Predictability matters: role of the hippocampus and prefrontal cortex in disambiguation of overlapping sequences

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Previous research has demonstrated that areas in the medial temporal lobe and prefrontal cortex (PFC) show increased activation during retrieval of overlapping sequences. In this study, we designed a task in which degree of overlap varied between conditions in order to parse out the contributions of hippocampal and prefrontal subregions as overlap between associations increased. In the task, participants learned sequential associations consisting of a picture, a face within the picture frame, and an outdoor scene. The control condition consisted of a single frame-face-scene sequence. In the low overlap condition, each frame was paired with two faces and two scenes. In the high overlap condition, each frame was paired with four faces and four scenes. In all conditions the correct scene was chosen among four possible scenes and was dependent on the frame and face that preceded the choice point. One day after training, participants were tested on the retrieval of learned sequences during fMRI scanning. Results showed that the middle and posterior hippocampus (HC) was active at times when participants acquired information that increased predictability of the correct response in the overlapping sequences. Activation of dorsolateral PFC occurred at time points when the participant was able to ascertain which set of sequences the correct response belonged to. The ventrolateral PFC was active when inhibition was required, either of irrelevant stimuli or incorrect responses. These results indicate that areas of lateral PFC work in concert with the HC to disambiguate between overlapping sequences and that sequence predictability is key to when specific brain regions become active.

We often rely on contextual information to retrieve specific events or sequences of events. Accumulating evidence indicates that interactions between the hippocampus (HC) and prefrontal cortex (PFC) are critical to the retrieval of events with overlapping elements that rely on contextual information (Rich and Shapiro 2009; Brown et al 2010, 2012). While some items or places have only one or two associations connected to them, other items may have multiple overlapping associations. For example, Chicago’s O’Hare airport may have two distinct events connected to it—the time you were forced to stay in the airport overnight due to a snowstorm and when you got food poisoning from the sushi restaurant in the food court—whereas the local hospital has many associations—it is where you gave birth to each of your children and had a broken leg and visited your dad when he was sick. Is there a difference in how we retrieve memories that have few versus multiple overlapping associations? In this study we were interested in investigating the following questions: As relational load increases how does activation of the HC and PFC and their associated networks change? How does ability to predict or think through an upcoming sequence affect neural activation at both regional and whole brain levels? Disambiguating between similar events or associations is thought to rely on pattern separation, a process dependent on the dentate gyrus of the HC and related medial temporal lobe structures. Pattern separation involves the creation of distinct neural representations for stimuli that are the same but used in different contexts (Yassa et al. 2011; Yassa and Stark 2011; Reagh and Yassa 2014; Bennett and Stark 2016). A series of studies in rodents have shown that the HC forms distinct representations for overlapping sequences in both spatial and nonspatial tasks (Frank et al. 2000; Wood et al. 2000; Agster et al. 2002). Human neuroimaging studies have also shown hippocampal activation during the encoding and retrieval phases of both spatial and nonspatial context-dependent memory tasks with overlapping stimuli, including single items and sequences (Bakker et al. 2008; Ross et al. 2009; Brown et al. 2010; Lacy et al. 2011; Newmark et al. 2013). Recently, multi-voxel pattern analyses of functional magnetic resonance imaging (fMRI) in the HC has shown that learning causes differentiation of hippocampal activation patterns for overlapping stimuli (LaRocque et al. 2013; Favila et al. 2016; Tompary and Davachi 2017). In addition to these functional differences, the ability to accurately and efficiently disambiguate between overlapping mazes is correlated with structural differences in the HC and rostral dorso-lateral PFC (Brown et al. 2014).

When items and associations are shared between sequences of events, the PFC is involved in synthesizing episodic memories into relational sets that evolve with experience and cognitive demands and contribute to behavioral flexibility (Miller and Cohen 2001; Eichenbaum 2004; Hasselmo 2005; Rich and Shapiro 2009). It is well established that the PFC plays an integral role in the retrieval of episodic memory, with different regions responding to different

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Participants learned sequences of contextual cues and associated face-scene combinations. Figure 1.

Based on previous studies of hippocampal function, we expected to find increased hippocampal activation in this task. In both animals and humans, memory for sequences of items has been shown to rely on the HC (Agster et al. 2002; Schendan et al. 2003; Brown et al. 2010, 2014; Hsieh and Ranganath 2015), and previous work in humans has suggested that different functions may be related to detailed (posterior) and gist-like (anterior) hippocampal representations (Poppenk et al. 2013). In our task, the low overlap condition contained a larger pool of stimuli than the high overlap condition. In contrast, the high overlap condition did not contain as many stimuli as the low overlap condition but there was an increased demand to separate, or disambiguate, each trial’s sequence from the three sequences with which it overlapped. Therefore, we expected to find increased activation in both the anterior and posterior HC where item and context information are combined and the anterior HC which is thought to relay context information to the PFC (Eichenbaum et al. 2012; Komorowski et al. 2013; Poppenk et al. 2013). We also expected to see increased activation in the anterior HC during the association period of the low overlap condition because it is at this point that the HC was able to relay information to the PFC pointing to the appropriate response to complete the sequence.

We expected to find differential activation in the vIPFC, based on differences in relational load and the degree of interference from overlapping items (Jonides and Nee 2007; Nee et al. 2006, Oren et al. 2017). Because the high overlap condition had the highest degree of interference from items not relevant to the current trial, we expected to see the most vIPFC activity during this condition, particularly at the association time point. In addition we anticipated differences in activation in the dIPFC between conditions. One proposed role of the dIPFC is executive control of neural regions processing task relevant representations involved in response preparation (Rowe et al. 2000). In our task, we would expect the dIPFC to be active when there is information present that determines the representations relevant to the trial. In the low overlap condition, this would be during both the cue and association periods and in the high overlap condition this would be during the association period. In the control condition, the participant should be able to predict the entire sequence at the cue period. However, due to the simplicity of the sequence used in the control condition and the number of times this trial is repeated, we did not expect to see activation in the PFC.

Results

Behavioral data
Participants were trained on the task 1 d prior to performing the task during fMRI scanning. The task consisted of one of 13 possible picture frames, each differentiated by color, pattern, and/or texture, presented on the computer monitor followed by the photograph of a face within the picture frame, followed by four outdoor scenes (see Fig. 1). The correct response changed with each frame/face combination that was presented.

The control consisted of a simple frame-face-scene sequence. The low overlap condition consisted of four pairs of sequences. Each frame was paired with two faces and two scenes. For example, when the green frame was shown before the old man the correct response was the vineyard but when paired with the younger man the correct response was the grassy field. However, when the old man was shown after the fire frame the correct
response was the grassy field and when the young man was shown after the fire frame the correct response was the vineyard (see Fig. 2).

The high overlap condition had the highest degree of overlap between associations and the lowest level of predictability. In the high overlap condition, the participant learned that each frame was associated with four different faces and four scenes. There were four frames in the high overlap condition and a total of sixteen overlapping associations (see Fig. 3).

We examined differences in accuracy and reaction times between the control, low overlap, and high overlap conditions while participants performed the task in the scanner. Participants responded correctly in the control condition on 96.7% of the trials (SD = 0.019), in the low overlap condition on 94.23% of the trials (SD = 0.049), and in the high overlap condition on 90.93% of the trials (SD = 0.102), A repeated measures ANOVA revealed a significant main effect of condition ($F_{(2,16)} = 4.00; P = 0.028$;
Post-hoc tests revealed that the main effect of condition was driven by differences in accuracy between the high overlap and control conditions ($P = 0.035$, Fisher’s Least Significant Difference (LSD) post-hoc test). No significant differences in percentage correct were found between the low and the high overlap conditions, indicating that the two overlapping sequences were performed with comparable accuracy ($P = 0.156$, Fisher’s LSD post-hoc test).

Mean reaction time for the control condition was 1.45 sec (SD = 0.404), for the low overlap condition was 1.55 sec (SD = 0.5116), and for the high overlap condition was 1.56 sec (SD = 0.19). A repeated measures ANOVA revealed a significant main effect of condition on reaction time ($F_{(2,16)}^2 = 8.811$, $P = 0.001$; $\eta^2_p = 0.355$). Post-hoc tests revealed that the main effect of condition was driven by differences between the overlapping and the control conditions. Responses in the control condition were significantly faster than in the low overlap condition ($P = 0.01$; Fisher’s LSD post-hoc test) and the high overlap condition ($P = 0.002$, Fisher’s LSD post-hoc test). The two overlapping conditions did not differ from one another in reaction time ($P = 0.097$, Fisher’s LSD post-hoc test).

### Functional MRI data

We used fMRI to assess activation during sequence retrieval in regions of interest: the HC, the dorsolateral PFC, and the ventrolateral PFC. To facilitate interpretation of our results, we averaged the beta values for each subject in each condition within ROIs and determined the overall mean beta value per condition at each timepoint and each region of interest (see Figs. 4 and 5 for bar charts).

**Figure 4.** Beta weights from each condition at both time points of interest (Cue and Association). The conditions are presented in order of relational load within ROI and timepoint: high overlap (red), low overlap (blue), and control (green).

**Figure 5.** A graph showing the average beta weight values for the dorsolateral and ventrolateral PFC during the cue and association periods. Although the ROI used for the small volume correction analysis was comprised of the whole lateral PFC, we discussed the results from this analysis separately for dorsolateral and ventrolateral PFC and therefore show their individual beta weight values.
graphs of beta averages within the ROI masks used in the small volume correction analyses. In addition, we examined activation at the whole brain level in an exploratory analysis. For both analyses, six contrasts were analyzed: the high and low overlap conditions compared to control, the high and low overlap conditions compared to each other, and the control condition compared to the high and low overlap conditions. The results are presented separately for the High Overlap, Low Overlap, and Control conditions. Results are summarized in Tables 1 (ROI based) and 2 (Whole brain).

**Small volume correction analyses**

**High overlap condition.** In the high overlap condition, the frame indicated to the participant which set of stimuli the trial belonged to but it was not until the face was shown within the frame that participants were able to think about which scene would accurately complete the sequence. The results from our small volume correction analyses showed that the high overlap condition was associated with increased activation in the posterior body of the HC compared to the control condition during the association period (see Fig. 6).

### Table 1. Small volume correction analysis complete results

| Condition                          | Region of interest     | Cluster size (voxels) | t-value | MNI peak coordinates |
|------------------------------------|------------------------|-----------------------|---------|----------------------|
| High overlap > control association | Right HC (body)        | 98                    | 5.27    | 24 – 28 – 8          |
|                                    | Right dorsolateral PFC | 2710                  | 9.41    | 48 26 – 28           |
|                                    | Left dorsolateral PFC  | 468                   | 4.80    | – 56 28 – 24         |
| Low overlap > control cue          | Left HC (tail)         | 43                    | 4.62    | – 12 – 36 0          |
| Low overlap > control association  | Right HC (tail)        | 20                    | 3.62    | 18 – 34 0            |
| Control > high overlap association | Right dorsolateral PFC | 1510                  | 7.35    | 48 18 32             |
| Control > low overlap association  | Left ventrolateral PFC | 48                    | 3.93    | – 52 10 12           |
|                                    | Left ventrolateral PFC | 27                    | 3.53    | – 54 10 12           |

**High load > control cue** Right HC (body) 1472 3.99 26 – 70 30
Left caudate 888 5.45 – 8 1 – 3
Left cuneus 683 4.98 – 18 – 58 26
Right intraparietal sulcus 402 5.39 – 30 – 66 34

**High load > control association**
Right HC (body) 98 5.27 24 – 28 – 8
Right dorsolateral PFC 2710 9.41 48 26 – 28
Left dorsolateral PFC 468 4.80 – 56 28 – 24
Left ventrolateral PFC 261 8.92 – 34 – 24 – 6

**Low overlap > control cue**
Left HC (tail) 43 4.62 – 12 – 36 0
Right HC (tail) 20 3.62 18 – 34 0

**Low overlap > control association**
Right dorsolateral PFC 1510 7.35 48 18 32
Left dorsolateral PFC 223 4.78 – 54 34 26

### Table 2. Whole brain analysis complete results

| Condition                          | Brain region                        | Cluster size (voxels) | t-value | MNI peak coordinates |
|------------------------------------|-------------------------------------|-----------------------|---------|----------------------|
| High load > control cue            | Right intraparietal sulcus          | 1472                  | 3.99    | 26 – 70 30           |
|                                    | Left caudate                        | 888                   | 5.45    | – 8 1 – 3            |
|                                    | Left cuneus                         | 683                   | 4.98    | – 18 – 58 26         |
|                                    | Right intraparietal sulcus          | 402                   | 5.39    | – 30 – 66 34         |
| High load > control association    | Right insula                        | 27,123                | 10.20   | 28 22 22             |
|                                    | Bilateral anterior cingulate cortex  | 1636                  | 8.76    | 4 30 42              |
|                                    | Right cerebellum                    | 1155                  | 6.95    | 40 – 30 – 32         |
|                                    | Bilateral posterior cingulate cortex| 2547                  | 5.49    | 0 – 20 30            |
|                                    | Left angular gyrus                  | 469                   | 6.36    | 44 – 70 34           |
|                                    | Left inferior parietal lobule       | 8752                  | 7.79    | – 34 – 50 40         |
| Low load > control cue             | Left lateral occipital gyrus        | 3432                  | 6.72    | – 34 – 86 – 2        |
| Low load > control association     | Right cerebellum                    | 3102                  | 7.98    | 20 – 36 – 48         |
|                                    | Right dorsal PFC                    | 1314                  | 6.67    | 38 20 24             |
| Low load > control cue             | Right supplementary motor area      | 1119                  | 6.97    | 2 14 54              |
| Low load > high load cue           | Left middle occipital gyrus         | 1726                  | 7.79    | – 30 96 2            |
| Low load > control association     | Right angular gyrus                 | 5406                  | 8.55    | 32 – 54 38           |
|                                    | Right fusiform gyrus                | 4075                  | 7.65    | 36 – 42 – 26         |
|                                    | Right dorsal PFC                    | 2881                  | 7.61    | 52 18 28             |
|                                    | Right inferior parietal lobule      | 584                   | 7.22    | 30 – 52 32           |
|                                    | Left dorsal PFC                     | 1691                  | 6.62    | – 40 22 20           |
| Low load > control association     | Bilateral supplementary motor area   | 589                   | 5.64    | 0 14 48              |
|                                    | Left angular gyrus                  | 586                   | 7.09    | – 36 52 40           |
|                                    | Right cerebellum                    | 615                   | 6.74    | 20 36 – 48           |
| Control > low load cue             | Left middle temporal cortex         | 808                   | 4.65    | – 58 – 54 18         |
| Control > low load association     | Right middle temporal cortex        | 761                   | 5.98    | 62 – 18 – 10         |
|                                    | Right angular gyrus                 | 594                   | 5.19    | 52 – 64 28           |
|                                    | Right dorsomedial PFC               | 538                   | 5.01    | 6 62 12              |
| Control > high load association    | Bilateral dorsomedial PFC           | 3648                  | 8.32    | 0 58 4               |
|                                    | Left insula                         | 3028                  | 5.91    | – 40 – 14 – 4        |
|                                    | Right supramarginal gyrus           | 2917                  | 6.31    | 60 – 36 42           |
|                                    | Bilateral middle cingulate gyrus    | 1945                  | 7.38    | 0 – 24 44            |
|                                    | Left middle temporal cortex         | 482                   | 5.09    | – 62 – 40 4          |
| Control > low load association     | Left insula                         | 2656                  | 8.59    | – 40 – 12 – 4        |
|                                    | Right insula                        | 2544                  | 8.75    | 40 – 16 – 4          |
|                                    | Bilateral dorsomedial PFC           | 2491                  | 6.90    | 0 58 6               |
|                                    | Left middle cingulate gyrus         | 1361                  | 6.38    | – 4 – 22 46          |

www.learnmem.org 339 Learning & Memory
During this timepoint, the high overlap condition also showed increased activation in the right dorsolateral PFC, the left dorsolateral PFC, and the left ventrolateral PFC compared to the control condition. At this point, there may be interference from overlapping elements, prompting recruitment of the PFC (Irlbacher et al. 2014 for review; Jonides and Nee 2006). In addition, at this stage, the participant has enough information to determine which scene accurately completes the sequence. The dlPFC may be needed at this timepoint to activate the regions needed to correctly respond.

There were no significant differences between the low and high overlap conditions during the cue period in either the MTL or lateral PFC. In addition, the high overlap and control conditions did not show differences in activation in either region of interest during the cue period. This may be due to the fact that in the high overlap condition, the cue period enables the participant to narrow down the upcoming sequences using the four possible scenes. When the contrasts created for the low overlap cue period were compared to those of the control cue period, significantly more activation was seen in the tail of the left and right HC in the low overlap condition (Fig. 7). We expected to see activation in the HC during this condition because the cue prompts retrieval of the two faces and scenes paired with it.

During the cue period, the low overlap condition also showed increased activation in the right dorsolateral PFC compared to the control condition. The high overlap condition showed increased activation in the right and left dlPFC compared to control. During both of these time points, the dlPFC may be active due to the fact that information is being obtained revealing the appropriate neural representations needed to make a correct response.

**Control condition.** The control showed increased activation in a small posterior region of the left ventrolateral PFC.

During this timepoint, the high overlap condition also showed increased activation in the right dorsolateral PFC, the left dorsolateral PFC, and the left ventrolateral PFC compared to the control condition. At this point, there may be interference from overlapping elements, prompting recruitment of the PFC (Irlbacher et al. 2014 for review; Jonides and Nee 2006). In addition, at this stage, the participant has enough information to determine which scene accurately completes the sequence. The dlPFC may be needed at this timepoint to activate the regions needed to correctly respond.

**Low overlap condition.** Presentation of the picture frame in the low overlap condition enables the participant to narrow down the upcoming sequence from 33 possible sequences to two sequences. At this point in the sequence, the participant can predict that one of two possible faces will follow and two of four possible scenes. When the contrasts created for the low overlap cue period were compared to those of the control cue period, significantly more activation was seen in the tail of the left and right HC in the low overlap condition (Fig. 7). We expected to see activation in the HC during this condition because the cue prompts retrieval of the two faces and scenes paired with it.

During the cue period, the low overlap condition also showed increased activation in the right dorsolateral PFC compared to the control condition. The low overlap condition showed increased activation in the right and left dlPFC compared to control. During both of these time points, the dlPFC may be active due to the fact that information is being obtained revealing the appropriate neural representations needed to make a correct response.

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**Control condition.** The control showed increased activation in a small posterior region of the left ventrolateral PFC.
compared to both the low overlap and high overlap conditions during the association period. There were no other differences between the control and overlapping conditions in any region of interest.

**Exploratory whole brain analysis**

To examine additional brain regions involved in disambiguating between overlapping elements in our task, we compared the contextual cue and association segments of the trials between the three conditions in an exploratory whole brain analysis. Table 2 shows a list of the complete results from the whole brain exploratory analysis. All reported results have been defined by cluster mass with a threshold set to $P < 0.01$.

**High overlap condition.** During the cue period, the high overlap condition showed increased activation in bilateral intraparietal sulcus, left caudate, and left cuneus compared to the control condition. During the association period, regions that are part of the cognitive control network showed increased activation in the high overlap condition compared to the control condition including the insula, the dlPFC, and the anterior cingulate cortices (Cole and Schneider 2007). Compared to the low overlap condition at the same time point, the high overlap condition showed increased activity in the posterior cingulate and angular gyrus (see Fig. 8).

**Low overlap condition.** During the cue period, the low overlap condition had similar regions of activation that the high overlap condition had during the association period when both were compared to control. The low overlap condition showed increased activation in regions that are part of the cognitive control network including the dlPFC and the SMA. During the association period, the low overlap condition showed increased activation in bilateral angular gyrus, fusiform gyrus, inferior parietal lobule, and SMA compared to the control condition (see Fig. 9).

**Control condition.** The control condition showed increased activation in regions that are part of the default mode network during both

Figure 8. (Top row) Whole brain analysis revealed additional regions of activation in the high overlap condition compared to the control during the cue period. (Middle and bottom rows) During the association period, whole brain analyses revealed increased activation in the anterior cingulate cortex, the insula, and dorsolateral PFC, all regions that are part of the cognitive control network, in the high overlap condition compared to the control condition. In addition, compared to the low overlap condition, the high overlap condition showed increased activation in parietal regions including the precuneus, the cuneus, and the angular gyrus.

Figure 9. (Top row) In the cue period of the low overlap condition, regions associated with memory retrieval (HC and angular gyrus) and cognitive control (dlPFC and SMA) showed increased activation compared to the control condition. (Bottom row) In the association period, the patterns of increased activation in the high and low overlap conditions compared to control were more similar than at the cue period. At this time point the low overlap condition showed increased activation in the SMA, the right dlPFC, and the inferior parietal lobules.
the cue and association periods compared to the overlapping conditions. These regions included the medial PFC and the temporal cortex. In addition the control condition showed increased activity in the insula and middle cingulate gyrus compared to the overlapping conditions during the association period.

Discussion

We manipulated the number of overlapping elements in a context-dependent sequencing task to examine the effects of degree of overlap and sequence predictability during memory retrieval. Our results showed that during retrieval of sequences with overlapping elements, the HC was more active at time points that enabled prediction of a correct response compared to sequences without overlapping elements at the same time points. In addition, the dorsolateral PFC was active when the pool of possible upcoming stimuli was reduced and the proper set of sequences could be delineated. The ventrolateral PFC, on the other hand, was active when responses had to be inhibited due to interference from overlapping associations.

Overlapping sequences and sequence predictability modulated hippocampal activity in a context-dependent memory retrieval task

During the contextual cue period, the small volume correction analyses showed increased activation in the left and right hippocampal tail when the low overlap condition was compared to the control condition. In the low overlap condition, there were four sets of sequences with stimuli overlapping within each set. The picture frame signaled to which set of sequences the trial belonged. Because the task was well learned, the frame (cue) enabled retrieval of the two faces and scenes associated with that frame through activation in the posterior HC. Although the contextual cue in the control also enabled retrieval of its associated sequence, the cue in the low overlap condition had more stimuli associated with it than the control condition, which may have resulted in the increased activity in posterior HC seen at this time point. This result was similar to that seen in Brown et al. (2010) in which the first hallway of an overlapping maze served as the contextual cue during which the participant was able to think ahead to the overlapping hallway and plan which way to turn. During the cue period of the overlapping mazes, Brown et al. (2010) found increased activation in the posterior HC compared to the nonoverlapping mazes.

During the association phase of the trial, when a face was shown within the picture frame, the cue was paired with the stimulus that determined the correct response. At this point in all conditions, the participant should have been able to predict the scene corresponding to that face-frame pairing. In the high overlap condition, this meant a narrowing down from four possible options to one. When the high overlap condition was compared to the control condition during the association phase, increased activation was seen in the body of the right HC. The increased activity in the body of the HC may have resulted from item and context information being combined at this phase, a process that involves the HC (Komorowski et al. 2009; Eichenbaum et al. 2012). In a human neuroimaging study, using representational similarity analysis to examine fMRI data, Collin et al. (2015) determined that the mid-portion of the HC was active during presentation of directly integrated event-pair associations that were part of a multiple event narrative. Unlike the posterior HC, which was active only during directly associated events, the body of the HC became active when multiple events were linked together to form a narrative. Interestingly, when participants in our study were asked to describe their learning strategies regarding the high overlap condition many stated they created stories that bound the frame, face, and scene together to keep track of the multiple overlapping sequences.

Different regions of lateral PFC responded when overlapping elements in a context-dependent memory task increased

Considerable converging evidence indicates that the role of the PFC in memory retrieval is strategic control of processes in other brain areas and the compiling of features of related memories to provide a context for related experiences (Buckner and Wheeler 2001; Miller and Cohen 2001; Dobbins et al. 2002; Blumenfeld and Ranganath 2007; Kuhl and Wagner 2009; Preston and Eichenbaum 2013). We found differences in activation of the dorsolateral PFC that were dependent on condition and phase of the trial. The overlapping conditions had a larger number of associations related to each stimulus (high > low > control) and previous research suggests the anterior dlPFC is involved in creating and integrating complex relationships between stimuli (Kroger et al. 2002; Blumenfeld and Ranganath 2006; Murray and Ranganath 2007). In addition, another proposed role of the dlPFC is top-down activation of the neural representations appropriate for completing a task (Roe et al. 2000). During the cue period, the low overlap condition showed increased right dlPFC activation compared to the control condition. The cue period in the low overlap condition prompted recall of two possible sequences whereas the control condition required retrieval of a single sequence. In the high and low overlap conditions, we found increased bilateral dlPFC activation in the association phase of the task compared to the control condition. The association phase in both the high and low overlap conditions enabled recall of the correct outdoor scene to complete the sequence. Therefore, it may be the recall of multiple associated stimuli combined with increased relational complexity inherent to a task with overlapping associations that required the dorsolateral PFC.

During the association phase of the task, the high overlap condition showed increased activation in the left anterior and middle ventrolateral PFC. Previous research has shown recruitment of the anterior vIPFC during tasks requiring control over memory access (Barredo et al. 2016) and recruitment of left vIPFC in particular when there is interference from overlapping associations (Nee et al. 2007; Irlbacher et al. 2014). At the association time point in the high overlap condition, the participant was required to recall the correct scene that completed the sequence and inhibit the three scenes irrelevant to the trial. This result supports the proposed role of vIPFC in activating correct sequences or associations while also inhibiting irrelevant associations (Petrides 2002; Badre and Wagner 2005, Nee et al. 2007). Barredo et al. (2015) showed that anterior vIPFC is part of a functionally connected network that includes the HC among other regions.

Whole brain analyses revealed regions in the cognitive control network and the default mode network active for different conditions during this task

In contrasts comparing the low overlap and control conditions, regions that are considered part of the cognitive control network, including the dorsolateral PFC and the rostral SMA, showed increased activation during both the cue and association periods. When the high overlap condition was compared to the control at the association time point, increased activation was seen in the dorsolateral PFC, the anterior cingulate cortex, and the anterior insula, also regions considered part of the cognitive control network.
The most striking result from comparisons between the control and overlapping conditions was the strong activation in the dorsomedial prefrontal cortices (dmPFC) in the control condition. At the cue, the activation in dmPFC was increased compared to control but by the association period there was a large region of activation in the dmPFC compared to both the high and low overlap conditions. The medial PFC is a major hub of the default mode network (Buckner et al. 2008).

Limitations and conclusions
In this study we sought to better understand the neural response to disambiguation of sequences with varying degrees of overlap and the interplay of predictability in episodic retrieval and its accompanying neural response. Initially the task was constructed like a more typical sequence in which the picture frame was shown, then a photograph of a face (without the frame), followed by the choice of four scenes. We sought to replicate animal studies in which context is temporally separated from its associations and determines the appropriate response (Komorowski et al. 2009; Navawongse and Eichenbaum 2013). However, when we piloted this task, participants were not able to learn the high overlap condition when the frame was separated from the face. We sought to even out the task difficulty and accuracy scores between conditions by enclosing the face with the frame in the second time point of the trial. Participants were able to learn the high overlap condition with this change but it made our results more difficult to interpret because the frame, or cue, was no longer separate from its associated face.

In interpreting the data, knowing the strategies participants used to remember the overlapping associations may have been enlightening. We did not conduct a formal interview of participants following the testing session on day two, but in the future would add this to similar studies. In addition, it would have been interesting to ask participants to return 1 wk after scanning to test how much they retained from the task, to see if there were differences in retention between the high and low overlap conditions and if neural activation was correlated with sequence retention.

When we designed this task we expected to see linear increases in activation that were dependent on the degree of overlap in each condition. We thought that because the high overlap condition had four times as many overlapping associations between each sequence than the low overlap condition we would see the highest level of activation in the MTL and PFC in the high overlap condition. However, what we found was that the extent to which stimuli revealed information about the upcoming sequence and increased the predictability of a correct response was the critical factor that determined which brain regions were active. Both our small volume correction and whole brain analyses demonstrated patterns of activation dependent on previously learned sequence predictability at each time point. In the low overlap condition, activation in the HC occurred during the cue period at which time the participant was able to whittle down the potential upcoming sequence from a large pool of possibilities to two sequences. In the high overlap condition, it was not until the association period that the HC became active and it was not until this time point that the participant was provided enough information to predict the upcoming sequence. At the association time point, participants were able to predict the correct response in both the high and low overlap conditions and both conditions showed similar patterns of activation in the dorsolateral PFC. The dIPFC may work with the HC to decipher the correct way to complete a sequence in a task with multiple possibilities by indicating the correct set of sequences and conveying that information back to the medial temporal lobe. The vIPFC may then inhibit the incorrect responses, a process integral to the participant ultimately accurately completing the sequence.

Materials and Methods

Participants
Participants were 18–34 yr old adults recruited from the Boston University community. Written informed consent was obtained from each participant before enrollment in accordance with the experimental protocol approved by both the Partners Human Research Committee and the Boston University Charles River Campus Institutional Review Board.

Twenty-one subjects underwent training followed 1 d later by testing in the fMRI scanner. Two subjects were omitted from the data analysis due to excessive motion in the scanner. Two subjects were excluded due to accuracy scores more than two standard deviations below the mean. Seventeen subjects (11 females) were included in the final data analysis (mean age ± SD = 22.84 ± 4.92).

Procedure

Task summary
After observing a frame and a face, the participant’s task was to select the scene that correctly completed the sequence. The initial time point was the picture frame, or contextual cue, which indicated to the participant which condition the trial was. In the low overlap condition this enabled the participant to narrow down the possible upcoming sequences from 16 down to two. In the high overlap condition this indicated to the participant that there were four possible upcoming sequences, out of a possible 16 in that condition. In the control condition, because there was only one sequence, the participant knew at this time point what the correct response was. The next time point was the presentation of the face in the picture frame, or the association period. At this point in all three conditions, the participant should be able to predict the correct response. In the low overlap condition this meant eliminating one of the two possibilities. In the high overlap condition this meant narrowing down the possibilities from four to one.

A more detailed description of the task design is provided below.

Task description
The task was programmed using E-Prime 2.0 software (Psychology Software Tools 2012). A picture frame appeared on the computer monitor for 2 sec followed by a 4 sec blank delay. Then a black and white photograph within the same picture frame was shown on the screen for 2 sec followed by a 4 sec blank delay. Then four photographs of outdoor scenes were presented on the screen for 4 sec and the participant was instructed to choose one of the scenes using the number pad on the keyboard. The correct response was determined by which picture frame and face were shown in the beginning of the sequence. During training the participants received feedback based on their response—either “Correct” or “Incorrect” appeared on the screen.

Stimuli consisted of images of 13 picture frames, each distinguishable by its unique color, texture, and/or pattern, 13 black and white photographs of faces with neutral expressions placed within the picture frame, and 16 color photographs of outdoor scenes. Each trial consisted of the contextual cue period (picture frame), the association period (photograph of face within the frame), and the test phase (photographs of four outdoor scenes) (see Fig. 1). The spatial location of the four outdoor scenes shown on the screen changed between trials and was counterbalanced across trials.

The task consisted of three conditions that varied by degree of overlap between sequences. The high overlap condition consisted of one set of sequences sharing four frames, four faces, and four scenes for a total of 16 frame/face/scene combinations. In this condition the stimuli included four unique frames, four unique faces, and four unique scenes.

The low overlap condition consisted of four sets of sequences sharing two frames, two faces, and two scenes. Each set consisted of four sequences for a total of 16 frame/face/scene combinations.
In this condition the stimuli included eight unique frames, eight unique faces, and eight unique scenes.

The control condition consisted of a single unique frame/face/scene sequence.

**Training one day prior to scanning.** During the training phase of the experiment, research participants learned each contextual cue (frame) and its corresponding association-test sequence (face-scene). During the first two trials of training, only the correct outdoor scene for that frame and face was shown during the test phase. In all subsequent trials, four outdoor scenes were presented and left on the screen during the test phase until the participant made a response. The participant used a numbered pad to indicate which of the four scenes was correct for that trial and was then given feedback ("Correct" or "Incorrect") that appeared on the computer screen for 1.5 sec. If the response was incorrect, the participant was not told what the correct answer was.

During training, participants learned one new sequence per training run. In all training runs after the first run, sequences previously learned were repeated for additional rehearsal. Training runs were balanced so that each unique sequence was presented a total of eleven times during training.

**Testing during fMRI scanning.** Participants returned for fMRI scanning 1 d after training. While in the scanner, participants completed six runs of 24 trials each. Each run included eight trials per condition that were identical to the training trials except that there was a 4 sec time limit for responding and participants were not given feedback on their performance. The three conditions were interspersed within each run. The order of trials was counterbalanced across runs, and the order of runs was randomized across participants. There were 48 trials of each condition per participant.

**Image acquisition.** Images were acquired using a 3 Tesla Siemens MAGNETOM TrioTim scanner with a 32 channel Tim Matrix head coil located at the Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Harvard Medical School. A high resolution T1-weighted multishot gradient-echo (MP-RAGE) structural scan was acquired using Generalized Autocalibrating Partially Parallel Acquisitions (GRAPPA) (TR = 2530 msec; TE = 3.4 msec; flip angle = 7°; slices = 176; resolution = 1 mm isotropic). T2*-weighted BOLD images were acquired using an Echo Planar Imaging sequence (TR = 2000 msec; TE = 30 msec; flip angle = 85°; slices = 33; resolution = 3.4 × 3.4 × 3.4 mm, interleave gap of 0.5 mm). Functional image slices were aligned parallel to the long axis of the HC.

**Image preprocessing.** Functional imaging data were preprocessed using the SPM8 software package (Statistical Parametric Mapping, Wellcome Department of Cognitive Neurology, London). All BOLD images were first reoriented so the origin (i.e., coordinate x y z = [0, 0, 0]) was at the anterior commissure. The images were then corrected for differences in slice timing and were realigned to the first image collected within a series. Motion correction was conducted and included realigning and unwarping the BOLD images (Andersson et al. 2001). The high-resolution structural image was then coregistered to the mean BOLD image created during motion correction and segmented into white and gray matter images. The bias-corrected structural image and coregistered BOLD images were spatially normalized into standard MNI space using the Diffeomorphic Anatomical Registration Through Exponentiated Lie (DARTEL) algebra algorithm (Ashburner 2007) for improved inter-subject registration. BOLD images were resampled during normalization to 2 mm³ isotropic voxels and smoothed using a 6 mm full-width at half-maximum Gaussian kernel. The normalized structural images of all 17 participants were averaged after normalization for displaying overlays of functional data.

**Behavioral analyses.** A repeated measures ANOVA was run to compare accuracy and reaction time between the low overlap, high overlap, and control conditions. When significant differences were found between the three groups, a Fisher’s LSDs post-hoc test was run to determine which groups significantly differed from one another. Analyses were completed using PASW Statistics (PASW Statistics for Windows, Version 18.0. Chicago: SPSS Inc.).

**fMRI analyses.** Nine regressors of interest were created for each participant to model the fMRI data. Separate regressors were created for the following three elements in each condition: contextual cue (presentation of the frame), association phase, (presentation of the face in the frame), and test phase (presentation of outdoor scenes). Incorrect trials and the six motion parameters calculated during motion correction were added to the model as additional covariates of noninterest. Regressors from the task were constructed as a series of square waves or "boxcars." Boxcar onsets were defined by the onset of each event and extended for the duration of the event (2 sec for the contextual cue and association phases, 4 sec for the test phase). These parameters were convolved with the canonical hemodynamic response function response in SPM8.

In total, six primary contrasts were created: the low and high overlap conditions each compared to the control conditions during the cue and association periods and the low and high overlap conditions compared to each other during the cue and association periods. The model was then analyzed using the general linear model approach.

All six primary contrasts were tested at the group level.

**Small volume correction analyses.** Due to our a priori hypothesis that hippocampal and prefrontal cortical activity would be modulated by degree of overlap, we defined regions of interest (ROIs) in the MTL and the PFC to perform voxelwise analyses within these areas. The MTL ROI consisted of the HC and parahippocampal gyrus in both hemispheres, as defined by the anatomical atlases of the Wake Forest University (WFU) Pick Atlas. The PFC ROI consisted of the dorsolateral PFC, defined as the middle frontal gyrus and Brodmann areas 9 and 46, and the ventrolateral PFC, defined as the inferior frontal gyrus and Brodmann areas 44, 45, and 47. The anterior cingulate and Brodmann area libraries within the WFU pick atlas were used to define the regions. All ROIs were defined bilaterally.

All region of interest, group-level analyses assessed statistical significance on the basis of cluster mass, with the cluster-defining threshold set to the nominal P < 0.01 level. Corrected P values were determined using permutation testing (FSL’s randomise; 5000 iterations), and results were thresholded at corrected P < 0.01. For one-sample t-tests, each iteration randomly sign-flipped individual subjects’ contrast coefficient maps and added the resulting maximum cluster mass to the empirical null-hypothesis distribution.

**Whole-brain analyses.** All whole-brain, group-level analyses assessed statistical significance on the basis of cluster mass, with the cluster-defining threshold set to the P < 0.01 level. Corrected P values were determined using permutation testing (FSL’s randomise; 5000 iterations), and results were thresholded at corrected P < 0.01. For one-sample t-tests, each iteration randomly sign-flipped individual subjects’ coefficient maps. Clusters were defined as having faces or edges or corners that touched and a minimum of 20 voxels. Peak activations within each cluster of activation were identified using AFNI software package (http://afni.nimh.nih.gov/afni). If a specific region of activity had multiple peaks within a cluster, the peak with the highest t-value was reported. Brain regions were identified using MRICron software and brain atlases (Damasio 2005; Petrides 2005).

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