Adverse Rearing Environments and Neural Development in Children: The Development of Frontal Electroencephalogram Asymmetry

Katie A. McLaughlin, Nathan A. Fox, Charles H. Zeanah, and Charles A. Nelson

**Background:** Children raised in institutional settings experience marked deprivation in social and environmental stimulation. This deprivation may disrupt brain development in ways that increase risk for psychopathology. Differential hemispheric activation of the frontal cortex is an established biological substrate of affective style that is associated with internalizing psychopathology. Previous research has never characterized the development of frontal electroencephalogram asymmetry in children or evaluated whether adverse rearing environments alter developmental trajectories.

**Methods:** A sample of 136 children (mean age = 23 months) residing in institutions in Bucharest, Romania, and a sample of community control subjects (n = 72) participated. Half of institutionalized children were randomized to a foster care intervention. Electroencephalogram data were acquired at study entry and at ages 30, 42, and 96 months. A structured diagnostic interview of psychiatric disorders was completed at 54 months.

**Results:** Children exhibited increases in right relative to left hemisphere frontal activation between the second and fourth years of life, followed by an increase in left relative to right hemisphere activation. Children reared in institutions experienced a prolonged period of increased right hemisphere activation and a blunted rebound in left frontal activation. Foster care placement was associated with improved developmental trajectories but only among children placed before 24 months. The development trajectory of frontal electroencephalogram asymmetry in early childhood predicted internalizing symptoms at 54 months.

**Conclusions:** Exposure to adverse rearing environments can alter brain development, culminating in heightened risk for psychopathology. Interventions delivered early in life have the greatest potential to mitigate the long-term effects of these environments.

**Key Words:** Brain development, childhood adversity, deprivation, electroencephalogram (EEG), frontal EEG asymmetry, institutionalization
predict internalizing psychopathology in early childhood. We examine these questions using data from the Bucharest Early Intervention Project (BEIP), a longitudinal study of children reared in institutional settings in Romania and a comparison sample of community-reared children. Institutional rearing is characterized by psychosocial deprivation and neglect (30) and has lasting effects on social development and mental health (30–36). Prior work in this sample suggests that institutionalization is associated with abnormal neural development, as indexed by high low-frequency power and low high-frequency power in the EEG signal of children exposed to institutionalization (37). If the environment does, in fact, alter the development of hemispheric activation patterns in the frontal cortex, we expect to observe these effects in this population.

Methods and Materials

Sample

The BEIP is a longitudinal study of a sample of children raised from early infancy in institutions in Bucharest, Romania. The BEIP is the only randomized controlled trial of foster care (FC) among institutionalized children (38). A sample of 136 children (aged 6–30 months) was recruited from institutions in Bucharest. An age- and gender-matched sample of 72 community-reared children was recruited from pediatric clinics in Bucharest. Assessments of health, cognitive ability, and brain development were completed at baseline and at 30, 42, and 96 months. Psychiatric disorders were assessed at 54 months.

Participants were selected from each of the six institutions for young children in Bucharest. Physical examinations were completed on 187 children residing in these institutions. Of this group, 51 were excluded for medical reasons (e.g., Down syndrome) (38). The remaining 136 children had lived in an institution for at least half of their lives (M = 89.0%). Following baseline assessments, half of the children (n = 68) residing in institutions were randomized to a FC intervention and half (n = 68) remained in institutional care. No differences were found between the intervention and control group in gender distribution, age, birth weight, or percentage of life spent in the institution (Table 1). The mean age at FC placement was 22.97 months. By the 96-month assessment, 19 children were lost to follow-up, primarily because of adoption or reintegration with their biological parents. The study design and methods are described elsewhere (38).

The BEIP was initiated with support from the Secretary of State for Child Protection in Romania. Study procedures were approved by local commissions on child protection in Bucharest, the Romanian ministry of health, an ethics committee including appointees from government and Bucharest University academic departments, and the institutional review boards of the institutions of the three principal investigators. A description of procedures employed to ensure ethical integrity has been published previously (39–41).

Intervention

Because FC was virtually nonexistent in Bucharest at the beginning of the study, investigators created a network of foster homes with Romanian collaborators (38,42). Foster parents were recruited through advertising and were trained using a manual adapted for the study. Foster care was supported by social workers in Bucharest who received weekly consultation from US clinicians. Social workers assisted foster parents in managing problem behaviors and facilitated the establishment of warm, supportive, and committed relationships with their foster children. The intervention is described in detail elsewhere (38,42).

<table>
<thead>
<tr>
<th>Table 1. Baseline Characteristics of Children in the Bucharest Early Intervention Project</th>
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<tr>
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<tr>
<td>Community Control</td>
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<tr>
<td>(n = 72)</td>
</tr>
<tr>
<td>Foster Care</td>
</tr>
<tr>
<td>(n = 68)</td>
</tr>
<tr>
<td>Usual Care</td>
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<td>(n = 68)</td>
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<table>
<thead>
<tr>
<th>Variable</th>
<th>Community Control</th>
<th>Foster Care</th>
<th>Usual Care</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female, No. (%)</td>
<td>41 (56.9)</td>
<td>33 (49.3)</td>
<td>35 (51.5)</td>
</tr>
<tr>
<td>Age, Mean (SD), Months</td>
<td></td>
<td></td>
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<tr>
<td>Age at institutionalization</td>
<td>–</td>
<td>2.6</td>
<td>2.4</td>
</tr>
<tr>
<td>Age at study entry</td>
<td>19.3 (7.1)</td>
<td>20.9 (7.1)</td>
<td>20.8 (7.7)</td>
</tr>
<tr>
<td>Birth Weight, Mean (SD), Grams</td>
<td>3333 (459.3)</td>
<td>2733 (576.2)</td>
<td>2847 (570.2)</td>
</tr>
<tr>
<td>Gestation Age, Mean (SD), Weeks</td>
<td>37.8 (1.4)</td>
<td>37.0 (2.4)</td>
<td>37.6 (1.5)</td>
</tr>
<tr>
<td>Head Circumference for Age Percentile, Mean (SD), cm</td>
<td>52.4 (25.6)</td>
<td>32.3 (28.1)</td>
<td>25.2 (22.7)</td>
</tr>
<tr>
<td>Height for Age Percentile, Mean (SD), cm</td>
<td>53.6 (27.7)</td>
<td>25.7 (22.5)</td>
<td>26.9 (23.1)</td>
</tr>
<tr>
<td>Weight for Age Percentile, Mean (SD), kg</td>
<td>47.2 (29.6)</td>
<td>18.2 (19.4)</td>
<td>22.7 (24.6)</td>
</tr>
<tr>
<td>Weight for Height Percentile, Mean (SD), kg</td>
<td>54.0 (28.0)</td>
<td>30.1 (27.2)</td>
<td>37.3 (29.6)</td>
</tr>
<tr>
<td>Duration of Institutionalization, Mean (SD), Weeks</td>
<td>–</td>
<td>85.2 (23.0)</td>
<td>87.9 (17.9)</td>
</tr>
</tbody>
</table>

SD, standard deviation.
child’s attention. The EEG signal was recorded for the entire 3-minute period, but only data from epochs in which the wheel was being spun were analyzed. At 96 months, EEG was recorded while the children sat quietly in a chair, alternating 1-minute epochs of eyes open and closed for 6 minutes. Data presented here are from epochs with eyes closed (results were unchanged with eyes open).

Processing and analysis of the EEG signal were performed using the EEG Analysis System from James Long Company (Caroga Lake, New York). Epochs containing blinks or eye movement or in which the EEG signal exceeded ± 250 μV were excluded from analysis. Electroencephalogram channels were re-referenced to an average mastoids reference and spectrally analyzed using a discrete Fourier transform with a 1-second Hannning window with 50% overlap between adjacent windows. Consistent with prior research specifying EEG frequency bands in early childhood (43–45), spectral power in the following frequency bands was computed at baseline and 30 and 42 months: theta (3–5 Hz), alpha (6–9 Hz), and beta (10–18 Hz). At 96 months, power was computed using different age-appropriate frequency bands: theta (4–6 Hz), alpha (7–12 Hz), and beta (13–20 Hz).

We computed a standard metric of FEA (1,5,11) using the left (F3) and right electrodes (F4) over frontal scalp regions. Asymmetry was calculated by subtracting the natural logarithm of power in the left frontal lead from that in the right lead [ln (F4) – ln (F3)]. Because alpha power is inversely associated with cortical activation, values above 0 reflect greater activation in left relative to right frontal regions and values below 0 reflect greater activation in right relative to left frontal regions. We computed an asymmetry index for homologous parietal leads [ln (P4) – ln (P3)] to evaluate whether observed associations were specific to frontal regions (5,26,27).

Electroencephalogram data at baseline were collected from 166 children who were at least 9 months of age, and from 105, 90, and 143 children at the 30-month, 42-month, and 96-month assessments, respectively. Electroencephalogram data were unavailable from a subset of children at each time point because of fussiness during placement of the cap, parent refusal, or excessive noise across channels. There were no significant group differences in the rate of attrition or data loss at any time point.

Statistical Analysis

We used multilevel modeling to characterize the development of FEA over time and to identify predictors. A series of two-level models (observations over time nested within persons) were estimated. This approach allowed us to simultaneously estimate the variance in FEA both within and between individuals over time (49). We first estimated an unconditional growth model that predicted FEA by time, with baseline coded as zero, and subsequent time points coded as the number of months from baseline. This model was specified as follows:

**Level 1:** \( \text{Asymmetry}_t = \pi_0 + \pi_1 \text{Time} + \epsilon_t \)

**Level 2:** \( \pi_0 = \beta_{00} + r_{0i} \)
\( \pi_1 = \beta_{10} + r_{1i} \)

In this model, \( \pi_0 \) (the intercept) represents the level of FEA at baseline, \( \pi_1 \) is the effect of time, and \( \epsilon_t \) represents time-specific residual variance in FEA for child \( i \) at time \( t \). \( \beta_{00} \) represents the average FEA at baseline across children, \( \beta_{10} \) represents the average slope (rate of change over time), and \( r_{0i} \) and \( r_{1i} \) represent the random effects or individual deviations from these mean values. We added a quadratic term for time to the model to determine the functional form of the growth trajectory and tested the difference in model fit between the linear and quadratic model using a chi-square test. The quadratic model was a better fit to the data (see Results) and was specified as follows:

**Level 1:** \( \text{Asymmetry}_t = \pi_0 + \pi_1 \text{Time} + \pi_2 \text{Time}^2 + \epsilon_t \)

**Level 2:** \( \pi_0 = \beta_{00} + r_{0i} \)
\( \pi_1 = \beta_{10} + r_{1i} \)
\( \pi_2 = \beta_{20} \)

In this model, \( \beta_{20} \) represents average quadratic growth across children, which was modeled as a fixed effect (i.e., did not vary across children). Including a random effect to the quadratic term did not significantly improve model fit. With a quadratic effect of time that is equal for all individuals, the linear slope is interpreted as the rate of linear change at the intercept (because it is the effect of \( \pi_1 \) when all other predictors equal zero). This effectively determines the tilt of the curve defined by the quadratic term. However, because the quadratic effect was found to be equal for all individuals, variability in the linear slope captures all individual differences in the rate of change in FEA over time.

We next examined level 2 (time-invariant) predictors of FEA. Specifically, we examined the association of institutional rearing with FEA intercepts (value at baseline) and slopes (linear change over time). This model was specified as follows:

**Level 1:** \( \text{Asymmetry}_t = \pi_0 + \pi_1 \text{Time} + \pi_2 \text{Time}^2 + \epsilon_t \)

**Level 2:** \( \pi_0 = \beta_{00} + \beta_{01} \text{Institutionalization} + r_{0i} \)
\( \pi_1 = \beta_{10} + \beta_{11} \text{Institutionalization} + r_{1i} \)
\( \pi_2 = \beta_{20} \)

In this model, \( \beta_{01} \) represents the effect of institutionalization on FEA at baseline, and \( \beta_{11} \) represents the effect of institutionalization on the rate of linear change in FEA over time. We also evaluated whether, among the institutionalized group, the FC intervention and timing of placement predicted FEA slopes. Multilevel modeling was conducted using the Hierarchical Linear Modeling software system (HLM6.0; Scientific Software International, Lincolnwood, Illinois) (50) using full maximum likelihood estimation with robust standard errors and heteroscedastic level 1 time-specific residuals.

Finally, we evaluated whether changes in FEA from baseline to 42 months were related to psychopathology at 54 months. Linear regression models were estimated that included terms for baseline and 42-month FEA to evaluate whether residualized change in FEA predicted 54-month psychopathology. Analyses controlled for birth weight and head circumference.

Results

Development of Frontal EEG Asymmetry

We plotted the average values for FEA at each assessment for the entire sample, which suggested a nonlinear growth trajectory
of FEA over time (Figure 1). To confirm this, we compared the fit of an unconditional model that included only time as a level 1 predictor with a model that included time and time$^2$. The model that included a quadratic term was a better fit to the data, $\chi^2 (3) = 31.7, p < .001$. The quadratic model indicated a significant effect of both time, $\beta_{10} = -.05, p < .001$, and time$^2$, $\beta_{20} = .01, p < .001$. This model also revealed significant variance components for the intercept, $\tau_0 = .08, p < .001$, and slope, $\tau_1 = .03, p < .001$, of FEA, indicating significant variability in baseline values and changes in FEA over time across children. Correlation coefficients for FEA at each assessment are shown in Table 2.

### Institutional Rearing and the Development of Frontal EEG Asymmetry

To determine whether institutionalization was associated with the development of FEA, we added a dummy variable coding whether each child had been raised in an institution at level 2. This model revealed significant associations between institutionalization and FEA intercepts, $\beta_{01} = .04, p = .024$, and slopes, $\beta_{11} = -.02, p = .014$. Although children raised in institutions had left greater than right frontal activity at baseline compared with control children, they exhibited a growth trajectory characterized by increasing frontal EEG activity in the right relative to the left hemisphere. Visual inspection of the growth trajectories revealed that children in both groups experienced an initial increase in right relative to left hemisphere frontal EEG activity, followed by an increase in left relative to right hemisphere activity (Figure 2). The initial increase in right hemisphere frontal EEG activity occurred for a longer period of time in institutionalized children, however, and the subsequent rebound in left frontal EEG activity was less marked for children exposed to institutionalization. By 96 months, children raised in the community exhibited fairly pronounced FEA characterized by left greater than right hemisphere EEG activity, whereas institutionalized children exhibited a pattern of right greater than left hemispheric activity (Table 3).

Institutionalization was unassociated with baseline values of parietal EEG asymmetry, $\beta_{01} = .01, p = .669$, or with changes in parietal asymmetry over time, $\beta_{11} = -.01, p = .774$, suggesting regional specificity.

### Foster Care Effects on the Development of Frontal EEG Asymmetry

We evaluated whether the intervention was associated with FEA growth trajectories by including a variable for FC at level 2. This model was estimated only among children exposed to institutionalization. We were interested only in the association of FC with the slope of FEA, because all children were still in institutional care at baseline. The FC intervention was unrelated to the growth trajectory of FEA over time, $\beta_{11} = .01, p = .663$. Next, we examined whether FC impacted the development of FEA among children placed at an early age. To do so, we created three dummy variables coding different cut points for timing of placement: 6, 12, and 24 months. These variables were added one at a time to the model. This analysis revealed a significant effect of the FC intervention on FEA among children placed before 24 months, $\beta_{11} = .02, p = .039$. Institutionalized children who were placed into FC before 24 months exhibited a more favorable growth trajectory of FEA than children placed after 24 months (i.e., a shorter increase in right frontal activity relative to left).

Foster care was unassociated with baseline values of parietal EEG asymmetry, $\beta_{01} = -.00, p = .980$, or with changes in parietal EEG asymmetry over time, $\beta_{11} = -.01, p = .531$. Timing of placement was also unassociated with changes in parietal EEG asymmetry ($p$ values $>.300$).

### Changes in Frontal EEG Asymmetry and Psychopathology

To determine whether changes in FEA were associated with the onset of psychopathology, we predicted internalizing symptoms (anxiety, depression) and externalizing symptoms (attention-deficit/hyperactivity disorder, oppositional defiant disor-

![Figure 1. Asymmetry in alpha power in frontal cortical areas (see Methods and Materials for details) in entire sample. Higher alpha power reflects lower cortical activity; asymmetry values greater than zero reflect left greater than right frontal activity and values less than zero reflect right greater than left frontal activity. EEG, electroencephalogram.](image1)

![Figure 2. Asymmetry in alpha power in frontal cortical areas (see Methods and Materials for details), separately for children reared in institutional settings and children raised by families in the community. Higher alpha power reflects lower cortical activity; asymmetry values greater than zero reflect left greater than right frontal activity and values less than zero reflect right greater than left frontal activity. EEG, electroencephalogram.](image2)

**Table 2. Correlations of Frontal Electroencephalogram Asymmetry Across Development**

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>30 Months</th>
<th>42 Months</th>
<th>96 Months</th>
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<tbody>
<tr>
<td>Baseline</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 Months</td>
<td>.22$^b$</td>
<td>.18$^b$</td>
<td>.12</td>
<td>.05</td>
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<tr>
<td>42 Months</td>
<td></td>
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<tr>
<td>96 Months</td>
<td></td>
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</tbody>
</table>

$^a$Children ranged in age from 6 to 30 months at baseline.

$^b$Significant at .05 level, two-sided test.
Institutionalized .010 (.08)
Institutionalized 3.60 (.61) 3.82 (.40) 3.85 (.41) 56.95 (31.07)
Community Control Subjects 3.82 (.45) 3.94 (.41) 3.83 (.47) 66.55 (28.48)

Table 3. Alpha Absolute Power in Frontal Scalp Regions and Frontal EEG Asymmetry in Children Exposed to Institutionalization and Children Raised in the Community Across Development

<table>
<thead>
<tr>
<th></th>
<th>Baselinea</th>
<th>30 Months</th>
<th>42 Months</th>
<th>96 Months</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>(SD)</td>
<td>Mean</td>
<td>(SD)</td>
</tr>
<tr>
<td>Institutionalized</td>
<td>3.64 (.57)</td>
<td>3.82 (.41)</td>
<td>3.78 (.39)</td>
<td>56.96 (32.4)</td>
</tr>
<tr>
<td>Foster care group</td>
<td>3.73 (.50)</td>
<td>3.88 (.39)</td>
<td>3.84 (.38)</td>
<td>60.48 (37.19)</td>
</tr>
<tr>
<td>Institutional care group</td>
<td>3.61 (.59)</td>
<td>3.81 (.40)</td>
<td>3.76 (.42)</td>
<td>53.38 (26.62)</td>
</tr>
<tr>
<td>Community Control Subjects</td>
<td>3.82 (.45)</td>
<td>3.94 (.41)</td>
<td>3.83 (.47)</td>
<td>66.55 (28.48)</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>(SD)</th>
<th>Mean</th>
<th>(SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Institutionalized</td>
<td>3.60 (.61)</td>
<td>3.82 (.40)</td>
<td>3.85 (.41)</td>
<td>56.95 (31.07)</td>
</tr>
<tr>
<td>Foster care group</td>
<td>3.69 (.48)</td>
<td>3.88 (.37)</td>
<td>3.89 (.41)</td>
<td>60.00 (33.55)</td>
</tr>
<tr>
<td>Institutional care group</td>
<td>3.58 (.65)</td>
<td>3.79 (.35)</td>
<td>3.81 (.45)</td>
<td>53.83 (28.32)</td>
</tr>
<tr>
<td>Community Control Subjects</td>
<td>3.80 (.47)</td>
<td>4.00 (.42)</td>
<td>3.83 (.41)</td>
<td>65.77 (29.37)</td>
</tr>
</tbody>
</table>

Institutionalized .016 (.04)
Foster care group .004 (.00)
Institutional care group .006 (.05)
Community Control Subjects .000 (.05)

Discussion

Asymmetrical hemispheric EEG activity recorded over frontal scalp regions is a well-established biological substrate of affective style and approach/withdrawal behavior associated with mental health and social functioning (1–3,5,6,11). Although asymmetrical EEG activation in the frontal cortex has been documented in young children (5,21,23), we are unaware of a previous study that examined the developmental trajectory of FEA or identified factors that influence patterns of development. We provide novel evidence for developmental changes in FEA from infancy through middle childhood. Specifically, we find an increase in right relative to left hemisphere frontal EEG activation between the second and fourth years of life followed by a subsequent increase in left relative to right hemisphere activation into middle childhood. Moreover, our findings suggest that the caregiving environment early in life influences developmental trajectories of FEA. The lack of associations between the rearing environment and parietal EEG asymmetry suggests regional specificity to neural circuits in the frontal cortex. Frontal EEG asymmetry development from infancy through early childhood is associated with internalizing symptoms, suggesting that the developmental course of relative hemispheric activation in the frontal cortex has important implications for mental health.

To our knowledge, we provide the first evidence documenting the developmental trajectory of FEA in early to middle childhood. Our results indicate that children exhibit a relatively greater increase in right relative to left frontal EEG activation during the second and third years of life. Electroencephalogram activation in the right frontal cortex underlies withdrawal behavior in response to aversive stimuli (1–3), and in infants this pattern has been observed during the approach of a stranger and following maternal separation (4,17). Although the specific neural mechanisms underlying FEA are unknown, it has recently been proposed that asymmetry is related to affective style because of the role of the left prefrontal cortex in inhibiting the amygdala (51). Plaisier et al. (52) find resting alpha activity in the left dorsolateral prefrontal and medial orbitofrontal cortex underlies motivated approach behavior in response to appetitive stimuli. Increased activation in the right frontal cortex beginning at age 2 may reflect an adaptive developmental change in withdrawal tendencies to novelty as the child’s experiences expand to encompass environments outside the home and interactions with people other than their primary caregivers. A concomitant increase in numerous aspects of effortful control—including delaying and inhibiting behavior—has also been observed among children during this developmental period (53) and is thought to reflect maturation of the prefrontal cortex (54). Together with our findings, these patterns suggest normative increases in behavioral inhibition in 2- to 3-year-olds that are mediated by developmental changes in the prefrontal cortex.

We also demonstrate an association between early-life caregiving experiences and the development of FEA. First, institutionalized children exhibited greater left than right frontal activation at baseline. This finding indicates a greater tendency for approach behavior among institutionalized children during infancy, a period in which responsive caregiving is critical to scaffold neural develop-

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ment. This pattern may reflect an adaptive response to the institutional environment where there is a scarcity of individualized social attention from caregivers. Second, children exposed to institutional rearing experienced a prolonged period of increased right relative to left hemisphere EEG activation in early childhood and a blunted rebound in left frontal EEG activity. By age 96 months, children raised in the community exhibited considerably greater left relative to right frontal activation, whereas children reared in institutions exhibited greater activation in the right frontal cortex compared with the left. These differences in brain development likely reflect a greater propensity for withdrawal behavior in response to novelty among institutionalized relative to noninstitutionalized children (5,8). As children raised in institutions become older, they accumulate increasing experiences of insensitive unresponsive caregiving (55); thus, withdrawal from novel stimuli may be an adaptive response to an aversive environment. The neural underpinnings of this pattern of withdrawal, however, are associated with increased risk of internalizing psychopathology. Although right greater than left activation appears normative in early childhood, a delayed rebound of left frontal activation may be a marker of psychopathology risk.

Our results are consistent with evidence linking poor maternal caregiving in infancy to FEA in children (26,27), as well as an extensive literature linking poor caregiving to elevated stress reactivity in animals (56–59). We extend this work by showing that an adverse rearing environment is associated with the developmental trajectory of relative hemispheric activation in the frontal cortex. These findings add to a growing literature documenting that early deprivation alters brain development in ways that increase risk for psychopathology. Abnormalities in the neural processing of facial emotion have frequently been observed in maltreated children (60,61), and we recently found that these deficits predict the onset of externalizing problems in institutionally reared children (Slopen N, et al.; Alterations in neural processing of facial emotions and psychopathology in children exposed to institutional rearing; unpublished data). In previous work, we also observed abnormal development of high-frequency brain electrical activity among children reared in institutions—suggesting a delay in maturation of the cerebral cortex—that predicted the onset of attention-deficit/hyperactivity disorder (62). That pattern raises the question of whether the frequency band used to define alpha is appropriate for the institutionalized group. Although our data cannot speak directly to this issue, the frequency band used to define alpha from baseline to 42 months incorporates normative alpha in children aged 5 to 51 months (43), increasing confidence that this frequency band is appropriate. The current findings build on previous work by documenting a neurodevelopmental mechanism linking early-life deprivation to the onset of internalizing psychopathology. Together, these studies provide compelling evidence suggesting that adverse environmental conditions in early life may become biologically embedded, resulting in lasting changes in neural development that may ultimately manifest as psychopathology.

Foster care placement had a positive effect on FEA trajectories but only among children placed before 24 months. This suggests that the brain circuits underlying hemispheric EEG activation in the frontal cortex are responsive to environmental input relatively early in life and become less susceptible to these influences around 2 years of age. Early-life experiences have been important predictors of FEA in previous studies (23,26,27). The beneficial effects of FC for children placed before 24 months is consistent with previous reports from this sample that also found substantial benefit for children placed before 24 months in other developmental domains, including cognitive ability and attachment security (63,64). Children placed before 24 months also exhibit greater increases in alpha power by age 8 relative to children placed later (65). Together, these findings suggest a possible sensitive period during which the environment has a particularly pronounced effect on brain development—particularly the development of high-frequency brain electrical activity and hemispheric activation patterns in the frontal cortex. An important caveat related to these findings is our inability to disentangle age at placement from duration of institutionalization. One explanation for the observed results is that the neural circuits underlying asymmetrical frontal EEG activation are most sensitive to environmental input during the first 2 years of life; an alternative explanation is that these neural systems are more profoundly affected in children who experience more chronic deprivation and therefore require additional time to catch up. These effects are difficult to disentangle in our sample but highlight an important topic for future research. At the very least, our findings highlight the importance of intervening as early as possible to reduce the long-term effects of adverse rearing environments on brain development.

Deprived rearing environments are associated with a developmental trajectory of FEA characterized by a prolonged period of greater activation in the right hemisphere as compared with the left. This developmental pattern predicts the onset of internalizing psychopathology in early childhood. Exposure to adverse caregiving environments early in life can alter the trajectory of brain development in children, culminating in heightened risk for psychopathology. Interventions delivered early in the life course have the greatest potential to mitigate the long-term effects of these adverse environments.

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