Testosterone during Puberty Shifts Emotional Control from Pulvinar to Anterior Prefrontal Cortex

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Increased limbic and striatal activation in adolescence has been attributed to a relative delay in the maturation of prefrontal areas, resulting in the increase of impulsive reward-seeking behaviors that are often observed during puberty. However, it remains unclear whether and how this general developmental pattern applies to the control of social emotional actions, a fundamental adult skill refined during adolescence. This domain of control pertains to decisions involving emotional responses. When faced with a social emotional challenge (e.g., an angry face), we can follow automatic response tendencies and avoid the challenge or exert control over those tendencies by selecting an alternative action. Using an fMRI-adapted social approach-avoidance task, this study identifies how the neural regulation of emotional action control changes as a function of human pubertal development in 14-year-old adolescents (n = 47). Pubertal maturation, indexed by testosterone levels, shifted neural regulation of emotional actions from the pulvinar nucleus of the thalamus and the amygdala to the anterior prefrontal cortex (aPFC). Adolescents with more advanced pubertal maturation showed greater aPFC activity when controlling their emotional action tendencies, reproducing the same pattern consistently observed in adults. In contrast, adolescents of the same age, but with less advanced pubertal maturation, showed greater pulvinar and amygdala activity when exerting similarly effective emotional control. These findings qualify how, in the domain of social emotional actions, executive control shifts from subcortical to prefrontal structures during pubertal development. The pulvinar and the amygdala are suggested as the ontogenetic precursors of the mature control system centered on the anterior prefrontal cortex.

Key words: adolescence; approach-avoidance task; fMRI; frontal pole; hormones; thalamus

Introduction

Adolescence is often characterized by reward- and sensation-seeking behaviors (Steinberg et al., 2008; Harden and Tucker-Drob, 2011). The increased occurrence of those behaviors during puberty has been linked to a relative imbalance in the maturation of prefrontal areas compared with already well developed striatal structures, resulting in reduced goal-directed control during decision-making processes (Braams et al., 2015; Defoe et al., 2015). However, it remains unclear whether developmental imbalances in the same frontostriatal circuits are also responsible for the altered control of emotional action tendencies observed during puberty (Ernst and Fudge, 2009). This study addresses the largely unexplored issue of how emotional action control is neurally implemented during puberty.
We know that, in adults, the lateral anterior prefrontal cortex (aPFC; also known as lateral frontal pole; Neubert et al., 2014) plays a crucial role in emotional action control by downregulating emotional action tendencies in the amygdala (Volman et al., 2011a, 2013). However, the aPFC and its connections to limbic structures develop relatively late in puberty (Tamnes et al., 2010). During that phase, emotional control might be coordinated by a number of subcortical structures showing faster maturation and strong responses to emotional processing. In addition to well known contributions from the amygdala and the striatum (Pfeifer et al., 2011; Scherf et al., 2013), recent work has shown that the dorsomedial pulvinar nucleus of the thalamus is necessary for emotional processing (Ward et al., 2007). Namely, by virtue of the colliculo-pulvino-amygdalar pathway (Morris et al., 1999; Tamietto et al., 2012) and its extensive connectivity with frontoparietal areas (Arcaro et al., 2015), the dorsomedial pulvinar is well placed for providing an alternative pathway to the frontal-amygdalar circuit controlling emotional action tendencies in adults (Arend et al., 2015; Barron et al., 2015). The neurophysiological properties of the pulvinar support this possibility, including rapid neuronal responses to visual facial features (Nguyen et al., 2013a), and a coordinating role across frontal, parietal, and temporal areas during movement selection (Wilke et al., 2010; Saalmann et al., 2012). However, it remains unclear whether and how the pulvinar is involved in emotional processing when the aPFC is not yet sufficiently mature to implement flexible goal-directed control of emotional action tendencies.

Here we address this issue by exploiting a task that has repeatedly been shown to robustly capture behavioral and cerebral correlates of emotional control (Rinck and Becker, 2007; Roelofs et al., 2009a; Volman et al., 2011a,b, 2013). The approach-avoidance (AA) task requires participants to evaluate the emotional expression (happy, angry) of faces and to respond by either pulling a joystick toward (approach) or away (avoidance) from themselves. Affect-congruent conditions involve automatic stimulus–response mappings (i.e., approach—happy and avoid—angry faces). In contrast, affect-incongruent conditions require participants to apply emotional control, namely, to override those emotional action tendencies and select an alternative action to meet task demands (approach—angry and avoid—happy faces) (Fig. 1). We use this task in a group of 14-year-old adolescents. This age marks a pubertal stage when sexual maturation is completing in both sexes, but behavioral, emotional, and cerebral development can still widely differ among individuals (Giedd et al., 1999; Crane et al., 2008; Monahan and Steinberg, 2011). Accordingly, we quantify pubertal development with an endogenous physiological marker, salivary testosterone, that is sensitive to those interindividual differences (Giedd et al., 2006; Forbes and Dahl, 2010; Berenbaum and Beltz, 2011) and is mechanistically involved in mediating neural development (Sisk and Zehr, 2005; Nguyen et al., 2013b; Herlting et al., 2014). We expect that relatively less mature 14-year-old adolescents control their emotional action tendencies by relying on a pulvino-amygdalar system. In contrast, developmentally more advanced 14-year-old adolescents might be able to access the prefrontal-amygdalar control system used by adults (Volman et al., 2011a).

Materials and Methods

Participants. Forty-nine 14-year-old right-handed adolescents (Table 1) participated in the study. All participants had normal or corrected-to-normal vision, no history of psychiatric disorders or neurological illness (as indicated by parent/guardian report). Participants were recruited from the Nijmegen Longitudinal Study on Infant and Child Development. Two participants were excluded due to technical problems (poor MRI image quality), resulting in 47 participants (21 males) who were included in the final analyses. Written informed consent was obtained from parents as well as from participants. The study was approved by the local ethics committee (CMO region Arnhem-Nijmegen, The Netherlands).

Experimental task. During the AA task, participants were presented with emotional faces, which they had to evaluate based on their affective expressions (happy, angry). They responded using a joystick, pushing it toward themselves (approach) or away from themselves (avoid). After making a response, participants had to return the joystick to its starting position (defined as the central area covering 15% along the sagittal plane) before the end of the intertrial interval (ITI; 2–4 s). If this did not happen, participants received visual feedback stating “return the joystick to the starting position”; the ITI was repeated after the joystick was returned to its correct position. Responses were considered to be valid when the joystick was displaced at least 65% along the sagittal plane and were delivered within 3 s following stimulus presentation. Invalid responses were signaled for 1 s with visual feedback indicating “you did not move your joystick far enough.”

The task consisted of 16 blocks, 12 trials each. After each block, there was a baseline period (21–24 s). There were two block types/response mappings (affect congruent, affect incongruent) and four affect × response combinations, namely, happy–approach, angry–avoid (affect congruent) and happy–avoid, angry–approach (affect incongruent). At the start of each block, the participant was instructed on the required response mapping. The block type of the first block was counterbalanced across participants, with the sequential blocks always switching between the two block types. Affective expressions and gender types (of faces) were pseudorandomly and evenly distributed within each block, with no more than three consecutive presentations of each. During the training...
part of the task, four blocks were presented with eight trials each. Each block consisted of the same affect × response combinations. The training phase did not contain the same visual stimuli as the experimental task.

Materials and apparatus. The fMRI data were acquired on a Siemens 3 tesla MAGNETOM Trio MRI scanner (Siemens Medical Solutions) using a 32-channel coil. The acquisition of the functional scans was performed with a multicathechoplanar imaging (EPI) sequence (TR = 2190 ms; TE = 93.20.9.32, and 44 ms; flip angle = 90°; 34 transversal slices; 3.3 × 3.3 × 3.0 mm voxels; FOV = 212 mm). This type of parallel imaging technique for functional images allows a significant reduction in the echo train length, which reduces motion artifacts and image distortion. It improves BOLD sensitivity, especially in brain regions that typically would be compromised by the use of a single short TE. Finally, the reduced distortion allows better coregistration of functional and anatomical data (Poser et al., 2006). Structural T1 images were acquired using an MPRAGE sequence (TR = 2300 ms; TE = 3.03 ms; 192 sagittal slices; 1.0 × 1.0 × 1.0 mm voxels; FOV = 256 mm).

An MR-compatible joystick (Fiber Optic Joystick, Current Designs), with a sampling rate of ~550 Hz, was placed on the abdomen of the participants to ensure comfortable push and pull movements. The visual stimuli consisted of faces from 36 models (18 male) taken from several databases (Ekman and Friesen, 1976; Matsumoto and Ekman, 1988; Lundqvist et al., 1998; Martinez and Benavente, 1998). Each model showed two affective expressions (happy, angry). The pictures were in grayscale, matched for brightness and contrast values, and displayed against a black background. To exclude influence from hair and nonfacial contours, the faces were trimmed (Roelofs et al., 2009a). The stimuli were presented at the center of a screen and viewed via a mirror above the participants head, with a visual angle of 4° × 6° (width × height). Stimuli presentation and acquisition of joystick positions were controlled by a PC running Presentation software version 10.2 (http://www.neurobs.com).

Procedure. Participants arrived at the laboratory with a parent. Before the experiment began, participants had the option of familiarizing themselves in a dummy scanner (a simulation scanner that lacks the magnetic field of the MRI scanner). Next, they completed several questionnaires dealing with mood, depression, and puberty. This was followed by the collection of saliva samples, after which the participants were positioned in the MRI scanner. The participants completed one additional nonemotional task in the MRI scanner before starting the experimental task. To familiarize them with the setup of the AA task, a short training session was completed (10 min). This was immediately followed by the fMRI session (20 min) and an anatomical scan (5 min). At the end of the fMRI session, saliva measurements were collected again, followed by two additional tasks outside the MRI scanner.

Pubertal development measures. Following Volman et al. (2011b), saliva samples for testosterone (Table 1) and cortisol measurements were collected at ~2–4 cm above the IBL by leveraging the Pubertal Development Scale (PDS; Petersen et al., 1988), a self-report questionnaire that contains questions on secondary sexual characteristics (Table 1). Participants indicated on a 4 point scale whether a physical characteristic (1) has not yet developed, (2) is slightly developed, (3) is moderately developed, and (4) is mature or not known. They were then stored at ~24°C. Testosterone concentration was measured using a competitive chemiluminescence immunoassay (CLIA) with a sensitivity of 0.0025 ng/mL (IBL). The intra-assay and interassay coefficients are between 10% and 12%. Cortisol concentration was measured using a commercially available CLIA with high sensitivity of 0.16 ng/mL (IBL). For this assay, the intra-assay and interassay coefficients are <8%. Participants were instructed to refrain from consuming any food, cigarettes, and drinks (except water) at least 1 h before the experiment. Testosterone levels undergo changes during the day/night cycle with the largest variations for mid-pubertal boys during the night (Albertsson-Wikland et al., 1997) and peak testosterone levels for girls occurring in the early morning (Ankærberg and Norrjaaera, 1999). The study design minimized the effects of those fluctuations on testosterone level estimation by sampling this hormone in duplicate, taken 2 h apart, after 10:00 A.M., resulting in consistent measurements across these two time points.

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Behavioral analysis. The behavioral data were analyzed with Matlab 2012 (MathWorks) and SPSS Statistics 19 (IBM). To obtain a reliable measure of the movement onset, joystick movement was reconstructed from the joystick displacement measures on each trial (Volman, 2011b). Reaction time (RT) was defined as the time from picture presentation to movement onset. Trials with no response or a joystick movement in the incorrect direction were excluded from further analysis, as were trials with an extreme RT (<100 and >1500 ms), an RT >-3 SDs from the mean, and an error rate (ER) above chance level in a block (in which instance, the entire block was excluded). To correct for a skewed distribution of the RT, a log transformation was applied. Mean RTs were calculated for each level of the two experimental factors [Valence (happy, angry) and Response (approach, avoid)]. The ER (as a percentage) analysis included trials with either no response or a joystick movement in the wrong direction. Mean ERs were also calculated for each level of the two experimental factors. Log-transformed and standardized per group (boys, girls) testosterone and cortisol levels from the first salivary measures were included in the model as covariates, with testosterone as a covariate of interest (Volman et al., 2011b). A three-way repeated-measures multivariate ANOVA with factors Group (boys, girls), Response (approach, avoid), and Valence (happy, angry) was conducted on the RT and ER separately; with testosterone and cortisol included in the model as covariates, testosterone being a covariate of interest. The α level was set at p = 0.05.

fMRI preprocessing and analysis. Functional data were preprocessed and analyzed using the Matlab toolbox SPM8 [Statistical Parametric Mapping (www.fil.ion.ucl.ac.uk/spm)]. The first four volumes were discarded to control for T1 equilibration effects. The multiecho sequence acquired four echoes per volume at every time point. The head motion parameters were estimated on the MR images with the shortest echo time (9.4 ms) because these images are the least affected by possible artifacts in the multiecho GRAPPA (generalized autocalibrating partially parallel acquisition) MR sequence (Poser et al., 2006). The calculated correction parameters, estimated using a least-squares approach with six rigid-body transformation parameters (translations, rotations), were applied to the remaining echoes of the same volume. Using an optimized echo-weighting method (the first 30 time points acquired before the actual experiment started), the four echoes were combined into a single volume. To correct for time differences in the acquisition of slices within a volume, the time courses of each voxel were realigned to the middle slice (slice 17). The T1-weighted image was co-registered to the mean of the functional images. Subsequently, the T1-weighted images were segmented into gray matter, white matter, and CSF using the “New Segment” tool in SPM8. To achieve more accurate spatial normalization given the relatively young age of the participants, the fMRI time series was normalized and smoothed using a group-specific template. Diffeomorphic anatomical registration through exponentiation of Lie algebra (DARTEL; Ashburner, 2007) was used for intersubject registration of the gray matter images. The fMRI time series was then transformed and resampled at an isotropic voxel size of 2 mm, resulting in spatially normalized, Jacobian scaled, and smoothed (8 mm FWHM Gaussian kernel) images in Montreal Neurological Institute (MNI) space.

Multiple regression fMRI analysis. The fMRI time series were analyzed in an event-related design within the general linear model. The vectors describing the onset and duration of each condition (approach—happy, approach—angry, avoid—happy, avoid—angry) were convolved with a canonical hemodynamic response function. Trials excluded from behavioral analysis were modeled with a separate regressor (“misses”), as well as those with instructions or feedback (“info”), resulting in six task-related regressors in the SPM multiple regression analysis. Residual head movement-related effects were modeled using the original, squared, cubic, first-order, and second-order derivatives of the movement parameters estimated with the spatial realignment procedure (Lund et al., 2005). Time courses of signal intensities of white matter, CSF, and the portion of the MR image outside the skull were included as the third and fourth level regressors. This procedure accounts for image intensity shifts due to movement of the hand within or near the magnetic field of the scanner (Verhagen et
Table 2. Reaction times and error rates in the AA task for approach and avoidance movements in response to happy and angry faces

<table>
<thead>
<tr>
<th></th>
<th>Approach</th>
<th>Avoid</th>
</tr>
</thead>
<tbody>
<tr>
<td>RT (ms)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Happy</td>
<td>622 (16)</td>
<td>679 (16)</td>
</tr>
<tr>
<td>Angry</td>
<td>673 (19)</td>
<td>679 (17)</td>
</tr>
<tr>
<td>ER (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Happy</td>
<td>6.4 (0.5)</td>
<td>8.4 (0.9)</td>
</tr>
<tr>
<td>Angry</td>
<td>8.7 (1.2)</td>
<td>6.7 (0.8)</td>
</tr>
</tbody>
</table>

Values indicate mean (SE) across participants.

Table 3. Significant clusters showing a larger effect during affect-incongruent than affect-congruent trials in the approach avoidance task modulated by testosterone

<table>
<thead>
<tr>
<th>Anatomical region</th>
<th>Side</th>
<th>BA</th>
<th>k</th>
<th>MNI coordinates</th>
<th>p</th>
<th>t</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive testosterone modulation of congruency effect</td>
<td>R</td>
<td>10/46</td>
<td>33</td>
<td>40 58 -6</td>
<td>0.045</td>
<td>3.55</td>
</tr>
<tr>
<td>Negative testosterone modulation of congruency effect</td>
<td>L</td>
<td>626</td>
<td>-8 20 16</td>
<td>&lt;0.001</td>
<td>5.39</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>37</td>
<td>272 -38 34 -2</td>
<td>0.013</td>
<td>6.56</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>6</td>
<td>196 8 50</td>
<td>0.046</td>
<td>4.91</td>
<td></td>
</tr>
</tbody>
</table>

BA, Brodmann area; k, number of voxels in a cluster; p, FWE-corrected cluster-level value; t, t-statistic at the peak voxel; R, right; L, left; SFS, superior frontal sulcus.

*VBM analysis. The anatomical images were manually checked for significant anatomical abnormalities and scanner artifacts, and were aligned to the anterior commissure at the origin. Next, the anatomical images were segmented into gray matter, white matter, and CSF using the “New Segment” tool in SPM8. DARTEL (Ashburner, 2007) was used for inter-subject registration of the gray matter images. The registered images were smoothed (8 mm FWHM Gaussian kernel, Jacobian scaled, threshold at 0.2) and transformed into MNI space. The gray matter images were entered into a second-level multiple regression analysis with testosterone (standardized and log-transformed per gender) as a covariate of interest. Gender and TBV (total gray matter and white matter volume) were included in the model as covariates of no interest. All statistical tests were performed on the voxel level, familywise error corrected for the whole brain or small volume corrected across all voxels in a region of interest. Our ROIs were those that were found to be significantly modulated by testosterone in the fMRI analysis. We used masks from the AAL atlas (Tzourio-Mazoyer et al., 2002) for the thalamus and amygdala, and the mask of the lateral frontal pole used in the fMRI analysis (Neubert et al., 2014).

**Results**

**Behavioral results**

Significant Valence $\times$ Response interactions for RTs ($F_{(1,43)} = 17.44, p < 0.001, \eta^2_p = 0.289$) and ERs ($F_{(1,43)} = 4.721, p = 0.035, \eta^2_p = 0.099$) confirmed the AA task-congruency effects as reported previously, driven by longer reaction times and more errors for affect-incongruent compared with affect-congruent responses (Table 2). These behavioral congruency effects did not significantly interact with testosterone ($F_{(1,43)} = 0.097, p = 0.757$) or gender ($F_{(1,43)} = 0.008, p = 0.928$). Additionally, there were significant main effects of Valence ($F_{(1,43)} = 38.366, p < 0.001$), Group ($F_{(1,43)} = 7.352, p = 0.01$), and Testosterone ($F_{(1,43)} = 4.745, p = 0.035$) for RTs. Furthermore, testosterone significantly interacted with Valence ($F_{(1,43)} = 6.8, p = 0.012$) and Response ($F_{(1,43)} = 5.018, p = 0.03$) on RTs, and with Valence ($F_{(1,43)} = 5.454, p = 0.024$) for ERs, indicating faster responses to happy faces, faster avoidance, and more errors on angry faces in individuals with lower testos-
testosterone levels. A correlational analysis between testosterone levels and PDS scores revealed a positive relationship ($r = 0.29, p = 0.048$), confirming that higher testosterone levels are associated with higher puberty scores (Huang et al., 2012).

fMRI results

Multiple regression analysis

The congruency effect in the right aPFC (local maximum: 40, 58, -6) was positively modulated by testosterone, indicating that adolescents with high testosterone levels showed a stronger aPFC effect for incongruent compared with congruent trials. In contrast, the congruency effect in the thalamus (local maximum: -8, -20, 16) and right amygdala (local maximum: 26, -2, -14) showed significant negative modulations by testosterone (for all effects, see Table 3). The Automated Talairach Atlas (Lancaster et al., 2000) and anatomical specifications by Arcaro et al. (2015) indicate that the activation cluster in the thalamus covered the dorsal pulvinar nucleus, the lateral dorsal nucleus, and the midline nucleus. A confirmatory ROI analysis of the pulvinar (mask extracted from the Automated Talairach Atlas) indicated a bilateral significant modulatory effect of testosterone (local maximum left: -10, -24, 16; voxel extent: 14; SVC $p_{FWE} = 0.028$; local maximum right: 24, -24, 12; voxel extent: 8; SVC $p_{FWE} = 0.037$). In summary, individuals with high testosterone levels showed relatively stronger aPFC congruency effects, whereas participants with low testosterone levels showed relatively stronger congruency effects in the pulvinar and amygdala (Fig. 2).

Next, we verified whether those individuals who have high aPFC congruency effects are also those with low subcortical congruency effects, and vice versa. We tested for a testosterone-modulated interaction between activity in aPFC and pulvinar. A two-way repeated-measures ANOVA with factors Congruency (incongruent, congruent) and Region (aPFC, pulvinar), and with testosterone as a covariate of interest, was conducted on parameter estimate values extracted from the peak coordinates of the given regions. An additional model considered parameter estimates from the amygdala instead of the pulvinar. A significant Congruency $\times$ Region $\times$ Testosterone interaction indicated that testosterone predicted stronger aPFC congruency-related effects in those participants with weaker congruency-related effects in the pulvinar ($F_{(1,45)} = 18.17, p < 0.001, \eta^2_p = 0.288$) and in the amygdala ($F_{(1,45)} = 17.72, p < 0.001, \eta^2_p = 0.282$; Fig. 2d). Post hoc confirmatory analyses verified this effect separately within each participant gender. For both boys and girls, significant Congruency $\times$ Region $\times$ Testosterone interactions were found for parameter estimates from the aPFC-pulvinar (boys: $F_{(1,19)} = 6.12, p = 0.023, \eta^2_p = 0.243$; girls: $F_{(1,24)} = 11.94, p = 0.002, \eta^2_p = 0.332$) and aPFC-amygdala (boys: $F_{(1,19)} = 5.02, p = 0.037, \eta^2_p = 0.209$; girls: $F_{(1,24)} = 13.40, p = 0.001, \eta^2_p = 0.358$).

A VBM analysis assessed whether the fMRI effects reported above could be accounted for by brain structural differences related to pubertal development. There were no significant testosterone-related differences in gray matter in the aPFC, pulvinar, and amygdala.

Functional connectivity

We performed a post hoc analysis to test for the presence of changes in coupling between the amygdala and pulvinar during emotional action control. We conducted a psychophysiological interaction analysis, with the right amygdala as the seed region and the congruency effect as a psychological factor. Following the study by Volman et al. (2011b), the connectivity strength for incongruent versus congruent trials was evaluated as a function of testosterone and, for the purpose of the present study, was combined with task performance. The latter was indexed with the congruency effects (incongruent vs congruent trials) observed in ERs and in RTs. We tested for the combined effects of testosterone and performance on amygdala–pulvinar connectivity as indexed by testosterone and ER and by testosterone and RT. There were no significant results when searching over the whole brain. We conducted an ROI analysis using a right thalamic mask, which indicated that the connectivity strength between the right amygdala and right pulvinar (local maxima: 18, -20, 18) changed as a function of the combined effect of testosterone and error rates (extent: 8 voxels; SVC $p_{FWE} = 0.043$). A confirmatory ROI analysis using a right pulvinar mask (Auto-
and amygdala to frontal cortex in relation to motor response inhibition (Rubia et al., 2000; Stevens et al., 2007), reward sensitivity (Forbes et al., 2010; Urošević et al., 2012; Braams et al., 2015), and perceptual processing of emotional faces (Monk et al., 2003; Hare et al., 2008; Somerville et al., 2011; Barbalat et al., 2013; Vink et al., 2014). This study indicates that, when control is exerted on decisions involving emotional responses, the pulvino–amygdalar pathway may be the relevant precursor of the mature aPFC emotional control system.

The pulvinar and emotional control

The pulvinar has been relatively overlooked in human neural development studies, yet an emerging literature points to its relevance for emotion and action control. Strong afferent and efferent connections with frontal, parietal, and temporal areas make the pulvinar suitable for integrating and coordinating the selection of a broad range of goal-directed actions (Wilke et al., 2010; Saalmann et al., 2012). Furthermore, the pulvinar connects the superior colliculus with the amygdala (Morris et al., 1999; Tamietto et al., 2012), and facilitates fast emotional saliency detection and threat recognition (Ward et al., 2005, 2007; Padmala et al., 2010; Nguyen et al., 2013a). These anatomical and neurophysiological properties indicate that the pulvinar can contribute to emotional action control (Barron et al., 2015). The current study indicates that this contribution is particularly relevant during pubertal development, when frontal–striatal–thalamic coupling has yet to be completed (Stevens et al., 2007; Barbalat et al., 2013).

Recent diffusion tensor imaging studies have shown that, during early development, connectivity among subcortical structures is more prevalent than during later development (Simmonds et al., 2014; Baker et al., 2015). This study qualifies that general observation by highlighting the role of the pulvino–amygdalar pathway in the control of emotional actions at a time when the aPFC is still developing. It remains to be seen whether the amygdala facilitates the transition toward mature emotional control by incorporating social context into the processing of emotional stimuli (Scherf et al., 2013) or whether the amygdala interferes with emotional control by enhancing bottom-up sensory influences on action tendencies (Hare et al., 2005). The developmental role of the amygdala may not in fact be uniform. The centromedial amygdala shows connectivity patterns with subcortical regions, including the pulvinar, whereas the lateral basal amygdala shows connectivity patterns with prefrontal areas (Roy et al., 2009; Bzdok et al., 2013). These subdivisions of the amygdala may undergo distinct circuit changes during adolescence. The findings of this study call for further investigation of these possibilities. On the one hand, developmentally more mature adolescents (as indexed by higher testosterone levels) used the same aPFC region recruited in adults (Volman et al., 2011a,b, 2013; Radke et al., 2015). When amygdala–pulvinar connectivity was stronger during emotional control, adolescents with higher testosterone levels made more errors. This observation tentatively suggests that amygdala–pulvinar connectivity may interfere with emotional control once that function is supported by the aPFC. On the other hand, regardless of pubertal maturation, adolescents had matched performance in exerting emotional control and, as such, irrespective of aPFC or amygdala–pulvinar circuitry involvement. This observation suggests that both prefrontal and amygdala–pulvinar circuits support emotional control equally well, at least within the rather mild emotional control demands evoked by the AA task.
Interpretational issues

Given the known differences in cerebral structure related to pubertal development and testosterone levels (Bramen et al., 2012; Nguyen et al., 2013b; GODDINGS et al., 2014; UROŠEVIĆ et al., 2014), it might be argued that the functional differences reported in this study are due to testosterone-related structural differences. However, there were no statistically significant differences in the structural features of the pulvinar, amygdala, or aPFC that were related to testosterone levels. This controlled situation is a major advantage of the current experimental design, focused on isolating puberty-related cerebral changes without generic age-related differences in executive control development.

The choice of testosterone levels as an objective and physiological marker of pubertal development is supported by previous work (Shirtcliff et al., 2009; Huang et al., 2012), and by the fact that this hormone is mechanistically involved in neuronal reorganization during adolescence (SISK and ZEHR, 2005; SCHULZ et al., 2009; HERTING et al., 2014). The study design minimized the effects of testosterone level fluctuations (ALBERTSSON-WIKLAND et al., 1997; ANKARBERG AND NORJAVAARA, 1999), as can be seen in consistent measurements across two time points, sampled 2 h apart. Additionally, we found that testosterone levels in our sample were positively correlated with puberty ratings for boys and girls (Shirtcliff et al., 2009). One might argue that testosterone is not only a proxy of puberty phase, but also of dominance behavior (ENTER et al., 2016). However, in this study testosterone was not related to anger–approach tendencies. In fact, previous work has shown that testosterone is more closely related to physicians’ ratings of pubertal maturation than self-reports, providing an objective and validated proxy of the actual pubertal stage (HUANG et al., 2012).

The anatomical specificity of the emotional control effect found in the pulvinar is limited by the characteristics of the cluster-level statistics since the effect includes other structures, namely the anterior and medial thalamic nuclei and the caudate nucleus. These regions are also involved in goal-directed and stimulus–response actions (Packard and Knowlton, 2002; OSTLUND and BALLEINE, 2008; BRADFIELD et al., 2013; PARNAUDEAU et al., 2015). Future studies might be able to differentiate among the specific contributions of those structures to emotional control in the early stages of puberty.

Based on this study, a broad extension to adolescent behavior in emotionally arousing situations is speculative and may need to be addressed with experiments where the social emotional challenge is not mild, such as the zooming version of the AA task (HEUER et al., 2007; RINCK and BECKER, 2007). The zooming effect (the angry face becomes larger–approach or smaller–avoid, depending on the response) has been shown to enhance the congruency effect and is especially sensitive in capturing failures in emotion action control in various psychopathologies (ROELOFS et al., 2009b, 2010; von BORRIES et al., 2012).

Conclusion

Developmentally more mature adolescents, in a manner similar to that of adults, recruit the aPFC during the control of emotional action tendencies, whereas less mature adolescents recruit the pulvinar and amygdala. These results are relevant for neurobiological models of pubertal development, qualifying the circuits implicated in the maturation of emotional action control. The findings are also relevant for understanding the neurobiological bases of emotion control alterations, given that the onset of affective disorders peaks during adolescence (Kessler et al., 2007). Chronic failures in emotion action control are common in various psychopathologies, such as social anxiety (ROELOFS et al., 2009b, 2010) and aggression-related disorders (von BORRIES et al., 2012). This study opens the possibility of replicating these findings in longitudinal studies testing whether affective disorders are mechanistically related to an altered developmental transition between the pulvino–amygdalar pathway and the mature aPFC emotional control system.

References


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