFC hypoactivation (Carter et al., 1997; Yucel et al., 2002). These inconsistencies may partly be the result of variability in task design, with some studies including neutral stimuli, presentation in block- or event-related designs, and modifications of the task in which the subject has to explicitly identify whether the word and color are congruent or not as opposed to identifying the ink color with a response. Stage of illness may also play a role as we have previously reported robust ACC decreases using a similar design to the present study in chronic patients (Kerns et al., 2005). Second, our data provide additional support for the dual mechanisms of control theory proposed by Braver et al. (2007). Notably, these data suggest relatively intact reactive control in patients with schizophrenia and imply that prefrontal control deficits in schizophrenia reflect a stronger loading on proactive control processes, reflected in decreased fronto-parietal recruitment. Reduced recruitment of PFC is consistent with the model we presented in a recent review, which theoretically links cellular abnormalities in the PFC with altered inter-regional cortical connectivity, cognitive control dysfunction, and disorganization (Lesh et al., 2011). Sustained activity in the PFC in non-human primates performing working memory tasks depends upon both dopamine (DA) and norepinephrine (NE), modulating neuronal activity through D1 and alpha 2 adrenergic receptors, respectively (Arnstien and Li, 2005; Arstniet et al., 1988; Brozoski et al., 1979; Cai and Arnstien, 1997). GABA-ergic interneurons are also integral to this process (Gonzalez-Burgos et al., 2010; Lewis et al., 2008). The
present data suggest that mechanisms supporting the sustained PFC activity needed for proactive control may be particularly disrupted in schizophrenia. Treatments that remediate these processes, including those that target catecholaminergic and GABA-ergic function in local circuits in this region of the brain, remain promising as interventions to enhance cognitive function in schizophrenia.

One limitation of the current study is the potential effect of medication on fMRI measures. Current evidence suggests that medication effects are not the primary contributor to reduced prefrontal BOLD response during higher-order cognition (Honey et al., 1999; Snitz et al., 2005). Additionally, in the present study, there were no differences between medicated and unmedicated patients in DLPFC and ACC activity across tasks. Finally, since the patients in the present study were very early in the course of illness, it is likely that decreased BOLD response of DLPFC during proactive control is robust and unrelated to the long-term effects of medication exposure or illness chronicity. Another potential limitation is the use of two different tasks, which may have differences in measurement properties that complicate interpretation of a differential deficit. To examine this issue, true score variance values were computed for each task and found to be markedly similar. Furthermore, when examining trial types that were most critical to evaluating proactive and reactive control (i.e., trial types used to compute difference measures), results indicated that Incongruent trials on the Stroop had much higher true score variance than BX trials on the AX-CPT. These data suggest that patients should be more likely to show a performance deficit on the reactive contrast of interest, which contradict our findings of more pronounced proactive deficits, and provide some evidence that our results are not due to differences in measurement properties of the tasks. Finally, differences in scanning parameters between the AX-CPT and Stroop raises the concern that reduced coverage and/or a decreased signal-to-noise ratio (SNR) in the Stroop may have influenced the results. In terms of coverage, within-group results for the Stroop show full coverage of PFC and parietal regions and only voxels that are present in all subjects across both tasks are included in the ANOVA. Differences in the TR and TE raise potential concerns about differences in SNR between the tasks. Although a theoretical SNR can be calculated, it remains difficult to quantify the end result of a lowered TR (which should lower SNR) and a lower TE (which should raise SNR) in the Stroop compared to the AX-CPT. While there may be concerns about decreased SNR in the Stroop, both groups showed robust activation on the Stroop I–C contrast, with peak t-values much higher than the AX-CPT B–A contrast. Additionally, our main findings represent a group by task interaction primarily driven by lower activity in the schizophrenia sample during the AX-CPT with high and relatively comparable activity in the Stroop for both groups. We might anticipate lower SNR in the Stroop to contribute to a main effect of task. However, there was no main effect of

Fig. 5. Correlations between disorganization symptom scores in patients with schizophrenia and a) AX-CPT accuracy difference score, b) DLPFC mean beta values during the AX-CPT task (B–A contrast), c) Stroop accuracy difference score, and d) DLPFC mean beta values during the Stroop task (I–C contrast).
task present in the whole-brain ANOVA or any of the a priori or post-hoc ROIs. Consequently, it would be unlikely for these differences in scanning parameters to account for the specific pattern of results we present.

4.1. Conclusions and future directions

In summary, the current study contributes to our understanding of cognitive control deficits in schizophrenia by highlighting dissociable processes that provide a more precise understanding of the cognitive and neural mechanisms underlying impaired cognition in the illness. Results suggest that proactive control and associated fronto-parietal dysfunction may represent a more robust marker of disease pathology associated with the clinical presentation of the disorder (i.e., disorganization) in first episode patients. Future studies may leverage these findings by tailoring cognitive training paradigms, neurostimulation and medication development to specifically target proactive control mechanisms in the prefrontal cortex. Future studies should also explore how stage of illness plays a role in the integrity of proactive and reactive control since alterations in elements of reactive control such as the ACC appear more robust in published studies involving chronic patients. Finally, future studies should focus on whether this profile of more pronounced proactive control deficits is also present in other psychiatric populations (i.e., mood disorders and autism) who present with impaired cognitive control (Pompei et al., 2011; Solomon et al., 2009; Strakowski et al., 2005; Wagner et al., 2006).

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