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RECEIVED 19 December 2023 ACCEPTED 30 May 2024 PUBLISHED 11 June 2024

#### CITATION

Croons H, Martens DS, Vanderstukken C, Sleurs H, Rasking L, Peusens M, Renaers E, Plusquin M and Nawrot TS (2024) Telomere length in early childhood and its association with attention: a study in 4–6 year old children.

Front. Pediatr. 12:1358272. doi: 10.3389/fped.2024.1358272

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# Telomere length in early childhood and its association with attention: a study in 4–6 year old children

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Telomere length (TL), a marker of cellular aging, has been studied in adults with regard to its connection to cognitive function. However, little is known about the association between TL and cognitive development in children. This study investigated the interplay between TL and cognitive functioning in 283 Belgian children aged four to six years of the Environmental Influence on Aging in Early Life (ENVIRONAGE) birth cohort. Child leukocyte TL was measured using qPCR, while cognitive functioning, including attention and memory, was assessed using the Cambridge Neuropsychological Test Automated Battery (CANTAB). Linear regression models were employed to examine the association between TL and cognitive outcomes, adjusting for potential confounders. We found an inverse association between TL and the spatial errors made during the Motor Screening task (p = 0.017), indicating a higher motor accuracy in children with longer telomeres. No significant associations were found between TL and other cognitive outcomes. Our results suggest a specific link between TL and motor accuracy but not with the other cognitive domains.

#### KEYWORDS

cognition, telomere, attention, childhood, ENVIRONAGE

### 1 Introduction

Cognitive development, the mental processes involved in acquiring, storing, processing, and using information (1), is crucial for learning and adaptative skills, and often translates to higher academic accomplishments, more promising career opportunities, contributing to an overall better quality of life (2). It has been well established that environmental factors such as lifestyle, fitness, and nutrition, as well as genetic factors, contribute to cognitive functioning (3).

Recently, research has focused on the potential role of the biological underpinnings of cognition, including telomere biology (4). Telomeres are the protective caps at the ends of human chromosomes that play a crucial role in cellular aging (5). Telomere length (TL), reflects the replicative capacity of cells (6–8) and leukocyte telomere length (LTL) has been shown to be associated with cognitive function in several studies (9–13), but results are inconclusive (14–17) Previous research primarily focused on the association between TL and cognition in adults, particularly the elderly (4, 11, 13, 18), while the relationship between TL and neurodevelopmental outcomes in children has received little to no attention. To date, only one study has examined the association between TL and cognition in 209 children at the age of 5, and no association was observed (19).

Given that childhood is a crucial period for cognitive development, this study aims to investigate the association between leukocyte TL and cognitive functioning in children aged four to six years participating in the prospective Environmental Influence on Aging in Early Life (ENVIRONAGE) birth cohort.

# 2 Methods

#### 2.1 Study design and population

ENVIRONAGE is an ongoing prospective birth cohort established in 2010 (Limburg, Belgium). Mother-child pairs, without planned cesarean section, and where the mother was able to fill out a Dutch language questionnaire, were recruited at birth at the East-Limburg Hospital (Genk, Belgium). When the child reached the age of 4 to 6 years old, they were invited for a follow-up visit at the university study center. Detailed information on the recruitment process at birth is provided elsewhere (20). Between October 2014 and October 2019, 439 mother-child pairs had a follow-up examination. The final ENVIRONAGE population for our analysis consists of 283 children (Supplementary Figure S1). The ENVIRONAGE study protocol is approved by the ethical committees of Hasselt University (Diepenbeek, Belgium, reference no. B371201216090 and B371201524537) and East-Limburg Hospital (Genk, Belgium). It has been carried out according to the Helsinki Declaration, and all mothers provided written informed consent.

#### 2.2 Measures

Mothers filled out detailed questionnaires at the in order to obtain household and lifestyle information. Data on child sex, maternal and child age, maternal education and average hours the child sleeps were retrieved. Maternal education is coded "low" when mothers did not obtain a diploma, "middle" when obtained a high school diploma, and "high" when obtained a college or university degree. Based on the date of the examination day, a seasonal scale was calculated (Winter is from the 21st of December to the 20th of March, spring from the 21st of March to the 20th of June, summer from the 21st of June to the 20th of September, and autumn from the 21st of September to the 20th of December). Maternal health status regarding the occurrence of preeclampsia, hypertension, infectious diseases or diabetes during pregnancy was retrieved via medical records.

In addition to the detailed questionnaires, mothers also completed the Perceived Stress Scale (PSS) questionnaire to assess their stress levels, and the Strengths and Difficulties Questionnaire (SDQ) developed by Goodman (21) to evaluate the child's behavior using SDQ sub-scales (peer relationship, emotional, conduct, and hyperactivity) and a Total Difficulties Score. Lastly, body anthropometrics of the children, including height (to the nearest centimeter) and weight (to the nearest 0.1 kilograms), were measured by trained staff. Body mass index (BMI) was calculated as the ratio of weight over squared height.

#### 2.3 Cognitive measurements

The children's neurocognitive functioning was assessed using the Cambridge Neuropsychological Test Automated Battery (CANTAB) on a touch-screen tablet (CANTAB, Cognitive assessment software, 2019). The CANTAB has been shown to provide reliable measurements of executive functions in children as young as four years old (22). Trained examiners administered four tasks according to standardized instructions. Two of these tasks focused on attention and psychomotor speed: the Motor Screening Task and the Big/Little Circle Task, which assessed response accuracy (in pixels) and/or response latency (in milliseconds). The other two tasks assessed visual recognition and working memory. The Spatial Span task evaluated the maximum number of squares the child could remember in the correct sequence (span length). The Delayed Matching to Sample task measured response latency (in milliseconds), percentage correct (%), and error probability (%) (Figure 1). More detailed information on the CANTAB procedures can be found in Supplementary Text S1.

#### 2.4 Relative average telomere length

Child blood was drawn and genomic DNA was isolated from the buffy coat of venous blood stored in EDTA tubes using the QIAamp<sup>®</sup> DNA mini kit (Qiagen GmbH, Hilden, Germany) following the manufacturer's instructions. DNA quantity and purity were assessed using the Nanodrop spectrophotometer (ND-1000; Isogen Life Science, De Meern, Netherlands) and the Quant-iT<sup>TM</sup> PicoGreen<sup>®</sup> dsDNA Assay Kit (Life Technologies, Foster City, CA, USA) using the Omega Fluostar plate reader (BMG LABTECH, Ortenberg, Germany). TL was measured using an adapted qPCR method described by Cawthon (23). Telomeres were measured in triplicate in a single batch. Telomere lengths were expressed as the ratio of telomere copy number to singlecopy gene number (T/S) relative to the mean T/S ratio of the entire sample. Detailed sample collection procedures and TL-assay specifications, and reliability parameters are provided in Supplementary Text S2 and have been previously provided for the ENVIRONAGE birth cohort (24).

#### 2.5 Statistical analysis

Data was analyzed using JMP Pro 16.2.0 software. Participants' characteristics were presented as means (SD) or frequencies (%). All latency data, i.e., response latency of the Motor Screening Task, Big/Little Circle task, and Delayed Matching to Sample task were log-transformed (log10), to better comply with assumptions of model linearity. The association between the different cognitive outcomes and TL was assessed using multivariable-adjusted general linear models. The threshold for statistical significance was set at a 95% confidence level (p < 0.05). We adjusted our models for *a priori*-selected covariates that may be associated with both the dependent (cognitive



outcomes) and independent variables (TL at 4–6 years old). In model 1, this includes the child's sex and age. In model 2, we further adjusted for the child's BMI, average sleep hours of sleep the child has on a typical day and night, maternal education, and the examination season. The normality of the residuals was visually evaluated by Q-Q plots. Estimates are presented as the difference in cognitive outcome for a 1-IQR increase in TL. For data that were log-transformed, we back-transformed the estimates and expressed them as a percentage difference.

To assess the robustness of our findings, we performed several sensitivity analyses on the computerized measures of cognitive function. First, we excluded children who showed any signs of disinterest or were distracted while taking the cognitive tests to account for possible response errors. Second, we adjusted our models for the child's behavior over the past six months by using four sub-scales of the SDQ questionnaire: the peer relationship and emotional subscales, and the conduct and hyperactivity subscales, each scored as normal vs. not normal. Further we additionally adjusted our models for PSS scores and maternal health conditions during pregnancy like diabetes, preeclampsia, hypertension or infections. Lastly, we stratified by sex.

### **3** Results

#### 3.1 ENVIRONAGE study population

Characteristics of the 283 participants are provided in Table 1. Children were on average (SD) 4.57 (0.41) years old, and approximately half of them were boys (51.9%). Most participants were of European ethnicity (94.0%), with an average BMI of 16.05 (1.26) kg/m<sup>2</sup>. The season in which the examinations took place was for 64 (22.6%) participants during autumn, 88 (31.1%) participants during spring, 71 (25.1%) participants during summer, and 60 participants (21.2%) during winter. The relative telomere length ranged from 0.64 to 1.47. Accompanying mothers were on average 30.1(4.4) years old, and the majority had a college education or higher (62.9%).

SDQ data was derived from 256 participants and categorized results into "normal" and "not normal" for each subscore. For the majority of participants (82%), the emotional symptom score falls within the "normal" range. The same is true for the hyperactivity score, where 77% exhibited behaviors classified as "normal", signifying a typical level of hyperactivity. Further, 78% of participants demonstrated "normal" conduct, indicating a standard range of behavioral conduct. Within the domain of peer relationships, 84% of participants showcased "normal" patterns, underscoring healthy social interactions. Moreover, 95% of participants exhibited "normal" prosocial behaviors, meaning they have a high degree of positive engagement and cooperation with others. The combination of these individual sub-scales resulted in a comprehensive assessment of participants' behavioral characteristics, quantified by the Total Difficulties Score. 86% of participants received a "normal" classification according to this measure.

### 3.2 Cognitive performance outcomes

The outcomes for the four CANTAB tasks are summarized in Table 2. The median (p25-p75) response latency of the Motor Screening task referring to the average time the child needed to

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Characteristics	Mean (SD) or frequency (%)			
Child				
Age at the visit, years	4.57 (0.41)			
Sex				
Male	144 (51.9%)			
Female	139 (49.1%)			
Ethnicity				
European	266 (94.0%)			
Non-European	17 (6.0%)			
BMI Child	16.05 (1.26)			
Sleep hours	10.88 (1.04)			
Season of examinations				
Autumn	64 (22.6%)			
Spring	88 (31.1%)			
Summer	71 (25.1%)			
Winter	60 (21.2%)			
Telomere Length (range)	0.64-1.47			
SDQ score <sup>a</sup>	'			
Emotional	211 (82%)			
Hyperactivity	198 (77%)			
Conduct	201 (78%)			
Peer problem	218 (84%)			
Prosocial	244 (95%)			
Total	222 (86%)			
Mother				
Age	30.1 (4.4)			
Education	·			
Low	27 (9.5%)			
Middle	78 (27.6%)			
High	178 (62.9%)			
PSS score	14.52 (6.06)			
Health conditions during pregn	iancy			
Diabetes	8 (2.8%)			
Preeclampsia	9 (3.2%)			
Hypertension	3 (1,1%)			
Infectious diseases	4 (1,4%)			

<sup>a</sup>SDQ scores based on 256 participants, frequency represents the participants with normal SDQ scores.

touch the target successfully, was 964.4 (803–1148.9) milliseconds. The median error, measured as pixel units from the target center made during the same task, was 14.3 (12.2–16.0). In the Big/ Little circle, the response latency was similarly calculated as the

TABLE 2 Characteristics CANTAB outcomes for the 283 ENVIRONAGE participants.

	Min	P25	P50	P75	Max			
Motor Screening Task								
Response Latency, ms	535.1	803	946.4	1148.9	1946.1			
Error, pixel units	6.65	12.2	14.3	16.0	21.5			
Big/Little Circle Task								
Response Latency, ms	683	965.7	1092.4	1228.6	2719.9			
Spatial Span Task								
Span length	0	2	3	3	5			
Delayed Matching to Sample Task								
Response Latency, ms	1376.6	3286.8	4142.9	5127.7	13994.7			
Percentage correct, %	10	35	45	55	85			
Probability of error given correct, %	0.1	0.4	0.6	0.7	1.0			



average time to select the correct circle and was observed to be 1092.4 (965.7–1228.6) milliseconds. During the Spatial Span task, participants demonstrated an ability to accurately reproduce a median of 3 (2–3) squares in the correct sequence. Finally, in the Delayed Matching to Sample task, participants needed a median time of 4133.1 (3282.8–5134.0) milliseconds to accurately select the correct pattern on the first try for all presented sample patterns. The percentage of trials answered correctly on the first try was 45.0 (35.0–55.0)% and the probability of error given a previous correct answer was 0.6% (0.4–0.7).

# 3.3 TL and cognitive performance outcomes

In unadjusted analysis, we only observed a negative association between TL and the error made during the Motor screening task (Figure 2). A 1-IQR increase in TL was associated with a 0.58pixel unit lower pixel error (95% CI: -1.04 to -0.12; p = 0.014). After adjustment for child sex and age the estimates were comparable and a 1-IQR increment in TL was associated with a 0.57-pixel unit decrease. In fully adjusted models this association remained significant (95% CI: -1.04 to -0.10; p = 0.017). No other cognitive outcomes related to attention and memory were found to be associated with childhood TL (Table 3).

#### 3.4 Sensitivity analysis

Results of the sensitivity analyses are shown in Supplementary Table S1. The association between TL and the error made during the Motor screening task remained robust after the exclusion of

	Unadjusted model		Model 1		Model 2				
	Estimate (95% CI)	р	Estimate (95% CI)	р	Estimate (95% Cl)	р			
Motor screening task									
Response Latency, ms	-0.28 (-4.28, 3.88)	0.89	0.30 (-3.41, 4.16)	0.88	0.42 (-3.33, 4.32)	0.83			
Error, pixel units	-0.58 (-1.04, -0.12)	0.014	-0.56 (-1.02, -0.10)	0.018	-0.57 (-1.04, -0.10)	0.017			
Big/little circle task									
Response Latency, ms	-1.31 (-4.57, 2.05)	0.44	-0.14 (-3.20, 2.99)	0.92	-0.20 (-3.25, 2.95)	0.90			
Spatial span task									
Span length	0.054 (-0.16, 0.27)	0.63	0.017 (-0.19, 0.23)	0.88	-0.01 (-0.22, 0.20)	0.94			
Delayed matching to sample task									
Response Latency, ms	3.6 (-2.58, 10.17)	0.26	4.08 (-2.23, 1079)	0.21	4.10 (-2.25, 10.85)	0.21			
Percentage correct, %	0.72 (-1.94, 3.39)	0.59	0.04 (-2.53, 2.60)	0.98	0.004 (-2.60, 2.60)	1.00			
Probability of error given correct, %	-0.002 (-0.04,0.03)	0.91	0.004 (-0.03,0.04)	0.84	0.006 (-0.03,0.04)	0.72			

TABLE 3 Association between child TL and cognition in children aged 4 to 6 years.

Estimates are presented as a difference in cognitive outcome for a 1-IQR increase in TL. Latency data is expressed as the percentage difference in cognitive outcome for a 1-IQR increase in TL.

Model 1 adjusted for sex and age.

Model 2 adjusted according to model 1 (sex, age) with additional adjustment for child BMI, average sleep hours, diploma of the mother, and the season of the examinations. Statistically significant values are highlighted in bold.

children who were disinterested or unfocused during the tests, and for additional adjustment for SDQ results, PSS results and maternal health conditions during pregnancy. After stratification for child sex, a 1-IQR increment in TL was associated with a 0.86 decrease in this error in boys and a and 0.49 decrease in this error in girls, and no significant interaction was observed (p-interaction = 0.93).

### 4 Discussion

In this study, we evaluated the association between leukocyte TL and cognitive performance in 4–6-year-old children specifically focusing on two domains of cognition: memory and attention. However, we found no evidence of a strong association. Nevertheless, we did observe that children with longer telomeres were more accurate in the Motor Screening Task, as shown by a decrease of 0.58-pixel unit distance from the target center, for a 1-IQR increase in TL. These findings were confirmed after adjustment for potential confounding factors.

Our study adds to the limited research examining the association between cognition and TL in children. In a study by Feiler et al. (19), no association was observed between leukocyte TL and neurodevelopmental outcomes in 209 5-year-old children, despite using a finger-tapping test similar to our motor screening task. It is important to note that the latter focused solely on latency and did not consider the distance between the object and the point of contact on the screen (19). In addition, several studies have found an association between TL and ADHD in children (7, 25, 26).

While early-life biological aging differences may not yet manifest as significant cognitive performance disparities in children, there is a growing recognition of the association between TL and various cognitive outcomes later in life. For instance, Valdes et al. found significant correlations between TL and CANTAB outcomes in a cross-sectional survey including 382 women aged 19 to 78 years (12). Similarly, TL has been linked to Modified Mini-Mental State Exam (MMSE) scores, a test widely used to assess global cognitive function, in 976 older men (11) and 17,052 adults (10). Additionally, associations have been found between TL and the composite score from six cognitive tests in 2,092 nurses (9). TL has also been linked to information processing speed, visual-spatial function, and memory, albeit to a lesser extent in the attention and executive domains (27).

Longitudinal investigations are crucial to capture the cumulative effects of early-life biological aging differences on cognition. Martin-Ruiz et al. found that longer telomeres at baseline was associated with less reduction in 2-year Modified Mini-Mental State Exam change scores in stroke survivors (28). Furthermore, elderly with longer TL exhibited better baseline attention and psychomotor speed and less decline in global cognitive functioning over seven years relative to elders with shorter TL (13). Other longitudinal studies have, however, failed to observe such effects in other populations (16, 17). Similarly, some cross-sectional studies did not observe significant associations of TL with general cognitive decline (14, 15) or specific cognitive domains (10, 14). Discrepancies in findings across these studies may be attributed to factors such as sample size, participant characteristics, TL assessment methods, and the types of cognitive measures used.

Children's brains demonstrate remarkable neuroplasticity, enabling rapid learning and adaptation. New neural connections readily emerge, fostering cognitive development. Consequently, the mechanisms linking telomere length to cognition may diverge between children and the elderly. At present, we can only speculate about possible mechanisms when investigating the link between TL and cognition. Possible mechanisms include: (i) oxidative stress and inflammation, which are commonly associated with aging, TL, and cognitive performance (29, 30). It is worth noting that brain tissue is particularly susceptible to oxidative stress due to its moderate levels of antioxidants, despite its high energy demands (31). Another hypothesized mechanism suggests (ii) a shared genetic basis for both the shortening of telomere length and the onset of cognitive decline (27). Studies examining the association between TL and gray matter atrophy concluded that shorter telomeres are associated with gray matter atrophy, mainly in the subcortical/limbic regions (32) which are found to be associated with children's cognitive abilities (33). Gampawar et al. stated that about half of the effect of TL on cognition in adults was mediated by the brain parenchymal fraction, which is the ratio of brain parenchymal (functional) volume to total brain volume (34). Better connectivity has also been observed with longer telomeres (35) and is a significant predictor of better performances in the attention/speed domain (36). An extensive meta-analysis on the role of telomere length in brain aging so far confirmed these results and showed that longer telomeres are indeed beneficial to both brain structure and cognition during aging (6).

Our study is among the first to investigate the relationship between TL and the cognitive functioning of children as young as four. Our study has several important strengths. First, we assessed neurocognition in children during the most critical period of brain development (37). Sensitivity analyses showed minimal changes to the observed associations, indicating robust and consistent results. We accounted for a potential lack of motivation causing inaccurate response validity in our sensitivity analysis. Furthermore, we also used leukocyte TL as a proxy for TL, since it was shown to correlate well with TL across different tissues within individuals (38). Nevertheless, we acknowledge the following limitations. First, this is a cross-sectional study, so we were unable to evaluate rates of change in these variables, which might be a more informative metric than a single measurement. Second, cognitive measurements are subject to random measurement error, such that we expect a certain amount of non-differential misclassification in our outcome, which might have attenuated our results. Even though results were measured in children during the most critical period of brain development, the pathology underlying cognitive impairments appears to begin decades prior to the onset of detectable symptoms, and thus our measurement of cognition might be quite early to be able to predict more obvious cognitive decline (9). In addition to that, our study population of 283 children might be relatively small to detect this possible decline, particularly if the effects are subtle and less pronounced in this early life stage as compared to later life cognitive effects. Last, we only found an association in one of the subdomains of attention. No associations were found for the other attention-related domains.

In conclusion, we found that LTL is only associated with higher motor accuracy in children aged 4 to 6 years. Currently, we have no evidence that memory or other domains of attention are associated with TL. Future studies with a larger sample size, prospective design, and other relevant biological markers (e.g., oxidative stress) are needed to clarify the role of TL in cognitive performances in children.

## Data availability statement

The data analyzed in this study is subject to the following licenses/restrictions: Requests to access these datasets should be directed to hanne.croons@uhasselt.be.

### Ethics statement

The studies involving humans were approved by Comité Medische Ethiek UHasselt (CME UHasselt). The studies were conducted in accordance with the local legislation and institutional requirements. Written informed consent for participation in this study was provided by the participants' legal guardians/next of kin.

## Author contributions

HC: Investigation, Writing – original draft. DM: Investigation, Supervision, Writing – original draft. CV: Investigation, Writing – review & editing. HS: Investigation, Writing – review & editing. LR: Investigation, Writing – review & editing. MP: Investigation, Writing – review & editing. ER: Investigation, Writing – review & editing. MP: Supervision, Writing – review & editing. TN: Supervision, Writing – review & editing.

# Funding

The author(s) declare financial support was received for the research, authorship, and/or publication of this article.

ENVIRONAGE is supported by grants of the Research Foundation, Belgium (Grant No. N1518119, No. G082317N and No. 1523817N). DM holds a postdoctoral grant by the Flemish Scientific Fund (FWO grant 12X9623N). LR acknowledges funding from the Special Research Fund (Bijzonder Onderzoeksfonds, BOF) for a doctoral fellowship: BOF20DOC15.

## Acknowledgments

The researchers would like to thank ENVIRONAGE participants, all the ENVIRONAGE follow-up study researchers, midwives, and the staff of the maternity ward of East-Limburg Hospital in Genk.

# Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

The author(s) declared that they were an editorial board member of Frontiers, at the time of submission. This had no impact on the peer review process and the final decision.

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## Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fped.2024. 1358272/full#supplementary-material

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