#### **RESEARCH ARTICLE**



# Differences in executive abilities rather than associative processes contribute to memory development

Nils C. J. Müller<sup>1</sup> | Nils Kohn<sup>1</sup> | Mariët van Buuren<sup>2</sup> | Nadia Klijn<sup>1</sup> | Helene Emmen<sup>1</sup> | Ruud M. W. J. Berkers<sup>1,3</sup> | Martin Dresler<sup>1,4</sup> | Gabriele Janzen<sup>1,5</sup> | Guillén Fernández<sup>1</sup>

<sup>1</sup>Donders Institute for Brain, Cognition and Behaviour, Radboud University Medical Centre, Nijmegen, The Netherlands

<sup>2</sup>Department of Clinical, Neuro and Developmental Psychology, Faculty of Behavioral and Movement Sciences, Institute for Brain and Behavior Amsterdam, Vrije Universiteit Amsterdam, Amsterdam, The Netherlands

<sup>3</sup>Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, Germany

<sup>4</sup>Max Planck Institute of Psychiatry, Munich, Germany

<sup>5</sup>Behavioural Science Institute, Radboud University Nijmegen, Nijmegen, The Netherlands

#### Correspondence

Guillén Fernández, Donders Institute for Brain, Cognition and Behaviour, Radboud University Medical Centre, Nijmegen, The Netherlands. Email: guillen.fernandez@donders.ru.nl

#### Funding information

NWO Research Talent Grant, Grant/Award Number: 406-13-008

#### Abstract

Children's learning capabilities change while growing up. One framework that describes the cognitive and neural development of children's growing learning abilities is the two-component model. It distinguishes processes that integrate separate features into a coherent memory representation (associative component) and executive abilities, such as elaboration, evaluation, and monitoring, that support memory processing (strategic component). In an fMRI study using an object-location association paradigm, we investigated how the two components influence memory performance across development. We tested children (10–12 years, n = 31), late adolescents (18 years, n = 29), and adults (25+ years, n = 30). For studying the associative component, we also probed how the utilisation of prior knowledge (schemas) facilitates memory across age groups. Children had overall lower retrieval performance, while adolescents and adults did not differ from each other. All groups benefitted from schemas, but this effect did not differ between groups. Performance differences between groups were associated with deactivation of the dorsal medial prefrontal cortex (dmPFC), which in turn was linked to executive functioning. These patterns were stronger in adolescents and adults and seemed absent in children. Thus, the children's executive system, the strategic component, is not as mature and thus cannot facilitate memory performance in the same way as in adolescents/adults. In contrast, we did not find age-related differences in the associative component; with activity in the angular gyrus predicting memory performance systematically across groups. Overall, our results suggest that differences of executive rather than associative abilities explain memory differences between children, adolescents, and adults.

#### KEYWORDS

executive abilities, fMRI, medial prefrontal cortex, memory development

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made. © 2021 The Authors. *Human Brain Mapping* published by Wiley Periodicals LLC.

#### 1 | INTRODUCTION

In virtually all contexts learners need to focus on what to learn, avoid distraction, relate information to each other or keep several types of information online to combine them. These abilities, executive functions, have been shown to strongly influence the efficiency of ones mnemonic system (Shing, Lindenberger, et al., 2010; Shing, Werkle-Bergner, et al., 2010; Simons & Spiers, 2003, see Ghetti and Bunge (2012) as well as Ofen and Shing (2013) for reviews on memory development). The maturation of the executive system-especially the prefrontal cortex-during adolescence (Bunge & Crone, 2009; Crone & Dahl, 2012; Luna, Marek, Larsen, Tervo-Clemmens, & Chahal, 2015; Maril et al., 2010, 2011; see Ofen, Tanga, Yua, & Johnson, 2019 for a recent review) makes it an excellent candidate to support the development of learning. The relation of executive functions with associative memory processes have been formalised and extended in a model explaining age-related differences in episodic memory: the two-component model of development (Shing, Werkle-Bergner, Li, & Lindenberger, 2008; Shing, Lindenberger, et al., 2010; Shing, Werkle-Bergner, et al., 2010). It postulates one associative and one strategic component with differential maturational trajectories. The associative component "refers to mechanisms of binding together different features of the memory content into coherent representations" (Shing, Lindenberger, et al., 2010; Shing, Werkle-Bergner, et al., 2010). The strategic component "refers to control processes that aid and regulate memory content at both encoding and retrieval" (Shing, Lindenberger, et al., 2010; Shing, Werkle-Bergner, et al., 2010). Whereas the strategic component is centred around the prefrontal cortex, the associative component is centred around the medial temporal lobe. However, the developmental interaction of the two systems and their underlying neurobiology are still poorly understood.

When we learn new information this usually involves prior knowledge. Almost nothing we learn is fundamentally new in all aspects but mostly relates to something we already know. This entails that when we form new memory representations, the different features that get integrated via the associative component of the two-component model also include prior knowledge. That prior knowledge benefits learning was first formulated by Bartlett (Bartlett, 1932) and Piaget (Piaget, 1936) in the context of schemas: Our knowledge is organised in schemas which can be used to readily assimilate new information about the world or provide a foundation that can be modified when we acquire new insight/perspectives. The idea of schemas had a strong influence on education and educational psychology (Thorndyke & Hayes-Roth, 1979). Throughout our life, we continuously acquire, modify, or enrich schemas. This difference in scope of schemas available to children versus adults might explain developmental memory differences (Brod, Werkle-Bergner, & Shing, 2013). Whereas adolescents and adults have a sophisticated net of knowledge spanning a large range of topics, children are still in the process of acquiring most of that. Thus, for new information, the children might have fewer opportunities to relate new information to their schemas. On the other hand, children might be superior in building new knowledge structures of previously unconnected information

due to their generally increased neural plasticity (Ismail, Fatemi, & Johnston, 2016).

Executive processes and the utilisation of schemas have both been linked to the prefrontal cortex, yet vary in their precise localisation (Miller & Cohen, 2001; Ridderinkhof, 2004; Tse et al., 2011; van Kesteren, Ruiter, Fernández, & Henson, 2012). Generally, the prefrontal cortex shows a protracted maturation trajectory, reaching a matured state only in the mid-twenties (Gogtay et al., 2004; Shaw et al., 2008). Brod, Lindenberger, and Shing (2017) observed that children aged 8-12 showed less medial prefrontal cortex (mPFC) activity when successfully retrieving schema-congruent events and stronger hippocampal activity when retrieving schemaincongruent events than young adults. Whereas most studies investigated the underlying neural correlates of memory development during mid- to late childhood, less attention has been given to early childhood below the age of 8. A recent study by Geng, Redcay, and Riggins (2019) observed that young children between 4 and 8 years also engage the hippocampus for encoding contextual details. Furthermore, a differentiation along the longitudinal axis of hippocampus increased with age. Thus, maturation of the mPFC and the hippocampus could play a significant role in the age-related variability in memory (DeMaster & Ghetti, 2013; Ghetti, DeMaster, Yonelinas, & Bunge, 2010; Keresztes et al., 2017; Keresztes, Ngo, Lindenberger, Werkle-Bergner, & Newcombe, 2018).

This relatively late maturation of the mPFC has previously been linked to the development of cognitive control as a crucial aspect of executive functions (Luna et al., 2015). While there is general agreement about the role of the mPFC for executive functions and the activation of prior knowledge, less consensus has been reached about the precise area within this region. While Brod and Shing (2019) were able to pinpoint the ventral mPFC as being central for the formation of schemas, Yu et al. (2018) could show that especially the dorsolateral PFC explains memory development. Müller et al. (2020) could not define a clear region within the mPFC.

Besides the mPFC and the hippocampus, the angular gyrus has been identified as a region that is not only involved in the retrieval of schemas (van Buuren et al., 2014), but especially in integrating or 'binding' prior learned information (Wagner et al., 2015). Wagner and coworkers showed that during retrieval different schema components converge within the angular gyrus. Interestingly, this convergence was only present after a consolidation period of 24-hr. The angular gyrus might be crucial for establishing combined single schema representations. However, in this study young adults aged 18–29 were investigated. The role of the angular gyrus for schema processing in children still needs to be explored.

The current study aims to investigate the two-component model and its role in memory development. Specifically, we further explore the role of the mPFC and the link to executive functions for the successful use of schemas during development. Based on that we tested three age groups that differ strongly in prefrontal maturation: children at mid- to late childhood between the age of 10–12 years, 18-yearold late adolescents, and adults over 25 years old. To measure memory performance we used a game-like object-location memory task (van Buuren et al., 2014). A strength of this paradigm is that it has no verbal requirements, which would have favoured the older groups as verbal abilities are still developing in children (Vakil & Blach stein, 2007). As associative component, we trained a schema during the first part of our study so that all groups have the same level of prior knowledge available to facilitate learning. For the strategic component, we included executive functions tests. We hypothesise that all age groups benefit from schemas, but that children perform worse on the executive function tests and engage the mPFC less than adolescents and adults.

#### 2 | MATERIALS AND METHODS

#### 2.1 | Participants

Ninety right-handed native Dutch-speaking volunteers participated in this study. As we investigated developmental differences related to differential maturation of the prefrontal cortex we tested three different age groups: Thirty adults aged between 25 and 32 years old (M = 26.9 years, SD = 21.9 months, 12 male), twenty-nine adolescents aged 18 (M = 18.5 years, SD = 3.1 months, 10 male) and thirtyone children aged between 10 and 12 years old (M = 11.0 years SD = 8.8 months, 8 male). All subjects had normal hearing and normal or corrected-to-normal vision. All participants were required to have no history of injury or disease known to affect the central nervous system function (including neuropsychological disorders such as dyslexia, autism, and attention deficit hyperactivity disorder) and to not have MRI contraindications. Assessment of these was based on selfreports by the participants. Adults and adolescents were recruited from the student population of Radboud University, Nijmegen, and from the surrounding community. Children were recruited through presentations and flyers at local schools. The study was approved by the institutional Medical Research Ethics Committee (CMO Region, Arnhem-Nijmegen). Written informed consent was obtained prior to participation from all participants who were at least 18 years old; for the children participating both parents signed the informed consent.

Of these 90 participants, 4 children had to be excluded (2 did not want to complete the study, 1 moved excessively in the scanner, 1 due to an experimenter error); 2 adolescents were excluded as they did not complete the training at home. All other participants completed the at home training satisfactorily. One adult had to be excluded due to an experimenter error. Of these 83 participants that completed the study, we excluded 11 (6 children, 1 adolescent, 4 adults) participants for the analysis based on their poor performance—see Section 2.13 for details. All analyses focussed on this final set of 72 participants (21 children, 26 adolescents, and 25 adults).

#### 2.2 | Summary of procedure

The study spanned 8 days in total. On Day 1, participants came to the lab for a first session. For the next 4 days, they performed additional

sessions at home. On Day 8, they returned to the institute for the final session. As a paradigm, we used an adapted version of the memory game task that was used in another study (van Buuren et al., 2014). Details of the paradigm are explained below. As additional measures, we utilised a short verbal memory task, a fractal *n*-back task (Ragland et al., 2002), the Wisconsin Card Sorting Task (WCST; Heaton, Chelune, Curtiss, Kay, & Talley, 1993), and the forward digit span task (Alloway, Gathercole, Kirkwood, & Elliott, 2008). The rationale for the additional measures is explained in a separate paragraph below.

On Day 1, participants came to the Donders Centre for Cognitive Neuroimaging and started in a behavioural lab with a practice of the *n*-back task, followed by the verbal memory task. Immediately after completing the verbal memory task, participants were taken into the MRI scanner where we first acquired a 10-min resting-state scan during which participants were instructed to lie still, think of nothing in particular, and look at a black fixation cross on a white background. After that, participants performed the *n*-back task and lastly, we acquired a structural scan. As the memory task required the use of a trackball—because of MRI-compatibility—participants had a practice session with the trackball (Kensington, Orbit Optical Trackball) that was used for all sessions of the memory task. During all uses of the trackball, we instructed participants to operate the trackball with two hands: the right dominant hand moves the cursor and the left-hand clicks.

After the practice, participants performed the first two sessions of the memory game. During the next 4 days, participants were instructed to "play" the memory game at home using a provided laptop and trackball. Participants were instructed to not skip a day and perform the task at roughly the same time of day. We monitored this online using the times the log files were created. Day 6 and 7 were free of any experimental tasks. On Day 8, the participants came back to the institute for the final session. The time of day during the two visits differed by maximally 2 hr to avoid time of day confounds. Day 8 started with the final two parts of the memory task in the MRI scanner. Between the parts, which took both roughly 17 min, there was a break during which the participant could leave the scanner. As a last scan, we acquired a second structural scan. Finally, we conducted the Wisconsin Card Sorting Test and the digit span in a behavioural lab. The total task time on Day 8 was around 1 hr.

#### 2.3 | Memory paradigm

The task mimics the card game "memory/concentration" on two boards of cards and is adapted from (van Buuren et al., 2014) to make it more suitable for children. It is a  $2 \times 2$  design (schema/noschema × paired/new paired associates): one board was the schema while the other was the no-schema condition. Each board contained 80 objects in total. Forty of these objects were learned during the first 4 days (paired associates). The remaining 40 objects (new paired associates) were added on Day 5 and filled the remaining empty positions on each board (see Figure 1 for an illustration of the task). On the schema board the place location associations stayed constant across the whole experiment, whereas on the other, on the no-schema



FIGURF 1 Task design and behavioural performance. (a) Participants needed to learn object-location associations (paired associates) in the memory game. For two boards (one schema, one no-schema board) there were two sets of associations to learn. During the first 4 days, participants learned the paired associates on both boards (40 associations each). For the schema board, participants could thus systematically learn the layout of the board. For the no-schema board at the start of every day, the paired associates switched places with each other, therefore preventing systematic learning. On Day 5, the new paired associates were added (again 40 associations per board). In the final session on Day 8. both the paired and the new paired associates were tested in a recall session in the MRI. The boards had coloured marks in the background to help participants to navigate the board. (b) During a trial, participants first saw an object (cue) at the bottom of the board. After 3 s the box in which the cue was presented turned green and a mouse cursor appeared. Participants then responded within 3 s with the location corresponding to the object. If the response was correct the object was only shown very briefly (0.5 s) whereas if they responded wrongly or not at all the object was shown for 3 s. Each object was repeated three times for participants to have ample opportunity to learn the layout. Additionally, at the start of each session, the whole board (during training the 40 paired associates, during new learning the whole 80 associations) Was presented. (c) During the training phase participants systematically learned the schema paired associates (sPA) on the schema board whereas the performance on the no-schema paired associates (nsPA) on the other board dropped at the start of every day due to the shuffling of locations. The schema new paired associates (sNPA) that were added during the new learning were better learned compared to the no-schema new paired associates (nsNPA) ( $F_{(1,69)} = 59.94$ , p < .001,  $\eta_p^2 = .47$ ). In the recall on Day 8, we observed a reduced performance in the children compared to both older groups ( $F_{(2,69)} = 5.33$ , p = .007,  $\eta_p^2 = .13$ ). Schema benefit refers to how many items participants had correct in the schema new paired associates over the no-schema new paired associates. All error bars indicate the standard error of the mean. A star indicates a significance of p <.05

board, the associations were randomly exchanged every day. Due to this manipulation, participants learned the associations on the schema board over the course of 4 days; forming a schema that contains the object place association of the first 40 objects. Participants could utilise this schema when learning the second set—the new paired associations—of associations on Day 5 for the schema board. The memory for all 160 associations across both boards was tested on Day 8 in the MRI scanner.

## 2.4 | Stimuli design, randomisation, and presentation

In total, we used pictures of 160 everyday objects. To ensure that especially the children could name all the objects effortlessly we selected the objects from a larger set of objects by asking an independent sample (n = 5) of younger children (below the age of 9) to name all the objects. Only objects all children were able to name were included in the set of 160 pictures. The objects were randomly distributed across the different conditions (paired associates or new paired associates on the schema or no-schema board). This randomisation was done individually per participant.

Differing from the initial paradigm (van Buuren et al., 2014) we did not use a  $10 \times 10$  board but a  $9 \times 9$  board, furthermore we arranged the  $9 \times 9$  board into nine visually segregated  $3 \times 3$  boxes, each containing nine cards, by increasing the spacing after each third row and column. These changes had two reasons. First, we aimed to reduce the difficulty and the time required for the task to make it more suitable for children. Second, we opted to have an additional, more sensitive, measure of memory; instead of only taking into account the objects where the chosen position was exact, we can also analyse objects where the response was in the right box. Thus, we would be able to pick on memories where only an approximate location can be recalled. The two boards were differentiated by the colour of the back of the cards that were placed on the board. Whether the schema board or the no-schema board was yellow or blue was randomly assigned per participant in a counterbalanced fashion for each group separately.

We randomised the coordinates of the cards in a pseudorandom fashion separately per participant. Each  $3 \times 3$  box of cards contained either four or five objects per condition (schema and no-schema) to ensure the cards were spread evenly across the board and there was no particular clustering. Furthermore, within each box there could not be a row of three objects, preventing particular easy structures from appearing within the box.

All of the memory tasks were implemented using Presentation version 18.1 (Neurobehavioural Systems, Berkeley, CA). The task at home utilised its web-based licence. For tracking compliance with the study protocol, we used Dropbox (Dropbox Inc., San Francisco, CA) on the laptops to automatically receive the log files. This communication was encrypted with a key only available to the researchers of this study to guarantee participants' privacy.

#### 2.5 | Day 1: basic training (in the lab)

Participants started by learning 40 paired associates on one board. The task started with a 1.5 min presentation of all the 40 objects on their respective location to increase learning speed. After this initial phase, the main task started and only the empty board was visible. One trial consisted of a cue-, a response-, and a feedback phase: participants saw an object at the bottom of the board in a red frame as cue. After 3 s this frame turned green and a cursor appeared at a random position of the board. To draw attention to the location of the cursor there was a short animation (around 120 ms) when it appeared. Participants now had to click within 3 s on the location that belonged to the cued object using the trackball. When the response was correct, the object was shown for 0.5 s at its location. If there was no or a wrong response the cursor turned red and the object was shown at the correct location for 2.5 s. After 40 trials, there was a self-paced pause with a black fixation cross being presented instead of the board. The task consisted of three cycles. During each of those cycles, every object was presented exactly one time. Participants, therefore, had three full training cycles for all items. After the three cycles were completed for the first board, the same procedure was repeated for the second board. Together both sessions took roughly 35 min. We used the laptops and the trackball they would also use at home, showed them how to start the task and explained that the laptop needs an internet connection. To ensure understanding we had participants start the second part of the task themselves. The order of the boards and their colours were randomised and counterbalanced across participants per group.

#### 2.6 | Day 2-4: training (at home)

During each of these 3 days, participants would perform training sessions at home. For all sessions from now onwards the boards were presented in an interleaved fashion. During the initial encoding phase, first, the one board and its 40 objects were presented for 1.5 min, then the other board was presented for 1.5 min. During the task, the board switched every five trials. This interleaved learning was used to reduce interference between the boards (McClelland, McNaughton, & O'Reilly, 1995). The start condition was randomised and counterbalanced across participants. Every day the associations on the noschema board were shuffled as described above, preventing learning across days. Participants were instructed to try as hard as possible to perform well at that board and we showed performance scores at the end of the task to motivate them.

On Day 4, after the task, the training of the 80 paired associates was concluded. Immediately afterwards, participants performed a recall task testing all the paired associates they had learned. The trial structure for the recall was identical to the training except that there was no feedback and there was only one cycle: each object was shown once. The purpose of the recall task was two-fold; first to have a measure how well the paired associations were learned up until now; second, to familiarise the participants with the recall task before they would do the final recall in the scanner on Day 8. Each session took around 35 min with day four being roughly 5 min longer.

#### 2.7 | Day 5: new learning (at home)

On Day 5, the 80 new paired associates were added, 40 to each board. As before, the no-schema board was shuffled. The session started as usual with an initial encoding phase, however, now all 80 objects per board were shown and participants had 3 min per board to memorise as many associations as possible. Aside from the number of associations, the session was identical to the previous training sessions. Each of the 80 objects per board was presented once per cycle leading to 480 trials in total. The boards were again presented in an interleaved fashion and the randomisation was done in such a way that there were never more than two paired associates trials or new paired associates trials in a row. The whole session took approximately 70 min. To reduce effects of exhaustion, participants were instructed to take a more prolonged self-paced break after 240 trials by standing up and moving around in the room, before resuming.

#### 2.8 | Day 8: recall (in the MRI)

Around 72 hr later participants returned to the lab for the final recall in the MRI scanner. Participants lay down in the MRI scanner with the trackball positioned on their abdomen or their right upper thigh at a comfortable distance. Participants familiarised themselves with using the trackball in the scanner. After the participant was proficient in using the trackball, we started with the recall task. One trial started as usual with a cue for 3 s, followed by a response window of 3 s followed by an inter-trial interval with only a black fixation cross on the screen for 2.5–7.5 s. The inter-trial interval was drawn from a uniform distribution. There was no feedback presented during recall. To keep the trial length and the visual input consistent across subjects the board would still be presented for the whole duration of the response window, independent of whether the response was already given. The boards were again interleaved every five trials. We split the task into two parts (balanced across conditions) so that participants could take a break from scanning. Each part took roughly 17 min.

#### 2.9 | Additional measures

Additional to the memory game we also conducted a short verbal memory task (Day 1, before scanning, outside the scanner), a fractal *n*-back task (Ragland et al., 2002; Day 1, in the scanner), the WCST (Heaton et al., 1993; Day 8, after scanning, outside the scanner), and a forward digit span task (Alloway et al., 2008; Day 8, after scanning, outside the scanner). The verbal memory task was used to investigate

links between the cortical thickness of the prefrontal cortex and verbal memory performance. The fractal *n*-back was planned as a control experiment for a planned model-free analysis of the memory game. The WCST was included as an established measure of executive function. Finally, we included the digit span measure to control for group differences unspecific to long-term memory processes.

#### 2.10 | Behavioural analysis

All statistical analyses of behavioural data were conducted in SPSS 21 (IBM Corp, Armonk, NY). Memory performance was measured as the number of correct responses per condition. The memory game consisted of four phases: training (Day 1 till Day 4), recall (Day 4), integration of the paired associates (Day 5), and integration of the complete set (Day 8). The phases were analysed separately. To analyse the training (Day 1 till Day 4) we used a repeated measures ANOVA with the factors schema (schema, no-schema), session (training Day 1-4), cycle (1-3), and group (children, adolescents, and adults). For the recall on Day 4, the repeated measure model included schema and group. For the integration on Day 5, we used a repeated measure model with the factors schema, cycle, group, and included only the new paired associates. The model for the recall on Day 8 was identical to the recall on Day 4 except that we now used the new paired associates instead of the paired associates. Whenever necessary, results were followed up with simple effect tests.

Complementarily, we repeated the analysis using the score for when participants clicked in the correct box (of the nine boxes). This analysis is more sensitive, as responding close to the correct location likely also indicates memory; this heightened sensitivity comes at the cost of a higher chance level (11 vs. 1.25%). We report only significant results with p < .05.

#### 2.11 | MRI data acquisition

Participants were scanned using a Siemens Magnetom Skyra 3T MR scanner equipped with a 32-channel phased-array head coil. The recall task comprised 935 volumes that were acquired using a T2\*-weighted gradient-echo, multiecho echoplanar imaging sequence with the following parameters: TR = 2,100 ms; TE<sub>1</sub> = 8.5 ms, TE<sub>2</sub> = 19.3 ms, TE<sub>3</sub> = 30 ms, TE<sub>4</sub> = 41 ms; flip angle 90°; matrix size = 64 × 64; field of view = 224 mm × 224 mm × 119 mm; voxel size = 3.5 mm × 3.5 mm × 3 mm; slice thickness = 3 mm; slice gap = 0.5 mm; 34 slices acquired in ascending order. As this sequence did not provide whole-brain coverage we oriented the FOV in a way that the hippocampus and the prefrontal cortex were fully inside and that only a small superior part of the parietal lobe was outside the FOV.

For the structural scans we used a T1-weighted magnetisation prepared, rapid acquisition, gradient echo sequence with the following parameters: TR = 2,300 ms; TE = 3.03 ms; flip angle 8°; matrix size =  $256 \times 256$ ; FOV =  $192 \text{ mm} \times 256 \text{ mm} \times 256 \text{ mm}$ ; slice thickness = 1 mm; 192 sagittal slices.

#### 2.12 | MRI pre-processing

Pre-processing was done using a combination of FSL tools (Jenkinson, Beckmann, Behrens, Woolrich, & Smith, 2012), MATLAB (The MathWorks, Natick, MA), and ANTs (Avants et al., 2011). From the two structural scans, we generated an average using rigid body transformations from ANTs (Avants, Tustison, Song, et al., 2011), this procedure removed small movement-induced noise. From the two scans and the average we always selected the scan with the least amount of ringing artefacts for all future analyses. If no difference was visible we used the average scan. These scans were denoised using N4 (Tustison et al., 2010) and generated a study-specific template with an iterative procedure of diffeomorphic registrations (Avants & Gee, 2004). For the registration of the functional volumes, we resampled the created template to a resolution of 3.5 mm isotropic. Using Atropos (Avants, Tustison, Wu, Cook, & Gee, 2011) the anatomical scans were segmented into six tissue classes: cerebrospinal fluid, white matter, cortical grev matter, subcortical grev matter, cerebellum, and brainstem. The segmentation also produced individual brain masks.

For the functional multiecho data, we combined echoes using inhouse build MATLAB scripts. It used the 30 baseline volumes acquired during the resting period directly before each part of the task to determine the optimal weighting of echo-times for each voxel (after applying a smoothing kernel of 3 mm full-width at half-maximum to the baseline volumes), by calculating the contrast-to-noise ratio for each echo per scan. This script also directly realigned the volumes using rigid body transformation. Afterwards, the volumes were smoothed using a 5 mm full-width at half-maximum Gaussian kernel and grand mean intensity normalisation was done by multiplying the time series with a single factor. Younger participants tend to move more than older ones. For a developmental study, it is thus important to minimise the effect of motion in the data. For this purpose, we applied AROMA, a state-of-the-art motion denoising algorithm that uses independent component analysis decomposition of the data to identify movement and other noise signals (Pruim, Mennes, Buitelaar, et al., 2015; Pruim, Mennes, van Rooij, et al., 2015). Variance in the BOLD signal that could only be explained by components identified in this manner was regressed out. Afterwards, we regressed out signal stemming from the cerebrospinal compartments and the white matter by extracting the signal from individual generated segmentations using ANTs (Avants, Tustison, Wu, et al., 2011). As the last step, a 100 s highpass filter was applied.

Boundary-based registration was first calculated from native functional to native structural space using FLIRT (Greve & Fischl, 2009). We then calculated nonlinear registration from native structural space to the study template with FNIRT (Smith et al., 2004). The warping was done in a way that every functional volume was only resliced exactly once after the initial realignment. For displaying the final results, we warped the final maps to MNI space using the nonlinear registration of ANTs (Avants & Gee, 2004).

#### 2.13 | fMRI data analysis

After pre-processing, the data were analysed using the general linear model framework implemented in FEAT (Jenkinson et al., 2012). On the first level, we included eight separate regressors: four regressors modelled correct responses for the separate conditions (paired associates vs. new paired associates on the schema or no-schema board). For the duration, we used the trial onset of the cue until the participant gave a response. As a correct response, we counted if the participant clicked in the correct one of the nine boxes. We used this way of scoring instead of using only the trials in which participants clicked on the correct card as we would have substantially more power due to the higher amount of trials for the MRI analysis while still maintaining a fairly low chance level (11%).

For all those conditions but the schema-paired associates, an additional regressor was included to model incorrect responses. For the schema paired associates condition the performance was designed to be as close to the ceiling as possible leading to only a few incorrect trials. These trials were modelled together with all the trials in which subjects failed to respond in time in a single "miss" regressor. For the miss trials, the full 6 s of the cue and response window was used. Regressors were then convolved with a double gamma hemodynamic response function. All regressors also included their temporal derivative. On the first level, the model was fitted separately per run. Using fixed effect modelling the runs were combined per subject and then the participant-specific contrasts were estimated. To calculate the group level statistics, we warped the participant level results into study template space and used mixed effect modelling implemented in FSL FLAME2. The results were thresholded using a cluster forming threshold of z > 2.3 (equal to p < .01) and a cluster significance threshold of p < .05 at the whole-brain level. Our central motivation for this study was to understand how the neural mnemonic processes differ across different stages of cortical maturation. Therefore, our imaging analysis was centrally guided by the behavioural results, illuminating the underlying neural architecture related to behavioural differences. Thus, the contrasts used will be explained while presenting the imaging results.

As a follow-up analysis, we conducted moderation analysis based on the results of the general linear model analysis. For this, we extracted the average betas from the significant clusters on a participant-by-participant basis. We then conducted the moderation analysis using the PROCESS macro (Hayes, 2013) for SPSS. This analysis tested whether age group (as a categorical variable) influenced the relationship between the extracted betas (representing neural activation) and memory performance. The moderation analysis used a regression model adding interaction terms to model the moderation.

Due to the scope of the study, we present the results of connectivity analysis in a separate article (Müller et al., 2020). That article focuses on the relation of the mPFC and its interaction with the default mode network in this dataset.

#### 3 | RESULTS

#### 3.1 | Training

As expected, during the recall on day four schema items were better recalled than no-schema items ( $F_{(1,69)} = 199.05$ , p < .001,  $\eta_p^2 = .74$ ). For the training, we observed a significant three-way interaction of schema × session × cycle ( $F_{(4.49,309.86)} = 27.84$ , p < .001,  $\eta_p^2 = .29$ ), reflecting the fact that in the schema condition the paired associates could be learned across days while the shuffling between days prevented this for the no-schema paired associates (Figure 1c).

#### 3.2 | New learning

The schema new paired associates were learned better compared to the no-schema paired associates ( $F_{(1,69)} = 59.94$ , p < .001,  $\eta_p^2 = .47$ ).

#### 3.3 | Recall Day 8

In the final recall, the schema new paired associates were better retrieved compared to the new no-schema paired associates  $(F_{(1,69)} = 17.09, p < .001, \eta_p^2 = .2)$ . However, there was a significant effect of group on the retrieval performance of the new paired associates overall  $(F_{(2,69)} = 5.33, p = .007, \eta_p^2 = .13)$ . Children performed worse than adolescents (MD = -5.97, p = .006) and adults (MD = -5.23, p = .005); whereas adolescents and adults did not differ significantly (MD = -.31, p = .881).

#### 3.4 | Difference between runs

A further control analysis showed a difference between the two runs with the second run being worse in performance ( $F_{(1,69)} = 2,361.9$ , p < .001). Importantly, this difference did not significantly differ between groups ( $F_{(1,69)} = 1.3$ , p = .28).

#### 3.5 | Precise location correct versus box correct

Analysing the data counting only trials as correct where the response was on the correct card instead of the correct box (3  $\times$  3 cards) did not alter the results: All of the reported effects were also significant for the box score.

#### 3.6 | fMRI: developmental differences

Our central behavioural finding is that children show lower memory performance than adolescents and adults while the latter two groups did not perform significantly differently. To understand the neural changes across development, we contrasted the activation during retrieval of the new paired associates for the correctly retrieved trials minus the trials in which a wrong response was given. As all groups seem to have profited to a similar degree from schema, we averaged across schema and no-schema trials to be more sensitive for developmental differences. The contrast between hits and misses was then compared between the children versus the average of the two older groups; this was done because the latter two groups did not differ in performance. As we used mixed modelling on the group level analysis the fact that the adolescent-adult group is twice as big as the children does not bias the results.

We observed increased activation in children in midline structures, including the dorsomedial prefrontal cortex (dmPFC). Increased activation in the adolescent-adult groups was most pronounced bilaterally in the angular gyrus (Figure 2). For a complete list of all clusters please see Table 1.

We further investigated this contrast in the medial temporal lobe region using a small volume correction. We included the hippocampus and parahippocampal gyrus bilaterally in the mask, none of the regions showed a significant activation.

As we hypothesised that performance differences might be due to differences in executive abilities in children, we tested whether there is a link between the (de)activation of the dmPFC and measures of executive function (Figure 2). Activation in the dmPFC was negatively correlated with performance in the WCST (r[70] = -.31, p = .008), as measured by the number of correct categories, but not significantly related to the forward digit span score (r[70] = .04, p = .72). To test the statistical difference between the magnitude of the dmPFC activation and WSCT performance correlational analyses were performed. The correlation of dmPFC activation and WCST performance was driven by a negative correlation across the two adult groups (r[49] = -.31, p = .026). This association between dmPFC activity and WCST performance for the children was reduced compared to the adolescent-adult group (z = 2.08, p = .038) and in itself not significant (r[19] = .26, p = .27).

#### 3.7 | fMRI: schema effect

Participants across all age groups remembered schema new associates better than the no-schema new associates. To illuminate the neural architecture behind this schema effect we calculated the contrast between the hits and the misses between the schema new paired associates (sNPA) and the no-schema new paired associates (nsNPA): sNPA (hits – misses) – nsNPA (hits – misses). However, there was no significant activation that survived whole-brain correction. More specifically, we tested the angular gyrus, as the region previously been found to be important for integrating different parts of a schema (Wagner et al., 2015) and in the paradigm utilised here in the same schema  $\times$  memory contrast (van Buuren et al., 2014). When separately contrasting sNPA (hits – misses) and nsNPA (hits – misses), we observed that both angular gyri were significantly activated in both contrasts (p <.05). To test whether there was activation specificity for schema, we extracted the betas for voxels that were significantly

MÜLLER ET AL.



**FIGURE 2** Age-related differences in mean memory performance for the new paired associates. During the recall of both the schema and the no-schema, new paired associates children showed increased activation in the dorsal medial prefrontal cortex (dmPFC) overlapping with the cingulate and paracingulate gyrus; a second cluster around the lateral occipital cortex showed the same effect. Adolescents and adults showed higher bilateral activation of the angular gyrus

activated for the sNPA's; we extracted both the values for sNPA (hits – misses) and nsNPA (hits – misses) (Figure 3). The difference between those contrasts was positively correlated (r[70] = .34, p = .003) to the magnitude of the schema benefit. There was no indication that this relation is significantly modulated by age group (F<sub>(2.66)</sub> = .95, p = .39).

#### 4 | DISCUSSION

We tested how differences in the associative and the strategic component of the two-component model of memory development (Shing et al., 2008; Shing, Lindenberger, et al., 2010; Shing, Werkle-Bergner, et al., 2010) contribute to memory performance differences between children, adolescents, and adults. We found that both the adolescent and adult group had higher memory performance than children at mid- to late childhood, independent of the conditions, while all groups profited equally from utilising schemas (Figure 1). Performance differences between groups were associated with deactivation of the dmPFC, which in turn was linked to executive function. In contrast, activation of the angular gyrus was consistently correlated with memory performance across all groups (Figure 3). This suggests that agerelated differences in memory are rather driven by differences in the strategic component, but not the associative component.

The two-component model helps us to test whether the agerelated differences we observed are driven by the immaturity of the associative or the strategic component. Memory differences linked to associative regions, such as the angular gyrus, or to the utilisation of schema would indicate differences in the associative component. Memory differences that are not linked to the associative memory regions but rather to regions involved in executive function would suggest a stronger role for the strategic component. To corroborate the links between task activation and executive function we used the independently acquired WCST performance as a general measurement of executive function (Greve, Stickle, Love, Bianchini, & Stanford, 2005): Participants with high levels of executive function in the WCST can likely use those functions strongly to facilitate their retrieval performance.

Children had a lower retrieval performance than adolescents and adults for the schema and the no-schema new paired associates. We found for both the adolescents and the adults, the level of the deactivation of the dmPFC during trials in which they recalled the correct location predicted their overall recall performance: the stronger the deactivation was, the better was the performance (Figure 3). This deactivation also predicted the WCST performance. Furthermore, the dmPFC cluster we found overlaps with a core cluster previously observed during performance of the WCST (Specht, Lie, Shah, & Fink, 2009) and is also contained within the Executive Control Network, a resting state network that is involved across many aspects of executive function (Smith et al., 2009). The link to the WCST, the involvement of our dmPFC cluster in the WCST and the dmPFC's important role in executive function (Domenech & Koechlin, 2015; Ridderinkhof, 2004), suggests to us that it reflects executive function benefitting retrieval performance. Participants with strong deactivation in the dmPFC could use executive function to improve their task performance, whereas participants that showed little or no deactivation could not. In contrast to the other groups, children did not seem to exhibit this behaviour: they neither showed a systematic

#### TABLE 1 Developmental differences in activation for the correct retrieval of the new paired associates

	MNI coordinates				
Region	x	у	z	z-Score	Voxels
Children > adolescents + adults					
R dorsomed. prefrontal cort.	3	15	46	4.04	243
L dorsomed. prefrontal cort	-7	-14	43	3.74	
R dorsomed. prefrontal cort	4	22	42	3.72	
L precentral gyrus	-40	3	40	3.54	
R middle frontal gyrus	-25	2	55	3.48	
L dorsomed. prefrontal cort	-4	5	55	3.34	
R lat. sup. occipital cort	25	-67	41	4.73	151
R precuneus	15	-58	25	3.5	
R lat. sup. occipital cort	42	-83	24	3.34	
R lat. sup. occipital cort	30	-80	29	3.3	
R Precuneus	9	-59	9	3.26	
R med. occipital cort	38	-76	19	3.17	
L lat. sup. occipital cort	-29	-84	32	4.2	103
L lat. sup. occipital cort	-26	-81	24	3.9	
L lat. sup. occipital cort	-30	-74	22	3.82	
L precuneus	-15	-72	38	3.39	
L precuneus	-19	-81	35	2.68	
L lat. sup. occipital cort	-45	-85	27	2.54	
Adolescents + adults > children					
L lat. sup. Occipital cort	-54	-61	44	4.89	227
L angular gyrus	-51	-59	32	4.14	
L lat. sup. occipital cort	-46	-72	38	4.01	
L par. operculum	-63	-34	22	3.85	
L lat. sup. Occipital cort	-58	-64	16	3.61	
L mid. temp. gyrus	-69	-48	6	2.83	
R lat. sup. occipital cort	60	-61	33	3.44	227
R lat. sup. occipital cort	48	-61	42	3.34	
R lat. sup. occipital cort	56	-60	42	3.27	
R angular gyrus	64	-53	25	3.07	
R angular gyrus	54	-47	23	2.82	
R lat. sup. occipital cort	60	-61	21	2.8	103

*Note:* The listed clusters here and their local maxima show differences between the children, the adolescent, and the adult groups for the retrieval of the new paired associates in which both older groups outperformed the children. The coordinates were always of the global/local maximum. The voxel-count as well as the *z*-score of the peak voxel were taken from study space. The MNI coordinates were obtained by warping the results into MNI space. All labels refer to regions on the cortex.

deactivation of the dmPFC nor was the dmPFC activity related to memory or WCST performance, in which children performed worse than adolescents and adults. We take all this as an indication that the strategic component in children is not as mature as in adolescents and adults: Whereas adolescents and adults can use their strategic abilities to enhance their memory performance, children did not seem to be able to do this. These results are nicely in line with work demonstrating that age-related increases in mnemonic strategies are linked to the development of the PFC in general (Yu et al., 2018). Furthermore, Yu and colleagues observed that especially the decrease in the volume of the right mPFC explained the relation between age related increase in strategic abilities. A recent study by Brod, Breitwieser, Hasselhorn, and Bunge (2020) confirms the link between prior knowledge and executive functions with demonstrating that children with better inhibitory control processes benefit from activating a prior schema.

The mPFC has consistently found to be involved in executive functions and prior knowledge, but no agreement exists on the precise role of subregions within the mPFC. Brod and Shing (2019)



**FIGURE 3** Developmental differences in brain-behaviour relation. For both the activation in the dmPFC and the angular gyri we found a relation to the mean memory performance across the age groups. For the dmPFC this relation was negative (r[70] = -.63, p <.001). For the angular gyri it was positive (r[70] = .75, p <.001). Most notably we found an age-related dissociation: whereas the brain-behaviour relation was consistent across age for the angular gyri; activation in the dmPFC showed a moderation with age  $F_{(1,68)} = 4.19$ , p = .045). Participants in both adult groups varied in the degree they deactivated the dmPFC, the stronger the deactivation the better the performance. Children showed neither a deactivation of the dmPFC nor a relation to recall performance. Mean memory performance refers to the average across the schema and no-schema paired associates. A star indicates a significance of p <.05

explored the ventral mPFC in three groups of primary school children, young adults, and older adults. Their findings indicate that the vmPFC plays a key role in schema-related memory processes across the entire life span. A study by Yu et al. (2018) found evidence that the dorsolateral PFC explains memory development, whereas Müller et al. (2020) could not identify a clear region within the mPFC. Further research is necessary to pinpoint the precise regions within the mPFC related to executive functioning and schema-related memory processing.

With regard to the associative component, we did not find any indications for differences between age-groups. The activity of the angular gyrus was correlated with successful memory performance consistently across groups. Additionally, the schema effect was indistinguishable across groups. All groups performed between 20 and 30% better in the schema over the no-schema condition. We interpret this absence of any developmental differences for associative processes as an indication for a weaker role of the associative component to explain age-related memory differences in our sample.

The consistent relation of the activation from the angular gyrus across groups suggests an important role in the task that is stable across the tested ages. This stability is consistent with previous work demonstrating that the angular gyrus has the same functional boundaries in school children (7-10 years old) compared to adults (Barnes et al., 2012); suggesting a relative early functional maturation. In recent years, the contribution of the angular gyrus to memory has received increased attention. There is now substantial evidence for it being an amodal convergence zone (Bonnici, Richter, Yazar, & Simons, 2016; Yazar, Bergström, & Simons, 2017) that integrates input from different modalities to create higher-level representations. With this facility, it lays the basis for abstract representations and thus semantic memory (Binder & Desai, 2011). The ability to combine several modalities seems ideally suited for the memory game task where spatial information (location) needs to be combined with semantic information (identity of the card). Another capacity of the angular gyrus that explains its involvement is the ability to guide attention during memory retrieval relying on retrieval cues or recovered memories (Cabeza, Ciaramelli, Olson, & Moscovitch, 2008; Vilberg & Rugg, 2008).

We replicated that schemas facilitate memory (Liu, Grady, & Moscovitch, 2017; Tse et al., 2007; van Buuren et al., 2014; van Kesteren, Rijpkema, Ruiter, & Fernandez, 2010) as indicated by the higher performance for the schema new paired associates. This effect did not show any differences across development, in line with a

previous study investigating children in a similar age range (Brod et al., 2017). A recent fMRI study by Brod and Shing (2019) could confirm this finding showing a strong, age-independent memory benefit for items encoded in a schema context as compared to non-schema items. While the behavioural results show a schema benefit on memory the fMRI findings diverge and show no schema-related memory difference in neural activity. A possible explanation could be that the subsequent memory effect is not sensitive enough to detect a schema by memory interaction. An alternative explanation is that the processes of memory formation are the same for schema congruent and incongruent trials, but that the likelihood that they occur are different (Uncapher & Rugg, 2005).

Neurally, we found that not the mPFC but the angular gyrus distinguished the retrieval of schema versus no-schema associations. Both the angular gyrus and the mPFC were activated in both the schema and no-schema condition, however, the angular gyrus was significantly more strongly activated whereas the activity of the mPFC did not differ significantly. This pattern is consistent with results previously found using this paradigm (van Buuren et al., 2014), but it appears at odds with the typical pattern that the mPFC orchestrates the utilisation of schemas (Fernández, 2017; Genzel & Battaglia, 2017; van Kesteren et al., 2012). We speculate that the mPFC did not differentially activate as there were too many associations at the same time that needed to be assimilated into the schema. If either there would have been fewer associations to learn or there would have been more time for learning the associations and stabilising their memories, we speculate that the mPFC would have been more strongly activated for the correctly retrieved schema new paired associates.

In summary, we investigated whether memory differences between children aged 10–12, adolescents, and adults would stem from developmental changes in executive abilities, the strategic component, or rather from differences in mechanisms related to binding different features together into a memory representation, the associative component. We found that adolescents and adults outperformed children in memory. The performance within the adolescents and adult group was associated with their individual executive abilities, thus providing evidence that maturation of the strategic component was driving the age-related differences we observed. In contrast, we did not find differences in the associative component that help to explain the differences in memory between the age groups.

#### ACKNOWLEDGEMENT

This research was funded by an NWO Research Talent Grant (406-13-008) to Nils C. J. Müller and Guillén Fernández.

#### CONFLICT OF INTEREST

The authors declare that they have no conflict of interest

#### DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

#### ORCID

Gabriele Janzen D https://orcid.org/0000-0002-5082-1777 Guillén Fernández D https://orcid.org/0000-0002-5522-0604

#### REFERENCES

- Alloway, T. P., Gathercole, S. E., Kirkwood, H., & Elliott, J. (2008). Evaluating the validity of the Automated Working Memory Assessment. *Educational Psycholology*, 28, 725–734.
- Avants, B. B., & Gee, J. C. (2004). Geodesic estimation for large deformation anatomical shape averaging and interpolation. *NeuroImage*, 23, 139–150.
- Avants, B. B., Tustison, N. J., Song, G., Cook, P. A., Klein, A., & Gee, J. C. (2011). A reproducible evaluation of ANTs similarity metric performance in brain image registration. *NeuroImage*, 54, 2033–2044. https://doi.org/10.1016/j.neuroimage.2010.09.025
- Avants, B. B., Tustison, N. J., Wu, J., Cook, P. A., & Gee, J. C. (2011). An open source multivariate framework for N-tissue segmentation with evaluation on public data. *Neuroinformatics*, 9, 381–400.
- Barnes, K. A., Nelson, S. M., Cohen, A. L., Power, J. D., Coalson, R. S., Miezin, F. M., ... Schlaggar, B. L. (2012). Parcellation in left lateral parietal cortex is similar in adults and children. *Cerebral Cortex*, 22, 1148– 1158.
- Bartlett, F. C. (1932). Remembering: A study in experimental and social psychology. Cambridge: Cambridge University.
- Binder, J. R., & Desai, R. H. (2011). The neurobiology of semantic memory. Trends in Cognitive Sciences, 15, 527–536.
- Bonnici, H. M., Richter, F. R., Yazar, Y., & Simons, J. S. (2016). Multimodal feature integration in the angular gyrus during episodic and semantic retrieval. *Journal of Neuroscience*, 36, 5462–5471.
- Brod, G., Breitwieser, J., Hasselhorn, M., & Bunge, S. A. (2020). Being proven wrong elicits learning in children—But only in those with higher executive function skills. *Developmental Science*, 23, e12916.
- Brod, G., Lindenberger, U., & Shing, Y. L. (2017). Neural activation patterns during retrieval of schema-related memories: Differences and commonalities between children and adults. *Developmental Science*, 20, e12475.
- Brod, G., & Shing, Y. L. (2019). A boon and a bane: Comparing the effects of prior knowledge on memory across the lifespan. *Developmental Psychology*, 55, 1326–1337.
- Brod, G., Werkle-Bergner, M., & Shing, Y. L. (2013). The influence of prior knowledge on memory: A developmental cognitive neuroscience perspective. *Frontiers Behavioral Neuroscience*, 7, 1–13.
- Bunge, S., & Crone, E. (2009). Neural correlates of the development of cognitive control. *Neuroscience Research*, 55, 22–37. https://doi.org/ 10.1017/CBO9780511757402.005
- Cabeza, R., Ciaramelli, E., Olson, I. R., & Moscovitch, M. (2008). The parietal cortex and episodic memory: An attentional account. *Nature Reviews Neuroscience*, *9*, 613–625.
- Crone, E. A., & Dahl, R. E. (2012). Understanding adolescence as a period of social—Affective engagement and goal flexibility. *Nature*, 13, 636–650.
- DeMaster, D. M., & Ghetti, S. (2013). Developmental differences in hippocampal and cortical contributions to episodic retrieval. *Cortex*, 49, 1482–1493.
- Domenech, P., & Koechlin, E. (2015). Executive control and decisionmaking in the prefrontal cortex. *Current Opinion in Behavioral Sciences*, 1, 101–106.
- Fernández, G. (2017). The medial prefrontal cortex is a critical hub in the declarative memory system. In *Cognitive neuroscience of memory consolidation* (pp. 45–56). New York, NY: Springer.
- Geng, F., Redcay, E., & Riggins, T. (2019). The influence of age and performance on hippocampal function and the encoding of contextual information in early childhood. *NeuroImage*, 15, 433–443.

### 6012 WILEY-

- Genzel, L., & Battaglia, F. P. (2017). Cortico-hippocampal circuits for memory consolidation: The role of the prefrontal cortex. In *Cognitive neuroscience of memory consolidation* (pp. 265–281). New York, NY: Springer.
- Ghetti, S., & Bunge, S. A. (2012). Neural changes underlying the development of episodic memory during middle childhood. *Developmental Cognitive Neuroscience*, 2, 381–395.
- Ghetti, S., DeMaster, D. M., Yonelinas, A. P., & Bunge, S. A. (2010). Developmental differences in medial temporal lobe function during memory encoding. *Journal of Neuroscience*, 30, 9548–9556.
- Gogtay, N., Giedd, J. N., Lusk, L., Hayashi, K. M., Greenstein, D., Vaituzis, C., ... Thompson, P. M. (2004). Dynamic mapping of human cortical development during childhood through early adulthood. Proceedings of the National Academy of Sciences, 101, 8174–8179.
- Greve, D. N., & Fischl, B. (2009). Accurate and robust brain image alignment using boundary-based registration. *NeuroImage*, 48, 63–72.
- Greve, K. W., Stickle, T. R., Love, J. M., Bianchini, K. J., & Stanford, M. S. (2005). Latent structure of the Wisconsin card sorting test: A confirmatory factor analytic study. Archives of Clinical Neuropsychology, 20, 355–364.
- Hayes, A. (2013). Introduction to mediation, moderation, and conditional process analysis (pp. 3–4). New York, NY: Guilford.
- Heaton, R., Chelune, G., Curtiss, G., Kay, G., & Talley, J. (1993). Wisconsin card sorting test manual: Revised and expanded. Odessa, FL: Psychological Assessment Resources Inc.
- Ismail, F. Y., Fatemi, A., & Johnston, M. V. (2016). Cerebral plasticity: Windows of opportunity in the developing brain. *European Journal of Paediatric Neurology*, 21, 23–48.
- Jenkinson, M., Beckmann, C. F., Behrens, T. E. J., Woolrich, M. W., & Smith, S. M. (2012). FSL. *Neuroimage*, 62, 782–790.
- Keresztes, A., Bender, A. R., Bodammer, N. C., Lindenberger, U., Shing, Y. L., & Werkle-Bergner, M. (2017). Hippocampal maturity promotes memory distinctiveness in childhood and adolescence. *Proceedings of the National Academy of Sciences*, 114, 9212–9217.
- Keresztes, A., Ngo, C. T., Lindenberger, U., Werkle-Bergner, M., & Newcombe, N. S. (2018). Hippocampal maturation drives memory from generalization to specifity. *Trends in Cognitive Sciences*, 22, 676–686.
- Liu, Z. X., Grady, C., & Moscovitch, M. (2017). Effects of prior-knowledge on brain activation and connectivity during associative memory encoding. *Cerebral Cortex*, 27, 1991–2009.
- Luna, B., Marek, S., Larsen, B., Tervo-Clemmens, B., & Chahal, R. (2015). An integrative model of the maturation of cognitive control. *Annual Reviews Neuroscience*, 38, 151–170.
- Maril, A., Avital, R., Reggev, N., Zuckerman, M., Sadeh, T., Sira, L. B., & Livneh, N. (2011). Event congruency and episodic encoding: A developmental fMRI study. *Neuropsychologia*, 49, 3036–3045.
- Maril, A., Davis, P. E., Koo, J. J., Reggev, N., Zuckerman, M., Ehrenfeld, L., ... Rivkinet, M. J. (2010). Developmental fMRI study of episodic verbal memory encoding in children. *Neurology*, 75, 2110–2116.
- McClelland, J. L., McNaughton, B. L., & O'Reilly, R. C. (1995). Why there are complementary learning systems in the hippocampus and neocortex: Insights from the successes and failures of connectionist models of learning and memory. *Psychology Review*, 102, 419–457.
- Miller, E. K., & Cohen, J. D. (2001). An integrative theory of prefrontal cortex function. Annual Reviews Neuroscience, 24, 167–202.
- Müller, N. C. J., Dresler, M., Janzen, G., Beckmann, C. F., Fernández, G., & Kohn, N. (2020). Medial prefrontal decoupling from the default mode network benefits memory. *NeuroImage*, 210, 116543.
- Ofen, N., & Shing, Y. L. (2013). From perception to memory: Changes in memory systems across the lifespan. *Neuroscience Biobehavioral Review*, 37, 2258–2267.
- Ofen, N., Tanga, L., Yua, Q., & Johnson, E. L. (2019). Memory and the developing brain: From description to explanation with innovation in methods. *Developmental Cognitive Neuroscience*, 36, 100613.
- Piaget, J. (1936). Origins of intelligence in children. London: Routledge & Kegan Paul.

- Pruim, R. H. R., Mennes, M., Buitelaar, J. K., & Beckmann, C. F. (2015). Evaluation of ICA-AROMA and alternative strategies for motion artifact removal in resting state fMRI. *NeuroImage*, 112, 278–287.
- Pruim, R. H. R., Mennes, M., van Rooij, D., Llera, A., Buitelaar, J. K., & Beckmann, C. F. (2015). ICA-AROMA: A robust ICA-based strategy for removing motion artifacts from fMRI data. *NeuroImage*, 112, 267–277.
- Ragland, J. D., Turetsky, B. I., Gur, R. C., Gunning-Dixon, F., Turner, T., Schroeder, L., ... Gur, R. E. (2002). Working memory for complex figures: An fMRI comparison of letter and fractal n-back tasks. *Neuropsychology*, 16, 370–379.
- Ridderinkhof, K. R. (2004). The role of the medial frontal cortex in cognitive control. *Science*, 306, 443–447.
- Shaw, P., Kabani, N. J., Lerch, J. P., Eckstrand, K., Lenroot, R., Gogtay, N., ... Wise, S. P. (2008). Neurodevelopmental trajectories of the human cerebral cortex. *Journal of Neuroscience*, 28, 3586–3594.
- Shing, Y. L., Lindenberger, U., Diamond, A., Li, S.-C., & Davidson, M. C. (2010). Memory maintenance and inhibitory control differentiate from early childhood to adolescence. *Developmental Neuropsychology*, 35, 679–697.
- Shing, Y. L., Werkle-Bergner, M., Brehmer, Y., Müller, V., Li, S.-C., & Lindenberger, U. (2010). Episodic memory across the lifespan: The contributions of associative and strategic components. *Neuroscience Biobehavioral Review*, 34, 1080–1091.
- Shing, Y. L., Werkle-Bergner, M., Li, S.-C., & Lindenberger, U. (2008). Associative and strategic components of episodic memory: A life-span dissociation. *Journal of Experimental Psychology General*, 137, 495–513.
- Simons, J. S., & Spiers, H. J. (2003). Prefrontal and medial temporal lobe interactions in long-term memory. *Nature Reviews Neuroscience*, 4, 637–648. https://doi.org/10.1038/nrn1178
- Smith, S. M., Fox, P. T., Miller, K. L., Glahn, D. C., Fox, P. M., Mackay, C. E., ... Beckmann, C. F. (2009). Correspondence of the brain's functional architecture during activation and rest. *Proceedings of the National Academy of Sciences*, 106, 13040–13045.
- Smith, S. M., Jenkinson, M., Woolrich, M. W., Beckmann, C. F., Behrens, T. E. J., Johansen-Berg, H., ... Matthews, P. M. (2004). Advances in functional and structural MR image analysis and implementation as FSL. *NeuroImage*, 23, S208–S219.
- Specht, K., Lie, C. H., Shah, N. J., & Fink, G. R. (2009). Disentangling the prefrontal network for rule selection by means of a non-verbal variant of the Wisconsin card sorting test. *Human Brain Mapping*, 30, 1734– 1743.
- Thorndyke, P. W., & Hayes-Roth, B. (1979). The use of schemata in the acquisition and transfer of knowledge. *Cognitive Psychology*, 11, 82–106.
- Tse, D., Langston, R. F., Kakeyama, M., Bethus, I., Spooner, P. A., Wood, E. R., ... Morris, R. G. M. (2007). Schemas and memory consolidation. *Science*, 316, 76–82.
- Tse, D., Takeuchi, T., Kakeyama, M., Kajii, Y., Okuno, H., Tohyama, C., ... Morris, R. G. M. (2011). Schema-dependent gene activation and memory encoding in neocortex. *Science*, 333, 891–895.
- Tustison, N. J., Avants, B. B., Cook, P., Zheng, Y., Egan, A., Yushkevich, P., & Gee, J. C. (2010). N4ITK: Improved N3 bias correction. *IEEE Transactions on Medical Imaging*, 29, 1310–1320.
- Uncapher, M. R., & Rugg, D. R. (2005). Effects of divided attention on fMRI correlates of memory encoding, effects of divided attention on fMRI correlates of memory encoding. *Journal of Cognitive Neuroscience*, 17, 1923–1935. https://doi.org/10.1162/089892905775008616
- Vakil, E., & Blachstein, H. (2007). Rey AVLT: Developmental norms for adults and the sensitivity of different memory measures to age. *Clinical Neuropsychology*, 7049, 37–41.
- van Buuren, M., Kroes, M. C. W., Wagner, I. C., Genzel, L., Morris, R. G. M., & Fernandez, G. (2014). Initial investigation of the effects of an experimentally learned schema on spatial associative memory in humans. *Journal of Neuroscience*, 34, 16662–16670.
- van Kesteren, M. T. R., Rijpkema, M., Ruiter, D. J., & Fernandez, G. (2010). Retrieval of associative information congruent with prior knowledge is

related to increased medial prefrontal activity and connectivity. *Journal* of Neuroscience, 30, 15888–15894.

- van Kesteren, M. T. R., Ruiter, D. J., Fernández, G., & Henson, R. N. (2012). How schema and novelty augment memory formation. *Trends in Neuroscience*, 35, 211–219.
- Vilberg, K. L., & Rugg, M. D. (2008). Memory retrieval and the parietal cortex: A review of evidence from a dual-process perspective. *Neuropsychologia*, 46, 1787–1799.
- Wagner, I. C., van Buuren, M., Kroes, M. C. W., Gutteling, T. P., van der Linden, M., Morris, R. G., & Fernández, G. (2015). Schematic memory components converge within angular gyrus during retrieval. *eLife*, 4, 1–28.
- Yazar, Y., Bergström, Z. M., & Simons, J. S. (2017). Reduced multimodal integration of memory features following continuous theta burst stimulation of angular gyrus. *Brain Stimulation*, 10, 1–16.

Yu, Q., McCall, D. M., Homayouni, R., Tang, L., Chen, Z., Schoff, D., ... Ofen, N. (2018). Age-associated increase in mnemonic strategy use is linked to prefrontal cortex development. *NeuroImage*, 181, 162–169.

How to cite this article: Müller, N. C. J., Kohn, N., van Buuren, M., Klijn, N., Emmen, H., Berkers, R. M. W. J., Dresler, M., Janzen, G., & Fernández, G. (2021). Differences in executive abilities rather than associative processes contribute to memory development. *Human Brain Mapping*, 42(18), 6000–6013. https://doi.org/10.1002/hbm.25665