Left Frontal Hypoactivation in Depression

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Baseline resting electroencephalogram activity was recorded with 3 different reference montages from 15 clinically depressed and 13 control subjects. Power in all frequency bands was extracted by fast Fourier transformation. There was a significant Group × Hemisphere interaction in the mid-frontal region, for the alpha band power only. Depressed subjects had less left-sided activation (i.e., more alpha activity) than did normal control subjects. This pattern of diminished left-sided frontal activation is interpreted as indicating a deficit in approach mechanisms in depressed subjects.

Many reports that indicate that depression is associated with a disruption of the normal pattern of cerebral laterality have recently appeared. This evidence comes from a number of different areas (for reviews, see Davidson, 1984, 1987; Silberman & Weingartner, 1986; Tucker, 1981). Investigations of mood disturbances in epileptics have found that left-sided epileptogenic lesions were associated with significantly higher levels of depression and anxiety than right-sided lesions (e.g., Perini & Mendius, 1984). Gainotti (1972) investigated stroke-induced lesions and reported that left-hemisphere lesions were associated with tears and anxiety whereas subjects with right-hemisphere lesions exhibited indifference and joking. Sackeim et al.'s (1982) review of cases of pathological laughing and pathological crying showed that left-hemisphere lesions were more prevalent in cases of pathological crying whereas pathological laughing was more often associated with right-hemisphere lesions. Other researchers have administered the Minnesota Multiphasic Personality Inventory to subjects with unilateral brain lesions and found that subjects with left-hemisphere lesions had significant elevations on the depression subscale whereas subjects who had lesions in the right hemisphere did not (e.g., Black, 1975; Gasparini, Satz, Heilman, & Coolidge, 1978).

Robinson and his colleagues (Robinson, Kubos, Starr, Rao, & Price, 1984; Robinson & Price, 1982) have used computerized tomography to clarify the relation between lesion location and poststroke mood changes. They found that the severity of poststroke depression was positively correlated with the lesion's proximity to the left frontal pole and negatively correlated with proximity to the right frontal pole (Robinson et al., 1984). Sinory et al. (1986) also found this positive relation between the severity of poststroke depression and proximity to the left hemisphere, but they found a curvilinear relation in the right hemisphere, such that both anterior and posterior lesions were associated with increased depression. If one assumes that stroke-produced lesions lead to decreased activation in the brain regions in which they are found (Burke et al., 1982; Takeuchi et al., 1986), these findings suggest that depression is associated with a decrease in left frontal, right posterior, and possibly right frontal activation. The issue of whether these different patterns tend to co-occur or whether they represent different subtypes of depression is unresolved.

On the basis of previous studies of both normal and depressed subjects, we propose that the left and right anterior zones of the cortex are differentially activated during approach- and withdrawal-related behavior, respectively (see Davidson & Tomarken, 1989, for a review). This leads to an interpretation of decreased left frontal activation as the proximal cause of deficits in approach-related behavior. At least some of the symptoms of depression are similar to the cognitive impairments of patients with left anterior lesions (Luria, 1973). For example, loss of initiative, impaired concentration, indecision, and shortened future perspective are all symptoms common to patients with left anterior lesions and certain subtypes of depression. In a detailed study of the specific symptomatology of endogenous depression, de Jonghe, Ameling, and Assies (1988) reported that these symptoms were present to a moderate or strong degree in at least 83% of the 46 patients studied.

It is important to note that not all studies of patients with unilateral brain lesions show increased depressive symptomatology with left anterior lesions (see Gainotti, 1989). The fact that some patients who show clear evidence of a left anterior lesion in the absence of depressive symptomatology have been identified indicates that decreased activation in this region is clearly not sufficient for the production of depressive symptomatology. We propose that left frontal hypoactivation, either naturally occurring or lesion-induced, represents a diathesis that increases a person's vulnerability to depression. Only when the requisite environmental stress occurs, however, is the vulnerability expressed. This view, therefore, recognizes the existence of persons with a depressogenic pattern of frontal activation who do not show any of the symptoms of depression. However, at least a
subset of persons who are already depressed ought to possess the diathesis and therefore may show decreased left frontal activation in comparison with controls.

Many studies to assess regional brain activation differences between depressed persons and normal control subjects have been performed. These studies have used a number of different methods to assess regional brain activation, including cerebral blood flow, glucose metabolism, and quantitative electroencephalography (EEG; see Henriques & Davidson, 1989, for review). Studies investigating cerebral asymmetries through either regional cerebral blood flow or cerebral glucose metabolism have produced inconsistent results. Guenther et al. (1986) examined changes in regional cerebral blood flow during a motor task that involved the right hand and found that more severely depressed subjects exhibited a lack of activation in the contralateral motor area, which suggests a left-hemisphere dysfunction. Kuhl, Mitter, and Riege (1983) found that the pattern of glucose metabolism in unipolar subjects at rest differed from control subjects in only one region: Depressed subjects had decreased metabolism in the left posterior-inferior frontal cortex. Baxter and his associates have also observed decreased left anterior activation in depressives in relation to that in control subjects (Baxter et al., 1985; Baxter et al., 1989). This finding has recently been confirmed by Martinot et al. (1990), but other investigators have not found this pattern of diminished left anterior activation (e.g., Gur et al., 1984; Uytdenhoef et al., 1983). It is not clear why these investigators found divergent results. Gur et al. used absolute regional values in computing their comparisons, whereas Baxter et al. (1989) computed the metabolic rate in each region relative to the entire ipsilateral hemisphere. However, Uytdenhoef et al. used relative values and found that their depressed subjects had greater left anterior blood flow than did the control subjects. Unfortunately, there were a number of problems with Uytdenhoef et al.'s study. The handedness of the patients and control subjects was not specified. The groups differed considerably in age, with the control subjects more than 10 years younger than the depressives. Uytdenhoef et al. reported a positive .35 correlation between relative left frontal blood flow and age within the control group. It is therefore unclear whether the group difference in left frontal blood flow is a function of the age difference between groups or is a genuine group difference. Without statistically partialing out the effects of age, the results from this study are inconclusive.

Investigations that have used quantitative EEG to examine asymmetries in activation also suggest left hemisphere involvement in depression. Some of the earliest work was done by d'Elia and Perris (1973, 1974), who examined the mean integrated amplitude and the within-patient variability of the integrated amplitude in depressed subjects. They found that the within-patient variability in the dominant left hemisphere was significantly lower in depression, and this variability in the left hemisphere increased at recovery. They interpreted this finding to suggest greater left hemisphere involvement in depression (d'Elia & Perris, 1973, 1974; Perris, 1975). Greater relative right hemisphere variability has also been found in neurotic depressives (Rochford, Swartzburg, Chowdhry, & Goldstein, 1976). Unfortunately, the functional significance of variability in brain electrical activity is not known, so that interpretation of these findings is problematic. Matousek, Capone, and Okawa (1981) found that endogenously depressed subjects had more relative left-sided alpha activity in the frontal region than did control subjects, though this difference was not significant. Work in our laboratory has found that depressed subjects differed from nondepressed subjects in measures of alpha power asymmetry in the anterior and posterior scalp regions (Davidson, Chapman, & Chapman, 1987; Davidson, Schaffer, & Saron, 1985; Schaffer, Davidson, & Saron, 1983). The most consistent finding we have obtained across studies is that depressed subjects differed from controls in the asymmetry of frontal activation. Compared with control subjects, depressives show more left frontal alpha power. In some studies, depressives have shown relatively more right-sided parietal alpha power in comparison with control subjects (e.g., Davidson et al., 1987). Note that decreases in alpha-band power reflect increases in cortical activation (see Davidson, 1988; Lindsley & Wicke, 1974). Thus, our electrophysiological data indicate that the most consistent difference between depressed and nondepressed subjects is that the former group shows less left-sided frontal activation in comparison with the latter group. A recent study found that this pattern of decreased left anterior activation also distinguished remitted depressives from never-depressed control subjects (Henriques & Davidson, 1990). Remitted depressives also showed decreased right-sided posterior activation in comparison with controls. In that study, in addition to assessing power in the alpha band, we examined power in the other EEG frequency bands and found that the group differences were specific to the alpha band.

In previous studies with acutely depressed subjects (Davidson et al., 1987; Davidson et al., 1985; Schaffer et al., 1983), the subjects consisted of subclinically depressed college students, selected on the basis of extreme scores on the Beck Depression Inventory (BDI; Beck, Ward, Mendelson, & Erbaugh, 1961). In addition to the usual criterion of high BDI scores that are maintained over at least a 1-month interval, subjects were required to report vegetative symptomatology on the final six items of the BDI (see Schaffer et al., 1983, for details). In this study we tested a group of subjects who met Research Diagnostic Criteria (Spitzer, Endicott, & Robins, 1978) for major unipolar depression, all but one of whom also met criteria for definite or probable endogenous depression. We compared these subjects with a group of control subjects who had no history of depression or any other psychopathology in themselves or their first-degree relatives. We focused primarily on endogenous depressives because we hypothesized that this subtype would be most likely to show deficits in approach-related behavior (e.g., show pervasive loss of pleasure and interest in objects and people) and therefore would show left frontal hypoactivity. We recorded EEG from the left and right hemisphere in several anterior and posterior scalp regions. Although there has been discussion in the electrophysiology literature about the appropriateness of different referencing strategies (Lehman, 1987; Nunez, 1981), there is currently no consensus in the literature about which approach is optimal. We adopted the strategy of recording the EEG with more than one reference montage and demonstrating consistency across different referencing procedures. Such cross-referencing consistency would support our suggestion of the localization of the significant effects to the frontal region because it would indicate that the group differences were not a function
of the particular reference used. Therefore, EEG data were recorded so that three different reference montages, (a) vertex, (b) computer-averaged ear lobes, and (c) average reference were available for analysis, (see Henriques & Davidson, 1990, for a more complete discussion). In addition to computing measures of power in the traditional EEG bands, we computed power in a high frequency (70–80 Hz) band (which presumably is purely myogenic in origin) to obtain estimates of muscle contamination. Power in this band was then used as a covariate in our analyses of EEG band power.

We predicted that the pattern of EEG asymmetry across the scalp would distinguish between depressed and control subjects and that this would be consistent across reference montage. On the basis of earlier work (Davidson et al., 1987; Henriques & Davidson, 1990; Schaffer et al., 1983), we predicted that depressed subjects, in comparison with control subjects, would have less left frontal activation. Although there is less information about parietal activation, we also predicted that depressed subjects would show less right parietal activation in comparison with control subjects.

Method

Subjects

Depressed subjects were recruited in connection with ongoing drug studies at the Center for Affective Disorders at the University of Wisconsin Hospital. Control subjects were recruited through advertisements in local newspapers. All subjects were screened with the Schedule for Affective Disorders and Schizophrenia (Endicott & Spitzer, 1978). Interviews were conducted by one of two laboratory members, both of whom had completed 40 hours of training on the Schedule for Affective Disorders and Schizophrenia. Depressed subjects were required to meet Research Diagnostic Criteria (Spitzer et al., 1978) diagnosis of unipolar major depressive disorder. In addition, subjects were required to have an absence of mania in their first-degree relatives. Of 54 possible depressed subjects interviewed, 16 (7 men and 9 women) met all criteria and were invited to participate in the study. Fifteen of the depressed subjects met criteria for endogenous depression (11 definite and 4 probable). Eight of the depressives met criteria for recurrent depression, and 3 were diagnosed as having chronic depression. The length of the current episode, at the time of testing, ranged from 10 to 364 weeks in duration. Five of the depressed subjects had begun treatment with either imipramine or fluvoxamine at the time that EEG was measured.

Because of insufficient artifact-free data, the EEG measures from 1 depressed and 2 control subjects were dropped. This resulted in a final group of 15 depressed and 13 control subjects. The two groups did not differ in age (depressed subjects, M = 40.40, range 33–57, and control subjects, M = 40.61, range 31–56), t(26) = −0.08, p > .05, or sex (p > .05, two-tailed, Fisher's exact test). The groups did differ in the amount of reported depression as assessed by the BDI, t(17) = 10.57, p < .0001, and the Hamilton Depression Rating Scale (Hamilton, 1960), t(15) = 13.68, p < .0001. The two groups also differed in socioeconomic status as assessed by Hollingshead's (1957) index, n(26) = 4.18, p < .001. Depressed subjects came from lower social classes than did the control subjects. Relevant subject variables are listed in Table 1.

Table 1

<table>
<thead>
<tr>
<th>Measure</th>
<th>Depressed</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (in years)</td>
<td>40.40</td>
<td>40.61</td>
</tr>
<tr>
<td>Socioeconomic status</td>
<td>3.93</td>
<td>2.61</td>
</tr>
<tr>
<td>Hamilton Depression</td>
<td>23.33</td>
<td>9.92</td>
</tr>
<tr>
<td>Rating Scale</td>
<td>6.23</td>
<td>1.12</td>
</tr>
<tr>
<td>Beck Depression Inventory</td>
<td>26.80</td>
<td>1.83</td>
</tr>
</tbody>
</table>

Note: Depressed women, n = 8; control women, n = 9; depressed men, n = 7; and control men, n = 4. Socioeconomic status is rated 1–7, wherein lower numbers reflect higher social class.

Procedure

Before the EEG recording, the subject was informed about the nature of the experiment and was asked to sign a consent form. The subject was then administered the Hamilton Depression Rating Scale and completed the BDI. On completion of the depression inventories, the subject was escorted to the experimental testing room where all further procedures took place.

The test session consisted of two 30-s baseline resting periods, and these baselines were followed by a series of emotion-eliciting film clips. This report will present only the data from the baseline periods. Baseline EEG was recorded during both an eyes-open and an eyes-closed rest period, the order of which was counterbalanced across subjects. The subjects were asked at the end of each baseline trial to rate their emotional state during the trial. This was done by rating emotional experience on seven emotion scales: interest, admiration, happiness, fear, sadness, disgust, and anger. Subjects used a 0–8 scale, wherein 0 indicated that the emotion was not experienced during the trial and 8 indicated that it was felt very strongly during the trial.

All subject instructions were presented on a video monitor controlled by computer. The subjects used a numeric keypad to advance through the instructions and to input their emotion ratings at the end of each trial. The subjects were instructed to use either their right or their left hand to enter their responses, and response hand was randomized across subjects.

Electroencephalography Recording

EEG was measured with a modified lyca electrode cap (Electro-cap International, Dallas, TX). The electrode cap was positioned on the subject's head at known anatomical landmarks. Elastic straps from the cap attached to a strap that traversed the subject's torso, and this enabled the subject to move comfortably without altering the placement. This procedure results in accurate electrode placements (Blom & Anneveldt, 1982). EEG was recorded from 14 scalp locations: F3, F4, F7, F8, T3, T4, T5, T6, P3, P4, C3, C4, Pz, and Fz (10–20 system). All

1 Six of the control subjects (2 men, 4 women) in this study were previously reported on in Henriques and Davidson (1990).

2 The reduced degrees of freedom for the tests of group differences on the Beck Depression Inventory and the Hamilton Depression Rating Scale are a function of the correction for unequal variances.
placements were referenced to Cz. Two additional channels, Cz-A1 and Cz-A2, were recorded in order to derive an averaged ears reference (see Davidson, 1988). Electrode impedances were all under 5,000  , and the impedances for homologous sites were within 500  of each other. Electrooculogram (EOG) was recorded from the external canthus to the supraorbit of one eye, in order to facilitate artifact scoring. EOG was only recorded on paper for the purpose of artifact scoring. We were thus unable to digitize EOG activity. EEG and EOG were amplified with a 20-channel Grass (Quincy, MA) Model 12 Neurodata System that had a bandpass of 1-100 Hz and a 60-Hz notch filter. All analog signals were passed through active low-pass filters (Rockland Systems, West Nyack, NY, Model 424) with a cutoff of 85 Hz and a 24 dB per octave roll-off (see Dumermuth & Molinari, 1987). The EEG was digitized at the rate of 250 samples/s. The EEG activity for each channels and the EOG activity were displayed on a Grass Model 7 nine-channel polygraph. This paper record was then used to identify those portions of data to be edited out because of eye blinks, gross muscle artifact, and movement artifact. A fast Fourier transform was applied to all chunks of artifact-free data that were 2.05 s in duration, with chunks overlapping by 75%. The two groups did not differ in the number of artifact-free chunks, t(24) = 0.45, p > .05. The mean number of chunks for the depressed group was 74.42 (SD = 27.69), and the mean for the control group was 69.36 (SD = 29.17). The fast Fourier transform output was then converted to power density (  ) in each of five bands: delta, 1-4 Hz; theta, 4-8 Hz; alpha, 8-13 Hz; beta 1, 13-20 Hz; and electromyogram (EMG), 70-80 Hz. This conversion was done by summing activity across all bins within a band and dividing by the number of 1-Hz bins. Power in the 70- to 80-Hz band was examined in an attempt to quantitatively evaluate the presence and amount of muscle artifact. Activity in this frequency range is presumed to be exclusively myogenic in origin and thus can be used to estimate the contribution of muscle artifact in each lead independent of EEG activity. In addition to the original recording montage (referencing to vertex), the EEG was recomputed off-line for two additional references, computer-averaged ears and an average reference. For the ears reference, the separate Cz-A1 and Cz-A2 channels were averaged and then added to the original vertex-referenced data. For the average reference the voltage at each electrode was expressed as a difference from the average voltage of all electrodes on the scalp. All power density values were log-transformed to normalize their distribution.

Results

We present the emotion self-report data first and then the EEG data. All analyses were computed as repeated measures multivariate analyses of variance (MANOVAs). Because of equipment malfunctions, 3 depressed subjects were missing data from one channel (T4); these subjects were not included in any MANOVA that examined activity at all sites, but they were included in all of the regional MANOVAs that were computed. The inclusion of these 3 subjects did not change the significance of any of the computed regional analyses.

Baseline Emotion Data

The subjects' self-report of experienced emotion was examined by computing separate two-way MANOVAs, with group and emotion as variables. There was a significant effect for group, $F(1, 26) = 5.13$, $p < .05$. This was because the depressed subjects reported more emotion than the controls during the baseline trials (Table 2). There was also a main effect for emotion, $F(6, 21) = 5.46$, $p < .01$. This was because subjects reported more interest, amusement, and happiness than sadness, fear, disgust, or anger. There was no significant interaction between group and emotion, $F(6, 21) = 0.91$.

Baseline Electroencephalography Data

Prior research in our laboratory with a large sample size has demonstrated that subjects do not differ in anterior asymmetry during eyes-open as compared with eyes-closed baseline periods. Moreover, a weighted average of eyes-open and eyes-closed data produces more stable estimates of EEG asymmetry (Tomarken, Davidson, Wheeler, & Kinney, in press) than either baseline type alone. In previous research to compare clinical samples with normal subjects, we failed to find Group X Baseline Type (i.e., eyes open or closed) interactions (Henriques & Davidson, 1990). In order to justify our use of a composite variable of the mean of the eyes-open and eyes-closed trials in this study, four-way MANOVAs were computed with group (depressed vs. control) as the between-groups variable and with hemisphere (left vs. right), region (midfrontal [F3 and F4] vs. lateral frontal [F7 and F8] vs. anterior temporal [T3 and T4] vs. posterior temporal [T5 and T6] vs. central [C3 and C4] vs. parietal [P3 and P4]), and condition (eyes open vs. eyes closed) as within-groups variables. These analyses revealed that there were no interactions with baseline condition. The data from the eyes-open and eyes-closed baselines were then averaged together by using the number of artifact-free chunks within each baseline as a weighting factor, and this composite variable was then used in all subsequent analyses.

Because earlier research in our laboratory has examined group differences in alpha power (e.g., Davidson et al., 1987; Davidson et al., 1985), we had specific hypotheses about activity in this frequency band. We predicted that depressed and control subjects would differ in asymmetry in both the midfrontal (F3 and F4) and parietal regions (P3 and P4). These analyses are presented first.

### Table 2

<table>
<thead>
<tr>
<th>Emotion</th>
<th>Depressed M</th>
<th>SD</th>
<th>Control M</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interest</td>
<td>2.26</td>
<td>1.72</td>
<td>1.54</td>
<td>1.75</td>
</tr>
<tr>
<td>Amusement</td>
<td>1.13</td>
<td>1.14</td>
<td>0.69</td>
<td>1.09</td>
</tr>
<tr>
<td>Happiness</td>
<td>1.60</td>
<td>1.80</td>
<td>1.50</td>
<td>1.53</td>
</tr>
<tr>
<td>Sadness</td>
<td>1.03</td>
<td>0.93</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Fear</td>
<td>0.83</td>
<td>1.13</td>
<td>0.15</td>
<td>0.43</td>
</tr>
<tr>
<td>Disgust</td>
<td>0.73</td>
<td>1.24</td>
<td>0.08</td>
<td>0.28</td>
</tr>
<tr>
<td>Anger</td>
<td>0.47</td>
<td>0.74</td>
<td>0.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>

Note. Each emotion was rated on a 0-8 scale.

3 We recorded the Cz channel referenced separately to each ear, rather than reference each ear to Cz, and thus the average of the Cz-A1 and Cz-A2 channels was added to the original data to derive the computer averaged ears reference. Thus if $a = (F3 - Cz)$ and $b = (Cz - A1)$, then $a + b = (F3 - A1)$.
**Midfrontal Region**

**Cz montage.** The analysis of alpha power in the midfrontal region referenced to Cz revealed a significant Group × Hemisphere interaction, $F(1, 26) = 6.71, p < .02$. Depressed subjects had a pattern of relatively right-sided activation (i.e., less right than left alpha power), whereas control subjects displayed relatively left-sided activation (more right than left alpha power). This group difference was because the depressed subjects had more left-sided alpha power (i.e., less activation) than did control subjects (Figure 1). Examination of individual subjects’ asymmetry scores, computed as Log Right Alpha Power − Log Left Alpha Power, revealed that only 1 depressed subject had an asymmetry score that was above the mean value for the control subjects and only 2 control subjects had asymmetry scores that were below the mean of the depressed group ($p < .0001$, two-tailed, Fisher’s exact test; see Figure 2).

**Average reference montage.** Analysis of alpha power referenced to an average of all the electrodes also revealed a significant Group × Hemisphere interaction in the midfrontal region, $F(1, 26) = 5.42, p < .03$. Depressed subjects had relatively right-sided activation in this region, and control subjects had relatively left-sided activation. Depressed subjects had more left-sided alpha power than the control subjects, but the two groups had comparable levels of activation in the right hemisphere (Figure 3). Once again we examined the individual subjects’ asymmetry scores and found that only 2 depressed subjects had asymmetry values that were above the mean of the control group and none of the control subjects had an asymmetry score below the mean for the depressives ($p < .0001$, two-tailed, Fisher’s exact test; see Figure 4).

**Averaged ears montage.** The patterning of alpha power in the midfrontal region, referenced to averaged ears, was similar to the data referenced to the two other montages. However, the Group × Hemisphere interaction was not significant, $F(1, 26) = 0.63$.

**Parietal Region**

Examination of alpha power in the parietal region did not reveal any group difference in asymmetry for any of the three reference montages. Both depressed and control subjects had relatively right-sided activation in this region.

**Other Analyses**

Because we had no specific hypotheses about activity in other bands or regions, we computed three-way MANOVAs for each frequency band with group as the between-groups variable and hemisphere and region as the within-groups variables.

**Delta.** There were no significant main effects for group or interactions with group for delta power referenced to Cz.

The three-way MANOVA for delta power referenced to the average reference montage revealed a significant Group × Region interaction, $F(5, 19) = 2.79, p < .04$. When this interaction was examined, it was found that control subjects had less delta power than depressives in the midfrontal, anterior temporal, posterior temporal, and central regions, but none of the univariate group differences for individual regions were significant.

The averaged ears montage analysis revealed a significant Group × Region × Hemisphere interaction, $F(5, 19) = 2.92, p < .04$. Separate Group × Hemisphere MANOVAs were computed.
When the analysis of power in the alpha frequency band included all 6 regions, there were no significant main effects for group or interactions with group for any of the three reference montages.

Beta. The analysis of power in the beta frequency band revealed no significant main effects for group or interactions with group for any of the three reference montages.

Electromyography

To investigate the possibility that the effects we observed in the midfrontal region were merely the result of underlying EMG asymmetries, we recomputed our analyses with EMG asymmetry (Log Right Power - Log Left Power) as a covariate in a series of one-way analyses of covariance. These analyses of covariance were computed separately for each reference, with group as the independent variable. An asymmetry metric was used rather than the raw power values so that there was only a single covariate for each analysis, rather than multiple covariates. The two groups did not differ in EMG asymmetry, $F(1, 25) = 0.02$ for Cz reference, and $F(1, 25) = 0.52$ for average reference. Also, the test for parallelism revealed no Group x EMG Asymmetry interaction, $F(1, 24) = 0.20$ and $F(1, 24) = 0.95$ for Cz and average references, respectively, which justified our use of EMG asymmetry in an analysis of covariance. The significant Group x Hemisphere interactions were unchanged, for the Cz reference montage, $F(1, 25) = 6.32$, and for the average reference montage $F(1, 25) = 5.82$.

Socioeconomic Status

Because the two groups differed in socioeconomic status, it is possible that the observed group differences in the midfrontal region are merely a function of this difference in social class. To investigate relations between alpha asymmetry (Log Right Power - Log Left Power) in the midfrontal region and social class, correlations were computed separately for each group. None of these correlations were significant. For the Cz referenced data, the correlation for the depressed subjects was -.29, indicating that lower social class (i.e., higher class numbers) was associated with more left-sided activation. The direction of this correlation goes against our hypothesized group difference. In the control subjects, the correlation was -.16. For the average reference data, the correlations for the depressed and control subjects were .05 and -.15, respectively. These correlations were -.17 and -.21 for depressed and control subjects, respectively, for data referenced to the average ears montage.

Medication Status

At the time of testing, 5 of the depressed subjects were receiving antidepressant medication (either imipramine or fluvoxamine). To investigate whether there were any differences in the midfrontal region between medicated and nonmedicated depressives, separate two-way Medication x Hemisphere MANOVAs were computed for each reference montage on the alpha power data in the midfrontal region. These revealed that there were no significant differences between these two groups of depressed subjects, for the Cz referenced data, $F(1,13) = .02$, for
Figure 3. Mean log-transformed alpha (8-13 Hz) power (in \(\mu V^2/Hz\)) for average-referenced electroencephalograms (across eyes-open and eyes-closed baselines), split by group and hemisphere, for the midfrontal region.

Relations Between Frontal Asymmetry and State Ratings of Emotion and Depression

Because the two groups differed in the amount of emotion reported during the baseline trials, it is conceivable that the observed differences in asymmetry reflect the group difference in self-reported emotion. To investigate this possibility we computed correlations between alpha frontal asymmetry scores (Log Right Power − Log Left Power) and self-reported emotion. None of these correlations were significant. The average correlation for data referenced to Cz was −.05, and the average correlation for data referenced to the average montage was .05. We also examined correlations between midfrontal asymmetry and the severity of depression for each group. None of these correlations were significant. For data referenced to Cz, the correlations between control subjects’ frontal asymmetry and severity of depression as measured by the BDI and Hamilton Depression Rating Scale, respectively, were −.18 and .24. For depressed subjects these correlations were −.11 and −.16. The correlations for data referenced to the average montage were −.19 and −.31 for control subjects and .05 and −.08 for depressed subjects. Negative correlations indicate that lower asymmetry scores (which reflect relatively greater right-sided activation) were associated with higher levels of depression.

Discussion

These data provide further support for our hypothesis that depressed subjects differ from normal control subjects in the patterning of anterior activation. Similar to the results of previous studies with subclinically depressed subjects (Davidson et al., 1987; Schaffer et al., 1983) and remitted depressives (Henriques & Davidson, 1990), depressed subjects in this study displayed a pattern of decreased left-sided frontal activation in contrast to normal control subjects. This pattern of decreased left-sided anterior activation in the depressed subjects was significant in two of the three reference montages.

This difference cannot be attributed either to the lower socioeconomic status of the depressed subjects or to the subgroup of depressed subjects who were receiving medication. Correlations between socioeconomic status and frontal asymmetry were not significant. In fact, the pattern in the depressed subjects was one in which lower social classes were associated with more relatively left-sided activation, a result which is at odds with the hypothesis that the decreased left-sided frontal activation in the depressed subjects was a result of their lower class. Comparisons between depressed subjects who were receiving medication and those who were drug free at the time of testing revealed no significant differences in the patterning of alpha asymmetry. The observed group differences cannot be attrib-

* Although the subjects in these studies were selected from a putatively normal population on the basis of elevated Beck Depression Inventory scores, it is likely that some of the subjects may have met rigorous diagnostic criteria for major depression.
Table 3
Log Transformed Delta Power (in $\mu V^2$/Hz) Referenced to Computer-Averaged Ears in the Left and Right Hemispheres by Group

<table>
<thead>
<tr>
<th>Region</th>
<th>Depressed</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$M$</td>
<td>SD</td>
</tr>
<tr>
<td>Lateral frontal</td>
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<tr>
<td>Left</td>
<td>0.152</td>
<td>0.603</td>
</tr>
<tr>
<td>Right</td>
<td>0.334</td>
<td>0.569</td>
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<tr>
<td>Anterior temporal</td>
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</tr>
<tr>
<td>Left</td>
<td>-0.347</td>
<td>0.415</td>
</tr>
<tr>
<td>Right</td>
<td>-0.204</td>
<td>0.392</td>
</tr>
</tbody>
</table>

Note. Averaged across eyes-closed and eyes-open baselines.

Figure 4. Individual subjects' log transformed alpha asymmetry scores ($\log \text{Right} - \log \text{Left}$ in $\mu V^2$/Hz) for average-referenced electroencephalograms (across eyes-open and eyes-closed baselines), split by group, for the midfrontal region.

We and other researchers (i.e., Davidson, 1984, 1987; Davidson, Ekman, Saron, Senulis, & Friesen, 1990; Kinsbourne 1978; Swartzburg, 1983) have proposed that the anterior region of the left and right hemispheres are important components of the circuitry that mediate approach and withdrawal behavior, respectively. Different asymmetrical patterns of activation in the anterior cortical zones are predicted to bias a person's emotional reactivity and to increase a person's vulnerability to particular types of psychopathology, in response to an appropriate situation or life event (Davidson & Tomarken, 1989). The model is explicitly a diathesis-stress model that holds that anterior asymmetry by itself is not sufficient to produce a different pattern of emotional behavior. Accentuated activation in the left anterior region is hypothesized to increase a person's vulnerability to approach-related emotion and behavior, whereas accentuated activation in the right anterior region is hypothesized to increase a person's vulnerability to withdrawal-related emo-
tion and behavior. Hypoactivation in the left anterior region is predicted to increase a person's vulnerability to behavior and emotion associated with deficits in the approach system. Sadness and depression are both expected to result from such approach-related deficits. This view is supported by studies that suggest that dispositional positive affect is decreased in depressed persons (Tellegen, 1985; Watson, Clark, & Carey, 1988) and in subjects selected from a normal population on the basis of extreme left frontal hypoactivation (Tomarken, Davidson, Wheeler, & Doss, in press).

In this study we found no relation between anterior activation asymmetry and reports of emotion at the time the baseline measures were obtained. The lack of a significant association between these classes of measures is consistent with previous data in normal subjects, for whom it was found that baseline frontal asymmetry predicted reactivity to emotional film clips but was not associated with measures of mood (identical to those used in this study) at the time the baselines were obtained (Tomarken, Davidson, & Henriquez, 1990). The fact that baseline measures of frontal asymmetry were not associated with concurrent measures of emotional state is also consistent with our theory that posits a diathesis–stress model. During the baseline periods, there is no situational provocation, and the level of baseline state–emotion is rather low. Under such circumstances we do not expect frontal asymmetry to account for variance in baseline emotional state.

Measures of frontal asymmetry were also not significantly associated with measures of severity of depressive symptomatology as indexed by either the BDI or the Hamilton Depression Rating Scale. This conclusion is based on the absence of any significant correlations within groups between the EEG measure of frontal asymmetry and scores on the two depression measures. Because these measures reflect more tonic dispositional characteristics in comparison with our measures of emotional state, we might expect them to be associated with frontal asymmetry. We offer two possible reasons to explain this lack of association. First, the lack of correlation between depression severity and frontal asymmetry may have arisen because of the truncated range of scores on the measures of depressive symptomatology within each group. We did not compute correlations across groups because subjects were selected on the basis of depressive symptomatology, and we therefore did not have the continuous range of depression severity scores necessary for correlation. If we had a full range of scores on the depression measures, we would expect to find significant relations between the depression measures and the frontal asymmetry measures. Second, according to our model, not all depressed subjects are expected to have the left frontal hypoactivation diathesis. We allow for the possibility that depression arises in other manners, and our depressed sample may have included subjects who lacked the diathesis (see Figures 2 and 4). In this regard, it will be of great interest in future research to determine if there are phenotypic differences between those depressed subjects with and without left frontal hypoactivation.

In light of the now available corpus of evidence that indicates that left frontal hypoactivation is present in acutely depressed persons, in remitted depressives, and in subjects with low dispositional positive affect drawn from a normal population, we propose that this pattern is a state-independent marker of vulnerability to affective disorders. If this proves to be a replicable finding, it will be of interest to examine its etiology. We know that individual differences in frontal asymmetry are present within the first year of life and predict important aspects of an infant's response to stressful challenges, such as a brief episode of maternal separation (Davidson & Fox, 1989). It is likely that the distal causes of individual differences in frontal asymmetry will be a complex mixture of early environmental and genetic effects. The proximal causes, or mechanisms which underlie the frontal asymmetry differences, are likely to reflect, at least in part, upstream influences from subcortical structures that have direct projections to the frontal lobes (Nauta, 1971). For example, catecholamine asymmetries in certain subcortical structures, such as the amygdala and the thalamus, may contribute to asymmetrical efferent outflow to anterior cortical regions (see Tucker & Williamson, 1984, for review). Alternatively, or possibly in addition, frontal cortical zones of the two hemispheres may be differentially sensitive to upstream influences from subcortical structures. The differential cortical sensitivity of the two hemispheres may also be a function of an asymmetrical distribution of neurotransmitters or asymmetries in receptor densities. The precise mechanisms that underlie the electrophysiological asymmetries we have found must await research with new methods to assess regional brain function, such as positron emission tomography, which can potentially reveal regional differences in neurotransmitter concentration and receptor densities.

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changes]. No To Shinke, 38, 1143-1149. (From Medline, Unique Identifier No. 87128641)


Received May 30, 1990
Revision received December 14, 1990
Accepted December 15, 1990

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