Cross-diagnostic comparison of visual processing in bipolar disorder and schizophrenia

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Abstract

Patients with Schizophrenia (SZ) show deficits across various stages of visual information processing. Whether patients with Bipolar Disorder (BD) exhibit these deficits is unclear. In this study, we conducted a detailed comparison of specific stages of early visual perception in BD and SZ. Forty-three BD patients, 43 SZ patients, and 51 matched healthy control subjects (HC) were administered three visual processing paradigms emphasizing: 1) an early stage of object formation (location backward masking), 2) a middle stage of object substitution (four-dot backward masking), and 3) a later stage at the perception—attention interface (rapid serial visual processing (RSVP) task eliciting the attentional blink). SZ performed significantly worse than BD and HC on location and four-dot masking. BD did not significantly differ from HC on either masking task. Both patient groups performed significantly worse than HC on the RSVP task; unlike SZ, BD did not show a significant attentional blink effect compared to HC. Our results indicate that BD patients were intact at the early and middle stages of visual processing (object formation and substitution) but intermediate between the SZ and HC groups at a later processing stage involving perceptual and attentional processes (RSVP task). These findings suggest that SZ is characterized by a diffuse pathophysiology affecting all stages of visual processing whereas in BD disruption is only at the latest stage involving higher order attentional functions.

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1. Introduction

Bipolar disorder (BD) and schizophrenia (SZ) share many features, including common genetic and psychosocial risk factors (Alloy et al., 2005; Lichtenstein et al., 2009). Impairment in cognition and functional outcome is evident in both disorders, although these deficits are milder and less widespread in BD than in SZ (Bora et al., 2009; Lee et al., 2013). SZ is characterized by deficits in early visual information processing (Butler et al., 2001; Rund et al., 2004). These deficits are associated with impaired higher-level cognition, negative symptoms, and poor functional outcome (Brittain et al., 2010; Green et al., 2012; Rassovsky et al., 2011), and are also linked to specific neural processes (Green et al., 2011a). While visual processing deficits are well-documented in SZ, few studies have examined visual processing integrity in BD.

Visual perception in psychopathology has been frequently assessed with two types of paradigms: visual masking and rapid serial visual processing (RSVP). Visual masking affects two different stages of processing defined by the timing of the mask’s effects on a target: an early object formation stage and a later object substitution stage (Enns, 2004). Masking at the object formation stage occurs roughly between 0 and 100 ms when the target and mask fuse together to create an integrated composite that is hard to identify. A later stage of processing (100–200 ms) has been assessed via object substitution masking, in which the target percept is replaced by the mask before it reaches awareness. Object substitution masking is thought to interrupt reentrant processing between lower and higher neural levels that are necessary to refine a percept (Enns and Di Lollo, 2000). Four-dot masking is thought to work exclusively through object substitution (Enns and Di Lollo, 1997). Finally, a later
stage of visual processing involves interactions between perception and attention and is assessed with RSVP paradigms. During this stage (approximately 200–500 ms) a working memory representation of the first target is established and there is temporary inattention or “attentional blink” (AB) for processing of a second target that appears later (Bachmann and Hommuk, 2005). While these paradigms are designed to emphasize a particular stage of processing they are not always able to rule out the influence of other perceptual or attentional stages of processing.

Visual processing abnormalities have been found in BD. For example, one study (MacQueen et al., 2001) reported impaired object identification and visual location masking in euthymic outpatients with BD. Another (Duffy et al., 2009) found that euthymic BD outpatients with a history of psychosis made more errors than those without a prior history of psychosis on masking. Note that MacQueen et al.’s study only included BD I patients who were not taking antipsychotic medications and that both studies used letters as target stimuli in their masking paradigms. These reports suggest that individuals with BD have deficits in visual perception, but they did not make comparisons with SZ.

Few studies have directly compared visual processing in BD and SZ, and the findings have been mixed. Some studies with hospitalized BD patients reported masking impairment in BD during and soon following a manic episode (Fleming and Green, 1995) that was comparable in magnitude to that seen in SZ (Green et al., 1994). Two studies compared stable outpatients with BD and SZ and reported that BD patients were indistinguishable from non.psychiatric controls on a location masking task (Goghari and Sponheim, 2008; Sponheim et al., 2013), but SZ patients were impaired. Another study reported that BD outpatients showed impairment on an object identification task that was comparable to that of SZ patients (Tam et al., 1998). A more recent study (Chkonia et al., 2012) found deficits on a shine-through masking paradigm in both SZ and BD.

Inconsistency in findings of visual impairment in BD may be due to differences in sample characteristics, task parameters, or the specific visual processing stage that is being assessed. Reliable deficits have been found in SZ, extending from early stages of object formation to later stages of object substitution (Green et al., 2011b) to the interface between perception and attention (Mathis et al., 2011). However, the earliest feedforward stage of visual processing (i.e., from retina to V1) appears to be intact (Jahshan et al., 2012). To our knowledge, no study has examined the pattern of performance of BD and SZ patients as one progresses through visual processing stages.

The aim of this study was to conduct a detailed cross-diagnostic comparison of specific stages of visual processing in stable outpatients with BD and SZ, relative to well-matched healthy controls. Three visual processing measures were administered targeting: 1) an early stage of object formation (location backward masking, first 0–100 ms), 2) a middle stage of object substitution (four-dot masking, 100–200 ms), and 3) a later stage at the perception–attention interface (RSVP task, 200–500 ms).

2. Method

2.1. Participants

The sample consisted of 43 patients with BD I or II, 43 patients with SZ (6 schizoaffective), and 51 healthy controls (HC). SZ and HC participants were a subset drawn from a much larger sample, and data from the complete sample were previously published (Green et al., 2012, 2011b; Mathis et al., 2011). These subsamples were selected to be matched to the BD sample on age, gender, and parental education. Data from the BD patients have not been published previously. The test parameters were identical for all samples.

SZ patients were recruited from outpatient clinics at the Veterans Affairs Greater Los Angeles Healthcare System (VAGLAHS) and board-and-care residences in the community. BD patients were recruited from mood disorder clinics at the University of California, Los Angeles (UCLA) and the VAGLAHS. Diagnosis was based on the Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I; First et al., 1997). Of the patients with BD, 29 were diagnosed with BD I (7 with a history of psychosis) and 14 with BD II. All patients were considered to be clinically stable, with no medication changes in the past six weeks, no inpatient hospitalization in the past three months, and no changes in housing in the past two months. Additionally, BD patients needed to be out of mood episode. Controls were recruited through internet advertisements and screened with the SCID-I and SCID-II (First et al., 1996). Exclusion criteria for controls were: any lifetime psychotic disorder, bipolar disorder, recurrent depression, substance dependence, paranoid, schizotypal, or schizoid personality disorder, or a reported history of psychotic disorder (including schizophrenia but not bipolar disorder) among first-degree relatives.

Additional exclusion criteria for all groups included being younger than 18 or older than 60 years, an IQ below 70 based on review of medical records, meeting diagnostic criteria for substance dependence in the past 6 months or abuse in the past month, having an identifiable neurological disorder, seizures, history of head injury or loss of consciousness for more than one hour, having less than 20/40 vision as assessed using the Snellen eye chart, or being insufficiently fluent in English, determined by the participant’s ability to understand the consent form. All participants gave written informed consent after receiving a detailed explanation of study procedures in accordance with procedures approved by the Institutional Review Boards at UCLA and VAGLAHS.

For SZ patients, 37 were receiving second-generation antipsychotic medications, 1 first-generation and 3 both types at the time of testing. One was not taking medication and 1 was missing medication information. For BD patients, most were receiving more than one psychoactive drug: 27 were taking antipsychotic medications, 21 were taking anticonvulsants, 12 were taking lithium, and 17 were taking antidepressants. Two were not taking medications and 1 was missing medication information.

2.2. Assessments

2.2.1. Clinical ratings

SZ received the 24-item UCLA version of the Brief Psychiatric Rating Scale (BPRS; Ventura et al., 1993b) and the Scale for the Assessment of Negative Symptoms (SANS; Andreasen, 1984). We report the positive symptoms and depression/anxiety factors, as well as the total score for the BPRS (Kopelowicz et al., 2008) in Table 1. BD received the Hamilton Depression Rating Scale (HDRS; Hamilton, 1960) and the Young Mania Rating Scale (YMRS; Young et al., 1978). Clinical assessments were conducted by interviewers trained to reliability through the Treatment Unit of the Department of Veterans Affairs VISN 22 Mental Illness Research, Education, and Clinical Center based on established procedures (Ventura et al., 1993a, 1998).

2.2.2. Visual perceptual measures

For all three tasks, participants sat 1 m from a 17 inch cathode ray tube monitor with a refresh rate of 160 Hz (6.25 ms per screen sweep). Stimuli were presented using E-Prime 1.1 (Psychological Software Tools, Pittsburgh, PA). Participants provided verbal responses to the experimenter who then recorded those responses.
2.2.2.1. Location backward masking. This procedure has been described in detail elsewhere (Green et al., 2011b). In this task the target consisted of a square with a notch that could appear at the top, bottom, or left side of the square. The target could appear at one of four different locations, arranged in a notional square, on the screen. The masking stimulus consisted of a pattern of squares that occupied every possible target location. Targets measured $0.27 \times 0.27 \text{C}$ visual angle and the mask measured $2.01 \times 2.01 \text{C}$. Examples of the stimuli are shown in Fig. 1A. The target was presented for 12.5 ms and the mask for 25 ms. Backward masking (mask following target) was assessed using 7 stimulus onset asynchronies (SOAs) ranging from 0 (simultaneous target and mask onset) to 75 ms. Twelve trials were presented for each SOA. Performance is lowest at the SOA of 0 ms and increases with increasing SOAs. Each trial started with a 400 ms fixation cross followed by a 500 ms blank screen. Participants reported in which one of the four quadrants the target appeared. Prior to the masking task, target contrast was set to each subject’s threshold so that all subjects identify the unmasked target with 84% accuracy. The thresholding procedure was a three up one down staircase procedure where the contrast of the target was increased or decreased.

Table 1
Demographic and clinical characteristics.

<table>
<thead>
<tr>
<th></th>
<th>HC (N = 51)</th>
<th>BD (N = 43)</th>
<th>SZ (N = 43)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (Mean/SD)</td>
<td>42.0 (9.1)</td>
<td>42.7 (10.7)</td>
<td>45.6 (7.8)</td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>32:19</td>
<td>25:18</td>
<td>24:19</td>
</tr>
<tr>
<td>Personal education</td>
<td>14.5 (1.7)</td>
<td>14.4 (2.4)</td>
<td>12.5 (1.8)</td>
</tr>
<tr>
<td>(Mean/SD)**</td>
<td>14.2 (2.7)</td>
<td>14.9 (3.1)</td>
<td>14.1 (2.8)</td>
</tr>
<tr>
<td>Race (% White)</td>
<td>54.9%</td>
<td>55.8%</td>
<td>37.2%</td>
</tr>
<tr>
<td>BPRS positive symptoms</td>
<td>–</td>
<td>8.0 (1.4)</td>
<td>15.6 (6.4)</td>
</tr>
<tr>
<td>(Mean/SD)**</td>
<td>–</td>
<td>8.3 (3.7)</td>
<td>8.1 (3.4)</td>
</tr>
<tr>
<td>BPRS depression/Anxiety</td>
<td>–</td>
<td>33.2 (7.3)</td>
<td>44.7 (10.5)</td>
</tr>
<tr>
<td>(Mean/SD)**</td>
<td>–</td>
<td>7.3 (6.8)</td>
<td>7.4 (3.5)</td>
</tr>
<tr>
<td>Total BPRS (Mean/SD)**</td>
<td>–</td>
<td>3.1 (4.2)</td>
<td>–</td>
</tr>
<tr>
<td>Total SANS (Mean/SD)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Total HDRS (Mean/SD)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Total YMRS (Mean/SD)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

BPRS = Brief Psychiatric Rating Scale; SANS = Scale for the Assessment of Negative Symptoms; HDRS = Hamilton Depression Rating Scale; YMRS = Young Mania Rating Scale. **p < 0.001.

Fig. 1. Examples of the stimuli used in the masking (A) and RSVP (B) paradigms.
2.2.2.2. Four-dot masking. This procedure has been described in detail elsewhere (Green et al., 2011b). In this task, four potential targets appeared in a notional square followed by a mask that surrounds one potential target and cues which target is to be identified. The targets were four squares with a notch missing from one side. The mask consisted of four dots that surround, but do not touch, one of the potential targets. Each potential target measured 1.55 x 1.55° of visual angle and was arranged in a square of 4.58 x 4.58°. The four-dot mask measured 2.23 x 2.23° of visual angle and each dot in the mask subtended 0.23 x 0.23°. Examples of the stimuli are shown in Fig. 1A. The target was presented for 25 ms and the mask was presented for 37.5 ms. Performance typically decreases with increasing SOAs, unlike location masking. Twelve trials were presented for each SOA. Each trial started with a 450 ms fixation cross ms followed by a 500 ms blank screen. Participants reported the direction of the notch (up, side, or down) of the target that was surrounded by the four-dot mask.

2.2.2.3. RSVP task. This procedure has been described in detail elsewhere (Mathis et al., 2011). This paradigm consists of a single target RSVP task that measures basic visual perception and a dual target task that elicits the attentional blink (AB) effect. The AB typically occurs when the interval between T1 and T2 is roughly 200–500 ms, resulting in decrease in accuracy for identifying T2 (Dux and Marois, 2009). For both tasks, each trial began with a 500 ms fixation cross followed by a 400 ms blank screen. Each stimulus subtended 2° of visual angle and was displayed for 62.5 ms with an inter-stimulus interval of 25 ms. In the single target task, a target (letter) was presented among a rapid stream of distracting stimuli (numbers) and participants were asked to identify the letter. In the dual target task, two targets (T1 and T2) were presented either with no intervening distractors (lag 1) or separated by 1 (lag 2), 2, 3, 4, 5, 7, 9, or 11 (lag 12) distractors. Ten trials were administered in the single target task and ten trials for each lag in the dual target task. Examples of the stimuli are shown in Fig. 1B. In both tasks after each trial a screen displaying all of the possible targets appeared.

2.3. Data analysis

One-way analyses of variance (ANOVA) and chi-square tests were used to assess group differences for continuous and categorical demographic variables, respectively. A one-way ANOVA was conducted for unmasked location performance. We performed a 3 (group) x 7 (SOA) ANOVA for backward masking and a 3 (group) x 8 (SOA) ANOVA for four-dot masking. For the RSVP task, we conducted a one-way ANOVA for the single target task, and a 3 (group) x 9 (lag) ANOVA for the dual target task. The dependent variable was the conditional probability for correctly identifying the second target given the correct identification of the first target: P(T2/T1) (Chun and Potter, 1995).

We also calculated the suppression ratio (SR) (cf. Cheung et al., 2002) at each lag to assess the AB effect after controlling for group differences on the single target RSVP task. The SR is the degree of performance change in the dual target task relative to the single target task, with scores ranging from 0 to 1. The SR was calculated as follows: SR at lag X = P(T2/T1) at lag X divided by [P(T1) on single target task + P(T2/T1) at lag X], and analyzed with a 3 (group) x 9 (lag) ANOVA.

Least significant difference (LSD) tests were conducted as post hoc analyses to follow up on significant main effects. We performed independent samples t-tests within the BD group to examine performance differences between 1) BD I versus BD II, 2) BD I with versus without history of psychosis, 3) those taking versus not taking lithium, and 4) those taking versus not taking antipsychotic medications. Lastly, we conducted Pearson’s correlations between each task and symptom ratings within each patient group.

3. Results

3.1. Demographic and clinical characteristics

Demographic and symptom ratings can be seen in Table 1. The groups were matched on age, gender distribution, race, and parental education. The groups differed on personal education (F[2, 134] = 14.32, p < 0.001) with SZ having significantly fewer years of education than both BD and HC (p’s < 0.001). SZ had more severe overall symptoms (BPRS: t[82] = 5.78, p < 0.001) than BD. BD had minimal to mild levels of depressive and manic symptoms. We examined gender as a factor but found no significant main effects of gender or group x gender interactions in any of the ANOVAs and decided to exclude it from further analyses.

3.2. Visual processing

3.2.1. Location backward masking

Results are shown in Fig. 2A and Table 2. The ANOVA revealed significant main effects of group (F[2, 134] = 4.79, p = 0.01) and SOA (F[6, 804] = 32.65, p < 0.001) but no significant group x SOA interactions in any of the ANOVAs and chi-square tests. The targets were four squares with a notch missing from one side. The mask consisted of four dots that surround, but do not touch, one of the potential targets. Each potential target measured 1.55 x 1.55° of visual angle and was arranged in a square of 4.58 x 4.58°. The four-dot mask measured 2.23 x 2.23° of visual angle and each dot in the mask subtended 0.23 x 0.23°. Examples of the stimuli are shown in Fig. 1A. The target was presented for 25 ms and the mask was presented for 37.5 ms. Performance typically decreases with increasing SOAs, unlike location masking. Twelve trials were presented for each SOA. Each trial started with a 450 ms fixation cross ms followed by a 500 ms blank screen. Participants reported the direction of the notch (up, side, or down) of the target that was surrounded by the four-dot mask.

We also calculated the suppression ratio (SR) (cf. Cheung et al., 2002) at each lag to assess the AB effect after controlling for group differences on the single target RSVP task. The SR is the degree of performance change in the dual target task relative to the single target task, with scores ranging from 0 to 1. The SR was calculated as follows: SR at lag X = P(T2/T1) at lag X divided by [P(T1) on single target task + P(T2/T1) at lag X], and analyzed with a 3 (group) x 9 (lag) ANOVA.

Least significant difference (LSD) tests were conducted as post hoc analyses to follow up on significant main effects. We performed independent samples t-tests within the BD group to examine performance differences between 1) BD I versus BD II, 2) BD I with versus without history of psychosis, 3) those taking versus not taking lithium, and 4) those taking versus not taking antipsychotic medications. Lastly, we conducted Pearson’s correlations between each task and symptom ratings within each patient group.
interaction \( F[12, 804] = 1.53, p = 0.11 \). The main effect of group was because SZ performed significantly worse than HC \( (p = 0.02, d = 0.49) \) and BD \( (p = 0.004, d = 0.67) \). BD did not significantly differ from HC \( (p = 0.43, d = -0.15) \).

Although all groups’ unmasked performance was near ceiling, there was a significant difference on this measure \( F[2, 134] = 3.35, p = 0.04 \) with SZ performing significantly worse than HC \( (p = 0.01, d = 0.54) \) and BD \( (p = 0.03, d = 0.49) \). The BD and HC groups did not differ \( (p = 0.70, d = 0.08) \). Means and standard deviations for the average across SOAs are reported in Table 2.

### 3.3. Effect sizes

To examine the magnitude of difference in performance between the patient groups and controls as one progresses through visual processing stages, we computed effect sizes for each task. The SZ group showed medium effect sizes on the masking tasks and large effect sizes on the RSVP tasks (Fig. 4). The BD group followed a different pattern, namely small effect sizes on the masking tasks, a medium effect size on the RSVP single target task, and a small-medium effect size on the dual target task.

### 3.4. Subgroup analyses with BD

We found no significant differences on any of the visual processing measures between BD I versus BD II patients (all \( p's > 0.20 \)), BD I patients with versus without history of psychosis (all \( p's > 0.25 \)), BD patients taking versus not taking lithium (all \( p's > 0.12 \)) and BD patients taking versus not taking antipsychotic medications (all \( p's > 0.19 \)).

### 3.5. Correlations with clinical symptoms

We found no significant associations between the visual processing measures and BPRS in both groups, SANS in the SZ group or HDRS/YMRS in the BD group.

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**Table 2**

<table>
<thead>
<tr>
<th>Location masking (%) accuracy</th>
<th>HC ((N = 51))</th>
<th>BD ((N = 43))</th>
<th>SZ ((N = 43))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Backward masking</td>
<td>50.96 (16.67)</td>
<td>53.50 (16.01)</td>
<td>43.53 (13.74)</td>
</tr>
<tr>
<td>Unmasked performance</td>
<td>96.57 (8.02)</td>
<td>95.00 (10.15)</td>
<td>91.28 (11.92)</td>
</tr>
<tr>
<td>Four-dot masking (# correct)</td>
<td>6.72 (1.45)</td>
<td>6.61 (1.29)</td>
<td>5.94 (1.43)</td>
</tr>
<tr>
<td>Attentional blink</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Single target task</td>
<td>0.85 (0.09)</td>
<td>0.78 (0.15)</td>
<td>0.73 (0.15)</td>
</tr>
<tr>
<td>Dual target task</td>
<td>0.59 (0.21)</td>
<td>0.54 (0.21)</td>
<td>0.40 (0.19)</td>
</tr>
<tr>
<td>Suppression ratio (lags 2–5)</td>
<td>0.39 (0.09)</td>
<td>0.36 (0.11)</td>
<td>0.32 (0.11)</td>
</tr>
</tbody>
</table>

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**Table 2 notes**

\( a \) Significantly different from HC group \((p < 0.05)\).

\( b \) Significantly different from HC group \((p < 0.001)\).

\( c \) Significantly different from BD group \((p < 0.05)\).
patients with BD and SZ on specific stages of visual processing. The effect sizes between the SZ and control groups were medium to large across tasks. The BD group did not significantly differ from the control group on the masking tasks but showed medium and small to medium effect-size deficits on the RSVP single target task and dual target task, respectively. Both patient groups showed larger effect sizes on the task with the latest stage of processing (RSVP), especially on the single target task. These results suggest that deficits in SZ are apparent at early and later stages of visual processing, whereas processing is intact in BD with deficits only appearing at later stages. When subgroups of BD were considered, there were no significant differences on any of the visual processing measures between patients with BD I versus BD II disorder, BD I with versus without history of psychosis, those taking versus not taking lithium, or those taking versus not taking antipsychotic medications.

The effects in BD patients are consistent with two other studies of visual processing showing intact early processing using a location masking task (Goghari and Sponheim, 2008; Sponheim et al., 2013); however, our results are inconsistent with those from two different studies that found impaired processing in BD using an object identification task (Tam et al., 1998) and a shine-through masking paradigm (Chkonia et al., 2012). The paradigms in those two studies differed from our masking tasks in several aspects (e.g., only 2 SOAs, pairs of digits as targets, discrimination threshold procedure). Moreover, Chkonia et al.’s BD sample was more symptomatic and included inpatients. Our findings extend the previous findings by showing that BD patients are comparable to controls on early and middle stages (i.e., object formation and object substitution, respectively).

In terms of the RSVP task (not included in the previous studies), the BD group significantly differed from the other two groups on the single and dual target tasks. BD patients’ performance did not seem to worsen with more complex task demands as seen by a smaller effect size for the dual target task versus the single target task. Although BD patients performed poorly on the RSVP tasks, the lack of a significant difference in the suppression ratio between the BD and control groups suggests that the more pronounced attentional blink effect in BD is largely due to deficits in the single target task. Therefore, the BD group’s reduced accuracy at detecting the second target is not related to the attentional blink itself but rather to a general deficit in sustaining attention or maintaining the target in a working memory stage that occurs after object formation.

BD patients’ deficient target identification in the single target task in the context of intact masking could also be explained by differences between the masking and RSVP paradigms. These measures clearly involve different demands as the SZ patients’ performance worsened (the effect size almost doubled) on the RSVP compared to the masking tasks. Masking may require more spatial information (e.g., locate a notch), whereas RSVP may require form discrimination. Moreover, RSVP tasks require sustained orientation or alertness to visual stimuli, whereas masking involves a single stimulus array preceded by a fixation point and followed by a mask.

Although SZ patients showed poor performance across all stages of visual processing, there is evidence that very early feedforward processing before the formation of percepts is intact in SZ (Green et al., 2009; Hahn et al., 2011; Jahshan et al., 2012). Taking into account these previous studies and the current study, it is likely that SZ patients are impaired at all subsequent stages beyond the initial sensory feedforward processing stage.

Our study is limited by the fact that SZ patients were taking antipsychotic medications at the time of testing. However, given that antipsychotic medications did not have an effect on visual processing in BD, and previous studies of visual masking in SZ found no effects of antipsychotic medications (Butler et al., 1996; Green et al., 1999), it is unlikely that these medications had a strong impact on the findings. Although lithium may have an effect on visual processing (Fleming and Green, 1995; Green et al., 2011b), only 28% of our BD patients were taking lithium and their performance was not significantly different than those not taking lithium.

In summary, the BD group’s performance was comparable to the control group at the earliest stages of perceptual processing (object formation and substitution) but intermediate between the SZ and control groups at a later processing stage involving the interface between perception and higher order processing. These results suggest that, in contrast to the widespread disruption of neural processes across early and later stages of the visual processing hierarchy observed in SZ, neural processes associated with early stage visual processing are intact in BD. However, reduced performance on the RSVP task is consistent with narrow disruption of higher order attentional networks in BD. Despite the commonly-reported neurocognitive and genomic similarities between SZ and BD (Bora et al., 2009; Purcell et al., 2009), deficient early visual processing does not seem to be a prominent characteristic in BD patients and does not represent a shared phenotype between the two patient groups. Given models of pathways from early visual perception to functional outcome in SZ (Green et al., 2012), future studies might benefit from evaluating cascade models in BD starting with the latest stage of visual information processing.

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Contributors

MFG and JKDW designed the study. CJ and JKDW analyzed the data. CJ, JKDW, AM, and MFG interpreted the findings and wrote the first draft of the manuscript with DCG and LLA providing feedback. All authors have contributed to and approved the final manuscript.

Conflict of interest

All authors declare that they have no conflicts of interest arising from this manuscript.

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