

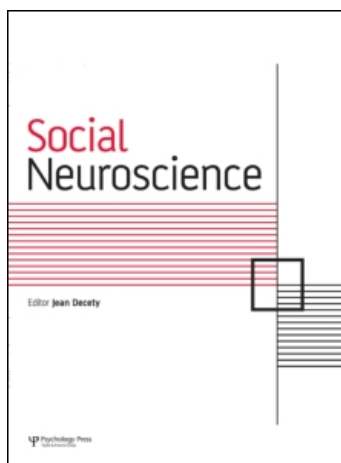
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Susan B. Perlman ^a; Kevin A. Pelphrey ^a

^a Yale University, New Haven, CT, USA

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Regulatory brain development: Balancing emotion and cognition

Susan B. Perlman and Kevin A. Pelphrey

Yale University, New Haven, CT, USA

Emotion regulation is a critical aspect of children's social development, yet few studies have examined the brain mechanisms involved in its development. Theoretical accounts have conceptualized emotion regulation as relying on prefrontal control of limbic regions, specifying the anterior cingulate cortex (ACC) as a key brain region. Functional magnetic resonance imaging in 5- to 11-year-olds during emotion regulation and processing of emotionally expressive faces revealed that older children preferentially recruited the more dorsal "cognitive" areas of the ACC, while younger children preferentially engaged the more ventral "emotional" areas. Additionally, children with more fearful temperaments exhibited more ventral ACC activity while less fearful children exhibited increased activity in the dorsal ACC. These findings provide insight into a potential neurobiological mechanism underlying well-documented behavioral and cognitive changes from more emotional to more cognitive regulatory strategies with increasing age, as well as individual differences in this developmental process as a function of temperament. Our results hold important implications for our understanding of normal development and should also help to inform our understanding and management of emotional disorders.

Keywords: Emotion regulation; Development; fMRI; Anterior cingulate; Temperament.

INTRODUCTION

Proficiency in the management of affective arousal is a critical aspect of early development. Infants move from a necessary dependence on the support of their caregivers to the development of more self-sufficient mechanisms for emotion regulation at school age (Denham, 1998). Grolnick, Bridges, & Connell (1996), for example, discuss emotion regulation in early childhood as developing along a continuum in which young children and infants are initially unable to remove focus from emotional distress, without adult assistance, but later develop self-soothing strategies and the ability to reorient attention towards a more pleasant stimulus. Behavioral studies in pre-school and school-age children have examined both

children's modulation of overt emotional behavior (Cole, 1986; Saarni, 1984) and subjective emotional state (Harris, Olthof, & Meerum Terwogt, 1981). To illustrate, Cole (1986) found that preschool children are able to suppress overt negative emotional facial and vocal expression in response to social demands, while Harris and colleagues (1981) documented that by later school years (age 11–15) children become adept at modifying internal emotional states.

Since this early research, the characterization of individual differences in the typical and atypical development of affective regulation has become a well-established area of psychological science (Cole, Martin, & Dennis, 2004). However, little is known about the biological mechanisms supporting the developmental transitions in emotion regulation during early

Correspondence should be addressed to: Susan B. Perlman, University of Pittsburgh Medical Center, 121 Meyran Ave., Room 203, Pittsburgh, PA 15213, USA. E-mail: perlmanb2@upmc.edu

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childhood or about the biological bases of individual differences in developmental pathways. This is unfortunate because information about the brain mechanisms underlying the typical and atypical development of emotion regulation could greatly inform our understanding of early social development as well as the disorders of emotion regulation (Lewis & Stieben, 2004; Lewis & Todd, 2007). Thus far, investigation of brain mechanisms for the development of emotion regulation has employed event-related potential (ERP) measures to implicate the prefrontal cortex in cognitive control of emotion. Lewis, Lamm, Segalowitz, Stieben, & Zelazo (2006), for example, discovered that a temporary loss of points towards a desired prize triggered an increase in the frontal P3 response, and this response changed with age to a more midline localization. This developmental change coincided with a decrease in self-reported negative emotions in children (ages 5–16 years). Here, we report the results of one of the first functional magnetic resonance imaging (fMRI) studies of emotion regulation process in school-age children.

The anterior cingulate cortex (ACC) is a specialized prefrontal region that has been consistently implicated in affective regulation in the adult neuroimaging literature (Allman, Hakeem, Erwin, Nimchinsky, & Hof, 2001; Bush, Luu, & Posner, 2000). Located between the neocortex and the limbic system, the ACC is well positioned to serve as an interface between cognition and emotion. This region contains spindle-shaped neurons allowing for widespread connections to other brain areas. Found only in humans and great apes (Nimchinsky et al., 1999), these cells become functionally mature at four to six months of age, coinciding with an infant's developing ability to divert attention and self-soothe (Grolnick, McMenemy, & Kurowski, 2006). Posner and Rothbart (1998, 2000) have noted an increase in cognitive control of emotion at the age of 4 years, which they relate to the child's emerging capacity to employ the ACC to manage impulsive emotional behavior through conscious regulatory strategy use. These theorists have hypothesized that variation in functional ACC maturation, and, consequently emotion regulation, is related to developmental and temperamental differences in the modulation of attention. Models of ACC function hypothesize divisions of a "cognitive" dorsal region, in proximity to dorsolateral prefrontal cortex (PFC), and an "emotional" ventral region close to orbitofrontal cortex and the amygdala (Bush et al., 2000). The dorsal division has been most often implicated in "cool" (Zelazo & Müller, 2002) executive function, activating during error monitoring and cognitive Stroop tasks, while ventral activity is most

often linked to "hot" executive function (Zelazo & Müller, 2002), including emotional Stroop tasks and recall of emotional events (Bush et al., 2000). Both divisions project to the amygdala and, via these connections, are thought to support affective downregulation. However, the dorsal pathway is hypothesized to be preferentially involved more in deliberate and consciously controlled regulatory processes (Lewis & Todd, 2007).

In a typically developing sample of 5- to 11-year-old children, we used fMRI to investigate the development of the dorsal and ventral divisions of the ACC and their relationships to developmental and temperamental differences in the modulation of emotion. We hypothesized that when contrasting periods of high and low regulatory demand, the change in blood-oxygen-level-dependent (BOLD) activity in the dorsal (cognitive) area of the ACC would increase as a function of increasing age, while change in the activity of the ventral (emotional) portion of this region would decrease with age. This pattern should reflect a developmental shift to cognitively oriented regulation strategies (Harris et al., 1981). We further hypothesized that decreased activation in the dorsal ACC would be related to fearful temperament, while increased ventral ACC activation would be correlated with fearfulness. This hypothesis was informed by previous electrophysiological studies reporting a negative influence of inhibited temperament on the brain's ability to control arousal in infancy and childhood (Fox et al., 1995).

METHOD

Subjects

Subjects were 20 (9 male) English-speaking, healthy, and typically developing children aged 5–11 years (average age = 8 years, 2 months) with normal or corrected-to-normal vision. An additional 6 children were scanned, but not included in data analyses due to excessive head movement (4 children), inability to complete the scan (1 child), or drowsiness (1 child). These children did not differ from children with usable data in age, $t(24) = -.19, p = .85$, or on any dimension of temperament, $t(24) = -.50-1.83, p = .08-.97$; (see below). Children were recruited from local community postings and internet advertisements and were paid for their participation. In addition, they earned an age-appropriate prize as part of their task participation (see below). A parent or legal guardian of each child provided written informed consent for their child's participation. Each child provided written assent.

Materials and procedure

Before data collection and practice of the task, children were told that they would be playing to win a prize: their choice of a toy from a large collection. After choosing their desired prize, children were given a few minutes to play with the toy and show it to their parent and/or siblings. They were then told that they would need to earn a large number of points during the game in order to keep their prize, and that they would earn points based on speed and accuracy. Children were not told the specific number of points that they would need to win, but if questioned, the experimenter replied, "You need a lot of points. Over a thousand, at least!" They were reminded that they would not be told whether they won or lost until the end of the game. Subjects then practiced the emotional go/no-go task outside of the scanner to ensure understanding during fMRI data collection.

Before scanning began, children participated in a "mock scanning" session to help ensure compliance with the requirement to remain motionless during data collection, and to help them to feel as comfortable as possible while participating in the actual experiment. Children were trained to remain still while watching a children's video inside a replica of our MRI scanner. During practice, custom-written software received input from a head motion sensor worn by the child and used that input to play a sound when the child moved outside of a set threshold (3 mm). In addition to viewing approximately 10 min of the video, children again practiced the go/no-go task inside the mock scanner. With the addition of realistic scanner sounds played during the children's practice session, we were able to closely reproduce the scanning environment in order to familiarize our young subjects with fMRI scanning procedures, thereby improving compliance and increasing the likelihood of obtaining high-quality data from the young children.

Subjects participated in a novel, event-related within a blocked design, emotion regulation task partially adapted from previous studies (Garavan, Ross, & Stein, 1999; Lewis et al., 2006). The emotional go/no-go task was presented using E-prime software (Psychological Software Tools, Pittsburgh, PA) and is depicted in Figure 1. Throughout the paradigm, a stream of pictures of common objects (e.g. balls, shoes, umbrellas) was presented at a rapid rate. Subjects were instructed to press a button when the object was presented in a green frame, but to inhibit their response when the object was presented in a red frame. Incorrect responses to

both go and no-go trials were followed by a large "X" on the center of the screen and the sound of a buzzer. Periodically, fearful faces from the *NimStim* set of facial expressions (Tottenham et al., 2009), appeared for a 1-s duration, accompanied by a short chime. Children were told that the presentation of the fearful face was unrelated to the game and did not reflect their task performance. Subjects were also notified of their total number of accumulated points approximately every 30 s throughout the task. As point totals were presented, subjects heard a "slide whistle" sound indicative of point loss or point gain.

This task was designed to maintain the same level of difficulty for all subjects. To this end, an algorithm was used to keep all children functioning at approximately the same error rate ($50 \pm 10\%$; Lewis et al., 2006). The algorithm was piloted with 25 5-year-olds (outside of the scanner) to ensure that children this young could successfully complete our task. The error rate for the task was maintained by adjusting the stimulus duration dynamically (see the top panel of Figure 1). Stimulus duration increased with each error made on a no-go trial and decreased with each correct response. Unknown to subjects, the task contained three blocks designed to induce different types of emotion (Lewis et al., 2006). Each block lasted approximately 4 min. During Block 1 (Winning), subjects saw their points steadily increase to well over 1000 with a stimulus interval set between 800 and 1150 ms. This block was designed to induce positive emotions related to success in our subjects, even though the task was designed to be challenging and engaging throughout. Changes in the point-adjustment algorithm and stimulus presentation speed (600–950 ms) caused increased task difficulty and point loss during Block 2 (Losing), which was intended to induce negative emotions of frustration. Children lost all of their previously earned points during this block. With a return to the more generous algorithm in Block 3 (Recovery), subjects regained their points and ultimately won their desired prize. Although there was notable point gain, we designed this block to represent a recovery period after an emotionally negative event. The experiment was designed with three long blocks in order to allow subjects the necessary time to feel the induced positive or negative emotions due to relatively lengthy periods of gaining or losing points, respectively. In order to ensure that subjects were unaware of our manipulation, there were no pauses between emotion induction blocks.

In our analysis, the 1-s presentations of the fearful faces were modeled as events. Each block contained

| | Block 1 - Winning | Block 2 - Losing | Block 3 - Recovery |
|---|-------------------|------------------|--------------------|
| Stimulus Presentation | 800-1150 ms | 600-950ms | 800-1150 ms |
| Correct No-Go Response | -20 ms | -50 ms | -20 ms |
| Incorrect No-Go Response | +70 ms | +50 ms | +70 ms |
| Points for Go Trial Correct/Incorrect | +3/0 | 0/-15 | +15/+1 |
| Points for No-Go Trial Correct/Incorrect | +7/0 | 0/-15 | +25/+1 |

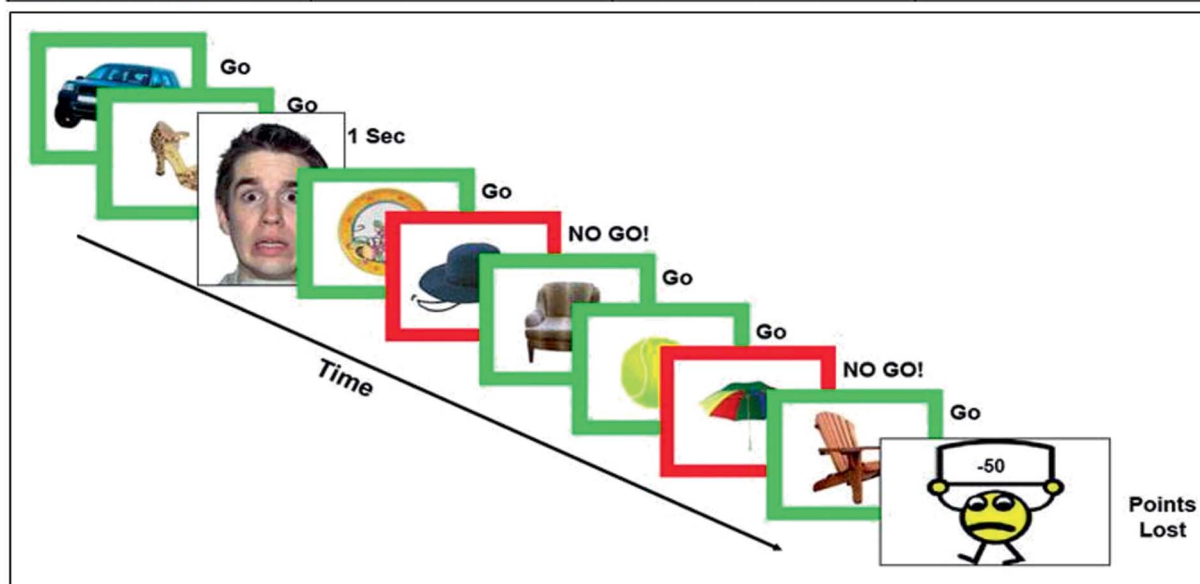


Figure 1. (Top) Stimulus presentation time and point adjustment algorithm for each of three emotion induction blocks. Subjects won points during Block 1, but then lost all earnings in Block 2. During Block 3, which had the same task difficulty level as Block 1, subjects regained their points to win their desired prize. (Bottom) A graphic representation of our go/no-go task. The paradigm was 12 min in length with three mood induction blocks and sporadic fearful faces presented as 1 s events.

20 fearful face events (60 total trials) that were presented 6–15 s apart regardless of emotion induction block or changes to the speed of the point adjustment algorithm. We compared the fearful face events between the Winning and Recovery blocks because they were essentially identical in speed and difficulty of the point adjustment algorithm, but offered very different regulatory contexts. In the Winning block, subjects were expected to be in a positive mood while rapidly gaining points compared to the Recovery block, which followed a significant point loss and the induction of negative emotions. We note that the go/no-go task in this study was employed simply as a mechanism to challenge children within an emotional context and was presented at speeds varying by participant success rate. We therefore did not model specific go and no-go trials in our analysis.

Manipulation check

Our child subjects were interviewed regarding their emotional experiences during the task. One hundred percent of children recalled losing all of their points during the game and 95% (all but one child) reported feeling negative emotions during that episode. Ninety percent reported feeling positive emotions when they discovered that they would, indeed, win their prize. Finally, 74% of children reported an overall enjoyment of the game.

fMRI data acquisition

Scanning was performed on a Siemens 3 T Allegra head-only scanner (Siemens, Erlangen, Germany).

High-resolution, T1-weighted anatomical images were acquired using an MPRAGE sequence (TR = 1630 ms; TE = 2.48 ms; FOV = 20.4 cm; $\alpha = 8^\circ$; image matrix = 256^2 ; voxel size = $0.8 \times 0.8 \times 0.8$ mm; 224 slices). Whole-brain functional images were acquired using a single-shot, gradient-recalled echo-planar pulse sequence (TR = 2000 ms; TE = 30 ms; $\alpha = 73^\circ$; FOV = 20.4 cm; image matrix = 64^2 ; voxel size = $3.2 \times 3.2 \times 3.2$ mm; 35 slices) sensitive to BOLD contrast. Runs consisted of the acquisition of 224 successive brain volumes beginning with two discarded RF excitations to allow for steady-state equilibrium.

The BrainVoyager QX software package (Brain Innovations, Maastricht, The Netherlands) was used for all analyses. The following preprocessing procedures were performed on the raw images: slice scan time correction (using cubic spline interpolation), high-pass temporal filtering to remove nonlinear drifts of three or fewer cycles per time course, three-dimensional motion correction to detect and correct for small head movements by spatial alignment of all volumes to the first volume by rigid body transformation, and spatial data smoothing using a Gaussian kernel with a 4 mm full width at half-maximum. Head movements never exceeded 3 mm for all participants included in this analysis. Functional data were coregistered to the anatomical volume by alignment of corresponding points to obtain optimal fit and were then transformed into Talairach space (Talairach & Tournoux, 1988). We employed the adult anatomical Talairach template. The literature suggests that after transformation into a common stereotactic space, anatomical differences between children and adults are small relative to the resolution of fMRI data (Burgund et al., 2002). Another study revealed minimal differences in the time courses and locations of functional activation foci between children and adults (Kang, Burgund, Lugar, Petersen, & Schlaggar, 2003).

A multiparticipant statistical analysis was performed by multiple linear regression of the time course of the BOLD response in each voxel. The general linear model of the experiment was computed for each participant's z -normalized volume time courses. Model predictors were defined by convolving an ideal boxcar response with a gamma-function model of the hemodynamic response (Friston et al., 1995). Boxcar values were equal to 1.0 during the face presentation and were 0.0 during presentation of objects. Activation maps were visualized on a Talairach-transformed brain of an 8-year-old male subject (see Figure 2), with only clusters of more than eight contiguous voxels displayed at 1 mm resolution.

Temperament data collection

Caregivers were asked to provide information on their children's temperament. They completed the Child Behavior Questionnaire (CBQ)–Long Form (Rothbart, Ahadi, Hershey, & Fisher, 2001), an experimentally validated and commonly used caregiver assessment of 15 dimensions of children's temperament. Our hypotheses focused on the fearful dimension of temperament due to its relationship to behavioral inhibition and anxiety, which have been implicated in regulatory deficits (Fox et al., 1995).

RESULTS

An examination of the main effect of condition (Winning, Losing, and Recovery) revealed several neural areas that were significantly active at the false discovery rate (FDR) $q < .05$ criterion (Genovese, Lazar, & Nichols, 2002). The ACC (BA 32) was the largest, and most significantly active, among them. All regions are presented in Table 1.

Next, brain responses to the presentation of fearful faces during the Winning period in which children gained points towards a desired prize were contrasted with activity in response to fearful faces presented during a Recovery period, which followed point loss. To focus our analysis on the ACC as an *a priori* region of interest, we used the Analysis of Functional Neuroimages anterior cingulate gyrus region of interest (ROI) (AFNI; National Institutes of Mental Health, Bethesda, MD) to define the area structurally. Across the ACC, the Winning condition was contrasted with the recovery condition and then correlated with our auxiliary variables, at the significance level of $p < .05$, $r(18) \geq .50$. For these image analyses, we used a statistical threshold criterion of $p < .001$ corrected for multiple comparisons using a cluster-size threshold adjustment, based on a Monte Carlo simulation approach extended to 3D data sets using the cluster threshold size estimator plug-in BrainVoyager QX (Forman et al., 1995). This procedure takes input regarding the functional voxel size (3 mm^3 native resolution), the total number of significant voxels within a map, and the estimated smoothness of a map and performs Monte Carlo simulations (1000 iterations) to estimate the probability of clusters of a given size arising purely by chance. By applying a spatial cluster threshold of 828 mm^3 , or 28 contiguous voxels, the family-wise (corrected) false probability rate was $p < .001$.

This analysis identified a section of the dorsal ACC in which greater positive change in the level of

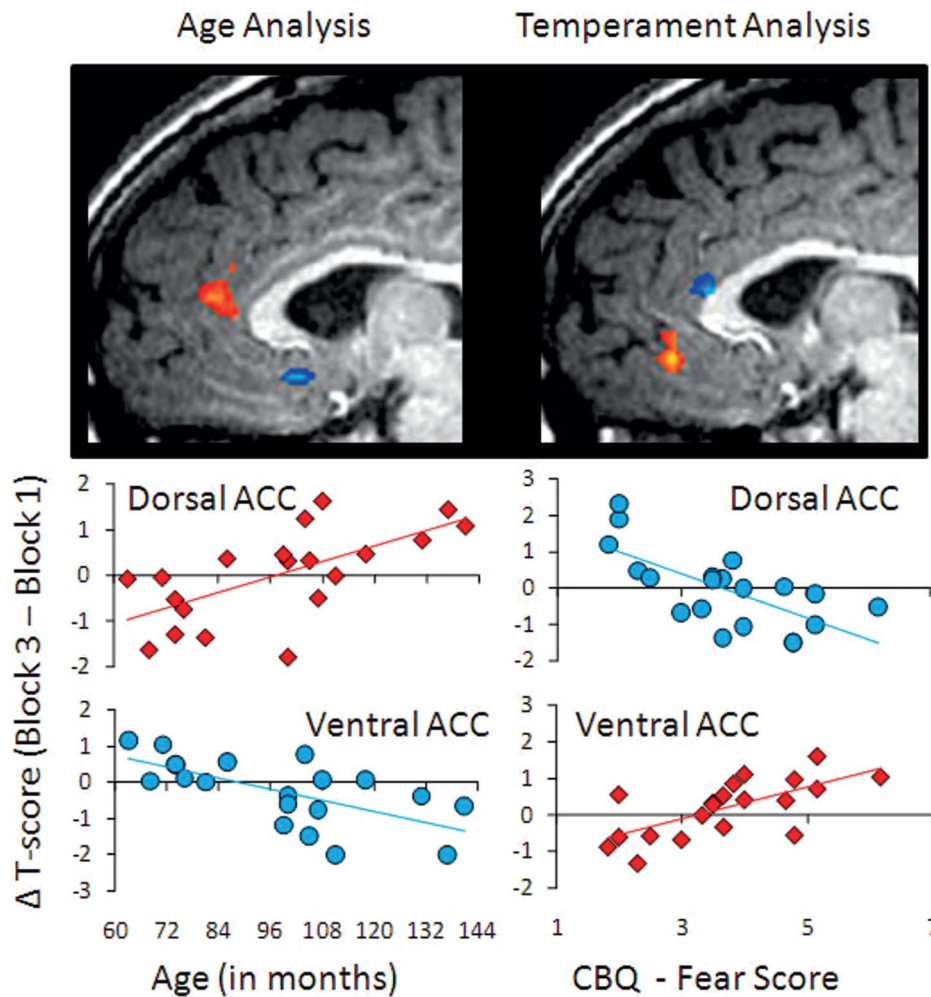


Figure 2. Correlations between age and fearful temperament and the difference in activation between pre-induction and post-induction blocks during 1 s presentation of fearful faces. Red represents positive correlation and blue represents negative correlation within the anatomically defined anterior cingulate cortex.

activity between Winning and Recovery correlated positively with a child's age, as well as a section of the ventral ACC in which greater positive change in activity levels between Winning and Recovery correlated negatively with age (see Figure 2). A regression analysis involving the change in activation from Winning to Recovery in both the dorsal and ventral areas simultaneously regressed onto child age revealed a significant overall model, $F(2, 19) = 20.375, p < .001$, with both the dorsal, $\beta = .545, p < .001$, and ventral, $\beta = -.543, p < .001$, regions of the ACC being significantly different from zero.

In addition, we examined the fearful dimension of temperament in relation to activity evoked by the presentation of faces during mood induction. Our analysis revealed a portion of the dorsal ACC in which

greater positive change in activity levels between Winning and Recovery correlated negatively with fearfulness and a portion of the ventral ACC in which greater positive change in activity from Winning to Recovery correlated positively with this dimension of temperament. In both our age and temperament correlations, clusters of activation appeared to be localized to the cognitive and emotional divisions of the ACC (Bush et al., 2000). Notably, child age and fearful temperament were uncorrelated in our sample, $r(18) = -.04, p = .87$, suggesting that age and temperament were contributing independently to the double dissociation that we observed. A regression analysis was completed with the change in activation from Winning to Recovery in both the dorsal and ventral areas simultaneously regressed onto child fearful temperament. This

TABLE 1
Differences in BOLD signal between winning, losing, and recovery

| Region (main effect of condition) | Hemisphere | Talairach coordinates | | | No. of voxels | F* |
|-----------------------------------|------------|-----------------------|-----|-----|---------------|------|
| | | X | Y | Z | | |
| Anterior cingulate (BA32) | R | 9 | 35 | 22 | 11445 | 18.2 |
| Middle temporal gyrus (BA20) | R | 57 | -43 | -8 | 3769 | 15.6 |
| Inferior frontal gyrus (BA9) | R | 42 | 8 | 25 | 1765 | 12.5 |
| Inferior parietal lobule (BA40) | R | 42 | -55 | 49 | 1057 | 8.8 |
| Medial frontal gyrus (BA8) | R | 6 | 20 | 43 | 703 | 9.4 |
| Cuneus (BA18) | R | 15 | -79 | 19 | 623 | 10.5 |
| Middle occipital gyrus (BA19) | L | -30 | -82 | 10 | 617 | 11.5 |
| Cuneus (BA19) | R | 24 | -82 | 37 | 599 | 10.2 |
| Middle frontal gyrus (BA10) | R | 39 | 38 | 16 | 570 | 12.1 |
| Paracentral lobule (BA5) | R | 15 | -31 | 52 | 367 | 12.3 |
| Fusiform gyrus (BA18) | L | -27 | -91 | -11 | 250 | 9.1 |
| Inferior temporal gyrus (BA20) | R | 54 | -28 | -14 | 168 | 9.9 |
| Middle occipital gyrus (BA37) | R | 48 | -67 | -5 | 88 | 7.04 |
| Inferior temporal gyrus (BA20) | L | -42 | -1 | -46 | 51 | 13.2 |

Notes: R, right; L, left; BA, Brodmann area; * $q < 0.05$ FDR corrected for multiple comparison.

analysis revealed a significant overall model, $F(2, 19) = 25.293$, $p < .001$, with both the dorsal, $\beta = -.555$, $p < .001$, and ventral, $\beta = .550$, $p < .001$, regions of the ACC being significantly different from zero.

DISCUSSION

Our results provide novel insights into the biological basis of regulatory mechanisms across child development. Extrapolating from functional models of the dorsal and ventral divisions of the ACC (Bush et al., 2000), older children in our experiment appeared to recruit the more dorsal “cognitive” areas of the ACC when required to emotionally regulate in the face of a frustrating task, while younger children engaged the more ventral “emotional” areas. We propose that this shift in prefrontal activation underlies the well-documented change to more cognitive regulatory strategy use (e.g. reappraisal) as children develop (Harris et al., 1981).

Our findings concerning age-related changes in ACC function are consistent with and serve to advance a broader literature concerning the development of the PFC and its relationship to the emergence of more effective executive functioning with age (Casey et al., 2000). The PFC is among the last brain areas to fully develop, with continued synaptogenesis (Rakic, Bourgeois, & Goldman-Rakic, 1994) and increasing white matter volume (Pfefferbaum et al., 1994; Shaw et al., 2006) until late adolescence or early adulthood. This protracted developmental course has been linked to improvements in various

complex cognitive skills (Caviness, Kennedy, Richelme, Rademacher, & Filipek, 1996; Diamond, 1988; Shaw et al., 2006), which are known to be important for the modulation of affective arousal (Gross, 1998). FMRI studies investigating the development of self-regulation have demonstrated that differential recruitment of regions of the PFC may underlie decreased cognitive control of emotion in children relative to adults (Bunge et al., 2002; Casey et al., 1997; Luna et al., 2001). Casey et al. (1997), for example, found that children recruited the same PFC regions as adults during a go/no-go task. Activation was greater in children than in adults, however, suggesting inefficient recruitment of the PFC in children. This finding was echoed by Lewis et al. (2006), who related their electrophysiological results to inefficient ACC functioning during their go/no-go task.

Independent of the age-related effects, the more fearful children preferentially recruited the less effective “emotional” areas of the ventral ACC in response to emotionally expressive faces, while less fearful children engaged the more “cognitive” dorsal regions of the ACC. These results suggest a neural mechanism for the influence of fearful temperament on the development of emotion regulation abilities and add to a growing understanding of the neurobiological basis of individual differences in emotion regulation. Prior research has probed the relationship between child temperament and the brain’s ability to control emotional arousal in infancy and childhood (Rothbart & Sheese, 2007). Fox and his colleagues (Fox, 1994; Fox & Calkins, 2003) have implicated the construct of electroencephalograph (EEG) asymmetry in individual

differences in children's temperament, emotional reactivity, and regulation. Specifically, right-lateralized frontal EEG asymmetry has been related to increased fearful temperament and impaired regulation of fear during social interaction (Fox et al., 1995). In a related line of fMRI research, adolescents who had been temperamentally classified as behaviorally inhibited displayed heightened amygdala activation while rating the emotional content of face stimuli (Pérez-Edgar et al., 2007). This amygdala hyperactivity might underlie difficulties in effective regulation during social engagement. Relatedly, Drevets and Raichle (1998) have reported competition between dorsal and ventral prefrontal activation in adult subjects using positron emission tomography (PET). Notably, these authors have found that increased anxiety is related to heightened ventral, at the expense of dorsal, ACC activity.

We found evidence that changes in the balance of dorsal and ventral anterior cingulate cortical activation during regulatory episodes may be related to differences in child age and temperament. However, our conclusions are tempered by some limitations to the current study. First, we did not include online, physiological measures of anxiety and/or arousal such as heart rate, pupil diameter, or skin conductance. Indeed, these measures would have helped to pinpoint moments of increase in task-related stress and thus provided additional precision for correlation with our brain and temperament data. Second, due to the need to induce negative emotions with point loss following a point gain, we presented the blocks in a fixed order. We note that this method could have resulted in subject "drift." For example, young children may have been losing track of the task in the recovery block, while older children may have remained engaged longer. We do not believe this to be the case because our correlational analysis of fearful temperament and ACC activity indicated that the dorsal ACC was most active in non-fearful children, who were represented across the observed age range. Nonetheless, this is an important limitation which should be considered.

Finally, the construct of emotion regulation is a widely debated topic in the field of child development. There is general disagreement on the definition of this construct, the nature of its development, and how to separate regulation from aspects of general affective reactivity (Cole et al., 2004). Our approach was designed to put children in a real-life situation in which they are typically required to regulate emotion: the loss of a desirable prize. Consequently, there was no condition in which participants were explicitly instructed to regulate emotion, which is a commonly employed technique in the adult literature (e.g. Ochsner,

Bunge, Gross, & Gabrieli, 2002). While commendable in that it is difficult to mimic ecologically valid emotional situations within the confines of the scanning environment, this method also has some drawbacks. Specifically, it is difficult to pinpoint exactly when and how children were regulating emotion during this task. Although the brain and self-report data are suggestive of the engagement of regulatory mechanisms during our go/no-go task, without the explicit instruction to regulate emotion, we cannot be sure that changes in activation between Winning and Recovery are due to increased regulation or persistent reactivity of negative emotions from point loss. We are, therefore, somewhat limited in the interpretation of our results. However, we believe that such a complicated theoretical debate (Campos, Frankel, & Camras, 2004; Cole et al., 2004) requires a variety of approaches to better characterize the nature of the neural mechanisms involved in affective self-regulation. In the future, experiments will be needed to better assess the specific regulatory strategies employed during implicit emotional tasks.

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