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Aberrant Processing of Deviant Stimuli in Schizophrenia Revealed by Fusion of FMRI and EEG Data

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Abstract

Background—Aberrant electrophysiological and hemodynamic processing of auditory oddball stimuli is among the most robustly documented findings in patients with schizophrenia. However, no study to date has directly examined linked patterns of electrical and hemodynamic differences in patients and controls.

Methods—In a recent paper we demonstrated a data-driven approach, joint independent component analysis (jICA) to fuse together functional magnetic resonance imaging (fMRI) and event-related potential (ERP) data and elucidated the chronometry of auditory oddball target detection in healthy control subjects. In this paper we extend our fusion method to identify specific differences in the neuronal chronometry of target detection for chronic schizophrenia patients compared to healthy controls.

Results—We found one linked source, consistent with the N2 response, known to be related to cognitive processing of deviant stimuli, spatially localized to bilateral fronto-temporal regions. This source showed significant between-group differences both in amplitude response and in the fMRI/ERP distribution pattern. These findings are consistent with previous work showing N2 amplitude and latency abnormalities in schizophrenia, and provide new information about the linkage between the two.

Conclusions—In summary, we use a novel approach to isolate and identify a linked fMRI/ERP component which shows marked differences in chronic schizophrenia patients. We also demonstrate that jointly using both fMRI and ERP measures provides a fully picture of the underlying hemodynamic and electrical changes which are present in patients. Our approach also has broad applicability to other diseases such as autism, Alzheimer's disease, or bipolar disorder.

Keywords

schizophrenia; fMRI; ERP; data fusion; independent component analysis; N2	

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INTRODUCTION

Key biological associations of schizophrenia include abnormalities in brain function as revealed by event-related potentials (ERPs) and functional magnetic resonance imaging (fMRI). These techniques are essential tools used to illuminate the temporal sequence and cerebral locations of cognitive and information processing deficits of schizophrenia patients. One of the most robust probes for aberrant brain function in schizophrenia is the auditory oddball task 1⁻⁴

The processing of auditory stimuli presented during an oddball task requires detection of infrequent target stimuli within the context of frequently presented standard stimuli 5, 6. Brain activity during oddball tasks is frequently measured by averaging task-related electroencephalogram (EEG) recordings to produce event-related potentials (ERPs). Over four decades ago it was reported that low probability, task relevant auditory stimuli elicited a characteristic ERP waveform that includes several meaningful components, including the mismatch negativity 7 related to sensory trace memory, the N2 related to matching stimuli to an internally-generated contextual template 8, 9, and the P3, which is generally thought to reflect directed, effortful processing 10 and contextual updating of working memory processes 11. ERPs reveal multiple abnormalities of sensory and cognitive processes in schizophrenia. The major ERP research measures associated with schizophrenia are the MMN (subcategory of the N2) 12, 13 and P3 4, 14-17. Also, the P3 abnormalities are identified as candidate biomarkers in other diseases, e.g. depression, Alzheimer's, alcoholism and epilepsy 18-20. Although P3 is well-studied, it lacks consistency and specificity to schizophrenia. In contrast, abnormality of the N2 component is much less explored or understood. The relative neglect by schizophrenia researchers of N2 is mainly due to the lack of knowledge regarding the clinical and neuropathological significance of N2 abnormalities 21. The N2 potential is a potentially important index of schizophrenia, because it is connected to initial stimulus categorization in the selective attention stream 9. 21-26. Under the N2 category (which peaks at approximately 200ms), the mismatch negativity (MMN or N2a) subcomponent originates in generators located in frontal-central and temporal lobes 27⁻29. And the N2b (or N2/P3) subcomponent, overlapping with MMN, is linked to neuronal activity of fronto-central, fronto-temporal, and parieto-temporal regions 25, 26, 30

EEG/MEG and functional magnetic resonance imaging (fMRI) have complementary strengths and weaknesses. The advantages of techniques such as EEG and MEG are their millisecond temporal resolution and ability to measure neuronal activity directly. In contrast, fMRI has excellent uniform spatial resolution but measures an indirect metabolic correlate of neuronal function - the blood oxygenation level dependent (BOLD) signal, over a considerably longer time period of seconds. To date, over a dozen fMRI experiments have employed oddball paradigms to examine target-related neural activity 31-33. A recent large scale (n=100) auditory oddball fMRI study found highly reliable activation in 38 regions for target detection 34. Regions activated during processing of target stimuli included portions of bilateral temporal, lateral frontal, and lateral parietal lobes, thalamus, amygdala, cerebellum, as well as motor-related areas 34. The temporal resolution of fMRI, though limited by the slow hemodynamic response, has been used to examine delay differences on the order of 100-200ms 35, 36 which is informative, but much less precise than the temporal information provided by EEG. In summary, though fMRI and ERP both provide spatial and temporal information, the strengths of each technique differ in a complementary manner. Thus an approach which combines fMRI and ERP can draw potentially on the strengths of each and provide additional information not afforded by either technique alone. Despite this obvious motivation combining these two measures has proven technically challenging, and is an ongoing effort that employs a variety of approaches.

Our work 37⁻⁴⁰ focuses on developing an effective multivariate fMRI/EEG data strategy aiming at a systematic group separation/diagnosis using independent component analysis (ICA), which maximizes the independence between components and has been used on both fMRI and EEG data separately. The ICA approach models spatiotemporal data as a linear combination of maps and time courses while attempting to maximize the independence between either the maps (spatial ICA) or the time courses (temporal ICA). The first application of ICA to fMRI data used spatial ICA 41 to determine spatially distinct brain networks. It is possible to perform ICA in either the temporal or spatial domain 42. For EEG and ERP data, it is more common to use temporal ICA whereas for fMRI data it is more common to use *spatial* ICA for various reasons, but primarily because of the larger number of data points in these domains 42. However, a joint estimation of the spatial components revealed by fMRI 31 and of the temporal components of the ERP response 43, 44 has seldom been attempted. In addition to extracting obvious joint sources that are also accessible to constraint or prediction based methods of multimodal integration, such an approach has the potential to reveal electrical sources which may not be readily visible in scalp ERPs or to expose brain regions that have participatory roles in source activity, but may not themselves be generators of the detected electrical signal 45.

This work extends our previous studies detecting the spatiotemporal relationship for healthy human subjects 38, 39 or for phantom simulation 37, in that we emphasize the detection of group differences between patients and controls, by jointly performing spatial ICA of fMRI data and temporal ICA of ERP data. We apply this approach to a group of twenty-three healthy participants and sixteen chronic schizophrenia patients in order to derive a spatiotemporal decomposition consisting of fMRI and ERP components, both indicating when the respective signal is changing with a specific focus upon joint differences between patients and healthy controls. Consistent with our previous work 2, 46, we hypothesized that a small number of joint components would capture differences between the patients and controls and reveal a joint network present in both groups. From previous work 2, 25, 46, 47, we predicted that this network would show both decreased activation and decreased ERP amplitude in patients. Our proposed approach provides a technique to examine linked hemodynamic and electrical sources which reveal significant differences between patients and controls.

METHODS

Participants

Participants were recruited via advertisements, presentations at local universities, and by word-of-mouth. Twenty-three healthy participants (15 males, 8 females, Age 41±14 years) and sixteen outpatients (11 males, 5 females, Age 38±11 years) with chronic schizophrenia, currently in complete or partial remission and on stable medication regimens, provided written, informed, Institutional Review Board (IRB)-approved consent at Hartford Hospital and were compensated for participation. See demographic details in Table 1. Prior to inclusion in the study, healthy participants were screened to ensure they were free from DSM-IV-TR Axis I or Axis II psychopathology, assessed using the SCID 48 and also interviewed to determine that there was no history of psychosis in any first-degree relatives. Patients met criteria for schizophrenia in the DSM-IV-TR Axis I disorders based on a structured clinical interview (SCID) and review of the case file. All participants had normal hearing (assessed by self-report), and were able to perform the task successfully during practice prior to the scanning session.

Experimental design

The auditory oddball task required subjects to press a button when they detected an infrequent sound within a series of regular and different sounds. Three stimuli were presented; frequent low-tone stimuli (standards), infrequent task-irrelevant stimuli (novels) and infrequent task-relevant stimuli (targets) requiring a button-press response. In the present experiment, the standard stimulus was a 500 Hz tone, the target stimulus was a 1000 Hz tone, and the novel stimuli consisted of non-repeating random digital noises (*e.g.*, tone sweeps, whistles) (Figure). Two runs of auditory stimuli were presented to each participant by a computer stimulus presentation system via insert earphones embedded within 30 dB sound attenuating MR compatible headphones for fMRI recording and standard headphones for EEG recording.

The target and novel stimuli each occurred with a probability of 0.10; the standard stimuli occurred with a probability of 0.80. The stimulus duration was 200 ms with a 2000 ms interstimulus interval (ISI). The intervals between stimuli of interest (target/novel) were allocated in a pseudorandom manner. All stimuli were presented at approximately 80 decibels above the standard threshold of hearing. All participants reported that they could hear the stimuli and discriminate them from the background scanner noise. Prior to entry into the scanning room or ERP booth, each participant performed a practice block of 10 trials to ensure each subject understood the instructions. The participants were instructed to respond as quickly and accurately as possible with their right index finger every time they heard the target stimulus and not to respond to the nontarget stimuli or the novel stimuli. An MRI compatible fiber-optic response device (Lightwave Medical, Vancouver, B.C.) was used to acquire behavioral responses for the task in both the fMRI and the ERP experiments. The stimulus paradigm, data acquisition techniques, and previously found stimulus-related activation are described more fully elsewhere 4· 31· 49.

Data acquisition

FMRI and ERP data were acquired on the same day in two different sessions, at the Olin Neuropsychiatry Research Center at the Institute of Living, using identical stimuli and counterbalancing the order of the fMRI or ERP sessions between individuals. The fMRI data were collected on a Siemens Allegra 3T dedicated head scanner equipped with 40mT/m gradients and a standard quadrature head coil. The functional scans were acquired using gradient-echo echo-planar-imaging with the following parameters (repeat time (TR)=1.50s, echo time (TE)=27ms, field of view=24cm, acquisition matrix=64×64, flip angle=70°, voxel size=3.75×3.75×4mm, gap=1mm, 29 slices, ascending acquisition). Six "dummy" scans were performed at the beginning to allow for longitudinal equilibrium, after which the paradigm was automatically triggered to start by the scanner. The ERP data was collected using an SA bioelectric amplifier system capable of amplifying electrical activity from 64 separate single-ended channels. Amplifiers were connected to a 16-bit A/D conversion using a custom program (Digitize) implemented on a Pentium II microcomputer running Solaris for Intel. The Digitize program recorded the EEG data and all stimulus and behavioral response codes for later analysis.

Preprocessing

FMRI—FMRI data were preprocessed using the software package SPM2 (http://www.fil.ion.ucl.ac.uk/spm/). Images were realigned using INRIalign – a motion correction algorithm unbiased by local signal changes 50. Next, data were spatially normalized into standard Montreal Neurological Institute space 51, spatially smoothed with a 12×12×12 mm full width at half-maximum Gaussian kernel. The data (originally acquired at 3.75×3.75×4mm) were slightly sub-sampled to 3×3×3 mm, resulting in 53×63×46 voxels.

ERP—Scalp potentials were recorded from tin electrodes (ElectroCap International) placed over 62 electrode sites according to the 10-20 electrode system placement and some supplemental sites. Vertical and horizontal electro-oculograms (EOG) were monitored from electrodes located on the lateral and supra-orbital ridges of the right eye. All electrodes were referenced to the nose. Electrical impedances were maintained below 10 k-ohms throughout the experiment. The EEG channels (SA instruments) were amplified (20,000 gain) with a bandpass of 0.1 to 100 Hz, digitized on-line at a rate of 500 samples per second, and recorded on computer hard disk. EEG data were preprocessed using ICA to remove ocular artifacts from the EEG data 52. Data were then digitally filtered with a 20 Hz low pass filter to reduce electromyographic activity and ERPs were constructed for trials in which participants correctly identified target stimuli. The recording epoch was 1400 milliseconds long with a 200 millisecond pre-stimulus baseline. Data from a midline central site (Cz) was included in the ICA fusion analyses because it appeared to be the best single channel to detect both anterior and posterior sources (results were nearly identical when scalp site Pz was used instead of Cz).

Joint FMRI/ERP Data Fusion

Joint ICA analysis tries to explain between-subject variations in data features in terms of underlying sources that cause those features. In this application, the data features include both spatial (fMRI) and temporal (EEG) components. The spatial features were contrast images of targets versus baseline for each subject generated using SPM2 software; providing a spatial map of oddball responses. The temporal data features were the ERP feature selected from the central electrode (Cz) from each subject; characterizing the temporal pattern of the oddball response. Furthermore, because we included both control subjects and schizophrenic patients in the analysis, an important source of variability in the underlying sources was the between-group differences in the neuronal expression of the oddball responses. It was these differences we wanted to assess.

We use an extended algorithm based upon the infomax principle 53, 54. The Infomax algorithm employs a natural gradient ascent algorithm to maximize the entropy of the output of a single layer neural network 54. We start with the assumption of joint spatial or temporal independence of the fMRI and ERP sources approximately, respectively, using the following generative model for the data, which maps from sources to observed data features: $\mathbf{x}^F = \mathbf{A}\mathbf{s}^F$

and $\mathbf{x}^E = \mathbf{A}\mathbf{s}^E$. For the case of two sources and two subjects, $\mathbf{x}^F = \begin{bmatrix} x_1^F & x_2^F \end{bmatrix}^T$ is the mixed data for the fMRI modality for the two subjects, $\mathbf{x}^E = \begin{bmatrix} x_1^E & x_2^E \end{bmatrix}^T$ is the mixed data for the ERP

modality for the two subjects, $\mathbf{A} = \begin{bmatrix} a_{11} & a_{12} \\ a_{21} & a_{22} \end{bmatrix}$ is a shared linear mixing matrix, and \mathbf{s}^F and \mathbf{s}^E are the respective fMRI and ERP sources. Instead of running ICA on each modality separately, we rewrite this as a single equation by forming a data vector for each subject as $\mathbf{x}_i = \begin{bmatrix} x_i^F & x_i^E \end{bmatrix}$, which includes both the spatial (fMRI) and temporal (EEG) features side by

side, and likewise for a source vector $\mathbf{s}_i = \begin{bmatrix} s_i^F & s_i^E \end{bmatrix}$. If we imagine that there are only two sources of the coupled data features, then the unknown sources would correspond to two row vectors ($\mathbf{s_1}$, $\mathbf{s_2}$) encoding the degree to which each coupled source was expressed in each subject. The mixing matrix \mathbf{A} (in this simple case) would be two column loading parameters encoding the spatiotemporal pattern this coupled source would cause over features. In reality, of course, there are many different sources of variability between subjects. The resulting update equation for the algorithm to compute the shared unmixing matrix \mathbf{W} (*i.e.* the inverse of \mathbf{A}) and the fused fMRI and ERP sources, \mathbf{u}^F and \mathbf{u}^E , is as follows:

$$\Delta W = \eta \left\{ I - 2y^F \left(u^F \right)^T - 2y^E \left(u^E \right)^T \right\} W, \tag{1}$$

where $\mathbf{y}^F = g(\mathbf{u}^F)$, $\mathbf{y}^E = g(\mathbf{u}^E)$, and $g(x) = 1/(+1 e^{-x})$ is the nonlinearity in the neural network 53. The basic idea underlying independent component analysis (ICA) is to assume that these sources are independent and distributed sparsely, encoded by the cumulative density function g. These assumptions allow one to estimate the unmixing matrix W in a maximum likelihood sense, without knowing the sources. When applied to the data features this provides estimates of the sources for each subject. Note that the sources have to explain both the data modalities and therefore this estimation represents the inversion of a multimodal fusion model, under sparsity and independence constraints. Notice also that this procedure is entirely data-led in the sense that the model does not know whether each subject belongs to one group or another. Therefore, it may identify group differences that would be missed in conventional between-group comparisons. Meanwhile, our reason for using a single optimal unmixing coefficient to maximize the joint likelihood function is that it makes intuitive sense not to compute the parameters independently, because the ICA results from the two different measures are derived from the same participant 2, 38, 46. We thus have a single W that fuses together the joint source (or alternatively, the basis vector common to the two measures). The main advantage of this approach is that maximizing the joint likelihood function provides a different (and more reasonable) solution from one that does not utilize the joint statistics.

Component Estimation

The number of independent components in the joint data was estimated to be twelve, using a method based on the minimum description length criteria 55, 56. Independent components were estimated, and ranked by their contribution to the average ERP time courses by first regressing the components onto the average ERP data, then computing the maximum absolute peak of the fitted time courses. A leave-one-out cross-validation approach 38, 57 was used to assess the robustness of the results; mean results are reported.

Analysis of Patient and Control Data

For comparison with the jICA results, we averaged the ERP data time locked to the target stimuli and also carried out a standard random-effects analysis within SPM2 by computing voxel-wise t-tests between the patient and control fMRI contrast images 58, 59. To ensure similar consideration (weights) for both measures, the fMRI and ERP features were normalized by dividing each modality by its average standard deviation and then the ERP data (timecourses) were upsampled to the same level as the fMRI (voxels). We interpolated the ERP data to provide a balanced representation of ERP and fMRI features. The fMRI dimension is about 75000 voxels, while the ERP dimension is about 1000 timepoints. We show in extensive simulations in 37 that this works well. The ¡ICA procedure was then performed on the fMRI/ERP joint data from the patient and control groups together. Correspondingly, within the comparison groups (patient or control), the identification of components with shared loading parameters, and the comparison of the associated maps, becomes a key means to identify couplings between brain image components of different measures of data; while between the comparison groups (patient and control), the differences of amplitude, latency and location of each data component become a significant evidence of the variation of schizophrenia patients from healthy controls. After the jICA analysis, we tested within each component for a significant difference between patients and controls using a two-sample t-test. Only one significant component was found and interpreted (p<0.001, see 'Joint ICA Analysis') (Figure 3). We further examined the joint data using a cross-modality 2D histogram analysis. Signals that were significant in the jICA

analysis for either of the two measures were used to generate a joint histogram of the fMRI and ERP data. These histograms were examined in group averages along with marginal distributions (Figure 4).

RESULTS

Behavioral Data

Performance on the auditory oddball task was nearly identical within the fMRI and ERP but differed between control and patient groups. The number of trials was 50 for both ERP and fMRI sessions. Mean and standard deviations are reported for a) reaction time, controls (fMRI 430.7 \pm 90.4ms; ERP 431.5 \pm 93.6ms, p>0.9 paired t-test) and patients (fMRI 518.3 \pm 116.5 ms; ERP 577.7 \pm 149.0ms, p>0.4 (0.3667) paired t-test) and b) accuracy for target detection, controls (fMRI 99.5 \pm 0.01 percent; ERP 99.5 \pm 0.02 percent, p>0.99 paired t-test) and patients (fMRI 88.9 \pm 16.5 percent; ERP 93.2 \pm 14.4 percent, p>0.4 (0.4390) paired t-test).

Group Analysis

FMRI SPM contrast images (controls: Figure 2 Left, patients: Figure 2 middle) and ERP group averaging (Figure 2 Right) are shown for the target stimuli. Translation and rotation corrections for each participant did not exceed half a voxel (i.e., 2mm) or 2.0 degrees, respectively. We also qualitatively examined each statistical contrast image to ensure there were no obvious motion artifacts (i.e., edge artifacts were not apparent) 38, 46. There was no significant difference in movement between patients and controls. We applied standard random-effects analyses by entering the features into a voxel-wise one-sample t-test for patients and controls separately (p<0.001, corrected for multiple comparisons using the false discovery rate (FDR; 60). The ERP plot was generated by averaging the data time locked to the target stimuli. Results are largely consistent with previous findings for both measures 4, 34, 61. The fMRI data show main reductions in bilateral frontal and temporal lobes, inferior parietal lobe, cerebellum, plus motor planning and execution regions. The ERP data show significant reductions at N1, N2 and P3 peaks (marked on the figure).

Joint ICA Analysis

Results from the jICA analysis of both measures are presented in Figure 3. Among twelve components for both groups, only one joint component was found to distinguish groups using a two-sample t-test (p<0.001) on patient and control loading parameters, which we interpret as a difference in the degree/magnitude of two linked brain functional features (fMRI/EEG) in the two groups. This identified component shows a clear difference in fMRI at bilateral fronto-temporal regions implicated in schizophrenia (Figure 3 left) and in ERP in times during the N2 (MMN/N2b, see in the discussion) peak (Figure 3 right) which have been previously implicated in patients. It is important to note that the maps of controls and patients separately show a main effect, whereas the statistical comparison for the joint analysis is testing for a difference between groups. Talairach coordinates for the fMRI/ERP jICA analyses are presented in Table 2.

To examine the joint task activity in more details, a joint histogram was computed as follows, a similar strategy that applied in our old studies 2, 46. Signals surviving the threshold for the fMRI of the joint source were sorted in descending order by the component voxel values (the same was done for signals in the ERP part of the joint source). This procedure resulted in two sets of signal coordinates. Histograms were then generated by pairing these two signal sets. For example, the first point for Participant 1 is the voxel value for the fMRI activation data (at the position that is maximum in the fMRI part of the jICA source) versus the signal value for the ERP activation data (at the position that is maximum

in the ERP part of the jICA source). Then, we computed the within-group average of the histograms for the control and the patient groups (shown in Figure 4 with the controls in the top middle and the patients in the top right). The 2-D histogram can be considered an estimate of the joint distribution function for the two measures (e.g., p(ffmri, ferp), where ffmri,erp indicates the signal amplitude for the fMRI or ERP, respectively). We also computed the marginal estimated distributions p(ffmri) and p(ferp) (Figure 4 top left, bottom). Note that both fMRI and ERP are showing a group difference as seen by the marginal distributions for the two modalities. In the marginal histograms it is clear that controls (yellow) tend to have higher fMRI activation, whereas patients (cyan) tend to have higher ERP (positive) values. This is also visible on the group average 2D histograms in Figure 4 (the control histogram is located above and to the left of the patient histogram).

DISCUSSION

Schizophrenia is hypothesized to be a disease involving impaired brain interaction 62. A number of explanatory models have been proposed, with many studies implicating regions in temporal lobe, cerebellum, thalamus, basal ganglia, and lateral frontal regions 62-66. Discoordination 63, heteromodal association 67, 68 as well as fronto-temporal disconnection models 69 have been suggested. Recent work attempts to capture both spatial and temporal properties of neuronal activity and shown promising results 70-75. Examining the correspondence between fMRI and EEG (ERP) combines the strengths of both techniques and provides a more detailed probe into human brain function. However, the gap between these two different measures still presents an issue due to the technical difficulty and computational complexity. Here, we demonstrate a new technique for studying the linked fMRI/ERP signals impacted by schizophrenia, using joint independent component analysis (iICA) of separate recordings of the same subjects. Our previous work 38 focused on using this approach for localization and chronometry of target detection in healthy participants. In this work, using the jICA algorithm, we focus upon joint ERP/fMRI sources which differentiate schizophrenia patients from healthy controls during the performance of an auditory oddball task. It is important to emphasize is that in this paper we are studying the linkage between ERP/fMRI signals (specifically we identify linear relationships between the two data sets using a data driven approach) and there has been very little work on this in schizophrenia.

Our novel joint spatiotemporal analysis revealed several interesting findings compared to traditional analyses. First, consistent with our hypotheses, the jICA results identified fMRI group differences in bilateral temporal and frontal regions activated by the auditory oddball target stimulus, tightly associated with the N2 complex in ERP time course. The N2 peak latency was between 180-200ms window, and likely contains contributions from both the MMN and the N2b subcomponents 25⁻30, 76, 77 as well as the possible latency shifts of the those peaks 78. Schizophrenia patients demonstrated significant decreased amplitude for the linked fMRI spatial component and ERP temporal component. This finding suggests that bilateral fronto-temporal neuronal activity may serve as a pathophysiological substrate for changes in the N2 peak of ERP as probed by the auditory oddball target stimuli in schizophrenia. This idea is consistent with previous research 21, 79, 80. Additionally, the region showing the largest group difference is associated with the N2 component in the jICA analysis (the second largest is associated with the P3 component, although it did not reach significance). This finding emphasizes the pathological importance of N2 generators. To date, schizophrenia, researchers have focused mainly on the P3 as the most significant biomarker of decision making, with many reports, especially in chronic schizophrenia 4, 14, 15, 17, 81. The N2 performs a variety of functions in mismatch detection and cognitive control 9, and although likely serving as a marker of psychosis classification, remains largely unstudied in this group 21. Our finding suggests that the N2 component is an

electrophysiological marker of disturbance in stimulus classification and attention processes in schizophrenia. Our results are consistent with O'Donnell and Kayser's reports 21, 25, 26.

Our approach provides several advantages. 1) Compared to the traditional fMRI activation region and direct EEG peak inspections, our method cuts down the spatial and temporal overlaps of different brain areas at different time course activities. 2) Contrasted with other common methods, e.g. PCA, the derived joint fMRI/ERP components from our approach have spatial/temporal projections that are maximally higher-order independent, distinct but not necessarily orthogonal. 3) Our approach does not require the use of a threshold for the fMRI data nor assumptions about the number of dipoles or the modeling of dipole fitting for EEG, which relaxes the constraints of the algorithm and is computationally straightforward 4) We illustrate an approach for combining data from fMRI and ERP measures, two methods with different strengths and advantages, in a symmetric analytic framework that does not favor one modality, but reveals changes which may manifest in fMRI only, ERP only, or both fMRI and ERP. 5) We perform a joint decomposition of both measures which are linked or fused by a common mixing parameter. This enables investigators to explore the relationship between electrophysiology and hemodynamic cognitive processes. 6) Our approach also separates the data into joint components, each with an fMRI and an ERP portion. Decomposing the data into joint components may provide a useful way to examine component specific differences, e.g., patient vs control groups or modified task versions. Here, we utilize a feature-based approach, providing a straightforward way to take advantage of data modeled at the subject level. These features are then queried for shared dependence, which is not detectable with a simple voxelwise subtractive or conjunction approach. 7) The shared mixing coefficient provides a way to examine individual or group differences in coupling. Currently, we chose a priori to analyze only the component(s) that revealed a statistical difference between groups. In future work it would be interesting to develop approaches for understanding the full ICA decomposition (e.g., to examine all the components). Additionally, given previous interest in laterality differences in schizophrenia 82, 83, it would be interesting to examine the laterality of these joint sources.

There are some limitations to our approach. First, given the heterogeneity of the spatial and temporal data in schizophrenia, it is important to address issues of statistical power for a joint analysis, which may be different than for an individual analysis. This is especially important if findings from a joint analysis are to become clinically relevant 84. Second, the potentially useful information from the first level may be not neglected in the source separation when we are carrying out a joint second-level (group) analysis of features, and we have begun working towards 1st level single trial analysis with parallel and joint ICA models (Eichele IJP 2008, Moosmann IJP 2008). Third, in the current framework, for practical reasons, we assume that both voxels and timepoints are independent and identically distributed. Although, most ICA models for fMRI/ERP data perform quite well under this assumption, it can be potentially useful to incorporate some additional prior information (such as correlation) as well as to include different distributions for different features into the model. We incorporated this attempt in our recent papers 85, 86. Fourth, in this work we analyze data from a single electrode. We have extended our methods to incorporate multiple EEG electrodes in some prior work 37, 39, 40. Fifth, we focus on analyzing target response in this study, it would be reasonable to compare the responses to target tones with those of non-target tones (standard and novel) in future, in order to delineate brain regions which are specically related to attending and responding to target tones. Sixth, the ERP and fMRI data were acquired in two separate sessions, instead of a concurrent session, which might induce the differences into the analysis. In order to minimize the difference, we collected the fMRI and ERP data on the same day, using identical stimuli and the order of the sessions was counterbalanced between individuals; and we verified that all subjects had nearly identical

performance on the task for both the ERP and fMRI sessions. The same strategy has been used successfully in several other published studies (see, e.g. 38).

It is interesting to observe the differences between separate and joint analyses. When analyzing the fMRI and ERP data separately, fMRI activity shows significant reductions in frontal and temporal lobes plus cerebellum, while the ERP data show significant reductions at N1, N2 and P3 peaks (see Figure 2). In the jICA analysis, we see only one significant N2 reduction associated with the fMRI activity reduction in fronto-temporal regions.

In summary, we have demonstrated a novel method for examining joint hemodynamic and electrical data to visualize the neural systems involved during different portions of the auditory "oddball" target detection response. This approach has enabled us to ask novel questions about fMRI/EEG data and revealed several interesting findings in an application to data collected from healthy controls and patients with schizophrenia that were missed by a standard analysis approach, which may greatly help us diagnose/understand the pathophysiology of schizophrenia. The development of models for jointly analyzing multimodal data has been largely overlooked and may be a useful tool for assessing how brain function during different cognitive probes and in different regions can vary systematically between measures.

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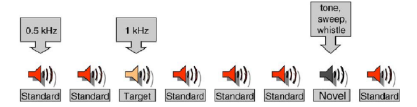


Figure 1. Auditory oddball paradigm: auditory oddball event-related fMRI task.

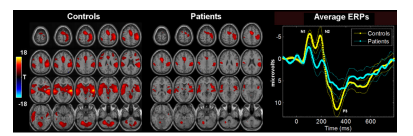


Figure 2. FMRI and ERP group analyses: Group fMRI SPM contrast images(left) and ERP (right) results for the target response. The SPM2 software was used to generate a contrast image of target-related activation for each subject. These images are then entered into a one-sample, voxelwise t-test and thresholded at p<0.001 (corrected for multiple comparisons) to produce the resulting image with T-values shown in color. The ERP plot was generated by averaging the data time locked to the target stimuli. Standard error is shown and peaks are labeled with the standard naming convention (e.g. N1 is the first negative peak).

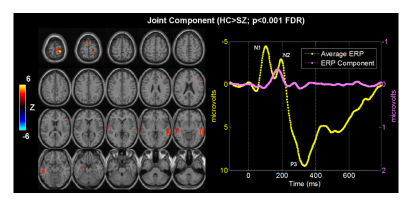


Figure 3. FMRI/ERP jICA analysis: The only joint component which showed significantly different loading parameters (p<0.001) for patients versus controls. Left: thresholded fMRI part of the joint component showing bilateral temporal and frontal lobe regions. Right: average ERP (yellow) plots along with the ERP part of the identified joint component (pink).

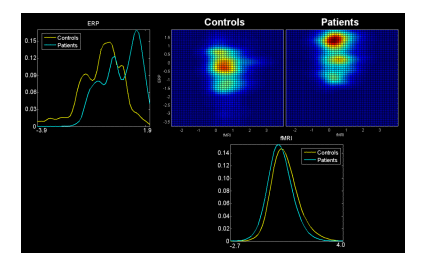


Figure 4. Cross-modality 2-D histograms: Joint 2-D histograms for ERP/fMRI identified in the joint independent component analysis (jICA). Group average histograms (with controls in the top middle and patients in the top right) are provided along with the marginal histograms for the ERP (top left) and for the fMRI (bottom). In the marginal histograms it is clear that controls (yellow) tend to have higher fMRI activation, whereas patients (cyan) tend to have higher ERP (positive) values.

Table 1

Demographic details.

	Controls	Patients
Age (years [mean (SD)]	41 (14)	38(11)
Gender (male : female)	15:8	11:5
Mean NART IQ (SD)	106.20 (6.09)	99.72 (9.37)
Handedness (R : L)	21:2	13:3

L, left; R, right; NART, national adult reading test; IQ, intellignece quotient.

 Table 2

 Talairach coordinates for auditory oddball fMRI/ERP jICA analysis

Anatomic label	Brodmann area*	Volume (cc): L/R	Max T (x,y,z): L/R
Increases			
Superior frontal gyrus	6	0.5/2.6	4.3 (-3, 38, 62)/5.5 (24, 1, 69)
Superior temporal gyrus	21, 22, 38, 13	1.9/2.8	4.0 (-53, 2 -10)/4.4 (62, -12, -4)
Middle temporal gyrus	21, 22	0.9/1.9	4.0 (-50, 2, -10)/4.4 (62, -15, -4)
Middle frontal gyrus	6, 46, 10, 47	0.0/1.1	na/4.0 (33, 0, 64)
Inferior frontal gyrus	45, 47, 46	0.9/0.3	3.9 (-65, 18, 2)/3.3 (56, 53, 6)
Precental gyrus	44	0.1/0.1	3.8 (-65, 15, 2)/3.2 (62, 12, 5)
Decreases			
Middle frontal gyrus	6, 9	0.0/0.2	ns/3.6 (50, 2, 41)
Precental gyrus	6	0.0/0.2	ns/3.4 (50, 2, 44)
Thalamus		0.1/0.0	3.1 (-3, -20, 7)/ns
Cingylate gyrus	31	0.0/0.0	3.0 (-24, -39, 38)/ns

Voxels above the threshold for Fig. 2 were converted from the Montreal Neurological Institute to Talairach coordinates and entered into a database to provide anatomic and functional labels for the left (L) and right (R) hemispheres. Both increasing (top) and decreasing (bottom) voxels are reported. The volume of activated voxels in each area is provided in cubic centimeters (cc). Within each area, the maximum t value and its coordinate are provided.

NS, not significant.

Brodmann areas (BA) are only approximate, based upon the Talairach Atlas.