Development of attention networks from childhood to young adulthood: A study of performance, intraindividual variability and cortical thickness

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Abstract

Human cognitive development is manifold, with different functions developing at different speeds at different ages. Attention is an important domain of this cognitive development, and involves distinct developmental trajectories in separate functions, including conflict processing, selection of sensory input and alertness. In children, several studies using the Attention Network Test (ANT) have investigated the development of three attentional networks that carry out the functions of executive control, orienting and alerting. There is, however, a lack of studies on the development of these attentional components across adolescence, limiting our understanding of their protracted development. To fill this knowledge gap, we performed a mixed cross-sectional and longitudinal study using mixed methods to examine the development of the attentional components and their intraindividual variability from late childhood to young adulthood (n = 287, n observations = 408, age range = 8.5-26.7 years, mean follow up interval = 4.4 years). The results indicated that executive control stabilized during late adolescence, while orienting and alerting continued to develop into young adulthood. In addition, a continuous development into young adulthood was observed for the intraindividual variability measures of orienting and alerting. In a subsample with available magnetic resonance imaging (MRI) data (n = 169, n observations = 281), higher alerting scores were associated with thicker cortices within a right prefrontal cortical region and greater age-related cortical thinning in left rolandic operculum, while higher orienting scores were associated with greater age-related cortical thinning in frontal and parietal regions. Finally, increased consistency of orienting performance was associated with thinner cortex in prefrontal regions and reduced age-related thinning in frontal regions.

Introduction

Three decades ago, Posner and Petersen (1990) proposed to divide attention into subsystems that carry out separable attentional functions. This theoretical framework was later used to develop the ANT (Fan et al., 2002), which combines the Eriksen flanker task (Eriksen & Eriksen, 1974) and the cued reaction time task (Posner, 1980) into a single behavioral paradigm. In the ANT, executive control is defined as resolving conflict among responses; orienting as the selection of information from sensory input; and alerting as achieving and maintaining an alert state (Fan et al., 2002, 2009). The efficacy of the attentional networks that carry out these functions during the ANT have been investigated by comparing the reaction times in the incongruent trials (executive control), spatial cue trials (orienting) and double cue trials (alerting) to the reaction times in congruent trials, center cue trials and no cue trials, respectively (Fan et al., 2002). Here, the efficacy of the executive control network is estimated by how fast the participant can resolve and react to the conflict induced by incongruent flankers, where lower executive control scores are indicative of better conflict resolution abilities. The efficacy of the orienting network is estimated by the reduced reaction time following spatial cues, where higher orienting scores indicate better ability to utilize spatial information to orient and react to the correct target location. The efficacy of the alerting network is estimated by the reduced reaction time following alerting cues, where higher alerting scores indicate higher vigilance before the appearance of a target stimuli or, alternatively, difficulty of maintaining alertness when there is no cue.

The ANT has been used to investigate age-related differences in attention in different parts of life (e.g. Mahoney et al., 2010; Rueda et al., 2004), as well as attentional functions in clinical groups, including individuals with ADHD (Adólfsdóttir et al., 2008; Mogg et al., 2015), schizophrenia (Spagna et al., 2015), and autism (Fan et al., 2012). Although there have been

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several developmental studies on the attentional components in middle and late childhood (Federico et al., 2017; Mezzacappa, 2004; Mullane et al., 2016; Pozuelos et al., 2014; Rueda et al., 2004), there is a limited number of longitudinal studies directly investigating developmental changes (but see Lewis et al., 2016; Suades-González et al., 2017). Thus, the late developmental trajectories of these attentional components remain poorly understood. Finally, while we know that the brain undergoes substantial structural changes across adolescence (Dennis & Thompson, 2013; Tamnes & Mills, 2020), we have limited understanding of how the development of specific attentional components relate to these structural changes.

Previous research suggests that the attentional components have different developmental trajectories during childhood (Federico et al., 2017; Lewis et al., 2016; Mezzacappa, 2004; Mullane et al., 2016; Pozuelos et al., 2014; Rueda et al., 2004). Executive control seems to develop until early adolescence (Baijal et al., 2011; Mullane et al., 2016; Rueda et al., 2004), possibly beginning to stabilize around late adolescence (Waszak et al., 2010), orienting seems to be stable across middle childhood (Lewis et al., 2016; Mullane et al., 2016; Rueda et al., 2004; Suades-González et al., 2017), while alerting has been shown to continue to develop from early childhood into late childhood (Lewis et al., 2016; Mullane et al., 2016; Pozuelos et al., 2014). However, it should be noted that the alerting scores typically decrease with age, possibly because younger children do poorly in no cue trials (Rueda et al., 2004). Moreover, the abovementioned studies have typically focused on overall reaction time and accuracy measures, but not intraindividual variability; i.e. variability across trials or sessions of the same task (MacDonald et al., 2009; Williams et al., 2005). Previous research has shown large age effects for intraindividual variability in both congruent and incongruent trials of the flanker task (Tamnes et al., 2012) and for choice reaction time and simple reaction time

tasks during development (Dykiert, Der, Starr, & Deary, 2012), documenting improved consistency with increasing age. By using the different trial types in the ANT, it is possible to investigate the development of the intraindividual variability in each of the three attentional components, henceforth referred to as executive control variability, orienting variability and alerting variability. Similar to the reaction times during the ANT, it seems plausible that intraindividual variability in incongruent trials, spatial cue trials and double cue trials will show a continuous reduction with age, suggesting that a decrease in executive control variability and an increase in orienting variability and alerting variability are likely to be observed with increasing age. This field of research is important as within-task intraindividual performance has been shown to provide insight into temporal fluctuations of cognitive functioning (Bielak et al., 2010; MacDonald et al., 2006), and is found to be larger among children with attentional problems compared to controls (Epstein et al., 2011).

In parallel to the development of the attentional networks, the brain also undergoes substantial structural changes (Fjell et al., 2015; Herting et al., 2018; Tamnes et al., 2017; Vijayakumar et al., 2016; Wierenga et al., 2014), but studies linking the two are lacking. Previous functional MRI and electroencephalography (EEG) studies have investigated ANT associated brain activity among both adults (Fan et al., 2005, Galvao-Carmona et al., 2014; Neuhaus et al., 2010) and children (Konrad et al., 2005, Gopalan et al., 2019), and have reported that the different attentional networks recruits different brain regions. Executive control has been associated with activity in the anterior cingulate and bilateral prefrontal cortical regions (Fan et al., 2005; Konrad et al., 2005), orienting with activity in bilateral superior parietal cortical regions, and alerting with activity in thalamic, inferior parietal and prefrontal cortical regions (Fan et al., 2005). Structural MRI studies are scarce, but a crosssectional study by Westlye et al (2011), which included 286 adults 20-84 years of age,

showed a positive association between executive control and cortical thickness of the anterior cingulate and inferior frontal cortical regions. They also reported negative associations between alerting and thickness in superior parietal cortical regions, while no relations between orienting and cortical thickness were found (Westlye et al., 2011). While this study provided insight into the structural correlates of the attentional networks in adults, the associations between the development of these attentional networks and structural brain maturation remain unknown.

The present study aimed to examine the 1) developmental trajectory, 2) intraindividual variability, and 3) structural cortical correlates of executive control, orienting and alerting from late childhood to young adulthood. Based on previous behavioral studies, we expect executive control performance improvements to decelerate (i.e. initial decrease that slows down) with age (Baijal et al., 2011; Mullane et al., 2016; Waszak et al., 2010), while we expect orienting to remain stable from late childhood to young adulthood (Rueda et al., 2004; Suades-González et al., 2017). Previous results have shown a decrease in alerting during childhood (Lewis et al., 2016; Pozuelos et al., 2014), possibly driven by poor performance in no cue trials among the youngest participants (Rueda et al., 2004). However, as alerting scores may reflect either difficulty in maintaining an alert state in cue trials or efficient use of alerting cues (Posner, 2008), and we do not know how these processes change across adolescence, no prediction about alerting could be made within the current age range. Furthermore, as intraindividual variability shows large age-related effects across childhood to young adulthood (Dykiert et al., 2012; Tamnes et al., 2012), we also expect a continuous development of the intraindividual variability measures of the attentional components across adolescence. Since we expect increased consistency in incongruent trials, spatial cue trials and double cue trials with age, we predict that the executive control variability scores will

decrease with age and that orienting variability will increase with age. However, if the previous notion that the alerting effect is highly influenced by the performance in no cue trials is true, it seems likely that this will be reflected in the variability scores as well, hence no prediction about the direction could be made. Finally, we hypothesize that the attentional components and their intraindividual variability to be related to cortical thickness and age-related cortical thinning in the ANT associated brain regions (Fan et al., 2005; Konrad et al., 2005; Westlye et al., 2011). Thus, we expect that thinner cortex and faster age-related cortical thinning in the anterior cingulate and lateral prefrontal regions to be associated with lower executive control and executive control variability scores. Furthermore, we predict that thinner cortex and faster age-related cortical thinning in the superior parietal region to be associated with higher orienting and orienting variability scores, and that cortical thickness and age-related thinning in parietal and prefrontal regions will be associated with alerting and alerting variability scores.

Materials and Methods

Participants

The sample was drawn from the three-wave accelerated longitudinal research project *Neurocognitive Development* (Ferschmann et al., 2019; Tamnes et al., 2010, 2013). The study was approved by the Regional Committee for Medical and Health Research Ethics. Initially, children and adolescents were recruited through newspaper ads and local schools, while new participants were recruited through social media platforms in the second and third waves to compensate for attrition and to increase the sample size and to account for practice effects. Written informed consent was obtained from all participants over 12 years of age and from a parent of participants under 16 years of age. Oral informed assent was given by children under 12 years of age. At each wave, participants aged 16 years or older and a parent of

participants under 16 years were screened with standardized health interviews to ascertain eligibility. Participants were required to be right handed, fluent Norwegian speakers, have normal or corrected to normal vision and hearing, not have history of injury or disease known to affect central nervous system (CNS) function, including neurological or mental disorders and serious head trauma, not be under treatment for a mental disorder, not use psychoactive drugs known to affect CNS functioning, not have had a premature birth (< 37 weeks), and not have MRI contraindications. All MRI scans were evaluated by a neuroradiologist and required to be deemed free of serious injuries or conditions.

A total of 287 participants (153 females) in the age-range 8.5–26.7 years (M = 17.6, SD = 4.1, across all observations) were included in the current study and had ANT data from one or two time points (ANT was not included in the test protocol in the first wave of the *Neurocognitive Development* project). Of these, 166 participants (92 females) had data from one time-point and 121 participants (61 females) had data from two time-points, yielding a total of 408 observations. For participants with data from two time-points, the mean interval between the observations was 4.4 years (SD= 0.4, range= 3.9–5.5). The length of the interval was not related to age (r=.08, p=.36).

MRI subsample

A subsample of 169 participants (83 females) had quality controlled MRI data (see below for details) from the same scanner and were included for MRI analyses, where 112 participants (60 females) had MRI data from two time points, yielding a total of 281 observations with valid MRI and ANT data. Of the full sample, 117 participants recruited in the third wave were not scanned on the same scanner as the rest of the sample due to transition to a new

MRI scanner, and one participant had missing MRI data, thus 118 participants were not included in the MRI analyses.

The Attention Network Test

We administered an adult version of the ANT (Fan et al., 2002), which consisted of a practice block with 24 trials and three blocks with 96 trials each, for a total of 288 test trials. During assessment, participants were seated in a comfortable chair at approximately 60 cm distance from a 19-inch monitor. For each trial, participants were required to press a key indicating whether a target arrow was pointing to the left or right. The arrow was presented either above or below a centrally located fixation cross. The target arrow was flanked by one of three different types of stimuli: 1) pairs of congruent arrows, 2) pairs of incongruent arrows, or 3) pairs of neutral lines. Each type of flanker stimuli was presented 32 times per block. Furthermore, each trial was preceded by one of four cue conditions, with each variant occurring 24 times in each block: 1) no cue, 2) center cue, 3) double cue, or 4) spatial cue. The cues, when presented, were single (center and spatial cue) or double asterisks replacing (center cue) or accompanying the fixation cross. The size of the fixation cross was approximately 0.5×0.5 cm (~0.5°), and the diameter of the asterisks used for cueing was about 0.3 cm (\sim 0.3°). Target arrows were horizontally centered 1.3 cm (\sim 1.2°) below or above the fixation cross. Each trial was initiated by the fixation cross for a random duration of 400-1600 ms. This was followed by the cue stimuli for 100 ms, the fixation cross for 400 ms and then the target stimuli, which remained visible on the screen until response or for 1700 ms. The participants were instructed to focus on both speed and accuracy throughout the session, and the length of the breaks between blocks were controlled by the participant. The experimental procedure was administered using E-prime software (Schneider et al., 2002), and responses were obtained on a PST Serial Response Box. The ANT adult version

was chosen instead of the ANT child version (Rueda et al., 2004), as pilot testing suggested that it was more suitable across the current age-range and since it includes more trials per time.

MRI acquisition

The MRI data was collected at Rikshospitalet, Oslo University Hospital with the same 12channel head coil on the same 1.5 T Siemens Avanto scanner (Siemens Medical Solutions, Erlangen, Germany) at both time-points. The sequence used for morphological analyses was a 3D T1-weighted magnetization prepared rapid gradient echo (MPRAGE) with the following parameters: repetition time/echo time/time to inversion/flip angle = 2400 ms/3.61 ms/1000 ms/8°, matrix = 192 x 192 x 160, sagittally acquired, field of view = 240 mm, bandwidth = 180 Hz/pixel, voxel size $1.25 \times 1.25 \times 1.2$ mm. Acquisition time of this sequence was 7 minutes and 42 seconds.

MRI processing

The MRI data underwent whole brain segmentation and cortical surface reconstruction and longitudinal processing in Freesurfer 6.0 (Dale et al., 1999; Fischl et al., 1999, 2002, 2012; Reuter et al., 2012). Cortical thickness was estimated based on the shortest distance between the gray matter/white matter boundary and the cortical surface. This was conducted at each vertex across the cortical mantle. Before statistical analyses, cortical thickness was resampled to 81,924 vertices and the thickness maps were smoothed with a Guassian kernel with a full-width of half-maximum of 10 mm. All MR images were manually inspected by a trained individual and a re-scan from the same session was used if available and if the images were deemed to be of insufficient quality (see Ferschmann et al., 2019 for details)

Statistical analyses

ANT components. To calculate ANT components, we used a ratio procedure with median reaction times based on correct trials. The ratio procedure was chosen to isolate the attention scores from the expected decrease in reaction times with age. We computed the following attention network scores (RT represents the reaction time):

$$Executive \ control \ = \ \frac{[RT \ incongruent \ trials - RT \ congruent \ trials]}{RT \ congruent \ trials}$$
$$Orienting = \ \frac{[RT \ center \ cue - RT \ spatial \ cue]}{RT \ spatial \ cue}$$
$$Alerting = \ \frac{[RT \ no \ cue - RT \ double \ cue]}{RT \ double \ cue}$$

Additionally, we utilized the same ratio procedure, replacing median reaction time with the standard deviation of the median reaction time to obtain a measure of intraindividual variability for each of the ANT components. We computed the following intraindividual variability scores (SD represents the standard deviation):

$$Executive \ control \ variability = \frac{[SD \ incongruent \ trials - SD \ congruent \ trials]}{SD \ congruent \ trials}$$

$$Orienting \ variability = \frac{[SD \ center \ cue - SD \ spatial \ cue]}{SD \ spatial \ cue}$$

$$Alerting \ variability = \frac{[SD \ no \ cue - SD \ double \ cue]}{SD \ double \ cue}$$

Development of the ANT components and their intraindividual variability. We used the nlme package (Pinheiro, Bates, DebRoy, Sarkar, & R Core Team, 2018) in R (R Core Team, 2018) to investigate the development of the attentional components and their intraindividual variability. All the continuous predictors in the general linear mixed models were centered

around their mean prior to the analyses. First, we assessed the best fitting developmental model by comparing the following models:

1) null model (Y = intercept + random (participant ID) + error)

2) linear age model (Y = intercept + random (participant ID) + age + error)

3) quadratic age model (Y = intercept + random (participant ID) + age + age^2 + error)

Second, if one of the age models was a significantly better fit than the null model, we examined whether the best fitting age model was further improved by including sex in the model (e.g., for linear model):

4) Main sex model (Y = intercept + random (participant ID) + age + sex + error)
5) Interaction age^xsex model (Y = intercept + random (participant ID) + age + sex + age*sex + error)

Models were fitted with maximum likelihood (ML) estimation method. To avoid overfitting, model selection was guided by *p*-values generated by the likelihood ratio (LRT) test, but the Akaike Information Criterion (AIC; Akaike, 1974) was emphasized. More specifically, the more complex model was chosen if the LRT *p*-value dropped below .05 and the AIC value decreased by at least 2. Since starting and ending points of samples can influence the age-related trajectory when using parametric models (Fjell et al., 2010), we used generalized additive mixed models (GAMM) with a cubic spline basis (Wood, 2017) for visualization of the results. The GAMM plots were produced using the following model in R: Y ~ s(age, bs = 'cr'), where "s" is the function used to define smooth terms, whereby the smooth is a function of "age", "bs" is the smoothing basis to use, and the "cr" refers to a cubic regression spline.

Note that the GAMM plots are used for visualizations only, while the plotted general linear mixed models can be found in the supplementary materials (Supplementary Figures 1 and 2).

ANT components and cortical thickness. We conducted vertex-wise analyses in SurfStat (http://www.math.mcgill.ca/keith/surfstat/) to investigate the relationship between behavioral performance in the ANT and cortical thickness. At each vertex, we computed a linear mixed model to investigate the effect of the attentional components and their intraindividual variability scores (ANT_{component}), interaction effect of ANT_{component} *age and ANT_{component} *age² using the following model: $Y_{thickness}$ = intercept + ANT_{component} + Sex + Age + Age² + ANT_{component} *Age + ANT_{component} *Age² + random(participant) + error in SurfStat All the continuous predictors were centered around their mean before running the analyses. To account for multiple comparisons, we used a random field theory correction of p < .05 and a cluster-defining threshold of p < .005. Here, we will report the significant clusters of the main effect of ANT_{component} and age².

Results

Overall behavioral performance. Behavioral data for each trial type are presented in Table 1. The results showed that the participants had > 91% accuracy across all trial types, indicating that the participants understood the task. The mean attentional component scores are presented in Table 2. Across all observations, the descriptive data showed lower RTs in the congruent, spatial cue and double cue compared to the incongruent, center cue and no cue condition, respectively. The positive values for the attentional components indicate that the sample showed the expected effect of the task conditions.

		Cue co	ondition		Flanker condition			
	Center	Double	No	Spatial	Congruent	Incongruent	Neutral	
Accuracy	95.43	95.57	97.00	97.24	99.05	91.29	98.69	
(SD)	(4.67)	(4.45)	(3.29)	(3.85)	(1.64)	(8.23)	(1.96)	
Mean	485.17	474.72	519.46	437.77	452.00	554.05	452.19	
(SD)	(81.27)	(76.14)	(82.97)	(74.78)	(72.78)	(94.33)	(70.06)	

Table 1. Accuracy and reaction times for the ANT trial types.

Note: Accuracy is presented in percentage. Mean reaction times are based on median reaction time on correct trials. Accuracy and reaction times are estimated across all observations. SD = standard deviation.

	To	tal sample	e	Females	Males
	Mean (SD)	Min	Max	Mean (SD)	Mean (SD)
Executive					-
control	.23 (.07)	.07	.56	.23 (.07)	.22 (.07)
Orienting	.11 (.05)	03	.26	.12 (.05)	.10 (.05)
Alerting	.10 (.06)	06	.34	.10 (.06)	.09 (.05)

Table 2. ANT performance for the sample as a whole, and females and males separately

Note: SD = standard deviation.

Development of the ANT components. For executive control, a quadratic age model, with no main effect of sex or age-sex interaction, was found to best fit the data (Table 3). For orienting, the best fitting model was a positive linear model with sex included as a main effect (Table 4), while alerting was best fitted by a negative linear model with no main effect

of sex or age-sex interaction (Table 5). The results from the mixed model analyses showed that the three attention networks had three different age models that best fitted the data (Table 6). The GAMM visualizations (Figure 1) indicate a decelerating negative effect of age on executive control, stabilizing around late adolescence, while the GAMM for orienting shows a linear increase across the sample range, with females having higher orienting scores than males. The GAMM slope for alerting, however, indicates a cubic pattern, with a decelerating negative effect of age during adolescence.

Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	-1050.57	-1038.53	528.28	-	-	-
Linear age model	2	4	-1055.78	-1039.74	531.89	1 vs 2	7.21	0.007
Quadratic age model	3	5	-1059.47	-1039.41	534.74	2 vs 3	5.69	0.017
Main sex model	4	6	-1058.29	-1034.22	535.14	3 vs 4	0.82	0.366
Interaction Sex*Age ² model	5	8	-1057.99	-1025.90	536.99	3 vs 5	4.52	0.211

Table 3. Comparison of models predicting executive control

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Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	-1293.55	-1281.52	649.78	-	-	-
Linear age model	2	4	-1300.10	-1284.05	654.05	1 vs 2	8.55	0.003
Quadratic age model	3	5	-1299.60	-1279.55	654.80	2 vs 3	1.50	0.220
Main sex model	4	5	-1309.51	-1289.46	659.76	2 vs 4	11.42	0.001
Interaction sex*age model	5	6	-1307.52	-1283.45	659.76	2 vs 5	11.42	0.003
Interaction sex*age model	5	6	-1307.52	-1283.45	659.76	4 vs 5	0.00	0.957

Table 4. Comparison of models predicting orienting

Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	-1198.03	-1185.99	602.01	-	-	
Linear age	2	4	-1204.53	-1188.49	606.27	1 vs 2	8.50	0.004
Quadratic age model	3	5	-1203.38	-1183.32	606.69	2 vs 3	0.85	0.357
Main sex model	4	5	-1203.75	-1183.70	606.88	2 vs 4	1.22	0.269
Interaction sex*age model	5	6	-1202.76	-1178.69	607.38	2 vs 5	2.23	0.328

Table 5. Comparison of models predicting alerting

Table 6. Model parameters for the best fitting model of the attentional components

Attention network	Value	S.E	S.E CI (95%)		p-value
			[Lower, Upper]		
Executive control					
Intercept	.219	.005	[.21, .23]	47.72	<.001
Age	002	.0008	[004,0005]	-2.64	<.01
Age ²	.0004	0.0002	[.0001, .0007]	2.39	<.05
Orienting					
Intercept	.119	.004	[.113, .126]	32.20	<.001
Age	.002	.0006	[.0004, 0.003]	2.66	<.01
Sex	018	.005	[029,008]	-3.40	<.001
Alerting					
Intercept	.096	.003	[.090, .102]	32.12	<.001
Age	002	.0007	[003,0006]	-2.93	<.01

Notes. S.E = Standard Error, CI = Confidence interval

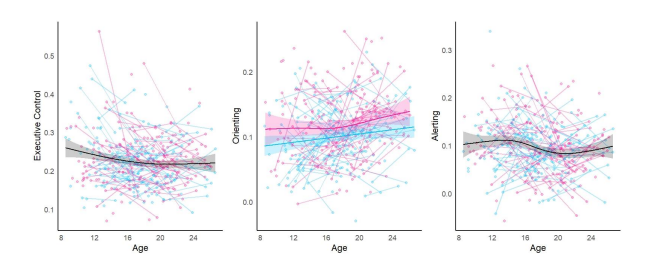


Figure 1. Developmental trajectories of the three attentional components. Left: a decelerating negative effect of age for executive control. Middle: a linear increase with age where females show higher scores in orienting than males. Right: a slight decrease in alerting during adolescence. Splines are plotted using generalized additive mixed modelling, and were plotted separately for each sex in orienting. Females are represented in pink and males in blue. Shaded areas correspond to the 95% confidence interval.

Development of intraindividual variability of the ANT components. For executive control variability, none of the models were found to improve model fit compared to the null model (Table 7). Orienting variability (Table 8) and alerting variability (Table 9) were both best fitted by a linear age model with no main effect of sex or age-sex interaction effect. Thus, the results of the best fitting model for intraindividual variability scores in the ANT indicated development in two of the three attention networks (Table 10). Specifically, orienting variability was positively associated with age, while alerting variability was negatively associated with age. The GAMM slopes were in line with the statistical linear models and showed a linear increase and a linear decrease for orienting variability and alerting variability, respectively (Figure 2).

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Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	247.77	259.80	-120.88	-	-	-
Linear age model	2	4	247.64	263.68	-119.82	1 vs 2	2.13	0.144
Quadratic age model	3	5	249.04	269.10	-119.52	1 vs 3	2.73	0.256

Table 7. Comparison of models predicting executive control variability

Table 8. Comparison of models predicting orienting variability

Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	195.95	207.97	-94.97	-	-	-
Linear age model	2	4	182.09	198.12	-87.04	1 vs 2	15.86	<0.001
Quadratic age model	3	5	183.30	203.34	-86.65	2 vs 3	0.78	0.376
Main sex model	4	5	183.91	203.96	-86.96	2 vs 4	0.17	0.678
Interaction sex*age model	5	6	185.83	209.88	-86.91	2 vs 5	0.26	0.880

Model	Model number	df	AIC	BIC	LogLik	Test	L.Ratio	р
Null model	1	3	140.19	152.22	-67.10	-	-	-
Linear age model	2	4	133.56	149.59	-62.78	1 vs 2	8.63	0.003
Quadratic age model	3	5	135.53	155.57	-62.76	2 vs 3	0.03	0.862
Main sex model	4	5	134.65	154.69	-62.33	2 vs 4	0.91	0.341
Interaction sex*age model	5	6	136.24	160.30	-62.12	2 vs 5	1.31	0.518

Table 9.	Comparison	of	models	predicting	alerting	variability

Table 10. Model	parameters for the	e best fitting model	of attention n	etworks variability
	p			

Attention network	Value	S.E	CI (95%)	t-value	p-value
variability	[Lower, Upper]				
Orienting variability					
Intercept	.190	.016	[.160, .221]	12.26	< 0.001
Age	.015	.004	[.008, .022]	4.02	<.001
Alerting variability					
Intercept	.063	.015	[.035, .092]	4.31	<.001
Age	010	.003	[017,003]	-2.95	< 0.01

Notes. S.E = Standard Error, CI = Confidence interval

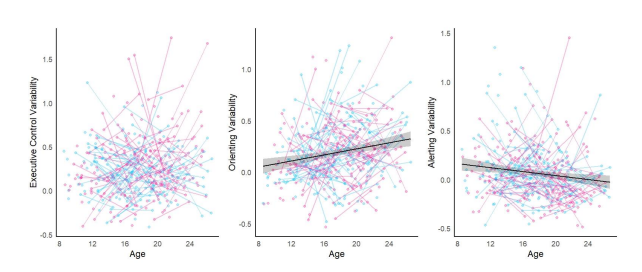


Figure 2. Developmental trajectories for the intraindividual variability measures of the three attentional components. Left: Spaghetti plot of executive control variability with no spline as no age terms improved the model fit of executive control variability. Middle: a linear increase with age on orienting variability. Right: a linear decrease with age on alerting variability. Splines are plotted using generalized additive mixed modelling. Shaded areas correspond to the 95% confidence interval.

Associations between ANT performance and cortical thickness. There were significant regional associations between two out of six behavioral measures and cortical thickness (Figure 3). Higher alerting scores were associated with thicker cortex in a cluster encompassing parts of the right anterior prefrontal cortex, while higher orienting variability scores were associated with thinner cortex in two clusters including parts of the left anterior and superior prefrontal cortex. These main effects were largely the same without the age interaction terms included in the mixed effect model. Neither executive control or orienting, or executive control variability or alerting variability were associated with cortical thickness.

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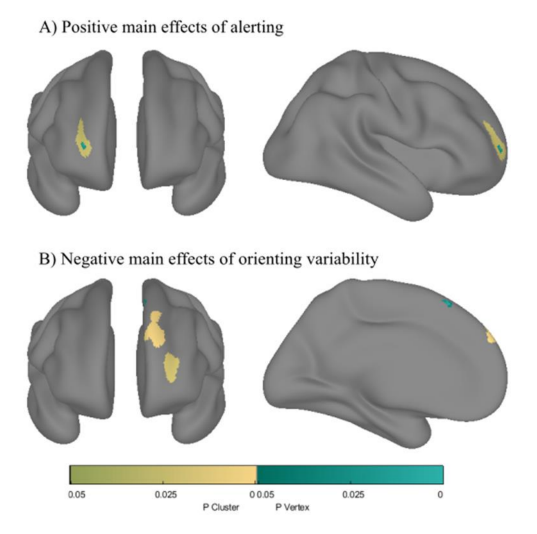


Figure 3. Main effects of the behavioral measures of the ANT on cortical thickness. A) A positive association between alerting and cortical thickness within the right anterior prefrontal cortex. B) A negative association between orienting variability and cortical thickness within the left anterior prefrontal cortex and left superior prefrontal cortex. The shades of yellow represent significant P-clusters, while the shades of turquoise represent significant vertices. The clusters and vertices are significant at p < .05 Random field theory corrected.

Interactions between ANT performance and age on cortical thickness. The results of the analyses testing the interaction term between ANT performance and age was significant for three out of six of the behavioral measures (Figure 4). A negative interaction between orienting and age yielded three significant clusters within the left inferior parietal lobule, left dorsolateral prefrontal cortex and left insula. Furthermore, a negative interaction between alerting and age was associated with cortical thickness within the left rolandic operculum. A positive interaction between orienting variability and age yielded two significant clusters in the right insula and the right supplementary motor area. No significant age interactions with executive control, executive control variability, nor alerting variability were found on cortical thickness. Finally, no association was found between cortical thickness clusters and any of the ANT measures interacting with age².

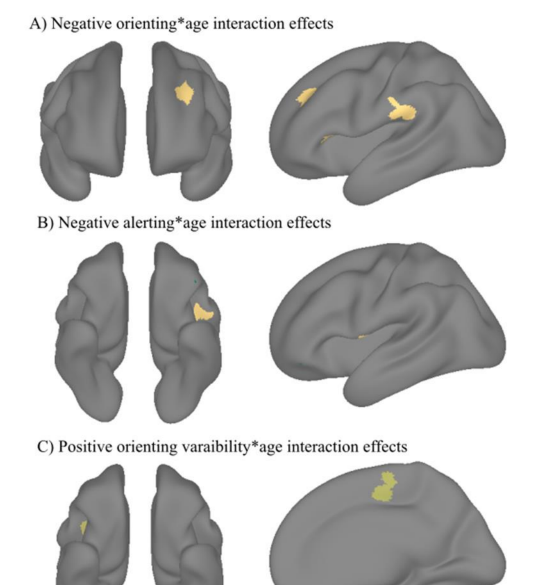


Figure 4. Interaction effect between the behavioral measures of the ANT and age on cortical thickness. A) Higher orienting scores were associated with greater age-related cortical thinning within the left dorsolateral prefrontal cortex, left insula and left inferior parietal lobule. B) Higher alerting scores were associated with greater age-related cortical thinning within left rolandic operculum. C) Lower orienting variability were associated with greater age-related cortical thinning within right insula and right medial supplementary motor area. The shades of yellow represent significant P-clusters, while the shades of turquoise represent significant at p < .05 Random field theory corrected.

P Cluster

0 0.05

P Vertex

0.025

0

0.05

0.025

Discussion

The present study investigated the developmental trajectories of three attentional functions and their intraindividual variability from late childhood to young adulthood, as well as their relationships to cortical thickness. Executive control showed a decelerating development from late childhood and into adolescence, with an initial improvement in late childhood and early adolescence, followed by a stabilization in late adolescence. Orienting continued to develop from late childhood to young adulthood, illustrated by a continuous increase in orienting performance, with females having higher orienting scores than males. Alerting did also exhibit a continuous development across the current age range, with a negative association with age. We also found a linear increase in orienting variability and a linear decrease in alerting variability with age. For the brain-behavior relationships, there was a main effect of two out of six of the behavioral measures on cortical thickness. Lower alerting scores were related to thinner cortex in a cluster within the right anterior prefrontal cortex, while higher orienting variability scores were related to thinner cortex in frontal regions. In addition, we identified an interaction effect with age on cortical thickness for three out of six of the behavioral measures. Here, higher orienting scores were related to greater age-related cortical thinning in the left inferior parietal and left prefrontal regions, higher alerting scores were related to greater age-related cortical thinning in the left operculum, and lower orienting variability scores were related to greater age-related cortical thinning in the right insula and the right motor area.

In line with our predictions and previous research that has shown age-related changes in executive control during adolescence (Waszak et al., 2010), our results support development of executive control that seem to stabilize around late adolescence. However, in contrast to previous research that has shown orienting to be fully developed by late childhood (Mullane

et al., 2016; Pozuelos et al., 2014; Suades-González et al., 2017), we found evidence for further age-related changes in orienting, where higher orienting scores were associated with higher ages. Most notably, we identified a main effect of sex with females having higher orienting scores compared to males, which could indicate that females benefit more from spatial cues than males. Furthermore, the results indicated a linear decrease in alerting with age, although the GAMM showed a more complex trajectory as the youngest participants seemed to profit from the alerting cues, while its beneficial effect seemed to decelerate during adolescence. As previously suggested, this might be due to poor performance in no cue trials among the youngest participants. (Rueda et al., 2004). It should also be noted that previous research on development of attention networks using ANT have included younger samples and used a modified version of ANT, such as the child version (Mezzacappa, 2004; Rueda et al., 2004; Suades-González et al., 2017). Hence, the discrepancies in the two versions could explain inconsistencies between the developmental trajectories of the attention networks reported with the child version and our results from the adult version.

It is well known that intraindividual variability of the reaction times in cognitive tasks show large age-effects across childhood and adolescence (Dykiert et al., 2012; Tamnes et al., 2012). While there have been several studies investigating the development of the attentional components, studies mapping the developmental trajectory of the intraindividual variability in the attentional components are lacking. To fill this knowledge gap, we substituted the median reaction time with standard deviation when calculating attention network performance, hence calculating estimates for intraindividual variability in the three attentional components. Although we expected a decrease in executive control variability with age, the results did not indicate any age-related effects. However, in line with our predictions, there was a linear increase with age on orienting variability, indicating increased

consistency in spatial cue trials with age. Further, the analyses also revealed that the alerting variability decreased with age and it seems plausible that this is due to faster increase in consistency in no cue trials, which would support the previous notion that youngest participants perform worse when there is no cues (Rueda et al., 2004), hence slower reaction times and larger trial-to-trial variability in no cue trials.

Intraindividual variability measures may reflect attentional lapses during cognitive tasks, which might be more common in children (Tamnes et al., 2012) and in individuals with ADHD (Adams et al., 2011). In fact, variability measures of reaction times often provide robust group differences between children with ADHD and control groups, irrespective of the cognitive task at hand, and have greater effect sizes compared to many other neuropsychological test outcomes (Epstein et al., 2011). Thus, we urge future studies to further utilize the present intraindividual variability measures obtained from the ANT as this could provide further insight into the development of performance consistency and differences in variability in participants with attentional problems.

We also investigated the relationship between behavioral measures obtained from the ANT and cortical thickness. First, we examined the main effects of the behavioral measures on cortical thickness. Although previous structural studies have found associations between executive control and anterior cingulate and the prefrontal cortex (Westlye et al., 2011), we did not observe any significant relationship between executive control and cortical thickness. However, in line with our prediction, we did observe a main effect of alerting in frontal regions, although no relations to thickness in parietal regions were found. Here, lower alerting scores were related to thinner cortex in a cluster within the right anterior prefrontal region. However, in contrast to our positive association between alerting and cortical

thickness, Westlye and colleagues (2011) did find a negative association between cortical thickness in parietal regions and alerting among adults. To reconcile these results, it seems plausible that lower alerting scores are indeed indicative of better attentional functioning, while cortical thinning is part of typical development in youths (Tamnes et al., 2017) and reflects aging processes in older adults (Fjell et al., 2009), which could explain why there is a positive association between alerting and cortical thickness early in life, and a negative association between alerting and cortical thickness later in life. In addition, assuming that reduced cortical thickness reflects better cognitive functioning among youths, this would further support the negative association between orienting variability scores and cortical thickness in left prefrontal regions, as higher orienting variability scores indicate increased consistency in spatial cue trials compared to the center cue trials.

The results also yielded significant clusters from the age interaction term with orienting, alerting and the orienting variability. It is interesting to note that orienting showed a different developmental pattern than predicted. Firstly, we did not expect any age-related differences during adolescence in the behavioral performance, while we predicted orienting performance to be associated with cortical thickness in superior parietal regions. However, in addition to exhibit age-related improvements, orienting was also related to two cortical clusters within prefrontal regions and one in the inferior parietal region. It is further interesting to note that increased orienting scores indicates increased age-related cortical thinning in the left insula, while the complete opposite pattern was observed for orienting variability, where higher orienting variability scores indicated less age-related cortical thinning in the right insula. Although speculative, this could indicate a role of the bilateral insula in the detection of relevant stimuli and in a speed-consistency trade off. Future studies should aim to investigate the role of the bilateral insula in orientation and possible role in speed and consistency during

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development. Finally, several of the significant clusters across alerting, orienting and orienting variability performance showed interaction with age in predominantly frontal regions, which might reflect a general improvement of higher attentional functioning during maturation of the frontal cortex. However, as we do not know the direction of causality or the underlying mechanisms of cortical thinning, we urge caution when interpreting these results. Indeed, it is important to note that the age-related cortical thinning should not be interpreted as thinning per se, but as MRI apparent cortical thinning of the distance between the estimated outer grey and white matter surfaces (Walhovd et al., 2017). Apparent cortical thinning with increasing age during development can partly be explained by increased myelination could result in more efficient neural processing, which is likely to be part of the underlying mechanism that allow for the observed continuous reduction in both reaction time and intraindividual variability during development.

To summarize, the current study included cross sectional and longitudinal data that provides important insight into the developmental trajectories of the efficacy of three attention networks. The behavioral results showed a decelerating age effect on executive control, stabilizing around late adolescence. Orienting showed continued development into young adulthood, with females having higher orienting scores than males, while alerting showed a positive linear age-effect, while alerting variability exhibited a negative linear age-effect across late childhood and young adulthood. Further, lower alerting scores were associated with thinner cortex within a cluster encompassing parts of the right anterior prefrontal cortex, while higher orienting variability scores were associated with thinner cortex in clusters within the left anterior prefrontal and left superior prefrontal cortex. Finally, higher orienting scores were

related to greater age-related cortical thinning in left dorsolateral prefrontal cortex, left insula and left inferior parietal lobule, higher alerting was related to greater age-related thinning in left rolandic operculum, while higher orienting variability scores were related to less agerelated cortical thinning in the right insula and right medial supplementary motor area.

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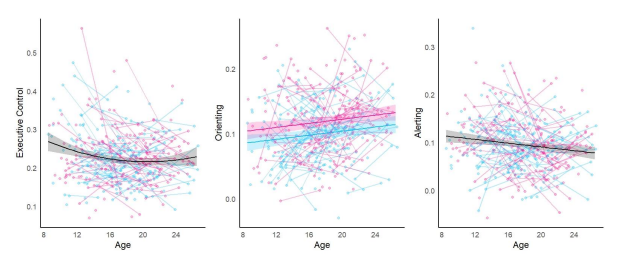
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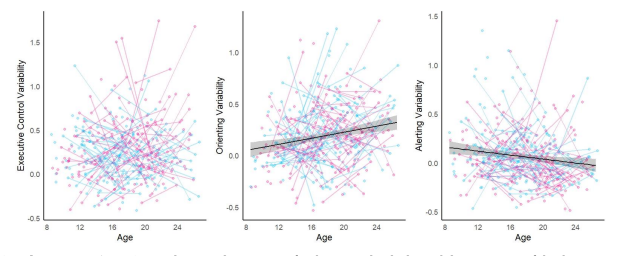
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Supplementary materials



Supplementary Figure 1. Developmental trajectories for the three attentional components. Left: a decelerating negative effect of age on executive control. Middle: a linear positive effect of age and a main effect of sex on orienting. Right: a negative linear effect of age on alerting. The regression lines are plotted using the mixed models that best fitted the data. Shaded areas correspond to the 95% confidence interval. Boys are represented in blue and girls are represented in pink.



Supplementary Figure 2. Developmental trajectories for the intraindividual variability measures of the three attentional components. Left: spaghetti plot of executive control variability with no regression line as no age terms improved the model fit of executive control variability. Middle, a positive linear effect of age on orienting variability. Right, a negative linear effect of age on alerting variability. Shaded areas correspond to the 95% confidence interval. The regression lines are plotted using the mixed models that best fitted the data.