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This will open up a panel down the right side of the document. The majority of tools you will use for annotating your proof will be in the Annotations section, pictured opposite. We’ve picked out some of these tools below:

1. **Replace (Ins) Tool** — for replacing text.
   - Strikes a line through text and opens up a text box where replacement text can be entered.
   - **How to use it**
     - Highlight a word or sentence.
     - Click on the Replace (Ins) icon in the Annotations section.
     - Type the replacement text into the blue box that appears.

2. **Strikethrough (Del) Tool** — for deleting text.
   - Strikes a red line through text that is to be deleted.
   - **How to use it**
     - Highlight a word or sentence.
     - Click on the Strikethrough (Del) icon in the Annotations section.

3. **Add note to text Tool** — for highlighting a section to be changed to bold or italic.
   - Highlights text in yellow and opens up a text box where comments can be entered.
   - **How to use it**
     - Highlight the relevant section of text.
     - Click on the Add note to text icon in the Annotations section.
     - Type instruction on what should be changed regarding the text into the yellow box that appears.

4. **Add sticky note Tool** — for making notes at specific points in the text.
   - Marks a point in the proof where a comment needs to be highlighted.
   - **How to use it**
     - Click on the Add sticky note icon in the Annotations section.
     - Click at the point in the proof where the comment should be inserted.
     - Type the comment into the yellow box that appears.
5. **Attach File Tool** – for inserting large amounts of text or replacement figures.

Inserts an icon linking to the attached file in the appropriate place in the text.

**How to use it**

- Click on the Attach File icon in the Annotations section.
- Click on the proof to where you’d like the attached file to be linked.
- Select the file to be attached from your computer or network.
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6. **Drawing Markups Tools** – for drawing shapes, lines and freeform annotations on proofs and commenting on these marks.

Allows shapes, lines and freeform annotations to be drawn on proofs and for comment to be made on these marks.

**How to use it**

- Click on one of the shapes in the Drawing Markups section.
- Click on the proof at the relevant point and draw the selected shape with the cursor.
- To add a comment to the drawn shape, move the cursor over the shape until an arrowhead appears.
- Double click on the shape and type any text in the red box that appears.
The ontogeny of relational memory and pattern separation

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Abstract

Episodic memory relies on memory for the relations among multiple elements of an event and the ability to discriminate among similar elements of episodes. The latter phenomenon, termed pattern separation, has been studied mainly in young and older adults with relatively little research on children. Building on prior work with young children, we created an engaging computer-administered relational memory task assessing what-where relations. We also modified the Mnemonic Similarity Task used to assess pattern discrimination in young and older adults for use with preschool children. Results showed that 4-year-olds performed significantly worse than 6-year-olds and adults on both tasks, whereas 6-year-olds and adults performed comparably, even though there were no ceiling effects. However, performance on the two tasks did not correlate, suggesting that two distinct mnemonic processes with different developmental trajectories may contribute to age-related changes in episodic memory.

RESEARCH HIGHLIGHTS

- Pattern separation, indexed by a new measure of mnemonic lure discrimination suitable for children, undergoes changes between the ages of 4 and 6, and continues to develop into adulthood.
- Relational memory also improves significantly between the ages of 4 and 6, with a non-significant difference between 6-year-olds and young adults.
- Pattern separation and relational memory may be separable cognitive components of episodic memory, with developmental changes in each likely contributing to episodic memory gains in early childhood.

1 | INTRODUCTION

Episodic memory is memory for events that occur within a specific spatiotemporal context (Tulving, 2002). For example, I can remember that the last time I visited my favorite restaurant with a friend, we sat by the window and I had salmon and she ordered steak, but that the time before, we sat in a back booth and both ordered steak.

Adults are better able to retrieve episodic memories like these than children younger than the age of 6 (Rubin, 2000), and some children fail to distinguish between real events and imagined events (Sluzenski, Newcombe, & Ottinger, 2004).

Remembering and discriminating between similar events presents at least two distinct challenges. First, I need to relate the various aspects of each event to form a cohesive episode (e.g., window seat with ate salmon). That is, I need to bind different elements, such as what, where, and when the event occurred, to form a memory of a complex event – that is, relational memory (Cohen & Eichenbaum, 1993; Eichenbaum & Cohen, 2001; Schacter & Tulving, 1994). Second, I need to discriminate between similar events and elements within them (e.g., type of steak at time 1 was rib-eye, type of steak at time 2 was sirloin). That is, I need to distinguish similar memories from one another – pattern separation (Complementary Learning Systems: Norman, 2010; Norman & O’Reilly, 2003). These two component processes of episodic memory are the focus of this paper.

1.1 | Relational memory

Episodic memory requires the formation of relational structures that bind information to the specific context. Thus, the ability to form, retain, and retrieve relational information has been thought to be a crucial component of episodic memory (Eichenbaum & Cohen, 2001). Relational memory is typically tested using paired-associates tasks, which assess memory for the co-occurrence of multiple items that
relational memory undergoes protracted development, whereas memory for single items develops relatively early and improves gradually (e.g., Lloyd, Doydum, & Newcombe, 2009; Riggins, 2014; Sluzenski, Newcombe, & Kovacs, 2006; Sluzenski et al., 2004). When asked to remember either single items (e.g., a tiger, a library) or unique item-place pairs (e.g., a tiger and a library), 6-year-olds outperformed 4-year-olds in relational memory test trials, but their item memory performances were comparable (e.g., Lloyd et al., 2009; Sluzenski et al., 2006).

More recent studies have employed episodic memory tasks that tap complex relational representations as opposed to unique pairings of items. For example, Newcombe, Balcom, Ferrara, Hansen, and Koski (2015) examined children's ability to remember two arbitrary associations, each occurring in a specific context. Children were shown two different rooms (e.g., a rainbow and a cloud room) by two different experimenters. Both rooms contained an identical set of four containers but their arrangements differed between the two rooms. Children witnessed a different toy being hidden inside a different container in each room, and were later asked to retrieve a given toy. This task required the use of contextual memory (e.g., the rainbow room) to retrieve the correct object pair association (the bubbles hidden in a basket). Children as young as 15–20 months old performed better than chance, with a significant increase in performance in 21- and 26-month-olds, 34- and 40-month-olds, and 64- and 72-month-olds.

In a similar paradigm, Richmond and Pan (2013) studied 3- to 5-year-old children who learned two series of animal-place pairs (e.g., the duck likes the train station), each presented in a storybook. Every place was associated with two animals, each learned from one of the two storybooks. This relational structure is referred to as ABAC format: A is paired with B in context 1 and paired with C in context 2. At test, children were asked to choose the correct object associated with an animal in a two-alternative force-choice test. Age significantly correlated with performance on this task: older children required fewer blocks to reach 70% accuracy, and were overall more accurate on this task, compared to younger children. Similar results were found by Yim, Dennis, and Sloutsky (2013). In this study, 4-year-olds, 7-year-olds, and young adults first learned two lists of object pairs; each list contained six pairs. Each pair consisted of an overlapping item – an object that occurred in both lists, and one unique item – an object that occurred only in one list (e.g., the bike was paired with a spoon in list 1 and paired with a mug in list 2). Participants first learned all object pairs in the first list until they reached 100% accuracy before learning the object pairs in the second list. Adults outperformed 7-year-old children, who outperformed 4-year-olds (Yim et al., 2013). Interestingly, increasing the saliency of the contextual cue improved 4-year-old children's performance, suggesting that poor attention to the context information at encoding may account for the age-related differences in learning relational information.

Together, these findings suggest important developmental changes in children's abilities to form and retrieve relational representations during early childhood. The marked improvement in relational memory between the ages of 3 and 7 coincides with the age window in which gains in episodic memory are robust (Peterson, Warren, & Short, 2011), consistent with the idea that the developmental changes in relational memory may contribute to the overall maturation of episodic memory in early childhood (reviewed in Olson & Newcombe, 2014). Given these findings, many researchers suggest that the age-related improvement in relational memory may be at the core of the development of episodic memory (Richmond & Pan, 2013; Yim et al., 2013).

1.2 Pattern separation

Relational memory has been used as a proxy for episodic memory across all age groups. While relational memory is crucial in the formation of episodic memory, it is equally important to consider how multiple episodic memories are discerned from one another. Accurate episodic memory also relies on the ability to form distinct memory representations that share overlapping elements. Returning to the restaurant example, the two visits to my favorite restaurant were highly similar: both took place in similar environments with the same company, but with subtle variations between the two episodes (e.g., which dish I ordered and, even more subtly, what type of steak my friend ordered). Memories for common day-to-day events share a considerable amount of feature overlap, which creates memory interference (Gómez & Edgin, 2015). A proposed process by which the overlap among similar memories is reduced to minimize memory interference is pattern separation (Complementary Learning Systems theory: Norman, 2010; Norman & O’Reilly, 2003; O’Reilly & McClelland, 1994). The hippocampus is thought to perform pattern separation, and rapidly assigns distinct representations to specific events by transforming similar memories into highly dissimilar and non-overlapping patterns of activation. Specifically, the dentate gyrus – a hippocampal subfield – is thought to play a crucial role in pattern separation by assigning non-overlapping representations even for highly similar inputs. The ability to remember similar, but not identical, memories as distinct from one another – that is, lure discrimination – is thought to represent the behavioral outcome of pattern separation (reviewed in Yassa & Stark, 2011).

Grounded in the idea that fine discrimination between similar memories depends on pattern separation, previous studies have used the Mnemonic Similarity Task (MST) – a recognition task that uses lure items, which are perceptually similar exemplars of some items in the study list (e.g., Bakker, Kirwan, Miller, & Stark, 2008; Bennett, Huffman, & Stark, 2015; Kirwan & Stark, 2007; Lacy, Yassa, Stark, Muftuler, & Stark, 2011; Stark Yassa, Lacy, & Stark, 2013; Stark Stevenson, Wu, Rutledge, & Stark, 2015; Toner, Pirogovsky, Kirwan, & Gilbert, 2009). In one version of the MST, participants perform an incidental outdoor/indoor task on a series of object images while scanned. Within this series, some objects are identical to previous objects within the series (repeat), some objects are similar exemplars of the previous objects (lure), and some are dissimilar to the rest of the objects within the series (novel). Separation-like activation profiles are only found in the hippocampal subregions CA3/dentate gyrus, but not in other hippocampal subregions (Bakker et al., 2008; Lacy et al., 2011), or other
MTL areas (Kirwan & Stark, 2007). These results are congruent with the idea that the dentate gyrus is responsive to small changes in input and may more readily assign similar inputs using non-overlapping representations despite only subtle differences. Together, there is evidence to suggest that the MST taxes hippocampal pattern separation and that this task can provide a behavioral index of pattern separation.

Importantly, pattern separation has been proposed as one key aspect of episodic memory that is sensitive to typical and atypical aging. In an MST overt recognition task variant, in which participants make either ‘old’, ‘similar’, or ‘new’ judgments to each test item, correctly identifying lures as ‘similar’ items is thought to rely on pattern separation. Lure discrimination is worse in old age (Bennett et al., 2015; Stark et al., 2015; Stark et al., 2013; Toner et al., 2009), correlates with standardized episodic memory performance (Bennett et al., 2015; Stark et al., 2013), and underlies memory dysfunction associated with amnestic mild cognitive impairment (Bakker et al., 2012; Yassa, Stark et al., 2010). Previous studies found that an age-related decrease in lure discrimination is accompanied by higher activation in CA3/dentate gyrus subfields (Yassa, Stark et al., 2010), and a decrease in white matter pathways within hippocampal circuitry (Yassa, Muftuler, & Stark, 2010; Yassa, Mattfeld, Stark, & Stark, 2011). Thus, the Complementary Learning Systems theory introduced a key mechanism by which episodic memory may change with age. Surprisingly, to our knowledge, there have been no assessments of pattern separation abilities in young children.

1.3 Present study

Both relational memory and pattern separation are critical component processes of episodic memory. Whereas relational memory has been positioned front and center within the episodic memory development literature, pattern separation has been largely overlooked. The goal of the current study was to identify developmental changes in both components – relational memory and pattern separation – that underlie the development of episodic memory during early childhood. Given that the ages of 4 and 6 are an age window in which gains in episodic memory are most robust (Peterson et al., 2011), we tested 4- and 6-year-olds, as well as young adults, on two tasks. Our relational memory task was inspired by Newcombe et al.’s (2015) and Yim et al.’s (2013) studies. Engaging, narrated animations were created that contained item–item associations tied to a specific context using an AB-AC paradigm (A is paired with B in one context, but paired with C in another context). Our pattern separation task was a variant of the MST, which requires participants to incidentally encode pictures of objects. The ability to mnemonicly discriminate between two similar items at test was used as a behavioral index of pattern separation. Developing these two tasks allowed us to test whether performances on the relational memory and pattern separation tasks correlate with one another within each age group. Although both relational memory and pattern separation have been ascribed to the hippocampal functions, they have been hypothesized to rely on distinct neural substrates, each of which follows a different maturation profile. Relational memory may rely on CA3 as well as frontal regions (e.g., Norman & O’Reilly, 2003; reviewed in Ghetti & Bunge, 2012), whereas pattern separation may rely on the dentate gyrus (Norman & O’Reilly, 2003). Previous work in non-human primates has shown that the dentate gyrus is a late-developing subfield relative to other hippocampal subfields such as CA1, CA2, and CA3 (Lavenex & Banta Lavenex, 2013; Serres, 2001). Given that the developmental trajectories of these regions are dissociable, it could be the case that relational memory and pattern separation abilities have an uneven development in young children.

To preview, we found a significant improvement in relational memory between the ages of 4 and 6, but 6-year-olds and young adults performed comparably. We also found a significant improvement in pattern separation, indexed by the MST lure discrimination, between the ages of 4 and 6, but a non-significant difference between 6-year-olds and young adults. Performances on the relational memory task and MST did not correlate for any age group.

2 | METHODS

2.1 Participants

A total of 39 4-year-old (23 females; \( M_{\text{month}} = 54.17 \pm 3.87; \text{range} = 48.39–59.47 \)) children and 30 6-year-old (15 females; \( M_{\text{month}} = 75.93 \pm 3.19; \text{range} = 72.11–83.35 \)) children recruited from the Philadelphia suburban areas participated in the study at the Temple Ambler Infant and Child Laboratory. All children were free of neurological damage and had no history of developmental disorders as reported by a parent. Of these children, nine 4-year-olds did not pass the training for the MST thus they were excluded from the MST analyses. All children completed the Relational Memory task. Informed consent was obtained from parents. All children received a small toy for their participation. The adult sample consisted of 52 undergraduate students from Temple University who participated for partial course credit. Two participants did not complete the experiment due to experimenter error. The remaining 50 young adults (25 females; \( M_{\text{age}} = 22.22 \pm 4.79; \text{range} = 18–49 \)) were included in the analyses. All participants gave informed consent and reported having normal or corrected-to-normal vision.

2.2 Relational Memory Task

2.2.1 Materials

We created a novel relational memory task that was engaging while also controlling for important psychologically important variables. Two animation sequences were created using images created in Microsoft PowerPoint or obtained from the internet and manipulated in Adobe Photoshop CS6. Each animation consisted of a tour to two locations (e.g., a red and a blue house), each containing four associations, totaling eight associations per animation (Cronbach’s \( \alpha \) of 16 associations = .82). Every association was made up of one common item (e.g., Pooh bear) – an item that appeared in both locations, and a unique item (e.g., book) – an item that only appears in one location (see...
A total of 16 animations were created to counterbalance the unique items and the order in which the locations were visited. Each animation lasted approximately 2.5 minutes. Example animations can be viewed at http://www.olson-lab.com/memory-test/.

2.2.2 Procedure

Young children

All children watched two videos: a house video and a park video. For each video, they followed an encoding-delay-test procedure. At the beginning of each video, the voice recording stated ‘We’re going on an adventure today. We’re going to see two different houses/parks. We will have to remember the things we see in each place, so let’s watch very carefully, okay?’ The house video toured two different color houses: a red house and a blue house. Four associations were presented in each house. Each association contained one common element and one unique element, that is, ABAC format (see Figure 1A).

For each association, the common element was first introduced: ‘Look, there is Pooh bear. What does Pooh bear like to do in the red house?’ The unique item then appeared on the screen (‘A book. Pooh bear likes to read a book the in red house’). Each association was presented statically for 5 s with 12 transition frames (100 ms/frame) before the next association appeared.

After the encoding phase of each video, there was 5-minute delay, during which the experimenter took the children into another room to play. The test phase consisted of eight, three-alternative forced-choice trials for each animation. In each test trial, children were presented with a static screen shot of the common item in a location (e.g., Pooh bear in the red house), with three options shown below: target, across-context lure, and foil. Targets were the correct unique items seen in the other location (e.g., paint). Foils were novel items that were not seen. Children were asked to choose one item that they saw with a given scene either by pointing or by verbal response (see Figure 1B). The experimenter recorded children’s responses on paper. The order of the test trials was randomized. The locations of the targets, across-context lures, and foils were counterbalanced across test trials. All unique items were counterbalanced such that they were assigned as targets, across-context lures, and foils an equal number of times across participants. The order of the two locations and the order of the animations were counterbalanced across participants.

2.3 Mnemonic Similarity Task (MST)

2.3.1 Materials

A total of 230 digital images of common objects (115 pairs of similar exemplars) were obtained from the Internet, and 46 pairs of object exemplars were sampled from Craig Stark’s laboratory MST stimuli database (http://faculty.sites.ucd.edu/starklab/mnemonic-similarity-task-mst/). We selected objects based on their appeal to children (e.g., toys, animals) and the likelihood that children would have had some familiarity with these objects (e.g., hat, bicycle). Each object pair was matched for size and orientation. The sizes of the objects were comparable across all objects (300 x 300 pixels). An independent sample of 31 young adults rated the levels of similarities for 161 pairs of objects on a 7-point Likert scale, ranging from completely different (1) to identical (7) (see Figure 2A), on Qualtrics (http://www.qualtrics.com/). The rating survey was interspersed with 42 catch trials (21 pairs of identical objects and 21 pairs of completely different objects) in order to ensure that participants did not respond randomly. All participants accurately rated the 21 pairs of identical objects as ‘identical’; therefore we included all participants’ rating results. The similarity ratings for 161 pairs of objects ranged from 4.66 to 6.00. They were binned into three levels of similarities based on the ratings: 20 level-1 items (4.66–4.99), 72 level-2 items (5.00–5.49) and 69 level-3 items (5.50–6.00). Of these, 88 pairs of objects were randomly selected (20 level-1, 34 level-2, and 34 level-3 items) in order to create a study and test set sizes of 66 items (see Figure 2B). The study and test set sizes were determined based on pilot work in order to ensure that 4-year-old children would be able to complete the task. To fully counterbalance all of the items, 88 object pairs were divided into four sets of 22 items, approximately matched on the level of similarities.

![Image](A) A schematic timeline of the associations encountered in one example of the house animation (A). Note that the animations are dynamic; thus Figure 1A illustrates an example of how the common and unique items were paired across two locations (ABAC format), not the actual images from the animations. An example of a test trial is shown in (B)
Each set contained an approximately equal number of stimuli from each semantic (e.g., animals, vegetables/fruit, toys). Eight versions of the study and test lists were created such that each set of objects was assigned as the target, lure, foil, and not-tested items an equal number of times across participants (mean Cronbach’s $\alpha$ of all test versions = .89, range = .84–.94). The experiment was programmed in Eprime 2.0 (Psychology Software Tools, Pittsburgh, PA).

### 2.3.2 Procedure

#### Young children

The first part of the study consisted of an incidental encoding task. Children were told that they would play a game about pictures of objects. The task was to look at pictures of objects presented one at a time on a computer screen and to make an indoor/outdoor judgment for each object (see Figure 3A). Responses were entered by depressing one of two large buttons on a wooden box which when pressed played the recorded word ‘inside’ or ‘outside’ (see Figure 3B). The experimenter instructed ‘I will show you a lot of pictures on this computer screen; whenever you see a picture of an object that usually belong inside, you should press the white button, which says ‘inside’, and whenever you see a picture of an object that usually belongs outside, you should press the yellow button, which says ‘outside’.

Preceding the encoding phase were two self-paced practice trials (a bird and a spoon) to acquaint the children with the general rule of the game. The experimenter informed the children that the pictures would appear and disappear quickly so they would have to press the buttons quickly. Sixty-six objects were presented sequentially in a randomized order for 3 s each followed by a .5 s ISI (see Figure 3A). The incidental encoding task lasted approximately 5 minutes.

The training phase immediately followed the incidental encoding task. The experimenter introduced a new game with a new toy box consisting of three buttons: a red, a green, and a blue button which when pressed played the audio phrases ‘exactly the same’, ‘kind of the same’, and ‘new picture’, respectively. Children were told to push each button by themselves and hear what each one said. The experimenter then explained the new game to the children: ‘Remember when we played the inside/outside game? Now we are going to play a different game with this toy box but with the pictures that we saw before, okay?’ Items assigned as ‘not-tested’ at encoding (e.g., the Lego in Figure 3A) were used for the training session to help children understand the task. During training, children were first shown one object that they had previously seen presented on the left side of the screen, and a triad of objects (one identical, one similar, and one completely different) shown on the right side of the screen (see Figure 4A). The experimenter then explained to the child while pointing that if the object on the right looked exactly the same as the one on the left, they should press the red button. If the object on the right looked kind of the same, but not exactly the same as the one on the left (e.g., they are both Legos, but not exactly the same Lego) they should press the green button. If the object on the right was completely different from the one on the left, they should press the blue button. After the first practice trial, children were encouraged to press the buttons by themselves for the following three trials with immediate feedback from the experimenter (see Figure 4B). After a total of four distinct examples, children then completed two ‘screening’ practice trials in which the object was shown by itself (see Figure 4C). If the child chose the correct button for all three objects for both practice trials (total of six trials), they continued onto the test phase. If the child was not 100% correct, they repeated the training phase with a different set of objects. Only when the child provided 100% correct responses for the two ‘screening’ practice trials could they enter the test phase. If not, the experimenter ended the MST and continued with the Relational Memory task. All lure items included in the ‘screening’ practice trials were level-2 items, ensuring that these lures would not be too easy or too difficult. Among the 69 children, nine 4-year-olds did not pass the training. Of these children, two pressed the same button throughout the training session; two children pressed all three buttons in a systematic order (e.g., red, green, blue), and the remaining five children made errors on a combination of test trials (targets, lures, and foils). The training session took approximately 3–6 minutes.

Immediately after the training session, children were given a self-paced test on 66 items: 22 targets, 22 lures, and 22 foils. Targets were identical items to those studied at encoding. Lures were similar exemplars of the studied items. Foils were novel items that were dissimilar from other objects in the stimuli set. For every trial, the experimenter asked, ‘Is this exactly the same, kind of the same, or completely new
NGO et al. compared with the ones you saw before? Once the children pressed a button on the toy box, the experimenter recorded their responses by pressing keys ‘o’, ‘s’, or ‘n’ for ‘old’, ‘similar’, and ‘new’ on the keyboard. The test image remained on the screen until the experimenter inputted the children's response. The order of the test items was randomized for each participant. The test phase took approximately 5–6 minutes.

FIGURE 3  The encoding phase consisted of 66 object images shown sequentially (A top), in which young children decided whether the object belonged indoors or outdoors by pressing a two-button toy box with audio recordings (B top). At test, children were tested on 66 object images (22 targets, 22 lures, and 22 foils) (A bottom). Children responded by pressing a three-button toy box with audio recordings analogous to ‘old’, ‘similar’, and ‘new’ for each test trial (B bottom).

FIGURE 4  Examples from the training session. (A) First, children were shown an item studied at encoding and a triad of objects, each corresponding to one of the buttons (target: red, lure: green, and foil: blue). Children were instructed when to use each button to respond to different type of items. This was repeated three more times with different items and immediate feedback was provided (B). Children were given two ‘screening’ practice trials in which they were asked to use the buttons to respond by themselves (C).
3 | RESULTS

3.1 | Relational Memory Task

These analyses included all children: 39 4-year-olds, 30 6-year-olds, and 50 young adults. First, we tested whether there was an age effect in overall memory accuracy, defined as the proportion of correct trials. A one-way ANOVA showed a significant age effect, $F(2, 116) = 9.55, p < .001$. Tukey post-hoc tests showed that 4-year-olds ($M = .69, SE = .04$) performed significantly worse than 6-year-olds ($M = .86, SE = .03$), $p = .002$, and young adults ($M = .86, SE = .03$), $p < .001$. There was no difference in accuracy between 6-year-olds and young adults, $p = .99$ (see Figure 5).

Next, we calculated a corrected hit rate for each participant by subtracting the proportion of lure-chosen trials from the proportion of correct trials. The corrected hit rate reflects participants’ accuracy corrected for their lure error rate. There was a significant age effect, $F(2, 116) = 3.39, p < .01$. Four-year-olds ($M = .44, SE = .06$) performed significantly worse than 6-year-olds ($M = .73, SE = .06$), $p = .004$, and young adults ($M = .75, SE = .06$), $p < .001$. There was no difference between 6-year-olds and young adults, $p = .98$.

It is worth noting that overall performance did not differ between the house and the park animations, or between the first and the second animations, all $p > .05$, suggesting that there were no unintended differences in difficulty between the two animations, and that participants did not improve or get worse from fatigue between the first and the second animations.

3.2 | Mnemonic Similarity Task

3.2.1 | Lure discrimination and item memory

The proportions of memory responses (old, similar, and new) for each item type (target, lure, and foil) were calculated for each participant (see Figure 6). Similar to Toner and colleagues (2009), lure discrimination index was calculated for each participant by subtracting the proportion of old responses to lures from the proportion of similar responses to lures. This value ranges from −1 to 1, and indexes the extent to which participants accurately identified lures corrected for the extent to which they falsely endorsed lures as targets. Positive values denote successful discrimination between targets and lures, whereas negative values denotes a higher tendency to over-generalize between two similar items. A lure discrimination index of zero denotes chance-level discrimination. A one-way ANOVA test revealed a significant age effect in the lure discrimination index, $F(2, 109) = 22.82, p < .001$, $MSE = 2.53, \eta_p^2 = .30$. Tukey post-hoc tests showed that 4-year-olds ($M = −.29, SE = .06$) performed significantly worse than 6-year-olds ($M = .07, SE = .06$), $p = .002$, and young adults ($M = .23, SE = .05$), all $p < .001$. There was no significant difference between the 6-year-olds and young adults, $p = .13$. One-sample t-tests (contrasting chance level at 0) revealed that 4-year-olds called lures ‘old’ significantly more often than they would at chance, $t(29) = −5.11, p < .001$, 6-year-olds did not differ from chance, $p = .29$, whereas young adults correctly identified lures above chance, $t(49) = 4.76, p < .001$.  

![Figure 5](https://example.com/figure5.png) Mean proportions of targets, across-context lures, and foils selected in the relational memory task in three age groups.

Young adults

The stimuli were identical to those used with the children. The procedure for young adults was similar to that used with the children. The main difference was that responses were entered by computer keyboard and the instructions were modified to be appropriate for adult listeners. Adult participants completed an incidental indoor/outdoor encoding task, similar to that described earlier, which included 66 items presented serially (2 each with the 0.5 s ISI). After the encoding phase, participants were given a self-paced memory test on 66 items (22 targets, 22 lures, and 22 foils). The test instruction was as follows: ‘if the objects were identical to the ones they saw, press key ‘o’ for old; if the objects were similar, but not identical to the ones they saw, press key ‘s’ for similar; if the objects were completely different from the ones they saw, press key ‘n’ for new.’

Overall procedure

All participants performed two tasks: the MST and the relational memory task. Both tasks were presented on a 29.5-inch Dell desktop. For 4- and 6-year-olds, the MST was always administered first, followed by the relational memory task. Based on our pilot work, the MST appeared to be more taxing than the relational memory task and thus was always administered first for young children to avoid low performance due to fatigue. Adults were first administered the relational memory task encoding phase, followed by the MST, followed by the relational memory task’s test phase. Our pilot work also showed that adults reached ceiling performance on the relational memory task if administered in the encoding-delay-test procedure for each video; thus the MST was flanked by the relational memory task encoding and the test phases to serve as a delay. All parts of the experiment took place in the same room. Children were only taken to the playroom during the delay period of the Relational Memory Task.
Age was the only factor that was significant, $F(1, 106) = 22.89, p < .001$, whereas item memory was not significant, $F(1, 106) = 2.90, p = .09$ (see Figure 7). These results show a significant improvement in lure discrimination between the ages of 4 and 6, but no increase was found between 6-year-olds and young adults.

### 3.2.2 Levels of similarity

As the exemplar pairs were more similar to each other, we expected lure discrimination to decrease. We tested whether lure discrimination differed across three levels of similarity as a function of age. We conducted a 3 (levels of similarity: 1, 2, 3) x 3 (age: 4, 6, adults) mixed factorial analysis. We found a main effect of levels of similarity, $F(2, 106) = 106.00, p < .001$, a main effect of age, $F(2, 107) = 25.16, p < .001$, and a significant interaction, $F(4, 105) = 5.53, p < .001$. To unpack the significant interaction, pairwise comparisons for the levels of similarities were conducted for each age group separately. In 4-year-olds, lure discrimination did not significantly differ between level-1 ($M = .17, SE = .08$) and level-2 ($M = .27, SE = .07$), $p = .21$, or between level-2 and level-3 ($M = .36, SE = .06$), $p = .16$, but the difference between level-1 and level-3 was significant, $p = .05$. In 6-year-olds, lure discrimination was significantly higher for level-1 items ($M = .51, SE = .07$) compared to level-2 ($M = .06, SE = .06$) and level-3 items ($M = .09, SE = .10$), all $p < .001$. The difference between level-2 and level-3 was not significant, $p = .06$. In young adults, lure discrimination was significantly higher for level-1 items ($M = .61, SE = .06$), followed by level-2 items ($M = .21, SE = .06$), followed by level-3 items ($M = .03, SE = .06$), all $p < .001$ (see Figure 8). Overall, these findings are consistent with the similarity ratings results such that the more similar the exemplars were to each other, the more likely the participants were to misremember them as the same.

Across all three levels of similarities, 4-year-olds’ lure discrimination indices were significantly lower than 0, all $p < .05$, suggesting that they were more likely to falsely endorse lures as targets than to correctly identify lures. Six-year-olds’ lure discrimination was only significantly above chance for level-1 items, whereas young adults’ lure discrimination was significantly above chance for level-1 and level-2 items, all $p < .001$. Together, these findings show that unlike 6-year-olds and young adults, 4-year-olds have a tendency to over-generalize, even for exemplar pairs in the lowest level of similarity. However, the fact that they were more likely to call a lure ‘old’ as the items were more similar is likely the participants were to misremember them as the same.

### 3.3 Correlation between pattern separation and relational memory

To examine whether relational memory is related to pattern separation, we correlated the corrected hit rate in the relational memory task with lure discrimination in the MST for each age group. Our normality tests showed that neither the MST lure discrimination nor the corrected hit (target – lure) of the relational memory task was normally distributed (Shapiro-Wilk < .001), thus violating one of the assumptions of the Pearson correlation. Therefore, we conducted
the Spearman rank correlation. There was no significant correlation between MST lure discrimination and the relational memory corrected hit rate in either the 4-year-olds, $r(28) = -.04, p = .83$, the 6-year-olds, $r(28) = .12, p = .52$, or adults, $r(48) = .10, p = .49$. Across all participants, MST lure discrimination and Relational Memory corrected hit rate significantly correlates, $r(108) = .23, p = .02$. However, when age is entered as a covariate, the partial correlation is non-significant, $r(107) = .15, p = .13$.

4 | DISCUSSION

Episodic memory is a complex and multifaceted construct. Thus, characterizing the development of episodic memory development requires an examination of its component processes. In a previous study, Cheke and Clayton (2015) aimed to delineate a single common underlying construct of episodic memory by administering various ‘episodic memory’ tasks to the same children including free and cued recall, what-where-when, and source memory tasks; an assessment of pattern separation was not included. Our goal is to fill this gap in the memory development literature. The current study examined the development of two crucial aspects of episodic memory: relational memory and pattern separation in early childhood. In our relational memory task, 4-year-olds performed significantly worse than 6-year-olds and young adults, whereas 6-year-olds and young adults performed comparably. In addition to choosing a lower proportion of targets, 4-year-olds showed higher false alarm rates for across-context lures – that is, the correct paired objects presented in the other context. Our results echo the findings of previous studies examining relational memory using different paradigms and stimuli (e.g., Lloyd et al., 2009; Sluzenski et al., 2006) in suggesting significant improvement in children’s ability to remember the relations among multiple items between the ages of 4 and 6. It is important to note that although 6-year-olds and young adults performed very similarly in our task, it is likely that relational memory continues to develop well into adolescence, although in a less dramatic fashion (Ghetti & Lee, 2011; reviewed in Ghetti & Bunge, 2012).

Compared to 4-year-olds’ performance in a previous study using a face-name relational memory task (Koski, Newcombe, & Olson, 2013), in which they chose targets 44% of the time in a three-alternative forced-choice test, 4-year-old children performed well in our relational memory task (69%). The use of engaging and dynamic presentation of the associations via animations in the current study may be more appropriate for children in this age range. In contrast to Yim and colleagues (2013) who found that children, but not adults, showed proactive interference such that learned associations from List 1 interfered with learning the associations from List 2, we did not find any difference in memory performance for the first location versus the second location for any age group. An important methodological difference between our relational memory task and those used Yim and colleagues (2013) and Richmond and Pan (2013) should be noted. In those studies, participants learned the associations in the first list until they reached a learning criterion before learning the associations on the second list. In our task, participants were only shown each association once, because episodic memory is often a single-trial learning phenomenon.

To assess pattern separation, we tested 4-year-olds, 6-year-olds, and young adults using the MST, a lure discrimination task, in which similar developmental effects were observed. We found a significant improvement in lure discrimination between 4- and 6-year-olds, and a non-significant difference between 6-year-olds and young adults. Unlike 6-year-olds and young adults, 4-year-olds frequently misidentified lures as ‘old’ more often than chance – an indicator of pattern separation failure. These results suggest that the ages of 4 to 6 may be an important age window in which pattern separation increases significantly. Although there was no significant difference in lure discrimination between 6-year-olds and young adults, 6-year-olds did not correctly identify lures above chance, whereas adults did. These results suggest that pattern separation likely undergoes improvement well beyond the age of 6. The age effect in lure discrimination held when controlling for item memory performance, suggesting that 4-year-olds’ poor performance on lure discrimination cannot be accounted for by lower general item memory alone. Four-year-olds did not discriminate between targets and lures for any levels of the similarity, even for lures that were least similar to targets (level-1 items). Six-year-olds, on the other hand, were successful at discriminating between targets and lures for level-1, but showed no discrimination when the items were more similar (level-2 and level-3 items), Young adults were successful at lure discrimination except for when the lures were very similar to targets (level-3 items). Importantly, in all age groups, the tendency to misidentify lures as ‘old’ items systematically increased as the similarity of the lure items increased, suggesting that young children’s poor performance is unlikely due to task procedural difficulty.

Although we did not find any differences between 6-year-olds and young adults in either the MST or the relational memory task, there are methodological differences between the tasks administered to children and adults that may account for the lack of findings. With regard to the relational memory task, children were shown the first animation, followed by a 5-minute delay, then tested. This procedure
was repeated for the second animation. In contrast, adults were shown both animations sequentially and then were tested on both. With regard to the MST, the encoding time and the delay differed for children versus adults. Children saw each item for 3s while adults saw each item for 2s. After encoding, children were given training, which lasts on average 3–4 minutes, whereas adults performed the relational memory task in between MST encoding and test phases, which lasts approximately 10 minutes. These methodological differences were intended to level the playing field for children and young adults, because these tasks were developed specifically for children. Thus it is likely that there are developmental changes that occur in both relational memory and pattern separation after the age 6 and well into adulthood, which were not detected in our paradigms.

In our study, although 6-year-olds performed better than 4-year-olds in lure discrimination, they did not correctly identify lures at above chance levels. On the other hand, 6-year-olds outperformed 4-year-olds at relational memory, and both age groups performed significantly above chance (target selected = .33). These findings suggest that pattern separation and relational memory may have dissociable developmental trajectories. Relational memory and pattern separation have been hypothesized to rely on distinct neural substrates, each of which follows a different maturation profile. Relational memory may rely on CA3 (Norman & O’Reilly, 2003) as well as portions of the frontal lobe (DeMaster & Ghetti, 2013; reviewed in Ghetti & Bunger, 2012), whereas pattern separation may rely critically on the dentate gyrus (Norman & O’Reilly, 2003). Findings in non-human primates show that the dentate gyrus is a late developing subfield relative to other hippocampal subfields such as CA1, CA2, and CA3 (Lavenex & Banta Lavenex, 2013; Serres, 2001). Consistent with this, we found that pattern separation and relational memory performance did not correlate with each other for any age group. One possible explanation for our findings is that relational memory and pattern separation tap different aspects of episodic memory. Relational memory may be important for forming an episodic memory (Cohen & Eichenbaum, 1993), whereas pattern separation allows for similar memories to be represented separately with reduced overlap in order to minimize interference at retrieval (Norman & O’Reilly, 2003).

It is important to note that the MST demands mnemonic distinction between similar items with similarity confined along the dimension of conceptual (object identity/category) and perceptual properties (e.g., color, shape). However, this is not the only way in which two items can share similarities. Perhaps what would be more critical for episodic memory are ways in which objects are similar on the ‘episodic’ dimension. Returning to our restaurant example, one defining difference between the two events is that I ordered salmon in one episode and steak in the other. In this example, salmon and steak, which possess numerous perceptual differences, are similar due to the overlap in the contexts in which they were encountered. It remains unclear whether similar items and similar episodes are handled differently by hippocampal pattern separation. This question has important theoretical implications and merits empirical investigation. A pivotal future direction for this line of research is to examine the development of pattern separation for contexts as it pertains to the development of episodic memory.

Another point worth noting is that our relational memory task, as well as others used in previous work, may require some degree of pattern separation. In tasks that employ the AB-AC paradigm, both associations can be said to be similar such that they share an overlapping component. In addition, these associations may also share other overlapping episodic features depending on the similarities between the contexts in which they were learned. In our task, although the contexts had distinct background colors (e.g., red versus blue house), as well as unique decorative details in each location, the structures of the house/park tours were made similar (e.g., the living room was first visited, followed by the kitchen in both houses). Similarly, tasks by Richmond and Pan (2013), Yim and colleagues (2013), and Newcombe and colleagues (2015) also employed variants of AB-AC relational structure, in which the to-be-remembered associations shared overlapping elements. This type of relational memory is not process-pure. That is, in addition to relational memory, they may place some demand on pattern separation. This can be contrasted to relational memory tasks that have an AB-CD relational structure, in which the to-be-remembered associations are always made up of unique components, and thus have fewer overlaps. However, given that the similarity of the associations and their encoding contexts was not manipulated in these studies, the development of pattern separation for associations or contextual memory in early childhood remains unknown.

Relational memory has been used as a proxy for episodic memory across all age groups. While relational memory is crucial in the formation of episodic memory, it is equally important to consider that new memories are formed against a backdrop of existing memories, and in many cases, these memories share overlapping elements with one another. However, the development of pattern separation has been overlooked in the memory development literature. The current study, to our knowledge, was the first to investigate the development of pattern separation, indexed by MST lure discrimination, in young children. We found that 4-year-olds exhibit a tendency to over-generalize, whereas 6-year-olds do not. Newcombe, Lloyd, and Ratliff (2007) previously argued that it may be more important to build semantic knowledge than to remember specific episodes in the early years of life. Aligned with this argument, generalizing across similar memories to learn about the regularities of the environment may be prioritized over fine-grained discrimination among them. That is, younger children may need to learn that objects such as chairs, regardless of subtle variations in colors and shapes, serve the same function. In parallel, we examined the development of relational memory – another key characteristic of episodic memory – in the same children. The current study provides important implications on the mechanisms of memory development. Our findings suggest that pattern separation undergoes a significant improvement in early childhood, and that it is crucial to consider the developmental changes of both relational memory and pattern separation to account for episodic memory development.

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