Coupling of theta activity and glucose metabolism in the human rostral anterior cingulate cortex: An EEG/PET study of normal and depressed subjects

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Abstract
In rodents, theta rhythm has been linked to the hippocampal formation, as well as other regions, including the anterior cingulate cortex (ACC). To test the role of the ACC in theta rhythm, concurrent measurements of brain electrical activity (EEG) and glucose metabolism (PET) were performed in 29 subjects at baseline. EEG data were analyzed with a source localization technique that enabled voxelwise correlations of EEG and PET data. For theta, but not other bands, the rostral ACC (Brodmann areas 24/32) was the largest cluster with positive correlations between current density and glucose metabolism. Positive correlations were also found in right fronto-temporal regions. In control but not depressed subjects, theta within ACC and prefrontal/orbitofrontal regions was positively correlated. The results reveal a link between theta and cerebral metabolism in the ACC as well as disruption of functional connectivity within frontocingulate pathways in depression.

Descriptors: Anterior cingulate cortex, Limbic system, Theta rhythm, Positron emission tomography, Electroencephalogram, Source localization

Following the initial description by Jung and Kornmüller (1938), much progress has been achieved regarding the study of neuronal generation of theta activity. Although most nonhuman studies have focused on the septo-hippocampal system, theta has been recorded in numerous other regions, including the anterior cingulate cortex (ACC), entorhinal cortex, hypothalamus, superior colliculus, medial septum, mammillary bodies, anterior thalamus, and amygdala (Bland & Oddie, 1998; Vinogradova, 1995). Theta activity may serve as a gating function on the information processing flow in limbic regions, as some evidence suggests it facilitates transmission between different structures (Vinogradova, 1995).

In humans, two different manifestations of theta rhythm have been described (Schacter, 1977). The first exhibits a widespread scalp distribution and has been associated with decreased alertness and impaired information processing, typically during drowsy states. The second is associated with a frontal midline distribution and has been linked to alert states characterized by focused attention, mental effort, and effective stimulus processing. Recent human studies have further suggested that either medial prefrontal cortex (PFC) sources centered on the ACC (Asada, Fukuda, Tsunoda, Yamaguchi, & Tonoike, 1999; Gevins, Smith, McEvoy, & Yu, 1997; Ishii et al., 1999), or sources in dorsolateral prefrontal regions (Sasaki et al., 1996) could account for scalp-recorded frontal midline theta activity. These results concur with animal data suggesting that theta associated with alert states arises locally within the ACC and not via volume conduction from the hippocampus (Feenstra & Holsheimer, 1979; Holsheimer, 1982).

Indirect evidence that the human ACC may be involved in the generation of frontal midline theta activity accrues from a comparison of the electrophysiological (EEG/MEG) and hemodynamic neuroimaging literature. In particular, focused attention (hemodynamic: Davis, Taylor, Crawley, Wood, & Mikulis, 1997; EEG: Asada et al., 1999), task difficulty (hemodynamic: Murtha, Chertkow, Beauregard, Dixon, & Evans, 2003).
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2-deoxyglucose method originally developed by Sokoloff et al., (1977), utilizing reflects synaptic (particularly, presynaptic) activity (Jueptner & Weiller, 1995) and scalp EEG signals reflect current associated with excitatory and inhibitory postsynaptic potentials (Nunez & Silberstein, 2000), a link between the two modalities is expected.

Although many EEG/MEG studies have described task-related changes in frontal midline theta, to date none have investigated the link between such activity and metabolism in the ACC during task or baseline recording, nor the relations between theta activity in the ACC and frontal regions, or whether putative functional connectivity within frontocingulate pathways may be disrupted in major depression. The present, methodologically oriented investigation aims to investigate these three topics. To this end, concurrent measurements of brain electrical activity and glucose metabolism were performed in a heterogeneous sample comprised of healthy control and depressed subjects during standard baseline (resting) recording conditions. The use of a tomographic source localization technique for the EEG data based on realistic head geometry and probabilistic brain atlases enabled the comparison between brain electrical and metabolic activity on a voxelwise basis. Based on the literature reviewed above, a positive association between theta electrical and metabolic activity in the ACC as well as between theta activity in the ACC and dorsolateral PFC were hypothesized. Also, in light of hypothesized dysfunctions in frontocingulate networks in depression, a disruption in functional connectivity between ACC and dorsolateral PFC was expected in subjects suffering from major depression.

Methods

Participants

Participants were recruited through advertisements in local media for a larger study investigating the functional neuroanatomy of major depression (Abercrombie et al., 1998; Pizzagalli et al., 2001). For the present analysis, all participants with both EEG and PET data were included: 17 participants with major depression (mean age: 33.3 years, SD: 10.5; 7 women), and 12 control subjects (35.2 ± 11.7; 6 women). Based on the Structured Clinical Interview for the DSM-III-R, Patient Edition (SCID-P; Spitzer et al., 1992), which was modified to make Diagnostic and Statistical Manual for Mental Disorders (DSM-IV) diagnoses, depressed subjects met DSM-IV criteria for major depressive disorder (American Psychiatric Association, 1994), and had no history of mania, psychosis, or other Axis I disorders, with the exception of specific phobias and social phobia secondary to major depressive disorder. Additionally, a family history of mania or psychosis led to study exclusion. For the depressed participants, the mean Beck Depression Inventory (BDI; Beck, Ward, Mendels, & Erbaugh, 1961) and Hamilton Depression Rating Scale (HDRS; 17-item version; Hamilton, 1960) scores were 30.13 (SD: 6.79) and 19.0 (SD: 5.39), respectively. According to the BDI, 8 of the patients were “severely depressed” (BDI > 30), and 8 were “mild/moderately depressed” (13 < BDI < 29; Beck & Steer, 1987). Control participants had no current or past Axis I pathology in themselves or first-degree relatives. Additional details about participant recruitment, assessment, and characteristics have been reported elsewhere (Pizzagalli et al., 2001).

Procedure

After obtaining written informed consent, participants underwent concurrent FDG–PET and EEG recording, following a protocol approved by the Human Subjects Committee of the University of Wisconsin. Injection of FDG was scheduled between 11:00 and 13:30 hr. Participants fasted for 5 hr prior to the injection. After application of scalp EEG electrodes, two 22-gauge intravenous catheters (injection and blood draw) were inserted for the PET procedure, one in the antecubital fossa of the right arm and the other in a vein in the posterior aspect of the left hand. To allow rapid sequential sampling of arterialized venous blood, the left hand was placed into a heated handwarmer.

The inclusion of both healthy control and depressed subjects likely led to a larger range of ACC activation, as depression has been associated with dysfunctional activity in the ACC (for review, see Davidson et al., 2002). A potentially large range of ACC activation appeared optimal for the correlational approach used in the current study.
(Phelps et al., 1979); the right hand was also heated to equate thermal stimulation on both sides. Participants were instructed to remain still, sit quietly, and relax but stay awake during uptake. To obtain an initial plasma glucose level, a 1- to 2-ml blood sample was drawn. Subsequently, approximately 5 milliCuries (range: 3.8–5.7 mCi) of FDG were administered by means of a bolus injection into the intravenous line in the right arm. Arterialized blood samples were drawn from the left hand throughout the 30 min following injection. Each blood sample was centrifuged, 0.5 ml of plasma was withdrawn, and the radioactivity level was measured in a well counter to obtain the blood time course of radioactivity.

EEG recording consisted of 10 contiguous 3-min trials (5 with eyes open and 5 with eyes closed, alternating according to a counterbalanced order across participants) covering the 30 min required for radiotracer injection and uptake. Consequently, the EEG and PET data reflect the same functional brain state. Alternation of eyes open and eyes closed trials was selected as previous studies from our laboratory (Tomarke, Davidson, Wheeler, & Kinney, 1992) have suggested that aggregating across eyes open and eyes closed trials results in the most reliable estimates of baseline EEG activation patterns. Following a 10-min break in which participants were encouraged to void their bladders, participants were positioned on the PET scanner.

Apparatus and Physiological Recording

EEG data. Twenty-eight-channel EEG was recorded referenced to the left ear with a modified lycra electrode cap (Electro-Cap International, Inc.). Horizontal and vertical electrooculogram (EOG) channels were also recorded. Impedances were below 5 KΩ (EEG) and 20 KΩ (EOG). Signals were amplified with a Grass Model 12 Neurodata system using Model 12C preamplifiers (bandpass: 1–300 Hz; 60-Hz Notch filter), filtered with MF6 digital anti-aliasing low-pass filters set at 100 Hz, and digitized at 250 Hz.

PET data. [18F]-fluoride was produced by a CTI RDS Cyclotron (Knoxville, TN, USA), and FDG was synthesized using the modified method of Hamacher (Hamacher, Coenen, & Stocklin, 1986). A General Electric/Advance PET scanner (Milwaukee, WI, USA) was used to acquire PET data (intrinsic in-plane and axial resolutions of approximately 5 mm full-width half-maximum [FWHM]). The PET scan started approximately 50 min after injection and consisted of a 30-min two-dimensional emission scan (15.2-cm field of view, 35 transaxial planes), a 10-min three-dimensional emission scan, and a 10-min transmission scan.

Data Reduction and Analyses

EEG data. Off-line, data were screened for artifacts caused by blinks, eye movements, muscle activity, or technical problems. Only epochs with artifact-free data across all channels were considered for the analyses. All available artifact-free 2,048-ms EEG epochs from both the eyes open and eyes closed trials were extracted (150.83 ± 73.22 vs. 168.21 ± 83.98, respectively, with no differences between eyes open and eyes closed trials, t26 = 0.82, n.s.), rederived to the average reference, and then subjected to standard spectral analyses via Discrete Fourier Transform (DFT) using a boxcar windowing (Brillinger, 1981). Subsequently, the three-dimensional intracerebral current density distribution for the theta band (6.5–8.0 Hz) was estimated using Low Resolution Electromagnetic Tomography (LORETA; Pascual-Marqui, Michel, & Lehmann, 1994; Pascual-Marqui et al., 1999). This technique has recently received important cross-modal validation, as summarized elsewhere (Pizzagalli et al., 2001). Without postulating a prespecified number of generating sources, LORETA estimates location(s) of electrical source activity by assuming similar activation among neighboring neuronal sources, an assumption implemented by computing the “smoothest” of all possible activity distributions. The present implementation uses a three-shell spherical head model (Ary, Klein, & Fender, 1981) and EEG electrode coordinates derived from cross-registrations between spherical and realistic head geometry (Towle et al., 1993). Both the head model and the electrode coordinates were registered to the digitized MRI available from the Brain Imaging Centre, Montreal Neurologic Institute (MNI305; Collins, Neelin, Peters, & Evans, 1994; Evans et al., 1993). The solution space (2,394 voxels; voxel size: 7 × 7 × 7 mm³) was restricted to cortical gray matter and hippocampi, as defined by the digitized MNI probability atlases. Consequently, all coordinates reported in the present study are in MNI space. Subsequently, the Structure–Probability Maps atlas (Lancaster et al., 1997) was used to label gyrus and Brodmann area(s). Because the Structure–Probability Maps atlas requires Talairach coordinates, and due to the imperfect match between the MNI and Talairach templates, MNI coordinates were converted to Talairach coordinates (Talairach & Tournoux, 1988) using the corrections proposed by Brett, Johnsrude, and Owen (2002; see also http://www.mrc-cbu.cam.ac.uk/Imaging/mnispace.html).

LORETA values represent the power, that is, squared magnitude, of the computed intracerebral current density (unit: amperes per square meter, A/m²).

PET data. Using the scanner manufacturer’s software (General Electric, Milwaukee, WI, USA), the two-dimensional PET data were reconstructed with calculated attenuation correction to 1.17 × 1.17 × 4.25 mm³ voxels, and converted to parametric images of an influx constant (Ki, 1/s) using the measured radioactive blood time course and a variation of the Sokoloff method (Phelps et al., 1979). They were then spatially normalized with SPM99 (Friston, Worsley, Frackowiak, Mazziotta, & Evans, 1994) to the template used by LORETA (MNI305), converted to 2 × 2 × 2 mm³ voxels, and smoothed with a 6 × 6 × 6 mm³ Gaussian kernel to match the estimated resolution of LORETA (~10 mm³). Using in-house software that considered weighting factors derived from the fractional volume of a PET voxel within a given LORETA voxel, PET data were resampled to voxels having the same size and center location as the LORETA voxels (Oakes et al., 2001). PET data are expressed in units of µg/min/100 cc.

Statistical Analyses

Voxelwise correlation analyses. Two sets of data were analyzed. In the first approach, which emphasizes subject-to-subject global variations in activation, neither the LORETA nor the PET data were intensity normalized. In the second approach, subject-to-subject global variations were removed in both LORETA and PET data by intensity normalization. The average value across voxels for each subject’s volume was computed, and then each image volume was scaled so its mean was the same as the overall mean. This approach emphasizes regional differences within the brain. At each voxel, a correlation was computed

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between current density for the theta band and glucose metabolism for both normalized and nonnormalized data. Due to different distributions of the LORETA and PET data, Spearman’s rank correlations ($\rho$) were computed.

Region-of-interest analysis. In addition to the whole-brain correlational analyses, a region-of-interest (ROI) analysis was performed. In a recent EEG study, we found that theta activity in a rostral ACC cluster was specifically associated with favorable treatment response in depression (Pizzagalli et al., 2001). In that study, this association was postulated because (a) metabolic rate in the rostral ACC had been previously associated with treatment response in PET studies (Mayberg et al., 1997), and (b) the ACC had been proposed as a possible neuronal source of theta activity in the human brain (Asada et al., 1999; Gevins et al., 1997; Ishii et al., 1999). In the present analysis, LORETA and PET activity was averaged from the rostral ACC cluster differentiating low and high treatment responders (Figure 1C). For both normalized and nonnormalized data, Spearman’s rank correlations assessed the relation between the ROI values of the two modalities.

Functional connectivity analyses. To test the role of theta activity on other regions, Spearman’s rank correlations were computed between theta activity in the ACC and other regions showing significant positive correlations between the EEG and PET data (see Results). For each pair of regions, Fisher’s tests for independent correlations (Fisher, 1921) were run to assess whether these patterns of correlations were similar for depressed and control subjects.

In addition, to investigate in more detail whether depression may be associated with dysregulation in frontocingulate networks, the functional connectivity between the ACC and the rest of the brain was interrogated for depressed and control subjects separately. To this end, Spearman’s rank correlations were run between theta current density within the ACC (averaged within cluster #1, Figure 2; see Results) and each voxel in the LORETA brain volume. Again, to test whether depressed and control

Figure 1. Sagittal slices centered on the ACC showing results of voxelwise Spearman’s rank correlation between current density in the theta band (6.5–8.0 Hz) and glucose metabolism in 29 subjects using (A) nonnormalized data and (B) normalized data. For display purposes only, maps have been thresholded at $p < .05$ (uncorrected). C: Location and spatial extent of the rostral ACC region (see green) previously found to be associated with treatment response in major depression (Pizzagalli et al., 2001).
Subjects may differ in functional connectivity between the ACC and other regions, Fisher’s tests for independent correlations were run at each voxel.

Overall, for regions with a priori hypotheses (e.g., ACC), significant results were accepted at \( p < 0.05 \). For regions without a priori hypotheses, a \( p \) value of .01 was selected, as previous LORETA studies using randomization procedures to estimate the false-positive rate under the null hypothesis have found that this threshold is adequate for protecting against Type I errors (Pizzagalli et al., 2002).

**Results**

*Voxelwise Correlation Analyses*

**Nonnormalized LORETA and PET data.** Figure 1A shows the results of the whole-brain correlation analysis for the entire sample \((n = 29)\) between theta current density and glucose metabolism at each voxel when no normalization was applied to either the EEG or PET data. Out of the 2,394 correlations, 2,317 were positive and 213 of them were significant at \( p < 0.05 \). No significant negative correlation emerged. Although positive correlations between theta current density and glucose metabolism were present in the ACC, they were not specific to this region. These data include the global activity for each subject, which is considered to be a nuisance variable; consequently, subsequent analysis focused on the normalized data.

**Normalized LORETA and PET data.** Figures 1B and 2 and Table 1 summarize the results of the whole-brain correlation analysis for the entire sample \((n = 29)\) between theta current density and glucose metabolism at each voxel when normalization was applied to both the EEG and PET data. A cluster in the rostral ACC (Brodmann areas (BAs) 24, 32) showed the largest (13 voxels, volume: 4.46 cm\(^3\)) and second strongest positive correlation between theta current density and glucose metabolism \((r = 0.52, p < 0.005; \ X = -3, \ Y = 45, \ Z = 1; \ BA \ 32; \ see \ cluster \ #1, \ Figure \ 2)\). The strongest positive correlation was observed in the right middle frontal gyrus \((r = 0.54, p < 0.0025; \ X = 39, \ Y = 17, \ Z = 29; \ BA \ 9; \ see \ cluster \ #2, \ Figure \ 2)\), in a cluster that was, however, comprised of only 2 voxels (0.69 cm\(^3\)). As listed in Table 1, only two further clusters showed reliable positive correlations, one involving the right middle and superior temporal gyri \((r = 0.48, p < 0.01; 4\) voxels, 1.37 cm\(^3\); see cluster #4, Figure 2) and the other the right superior frontal gyrus \((r = 0.48, p < 0.01; 4\) voxels, 1.37 cm\(^3\); see cluster #4, Figure 2). For each of these clusters, the highest correlation was associated with a \( p \) value smaller than .01, suggesting adequate protection against Type I errors. The strongest negative correlation was observed in the right precentral gyrus \((r = -0.67, p < 0.0001; \ X = 46, Y = 3, \ Z = 8; \ BA \ 44; \ 4\) voxels, 1.37 cm\(^3\); see cluster #5, Figure 2), whereas the largest cluster with negative correlations encompassed the cuneus, precuneus, culmen, posterior cingulate and lingual gyri (78 voxels, 26.75 cm\(^3\); BAs 18, 19, 23, 29, 31; see cluster #6, Figure 2).

**Region-of-Interest Analysis**

As shown in Figure 1C, the rostral ACC cluster (14 voxels, 4.80 cm\(^3\)) previously found to be associated with treatment response in depression (Pizzagalli et al., 2001) overlapped with the region with the reliable positive correlations between LORETA and PET data (Figure 1B). For both the normalized
superior (and each voxel in the brain volume to examine putative
significantly lower functional connectivity between the ACC and
tions. Thus, compared to controls, depressed subjects showed
depressed and control subjects, those for the right superior
temporal gyri (.19 vs. .52), right middle (7)
were similar for depressed (7)
showing significant positive correlations between the EEG and
Functional Connectivity Analyses
To test the role of theta activity in the ACC on other regions
showing significant positive correlations between the theta current
and glucose metabolism in the rostral ACC, no reliable
reductions emerged for control or depressed subjects, or for
the entire sample, when repeating the ROI analyses for other
frequency bands, such as delta (1.5–6.0 Hz), alpha1 (8.5–
10.0 Hz), alpha2 (10.5–12.0 Hz), beta1 (12.5–18.0 Hz), beta2
(18.5–21.0 Hz), beta3 (21.5–30.0 Hz), and gamma (36.5–44 Hz),
all p’s > .05.

Table 1. Summary of Significant Positive Spearman’s Rank Correlations between Theta Current Density and Glucose Metabolism (n = 29)

<table>
<thead>
<tr>
<th>Region (cluster in Figure 2)</th>
<th>BA Side</th>
<th>No. voxels</th>
<th>Spearman ρ</th>
<th>X</th>
<th>Y</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anterior cingulate (#1)</td>
<td>24/32</td>
<td>Right</td>
<td>5</td>
<td>.433±.03</td>
<td>5.8±3.5</td>
<td>41.5±4.0</td>
</tr>
<tr>
<td>Middle frontal gyrus (#2)</td>
<td>6</td>
<td>Right</td>
<td>2</td>
<td>.473±.09a</td>
<td>42.5±4.9</td>
<td>17.0±0.0</td>
</tr>
<tr>
<td>Middle temporal gyrus (#3)</td>
<td>21</td>
<td>Right</td>
<td>7</td>
<td>.426±.04a</td>
<td>61.0±6.3</td>
<td>-11.0±12.1</td>
</tr>
<tr>
<td>Superior temporal gyrus (#3)</td>
<td>22</td>
<td>Right</td>
<td>4</td>
<td>.422±.03</td>
<td>53.0±5.7</td>
<td>3.0±5.7</td>
</tr>
<tr>
<td>Superior frontal gyrus (#4)</td>
<td>6</td>
<td>Right</td>
<td>4</td>
<td>.421±.03a</td>
<td>14.5±9.0</td>
<td>31.0±0.0</td>
</tr>
</tbody>
</table>

The mean (±SD) coordinates and mean (±SD) correlation of clusters showing significant positive Spearman’s rank correlations (ρ) are listed. Note that these correlations differ from the one reported in the main text, where the highest correlation within a given cluster is provided. The spatial extent (number of voxels exceeding the p<.05, uncorrected, n = 29), hemisphere, and Brodmann areas of these clusters are also provided. Coordinates in millimeters (MNI space), origin at anterior commissure; (X) = left (−) to right (+); (Y) = posterior (−) to anterior (+); (Z) = inferior (−) to superior (+).

*Highest correlation within the cluster is associated with p<.01, Spearman ρ>.475.

(p = .34, n = 29, p = .067) and the nonnormalized (p = .44,
n = 29, p<.025) data, a positive correlation emerged.

Highlighting the specificity of a link between theta activity and
and glucose metabolism in the rostral ACC, no reliable correlations emerged for control or depressed subjects, or for
the entire sample, when repeating the ROI analyses for other
frequency bands, such as delta (1.5–6.0 Hz), alpha1 (8.5–
10.0 Hz), alpha2 (10.5–12.0 Hz), beta1 (12.5–18.0 Hz), beta2
(18.5–21.0 Hz), beta3 (21.5–30.0 Hz), and gamma (36.5–44 Hz),
all p’s > .05.

Functional Connectivity Analyses
To test the role of theta activity in the ACC on other regions
showing significant positive correlations between the EEG and
PET data, Spearman’s rank correlations were computed between
theta activity in the ACC, on one hand, and, in the right middle
frontal gyrus (cluster #2, Figure 2), right middle and superior
temporal gyri (cluster #3, Figure 2), and the right superior frontal
 gyrus (cluster #4, Figure 2), on the other hand. ACC theta activity was significantly correlated with theta activity in the right
superior frontal gyrus (ρ = .66, n = 29, p<.001) and right middle
temporal gyrus (ρ = −.38, n = 29, p<.05), but not with theta activity in the right superior temporal (ρ = −.32, n.s.) or right
middle frontal (ρ = .30, n.s.) gyrri. These patterns of correlations were similar for depressed (n = 17) and control (n = 12) subjects, with one exception. Whereas the correlations for the right middle
frontal gyrus (.19 vs. .52), right middle (−.49 vs. −.16), and
superior temporal gyr (−.33 vs. −.13) did not differ between
depressed and control subjects, those for the right superior
frontal gyrus were significantly different (.37 vs. .89, z = −2.39,
p<.02), as assessed with Fisher’s test for independent correlations.
Thus, compared to controls, depressed subjects showed
significantly lower functional connectivity between the ACC and
the right superior frontal gyrus.

Spearman’s rank correlations were run between theta current
density within the ACC (averaged within cluster #1, Figure 2)
and each voxel in the brain volume to examine putative
dysregulation involving the ACC in depression. As evident in
Figure 3A, control subjects showed significant correlations
(p<.01) between the ACC and various regions in the prefrontal
and orbitofrontal cortex, such as the inferior (BAs 11/47), middle
(BAs 8/9/11), superior (BAs 8/9/10/11), and medial (BAs 6/8/9/
10/11) frontal gyrus, as well as the orbital gyrus (BA 11).2 In
control subjects, most of these significant correlations were
located in the right hemisphere (Figure 3A). To test whether this
laterality effect was statistically supported, the formula proposed
by Meng, Rosenthal, and Rubin (1992) was used to assess
whether the correlations between the ACC and right-sided
regions were significantly different from the correlations between
the ACC and left-sided regions. Right-sided regions were defined
as follows: first, left-sided regions displaying significant correlations were flipped by reversing the X coordinates (Pizzagalli et
al., 2002); second, those flipped left-sided regions were sub-
tracted from right-sided regions showing significant correlations
to eliminate spatial overlap between regions in the two hemi-
spheres with significant correlations. Subsequently, this spatially
unique cluster was projected back to the left hemisphere. Finally,
for both clusters, correlations were averaged and entered in the
formula proposed by Meng et al. (1992; see p. 173). Among the
four separate clusters tested, only the OFC cluster (see yellow
circle in Fig. 3A) showed significantly higher correlations than
the homologous region in the left hemisphere (pACC-right
OFC = .83, p<.01; pACC-left OFC = .49, n.s.; Z = 2.66, n = 12,
p<.02; note: pOFC-right OFC = .85, p<.01).

Depressed subjects, on the other hand, were characterized by
significant correlations that were restricted to more medial
regions (Figure 3B). Voxel-by-voxel Fisher’s tests confirmed that
the correlations between the ACC, on one side, and right
dorsolateral PFC, ventrolateral PFC, and orbitofrontal cortex,
on the other side, were significantly lower for depressed than control subjects (Table 2). Notably, apart from a single voxel in the
inferior occipital gyrus (X = 39, Y = −88, Z = −13; BA 18)
at which depressed subjects showed significantly higher correlations
than controls (z = 1.99, p<.05), significant differences between control and depressed subjects were restricted to anterior
regions of the brain (Y > 31) that involved the
dorsolateral PFC and orbitofrontal cortex. A closer look at

2In these analyses correlating theta activity averaged within a
subregion of the ACC (see cluster #1, Figure 2) with the rest of the
brain, significant results involving the ACC should not be further
considered. Due to the redundancy of these data and the smoothness
assumption implemented in the LORETA algorithm, neighboring voxels
in the ACC are highly correlated, which led to inflated correlations.
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Figure 3 reveals again that group differences in correlations between the ACC and PFC/OFC regions were predominantly restricted to the right hemisphere. To formally test whether this laterality effect was statistically supported, we extracted \( z \) values from voxels in the left hemisphere that were homologous to the ones listed in Table 2 (by reversing the \( X \) coordinates). Subsequently, \( z \) values at right- and left-sided clusters were tested by using the formula proposed by Rosenthal (1987; p. 182), \( Z' = \frac{(Z_{\text{right}} - Z_{\text{left}})}{\sqrt{2}} \). No significant differences emerged (all \( |z| < 1.30, \text{n.s.} \)), suggesting that the differences between depressed and control subjects in correlations between the ACC and right-sided regions were not significantly different from the differences between depressed and control subjects in correlations between the ACC and left-sided regions.

**Discussion**

Using an analytic approach involving a distributed source localization technique for EEG data based on realistic head geometry and probabilistic brain atlases, we found positive correlations between theta current density and glucose metabolism in rostral regions of the ACC (BAs 24, 32), as well as in right fronto-temporal regions, including the right middle (BA 9) and superior (BA 6) frontal gyri, and the right middle (BA 21) and superior (BA 22) temporal gyri. For the rostral ACC, the association between brain electrical and metabolic activity was absent for other classic EEG bands. Under normal conditions, glucose is the primary source of energy for the brain, and the coupling between neuronal activity and glucose utilization implies that spatially localized and temporally delimited increases in glucose utilization occur in association with neuronal activity in circumscribed brain regions subserving a specific function (Magistratti, 2000; Sokoloff, 1989). Consistent with the notion that neuronal activity leads to localized increases in glucose metabolism, the relations between strength of theta current density and glucose metabolism emerged especially when relative rather than absolute glucose metabolism was considered, where subject-to-subject global variations in both data sets have been removed and thus regional activations have been emphasized. Overall, these results suggest that in the alert resting state, higher cerebral metabolism is associated with greater EEG theta in territories of the human cortex that, on the basis of the animal neurophysiology, are expected to be regulated by limbic theta. Direct or indirect evidence for a critical role of the ACC in the

**Table 2. Summary of Significant Differences between Depressed and Control Subjects in Functional Connectivity between the ACC and Various Prefrontal and Orbitofrontal Regions**

<table>
<thead>
<tr>
<th>Region</th>
<th>BA</th>
<th>Side</th>
<th>( Z ) value</th>
<th>( X' )</th>
<th>( Y' )</th>
<th>( Z' )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superior frontal gyrus</td>
<td>11</td>
<td>Right</td>
<td>-3.04</td>
<td>18</td>
<td>45</td>
<td>-20</td>
</tr>
<tr>
<td>Inferior frontal gyrus</td>
<td>47</td>
<td>Right</td>
<td>-2.97</td>
<td>25</td>
<td>31</td>
<td>-13</td>
</tr>
<tr>
<td>Middle frontal gyrus</td>
<td>8</td>
<td>Right</td>
<td>-2.88</td>
<td>18</td>
<td>31</td>
<td>50</td>
</tr>
<tr>
<td>Superior frontal gyrus</td>
<td>11</td>
<td>Right</td>
<td>-2.71</td>
<td>25</td>
<td>45</td>
<td>-20</td>
</tr>
<tr>
<td>Inferior frontal gyrus</td>
<td>11</td>
<td>Right</td>
<td>-2.59</td>
<td>25</td>
<td>38</td>
<td>-20</td>
</tr>
</tbody>
</table>

The coordinates, Brodmann areas, hemisphere, and \( z \) values of voxels showing significant differences between depressed and control subjects are listed (all \( p's < .01 \)). At each voxel in the brain volume, Fisher’s test for independent correlations were run to assess whether the Spearman’s rank correlations between theta current density within the ACC (averaged within cluster #1, Figure 2) and that voxel differed between depressed and control subjects. Coordinates in millimeters (MNI space), origin at anterior commissure; \((X) = \text{left} (-) \text{to right} (+); (Y) = \text{posterior} (-) \text{to anterior} (+); (Z) = \text{inferior} (-) \text{to superior} (+).
generation of theta rhythm can be found in prior invasive animal and human research as well as EEG/MEG studies using various source localization techniques. First, in rodents, theta rhythm has been directly recorded from anterior and posterior regions of the cingulate gyrus (Borst, Leung, & MacFabe, 1987; Destrade & Ott, 1982; Feenstra & Holshheimer, 1979; Leung & Borst, 1987). Whereas some studies have suggested that this activity may be caused by volume conduction from other regions (Leung & Borst, 1987), others have reported polarity inversion (which implies theta generation) in the cingulate cortex (Feenstra & Holshheimer, 1979; Holshheimer, 1982). Second, in patients with intracranial recording for epilepsy assessment, electrical stimulation of the ACC (midportion of BA 24) has been found to induce a 3–8 Hz rhythm in frontomedial recordings as well as autonomic changes (Talairach et al., 1973). Similarly, scalp and intracranial ictal recordings in patients with cingulate cortex seizures showed rhythmic frontal theta activity (Devinsky, Morrell, & Vogt, 1995). Third, experimental manipulations involving focused attention, working memory, and regulation of the autonomic nervous system, for example, elicited both ACC activation and frontal midline theta activity (Asada et al., 1999; Davis et al., 1997; Dietl et al., 1999; Gevins et al., 1997; Jansma et al., 2000; Tesche & Karhu, 2000; Williams et al., 2000). Finally, single- or distributed sources modeled in medial PFC or dorsal ACC regions have been shown to account for theta activity recorded maximally over frontal midline scalp regions (Asada et al., 1999; Gevins et al., 1997; Ishii et al., 1999). The present results achieved with voxelwise comparisons of concurrently recorded EEG and PET data agree with and extend these findings by showing that tonic ACC activation during task-free resting conditions was associated with current density of theta activity in the same region.

Results from the current data set cannot establish whether theta activity in the ACC relies on intrinsic local properties, patterns of afferent inputs, or passive propagation from brain regions not included in the LORETA solution space. In rodents, theta has been recorded from the hippocampus and entorhinal cortex, but its generation is dependent on the medial septum. Occurrence of hippocampal theta is crucially dependent on afferents from the medial septum/vertical limb of the diagonal band of Broca complex (MS/vDBB), which is considered the pacemaker of hippocampal theta (Vertes & Kocsis, 1997; Vinogradova, 1995). Whereas lesions or pharmacological blockade of the MS/vDBB led to abolition of hippocampal activity, the same was not true for cingulate theta activity (Borst et al., 1987; Destrade & Ott, 1982). Thus, theta can be generated in the cingulate cortex independently of the hippocampal system. Alternatively, because the horizontal limb of the diagonal band projects over the genu of the corpus callosum to the anterior limbic area of the cingulate cortex (BA 24; Swanson & Cowan, 1979), theta in the ACC may be driven by reciprocal anatomical connections with other subcortical areas involved in its generation.3 Despite these anatomical details, the intracellular mechanisms governing theta rhythm in the ACC are unknown. At least in the hippocampus, it seems that inhibitory postsynaptic potentials are not involved in the generation of theta rhythm but that two excitatory synaptic influences contribute to the intracellular theta rhythm: an acetylcholine-mediated sustained depolarization of septal origin and glutamate-mediated EPSP (Steriade, Gloor, Llinàs, Lopes da Silva, & Mesulam, 1990). If these mechanisms extend to the ACC, this would explain the positive relation between electrical and metabolic activity.

Frontocingulate Network and Depression

In the present study, strong positive correlations were found in control subjects between theta current density in the rostral ACC and various regions within the prefrontal and orbitofrontal cortices, particularly in the right hemisphere. Conversely, depressed subjects showed no statistical evidence for such correlations, and, compared to control subjects, were characterized by significantly lower correlations between ACC, on one hand, and right PFC and orbitofrontal regions,4 on the other hand. In control subjects, these indices of functional connectivity between the ACC and frontal regions are consistent with anatomical evidence suggesting that the dorsolateral prefrontal and orbitofrontal cortices send to and receive dense projections from the ACC (Barbas, 1992; Petrides & Pandya, 1999), particularly area 24 and rostral area 32. In agreement with these anatomical data, a recent meta-review of 107 PET studies revealed that changes in blood flow in dorsolateral and orbitofrontal cortices were more likely to occur in tasks evoking ACC activation than in those without ACC activation (Koski & Paus, 2000). Moreover, in a recent study, modulations of dorsolateral PFC activation through repetitive transcranial magnetic stimulation (rTMS) to the left mid-dorsolateral PFC were associated with similar blood flow changes in the ACC (Paus, Castro-Alamancos, & Petrides, 2001).

Evidence of disrupted functional connectivity between the ACC and PFC regions in major depression is intriguing. According to an influential theory of ACC function (Botvinick, Braver, Barch, Carter, & Cohen, 2001), the ACC plays a key role in monitoring conflicts among brain regions. In this theory, the ACC serves an evaluative function reflecting the degree of response conflict elicited by a given task. When conflict is detected, the ACC issues a call for further processing to the dorsolateral PFC, which is assumed to be critical for controlled processing and for guiding behavior toward a goal by adjudicating among various response options. Thus, the dorsolateral PFC represents and maintains task demands necessary for such control and modulates neural activity in brain regions implicated in the conflict. Interestingly, the region of the ACC involved in the present study is slightly more rostral compared to the one included in conflict monitoring, and has been implicated in error detection and emotional evaluation of errors, two further situations requiring executive control of actions (Kiehl, Liddle, & Hopfinger, 2000; Luu, Tucker, Derryberry, Reed, & Poulsen, 2003; Milham et al., 2001). In recent reviews, abnormalities in
cross talk between the ACC and PFC have been emphasized in the pathophysiology of depression (Davidson et al., 2002; Mayberg, 1997). By highlighting diminished functional connectivity during task-free resting conditions between the ACC and various prefrontal regions, including the dorsolateral prefrontal and orbitofrontal cortex, the present findings provide initial empirical evidence for this assumption. Whether in depression such abnormalities in functional connectivity reflect ACC-based impairments in conflict monitoring, particularly in the affective evaluation of conflicts and errors, or PFC-based impairments in controlled processing, or both, is currently unclear.

**Limitations and Future Directions**

Although both cerebral glucose metabolism and neuroelectric activity reflect synaptic activity, several factors contribute to an ill-posed comparison between the two modalities that limits the interpretation of any EEG–PET analysis. First, EEG signals recorded at the scalp are highly dependent on synchronization/desynchronization mechanisms, and their relation to glucose utilization is unclear (Nunez & Silberstein, 2000). For instance, increased neuronal firing associated with reduced synchrony would lead to a small signal recorded at the scalp, yet it would require substantial glucose supply (e.g., alpha blockade after eye opening). Further, it is well known that both excitation and inhibition require glucose utilization (Ackermann, Finch, Babb, & Engel, 1984). For example, loss of inhibitory synaptic action in specific conditions (e.g., interictal periods in epileptic patients) would lead to large EEG spikes but low metabolic utilization. Second, whereas FDG–PET measures glucose metabolism irrespectively from the spatial arrangement of the activated neurons, scalp EEG cannot detect activity arising from assemblies arranged in closed electrical fields (Nunez & Silberstein, 2000). This may explain why no reliable correlations were observed in the present study within the hippocampus—the region most commonly investigated in relation to the theta rhythm in nonhuman studies—which, due to its structural features, represents a closed field. Also, spatial configurations involving opposing dipoles located in sulci generate activity that is opaque to scalp EEG recordings, as the dipoles would cancel. Third, coupling of synaptic activity and energy metabolism is assumed to primarily occur in astrocytes (Magistretti, 2000), glial cells with a “closed field” structure that would be, again, electrically invisible. Finally, perhaps the most important limitations of the present study are its correlational nature and the fact that the reported relation between theta activity and regional glucose metabolism pertained to resting-state data only. Although we expect that such results would extend to task-elicited theta, with different subdivisions of the ACC involved (“cognitive” vs. “affective”) depending on the nature of the task (Paus, 2001), future studies using experimental manipulations known to activate the ACC are needed for testing whether such task-induced changes produce systematic variation in both the theta signal and simultaneously measured hemodynamic or metabolic changes in this region. Importantly, due to the correlational nature of the study, the present findings—though not inconsistent with it—cannot provide direct evidence that theta activity is generated in or modulated by the ACC. Despite these limitations, the present study showcases a novel approach to compare metabolic and electrical activity on a voxelwise basis, and suggests functional coupling between theta activity and glucose metabolism in the rostral ACC as well as functional connectivity within frontocingulate networks that appears to be disrupted in major depression.

**REFERENCES**


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