Classification of Schizophrenia Patients and Healthy Controls Using P100 Event-Related Potentials for Visual Processing

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Discriminant analysis · Event-related potentials · Information processing · Schizophrenia

Abstract
The study of event-related potentials (ERPs) is capable of elucidating the abnormalities in brain network dynamics relevant to the information-processing deficits in schizophrenia patients. In contrast to P50 and P300 ERPs, however, the results of P100 ERP studies in schizophrenia patients are less consistent. We have previously reported that P100 amplitudes did not differ significantly between patients with schizophrenia and healthy subjects. This result raised a question as to whether P100 ERPs carry information on brain network dynamics in schizophrenia patients that is distinct from normal controls. To answer this question, in this study we performed discrimination analysis on the P100 data. The rate of correct classification of patients and controls was high (80–90\% depending on stimulus categories), indicating that patients have spatial patterns of P100 amplitudes that are distinguishable from those in healthy subjects. To further explore this possibility, we performed principal component analysis on the P100 data. For the patients, the first principal component represented global activity, the second component represented the reciprocal anterior-posterior activation, and the third component represented the hemispheric reciprocity in activity. The first and second components were similar to those of the control group; however, the third component in control subjects showed activation of the center versus anterior and posterior regions. This result is consistent with the notion of abnormalities in hemispheric asymmetries during the processing of sensory information in schizophrenia. In conclusion, this ERP study demonstrated that P100 amplitudes have information that can successfully classify patients and controls.

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Introduction
Neuroimaging is a promising method of assessing the normality of information processing in the brain because it permits the objective assessment of neural activation. Since the development of neuroimaging techniques such as event-related potentials (ERPs) and functional magnetic resonance imaging, a number of studies using these new techniques have detected abnormalities in patients with schizophrenia and other diseases. In many ERP studies, P50 and P300 ERPs have been robust indicators of disease \cite{1, 2}. These measures have assessed several aspects of information-processing deficits that have long
been proposed as having a central role in schizophrenia [3]. Along with prepulse inhibition, P50 suppression is reduced in patients with schizophrenia, which is suggestive of sensory gating deficits [4, 5]. Interestingly, the prefrontal cortex has been suggested to be involved in the generation of P50 ERPs, which occurs at an early stage in sensory processing [6]. The auditory P300 ERP has been recorded in many studies using oddball tasks. The P300 amplitude increases as the probability of encountering an infrequent deviant stimulus decreases [7]. Patients with schizophrenia showed reduced amplitudes and prolonged latencies of P300 ERPs [1, 2]. Recent studies using visual oddball tasks reported that patients with schizophrenia showed significant reductions of P300 amplitudes [8]. These diminished P300 ERPs, both auditory and visual, were recovered by the administration of an antipsychotic drug [9].

In contrast to P50 and P300, the studies of P100 ERPs in schizophrenia patients are less consistent. There are studies reporting reduced P100 amplitudes in patients with schizophrenia [10–12]; however, other studies have reported inconsistent results [13, 14] or found no differences [15–17]. The P100 amplitude correlated positively with both the Brief Psychiatric Rating Scale and the Scale for the Assessment of Negative Symptoms, exhibited variability across the electrodes, and was higher in patients than healthy controls at approximately 110 ms [13]. We recently recorded peaks at approximately 110 ms that were higher in patients than in healthy controls, although the differences did not reach significance [15]. The P100 latencies for a no-contour stimulus correlated with the cognitive subscales of the Positive and Negative Syndrome Scale [15], being consistent with cognitive impairment in patients [15, 18]. On the other hand, the relationships with early sensory processing deficits have been obscure. Given that early-stage visual processing, which is typically in the range of less than 200 ms after stimulus onset, is also dysfunctional in schizophrenia patients [11, 12, 19, 20], P100 ERPs might reflect altered brain network dynamics that is relevant to early sensory processing deficits. To explore this possibility, we investigated P100 ERPs in the following two steps. First, we examined whether multielectrode P100 amplitudes could discriminate patients from healthy controls. Successful discrimination with the multielectrode P100 amplitudes would indicate that the ERP data contain information on the network dynamics of patients with schizophrenia that differs from healthy controls. Second, we attempted to extract principal components from the ERP data that might be relevant to the discrimination. The extracted component structure would clarify spatial characteristics of network dynamics reflecting information processing in the brain of patients as well as controls.

### Methods

#### Participants

A total of 20 medicated, right-handed, schizophrenic patients (13 males) between the ages of 17 and 49 (mean ± SD: 31.3 ± 7.2) were recruited at the Juntendo Koshigaya Hospital in Saitama, Japan (table 1). The patients who participated in this study met the Diagnostic and Statistical Manual of Mental Disorders fourth edition (DSM-IV) criteria for schizophrenia and had relatively mild psychotic manifestations. The majority of the patients had intact social adjustments, either having an occupation or being a student, until the manifestation of psychotic episodes. The Positive and Negative Syndrome Scale [21] scores were recorded on the basis of a structured psychiatric interview and by a review of the medical charts of the patients. The ages at onset were 21.4 ± 6.0 and the durations of the disease were 9.8 ± 6.6 years. All patients were medicated with typical (n = 6) or atypical antipsychotic drugs (n = 14). The patient and healthy control groups were matched for age, handedness and gender. The demographic and clinical characteristics of the participants are summarized in table 1. All patients were judged to be in good physical health on the basis of medical history, physical examination and laboratory measures. None of the subjects had a history of electroconvulsive shock treatment.

| Patients | Healthy controls |
|----------------|-----------------|
| Age, years | 31.3±7.2 | 31.8±5.0 |
| Sex, n | | |
| Male | 13 | 13 |
| Female | 7 | 7 |
| Age at onset, years | 21.4±6.0 | 9.8±6.6 |
| Illness duration, years | 72.3±18.6 | |
| Total PANSS score | | |
| Subtypes, n | | |
| Disorganized | 7 | |
| Paranoid | 4 | |
| Catatonic | 2 | |
| Residual | 6 | |
| Unclassified | 1 | |
| Amount of episodes including that at the tests | 1.5±1.6 | |
| Antipsychotic medication, n | | |
| Typical | 6 | |
| Atypical | 14 | |

Values are presented as mean ± SD unless otherwise indicated. PANSS = Positive and Negative Syndrome Scale.
Patients and Healthy Controls

Discriminant Analysis

Discriminant analysis attempts to predict group membership from a set of predictors [22]. In linear discriminant analysis, the discriminant function is defined by a linear combination of independent variables or predictors as follows:

\[ D = a_1X_1 + \ldots + a_pX_p \]  

where \( D \) is the discrimination function, \( X_1, X_2, \ldots, X_p \) are the predictors, and \( a_1, a_2, \ldots, a_p \) are the standardized discrimination function coefficients. In this study, the predictors are P100 amplitudes at the corresponding electrodes, and \( p = 19 \), which is the number of electrodes. The discrimination function \( D \) was calculated for each group (patients and healthy controls) and for each stimulus category (RC, NC and IC). The mean of the discrimination function over the two groups for each stimulus category was zero because the predictors were standardized. If the score of the discrimination function for a subject was less than zero, the subject was classified as a patient with schizophrenia. The discriminant analysis was performed with SPSS Statistics for Windows, version 20 (IBM Corp., New York, N.Y., USA).

Principal Component Analysis

The P100 amplitude data for the three categories of stimuli were subjected to principal component analysis (PCA). From the data, the PCA extracted principal components, which had characteristic scalp distributions. The analysis was performed for the patient group and the control group separately, so that the analysis yielded different sets of principal components for the two groups. In each group, the principal components whose eigenvalues were greater than 1.0 were subjected to interpretation. The PCA was performed with R (the R Statistical Computing Environment).
**Results**

*Classification of Patients and Controls*

From the observed P100 amplitudes, which were the predictors of the discriminant function, given by equation 1, the discriminant analysis estimated the scores for individual subjects, D. The distributions of the scores for both groups of subjects are shown in figure 2. Negative values of the discriminant score classified the subjects as patients, while positive values classified the subjects as healthy controls. Although there was some overlapping, the distributions of the scores were well separated (p < 0.00001 for RC, NC and IC). For the RC stimuli, 2 of the 20 healthy controls were mistakenly classified as patients, while 2 of the 20 patients were mistakenly classified as healthy controls. The rate of correct classification was 90.0%. For the NC stimuli, 5 controls and 3 patients were mistakenly classified; the correct classification rate was 80.0%. For the IC stimuli, 3 controls and 4 patients were mistakenly classified; the correct classification rate was 82.5%.

*Principal Component Analysis*

If the P100 amplitudes differed significantly between the groups, the differences of the P100 amplitudes were the primary contributor to the discrimination of patients from controls. However, this was not the case in this study. One might consider that just an unweighted collection of P100 amplitudes at 19 electrodes would have increased the power of discrimination. This inference was denied because the average P100 amplitude over the electrodes did not differ between the groups (p = 0.19 for RC, 0.45 for NC and 0.27 for IC). On the contrary, the rather high discrimination rates indicate that patients had characteristic spatial patterns of the P100 amplitudes over the electrodes that were distinctive from those of healthy subjects. Then, to explore these patterns, we performed PCA on the P100 amplitudes. The PCA extracted the principal components (table 2). Among them, the first three components that had eigenvalues greater than 1.0 were retained (the eigenvalues for the first, second and third components were 9.60, 4.23 and 1.18, respectively). These three components explain 84.8% (for the patients) and 74.0% (for the controls) of total variance. All of the coefficients for the first principal component (PC1) had positive values (table 2), which indicates that this component represents global activity. This feature was the same in both patients and controls. However, although the coefficients for PC1 had the same sign, they were relatively higher in the frontocentral region than in the posterior region in the patient group. This feature was not seen in the control group. The coefficients for the second principal component (PC2) had negative values at the anterior electrodes and positive values at the posterior electrodes, while they were smaller with mixed signs at the central electrodes. Therefore, PC2 represents reciprocal anterior-posterior activation. In contrast to the first and second components, the third principal component (PC3) was essentially different between the groups. This component in the patient group showed the hemispheric reciprocity of activation, as the coefficients for PC3 had positive values in the left hemisphere and negative values in the right hemisphere. In contrast, PC3 in the control group did not show such hemispheric differences but rather showed the activation of the central versus the anterior and posterior regions. These results are visualized in figure 3.
Discussion

Because our previous analysis showed insignificant group differences in the P100 amplitudes [15], the first question we addressed was whether the P100 amplitude data could discriminate patients from healthy controls. The discriminant analysis provided the scores for the individual subjects, which differed significantly between the groups. As a result, the rates of successful classification of patients and healthy controls were 80–90% depending on the categories of visual stimuli. This result indicates that, although the ERP at each electrode did not carry sufficient information on the pathological dynamics of the brain of patients with schizophrenia, the assembly of information at individual electrodes increased the power of the classification. In fact, reducing the numbers of electrodes in the analysis decreased the discrimination rate monotonically. Note, however, that this increased power was not due to just the averaging of the amplitudes over the electrodes because the group differences in the averaged amplitudes did not reach significance. Instead, we inferred that there were spatial patterns in the distribution of ERP amplitudes that were specific to schizophrenia. To confirm this inference, we performed PCA on the ERPs. If this inference was correct, the PCA would extract components with spatial profiles that differed between patients and controls. If the PCA extracted only one significant component that had no specific spatial profile, so that the loadings on this component were somewhat homogeneously distributed over the electrodes, this inference was to be denied. The result of the PCA showed that three significant components were extracted. Two of the extracted components had obvious spatial profiles; one was anterior-posterior reciprocity and the other was left-right reciprocity. Therefore, we conclude that there were spatial patterns in the distribution of ERP amplitudes that could be specific to schizophrenia.

Spatial Components of P100 Amplitudes

The next question we addressed regarded the type of information the P100 ERPs carried that was utilized for the discrimination of patients from controls. The spatial components extracted by the PCA were the anterior-posterior and left-right reciprocity. The anterior-posterior

| Table 2. Principal component coefficients or eigenvectors |
|----------------------------------------------------------|
| Patients with schizophrenia | Healthy controls |
|-----------------------------|-------------------|
| PC1  | PC2  | PC3  | PC1  | PC2  | PC3  |
| Fp1  | 0.211 | -0.257 | 0.075 | Fp1  | 0.202 | -0.308 | 0.395 |
| Fp2  | 0.224 | -0.265 | -0.013 | Fp2  | 0.212 | -0.336 | 0.249 |
| Fz   | 0.269 | -0.172 | -0.194 | Fz   | 0.244 | -0.166 | -0.209 |
| F3   | 0.273 | -0.191 | 0.057  | F3   | 0.248 | -0.203 | 0.054  |
| F4   | 0.270 | -0.188 | -0.159 | F4   | 0.224 | -0.226 | -0.202 |
| F7   | 0.256 | -0.211 | 0.195  | F7   | 0.254 | -0.241 | 0.111  |
| F8   | 0.277 | -0.120 | -0.084 | F8   | 0.184 | -0.299 | 0.284  |
| Cz   | 0.288 | -0.018 | -0.143 | Cz   | 0.257 | -0.022 | -0.326 |
| C3   | 0.271 | 0.017 | 0.266  | C3   | 0.276 | 0.033 | -0.188 |
| C4   | 0.280 | 0.042 | -0.284 | C4   | 0.245 | -0.038 | -0.353 |
| T3   | 0.248 | -0.077 | 0.401  | T3   | 0.200 | 0.062 | -0.158 |
| T4   | 0.255 | 0.083 | -0.207 | T4   | 0.232 | -0.101 | -0.114 |
| T5   | 0.185 | 0.228 | 0.452  | T5   | 0.214 | 0.316 | 0.107  |
| T6   | 0.175 | 0.324 | -0.156 | T6   | 0.220 | 0.159 | 0.125  |
| Pz   | 0.196 | 0.272 | -0.200 | Pz   | 0.235 | 0.229 | -0.081 |
| P3   | 0.163 | 0.311 | 0.138  | P3   | 0.250 | 0.276 | 0.054  |
| P4   | 0.156 | 0.335 | -0.278 | P4   | 0.248 | 0.257 | -0.120 |
| O1   | 0.121 | 0.343 | 0.227  | O1   | 0.196 | 0.298 | 0.392  |
| O2   | 0.110 | 0.361 | -0.097 | O2   | 0.192 | 0.313 | 0.304  |

In both groups, the first 3 components had eigenvalues greater than 1.0. These 3 components explain 84.8% (for the patients) and 74.0% (for the controls) of total variance.
reciprocity might be relevant to hypofrontality or the abnormality of top-down processing in schizophrenia [23]. However, the PCA in this study extracted the anterior-posterior component in both groups, and the profiles of this component over the electrodes were similar. Given that the frontoparietal network plays a role in attentional control [24–26], this result may suggest that both groups utilized this network to perform the task. Similarly, it might be expected that both groups had the laterality component because sensory processing is asymmetrical even in healthy subjects [27, 28]. However, the laterality component was extracted only from the patient group.

**Fig. 3.** Spatial distribution of the coefficients of the extracted principal components. Red (color refers to online version only) indicates a positive value, while blue indicates a negative value of the coefficient at each electrode. Only the coefficients greater than 0.1 or less than –0.1 are colored. **a–c** Patients with schizophrenia. **d–f** Healthy controls. **a, d** The first principal component (PC1). **b, e** The second principal component (PC2). **c, f** The third principal component (PC3). Note that the PC1 coefficients are not colored (in **a** and **d**) because all of them had values greater than 1.0 (see table 2).
This result suggests that, although hemispheric asymmetries of brain sensory information processing existed in both groups, the asymmetries were distorted further in patients with schizophrenia, being consistent with previous studies [29–31].

Cortical Network Dynamics

Although the difference at each electrode was subtle, the spatial components of P100 amplitudes as a whole had the power to discriminate patients from healthy subjects. The differences of the ERP spatial components between the groups would reflect the alterations of network dynamics in the patients, which would be relevant to the information-processing deficits in schizophrenia. A computational study suggested that the dynamics of the dorsolateral prefrontal cortex tends to be unstable under hypodopaminergic conditions and that processing speed becomes slower under the same conditions [32]. If top-down processing contributes to early visual processing, as suggested previously, the altered dynamics of the prefrontal cortex would influence the dynamics of the cortical areas for visual information processing. This possibility was tested in previous studies [15, 18], in which the observed association between cognitive factor in the Positive and Negative Syndrome Scale scores and P100 latencies suggested the abnormality in top-down rather than bottom-up processing in patients with schizophrenia. To date, however, how schizophrenia influences network dynamics of the brain has not been elucidated. Therefore, for further exploration of distinctive network dynamics of schizophrenia and its relationship to psychopathology, a strategic approach integrating an ERP study with a computational study would be needed. Because all the patients in this study were medicated by either typical or atypical antipsychotic drugs, the medication effects on brain network dynamics are also to be examined. It is, nevertheless, meaningful that the analysis in this study showed successful classification of patients and healthy controls. In conclusion, multielectrode P100 ERPs, based on weighted combination of P100 amplitudes at individual electrodes, provides diagnostic classification power, which will be useful as a biomarker of the disease.

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Disclosure Statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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