Episodic memory formation in naturalistic viewing

Andrey R. Nikolaev^{1,2*}, Inês Bramão¹, Roger Johansson¹, Mikael Johansson¹

¹Lund Memory Lab, Department of Psychology, Lund University, Lund, Sweden ²Brain & Cognition Research Unit, KU Leuven, Leuven, Belgium

* Corresponding author Department of Psychology, Lund University, Box 213 22100 Lund Sweden andrey.nikolaev@psy.lu.se

Abstract

The brain systems of episodic memory and oculomotor control are tightly linked, suggesting a crucial role of eye movements in memory. But little is known about the neural mechanisms of memory formation across eye movements in naturalistic viewing behavior. Here, we leverage simultaneous recording and analysis of eye movements and EEG to examine the formation of episodic memory in free viewing. Participants memorized multi-element events while their EEG and eye movements were concurrently recorded. Each event comprised elements from three categories (face, object, place), with two exemplars from each category, in different locations on the screen. A subsequent associative memory test assessed participants' memory for the between-category associations that specified each event. We overcame the problem of overlapping EEG responses to sequential saccades in free viewing using a deconvolution approach. Brain activity was segmented relative to the fixation onsets and we examined EEG power in the theta and alpha frequency bands, the putative oscillatory correlates of episodic encoding mechanisms. High subsequent memory performance was predicted by three modulations of fixation-related EEG during the event memory formation: 1) theta synchronization at fixations after between-category gaze transitions, 2) theta synchronization at fixations after within-element gaze transitions, 3) alpha desynchronization at fixations after between-exemplar gaze transitions. Thus, event encoding with unrestricted viewing behavior was characterized by three neural mechanisms, manifested in fixation-locked theta and alpha EEG activity that rapidly turned on and off during the unfolding eye movement sequences. These three distinct neural mechanisms may be the essential building blocks that subserve the buildup of coherent episodic memories during naturalistic viewing behavior.

Introduction

Life events consist of visual, semantic, contextual, and other features associated and stored as coherent episodic memory representations. Episodic memory allows us to mentally travel in time to relive past events with considerable detail and specificity (Tulving, 1983). A critical aspect when building such memory representations is that the visual field of our perception is spatially limited. At any given time, we can apprehend information in full acuity only in a small spot, determined by the size of the fovea. We overcome this limitation by constantly shifting our visual focus with eye movements. In effect, these visual "samples" of the world constitute the pieces that we successively bind together into meaningful, coherent episodic memory, memory formation has primarily been studied in experimental paradigms where the study material is presented in a single location on the screen and where eye movements typically are treated as artifacts. Thus, little is currently known about the neural mechanisms of episodic memory formation across eye movements. In the present study, we set out to capture the neural signatures of building coherent episodic memories via eye movements at the level of individual gaze fixations. To this end, we apply a state-of-the-art approach of simultaneous eye-tracking and EEG recording and analysis in a free-viewing memory task.

Forming new episodic memories critically hinges upon binding together the different elements that constitutes an event as a whole. Sequential saccadic eye movements organize these relationships, acting themselves as a mechanism that links different aspects of the environment into coherent memory representations (for overviews, see Ryan et al., 2020; Voss et al., 2017). Eye movements also serve an important role during memory retrieval (Damiano and Walther, 2019; Johansson and Johansson, 2014; Johansson et al., 2022; Wynn et al., 2020), and a longstanding theory holds that the stored representations include the sequences of eye movements established during memory formation, which supposedly carry information about linked event elements (Brandt and Stark, 1997; Noton and Stark, 1971).

The clock frequency of this 'linking' is the theta rhythm (4-7 Hz). We typically make about 3-4 saccades in natural viewing per second. Such visual sampling frequency corresponds to the theta rhythm (Amit et al., 2017; Otero-Millan et al., 2008). The saccadic theta rhythm is part of a periodic mechanism of attentional deployment (Hogendoorn, 2016). It also corresponds to theta oscillations of the hippocampus, which is the main dispatcher of associative memory formation (Cohen and Eichenbaum, 1993). The hippocampus also has tight anatomical connections with brain structures that control oculomotor behavior (Pierrot-Deseilligny et al., 2004; Shen et al., 2016). Recent data suggest that hippocampal theta oscillations support saccade guidance during memory formation on a fixation-by-fixation basis (Jutras et al., 2013; Kragel et al., 2021). In this way, the theta cycle may organize sequential visual inputs and enables the binding of event elements into a coherent memory representation (Herweg et al., 2020).

The relevance of theta oscillations for memory processes has long been established in studies using stimulus-response paradigms with fixed stimulus presentation. For instance, increased frontal theta power during episodic memory encoding predicts subsequent retrieval performance (Hsieh and Ranganath, 2014; Klimesch, 1996). The increase in theta power, i.e., theta synchronization, is often accompanied by a decrease in alpha (8-13 Hz) power, i.e., alpha desynchronization (HansImayr et al., 2016; Waldhauser et al., 2012). Alpha desynchronization may represent the fidelity of memory representations for the constituent elements of episodes (HansImayr et al., 2016). Thus, alpha and theta oscillations serve essential but distinct functions of episodic memory formation: representing elements (alpha) and binding them together into a coherent whole (theta). It is currently unknown how these mechanisms engage during the sequential sampling of event elements across saccades to allow later episodic remembering of the event as a whole. In the current study, we track how these

neural mechanisms subserve the buildup of episodic memories during unrestricted viewing behavior.

To study neural activity across eye movements, it is necessary to record and analyze brain activity and eye movements simultaneously. Over the last decade, the combination of EEG and eye-tracking has become a powerful tool for studying perceptual and cognitive processes in free-viewing behavior (Coco et al., 2020; Devillez et al., 2015; Dimigen et al., 2011; Fudali-Czyz et al., 2018; Körner et al., 2014; Tyson-Carr et al., 2020), including attention and memory across saccades. For instance, presaccadic EEG activity has been found to predict successful encoding to working memory of taskrelevant scene elements in a change blindness task (Nikolaev et al., 2011). However, despite the apparent benefit of utilizing EEG-eye movement coregistration, attempts to apply it are virtually absent in the episodic memory literature. One explanation may be the vast methodological challenges that come with these techniques, including the effects of overlapping saccades on EEG activity in free viewing (Dimigen and Ehinger, 2021; Dimigen et al., 2011; Nikolaev et al., 2016). Moreover, low-level oculomotor effects of eye movement characteristics can systematically differ between experimental conditions and distort the EEG, creating differences that are likely to be confounded with effects of the experimental conditions. To overcome these and other challenges as described below, the present study applied a regression-based deconvolution approach that recovers unknown isolated neural responses given the measured EEG and the latencies of experimental events (Dimigen and Ehinger, 2021; Ehinger and Dimigen, 2019). This approach ultimately allowed us to isolate and track the neural correlates of episodic memory formation across eye movements.

The present study introduces an associative memory task in which participants encoded multiple events, each comprising three pairs of images from three different categories (faces, places, objects) in different locations on the screen (Fig. 1). Two exemplars from the same category appeared close together on the screen, forming a pair, while the distances between pairs, i.e., between categories, were much greater. This allowed us to distinguish gaze transitions between different categories (between-category saccades) and between two exemplars from the same category (betweenexemplar saccades). In a subsequent associative memory task, we tested episodic memory for between-category associations (i.e., associations between exemplars from different categories) within each event, the formation of which supposedly depends on the corresponding betweencategory saccades during encoding. The within-category associations (i.e., associations between two exemplars from the same category) were not tested. Thus, between-category and betweenexemplar saccades correspond to associations between event elements that differ in their taskrelevance. Additionally, we examined scrutinizing saccades within an exemplar (within-element), as it is well-known that the cumulative number of fixations on a visual stimulus predicts subsequent memory of that stimulus (Liu et al., 2017; Loftus, 1972; Olsen et al., 2016). This allowed us to examine the neural correlates of two types of memory mechanisms: the first related to eye movements that bind the elements into a coherent event, the second related to eye movements that sample visual information from individual elements.

Moreover, for between-category and between-exemplar saccades, we distinguished between saccades associated with the first "visit" to a category/exemplar and "revisits" to them. Revisits, i.e., refixations of previously visited locations, have proven to be of particular interest in memory-related gaze behavior. Specifically, refixations can increase the sampling of information necessary for successful memory formation (Mirza et al., 2016; Turk-Browne, 2019; Voss et al., 2011) and recover information lost or missed during scanning (Gilchrist and Harvey, 2000; Körner and Gilchrist, 2008; Meghanathan et al., 2019; Tatler et al., 2005). Our previous coregistration studies have shown that refixations differ in the allocation of attention during saccade planning from ordinary fixations to locations visited once (Nikolaev et al., 2018) and that refixation planning is goal-dependent (Meghanathan et al., 2020). Moreover, we found that information acquisition at fixations to

locations that are revisited later is distinct from that at refixations and ordinary fixations (Nikolaev et al., 2022). Critically, a recent study combining eye-tracking with intracranial EEG found that presaccadic hippocampal theta activity predicts whether an upcoming saccade will be a first visit or a revisit (Kragel et al., 2021). Based on these previous findings, we expected EEG activity during between-category revisits to be predictive of the successful binding of the task-relevant event elements.

Methods

Participants

Thirty-six healthy adults took part in the experiment in exchange for a gift card in a shopping mall (approx. \leq 10). Three participants were excluded because of system crashes during the recording and another three due to below-chance memory performance (53.2%). Finally, two participants were excluded due to ceiling performance (> 90%), i.e., too few incorrect trails for the planned analysis. The final group included 28 participants (20 females; mean age 23.3; age range 18-32).

The study was conducted in accordance with the Swedish Act concerning the Ethical Review of Research involving humans (2003:460). All participants gave written informed consent, and the study followed the local ethical guidelines at Lund University. Moreover, the study followed the Lund University safety protocol for conducting EEG, eye-tracking and behavioral experiments in the context of the COVID-19 pandemic.

Stimuli

A total of 324 photo images, from three different categories: faces, places, and objects, were used. Faces were selected from the Oslo Face Database (Chelnokova et al., 2014), which provided consent for the publication of the faces that were used in the figures. Objects were selected from the Bank of Standardized Stimuli (Brodeur et al., 2010). Places were selected from the Michael Tarr lab scene database (http://www.tarrlab.org/). The images were converted to black-and-white, and each image's pixel intensity was normalized to the average pixel intensity of all images. The images were combined into 54 events. Each event comprised three image pairs (two exemplars from each category) distributed across distinct locations on the display (Fig. 1). Exemplars from the same category were close to each other, whereas the different categories were more separated. Specifically, each image had a square shape with a side of 4.2 degrees of visual angle (dva). The width of a pair (two exemplars from one category) was 8.5 dva, and the distance between the centers of the three pairs was 17.9 dva. The three pairs were placed on an invisible circle with a radius of 9.2 dva centered at the center of the screen. The location of the pairs on the circle was random for each event, but the pairs were always separated by arcs of 120 degrees. The distribution of images and their pairing across events was random for each participant. The images were presented on a grey background.

Procedure

Participants sat in a dimly lit room, with their heads stabilized with a chin rest. Stimuli were presented at a viewing distance of 62 cm on an EIZO FlexScan EV2451 monitor with a resolution of 1920 x 1080 pixels and a refresh rate 60 Hz, which was part of the Tobii Pro Spectrum eye tracker.

The experiment comprised six blocks with the possibility to take a short break between each block. Each block included two main stages: memory encoding and memory retrieval with a distractor task in-between (Fig. 1). Participants self-initiated each block by pressing the space bar. During encoding, nine events were presented for 10 s each. Before each event, a fixation cross was presented for a random interval between 1 and 1.5 s. Participants were instructed to memorize the faces, places and objects that appeared together on the same event. The encoding stage was followed by a distractor task, where participants were asked to count backwards in steps of 7 from a random 3-digit number during 30 s.

Three seconds after the distractor task, a cued recall task followed, which assessed episodic memory by testing retrieval of the between-category associations of each event. A test screen with three images, one at the top and two at the bottom, was presented for 2 s (Fig. 1). Each test screen was preceded by the presentation of a fixation cross randomly jittered between 1 and 1.5 s. The image at the top served as the retrieval cue, and participants were asked to indicate which of the two images at the bottom that had appeared together with the cue during encoding, i.e., to indicate the target. The distractor was an exemplar from the same category as the target but selected from a different event studied no more than two events before or after the target event. Participants responded by pressing the left or right arrow keys of a standard keyboard. After response, confidence was measured. Participants were asked to indicate how certain they were in their response. The question "Are you sure?" was displayed in the screen and participants had three response options: "Sure", "Maybe", "Guess", which were given by pressing the left, down and right arrow keys, respectively.

Between-category associations for each image in each event were tested on four trials. For example, a face image was tested twice with place images (once as cue and once as target), and twice with object images (once as cue and once as target). The cue-target status of categories was counterbalanced across blocks, such that in the odd blocks the cue-target status was faces-places, places-objects, objects-faces, and in the even blocks the corresponding cue-target status was places-faces, objects-places, faces-objects. Thus, memory for each event was tested for 12 different associations. In total, 108 associations were tested in each block (12 x 9 = 108 tests for the nine events within a block) with randomized order of presentation.

Prior to the main experiment, two practice sessions were conducted in which participants received feedback on their performance after each test trial. The practice session ensured that participants were familiar with the task and had understood the instructions. The overall duration of an experiment was about two hours.

The experiment was programmed in PsychoPy v.2020.2.4 (Peirce et al., 2019).



Fig. 1. At encoding, nine events were sequentially presented. Each event comprised two exemplar images from three different categories (faces, places, objects) and was presented for 10 s. After a 30-s distractor task, episodic memory was assessed by testing all between-category associations from each event. The test screen (presented for 2 s) included a cue image at the top with two images at the bottom from a different category. Participants were asked to indicate which of the two images that had appeared together with the cue in the previously encountered events. In this example the correct response is the lock to the right, as it appeared together with the face cue in Event 1, whereas the binoculars to the left appeared in Event 2. Note that on the actual encoding screen the angular distances between categories were much larger.

Simultaneous EEG and eye movement recording

EEG and eye movements were recorded concurrently throughout the experiment. EEG data were recorded using a Grael 4K EEG amplifier (Neuroscan, Compumedics Limited, Australia) at a sampling frequency of 2048 Hz using 31 Ag/AgCl electrodes positioned according to the extended 10-20 International system (Easycap GmbH, Germany). Two electrodes were mounted over the left and right mastoids, using the latter as recording reference. The ground electrode was AFz. Additional electrodes recorded the vertical and horizontal electrooculograms (VEOG and HEOG): the VEOG electrodes were placed above and below the right eye, the HEOG electrodes were placed at the left and right outer canthi of two eyes. Electrode impedances were kept below 5 k Ω .

The Tobii Pro Spectrum eye tracker (Tobii, Stockholm, Sweden) recorded movements of both eyes with the sampling frequency of 600 Hz. 9-point calibration and validation routines were conducted before the first block. An error of 1 dva between calibration and validation was tolerated: if it was larger, the calibration was repeated. The eye tracking was controlled from the PsychoPy experimental program via the open-source toolbox Titta (Niehorster et al., 2020). The experimental program ran on the same computer that controlled the eye tracker. To synchronize stimulus presentation and EEG recording, a transistor-transistor logic (TTL) signal was sent via a parallel port from the stimulus presentation computer to the EEG system at the beginning and end of each block, encoding event, and memory test.

Data analysis

Only data from the encoding stage were analyzed. Analysis was conducted with custom scripts for R (Team, 2020) as well as for MATLAB R2020a (The MathWorks Inc., Natick, Massachusetts), using the EEGLAB toolbox (Delorme and Makeig, 2004), the EYE-EEG extension (Dimigen et al., 2011) for EEGLAB, and the Unfold toolbox (Ehinger and Dimigen, 2019).

Behavioral analysis

Memory performance for each event was assessed by retrieval accuracy weighted by confidence, such that correct responses were scored as 3 for "Sure" responses, as 2 for "Maybe" responses, and as 1 for "Guess" responses. Each image from each event occurred either as a cue or a target in four memory tests trials. Thus, each image can in total receive a score ranging from 0 (incorrect responses in all four tests) to 12 (correct responses accompanied by four "sure" responses in all four tests). To calculate a memory score for the whole event, we then summed the scores for the six elements of each event (range from 0 to 72). For the EEG analysis, events were divided into the two categories "high memory" and "low memory" according to the subsequent memory scores. For each participant, high and low memory was calculated by a median split of that participant's memory scores for the whole events.

Eye movement analysis

Fixations and saccades were detected from participants' right eye using the velocity-based algorithm for saccade detection (Engbert and Mergenthaler, 2006) implemented in the EYE-EEG extension. Fixations were considered to be on an image if they were located within 0.5 dva of the outer contour of the image. Since participants cannot adequately perform a memory test if they did not visually process (fixate) all images while event encoding, we excluded from all analyses the events where fixations were detected only on two of the three categories (on average 3±4 (mean ± SD) events per participant). Such fixations and fixations outside the images on different parts of the display were assigned to an 'other' saccade type, which were used as a reference in the EEG analysis (see the Deconvolution section below).

Saccades were categorized according to whether they occurred within an exemplar (within-element saccades), between the two exemplars of a category (between-exemplar saccades), and between categories (between-category saccades). In addition, we distinguished between the first visit to an image and revisits to it. Altogether, we defined five different types of saccades (Fig. 3A):

- 1. Between-category saccades: first visit
- 2. Between-category saccades: revisits
- 3. Between-exemplar saccades: first visit
- 4. Between-exemplar saccades: revisits
- 5. Within-element saccades

The fixation-related EEG analysis examined EEG activity in the fixation intervals that succeeded these five saccade types.

EEG preprocessing

The goal of our study required fixation-related analysis of EEG in short epochs time-locked to fixation onsets in the encoding interval. This analysis used deconvolution modelling to correct for the overlapping effects of saccades on EEG. Deconvolution modelling involves time regression, which requires continuous EEG, therefore, preprocessing was performed on continuous EEG. In addition, we analyzed EEG time-locked to the event onset during the entire 10-s encoding interval. The preprocessing for this analysis was the same as for the fixation-related analysis up to the segmentation step. We used the following preprocessing and deconvolution pipelines, which we developed earlier (Nikolaev et al., 2022).

We down-sampled EEG signals to 256 Hz. The EEG and eye-tracking data were synchronized using the function *pop_importeyetracker* from the EYE-EEG extension. Saccades, fixations, event onsets, and bad eye-tracking intervals were inserted into the EEGLAB data structure.

The preprocessing pipeline included a series of cleaning functions from EEGLAB. The *pop_cleanline* function removed power line noise using multi-tapering and a Thompson F-statistic. The *clean_artifacts* function removed flat-line channels, low-frequency drifts, noisy channels, short-time bursts. This function is based on artifact subspace reconstruction (ASR), which compares the structure of the artifactual EEG to that of known artifact-free reference data (Kothe and Jung, 2016). The tradeoff between artifactual and retaining brain activities depends on the ASR parameter, which we set to 20 according to the recommendations by (Chang et al., 2020).

Ocular artifacts were removed with the *OPTICAT* function (Dimigen, 2020). First, EEG was high-pass filtered at 2 Hz to suppress large deviations from baseline due to summation of the corneo-retinal artifacts during sequential eye movements. Next, 30-ms segments around saccade onsets were obtained and re-appended to EEG to 'overweight' the contribution of the saccadic spike activity in the EEG input to independent component analysis (ICA). Then, the ICA weights obtained after ICA training on these filtered data were transferred to the unfiltered version of the same data. Finally, the ratio between the mean variance of independent components during saccade and fixation intervals was calculated (Plöchl et al., 2012). If this ratio was higher than 1.1, the corresponding independent components were considered as saccade-related and removed.

The automatic classifier (*pop_iclabel*) revealed independent components related to remaining artifacts; these components were removed. EEG was re-referenced to average reference. The removed channels (mean = 1.1, SD = 1 per participant) were interpolated with spherical spline interpolation.

Theta and alpha power were extracted by applying the procedure from (Ossandón et al., 2020). We filtered the continuous EEG with a low cut-off of 4 Hz and high cut-off of 7 for the theta band and with a low cut-off of 8 Hz and high cut-off of 13 for the alpha band using the *pop_eegfiltnew* function with default settings. We Hilbert-transformed the filtered signal and obtained the instantaneous power. We normalized the EEG power at each sampling point to its ratio with the respective channel mean power in a baseline interval. In the analysis of the encoding interval the baseline interval was set from -200 to 0 ms before the onset of an encoding event. In the fixation-related analysis, to avoid an interval of saccade execution just before fixation onset, the baseline interval was set from -200 to -100 ms before the fixation onset. The power values were log-transformed.

Deconvolution

To overcome the problems associated with EEG-eye movement coregistration in free viewing behavior described in Introduction, we used deconvolution modelling implemented in the Unfold toolbox (Ehinger and Dimigen, 2019). The toolbox performs regression-based EEG analysis that includes massive univariate modeling, linear deconvolution modeling, and generalized additive modeling. The outputs of the toolbox are the partial effects (i.e., beta coefficients) for the predictors of interest, adjusted for the covariates included in the model.

The toolbox allowed us to correct overlapping EEG responses to sequential saccades in free viewing, as well as to account for several possible detrimental covariates. These covariates included the low-level oculomotor effects of eye movement characteristics and the ordering factors in time courses of an encoding trial, a block, and an entire experiment. Specifically, we considered the low-level

oculomotor characteristics of fixation duration and saccade size and direction, which are known to affect the fixation-related EEG (Dimigen et al., 2011; Nikolaev et al., 2016). Furthermore, we considered three ordering factors. Since time course of a trial with multiple eye movements may affect the fixation-related EEG activity (Fischer et al., 2013; Kamienkowski et al., 2018), we considered the rank of a fixation in relation to all other fixations that occurred during in the whole encoding interval of an event. The rank was assigned to each fixation as an ordinal number of this fixation in the encoding interval. Behavioral results (see below) revealed a nonlinear effect of event order within a block and a strong linear effect of block order on subsequent memory. We accounted for these factors by including them in the deconvolution model.

The Unfold toolbox involved the following procedures, which were sequentially applied to continuous EEG of each participant.

To account for the experimental conditions of interests and possible confounding factors in the same model we used the following Wilkinson notation of the model formula:

```
Fixation: y ~ 1 + cat(MemoryPerformance) + rank + iBlock + spl(iEventPerBlock,4) +
spl(duration,5) + spl(sac_amplitude,5) + circspl(sac_angle,5,-180,180)
Stimulus: y ~ 1
Levels of MemoryPerformance: `other' (reference level), `low memory', `high memory'
```

This formula includes the multiple effects at fixations, as well as the onset of the event screen. Specifically, for fixations, the formula considers the fixation onset ($y \sim 1$, i.e., the intercept), a categorical predictor of the subsequent memory effect ('MemoryPerformance') and continuous predictors: fixation rank ('rank'), a block order in an experiment ('iBlock'), an event order in a block ('iEventPerBlock'), fixation duration ('duration'), size ('sac amplitude') and direction ('sac angle') of the preceding saccade. In this example formula, the 'MemoryPerformance' predictor includes three levels: 'other' fixations (see Eye movement analysis above), 'low memory' and 'high memory' (see Behavioral analysis above). We modified the formula by defining the categorical predictors depending on the analysis goals, as specified below in each section of Results. In all analyses we used treatment coding with 'other' fixations as the reference level. Since fixation duration, and saccade size and direction have nonlinear effects on EEG (Dimigen and Ehinger, 2021; Nikolaev et al., 2016), we modeled them with a basis set of five spline predictors (splines were circular for saccade direction). Since we found that the event order in the block has a nonlinear effect on the memory score (Fig. 2B), and we assumed its nonlinear effect on the EEG, we modeled it also with spline predictors. Spline knots were placed on the participant-specific percentiles of the covariates. For events, we considered only their onsets (y \sim 1), i.e., the intercept that described the overall shape of EEG evoked by the event screens.

To recover isolated EEG responses (betas) to each fixation and to each event screen, which best explained the continuous EEG, we created a design matrix and time-expanded it in a time window between -200 and +400 ms around fixation and event onsets. Time expansion involved duplicating and shifting the predictors of the mass univariate linear regression design matrix for each time lag. The time-expanded design matrix spanned the duration of the entire EEG recording. As a result, instead of estimating multiple linear models for all time points, we estimated one linear model, simultaneously estimating all fixation and event betas. The variable temporal distance between sequential fixations allowed separation of the overlapping effects.

Before fitting the model, we excluded irrelevant and artifactual intervals from modelling by setting the entire rows of the time-expanded design matrix to zero. Specifically, we removed intervals of the distractor task, memory tests, intervals between events, breaks between blocks, and bad eye-tracking intervals, leaving only intervals of free viewing exploration of event screens during

encoding. We fitted the deconvolution model to each of the 31 electrodes using the iterative Least Squares Minimal Residual algorithm for sparse design matrices (Fong and Saunders, 2011).

To extract EEG power for the statistical analysis, we reconstructed the regression-based power waveforms from the beta coefficients of the model for various categorical conditions (specified in each section of Results) in epochs from –200 to +400 ms after the fixation onset. This reconstruction included adjusting the covariate effects of the eye movement characteristics and ordering factors (specified above) by adding the marginal effects of their means. Specifically, for each participant we calculated the average value for each covariate and evaluated the effect-power at this value. This value was then added to all other predictors, effectively adjusting all regression-based power waveforms to the same covariate values. The resulting regression-based power waveforms were equivalents to participant-level averages in a traditional EEG analysis.

Statistical analysis

We estimated theta and alpha oscillations at the first peak of their cycle about 100 ms after the fixation onset. In the statistical analysis, we used the average power between 60 and 140 ms after the fixation onset, that is, 40 ms before and after the peak. To capture effects with potentially different topographical distributions we divided all electrodes (except two temporal electrodes T7 and T8, which were typically noisy) into three broad groups:

- 1. 12 frontal electrodes: FP1,FP2,F7,F8,F3,F2,F4,FC1,FC2,FC5,FC6
- 2. 12 centro-parietal electrodes: C3,CZ,C4,CP5,CP1,CP2,CP6,P7,P3,PZ,P4,P8
- 3. 5 occipital electrodes: PO9,PO10,O1,IZ,O2

In each group, the electrodes were symmetrically distributed over the scalp relatively to the sagittal axis. Since regional averaging of electrodes provides a more reliable estimate of activity in a region than a single measurement (Dien and Santuzzi, 2005), we averaged power waveforms across the electrodes of each group. The statistical analysis was done for these areas separately.

Unless indicated otherwise, for the statistical analysis we used a repeated measures ANOVA. To handle violation of sphericity we applied the Huynh-Feldt correction for p-values associated with two or more degrees of freedom. Fisher's LSD (least significant difference) test was used for posthoc analyses. The statistical analyses were performed with STATISTICA 10 (StatSoft Inc., Tulsa, OK, USA) and R (R Core Team, 2020).

Results

Behavioral results

Average memory performance across 28 participants was $68.6 \pm 8.33 \%$ (mean \pm SD). The memory score weighted by confidence for the low memory condition was 27.8 +/- 6.8 and for the high memory condition 48.8 +/- 9 (mean +/- SD). Memory performance changed in the time course of a block and experiment (Fig. 2). We evaluated these changes with a repeated measures ANOVA on the memory scores weighted by confidence with the factors of Event (nine levels) and Block (six levels). There were significant main effects of Event (F(8, 216) = 3.7, p = .004, $\varepsilon = .62$) and Block (F(5, 135) = 9.8, p < .001, $\varepsilon = .95$)), but no interaction between them. To characterize the effects of Event and Block, we used planned comparisons with the linear and quadratic contrasts. The contrast analysis revealed significant effects for the linear contrast for Block (F(1, 27) = 31.2, p < .001) and for the quadratic contrast for Event (F(1, 27) = 16.8, p < .001), but not for the linear contrast for Event (F(1, 27) = 0.7, p = .4). These results indicate a prominent learning effect during the experiment, as well as possible primacy and recency effects in time course of a block.



Fig. 2. Memory performance weighted by confidence in the time course of an experiment (A) and a block (B). The thin lines indicate significant linear (A) and quadratic (B) fits.

Eye movement results

The dynamics of eye movement behavior during the 10-s encoding interval is illustrated in Fig. 3B by the fixation rank probability density for each type of preceding saccade. At first, participants tended to visit all categories quickly with between-category saccades. Then, between-exemplar saccades became predominant. The number of within-element saccades was relatively stable throughout the whole encoding interval. Towards the latter part of the interval, between-category revisits became more frequent.

To determine if, and to what degree, the saccade categories predicted subsequent memory performance, we analyzed the relationship between the cumulative number of saccades (in each category) within an event and subsequent memory performance for that event. The between-category saccades for first visits were not analyzed since there were always 3 per event. We computed the number of remaining four saccade types for each event and pooled these numbers with the corresponding event memory scores over 28 participants. We applied a one-way ANOVA on the event memory score with the number of the saccades as a categorical predictor (bins with less than 20 values at the left and right ends of the distributions were excluded). The results showed that memory for the event significantly increased with the cumulative number of between-category revisits (F(8, 1392) = 3.8, p < .001) and with the cumulative number of within-element saccades (F(14, 1360) = 3.1, p < .001) (Fig. 3C).

To be successful in the retrieval task, association from all three categories need to be considered during event encoding. This can be achieved by two different gaze patterns: (1) engaging in active gaze transitions between each of the three categories, or (2) engaging in active gaze transitions between only two of the categories, i.e., when the association between the first visited category and the last visited category is inferred without an active gaze transition between the categories. In the present data set, gaze transitions involving only two categories characterized approximately one-

third of the events. To evaluate whether the two eye movement patterns are associated with different subsequent memory performance, we compared event memory scores between these pattern types with a paired t-test. We found no difference in memory between the two patterns (t(27) = 0.05, p = .96), suggesting that event memory does not require gaze transitions between all three categories.



Fig. 3. A: an example scanpath illustrating fives types of saccades. B: Probability density estimation of the fixation rank in a 10-s event interval pooled across 28 participants. The vertical lines indicate mean ranks of the corresponding fixation type. C: Dependence between the occurrence frequency of saccade types and the memory score. Significance of the dependence obtained with one-way ANOVA is indicated at each panel. Error bars indicate standard errors of means across 28 participants.

In sum, the eye movement results show that between-category saccades (revisits) and withinelement saccades are predictive of subsequent memory performance, whereas between-exemplar saccades are not. These dependencies were determined by our memory task, which required strong associations between the categories whereas associations between the exemplars of a category were irrelevant to the task. Within-element saccades are relevant for acquiring sufficient visual information about each image, a prerequisite for establishing a strong representation of each element and for recognizing them during the retrieval task.

EEG results at the event level

To assess how theta and alpha power at encoding predicts subsequent memory at the time scale of the entire encoding interval we segmented the (not deconvolved) EEG power from -0.2 to 10 s relatively to the event onset and baseline corrected it to the interval from – 0.2 to 0 s before the event onset. We smoothed the individual power waveforms with a Savitzky-Golay filter (R package 'signal'). Events were split between low and high memory performance (the number of low memory events was 28.5 (1.35); the number of high memory events was 25.1 (1.74) (mean (SD) per participant). To investigate the modulation across a 10-s encoding interval, we created ten time-bins of 1 s each and averaged the power values in these bins. We applied a repeated-measures ANOVA on the power values with the factors of Memory (low vs. high) and Time bins (10 levels). For theta power, we found interactions between Memory and Time bins for the frontal (F(9, 243) = 2.4, p =.02, $\varepsilon = .95$) and centro-parietal (F(9, 243) = 2.1, p = .04, $\varepsilon = .92$) areas (Fig. 4A). Post-hoc tests revealed that these interactions occurred due to higher power for the high than low memory events starting for the frontal areas from the 7th time bin (all p < .05) and for the centro-parietal areas from 8^{th} time bin (all p < .03) until the end of the encoding interval. For alpha power, there was a clear trend towards lower power in high-memory versus low-memory events, most noticeably over the occipital areas (Fig. 4B), which, however, did not reach significance.



Fig. 4. Theta (A) and alpha (B) power time-locked to the event onset during a 10-s encoding interval for low and high subsequent memory performance. The time course of the grand-averaged power waveforms and the mean-error plots (mean +/- SEM) for ten 1-s time bins are shown. In (A) the p-levels of the interactions between Memory and Time bins and significance of the post-hoc tests (with the asterisks) are indicated.

In sum, EEG analysis of the entire encoding interval revealed the expected neural signatures, where higher subsequent memory performance was associated with increased theta synchronization and alpha desynchronization during encoding (Hanslmayr et al., 2016). Consistent with the idea of a gradual buildup of an episodic memory representation, theta synchronization was more prominent in the second half of the encoding interval over the frontal and centro-parietal areas. Theta synchronization was accompanied by a tendency for alpha desynchronization, which was most

prominent over the occipital areas (Fig. 4B), although there was no significant difference between high and low memory.

EEG results at the fixation level

Since the deconvolution framework is a novel approach to free-viewing behavior in tasks of episodic memory formation, we first evaluated how it changes the power waveforms (see Supporting Information for details).

The general memory effects of theta and alpha activity

Before analyzing the data for the different saccade categories, we wanted to assess if the EEG findings at the event level could be reproduced at the fixation level. Thus, we tested the subsequent memory effect on fixation-related theta and alpha power by contrasting the fixation-related EEG of high memory events with low memory events over all saccades within an event. Since the event analysis showed a significant memory effect only in the second part of the event (Fig. 4), we divided the fixations into two bins according to fixation rank, which roughly correspond to the first and second half of the 10-s interval and restricted our analysis to the second bin (the second half of the event). To accurately account for the overlapping effects of eye movement sequences on EEG, the deconvolution formula should include fixations of the entire encoding interval. Therefore, in the formula for deconvolution modelling we used five levels of categorical predictors as follows: Other, First half/Low memory, First half/High memory, Second half/Low memory, Second half/High memory. In the second half, the number of the fixation-related epochs per participant was 258 (65.2) in the low memory condition and 222 (58.6) in the high memory condition (mean (SD)).

In a statistical analysis of power averaged in the window from 60 to 140 ms after the fixation onset, we found higher theta power over the frontal areas for encoding leading to high than low memory performance (t(27) = 2.4, p = .03) (Fig. 5A). No significant effects for other areas were found. Additionally, we observed lower alpha power for encoding leading to high than low memory performance over the frontal areas (t(27) = 3.3, p= .003) and the occipital areas (t(27) = 3.0, p = .006) (Fig. 5B).



Fig. 5. Theta (A) and alpha (B) power over the frontal areas for fixations in the second half of the encoding interval. The grand-averaged (N=28) power in low and high memory conditions. The mean-error (mean +/-SEM) plot of the power in the interval from 60 to 140 ms after the fixation onset. The difference power map: high minus low memory.

To summarize, the fixation-related EEG results corroborate and extend what was found at the event level, i.e., theta synchronization and alpha desynchronization during encoding covary with subsequent memory performance. Next, we analyzed EEG power at fixations succeeding the five saccade types.

The EEG results across saccade types

Here, we examined EEG power in the fixation intervals succeeding the five saccade types (Fig. 3A). For this purpose, we specified ten levels of categorical predictors in the deconvolution model formula: the five preceding saccade types x memory performance (low, high). "Other" fixations were the reference level, as in the previous model. The number of the obtained fixation-related epochs is presented in Table 1.

Table 1. The number of fixation-related epochs succeeding each saccade type per participant (mean and SD across 28 participants)

	low memory	high memory
Between-category saccades: first visit	85 (9.8)	67 (8.8)
Between-category saccades: revisits	83 (36.8)	72 (31.3)
Between-exemplar saccades: first visit	67 (12.4)	52 (12.5)
Between-exemplar revisits: revisits	120 (55.2)	90 (40.5)
Within-element fixations	195 (73.2)	155 (58.5)

Between-category saccades

Here, we examined EEG power in the fixation intervals succeeding the between-category saccades that we predicted to support task-relevant associations between categories. We applied a repeated measures ANOVA with the 2x2 design: Gaze transition status (First visits vs. Revisit) x Memory performance (Low vs. High) on theta and alpha power (Fig. 6A). For theta power, we found a significant main effect of Memory performance over the *centro-parietal* areas: theta power was higher for high subsequent memory performance, F(1,27) = 5.6, p = .03 (Fig. 6B-D). In addition, we found a significant main effect of Gaze transition status over the frontal areas (F(1,27) = 5.7, p = .02): theta power was higher during the first visit. For alpha power, no significant effects were found.

Thus, theta power over the centro-parietal areas, after between-category saccades, predicts subsequent event memory performance. This theta memory effect is consistent with the eye-movement analyses, where subsequent memory performance increased with the cumulative number of between-category revisits during encoding (Fig. 3C).



Fig. 6. A: An example scanpath illustrating a subset of between-category (colored) saccades preceding the fixation intervals used in the EEG analysis. The grey arrows are saccades not included in this analysis. B: The difference power maps: high minus low memory. C: The grand-averaged (N=28) theta power over the centro-parietal areas for low and high memory for fixations after the between-category saccades. D: The mean-error (mean +/- SEM) plot of the power in the interval from 60 to 140 ms after the fixation onset.

Between-exemplar saccades

Next, we examined EEG power in the fixation intervals succeeding between-exemplar saccades, i.e., gaze transitions that may be considered irrelevant when building the task-relevant associations between categories (Fig. 7A). As in the previous analysis, we used a repeated measures ANOVA with a 2x2 design: Gaze transition status (First visit vs. Revisit) x Memory performance (Low vs. High). For theta power, no significant effects were found. But we found lower alpha power for high than low memory performance over the centro-parietal (F(1,27) = 5.0, p = .03) and occipital (F(1,27) = 6.4, p = .02) areas (Fig. 7B-D).

Thus, as expected, the theta power after task-irrelevant between-exemplar saccades did not predict subsequent event memory performance. This is consistent with the eye-movement result, where the cumulative number of between-exemplar saccades did not influence subsequent memory performance (Fig. 3C). The alpha desynchronization effect over posterior areas was unexpected, but may reflect other mechanisms, apart from binding per se, that are important for optimal memory formation.



Fig. 7. A: An example scanpath illustrating a subset of between-exemplar (colored) saccades preceding the fixation intervals used in the EEG analysis. The grey arrows are saccades not included in this analysis. B: The difference power maps: high minus low memory. C: The grand-averaged (N=28) theta power over the occipital areas for low and high memory for fixations after between-exemplar saccades. D: The mean-error (mean +/-SEM) plot of the power in the interval from 60 to 140 ms after the fixation onset.

Within-element saccades

The analysis of the fixation intervals after within-element saccades revealed higher theta power for high than low memory performance over the frontal areas (t(27) = 3.02, p = .006) (Fig. 8). No significant effects were found for other areas or for alpha power.



Fig. 8. A: An example scanpath illustrating a subset of within-element (colored) saccades preceding the fixation intervals used in the EEG analysis. The grey arrows are saccades not included in this analysis. B: The difference power map: high minus low memory levels. C: The grand-averaged (N=28) theta power over the frontal areas for low and high memory for fixations after the within-element saccades. D: The mean-error (mean +/- SEM) plot of the power in the interval from 60 to 140 ms after the fixation onset.

Thus, the theta power related to within-element saccades predicts subsequent memory performance similarly as for the between-category saccades. Moreover, for both saccade types, subsequent memory increased as a function of the cumulative number of saccades during encoding (Fig. 3C). Importantly, however, the topographies of the theta memory effects for the two saccade types were *distinct*: a centro-parietal maximum for between-category saccades (Fig. 6B) and a frontal maximum for within-element saccades (Fig. 8B), suggesting dissociable neural mechanisms.

To test the reliability of the topographical dissociation of the two theta effects we extracted the 'within-element effect' as the difference between theta power in high and low memory performance for fixations after within-element saccades. We extracted the 'between-category effect' as the difference between theta power in high and low memory performance for fixations after between-category saccades (averaged across first visits and revisits). We scaled the power values across the frontal, centro-parietal and occipital areas by dividing the value of each area by the square root of the sum of squares for all areas (McCarthy and Wood, 1985). A repeated-measures ANOVA on the scaled theta power with factors of Theta effect ('within-element' vs. 'between-category') and Areas (frontal, centro-parietal and occipital) revealed an interaction between these factors (F(2, 54) = 4.8, p = .02, $\varepsilon = .81$). The post-hoc test revealed higher power for the between-category effect over the centro-parietal area (p = .01) and a reversed tendency for the frontal area (p = .08) (Fig. 9). This finding indicates topographical differences for the two theta effects and suggests that they are associated with distinct neural mechanisms, operating in the same frequency band.

In sum, the correspondence of theta and eye movement results for between-category saccades and within-element saccades, which may be based on distinct mechanisms, indicates a diversity of relationships between theta oscillations, eye movements, and episodic memory formation, as will be discussed below.



Fig. 9. The scaled theta power for the 'within-element' and 'between-category' theta memory effects. The mean-error (mean \pm SEM) plot of the power in the interval from 60 to 140 ms after the fixation onset. The plevels indicate the results of post-hoc comparisons between low and high memory performance for each area.

Discussion

The present study set out to advance our current understanding of the relationship between eye movements and episodic memory encoding. In particular, we aimed at elucidating the neural mechanisms that mediate the formation of coherent episodic memories as a function of saccades to event elements during unrestricted viewing behavior. To this end, we applied a state-of-the-art analytical approach to EEG coregistered with eye movements during a free-viewing episodic memory task. Our approach provided a unique possibility to capture neural encoding mechanisms across eye movements at the level of gaze fixations while overcoming confounding effects of sequential saccades on brain activity. We identified neural signatures that are associated with the buildup of coherent episodic memories as a function of visual sampling behavior. The encoding of task-relevant associations between event elements was accompanied by simultaneous modulation of eye movements and theta oscillations that were predictive of subsequent memory. This modulation likely reflects the binding of event elements into a coherent representation across eye movements. In addition, a subsequent memory effect was also observed for theta oscillations related to small scrutinizing saccades within elements. The topographies of these theta effects were distinct, revealing the contributions of dissociable neural mechanisms subserving task performance. Although association between category exemplars was incidental to the task, alpha oscillations related to between-exemplar saccades also proved predictive of later episodic remembering. Overall, we have identified the engagement of three distinct neural mechanisms that may be the essential building blocks in formation of episodic memory during naturalistic viewing.

Theta and alpha oscillations supposedly serve important but distinct functions of episodic memory formation: binding individual elements together into a whole (theta) and representing individual elements (alpha) (Hanslmayr et al., 2016). But it has not yet been established how these

mechanisms are engaged when coherent episodic memories are formed in a piecemeal procedure across eye movements. In our study, we tracked how these neural mechanisms subserve the buildup of episodic memories during unrestricted viewing behavior.

Saccade types that predict subsequent memory

It is well known that eye movement behavior during free viewing is important for successful formation of episodic memory (for overviews, see Ryan et al., 2020; Wynn et al., 2019). We replicate and extend these findings by showing that the number of within-element saccades and between-category revisits positively correlated with performance in an associative memory task (Fig. 3C). Short within-element saccades scrutinize the visual features of single exemplars, increasing the visual sampling of different details, which leads to better subsequent memory of the stimuli (Liu et al., 2017; Loftus, 1972; Olsen et al., 2016). Additionally, as expected, the cumulative number of gaze transitions between categories during encoding increase memory performance. Likely, the between-category saccades contribute to the binding of separate elements into a coherent representation of the event, in which the hippocampus is involved (Herweg et al., 2020). Corroborating this idea, previous studies have shown that hippocampal activity and memory performance increase with the number of gaze fixations (Liu et al., 2017) and especially with the number of revisits (Kragel et al., 2021; Voss et al., 2011), as in our study. In contrast, the between-exemplar saccades were not relevant for successful task performance, which may explain why between-exemplar saccades were not predictive of memory performance.

Theta and alpha activity throughout the encoding interval predict subsequent memory

First, the EEG analysis confirmed the presence of theta and alpha oscillations predictive of episodic remembering, extending previous work where the to-be-remembered material was presented centrally without the need for visual exploration via eye movements (Hanslmayr and Staudigl, 2014; Klimesch, 1996). Our results revealed that, towards the end of the encoding interval (7-10 s) the theta power was higher for high than low subsequent event memory (Fig. 4A). The predominantly anterior topography of this theta effect suggests the contribution of frontal midline theta oscillations, which are related to intrinsic episodic memory processes (Hsieh and Ranganath, 2014).

Consistent with previous research (Griffiths et al., 2019; Hanslmayr and Staudigl, 2014; Klimesch et al., 1996), we also observed a tendency for subsequent memory to be associated with a decrease in alpha power. Fig. 4B shows that this tendency was mostly noticeable over the occipital areas (characteristic of alpha activity).

Overall, in good agreement with the expected role of theta and alpha oscillations in episodic memory formation, we observe opposite trends in theta and alpha activity predictive of subsequent memory. These theta synchronization and alpha desynchronization are likely complementing each other, reflecting the functioning of the hippocampal and cortical memory systems, respectively (Hanslmayr et al., 2016).

Fixation-related EEG predicts subsequent memory

The fixation-related EEG corroborated the findings reported at the event level. We observed statistically significant higher theta and lower alpha power for high than low memory performance (Fig. 5). The significance of both these effects at the fixation level, but not at the event level, suggest that segmentation of the EEG relative to fixations is potentially a more sensitive way to isolate the features characteristic of the subsequent memory effects.

The time window in the fixation-related EEG analysis was centered on the first power peak about 100 ms after the fixation onset. The latency of this peak coincides with the latency of lambda activity, which is associated with early perceptual processes at gaze fixation (Kazai and Yagi, 2003; Ossandón et al., 2010; Ries et al., 2016; Thickbroom et al., 1991). However, it seems unlikely that the observed memory effects are solely related to perceptual processes given that we observed clear modulations between these effects and subsequent memory performance. Additionally, the effects here described have a quite distinct topography from that of a typical lambda activity, over occipital regions (Ossandón et al., 2010; Thickbroom et al., 1991). Moreover, the peak latency coincides with the peak frontal theta power, which predicts memory performance in working memory tasks (Adam et al., 2018). As such, the time window used in our analysis captures the cycle of ongoing brain oscillations that carry out the rhythmic sampling of external information. Indeed, oscillations at the theta frequency drive perceptual sampling by organizing sequential inputs (Herweg et al., 2020). In free-viewing behavior, perceptual sampling is performed by saccadic eye movements generated at theta frequency (Amit et al., 2017; Otero-Millan et al., 2008). Eye movements are accompanied with phase resetting in a wide frequency range of oscillations observed in the visual cortex of monkeys (Ito et al., 2011; Rajkai et al., 2008), as well as in the scalp EEG in humans (Nikolaev et al., 2016). Since phase resetting of hippocampal theta oscillations supports successful memory formation (Jutras et al., 2013; Kota et al., 2020), rhythmic gaze sampling may synchronize brain areas and optimize the processing and encoding of visual information (Jutras and Buffalo, 2010; Voss et al., 2017). Importantly, theta oscillations represent several items within the same cycle, where the current item is strongly represented and previous items concurrently represented in a weaker form (Herweg et al., 2020). Transferred to our paradigm, theta activity may represent not only the fixated information but also previously visited exemplars (as well as previously presented events). Our time window thus may capture individual perceptual moments at fixations as well as the sum of active and latent representations accumulated over a series of eye movements.

EEG memory effects for saccades between event elements

Perhaps the most fascinating results were obtained when analyzing EEG in fixation intervals after different saccade types during the 10-s encoding interval. Next, we analyzed EEG power at fixations that followed between-category, between-exemplar, and within-element saccades. In our paradigm, between-category saccades supported task-relevant associations between event exemplars. The theta power was higher for high than low subsequent memory at fixations after between-category saccades (Fig. 6), and the cumulative number of such gaze transitions predicted subsequent memory (Fig. 3C). The alpha power was lower for high than low memory at fixations after between-exemplar saccades, i.e., task-irrelevant associations (Fig. 7). The cumulative number of such gaze transitions did not predict subsequent memory (Fig. 3C).

It is conceivable that gaze transitions between categories link the event elements together, creating a holistic representation of the entire event. These gaze transitions are thus of critical importance to episodic memory as they support the binding of information experienced separately in space and time (Cohen and Eichenbaum, 1993; Voss et al., 2017). Previous research has shown that this binding mechanism relies on the transfer of information between distant brain regions and is promoted by the synchronization of their activity at a theta rhythm (Clouter et al., 2017; Fell and Axmacher, 2011). Our results fit well with this previous literature. We observed high theta power after between-category saccades, predictive of high event memory, suggesting the involvement of theta in the binding of task-relevant elements into a coherent episodic representation. It may be the case that theta reflects an optimized oculomotor control that guides this binding mechanism. In fact, it has been shown that the theta rhythm also underlies viewing behavior optimal for memory formation (Herweg et al., 2020; Jutras and Buffalo, 2010; Voss et al., 2017). The key role of the hippocampal theta oscillations in memory-related gaze behavior has recently been demonstrated using intracranial EEG recording combined with eye tracking in free-viewing memory tasks (Kragel et al., 2021; Kragel et al., 2020). Kragel and colleagues (2021) described how revisits support the efficiency of visual sampling in the formation of memory for scenes. Hippocampal theta power decreased before revisits and increased after revisits, indicating rapid switches between memory-based saccade guidance and perceptual processing at fixation. Thus, the theta increase after revisits may reflect the mechanism by which contents attended to at multiple fixations become bound into a coherent representation. In the present study, the number of between-category *revisits* was predictive of event memory, whereas the EEG analysis contrasting between-category first visits vs. revisits failed to reveal any interaction with memory. This suggests that the between-category saccades for first visits and revisits are comparably important for the buildup of a coherent representation in our paradigm.

As expected, the cumulative number of between-exemplar saccades that were incidental to the task did not predict subsequent memory. Nevertheless, alpha power at fixations after these saccades was lower for high than low subsequent memory over the centro-parietal and occipital areas (Fig. 7). Although this alpha effect thus reflects a mechanism that supports successful memory formation, its exact nature remains unclear. For example, in episodic memory encoding, alpha activity indicates cortical information processing during the perception of an event (Jensen and Mazaheri, 2010; Klimesch, 2012), and its decrease predicts successful formation of episodic memory (Hanslmayr et al., 2016). Specifically, alpha desynchronization is thought to increase the capacity of local cell ensembles to process information by reducing local synchronization (Hanslmayr et al., 2016). Thus, the occipital predominance of memory-related alpha desynchronization in our visual memory task may reflect an increase in the capacity of cell ensembles in the visual system, which is beneficial for category-specific processing. Visual perception in our task involves discriminating between two similar exemplars from the same category and the between-exemplar saccades may facilitate a comparison of the two exemplars. Alpha activity has been shown to predict behavioral performance in visual discrimination tasks (Hanslmayr et al., 2005; Nelli et al., 2017). Thus, the here observed alpha desynchronization after between-exemplar saccades may indicate efficient discrimination of two similar exemplars that promote the successful encoding of the task-relevant associations.

Theta memory effects for within-element saccades

Another theta memory effect was observed at fixations after within-element saccades (Fig. 8), the number of which predicted subsequent memory (Fig. 3C). Remarkably, the theta memory effect for within-element saccades had a frontal maximum, whereas the theta effect for between-category saccades had a centro-parietal maximum (Fig. 9). The different topographies suggest distinct neural mechanisms underlying these theta effects.

Short within-element saccades, which serve to scrutinize visual features of images, increase visual sampling, which leads to better subsequent memory (Loftus, 1972; Olsen et al., 2016). Given this, we would have predicted within-element saccades to be associated with alpha desynchronization. This was however not found; instead, we found a frontal theta synchronization to be predictive of high event memory. Increased visual sampling results in high memory load, which is reflected in the high frontal theta power (Jensen and Tesche, 2002; Sederberg et al., 2006). In our paradigm, an increased memory load across encoding time corresponds to the accumulation of more visual details about event exemplars, which is favorable for later recall. Alternatively, frontal theta power has also been associated with the detection of interference during selective retrieval in episodic memory (Ferreira et al., 2014; Kerrén et al., 2021; Staudigl et al., 2010) and during multiple-list learning in working memory (Kliegl and Bauml, 2021). In the present study, encoding exemplars from multiple events may be accompanied by interference from exemplars of the same categories in previously acquired events. Therefore, detection of interference may be crucial for encoding distinct memories of each

event and may trigger adaptive inhibition of exemplars from previous events, ensuring that each event is stored separately to succeed in the memory test. It is also conceivable that the frontal theta reflects the complementary effects of interference detection and memory load, as high memory load can lead to increased interference.

Conclusions

We used a novel method of EEG-eye movement coregistration to examine episodic memory encoding in naturalistic viewing with a precision unavailable thus far. Our findings confirm a crucial role of eye movements in the formation of episodic memory (Voss et al., 2017). We demonstrate that the functional specialization of eye movements manifested in various types of saccades is associated with different perceptual and cognitive processes, as evidenced by fixation-locked theta and alpha EEG power. Specifically, two distinct processes reflected in theta synchronization and one process reflected in alpha desynchronization predict the success of episodic memory formation. Episodic memory about an event is constructed through rapid switching between these processes in a sequence of fixations. These processes may indicate the complementary functioning of the hippocampal theta system for binding episodes and the cortical alpha system for representing the content of these episodes (Hanslmayr et al., 2016). Our study extends previous literature by elucidating how these systems engage across eye movements in the service of building coherent episodic memories for personal experiences.

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