Patients with schizophrenia demonstrate reduced cortical sensitivity to auditory oddball regularities

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A B S T R A C T
Background: Individuals with schizophrenia demonstrate deficits in context processing. These deficits can be characterized by examining the influence of auditory context on ERP responses to rare target tones. Previous studies demonstrate that target ERP deficits in schizophrenia depend on the number of non-targets that precede the target ERP. Our goal was to extend these findings by examining whether patients with schizophrenia demonstrate a reduced sensitivity to subtle differences in the auditory context preceding rare target stimuli, as quantified by Itti and Baldi’s Bayesian prediction error model.

Methods: Cortical responses to auditory oddball tones were measured within 59 individuals with schizophrenia (SZ) and 59 controls (HC). Individual trial amplitudes were estimated by conducting group ICA on the EEG time series and analyzing the reconstructed individual temporal sources. We quantified the auditory context of target tones using the Bayesian prediction error model and determined whether ERP amplitudes to tones were sensitive to this measure of context, or the number of preceding non-targets directly, within HC and SZ.

Results: Individuals with schizophrenia show a significant reduction in ERP response amplitudes to targets approximately 244–412 ms following target onsets. Individual amplitudes within this window showed significantly greater sensitivity to the modeled prediction error within the controls than in individuals with schizophrenia. These differences approached significance when examining differences in amplitudes as a function of the number of preceding non-targets.

Conclusions: These findings further clarify differences in HC and SZ with regard to their attentional and perceptual sensitivity to subtle environmental regularities.

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1. Introduction

A prominent characteristic of schizophrenia is a perceived loss of environmental continuity or a reduced ability to integrate multiple stimuli coherently (Cox and Leventhal, 1978). These deficits in perceptual organization appear to underlie a number of symptoms of schizophrenia, including a reduced ability to integrate visual elements into a coherent whole (Place and Gilmore, 1980; Butler et al., 2001) and a reduced ability to utilize auditory context when detecting auditory targets (Silverstein et al., 1996).

Perceptual organization deficits can be examined in greater detail by examining event-related potential (ERP) responses to a series of frequent and infrequent auditory stimuli, as in auditory oddball tasks. It has long been known that individuals diagnosed with schizophrenia demonstrate reduced ERP amplitudes ~300 ms following rare auditory targets (Roth and Cannon, 1972; Levit et al., 1973). These responses, termed the P3, appear sensitive to the sequence of auditory tones that precede the behaviorally relevant target. For example, Gonzalez et al., 1995 demonstrate P3 amplitude differences between controls and schizophrenia patients when targets appear after 3–7 non-targets, but not after a shorter (1) or longer (9) sequence of non-targets. Javitt et al. (1998) demonstrate greater ERP amplitude differences in schizophrenia in blocks with a reduced probability of rare stimuli (i.e. in situations where rare stimuli generally appear following a greater number of non-targets).

These oddball ERP deficits in schizophrenia patients suggest sensitivity differences between patients and controls with respect to the acoustic context (Gonzalez et al., 1995; Javitt et al., 1998; Shelley et al., 1999; Gilmore et al., 2005). In addition to manipulating rare target probability or the number of preceding non-targets, the acoustic context can be further characterized by modeling the level of Bayesian surprise or prediction error of rare stimuli (Itti and Baldi, 2005, 2009; Baldi...
and Itti, 2010). According to this model, individuals have a prior expectation of the probability in which each stimulus occurs. This prior expectation is updated when a new stimulus is encountered, and the prediction error is reflected by the degree in which the probability is updated. Stimuli that are more unexpected correspond to greater differences between the two probabilities, which corresponds to greater prediction error (Itti and Baldi, 2005, 2009; Baldi and Itti, 2010) and a greater ERP modulation (Lieder et al., 2013).

Our goal was to examine whether patients with schizophrenia demonstrate a reduced sensitivity to differences in the context of rare target stimuli, as quantified by the Bayesian prediction error model. This model allows estimates of ERP amplitudes for individual targets even when the number of preceding non-targets is fixed and when a target follows another target. Thus, the model allows additional flexibility in the complexity of the auditory environment (i.e., with non-target intervals between 0 and 20), in contrast to previous studies which focus on a comparatively reduced range of non-target intervals (Gonsalvez et al., 1995; Gilmore et al., 2005), or broad measures of regularity such as target probability (Javitt et al., 1998; Shelley et al., 1999). In order to explore differences in auditory regularity sensitivity between the two groups, individual target ERP responses were linearly fit to the modeled prediction error, or the number of preceding non-targets, and differences in the linear fit between the HC and SZ groups were demonstrated. These results help further clarify differences between schizophrenia patients and healthy controls with respect to their sensitivity to auditory regularities.

2. Methods

2.1. Participants and procedures

One hundred and eighteen individuals participated at the Institute of Living at Hartford Hospital. The study was conducted in accordance with an experimental protocol approved by the Institutional Review Board (IRB). Trained clinicians acquired medical history, conducted Structured Clinical Interviews for DSM-IV diagnosis (First et al., 1997), and acquired symptom information (Positive and Negative Syndrome Scale (Lancot et al., 2000)). Further details on recruitment, diagnosis, and exclusion criteria can be found in Ethridge et al. (2012) and Supplementary methods.

Among the 118 participants, 59 were healthy controls, 45 were diagnosed with schizophrenia, and 14 were diagnosed with schizoaffective disorder, depressed type using DSM-IV criteria based on a SCID interview. The two diagnostic groups were combined into a single group with schizophrenia spectrum psychotic disorders (SZ group), since the schizoaffective depressed type patients demonstrate similar overall cognitive impairment as patients diagnosed with schizophrenia (Hill et al., 2013). Medication information and mean symptom scores are provided in the Supplementary methods and Supplementary Table 1. The healthy control group consisted of 27 males (mean age = 35 ± 11 years) and 32 females (mean age = 36 ± 11 years). The SZ group consisted of 42 males (mean age = 30 ± 10 years) and 17 females (mean age = 36 ± 13 years). The groups did not significantly differ in age (T(116) = 1.656; p = 0.100), but showed significant differences between the two probabilities, which corresponds to greater prediction error (Itti and Baldi, 2005, 2009, 2010), implemented in Nathan Mundhenk’s Bayesian Surprise Matlab Toolkit (http://sourceforge.net/projects/surprise-mltk/). In order to calculate the Bayesian surprise or prediction error, we assume that the individuals probabilistically model the frequency in which a relevant tone occurs in healthy controls. Group temporal ICA was used to decompose the multiplexed ERP response into distinct components which potentially reflect the distinct ERP peaks (Eichele et al., 2011; Supplementary Fig. 2 and Supplementary methods).

2.2. Oddball task

Individuals pressed a button when an infrequent target tone (1000 Hz) appeared within a stream of frequent standard tones (1500 Hz). The tones were delivered by two 8-ohm speakers 50 cm away from the individual. The tones were presented every 1.3 s over the course of the 14 min 50 s experimental session. The tones were presented in a pseudorandom order with target tones appearing 15% of the time (n = 98).

2.3. EEG acquisition and preprocessing

EEG was recorded with a 66-channel Neuroscan system (Compumedics, Charlotte, NM). Silver/silver chloride electrodes were placed according to the International 10–10 system with a midforehead ground and nose reference (sampling rate = 1000 Hz; impedance ≤ 5 kΩ). An additional four electrodes were used to record horizontal and vertical eye movements. EEG preprocessing was conducted in Matlab (http://www.mathworks.com) using custom functions, built-in functions, and the EEGLAB toolbox (http://sccn.ucsd.edu/eeGLab). The EEG data was linearly detrended, forward and backward filtered with a Butterworth filter (bandpass: 0.01 to 50 Hz), downsampled to 250 Hz, and average referenced. Eye blink and muscle artifacts were attenuated by conducting a temporal ICA decomposition on the individual recordings (see the Supplementary methods).

2.4. Group temporal ICA of EEG

The average difference between the target and standard responses is indicated in Supplementary Fig. 1 for the HC (N = 59) (in red) and SZ (N = 59) (in blue) groups. The results are consistent with those of the previous studies indicating an increased ERP modulation to the rare relevant tone within healthy controls. Group temporal ICA was used to decompose the multiplexed ERP response into distinct components which potentially reflect the distinct ERP peaks (Eichele et al., 2011), using the same temporal window as the group component.

2.5. Auditory statistical regularities

The prediction error of each tone was modeled using the model of Baldi and Itti (Itti and Baldi, 2005; Einhäuser et al., 2007; Itti and Baldi, 2009, 2010), implemented in Nathan Mundhenk’s Bayesian Surprise Matlab Toolkit (http://sourceforge.net/projects/surprise-mltk/). In order to calculate the Bayesian surprise or prediction error, we assume that the individuals probabilistically model the frequency in which a particular tone will appear based upon the sequence of previous tones. This probability, or model M, is updated with a new observation, D = λ, generating a posterior distribution of the expected stimulus frequency. The prior and posterior densities are characterized by

<table>
<thead>
<tr>
<th>Component</th>
<th>Peak time (ms)</th>
<th>Peak window (fwhm) (ms)</th>
<th>Spatial peak Min (z-score)</th>
<th>Spatial peak Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>96</td>
<td>68-136</td>
<td>O/T (−1.7)</td>
<td>F/C (1.2)</td>
</tr>
<tr>
<td>2</td>
<td>144</td>
<td>120-168</td>
<td>T (−1.2)</td>
<td>F/C (0.7)</td>
</tr>
<tr>
<td>3</td>
<td>196</td>
<td>148-264</td>
<td>O (−0.7)</td>
<td>F/C (1.3)</td>
</tr>
<tr>
<td>4</td>
<td>208</td>
<td>172-240</td>
<td>F/C (&lt;0.5)</td>
<td>F/O (0.6)</td>
</tr>
<tr>
<td>5</td>
<td>340</td>
<td>244-412</td>
<td>O/P (−1.1)</td>
<td>F (1.6)</td>
</tr>
<tr>
<td>6</td>
<td>484</td>
<td>372-656</td>
<td>O/P (&lt;1.0)</td>
<td>F (1.3)</td>
</tr>
</tbody>
</table>

O = occipital, T = temporal, P = parietal, C = central, F = frontal.
gamma densities $\gamma(\lambda; \alpha, \beta)$ and $(\lambda; \alpha', \beta')$, respectively. The posterior density is adjusted given a new observation $\lambda$ by

$$\alpha' = \zeta \alpha + \bar{X}$$ and $\beta' = \zeta \beta + 1$

where the “working memory” parameter $\zeta$ adjusts $\alpha$ and $\beta$ to reduce the influence of the less recent stimuli. Zeta ($\zeta$) was set to $0.7$ in the current study, which is consistent with Itti and Baldi (2009) and estimates of neural adaptation (Muller et al., 1999).

The difference between each density is given by the Kullback–Leibler (KL) divergence (Kullback, 1959) between the posterior and prior gamma ($\gamma$) densities as given by:

$$S(D, M) = KL(\gamma(\lambda; \alpha, \beta), \gamma(\lambda; \alpha, \beta)) = -\alpha + \alpha \log \frac{\beta}{\beta'} + \log \frac{\Gamma(\alpha)}{\Gamma(\alpha')} + \beta \frac{\alpha'}{\alpha} + (\alpha' - \alpha) \Psi(\alpha')$$

where $\Gamma$ and $\Psi$ are the Euler gamma and digamma functions, respectively (see Itti and Baldi, 2009 for additional details). Frequent (i.e. standard) tones were coded as $\bar{X} = 1$ and infrequent target tones were coded as $\bar{X} = 2$. The greater the stimulus deviates from expectations, the greater difference between the posterior probability density and the prior probability density, and the larger the prediction error $S$. The prediction error for individual stimuli in a sequence is demonstrated in Fig. 1. Supplementary Fig. 3 demonstrates the stimulus sequence for a subset of the targets, organized by their level of prediction error.

3.2. Group temporal ICA of EEG

The average difference between the target and standard responses is indicated in Supplementary Fig. 1 for HC ($N = 59$) (in red) and SZ ($N = 59$) (in blue). The results are consistent with those of the previous studies indicating a reduced ERP modulation to the rare relevant tones within SZ, and with previous studies analyzing a larger dataset which included individuals within the current group (Ethridge et al., 2012, 2014). The predominance of noise within unaveraged ERP responses motivated further processing steps for single trial analysis. Group ICA was used to decompose the multiplexed ERP response into distinct components which potentially reflect the distinct ERP peaks. The components were averaged across epochs for target and standard tones and 6 out of 15 components were selected for further analysis since they demonstrate an average peak to either the standard or target tone.

The average event-related response is indicated in Fig. 2a and c for target (red) and standard (blue) tones along with the average topography across all subjects. The peak time, window, and topographic peak locations are indicated in Table 1. The individual time courses were determined by back reconstructing the group components on the individual data. Fig. 2b and d demonstrates the reconstructed response averaged across targets. Each row indicates the response for a particular subject, with HC above the solid line and SZ below.

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The average difference between the target and standard responses is indicated in Supplementary Fig. 1 for HC ($N = 59$) (in red) and SZ ($N = 59$) (in blue). The results are consistent with those of the previous studies indicating a reduced ERP modulation to the rare relevant tones within SZ, and with previous studies analyzing a larger dataset which included individuals within the current group (Ethridge et al., 2012, 2014). The predominance of noise within unaveraged ERP responses motivated further processing steps for single trial analysis. Group ICA was used to decompose the multiplexed ERP response into distinct components which potentially reflect the distinct ERP peaks. The components were averaged across epochs for target and standard tones and 6 out of 15 components were selected for further analysis since they demonstrate an average peak to either the standard or target tone.

The average event-related component response is indicated in Fig. 2a and c for target (red) and standard (blue) tones along with the average topography across all subjects. The peak time, window, and topographic peak locations are indicated in Table 1. The individual time courses were determined by back reconstructing the group components on the individual data. Fig. 2b and d demonstrates the reconstructed response averaged across targets. Each row indicates the response for a particular subject, with HC above the solid line and SZ below.

3.3. Behavioral relevance and regularities

We found that the component amplitude to behaviorally relevant infrequent tones was significantly larger than the component amplitudes to the irrelevant frequent standard tones for all 6 selected components (Table 2, rows 2 and 3). Differences in amplitudes between HC and SZ were examined with an ANOVA analysis with the difference in reconstructed amplitudes between targets and standards as the independent variable and diagnosis and sex as dependent variables. Amplitudes significantly differed between the HC and SZ groups for components 5 and 6 ($F(1,116) = 18.6; p = 0.00003, F(1,116) = 10.0; p = 0.002$), with target responses significantly larger (i.e. more positive) than standard responses for HC compared to SC (Table 2, rows 6 and 7). These results indicate that HC and SZ show differences in auditory processing ~340 and ~484 ms following an auditory stimulus.

In addition to differences in the response to targets and standards, we also examined whether the response to individual target tones was modulated by the subtle auditory context, as indicated by the tone prediction error. The modeled prediction error was plotted for each target as a function of the number of preceding non-targets (Fig. 3a). Similar trends are observed when comparing the prediction error (Fig. 3a) and the average amplitude for component 5 (Fig. 3b) and component 6 (Supplementary Fig. 4b) against the number of preceding non-targets. In general, these figures suggest a non-linear relationship between the amplitude and the number of preceding non-targets. This relationship appears linear when amplitudes are plotted against the modeled prediction error (Fig. 3c and Supplementary Fig. 4c), suggesting that the prediction error may be a more appropriate dependent variable in linear statistical tests.

We found that the two group components with the greatest delay (e.g. components 5 and 6) each demonstrate a significant correlation with the prediction error ($r(96) = 0.54; p < 0.001$;
\( r(96) = 0.45; p < 0.001 \), with increases in the prediction error associated with increased ERP amplitudes (Table 2, rows 4 and 5). Differences in the sensitivity to the prediction error between HC and SZ were examined with an ANOVA analysis. The individual subject amplitudes to target tones were linearly fit to the prediction error for each tone, and the slope of the fit served as the independent variable, with diagnosis and sex as dependent variables. Slopes significantly differed between the HC and SZ groups for component 5 (\( F(1,116) = 10.6; p = 0.002 \)) (which overlaps with the P3 ERP component, see Supplementary Fig. 5 and Supplementary results). This difference is characterized by a greater slope in the linear fit between the amplitude and the prediction error within the HC group compared to the SZ group. The relationship between the component amplitude and the target prediction error is indicated in Fig. 4, along with the average amplitude for each target separately for HC and SZ. This relationship approached uncorrected
The ERP components which peaked at 340 and 484 ms each showed sensitivity to the prediction error at the group level. However, healthy controls showed a greater sensitivity to the prediction error within component 5, which peaked at 340 ms, while the two groups did not significantly differ within component 6, which peaked at 484 ms. It is interesting to note that these components appear to correspond to the P3 (see the Supplementary results) and P3b ERP response, respectively, and that these responses previously demonstrated differences with regard to attentional modulation and their relationship to gray matter volumes. For example, the P3 response appears when rare stimuli are either attended or ignored and is associated with frontal gray matter volume, while the P3b appears primarily when stimuli are attended and is associated with parietal gray matter volume (Squires et al., 1975; Ford et al., 1994). The present study further clarifies differences between these responses with respect to the sensitivity to environmental regularities in healthy controls and individuals diagnosed with schizophrenia.

4.1. Comparing Bayesian prediction error and the number of preceding non-targets

Within auditory oddball tasks, context processing deficits have primarily been explored by considering the number of non-targets that precede rare stimuli. For example, ERP amplitudes have been compared separately for rare stimuli that follow between 1 and 9 non-targets (Gonsalvez et al., 1995; Gilmore et al., 2005), or as a function of rare stimulus probability (Javitt et al., 1998; Shelley et al., 1999). In some cases, the number of preceding non-targets may be an incomplete summary of acoustic regularities. According to the Bayesian prediction error model, this may apply in particular for targets that precede 1–3 non-targets (see Supplementary Fig. 3). For example, all rare stimuli that follow 1 non-target are assumed to have the same ERP amplitude, disregarding the sequence of stimuli that appeared prior to the non-target. The prediction error model includes a weighted contribution of past events (e.g. incorporating biologically plausible assumptions about working memory), which improves predictions of fluctuations in rare stimulus amplitudes in instances with a fixed number of preceding non-targets, or in instances where a target precedes another target (which occurred within 17% of targets within the current study).

Individual’s sensitivity to auditory context was summarized in the current study by calculating the slope of the linear fit between single trial rare target amplitudes and the prediction error, or the number of preceding non-targets. It is interesting to note the close correspondence between the modeled prediction error and the observed amplitude, when plotted against the number of preceding non-targets (Fig. 3a and b). This result cautions against assuming a linear relationship between ERP amplitudes and the number of preceding non-targets, further supports the utility of the prediction error model for predicting ERP amplitudes (Lieder et al., 2013), and is consistent with the observation of significant differences between HC and SZ when linearly fitting amplitudes with the prediction error, compared to with the number of preceding non-targets.

4.2. Context processing deficits and symptoms

The reduced sensitivity to environmental regularities in schizophrenia could potentially be related to the frequency of hallucinations (e.g. internal distractions), be related to the influence of medication, or result due to a general cognitive deficit. In an exploratory analysis, the component which differed in the prediction error between the groups was not significantly associated with positive (r = −0.07) or negative (r = 0.13) PANSS scores, or with total current antipsychotic dosage (r = −0.16). The absence of a significant relationship may result due to the relatively modest sample size. Thus, a larger heterogeneous sample could disentangle the underlying factors that contribute to these results.

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### Table 2

Statistical tests for differences in the components and reconstructed responses. The component difference section displays the results for component differences between target and standard amplitudes (relevance) and correlations between component amplitudes and the prediction error (regularity) for targets. The reconstructed difference section displays the results for relevance and regularity differences between HC and SZ. The results are displayed for 6 selected components.

<table>
<thead>
<tr>
<th>Group</th>
<th>Component difference</th>
<th>Reconstructed difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Relevance</td>
<td>p</td>
</tr>
<tr>
<td></td>
<td>T(654)</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>7.9</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>2</td>
<td>9.6</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>3</td>
<td>3.56</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>4</td>
<td>4.8</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>5</td>
<td>71.3</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>6</td>
<td>65.7</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

Bold data indicate P-values below the Holm-Bonferroni corrected threshold.
4.3. Conclusion and implications

The present findings further clarify differences in HC and SZ with regard to their sensitivity to subtle environmental regularities. Rare target ERP amplitudes in schizophrenia were less influenced by the context of acoustic events before each target, indicating a relative ‘uncoupling’ of the response to a particular target from its acoustic context. This is consistent with impairments in working memory (Park and Holzman, 1992) and a more fragmented perceptual experience (Cox and Leventhal, 1978; Silverstein et al., 1996) in schizophrenia.

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Contributors
GP contributed to the experimental design. DB conducted the analysis and wrote the manuscript. GP, KK, and VC edited the manuscript and assisted with interpretation of the results. All authors have contributed to and approved the final manuscript.

Conflict of interest
Dr. Pearson reports serving as a consultant for Bristol-Myers Squibb in 2012. Dr. Bridwell, Dr. Kiehl, and Dr. Calhoun have no conflict interest to declare.

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Appendix A. Supplementary data
Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.schres.2014.06.037.

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