Moderation of the relationship between the error-related negativity and anxiety by age and gender in young children: A preliminary investigation

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ABSTRACT

The error-related negativity (ERN) is a neurophysiologic response to errors that associates with anxiety. Despite the potential relevance of the ERN for understanding mechanisms of early anxiety problems in the developing brain, the relation between ERN and anxious symptoms in young children remains poorly understood. Emerging evidence suggests that ERN-anxiety associations could vary by developmental stage, but this work requires replication and consideration of gender effects, given earlier maturation of the ERN and higher rates of anxiety problems in girls relative to boys. To address this gap, the ERN was collected in 49 preschool- to school-aged children (ages 4–9; 26 girls) sampled across a wide range of anxiety severity. Regression analyses revealed that ERN-anxiety associations depended on age and gender. Specifically, larger (more negative) ERN associated with more anxiety in older girls, whereas smaller ERN associated with more anxiety symptoms in younger girls. No ERN-anxiety association was found in boys. These findings suggest that age and gender moderate the direction of the relation between ERN and anxiety in early childhood and could have important implications for the development of ERN-based risk identification and targeted treatment strategies tailored to individual children.

1. Introduction

Early anxiety, even at subclinical levels, increases risk for clinically significant internalizing problems (i.e., anxiety and depressive disorders) in mid-childhood and into adult life (Kroes et al., 2002; Mesman et al., 2001; Moffitt et al., 2007; Petty et al., 2008). As a result, early detection and prevention efforts have been widely endorsed (Novins et al., 2014; Woody and Gibb, 2015). For instance, the error-related negativity (ERN) is a neurophysiological marker of cognitive control associated with anxiety (Vaidyanathan et al., 2012) that may differentially associate with anxiety symptom severity in children compared to adolescents and adults (Meyer, 2017; Moser, 2017), and in women compared to men (Moser et al., 2016). Examining the effects of age and gender on ERN-anxiety associations in early childhood, when anxiety symptoms first present, represents an important step towards characterizing the ERN for developmentally informed application within the RDoC framework.

1.1. ERN as an RDoC biomarker associated with anxiety

To realize this promise, candidate biomarkers of RDoC constructs and related neural circuits must be understood in the context of age and gender differences known to characterize psychopathology (Casey et al., 2014; Woody and Gibb, 2015). For instance, the error-related negativity (ERN) is a neurophysiological marker of cognitive control associated with anxiety (Vaidyanathan et al., 2012) that may differentially associate with anxiety symptom severity in children compared to adolescents and adults (Meyer, 2017; Moser, 2017), and in women compared to men (Moser et al., 2016). Examining the effects of age and gender on ERN-anxiety associations in early childhood, when anxiety symptoms first present, represents an important step towards characterizing the ERN for developmentally informed application within the RDoC framework.

The ERN is a negative deflection in the event-related brain potential (ERP) that occurs within 100 ms following an erroneous response (Gehring et al., 1993), is typically maximal at frontocentral scalp locations, and is believed to be generated by anterior cingulate cortex
Moreover, Torpey and colleagues (Torpey et al., 2013) found that erasure of anxiety symptoms from early adolescence into adulthood (Fitzgerald and Taylor, 2015), or hyperactivity to the importance of mistakes (Weinberg et al., 2016a). The ERN has gained attention as an RDoC biomarker for anxiety disorders given that clinically-affected adolescents and adults, particularly those who experience high levels of anxious apprehension (i.e., worry), exhibit an enlarged ERN compared to healthy individuals (Vaidyanathan et al., 2012; Cavanagh and Shackman, 2015; Moser et al., 2013). Indeed, consistent with the RDoC framework (Cuthbert, 2014), the ERN tracks anxiety symptoms across the normal to abnormal range in community samples, with larger ERN relating to more anxiety symptoms in early childhood into adulthood (Bress et al., 2015; Moabad et al., 2010; Moran et al., 2015; Olvet and Hajcak, 2008). While increased ERN amplitude has been most consistently associated with anxiety (Carrasco et al., 2013; Hajcak et al., 2008; Ladouceur et al., 2006; Meyer et al., 2013; Santesso et al., 2006), some studies have also found increased ERN in relation to depression (Chiu and Deldin, 2007; Holmes and Pizzagalli, 2010), whereas others show decreased ERN in depression (Weinberg et al., 2016a; Ladouceur et al., 2012; Weinberg et al., 2012a, b) or no relation between ERN and depressive symptoms (Bress et al., 2015).

1.2. ERN, anxiety and development

A gradual increase in ERN magnitude with age has been well-documented (Tamnes et al., 2013), but the influence of development on ERN-anxiety associations is less understood. When studied dimensionally in a community sample, greater anxiety was related to a larger ERN in older children (11–13 years), but to a marginally (i.e., trend-level significant) smaller ERN in younger children (6–10 years) (Meyer et al., 2012), suggesting a reversal of the adult-like associations between anxiety symptoms and ERN in younger children. Consistent with this suggestion, higher levels of separation anxiety were related to a smaller ERN in a community sample of children ages 5–7 years (Lo et al., 2016).

Moreover, Torpey and colleagues (Torpey et al., 2013) found that fearful temperament at aged 3 was associated with a smaller ERN among 6-year-old children, but with a larger ERN when children were re-assessed at aged 9 (Meyer et al., 2018). However, when anxiety was examined categorically, six-year-olds who met criteria for clinically significant anxiety demonstrated larger ERN compared to healthy children, consistent with the pattern of group differences observed when older patients are compared to healthy controls (Meyer et al., 2012). Collectively, these findings raise the possibility that ERN-anxiety associations may shift with age and symptom severity in childhood (Meyer, 2017; Moser, 2017; Meyer et al., 2018), and underscore the importance of examining the relationship between ERN and anxiety symptoms in young children across the full spectrum of severity.

1.3. Gender effects on the ERN and early expression of anxiety problems

Converging lines of evidence suggest that characterization of the ERN as a developmentally sensitive biomarker of anxiety symptoms should consider interactive effects of age and gender. Not only is there evidence to suggest earlier maturation of the ERN in girls than boys (Davies et al., 2004) but anxiety problems are more common in females than males across the lifespan. It is well-documented that fear and anxiety affect more girls than boys beginning in childhood (Craske, 2003; Ollendick et al., 2002), and foreshadow higher rates of anxiety and depression in females compared to males from adolescence (Costello et al., 2005; Fine et al., 1998) into adulthood (Beesdo et al., 2009). The higher frequency of internalizing problems in females than males may be influenced by gender differences in neural circuitry (Bangasser and Valentino, 2014). While it remains unknown whether the ERN indexes a process that is responsible for higher rates of anxiety disorders in females than males, recent meta-analytic evidence suggests that the association of larger ERN with anxiety in adulthood is characteristic of women, not men (Moser et al., 2016; Moran et al., 2012). Gender effects on the relation between ERN and anxiety symptoms have yet to be examined in children as a function of age.

1.4. The current study

The ERN has been previously posited as an RDoC-relevant biomarker of anxiety symptoms (Vaidyanathan et al., 2012; Weinberg et al., 2015); but, despite the common emergence of anxiety in early to middle childhood (Beesdo et al., 2009) and developmental change in ERN magnitude during this period (Tamnes et al., 2013), the association between ERN and anxiety symptoms in young children is not well-characterized. Better understanding how ERN-anxiety associations shift with age, gender and symptom severity will be important for refining the ERN as an early biomarker of anxiety risk and potential treatment target. Thus, we sought to examine age and gender effects on the relationship between ERN and anxiety symptoms in children, ages four to nine, sampled across the non-clinical to clinical range of severity. Prior work has suggested a reversal of the adult pattern of ERN-anxiety associations at approximately age 10 years (Meyer et al., 2012), however, neural networks for cognitive control (Tamnes et al., 2013) and behavioral capacity for this function (Diamond, 2013) undergo dramatic development between early childhood and preadolescence. Thus, we tested for an age-related reversal in the association between ERN and anxiety between ages 4–9 years, remaining agnostic to precisely when this reversal might occur. In addition, based on findings from an adult meta-analysis (Moser et al., 2016), we hypothesized that the relationship of ERN with anxiety might be further moderated by gender such that ERN-anxiety associations would be stronger in girls than boys.

2. Method

2.1. Participants

Participants included 56 children (30 girls) sampled from the community and the University of Michigan Child and Adolescent Psychiatry Clinic to capture the full spectrum of anxiety symptoms severity. Participants were 6.87 years old on average (SD = 1.39, range = 4.1–9.7); 77% Caucasians, 5% African American, 16% bi-racial and 2% “Other”. To be eligible for participation, children had to be between 4–9 years old and to have no history of head injury, serious medical illness, neurodevelopmental delay (autism spectrum disorder or mental retardation) and not taking medications that affect central nervous system functioning.

After data cleaning, the final sample consisted of 49 participants (26 girls; mean = 6.99 +/- 1.32 years, range 4.1–9.7). Six children (mean = 5.44 +/- .76 years, 4 girls) made fewer than six errors on the ERN-eliciting task and were excluded from further analysis based on standard convention (Olvet and Hajcak, 2009). One additional child, whose ERN amplitude was more than three standard deviations from the mean, was excluded (age = 9.42; male). No differences with respect to age, gender and anxiety symptoms were found between children who were in the final sample compared to those excluded (all ps > .05). The age distribution was comparable across gender (girls: mean age in years 6.63 +/- 1.11.; boys: mean age in years 7.31 +/- 1.43, t(47) = 1.82, p = n.s.).

Based on the effect size (partial r = .28) observed previously in a community sample of 55 children, 8- to 13-years old (Meyer et al., 2012), power analysis conducted in G*Power suggested a sample size of 75 is needed for detecting interactive effects of age x ERN on anxiety in that age group (with a power of .80). However, no prior developmental
studies have investigated the relation between ERN and age on anxiety in younger, preschool- to school-aged children or the moderating role of gender, thus precluding direct comparison with the present study. Moreover, inclusion of anxious children recruited from a Child and Adolescent Psychiatry Clinic enabled sampling into the clinical range of anxiety severity (i.e., increased variance), distinguishing the sample presented here from prior work. Thus, analyses were conducted to test a priori hypotheses involving both age and gender effects on ERN-anxiety associations to generate preliminary results in the unique sample collected here.

2.2. Task

Participants performed the child-friendly Go/No-go “Zoo” task (Grammer et al., 2014). In the Zoo task, children were asked to help a zookeeper return loose animals to their cages, except three friendly orangutans who are the zookeeper’s “helpers” and should remain free. Children were asked to put the loose animals back in their cages by pressing a button as quickly as they could every time an animal picture was presented (Go Trials), but to withhold their response each time they saw an orangutan (No-Go trials). No-Go trials on the Zoo Task have been previously shown to produce error rates that are sufficient to elicit the ERN (Grammer et al., 2014).

Children completed 8 blocks of the task, each including 30 Go trials and 10 No-Go trials for a total of 320 trials. For each trial, a fixation cross was presented for 200–300 milliseconds (ms), followed by an animal image presented for 750 ms, and a blank screen for 500 ms. Responses could be made during the animal image and blank screen presentation. Each block consisted of novel sets of animal images, balanced on color, animal type, and size. The task was presented using Eprime software (Psychology Software Tools, Inc.: Pittsburgh, PA). Before the experimental trials of the Zoo task, children practiced on a set of 12 trials, 3 with orangutans and 9 with other animals and could practice multiple times until they understood the task.

2.3. Procedure

The study was approved by the University of Michigan Medical School Institutional Review Board. Initially, phone screening was conducted to determine that the child met study inclusion criteria. After written informed consent and oral assent were obtained from parents and children, respectively, children were brought to a child-friendly EEG booth by experimenters while parents filled out questionnaires. EEG experiments were conducted using the BioSemi ActiveTwo recording system (see below). Children were seated on a comfortable chair in front of a computer screen while watching cartoons during experimental set-up. To reduce fidgeting and increase compliance during ERP recording, children were given brief breaks between blocks and animal stickers as tokens for every block they completed. Verbal and visual feedback (in the form of a zoo map) were provided between blocks to remind children to stay still during blocks and monitor their progress through the zoo. Families received monetary incentives and children received toys for their participation.

2.4. Electrophysiological recording, data reduction and analysis

The EEG was recorded from 34 Ag/AgCl scalp electrodes and two mastoid electrodes, using BioSemi ActiveTwo recording system. Electro-oculogram (EOG) data were recorded from electrodes placed above and below the right eye and at the outer canthi of both eyes to capture vertical EOG and horizontal EOG, respectively. Data were referenced to a ground formed from a common mode sense active electrode and driven right leg passive electrode (see http://www.biosemi.com/faq/cms&drl.), and sampled at 1024 Hz. For analysis, EEG data were referenced to averaged mastoid electrodes, and band-pass filtered 0.05–30 Hz using zero-phase shift butterworth filters. EEG data were screened using automated algorithms that rejected epochs in which the absolute voltage range exceeded 500 μV for midline channels (Fz, FCz, Cz, and Pz), consistent with prior work (Grammer et al. (2014)). Ocular movement artifacts were then corrected using a regression-based algorithm (Gratton et al., 1983). After ocular correction, individual trials were rejected if any amplitudes were greater than 100 μV, differed by more than 50 μV from the previous time point, or were less than 0.5 μV in magnitude in any midline electrode.

2.4.1. Behavioral measures

Correct trials included correct response to Go trials (button press when viewing any animal that was not an orangutan) and correct inhibition of response to No-Go trials (withholding button press to orangutan stimuli). Only the number of correct Go trials were evaluated. Errors were evaluated only for No-go trials, defined as errors of commission when children incorrectly responded to an orangutan (Grammer et al., 2014). Response times were evaluated for correct Go trials.

2.4.2. ERP measures

Response-locked ERP components were quantified using mean amplitude measurements relative to a pre-response baseline -200 to −100 ms, consistent with prior work in young children (Grammer et al., 2014). The mean amplitude of the ERN was computed for commission errors in a window 0–50 ms after the incorrect button response on No-Go trials (Grammer et al., 2014). ERN was measured at Fz (mean amplitude: -5.0 ± 5.1), FCz (mean amplitude: -5.4 ± 4.7) and Cz (mean amplitude: -3.6 ± 5.1). Overall amplitude at each of these locations was more negative on error relative to correct trials measured in the same time window (i.e., ERN effect, p < 0.001). As with prior work in this age group (Grammer et al., 2014), ERN at FCz (Fig. 1) had the highest mean amplitude and increased with age (Tannnes et al., 2013); thus, ERN measured at FCz was used in all subsequent analyses. When participants with higher numbers of errors were considered (e.g., > 16), the split-half reliability of the ERN increased and was comparable with prior studies (e.g., Schroder et al., 2017). Importantly, main findings remained significant in secondary analysis of subjects who made more errors (e.g., > 16) and therefore we report results with

Fig. 1. Panel A: ERN and CRN waveforms at FCz electrode. Panel B: ERN amplitude increased (more negative) with age.
the full sample (> 6 errors; N = 49) and provide sub-sample analyses (> 16 errors; N = 33) in Supplementary Table 2.

2.4.3. Child anxiety symptoms

Parent report, using the Child Behavior Checklist (CBCL/1-5 years and CBCL/6-18 years) (Achenbach and Rescorla, 2001), is commonly used to measure psychopathology in young children who struggle to provide accurate self-report (Tandon et al., 2009). Thus, given the inclusion of the children as young as 4 years and the lack of validated self-report anxiety measures for children at this age (Birmaher et al., 1997; March et al., 1997), child anxiety was measured by parent report on the CBCL DSM-oriented Anxiety Problems subscale (CBCL-AP; α = .79) (Achenbach et al., 2003) combined with the CBCL Somatic Problems (α = .64) subscale (Kendall et al., 2007) by averaging the T-scores from the two subscales. The composite of these scales was used because young children often express anxiety as somatic complaints, and combination of the Anxiety and Somatic Problems subscales of the CBCL has been found to better capture anxiety severity than the CBCL-AP subscale alone (Kendall et al., 2007). Moreover, the correlation between the two subscales were moderately high (r = .48, p = .001) and reliability showed that the combined scale (with all the anxiety and somatic subscale items together) showed adequate reliability (α = .78). Of the 49 children who provided ERN data, CBCL data was missing for one (7.3-year-old boy), but was imputed using the Expectation-Maximization (EM) algorithm (Dong and Peng, 2013); therefore, the final sample remained 49 children.

2.4.4. Child depressive symptoms and attention problems

Parent report on the DSM-oriented Depression Problems (α = .64) and Attention Problems (α = .76) subscales from the CBCL were also examined given prior work suggesting the ERN may be differentially modulated by these phenomena (Weinberg et al., 2016a; Ladouceur et al., 2012; Weinberg et al., 2012a; Albrecht et al., 2008; Weinberg et al., 2012b).

2.5. Data analysis plan

All predictor variables were mean centered prior to the analyses. Pearson correlation was first conducted to examine inter-correlation among all study variables (Table 1). Hayes’ (2013) PROCESS (Model 3) was used to test for conditional effects of age and gender as moderators of the relation between ERN and child anxiety symptom severity. Specifically, the PROCESS model considered all two-way (Age X ERN, Gender X ERN, Age X Gender) and the three-way (Age X Gender X ERN) interactions as predictors of child anxiety symptoms. The main effect of age was treated as a continuous variable in the model.

Two additional PROCESS models (Model 3) were also conducted to test for conditional effects of age and gender on associations of ERN with child depressive and attention problems. To aid interpretation of results, linear regression was used to test for effects of age, gender and the interaction on behavioral measures; if not significant, the interaction term was dropped from the model. For comparison with prior work, effects of age, gender and performance on ERN were also assessed. Unstandardized beta coefficients were reported in all PROCESS and regression models.

3. Results

3.1. Behavioral

Participants committed an average of 23.98 (SD = 11.40; range = 6–63) commission errors on 80 total No-Go trials, and responded correctly to 231.59 (SD = 8.94; range = 194–240) of 240 total Go trials. On No-go trials, the number of commission errors were fewer in girls than boys (M = 18.6; SD = 6.1 for girls, M = 30; SD = 13 for boys; B = -11.19, p = .000), but did not vary by age (B = -.36, p = .75). On Go trials, a greater number of correct trials (B = 2.88, p = .003) and faster response times (B = -.29, p = .000) occurred with older age, but there was no effect of gender on either measure (number of correct Go trials: B = 1.47, p = .55; response times: B = 21.25, p = .20). There were no age X gender interactions on any measure of behavioral performance.

3.2. Age, gender and performance effects on ERN

Consistent with prior work (Tamnes et al., 2013), older age associated with larger (i.e. more negative) ERN amplitude (B = -1.17, p = .03, total R² = .14; Fig. 1), controlling for the effects of gender and performance. There were no significant gender or age X gender interaction effects on ERN amplitude (ps > 0.05). Performance (i.e., number of No Go commission errors, number of correct Go trials and Go response time) did not associate with ERN amplitude (Table 1) and there were no age X performance interaction effects on ERN amplitude (ps > 0.05).

3.3. Differential relation of ERN with anxiety by age and gender

As shown in Table 2, there were no significant main effects of age, gender, ERN or two-way interaction terms as predictors of child anxiety symptoms (all p > .05). However, there was a significant 3-way interaction effect of Age X Gender X ERN on child anxiety symptoms (B = -.91, p = .002) that explained 13% of the variance (F (1, 41) = 11.19, p = .002; effect size: partial r = .35; observed power = .92) in the model (total R² = .24; F (7, 41) = 4.85, p = .0005).

To explore the 3-way interaction on child anxiety, Johnson-Neyman (J-N) analyses were first conducted using PROCESS (Model 1) to characterize the conditional effects of age (in years) on the relationship of ERN with anxiety, separately for each gender. The J-N analysis for girls revealed a significant shift in the directionality of the ERN effect on anxiety between 7.2 and 8.6 years (Fig. 2). There was no J-N significance region (or significance transition points) across the study age range (all ps > .05) in boys. Next, based on the J-N defined age split, post-hoc simple slope analyses within PROCESS were conducted. Partial residuals from this analysis were plotted using the Visreg package in

Table 1 Descriptive statistic and Pearson correlation among all study variables.

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>SD</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Age</td>
<td>6.99</td>
<td>1.32</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Correct Go Trials</td>
<td>231.59</td>
<td>8.94</td>
<td>.45</td>
<td>**</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Correct Go RT</td>
<td>544</td>
<td>65.9</td>
<td>-.55</td>
<td>**</td>
<td>-.45</td>
<td>***</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. NoGo Errors</td>
<td>23.98</td>
<td>11.4</td>
<td>-.17</td>
<td></td>
<td>-.32</td>
<td>*</td>
<td>-.29</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. ERN</td>
<td>-5.40</td>
<td>4.70</td>
<td>-.34</td>
<td>*</td>
<td>-.12</td>
<td>.21</td>
<td>.19</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Anxiety</td>
<td>53.63</td>
<td>5.21</td>
<td>-.18</td>
<td>.08</td>
<td>-.09</td>
<td>.09</td>
<td>.00</td>
<td>-.02</td>
<td></td>
</tr>
<tr>
<td>7. Depression</td>
<td>53.37</td>
<td>5.19</td>
<td>-.03</td>
<td></td>
<td>-.03</td>
<td>-.03</td>
<td>.11</td>
<td>.67</td>
<td>***</td>
</tr>
<tr>
<td>8. Attention Prob.</td>
<td>54.47</td>
<td>5.79</td>
<td>-.09</td>
<td>-.28</td>
<td>.15</td>
<td>-.05</td>
<td>.09</td>
<td>.38</td>
<td>*</td>
</tr>
</tbody>
</table>

Note. RT is reaction time in millisecond. #No Go Errors is the number commission errors on No Go trials. *p < .05, **p < .01, ***p < .001.
Table 2
Regression predicting child’s anxiety symptoms.

<table>
<thead>
<tr>
<th>Variables</th>
<th>B</th>
<th>SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>−1.04</td>
<td>.68</td>
</tr>
<tr>
<td>Gender</td>
<td>−2.97</td>
<td>1.68</td>
</tr>
<tr>
<td>ERN</td>
<td>.08</td>
<td>.17</td>
</tr>
<tr>
<td>Age X Gender</td>
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<td>1.44</td>
</tr>
<tr>
<td>Gender X ERN</td>
<td>.59</td>
<td>.36</td>
</tr>
<tr>
<td>Age X ERN</td>
<td>.05</td>
<td>.13</td>
</tr>
<tr>
<td>Age X Gender X ERN</td>
<td>−.91</td>
<td>**</td>
</tr>
<tr>
<td>$R^2$</td>
<td>.24</td>
<td></td>
</tr>
<tr>
<td>$F$</td>
<td>4.85</td>
<td>***</td>
</tr>
</tbody>
</table>

Note. *p < .05, **p < .01, ***p < .001. Male gender is the reference group.

R to visualize how JN-defined age groups and gender moderate the relation of ERN and anxiety symptoms (Fig. 3). As shown in Fig. 3, smaller (i.e., less negative) ERN associated with more anxiety symptoms in younger girls (effect = .90, SE = .30, p = .004) whereas, in older girls, larger (i.e., more negative) ERN associated with more anxiety symptoms (effect = −.16, SE = .07, p = .03). In boys, ERN-anxiety associations were not significantly moderated by age (younger: effect = −.53, se = .42, p = .21; older: effect = −.13, se = .42, p = .74). Including the number of No go errors in the PROCESS models did not change primary results, and did not reveal any main effect of error rates. For comparison, Supplementary Fig. 1 plots raw data showing the ERN plotted against anxiety for each gender and age group.

Finally, there were no significant associations between any behavioral measures (number of correct trials or reaction time) and anxiety symptoms, and no two- or three-way interaction effects of any behavioral measure with age and/or gender on anxiety.

3.4. Relation of ERN with depression and attention problems by age and gender

No significant model was found for predictors of child depressive symptoms ($F$ (1, 41) = 1.11, $p = .37$) and child attention problems ($F$ (1, 41) = 1.29, $p = .28$).

4. Discussion

As operationalized in the RDoC framework (Cuthbert, 2014), characterizing the neural circuits that relate to psychopathology should yield synergistic advances in the understanding and treatment of psychiatric illness, including anxiety (Pine, 2007). However, to realize this promise, converging lines of evidence suggest that effects of age and gender on neural circuits must be considered (Casey et al., 2014; Woody and Gibb, 2015). Towards this end, the current study examined whether age and/or gender moderated the association between the ERN and anxiety in girls and boys, ages 4–9 years, when anxiety often first presents (Beesdo et al., 2009). Results support prior work implicating ERN as a neural correlate of anxiety in children (Meyer et al., 2013, 2012; Lo et al., 2016) and substantiate the possibility of an age-related reversal in the association of ERN and anxiety (Meyer, 2017; Moser, 2017). Specifically, the adult-like pattern of larger ERN with greater anxiety severity (Moser et al., 2013; Olvet and Hajcak, 2008; Weinberg et al., 2015; Endrass and Ullsperger, 2014; Weinberg et al., 2016; Xiao et al., 2011) was observed in older girls, but reversed in younger girls in our sample. In contrast, there were no significant associations between the ERN and anxiety in younger or older boys, consistent with recent work showing ERN-anxiety association in women, but not men (Moser et al., 2016). These results suggest that the ERN-anxiety link is not only developmentally sensitive, but may also be gender specific. Thus, our findings highlight the importance of considering both age and gender differences known to characterize psychopathology when examining neural markers of RDoC constructs (Casey et al., 2014; Woody and Gibb, 2015).
4.1. Cognitive control theory of ERN and anxiety

Cognitive control functions begin to develop in early childhood (Diamond, 2013). Moreover, age-related change in cortical networks has been found to contribute to increased neural capacity for cognitive control (Bunge and Wright, 2007; Casey et al., 2005; Menon, 2013; Posner et al., 2014). Accumulating evidence shows that, while neural networks that support cognitive control continue to mature from childhood into adolescence and early adulthood (Luna et al., 2015), key components of these networks (e.g., ACC), are already involved in the implementation of control functions in preschool-aged children (Petrican et al., 2017). A widely accepted index of cognitive control, the ERN, occurs in response to errors, localizes to the ACC, and can be elicited as early as 3 years of age (Tamnes et al., 2013; Grammer et al., 2014; Ferdinand and Kray, 2014). Consistent with maturational trajectories for cognitive control, prior studies have found developmental increase in ERN amplitude from childhood (Torpey et al., 2012) and throughout adolescence (Tamnes et al., 2013; Davies et al., 2004; Lo, 2018). Although we did not find a relationship between larger ERN and better performance (nor age moderation of this relationship) in the current study, likely due to insufficient power, prior work has found that age-related increases of ERN magnitude associates with faster RTs and higher accuracy (Torpey et al., 2012). Taken together, this prior work suggests that the age-related increase in ERN amplitude may index the maturation of neural substrate for cognitive control, especially in performance monitoring of errors (Tamnes et al., 2013; Ferdinand and Kray, 2014 (Torpey et al., 2012).

Cognitive control may play a key role in facilitating the inhibition and/or regulation of negative/threatening thoughts (i.e., rumination and worry) (Derryberry and Rothbart, 1997; Eisenberg et al., 2009), and behavioral adaptation to reduce anxiety problems (Ip et al., 2019; Lemery-Chalfant et al., 2008; Lengua, 2003; Rigg et al., 2004). The directional shift of the ERN-anxiety relationship from younger to older girls may therefore mark the developmental transition of increasing neural capacity of using ERN to signal the need for cognitive control (Moser, 2017), such that an enlarged ERN allows greater recruitment of cognitive control (Gehring et al., 1993; Debener et al., 2005; West and Travers, 2007). In theory, low levels of ERN-indexed cognitive control in younger children may leave early anxiety symptoms unchecked, whereas the reversal of this relationship at older ages may reflect a compensatory process by which increasing neural capacity for cognitive control is leveraged to maintain adequate performance on task (Moser, 2017) and/or reduce anxiety symptom severity (Fitzgerald and Taylor, 2015).

Notably, we found a significant relationship between ERN-anxiety problems in girls but not in boys. Our gender specific finding is consistent with a meta-analysis in adults showing the presence of ERN-anxiety problems in women, but not men (Moser et al., 2016). In epidemiologic work, anxiety disorders have been demonstrated to be more common in girls than boys from childhood, throughout adolescence and into adulthood (Ollendick et al., 2002; Costello et al., 2005; Beesdo et al., 2009). Indeed, it has been suggested that biological mechanisms for processing threat and anxiety-inducing stimuli may be more sensitive in girls than boys from the earliest stages of development (Lebrun-Millard et al., 2012; Ruigrok et al., 2014). With a greater biological sensitivity to threatening stimuli, girls may have to recruit greater cognitive control (as indexed through ERN) to maintain task goals and/or reduce anxiety than boys. Therefore, the association of ERN with greater anxiety severity in girls (but not boys) may suggest that girls are more dependent on ERN-indexed ACC network to suppress sensitivity to and/or interference from anxiety.

On experimental tasks requiring cognitive control, girls exhibit better performance than boys (e.g., higher accuracy, fewer commission errors), consistent with the greater cognitive control capabilities that have been previously demonstrated in preschool-aged girls compared to boys (Else-Quest et al., 2006). It is possible that ERN-indexed systems for cognitive control are more available to manage behavior, including behaviors related to fear and anxiety, in females than males over the course of development, which may be related to gender differences in the functional maturation of ACC (Christakou et al., 2009; Liu et al., 2012). Further research, across a broader age range, is needed to understand whether the maturation of ACC influences the relationship between the ERN and anxiety symptoms and whether this relationship differs by gender.

4.2. Developmental transition of anxiety phenomenon and ERN

We hypothesize that the developmental shift in ERN-anxiety relationship in girls may reflect a developmental change in neural capacity for cognitive control, however, other interpretations are possible. A larger ERN has been found to have a stronger association with worry/anxious apprehension than other forms (e.g., fear) of anxiety-related symptoms in meta-analysis (Moser et al., 2013). Some have argued that with the natural transition of anxiety phenomena from more fear-based disorders at younger ages (e.g., phobias) to worry-related disorders at older ages (e.g. generalized anxiety), a smaller ERN in younger, more anxious children may reflect sensitivity to acute, external threat (i.e., fear), whereas a larger ERN in older, anxious children may relate to the greater relevance of internal threat (i.e., worry (Meyer, 2017; Weinberg et al., 2016a)). Thus, the shift of the ERN-anxiety relationship across ages may reflect the changing nature of anxiety symptom phenomenology with age (Meyer, 2017; Meyer et al., 2018).

Finally, it is important to note that although we do not find a significant ERN-anxiety link in boys, a non-significant pattern of larger ERN with greater anxiety severity was observed in younger boys (b = - .53, p = .21, see Fig. 3). This pattern is consistent with previous work showing a larger ERN in clinically anxious compared to healthy 6-year-old children in a sample comprised of more boys (2/3) than girls (1/3) (Meyer et al., 2013). It is possible that these previously reported findings were male-driven. Alternatively, ERN-anxiety association may also be influenced by anxiety severity (i.e., clinical vs subclinical) and further study in larger samples will be needed to understand whether age and gender effects on ERN-anxiety association shift with anxiety severity.

4.3. Limitations

Our findings should be viewed with caution given our small sample size. Power analysis based on the effect size (partial r = .28) observed in prior work suggests that a sample size of 75 (power = .80) is needed to detect the interactive effect of age and ERN on anxiety in 8 - 13-year-old children. However, no prior developmental studies have investigated moderating effects of age and gender on ERN-anxiety relationships among younger children, precluding direct comparison. In fact, in the current study, we observed a larger effect size (partial r = .35; observed power = .92) than previously reported (Meyer et al., 2012), raising the possibility that a smaller sample may be appropriate when considering the interactive effects of age and gender on ERN-anxiety associations in preschool- to school-aged children (ages 4–9 years). On the other hand, small samples can produce statistically significant results that do not reflect a true effect due to outliers (Button et al., 2013). Thus, even though we did not observe any outlier(s) driving the findings presented here, it is important to view our results as preliminary, requiring confirmation with a larger sample. Nonetheless, our results should guide future research to consider both age and gender when examining the ERN as a neurophysiological marker of anxiety in children.

Several other limitations deserve consideration. First, our study used a cross-sectional design and therefore is unable to infer longitudinal relationship or causality of ERN and anxiety symptoms. Second, our study relied on parent-report of anxiety symptoms as there is no validated self-report measure that reliably assesses anxiety...
symptoms in preschool-aged children. Third, we did not include a cognitive control measure outside of performance on the Go No Go task used to elicit the ERN. Further research that includes other behavioral measures of cognitive control is needed to more fully understand the relations among ERN, cognitive control and anxiety in children and to test the prediction that a smaller ERN may reflect less capacity for cognitive control in more anxious, younger girls (i.e., a 4-way interaction).

4.4. Conclusion

Our findings represent an important first step towards clarifying the relation of the ERN and anxiety symptoms in children. If, as suggested by RDoC, indices of error-processing (e.g., ERN) in children with subclinical anxiety symptoms fall on a continuum between clinically affected children at one end and healthy children at the other, this information would justify future testing of strategies to shift ERN along the continuum to reduce subclinical symptoms and prevent the progression to illness. Specifically, cognitive control training or other interventions designed to increase ERN might help to reduce anxiety symptoms among preschool-aged girls. By contrast, given the opposite pattern of the ERN-anxiety relationship in school-aged girls, strategies to decrease ERN may hold more promise for anxiety symptom reduction (Schrod et al., 2018). On the other hand, if greater ERN is an adaptive response to reduce anxiety, then interventions should be designed to increase ERN across early to late childhood and across genders. Future longitudinal work is needed to understand how changes in ERN associate with changes in anxiety symptoms over time and should consider age and gender to identify whether the ERN should be targeted differently in children to reduce and prevent anxiety problems.

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