Volumetric development of hippocampal subfields and hippocampal white matter connectivity: Relationship with episodic memory

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Funding information
Eunice Kennedy Shriver National Institute of Child Health and Human Development, Grant/Award Number: R01HD098152

Abstract
The hippocampus is a complex structure composed of distinct subfields. It has been central to understanding neural foundations of episodic memory. In the current cross-sectional study, using a large sample of 830, 3- to 21-year-olds from a unique, publicly available dataset we examined the following questions: (1) Is there elevated grey matter volume of the hippocampus and subfields in late compared to early development? (2) How does hippocampal volume compare with the rest of the cerebral cortex at different developmental stages? and (3) What is the relation between hippocampal volume and connectivity with episodic memory performance? We found hippocampal subfield volumes exhibited a nonlinear relation with age and showed a lag in volumetric change with age when compared to adjacent cortical regions (e.g., entorhinal cortex). We also observed a significant reduction in cortical volume across older cohorts, while hippocampal volume showed the opposite pattern. In addition to age-related differences in gray matter volume, dentate gyrus/cornu ammonis 3 volume was significantly related to episodic memory. We did not, however, find any associations with episodic memory performance and connectivity through the uncinate fasciculus, fornix, or cingulum. The results are discussed in the context of current research and theories of hippocampal development and its relation to episodic memory.

KEYWORDS
hippocampus development, episodic memory, fornix, uncinate fasciculus, cingulum, medial temporal lobe

1 INTRODUCTION

The hippocampal formation and adjacent cortices that make up the medial temporal lobe (MTL) have long been known to play an important role in learning and memory (Scoville & Milner, 1957). Development of the MTL system, its maturation with respect to overall cerebral cortical development, and related functional consequences are central motivating questions. Here, we attempt to (1) establish whether MTL development, with a focus on the hippocampus and respective subfields—both structurally and through white matter connectivity—continues beyond early childhood; (2) evaluate the relation between age-related MTL volumetric differences with respect to the rest of the cerebral cortex; and (3) lastly assess how these age-related differences in volume and white matter connectivity relate to performance on a standardized memory test that assesses mechanisms important for episodic memory.

The MTL is a heterogeneous set of structures with unique functional attributes related to learning and memory composed of the...
hippocampal formation and adjacent entorhinal, perirhinal, and parahippocampal cortices (Lech & Suchan, 2013; Squire, Stark, & Clark, 2004). The hippocampal formation can be further subdivided into distinct subfields defined by cytoarchitecture: cornu ammonis (CA) fields 1–3, dentate gyrus (DG), and subiculum (Winterburn et al., 2013). The unique functional contributions of the adjacent MTL cortices arise as a result of segregation of information, with spatial information predominately represented by the parahippocampal and medial entorhinal cortices and object information largely represented by the perirhinal and lateral entorhinal cortices (Lee, Hunsaker, & Kesner, 2005; Ranganath, 2010). The two streams of information subsequently converge on neurons in the hippocampal formation.

Distinct cytoarchitecture of the hippocampal subfields further confers functional attributes to the MTL system (Marr, 1971). The DG is composed of a high volume of largely quiescent cells. The large number of neurons, combined with a low firing rate, provides the ideal substrate for performing a computation known as “pattern separation,” which is proposed to orthogonalize or make overlapping representations distinct (Rolls, 2013; Rolls, 1996). Neurophysiological evidence of pattern separation has been identified in both rodent and human studies (Bakker, Kirwan, Miller, & Stark, 2008; Leutgeb, Leutgeb, Moser, & Moser, 2007). Neurogenesis, which occurs in only a handful of places in the adult brain, including the DG, is also thought to play an important role in pattern separation (Pereira et al., 2007). In contrast, the most notable cytoarchitectural feature of the CA3 subfield is its recurrent collaterals (Bains, Longacher, & Staley, 1999). The structure of the CA3 makes this subfield adept at a computation called “pattern completion” or the reinstatement of a prior representation from partial input (Rolls, 1996). Lastly, the CA1 subfield receives notable input from both the CA3 and entorhinal cortex (Brun et al., 2008), distinguishing this subfield as a comparator of current information (incoming from the entorhinal cortex) with previously encoded information (retrieved through the CA3 process of pattern completion; Lisman & Grace, 2005). Together these regions and their unique functional attributes support the ability to discriminate overlapping memories, generalize across similar episodes, and lay down new memories.

1.1 Differential development of the hippocampus, its subfields, and its connectivity

In addition to investigating the functional contributions to subdivisions of the hippocampus, histological studies in nonhuman primates (Lavenex & Lavenex, 2013; Lavenex, Lavenex, & Amaral, 2007) combined with cross-sectional and longitudinal human MRI studies (see Jones & McHugh, 2011 for a review) have observed distinct developmental trajectories of the cortex and hippocampal formation as well as the different hippocampal subfields. Several important questions have been raised in the literature, and some have been answered to varying degrees. The present study focuses on a number of questions that remain unanswered, mainly due to the presence of low statistical power in the extant literature, or to the investigation of restricted age ranges (Joshua K Lee, Johnson, & Ghetti, 2017; Uematsu et al., 2012). These open questions are as follows: (1) does the structure of the hippocampus continue to develop substantially beyond early childhood? (Joshua K Lee, Ekstrom, & Ghetti, 2014); (2) how does the structure of the hippocampus develop in relation to the rest of the cerebral cortex; (3) does the white matter connectivity of the hippocampus develop substantially beyond early childhood; and (4) how do these factors relate to the development of episodic memory at the behavioral level?

1.2 Hippocampal structural development

There are a number of studies that have examined the hippocampus as a whole structure, yet these studies have failed to come to a consensus on whether the hippocampus continues to show age-related differences throughout development (Goddings et al., 2014; Lee et al., 2015; Østby et al., 2009; Riggins et al., 2018; Tannes et al., 2013; Uematsu et al., 2012; Yurgelun-Todd, Killgore, & Cintron, 2003). Some studies have found that development of the hippocampus peaks earlier in childhood or have otherwise failed to find age-related differences, even when substantial age ranges are investigated (Barnea-Goraly et al., 2014; Giedd et al., 1996; Gogtay et al., 2006; Joshua K Lee et al., 2015; Riggins, Blankenship, Mulligan, Rice, & Redcay, 2015; Uematsu et al., 2012). For example, in a longitudinal study of 31 children with repeated scans, Gogtay and colleagues (2006) reported that the total volume of the hippocampus remained unchanged bilaterally from age 4 to 25 years. However, other studies investigating similarly broad age ranges have found substantial age-related differences (e.g., Østby et al., 2009). For example, in a large cross-sectional study of 171 participants aged 8–30 years, Østby and colleagues (2009) reported nonlinear decreases in gray matter in cerebral cortex, with concomitant increases in overall volume in amygdala and hippocampus, peaking during adolescence. This nonlinear developmental trajectory was later replicated in a large cross-sectional morphometric MRI study of the amygdala and hippocampus from 1 month to 25 years of age (Uematsu et al., 2012). Thus, there remain considerable discrepancies in the literature regarding how the structure of the hippocampus changes with development.

More recently, a growing body of literature has investigated structural development of the hippocampus with increased attention to the possibility of differential development of hippocampal subfields. Riggins and colleagues (2018) examined younger children between 4 and 9 years, and found age effects only in CA1. Lee and colleagues (2014) demonstrated age-related increases, in a sample of 39, 8– to 14-year-olds, in CA3 and DG between middle childhood and early adolescence, consistent with previous histologic studies (c.f. Insauti, Cebada-Sanchez, & Marcos, 2010). Similarly, Krogsrud and colleagues (2014) demonstrated in 244 healthy participants, aged between 4 and 22 years, that there is a rapid development of most of the hippocampal subfields until around 13–15 years. In a recent update to their earlier work, Canada and colleagues (Canada, Botdorf, & Riggins, 2020; Canada, Hancock, & Riggins, 2021) examined the development...
METHOD AND DESIGN

2.1 Participants

The current cross-sectional study was conducted using MRI data from the publicly available database, Pediatric Imaging, Neurocognition, and Genetics (PING) (http://pingstudy.ucsd.edu). Data are thus available to the scientific community through PING. Data were provided on an external hard drive that included a virtual machine and Linux mountable partition. Data from the Linux partition were uploaded to our local high-performance computing cluster. The PING database consists of 1493 typically developing participants between the ages of 3.0 and 21.0 years (713 female). Participants were scheduled for two separate visits. The first included neuropsychological assessments and the second included MRI data acquisition. Of these 1493 participants, our analysis was conducted on a total of 830 participants (380 female), with an age range of 3.0–21.0 years (M = 12.6; SD = 4.9). Age was recorded at the time of the MRI scan. The discrepancy between the number of participants available and the number of participants in our analysis is due to the fact that 830 participants had an available MRI data that had been processed through FreeSurfer so that we could conduct the hippocampal segmentation (see below).

Recruitment of participants is described in detail in Jennigan et al. (2016). Briefly, participants were recruited from multiple cities in the United States (Baltimore, Boston, Honolulu, Los Angeles, New Haven, New York, Sacramento, and San Diego). Exclusion criteria included (a) neurological disorders; (b) history of head trauma; (c) preterm birth (less than 36 weeks); (d) diagnosis of an autism spectrum disorder, bipolar disorder, schizophrenia, or mental retardation; (e) pregnancy; (f) daily illicit drug use by the mother for more than one trimester; and (f) contraindication to MRI. There were 380 females and 450 males in our subsample. The mean age was 12.56 years (SD = 4.91 years; range = 3–21 years). Highest education was reported and distributed across seven categories: 1 (Less than 7 years of school) = 0.7%; 2 (7–9 years of school) = 0.4%; 3 (10–11 years of school) = 1.9%; 4 (high
school graduate) = 9.6%; 5 (1-3 years of college) = 21.4%; 6 (4-year college graduate; BA, BS, BM) = 28.3%; 7 and (Professional; MA, MS, ME, MD, PhD, LL.D) = 35.5%. Household income was measured on a 12-point categorical scale (from <$5000 to >$300,000): 1-2 = 2.6%; 3-4 = 6.1%; 5-6 = 6.9%; 7-8 = 24.8%; 9-10 = 6.6%; and 11-12 = 2.1% (see Khundrakpam et al., 2020). Genetic ancestry factors were derived for five categories, and participants were probabilistically assigned to each category based on genetic markers (Jernigan et al., 2016).

The percentage of participants showing a nonzero probability is shown in parentheses: African (11.9%), Native American (11.7%), East Asian (16.9%), Oceania (4.5%), Central Asian (11.2%), and European (69.6%).

As part of the PING study, participants were administered multiple neuropsychological tests, including social and emotional assessments from the Phoenix toolkit (https://www.phenxtoolkit.org) and cognitive assessments from the NIH Toolbox (http://www.nihtoolbox.org). In this study, we focused on the NIH Toolbox Picture Sequence Memory Task (PSMT), which is based on the Imitation-Based Assessment of Learning (IBAL), a measure of episodic memory.

2.2 Image acquisition and data analysis

Image acquisition and image data postprocessing were conducted by the PING team as described in Jernigan et al. (2016). We provide only cursory details here. Across 10 sites and 12 scanners, a standardized, multiple modality high-resolution MRI protocol was implemented using the following scanner manufacturers: GE Discovery MR 750, GE Signa HDX, Siemens Trio TIM, and Phillips Medical Systems Achieva. T1-weighted structural and diffusion-weighted imaging (DWI) data were used for the present study. The scanning protocols are summarized as follows: Siemens (T1-weighted: TR = 2170 ms, TE = 4.33 ms, flip angle = 7°, matrix size = 256 × 256 mm, voxel size: 1 × 1 × 1.2, acquisition time = 8:06 min; Diffusion-weighted: Isotropic [2.5 × 2.5 × 2.5 mm], single-shot echo-planar sequence protocol: slices = 68, slice thickness = 2.5 mm, FOV = 240 × 240 × 170 mm, TR = 9000 ms, TE = 91 ms, directions = 30, b = 1000 s/mm², acquisition time = 10:00 min, acquisition matrix = 240 × 240 × 170); Phillips (T1-weighted: TR = 6.8 ms, TE = 3.1 ms, flip angle = 8°, matrix size = 256 × 240 mm, voxel size: 1 × 1 × 1.2, acquisition time: 9:19 min; Diffusion-weighted: Isotropic [2.5 × 2.5 × 2.5 mm], single-shot echo-planar sequence protocol: slices = 60, slice thickness = 2.5 mm, FOV = 240 × 240 × 150 mm, TR = 9000 ms, TE = 91 ms, directions = 30, b = 1000 s/mm², acquisition time = 5:15 min, acquisition matrix = 96 × 95); and GE (T1-weighted: TR = 8.1 ms, TE = 3.5 ms, flip angle = 8°, matrix size = 256 × 192 mm, voxel size: 1 × 1 × 1.2, acquisition time: 8:05 min; Diffusion-weighted: Isotropic [2.5 × 2.5 × 2.5 mm], single-shot echo-planar sequence protocol: slices = 60, slice thickness = 2.5 mm, FOV = 240 × 240 × 150 mm, TR = 9000 ms, TE = 91 ms, directions = 30, b = 1000 s/mm², acquisition time: 5:15 min, acquisition matrix = 96 × 95).

2.3 Postprocessing and quality control of MRI data

All postprocessing of the volumetric and diffusion data was conducted by the PING study team and is described in Jernigan et al. (2016) with the exception that the hippocampal segmentation was conducted by us (described below). A standard PING scan session included (1) a three-dimensional T1-weighted inversion-prepared RF-spoiled gradient echo scan using prospective motion correction (PROMO; White et al., 2010) for cortical and subcortical segmentation; (2) a three-dimensional T2-weighted variable flip angle fast spin echo scan, which also used PROMO for detection and quantification of white matter lesions and segmentation of CSF; and (3) a 30-direction DWI scan, with integrated B₀ distortion correction (DISCO), for segmentation of white matter tracts and the measurement of diffusion parameters.

Raw image quality control was conducted for each scan session of the PING project. Images were automatically checked for completeness and protocol compliance, and were reviewed for image quality by technicians. All images were screened for motion artifacts, excessive distortion operator error, or scanner malfunction. Images were rated with either, good, average, and bad. T1-weighted images were examined slice by slice for excessive motion. Each volume was rated as either as acceptable or recommended for rescans.

Because we wanted to take advantage of this extensive postprocessing pipeline, we began our investigation with T1-weighted scans that had been processed through FreeSurfer 5.3 and which were part of the public release. FreeSurfer (a) segments the white and gray matter of anatomical volumes; (b) inflates the cortical surfaces separately for each hemisphere (Dale, Fischl, & Sereno, 1999; Fischl, Sereno, Tootell, & Dale, 1999); and (c) segments the subcortical structures, including the hippocampus. We further applied the hippocampal segmentation in FreeSurfer v6.0, as described below.

2.4 Segmentation of the hippocampus

For the structural analysis of hippocampal volume, we re-ran the reconstruction of the v5.3 public release data using FreeSurfer v6.0, including the hippocampal segmentation step to segment the hippocampus into subfields. This step follows the routine outlined in Iglesias et al. (2015) and was invoked by applying the -hippocampal-subfields-T1 flag to the standard FreeSurfer reconstruction. This reconstruction was applied to all 830 participants who had available FreeSurfer v5.3 reconstructions as part of the public release. The entire recon-all procedure was applied in v6.0, beginning with the T1.mgz that was provided with the release (thus, it was already subjected to the quality control steps through PING). The hippocampal subfield segmentation in v6.0 employs a number of substantial improvements over v5.3, which was the first version of FreeSurfer to include a hippocampal subfield segmentation option. These improvements, which specify the training atlas used for automated segmentation, include use of
ultrahigh-resolution (0.1 mm isotropic) ex vivo MRI data collected at 7T in 15 brains (employing 0.13 mm isotropic resolution, allowing delineation of a molecular layer), delineation of a large number of structures (i.e., 15), and modeling of surrounding structures to constrain labeling within the hippocampus. An unbiased segmentation of each brain in our sample was computed based on this training atlas using a Bayesian inference algorithm (Iglesias et al., 2015). The procedure was validated on three publicly available datasets and has been found to have good agreement to manual labeling of ex vivo scans (Iglesias et al., 2015). Using this automated tool, we segmented the hippocampal formation into the following subdivisions, with DG and CA3 combined: CA1, DG/CA3, subiculum, and entorhinal cortex (not part of hippocampus proper) in both hemispheres.

2.5 DWI data postprocessing

Diffusion data were postprocessed by the PING team following the steps elaborated upon in Hagler et al. (2009). The procedure included removal of head motion by rigid body registration between $b = 0$ images of each scan, removal of within scan motion, eddy current correction with a nonlinear estimation procedure, and removal of spatial and intensity distortions by application of nonlinear registration to a reverse phase-encoded scan (Holland, Kuperman, & Dale, 2010). See Jernigan et al. (2016) for additional details.

2.6 Diffusion tensor imaging analysis

All of the fiber pathways in the current study were traced by the PING team. To trace these pathways, the PING team employed the automated labeling process developed by Hagler et al. (2009). This method uses a probabilistic atlas containing 23 fiber tracts that were constructed by manually identifying fiber tracts in 21 healthy controls and 21 patients with temporal lobe epilepsy (for additional details please, see Hagler et al., 2009). The tracking methods include the principal diffusion orientation, FA, mean, longitudinal, and transverse diffusivity (MD, LD, and TD). To label long-range white matter tracts, AtlasTrack was used to automatically label fiber pathways based on a probabilistic atlas of fiber tract locations and orientations (Hagler et al., 2009). The atlas probability was applied after resampling, rotating, and warping from the atlas to the single subject space.

2.7 Behavioral methods

To examine the relation between volumetric and diffusion-weighted images and episodic memory performance, we used the NIH Toolbox PSMT, conducted as part of the NIH Toolbox Cognition battery (http://nihtoolbox.com), and gathered as part of the PING data collection efforts. In this task, images of pictured objects and activities are presented on an iPad in an arbitrary order. The participant’s task is to view the presentation of the images, and then to reproduce the ordering of the images after they leave the screen. The sequence length is varied to manipulate task difficulty, from six to 15 pictures (Bauer et al., 2013). The participant’s score is derived from the cumulative number of adjacent pictures remembered correctly over three learning trials. The task has good test–retest reliability (intraclass correlation coefficient $= .76$) and convergent validity with other measures of episodic memory (Bauer et al., 2013).

2.8 Statistical analysis

Three separate analyses were performed. The first set of analyses examined age-related differences in the volume of the whole hippocampus and entorhinal cortex, and the hippocampus segmented into three subregions (CA1, DG/CA3, and subiculum). The second set of analyses examined age-related differences in FA and MD of the three fiber pathways of interest: the fornix, the uncinate fasciculus, and the hippocampal cingulum. The third set of analyses examined the relation of these morphologic and diffusion metrics to performance on the NIH Toolbox PSMT.

2.9 Combat correction for cross-site differences

Before conducting the analysis, we also corrected the data (both morphologic and diffusion) for cross-site differences in scanner type and acquisition protocols. To make this correction, we employed the ComBat method developed by Johnson and colleagues (2007) and applied to imaging data by Fortin and colleagues (2017). This method uses an empirical Bayes framework to calculate a location and scale adjustment to the data. It estimates an empirical statistical distribution of the location and scale parameters under the assumption that all voxels share a common distribution. Within a regression framework, an adjusted value is computed for each data point. The ComBat-adjusted data were used for all analyses going forward.

2.10 Analysis 1: Age-related differences in volume of hippocampus and entorhinal cortex

To analyze age-related differences in hippocampal, entorhinal cortex, and hippocampal subregion volume (for both hemispheres), we followed the procedure from Østby and colleagues (2009) and corrected for residual intracranial volume (ICV). This is done because brain structures tend to scale with head size, and can “obscure the unique effects of age” (Østby et al., 2009, p. 11774). We are interested in size differences of the structures of interest that are not confounded with head-size differences. The estimate of ICV was taken from the FreeSurfer output, and was calculated by using atlas normalization as a proxy for ICV (see Buckner et al., 2004).

We used the residual method to correct for differences in head size, as is recommended by empirical investigation (Sanfilipo, Benedict, Zivadinov, & Bakshi, 2004). Each volume of interest (dependent vari-
able) was subjected to a regression, with ICV predicting the dependent variable. The residuals of this regression were standardized, and passed to the next analysis (i.e., the new dependent variable is a scaled ICV-corrected volume estimate for each structure of interest). A generalized additive model (GAM; R 3.4.3; package gam; R Core Team, 2017) was fit to each variable of interest, with age at exam as the predictor. Integrated smoothness estimation was applied (of the form \( \text{gam}(dv \sim predictor) \)). We computed Akaike information criterion (AIC) values for each model. In addition, we computed the first derivative of the data to identify the “peak” of the curve, in cases where the function was nonlinear.

We also conducted an analysis of the hippocampus proper in relation to development of the cortex. To do so, we computed the ratio of hippocampal volume to volume of the cortex (hippocampal volume/total cortical volume). Cortical volume was derived from FreeSurfer by taking the volume of tissue inside the pial brain surface minus the volume of tissue inside the white matter surface, and minus the volume of noncortical gray matter (e.g., the hippocampus). This variable was scaled and subjected to the same GAM analysis as above. Each variable (region) was entered into its own regression. We applied a false discovery rate (FDR; Benjamini & Hochberg, 1995) correction across regions.

### 2.11 Analysis 2: Age-related differences in FA or MD of the fornix, uncinate fasciculus, and hippocampal cingulum

We used a similar approach as we described above, except that whole brain FA or MD was residualized in place of ICV. Thus, in each case, for each hemisphere, we specified age as the predictor in the model, and residualized scaled FA or MD of each pathway as the outcome variable. The first derivative was similarly computed for each of the pathways, for each hemisphere. Each variable (tract) was entered into its own regression. We applied an FDR correction across tracts.

### 2.12 Analysis 3: Relation of morphologic and diffusion metrics to performance on the NIH Toolbox PSMT

To explore the relation between morphologic and diffusion metrics and episodic memory, we conducted linear mixed effects models (R package lme) using each hippocampal volume measure, FA and MD in each pathway of interest, and entorhinal cortex volume, for each hemisphere. Age, gender, ethnicity, parent highest level of education, household income, and device serial number (to control for scanner type) were entered as covariates. In addition, ICV (for the morphologic variables) and whole brain FA or MD (for the diffusion variables, which was the average of all streamlines in the atlas) were entered as covariates. Finally, subject ID was entered as a random effect. We also ran these analyses using hippocampal–cortical ratio as a predictor. Each variable (tract or region) was entered into its own regression. We applied an FDR correction across regressions.

### 3 RESULTS

#### 3.1 Results of analysis 1: Age-related differences in volume of hippocampus and entorhinal cortex

The results of Analysis 1 are presented in Figure 1 and Table 1. These results reflect the volume of the regions of interest after controlling for ICV. In initial models, we also examined the data for effects of sex, but finding none, we removed sex from the models. The analysis showed that for the whole hippocampus, in both hemispheres, there is a similar nonlinear relation such that the volume of the hippocampus is elevated with increasing age up until 15 years old and then plateaus. Similar trajectory were demonstrated for CA1, DG/CA3, and subicum, with volumes peaking in the 13- to 15-year age range. The entorhinal cortex did not show such a trajectory, and instead evidenced a flat trend, indicating it did not show age-related differences that were dissociable from the differences seen in the rest of the cortex. Results of the analysis of hippocampal–cortex volume ratio are presented in Figure 2 and Table 1. These are plotted against the age-related differences seen at the whole-brain level (gray line). For the ratio, the age-related difference trajectory was linear, with no evidence of an asymptote. Thus, the ratio of the hippocampal volume relative to the cortex reflects a continued increase with age.

#### 3.2 Results of analysis 2: Age-related differences in FA and MD of the hippocampal cingulum, fornix, and uncinate fasciculus

The results of Analysis 2 are presented in Figure 3 and Table 1. These results reflect the FA and MD of the pathways of interest after controlling for whole-brain FA and MD. The analysis showed (bilaterally) a positive slope for the cingulum, a negative slope for the fornix, and no discernible age-related differences for the uncinate fasciculus. However, even for the cingulum and fornix, the differences over this age range were small, and no asymptote was apparent. In order to facilitate comparison with prior work, we report plots of uncorrected FA and MD in Figure S1. In these plots, we did not control for the whole-brain FA and MD values.

#### 3.3 Results of analysis 3: Relation of morphologic and diffusion metrics to performance on the NIH Toolbox PSMT

The results of Analysis 3 are presented in Figure 4 and Table 2. The volume of bilateral hippocampus, bilateral DG, left CA1, and right DG/CA3 were positively associated with performance on the PSMT, our episodic
memory task. The ratio of hippocampal to cortical volume (bilaterally) was also positively associated with performance on the PSMT. For analyses of relations between FA or MD of any of the fiber pathways and episodic memory performance, we found only right fornix MD was statistically associated with episodic memory, but this did not survive correction for multiple comparisons.

4 DISCUSSION

The hippocampus is a complex structure composed of several distinct subfields and has been at the center of scientific study examining the neural foundations of episodic memory. In order to understand episodic memory, it is necessary to establish the boundaries that define typical development of the structures that support it, namely, the hippocampus. Here, in a large sample of 830, 3- to 21-year-olds from a unique and publicly available dataset, we examined age-related differences in the volume of the hippocampus, its subfields, and in the diffusion properties of the fiber pathways connecting hippocampus to other cortical and subcortical structures. We found greater hippocampal and subfield volume exhibiting a nonlinear relation with age peaking in the mid-teen years. A similar relation between age and volume in an adjacent cortical region, the entorhinal cortex, was not observed. We
Table 1: Model fit for the linear and quadratic models for each region and pathway

<table>
<thead>
<tr>
<th>Region</th>
<th>Hemisphere</th>
<th>Linear model AIC</th>
<th>Quadratic model AIC</th>
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<tr>
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<td>2325.7***</td>
<td>2313.7***</td>
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<tr>
<td></td>
<td>Right</td>
<td>2348.30 n.s.</td>
<td>2342.80 n.s.</td>
</tr>
</tbody>
</table>

Note: Bold denotes the smaller Akaike information criterion (AIC), indicating better model fit. 
*p < .05; **p < .01; ***p < .001; n.s. = nonsignificant; † p value survived false discovery rate correction.

also found that while cortical volume decreased with age, hippocampus volume continued to increase. Age-related elevations in gray matter volume in right DG/CA3 was significantly positively related to episodic memory performance. Finally, we examined how MTL white matter pathways were associated with episodic memory, but did not find any evidence, in this dataset, of a significant relation between these pathways and age nor episodic memory performance. We discuss each of our findings in turn.

4.1 Hippocampal subfield and entorhinal cortex gray matter volume development

We first examined the basic volumetric differences associated with age in regions supporting episodic memory, including the entorhinal cortex, and the hippocampus and its subfields. We found nonlinear developmental trajectories that continue to evidence greater volumes with age well into the mid-teen years for the whole hippocampus, as well as for the hippocampal subfields CA1 and DG/CA3. We did not see the same protracted increase in volume in either the subiculum or the entorhinal cortex, the latter a major cortical input to the hippocampus.

Whole-hippocampal volume exhibits a protracted developmental trajectory. Many studies have examined the structural development of the hippocampus (Goddings et al., 2014; Lee et al., 2015; Østby et al., 2009; Tamnes et al., 2013; Uematsu et al., 2012; Yurgelun-Todd et al., 2003). Our findings—greater age-related hippocampal volumes well into mid-teen years—are consistent with prior work that similarly observed nonlinear age-related increases in volume that peaked in adolescence (Østby et al., 2009; Uematsu et al., 2012). However, examining the hippocampus as a whole structure has yielded somewhat inconsistent results, especially in younger samples. For example, Gogtay and colleagues (2006) reported no age-related differences in overall hippocampal volume between 4 and 25 years, while Tamnes and colleagues (2013) observed a slight decrease in volume between 8 and 22 years. Discrepancies in findings across studies may arise due to differences in segmentation methods (e.g., manual vs. automated),
The hippocampus is not a uniform structure. Notable differences in anatomical connectivity and genetic, molecular, and functional contributions have been identified along its longitudinal axis (Fanselow & Dong, 2010; Poppenk, Evensmoen, Moscovitch, & Nadel, 2013; Strange, Witter, Lein, & Moser, 2014; Vogel et al., 2020). Further, the hippocampal formation is composed of histologically distinct subfields thought to make unique functional contributions to episodic memory (Marr, 1971). Accordingly, more recent studies have complimented their examination of whole hippocampal development by segmenting the hippocampus into subregions along the long axis (i.e., head, body, and tail or by anterior and posterior hippocampus; see Canada et al., 2020; Canada et al., 2021; DeMaster et al., 2013; Lee et al., 2015; Riggins et al., 2015) or subfields. Riggins and colleagues (2015) showed that 6-year-olds had marginally larger overall hippocampal volume than 4-year-old children, but in a follow-up longitudinal study with 4- to 8-year-olds showed a quadratic pattern of change in the head of the hippocampus, with a more monotonic pattern in the body and tail (Canada et al., 2020). As noted earlier, Gogtay and colleagues (2006) did not observe age-related differences in overall hippocampal volume, but when examined along the long axis (i.e., anterior/posterior hippocampus) the anterior hippocampus showed an increase in volume with age, while the posterior hippocampus decreased in volume with age between 4 and 25 years.

Differences in hippocampal subfield volume have been identified across a broad range of ages. When examined from early-to-mid-childhood, the CA1 increased in volume between 4 and 5 years, while the CA2–4/DG and subiculum peaked in volume between 5 and 6 years (Canada et al., 2021). Across a broader age range, all studies have identified nonlinear positive associations with age across hippocampal subfield peaking in adolescence (Krogsrud et al., 2014; Lee et al., 2014; Tamnes et al., 2018). Similarly, we observed nonlinear increases in CA1 and DG/CA3 volumes with age that peaked in mid-teen years, but this pattern was not observed for the subiculum in our sample. Our results, taken together with prior studies, suggest that segmentation of the hippocampus into subfields or regions along its long axis provides a more accurate picture of developmental differences that can be masked when the hippocampus is examined as a whole.
TABLE 2  Summary of regression results for regions and pathways of interest predicting episodic memory performance

<table>
<thead>
<tr>
<th>Region</th>
<th>B</th>
<th>SE (B)</th>
<th>β</th>
<th>df</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Left hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left hippocampus</td>
<td>0.00</td>
<td>0.00</td>
<td>0.08</td>
<td>729</td>
<td>.03*</td>
</tr>
<tr>
<td>Left entorhinal cortex</td>
<td>0.00</td>
<td>0.00</td>
<td>0.01</td>
<td>729</td>
<td>.79</td>
</tr>
<tr>
<td>Left CA1</td>
<td>0.01</td>
<td>0.02</td>
<td>0.08</td>
<td>729</td>
<td>.02*</td>
</tr>
<tr>
<td>Left DG/CA3</td>
<td>0.02</td>
<td>0.01</td>
<td>0.07</td>
<td>729</td>
<td>.04*</td>
</tr>
<tr>
<td>Left subiculum</td>
<td>0.02</td>
<td>0.02</td>
<td>0.04</td>
<td>729</td>
<td>.26</td>
</tr>
<tr>
<td>Left fornix FA</td>
<td>1.16</td>
<td>11</td>
<td>0.00</td>
<td>729</td>
<td>.92</td>
</tr>
<tr>
<td>Left cingulum FA</td>
<td>-14.06</td>
<td>7.88</td>
<td>0.06</td>
<td>729</td>
<td>.08</td>
</tr>
<tr>
<td>Left uncinate fasciculus FA</td>
<td>5.38</td>
<td>12.80</td>
<td>0.01</td>
<td>729</td>
<td>.67</td>
</tr>
<tr>
<td>Left fornix MD</td>
<td>3.75</td>
<td>3.31</td>
<td>0.03</td>
<td>729</td>
<td>.26</td>
</tr>
<tr>
<td>Left cingulum MD</td>
<td>4.54</td>
<td>6.08</td>
<td>0.02</td>
<td>729</td>
<td>.46</td>
</tr>
<tr>
<td>Left uncinate fasciculus MD</td>
<td>-15.21</td>
<td>15.96</td>
<td>-0.03</td>
<td>729</td>
<td>.34</td>
</tr>
<tr>
<td><strong>Hippocampal cortical ratio</strong></td>
<td>518.93</td>
<td>221.23</td>
<td>0.08</td>
<td>730</td>
<td>.02*</td>
</tr>
<tr>
<td><strong>Right hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right hippocampus</td>
<td>0.00</td>
<td>0.00</td>
<td>0.07</td>
<td>729</td>
<td>.04*</td>
</tr>
<tr>
<td>Right entorhinal cortex</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>729</td>
<td>.93</td>
</tr>
<tr>
<td>Right CA1</td>
<td>0.01</td>
<td>0.00</td>
<td>0.05</td>
<td>729</td>
<td>.14</td>
</tr>
<tr>
<td>Right DG/CA3</td>
<td>0.03</td>
<td>0.01</td>
<td>0.08</td>
<td>729</td>
<td>.001***</td>
</tr>
<tr>
<td>Right subiculum</td>
<td>0.02</td>
<td>0.02</td>
<td>0.03</td>
<td>729</td>
<td>.36</td>
</tr>
<tr>
<td>Right fornix FA</td>
<td>-13.07</td>
<td>11.18</td>
<td>0.03</td>
<td>729</td>
<td>.24</td>
</tr>
<tr>
<td>Right cingulum FA</td>
<td>-9.46</td>
<td>8.87</td>
<td>0.03</td>
<td>729</td>
<td>.29</td>
</tr>
<tr>
<td>Right uncinate fasciculus FA</td>
<td>-19.49</td>
<td>14.87</td>
<td>0.05</td>
<td>729</td>
<td>.19</td>
</tr>
<tr>
<td>Right fornix MD</td>
<td>8.64</td>
<td>3.72</td>
<td>0.07</td>
<td>729</td>
<td>.02*</td>
</tr>
<tr>
<td>Right cingulum MD</td>
<td>-2.79</td>
<td>6.95</td>
<td>-0.01</td>
<td>729</td>
<td>.69</td>
</tr>
<tr>
<td>Right uncinate fasciculus MD</td>
<td>-17.60</td>
<td>14.74</td>
<td>-0.05</td>
<td>729</td>
<td>.23</td>
</tr>
<tr>
<td><strong>Hippocampal cortical ratio</strong></td>
<td>426.03</td>
<td>216.35</td>
<td>0.07</td>
<td>730</td>
<td>.049**</td>
</tr>
</tbody>
</table>

*p < .05; **p < .01; ***p < .001; † p-value survived false discovery rate correction.

*FIGURE 4  Association between right DG/CA3 volume and episodic memory performance. Only the false discovery rate-corrected association from Table 2 is shown. **p < .001. Uncorrected nominally significant associations from Table 2 are shown in Figure S2.

Our findings add to the body of work suggesting a protracted hippocampal development of the hippocampus and subfields (with the exception of subiculum) at least into adolescence (Goddings et al., 2014; Lee et al., 2015; Østby et al., 2009; Riggins et al., 2018; Tamnes et al., 2013; Uematsu et al., 2012; Yurgelun-Todd et al., 2003). While some studies have suggested that there is an early peak in hippocampal development or found no age-related differences (Barnea-Goraly et al., 2014; Giedd et al., 1996; Gogtay et al., 2006; Joshua K Lee et al., 2015; Riggins et al., 2015; Uematsu et al., 2012), the differences in findings are potentially related to the variety of different age ranges studied. However, when more extended age ranges are included, hippocampal development seems to extend into mid-adolescence. Indeed, one shortcoming of previous research is that the present study alleviates is the examination of varying ages within truncated time frames (DeMaster et al., 2013; Joshua K Lee et al., 2014; Stine K Krogsrud et al., 2014; Tamnes, Bos, van de Kamp, Peters, & Crone, 2018). The major contribution of the present study is the examination of a wide age range in a large sample with a good representation at each age range. This allows us to zoom out and take a “birds-eye view” of the development of the
hippocampus, beginning from a very young age (3 years) through the adolescent years and into early adulthood.

Interestingly, even with our large sample we did not find that the entorhinal cortex followed the same trajectory of volumetric change over age as the hippocampus and its subregions. From one perspective, this may be surprising, given the extensive connectivity of the entorhinal cortex to the rest of the hippocampus. However, the results also suggest that volumetric differences can occur along differential timelines, even for highly connected regions. This is consistent with data from Alzheimer’s studies showing that changes in entorhinal cortex volume as a result of the disease do not immediately track with changes in hippocampal volume (Killiany et al., 2002). In terms of typical development, it is possible that pruning of hippocampal synapses follows a different timeline than pruning of neocortical synapses. Significant work has investigated axonal and dendritic pruning in the rodent hippocampus, and quite a bit is known from those studies (Faulkner, Low, & Cheng, 2007). However, these processes are far more difficult to investigate in the human hippocampus, and thus differences in MRI volume must be used. However, even here we must be cautious. A recent paper has suggested that differences in MRI measurements over development (in this case, cortical thinning) may be more associated with changes in myelination, and not changes in gray matter morphology, as was previously assumed (Natu et al., 2019). We speculate that it is possible that myelination is proceeding differently in the hippocampus than it is in the cortex, which could drive differences in volume as measured by MRI. However, further research would need to be conducted to support this claim. Thus, these two possible explanations may explain why we see developmental differences in the volume of hippocampus compared with the neighboring entorhinal cortex.

4.2 Hippocampal development in relation to the cortical gray matter

Hippocampal development progresses in the context of broader maturation changes across the brain. In the interest of investigating the relation between the hippocampal and cortical developmental trajectories, we examined differences in the ratio of hippocampal volume to whole brain cortical gray matter volume with age. If hippocampal development was following in lockstep with the developmental course of the cortex, then no discernable pattern would be evident across age. The linear pattern that emerged when examining the hippocampal–cortical ratio, on the contrary, highlights a tight relation between hippocampal and cortical development. Research on differences in whole brain cortical gray matter volume has demonstrated that cortical volume peaks earlier in childhood, and decreases in late childhood and throughout adolescence (Lebel & Beaulieu, 2011; Tanné et al., 2013; Wierenga et al., 2014). The findings of the current study suggest that the hippocampus follows a somewhat inverse trajectory. This a first step, however, and the issue of how the hippocampus develops in relation to the broader cortical network to which it is connected deserves additional scrutiny.

4.3 Hippocampal volumetric associations with behavior

The hippocampus is an important region for episodic memory; thus, development of this structure should be related to memory performance in tasks that tax functions related to episodic memory. We found that bilateral hippocampus and subfield volumes (CA1 and DG/CA3) were all positively associated with performance on the PSMT, even after controlling for a number of relevant covariates (age, gender, ethnicity, parent highest level of education, household income, scanner type, and ICV). Notably, only DG/CA3 survived the correction for multiple comparisons, and not all subfields exhibited a similar relation between volume and PSMT performance. It may be that specific subfields of the hippocampus play differential roles in the mechanics of episodic memory (Kim & Yassa, 2013; Wendelken et al., 2014), and this may differ across development (Lee et al., 2020). For example, research in patients with damage to the CA1 shows that they have extensive episodic memory loss suggesting an important role for CA1 in episodic memory (Zola-Morgan, Squire, & Amaral, 1986). In children, DG/CA3 volume has been positively related to episodic memory (Lee et al., 2014) and greater volume in CA1 and CA2–3 has been related to recall and retention after an extended delay (Krogsgrud et al., 2014). Our results are somewhat consistent with these previous studies. While we found that both DG/CA3 and CA1 volumes (the latter uncorrected) predicted PTSM scores, we did not find the same relations for subiculum.

We also showed that the ratio of the hippocampus to the rest of the cortex was positively associated with PTSM scores, although this did not survive correction for multiple comparisons. Considered in the context of this caveat, this finding suggests that children who retain or increase hippocampal volume as cortical volume declines exhibit better episodic memory. It remains to be seen whether the association between behavior and hippocampal development in the context of cortical maturation is specific to retention of hippocampal volume with developmentally appropriate cortical reduction, increase of hippocampal volume in relation to steady cortical volume across age, or more likely a manifestation of the combination of hippocampal increases and cortical reductions across development—with those individuals exhibiting the steepest divergence between these two measures reaping the largest mnemonic enhancement.

4.4 White matter connectivity with the rest of the brain

In our final analysis, we examined how several white matter pathways, which have been identified as playing important roles in episodic memory, contribute to episodic memory development. Unexpectedly, we found almost no evidence for an association between diffusion indices of the three pathways and PTSM scores. An exception was an association between right fornix MD and PTSM performance; however, this did not survive a multiple comparison correction. A number of
studies (Krogsrud et al., 2016; Loenneker et al., 2011; Moon et al., 2011; Simmonds, Hallquist, Asato, & Luna, 2014) have shown a protracted timeline of white matter development from early childhood until adulthood, with different rates of maturation in different pathways (for review, see Lebel, Treit, & Beaulieu, 2017). We expected that such differences would be associated with behavioral differences in episodic development in hippocampal connectivity. Indeed, this has been found in some other studies (Mabott et al., 2009; Ngo et al., 2017). For example, Ngo and colleagues (2017) examined the fornix and uncinate fasciculus in a small sample of 4- and 6-year-olds. Their results revealed that white matter pathways connecting hippocampus and inferior parietal lobule significantly predicted episodic memory performance. In a more recent paper examining a sample of 66, 4- to 8-year-olds, Hoffman and colleagues (2022) showed robust associations between fornix diffusion metrics and several episodic memory measures, including story recall, source memory, and temporal memory measures. They did not find a similar association with the uncinate fasciculus. Surprisingly, the current data are not consistent with these studies that do find significant associations. One possible explanation for this may be that prior studies did not control for the covariates that we controlled for, including whole brain FA. This may have masked associations that were found in previous studies. However, we argue that such controls are necessary. For example, if whole brain white matter development is largely explanatory, it does not tell us anything about the specific fiber pathways connected to the hippocampus. In addition, studies based on small samples may actually inflate the reporting of effect sizes, especially in studies addressing brain–behavior associations (Dick et al., 2021). It is therefore possible that such relations reported in smaller samples are not reliable when larger samples are investigated. At the same time, the null findings for all three pathways were surprising, especially given the increasingly replicated associations between fornix white matter and episodic memory. Thus, other shortcomings of the present study may explain the lack of findings. For example, the measures used in other studies may be more sensitive measures of episodic memory development. It is also possible, in the case of the fornix, that quantification of diffusion metrics was contaminated by neighboring cerebrospinal fluid (Hoffman et al., 2022). Future studies should take into account these possible sources of contamination and incorporate a wider range of episodic memory measures.

4.5 Limitations

There are several important limitations of the current study that must be considered. First, the study is cross-sectional instead of longitudinal. This limits the degree to which we can make inferences about age-related changes in hippocampus and related connections. This fact was highlighted in a recent paper by Keresztes and colleagues (2022). In that paper, the authors investigated both longitudinal and cross-sectional differences in a sample of 109 6- to 10-year-olds. The longitudinal and cross-sectional results did not necessarily agree. However, the authors point out that only two time points were examined for the longitudinal change, across a narrow age range. As they indicate, if change is nonlinear, as we found in the present study for some of our effects, the use of only two time points would preclude analysis of nonlinear effects. They also pointed to the importance of examining both longitudinal and cross-sectional effects in future studies, as the nature of the differences must be examined in large samples with multiple longitudinal time points.

A second limitation is the fact that participants were collected at multiple sites on different scanners. One advantage of this is that the sample is more diverse than is typical, representing lived experiences across, in this case, multiple cities in the United States. However, the fact that these are MRI research centers means they tend to be localized to cities, underrepresenting rural participants and the potential effects of those developmental environments. In addition, different scanners contribute sources of noise. We have attempted to deal with some of this using COMBAT correction (Fortin et al., 2017; Johnson et al., 2007) and by including device serial number as a regressor. These methods are shown to be effective in reducing substantial sources of noise in imaging data, thus increasing statistical power (Radua et al., 2020). However, we cannot remove all sources of noise, and the results should be considered in the context of this limitation. Thus, the sampling strategy of the PING study does limit, to some degree, the external validity of the results, and notably whether the results can be generalized to populations who did not have the same rearing experience as those raised in cities in the United States.

Third, methodological differences with prior studies can lead to different results, especially when it comes to examination of subfields of the hippocampus. The most ideal method of subfield segmentation is manual segmentation, but a field-wide harmonized protocol for both manual and automated segmentation has yet to be developed (Wisse et al., 2017; Yushkevich et al., 2015). Although the automated method allowed us to exploit a dataset with a larger sample size than would have been feasible using manual segmentation methods, it is important to acknowledge its limitations. Giuliano et al. (2017) recently investigated this issue and showed that, indeed, different volumetric estimates are reported when using different methods. But the authors also showed that automated methods can give excellent results, especially for the FreeSurfer 6.0 method we used here (Iglesias et al., 2015; Mueller et al., 2018). Indeed, several studies report that the FreeSurfer 6.0 method has high test–retest reliability within and across magnets, ranging from .5 to .9 intraclass correlation coefficient (Brown et al., 2020; Elvashagen et al., 2016; Quattrini et al., 2020; Whelan et al., 2016; Worker et al., 2018). Samann and colleagues (2022) provided a recent, comprehensive review of the FreeSurfer 6.0 method, suggesting that it is a valuable tool with good measurement reliability and validity, with the explicit advantage of reducing methodological heterogeneity for large-scale collaborative studies, such as PING. That said, the method is not perfect (Mueller et al., 2018; Seiger et al., 2021; Wisse, Biessels, & Geerlings, 2014). For example, segmentation of the subiculum from CA1 differs substantially among different protocols, and indeed differences among protocols are greatest in the anterior hippocampus relative to the body and tail portions (Yushkevich et al., 2015). Our reported results for subiculum should be especially...
interpreted with caution. Although some of the issues raised about the automated FreeSurfer method have been mitigated by updates in more recent versions (especially going from 5.0 to 6.0; Iglesias et al., 2015; Mueller et al., 2018), the results we report should be considered in the context of the method.

5 | SUMMARY

In summary, we found in a large sample of 830, 3- to 21-year-olds nonlinear volume increases in the hippocampus and related subfields that peaked in the midteen years. These trajectories controlled for differences in the general size of the brain that also occur during childhood and adolescence. We also found significant age-related differences in global hippocampal volume when considered in the context of cortical gray matter volume. That is, we found that as cortical volume decreased with age, hippocampus volume continued to increase. In addition to age-related differences in gray matter volume, we found that several distinct subfields, specifically CA1 and DG/CA3, are significantly related to episodic memory as assessed using the PSMT. Finally, we examined how white matter pathways were associated with episodic memory, but did not find any evidence in our large sample to suggest that these pathways significantly influenced episodic memory development. These findings contribute to further understanding of hippocampal development over an extended age range, and its associated effects on episodic memory development.

AUTHOR CONTRIBUTIONS

Anthony Steven Dick and Aaron T. Matfeld conceived of the study. Anthony Steven Dick and Kristafor Farrant wrote the initial draft, conducted initial data analysis, constructed figures, and revised the manuscript. Yvonne Ralph, Bethany Reeb-Sutherland, Shannon Pruden, and Aaron T. Matfeld contributed to manuscript revisions.

ACKNOWLEDGMENT

This study is supported by the Eunice Kennedy Shriver National Institute of Child Health and Human Development (grant number R01HD098152 to S.P., B. R-S., and A. M.).

CONFLICT OF INTEREST

The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

The current cross-sectional study was conducted using MRI data from the publicly available database, Pediatric Imaging, Neurocognition, and Genetics (PING) (http://pingstudy.ucsd.edu). Data are thus available to the scientific community through PING.

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