



**University of
Zurich**^{UZH}

**Zurich Open Repository and
Archive**

University of Zurich
University Library
Strickhofstrasse 39
CH-8057 Zurich
www.zora.uzh.ch

Year: 2023

Cortico-ocular coupling in the service of episodic memory formation

Popov, Tzvetan ; Staudigl, Tobias

Abstract: Encoding of visual information is a necessary requirement for most types of episodic memories. In search for a neural signature of memory formation, amplitude modulation of neural activity has been repeatedly shown to correlate with and suggested to be functionally involved in successful memory encoding. We here report a complementary view on why and how brain activity relates to memory, indicating a functional role of cortico-ocular interactions for episodic memory formation. Recording simultaneous magnetoencephalography and eye tracking in 35 human participants, we demonstrate that gaze variability and amplitude modulations of alpha/beta oscillations (10–20 Hz) in visual cortex covary and predict subsequent memory performance between and within participants. Amplitude variation during pre-stimulus baseline was associated with gaze direction variability, echoing the co-variation observed during scene encoding. We conclude that encoding of visual information engages unison coupling between oculomotor and visual areas in the service of memory formation.

DOI: <https://doi.org/10.1016/j.pneurobio.2023.102476>

Posted at the Zurich Open Repository and Archive, University of Zurich

ZORA URL: <https://doi.org/10.5167/uzh-252752>

Journal Article

Published Version

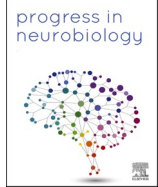


The following work is licensed under a Creative Commons: Attribution 4.0 International (CC BY 4.0) License.

Originally published at:

Popov, Tzvetan; Staudigl, Tobias (2023). Cortico-ocular coupling in the service of episodic memory formation. *Progress in Neurobiology*, 227:102476.

DOI: <https://doi.org/10.1016/j.pneurobio.2023.102476>



Cortico-ocular coupling in the service of episodic memory formation

Tzvetan Popov^{a,b,*}, Tobias Staudigl^c

^a Methods of Plasticity Research, Department of Psychology, University of Zurich, Zurich, Switzerland

^b Department of Psychology, University of Konstanz, Konstanz, Germany

^c Department of Psychology, Ludwig-Maximilians-Universität München, Munich, Germany

ARTICLE INFO

Keywords:

Episodic memory
Brain oscillations
Eye movements
MEG

ABSTRACT

Encoding of visual information is a necessary requirement for most types of episodic memories. In search for a neural signature of memory formation, amplitude modulation of neural activity has been repeatedly shown to correlate with and suggested to be functionally involved in successful memory encoding. We here report a complementary view on why and how brain activity relates to memory, indicating a functional role of cortico-ocular interactions for episodic memory formation. Recording simultaneous magnetoencephalography and eye tracking in 35 human participants, we demonstrate that gaze variability and amplitude modulations of alpha/beta oscillations (10–20 Hz) in visual cortex covary and predict subsequent memory performance between and within participants. Amplitude variation during pre-stimulus baseline was associated with gaze direction variability, echoing the co-variation observed during scene encoding. We conclude that encoding of visual information engages unison coupling between oculomotor and visual areas in the service of memory formation.

1. Introduction

Encoding of visual material is a prerequisite for most of our episodic memories and typically begins with the exploration of the environment. Relying heavily on vision, humans tend to explore the environment by eye movements. These typically include, but are not limited to, micro- and macro saccades as well as fixations jointly contributing to the overall gaze pattern (Otero-Millan et al., 2008). Evidence suggests a strong association between gaze patterns and episodic memory formation (Damiano and Walther, 2019; Fehlmann et al., 2020; Molitor et al., 2014; Olsen et al., 2016; Kragel and Voss, 2022; Broers et al., 2022). Visual scenes that were explored with more eye movements are more likely to be remembered than scenes explored with less eye movements (Voss et al., 2017). These associations between memory encoding and gaze are complemented by studies showing that gaze pattern reinstatement during retrieval supports successful recollection (Johansson et al., 2022; Wynn et al., 2020, 2019; Brandt and Stark, 1997). Similarly, gaze patterns are reinstated during visual imagery (Brandt and Stark, 1997; Bochynska and Laeng, 2015; Laeng and Teodorescu, 2002; Gbadamosi and Zangemeister, 2001) even without the guidance of retinal input, i.e. during full darkness (Johansson et al., 2006).

Electro- and magnetoencephalographic studies confirm a robust

power modulation of alpha/beta oscillatory activity (appr. range 10–20 Hz) during the encoding of items that is predictive of later memory performance: alpha/beta power during later remembered items is reduced as compared to later forgotten items, an observation termed “subsequent memory effect” (SME (Paller and Wagner, 2002)). In electro- and magnetoencephalographic examinations, the time-course of the alpha/beta decrease is usually observed after ~0.5 s and can endure multiple seconds depending on the duration of stimulus presentation. The topography and underlying source generators vary across studies, typically including wider networks of posterior (e.g., (Griffiths et al., 2021a)) but also frontal areas (see e.g., (Hanslmayr et al., 2011)). Alpha/beta SMEs have been replicated consistently (Vogelsang et al., 2018; Fellner et al., 2013; Khader et al., 2010; Hanslmayr et al., 2008; Klimesch et al., 1996; Sederberg et al., 2003), with remarkable specificity of alpha/beta power decreases during successful memory encoding (Hanslmayr et al., 2012; Griffiths et al., 2019, 2021a; Strunk and Duarte, 2019; Sander et al., 2020; Griffiths et al., 2021b; Hanslmayr and Staudigl, 2014). This is of particular relevance as these alpha/beta effects have been interpreted as the mechanism by which the cortex tracks and organizes representations of the stimulus input to be fed forward to downstream memory circuits. According to these ideas, alpha/beta activity is functionally involved in successful episodic memory formation

* Correspondence to: Methods of Plasticity Research Laboratory, Department of Psychology, University of Zurich, Binzmühlestrasse 14, CH-8050 Zurich, Switzerland.

E-mail address: tzvetan.popov@uzh.ch (T. Popov).

<https://doi.org/10.1016/j.pneurobio.2023.102476>

Received 27 January 2023; Received in revised form 25 May 2023; Accepted 30 May 2023

Available online 1 June 2023

0301-0082/© 2023 The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY license (<http://creativecommons.org/licenses/by/4.0/>).

(Hanslmayr et al., 2012; Parish et al., 2018; Hanslmayr et al., 2016).

Complementing such cognitive interpretations, a recent account of alpha/beta activity argues for a view closer to the biological implementation of the association between alpha/beta power modulations and oculomotor action (Popov et al., 2021a). Recent evidence demonstrates topographic modulation of alpha power (i.e. power reduction) associated with consistent eye movements during in a variety of tasks, including visually guided and microsaccades tasks, visual and auditory covert spatial attention, spatial working memory and natural speech comprehension (Staudigl et al., 2017a; Popov et al., 2021b; Printzlau et al., 2022). The topographic modulation of alpha power (i.e. power reduction) was associated with consistent eye movements towards the contralateral visual hemifield (Printzlau et al., 2022; Popov et al., 2022; Liu et al., 2022a). This evidence portrays a fundamental and domain-general functional organization linking visual cortical signals to oculomotor action. We here set out to test the link between eye movements and alpha/beta activity in the context of episodic memory formation.

We hypothesize that alpha/beta power modulations link to episodic memory formation through consistent biases in gaze patterns. Thirty-five participants performed a memory task while simultaneously tracking eye movements and monitoring brain activity using magnetoencephalography (MEG). Gaze bias analyses revealed a robust gaze-

related SME confirming that higher levels of visual exploration are associated with better memory performance. Crucially, high subjective confidence in correctly memorizing an item was predictive of gaze-related and alpha/beta SME. Moreover, splitting trials according to high versus low gaze bias revealed the typical alpha/beta SME over posterior sensors. During the baseline interval prior to scene encoding, spontaneous fluctuations of alpha/beta power reliably predicted the gaze direction variability, akin to the one observed during scene encoding. Exploratory analyses of the covariation between gaze direction and alpha/beta power confirmed a consistent relationship present throughout the entire recording and evident between and within participants.

2. Results

2.1. Memory performance evident in both gaze variability and power modulation of visual cortex activity

During the study phase, participants viewed visual scenes presented for 4 s. For each trial and participant, the simultaneously recorded eye tracking data were extracted and converted into 2D density heat maps with the x-axis representing the horizontal, y-axis vertical visual eccentricity. The density of gaze direction locations is color coded.

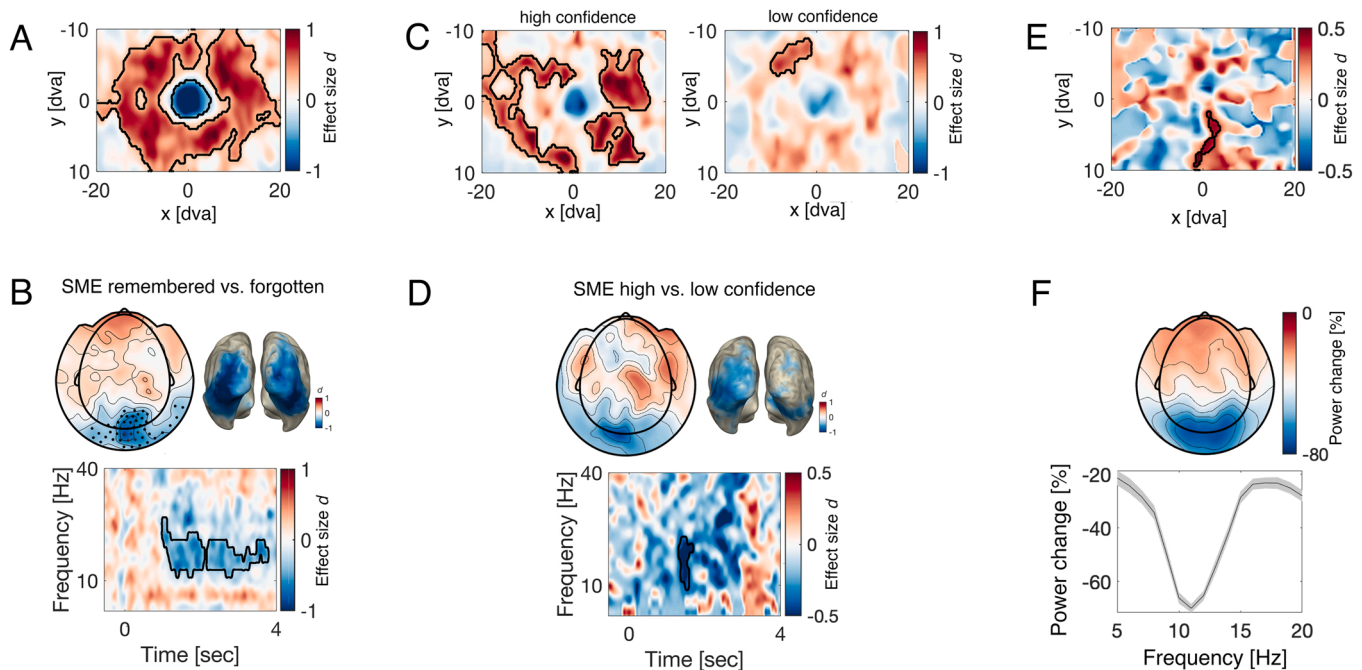


Fig. 1. Coinciding subsequent memory effects (SME) reflected in gaze biases and modulation of visual alpha/beta power: A-Gaze density contrast between later remembered vs. later forgotten items. The 2D densitymaps were calculated per trial and latency between 0- 4sec following scene presentation during the study phase. Red color indicates increased biasedtowards the respective visual screen locations for later remembered items and blue color decreased gaze density or gaze direction away from the respective location. Color code is expressed in units of effect size Cohen's d obtained after cluster-permutation test ($p < 0.025$). B- Scalp topography and time-frequency representation of power (TFR) of the contrast between later remembered vs. later forgotten items during the study phase. Color code indicates effect size Cohen's d obtained after cluster-permutation approach at $p < 0.025$ (sensors belonging to the cluster marked with black dots). The black outline in the TFR highlights the cluster supporting the rejection of the null hypothesis of no effect between later remembered and later forgotten items. TFR spectrogram is averaged across the sensors marked in the topography. Source reconstruction of the alpha/beta effect is illustrated highlighting the recruitment of early visual, parietal and ventral cortical areas. Source maps are thresholded using cluster-permutation test ($p < 0.025$) and expressed in units of effect size Cohen's d . C- Similar to A but split according to the participants' confidence as of how certain their response is for an item being "old" or "new". Gaze bias for high confidence trials (left) versus gaze bias for low confidence trials (right). D- 2x2 Interaction SME (remembered, forgotten) x Confidence (high, low) illustrating the scalp topography, the TFR and the corresponding source map expressed in units of effect size Cohen's d (cluster-permutation test, $p < 0.025$). E- Contrast in gaze density between trials dominated by low alpha/beta power minus high alpha/beta power during the prestimulus baseline (-1 to 0 sec) in the study phase. Color code is expressed in units of effect size Cohen's d obtained after cluster-permutation test ($p < 0.025$). Red color indicates increased bias towards the respective visual screen locations for trials dominated by low alpha/power power as compared to high alpha/beta power. F- Scalp topography of the relative alpha/beta power change between the same trials as in (E) dominated by low vs. high alpha/beta power. Color code denotes the relative change in %. Statistical contrast and source maps are inappropriate in this case as they will be highly significant per construction. This is however not the case for the evaluation of the gaze biases illustrated in E. The power spectrum of the difference averaged across occipital sensors is provided with shading denoting SEM.

Subsequently, a statistical contrast using non-parametric testing with clusters was performed (Maris and Oostenveld, 2007). This analysis revealed a robust gaze-related SME (Fig. 1A, cluster-permutation test, $p < 0.025$; corrected for multiple comparisons across time and frequencies), indicating that, during the study phase, items that will be later remembered are characterized by a strong gaze bias away from fixation and towards various locations of the viewed scene. Quantification of the different eye movement types such as saccades and fixations using the EEG-EYE toolbox (Dimigen et al., 2011; Engbert and Mergenthaler, 2006) yielded similar results (supplemental Fig. S1). For later remembered items, the average eye events per scene were fixations (M/STD = 13.8/1.8) and saccades (M/STD = 14.1/1.9) and for forgotten items fixations (M/STD = 13.2/2.6) and saccades (M/STD = 13.4/2.7). A significant condition difference was confirmed for both, fixations ($t_{34} = 2.3$, CI [0.08 1.23], $p = 0.026$, Cohen's $d = 0.393$) and saccades ($t_{34} = 2.3$, CI [0.08 1.24], $p = 0.026$, Cohen's $d = 0.391$). In the test phase, participants correctly recognized 72.83% (+/- 2.51 SEM) of the scenes on average, yielding a D -prime of 2.24 (+/- 0.09 SEM).

Applying a standard (i.e. in accordance with previous work in the field) subsequent memory analyses on the MEG data confirmed the well documented alpha/beta SME (Fig. 1B). Encoding of later remembered material was associated with a stronger alpha/beta decrease over occipital sensors (cluster-permutation test, $p < 0.025$; corrected for multiple comparisons across time and frequencies). The variability of the SME effect size observed here and enclosed within the black outline of Fig. 1 B is within the range of -0.8 to -0.5 (or 5 to 8 in absolute values). These effect sizes are comparable to earlier reports estimating an SME effect size of 0.56 (Griffiths et al., 2019) and 0.6 (Griffiths et al., 2021a). Scene exploration (i.e., stronger gaze biases away from fixation for later remembered items) as well as the modulation of visual alpha/beta power (i.e., stronger decrease for later remembered items) were informative for the participant's memory confidence. Items reported with high confidence in being correctly memorized displayed stronger gaze biases (exploration) during encoding (Fig. 1 C left, cluster-permutation test, $p < 0.025$) and alpha/beta SMEs (Fig. 1 D, cluster-permutation test, $p < 0.025$), as compared to items with lower confidence (Fig. 1 C right; see Fig. S6, for a time-resolved version of this analysis). Source reconstruction of the SME effect using beamforming indicated the recruitment of a network of visual, parietal and ventral-temporal cortical areas (Fig. 1B). Largely similar, albeit less pronounced source networks are indicated for the low vs. high confidence SME (Fig. 1D).

A complementary interpretation of the apparent relationship between eye movements and the modulation of visual alpha/beta power might be the subjective saliency of the stimulus material. Saliency is hard to quantify objectively and strongly depends on subjective experience. Hence, some stimulus material could trigger both, variability in eye movements and alpha/beta power modulations. It could allow the conjecture that the apparent correlation between the two is driven by external stimulus factors. In order to test this alternative, we set out to explore a potential relationship between gaze variability and power modulation of ongoing alpha/beta activity using the data of the pre-stimulus (e.g. baseline) intervals during the study phase, where only a fixation cross instead of a visual scene was displayed. Participants were required and were successful in maintenance of central fixation, ranging within $\pm 5^\circ$ of horizontal and vertical visual angle (see Fig. S2). We reasoned that, within participants, one could split the trials into high and low alpha/beta power (e.g. based on median split) during the pre-stimulus baseline interval. Subsequently, one could evaluate the distribution of gaze density around the required fixation. If the above conjecture is true, there should be no difference in the gaze variability between high and low alpha/beta power conditions, since the stimuli in the baseline period (that is, the fixation cross) did not vary in saliency. Conversely, differences in gaze bias would further confirm a co-varying relationship between the direction of gaze and modulation of visual alpha/beta power, independent of the task context and stimulus

properties. The results of this analysis are illustrated in Fig. 1 E and F. Within participants, trials with less alpha/beta power were also associated with a gaze pattern that deviated more from fixation (Fig. 1 E). Conversely, high alpha/beta power during baseline was associated with reduction of the degrees of freedom in eye movements, resulting in less variability of gaze around the targeted (fixation) direction. This result echoes the observations made during stimulus presentation (e.g. Fig. 1A, B), both in terms of the direction of the relationship (i.e. alpha/beta power decrease relates to higher gaze variability) and scalp topography (Fig. 1F). Hence, the complementary interpretation that the relationship between alpha/beta SEM and gaze bias is dependent on saliency appears less likely.

2.2. Gaze variability predicts both modulation of visual cortical activity and memory performance

Next we asked to what extent the inverse direction of high vs. low gaze variability will predict alpha/beta SMEs. Within participants we median split the trials during encoding into high and low gaze variability within the visual display area of high variability identified in Fig. 1A (i.e. positive cluster). Trivially, the group difference in gaze variability is significant per design (Fig. 2A). Importantly however this should not necessarily hold for cortical alpha/beta power and/or memory performance. We observed that trials dominated with high gaze variability were also associated with stronger modulation of alpha/beta power over occipital sensors, akin to the alpha/beta SME topography observed in Fig. 1B (cluster-permutation test, $p < 0.02$). Moreover, high gaze variability during encoding was associated with an increase in the number of later remembered items as compared to low gaze variability (Fig. 2C, left). The opposite relationship was observed for forgotten items (Fig. 2C, right). An alternative illustration of the same result is depicted in Fig. 2D. The effect size of the memory contrast (# items remembered vs. forgotten) was dependent upon the gaze exploration during encoding: high gaze variability is linked to better memory performance as compared to low gaze variability (Fig. 2D). In a supplementary analysis (Fig. S8), we show that gaze variability covaries with the "memorability" of the stimuli (i.e., some stimuli are consistently better remembered than others across observers; see, e.g., Bainbridge et al. (2017) and Bainbridge et al. (2013)), in line with previous findings indicating that eye movement patterns reliably predict this effect (Bylinskii et al., 2015).

2.3. Gaze variability and visual cortical activity are continuously coupled

Given the results described above, we next asked whether a co-fluctuation of gaze bias and alpha/beta power would be restricted to the memory task or could be a more general pattern. Specifically, we asked whether alpha/beta power and gaze bias would be correlated over the whole period of the experiment. The results of this analysis are summarized in Fig. 3.

First, we computed the time-frequency representation of power (TFR) for the entire recording per participant (M/SEM = 60.59/ ± 2.01 min) and accounted for the presence of aperiodic activity (Donoghue et al., 2020a). For illustrative purposes, Fig. 3A illustrates the first 6 min in the recording of a representative participant. The TFR depicts the spontaneous power modulation of alpha (predominantly in 10–14 Hz) activity over occipital sensors. In addition, we extracted the fixation density for the horizontal direction averaged across the vertical direction (see Fig. S5, for complementary analyses using fixation density for the vertical direction averaged across the horizontal direction). This fixation density signal can be visualized as a function of time with the horizontal viewing direction now illustrated on the y-axis (Fig. 3B). In both panels (Fig. 3A and B), the power of the signals is range corrected ($[x - \min]/[\max - \min]$) to vary between 0 and 1. The time series of alpha/beta power and fixation density can be extracted and overlaid (Fig. 3 C) conveying a (descriptive) similarity between the variation in

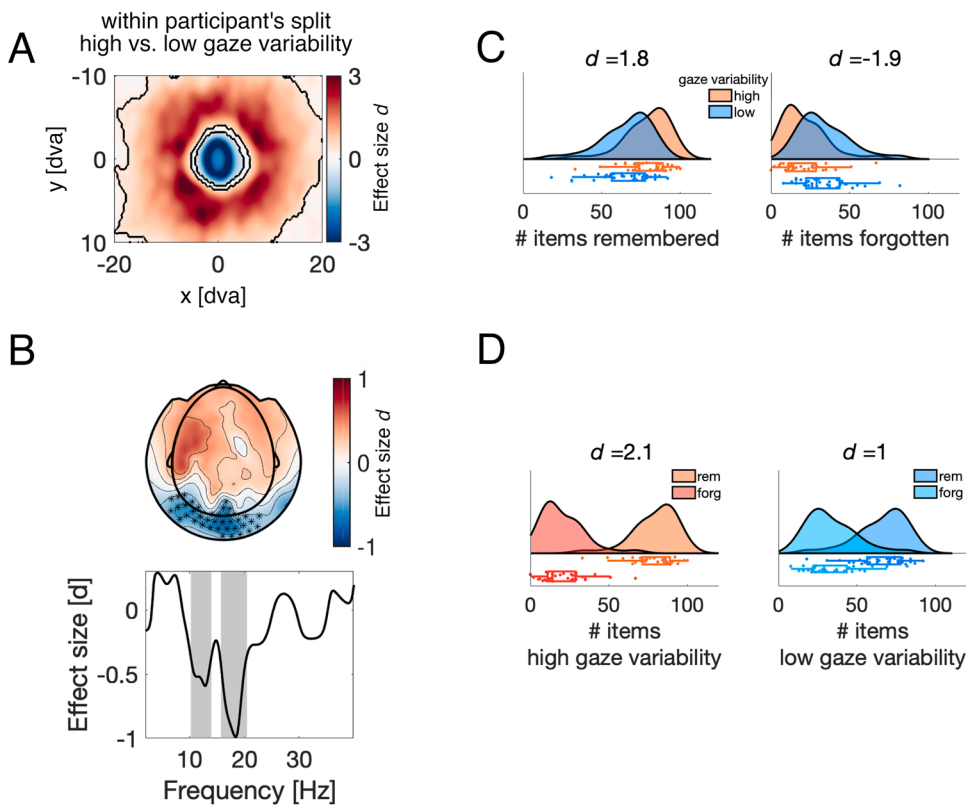


Fig. 2. Gaze variability affects cortical alpha/beta power modulation and the effect size of memory performance. A- Gaze density contrast between trials dominated by high vs. low gaze variability (median split within participants). The 2D density maps were calculated per trial and latency between 0 and 4 s following scene presentation during the study phase. Red color indicates increased biased towards the respective visual screen locations for later remembered items and blue color decreased gaze density or gaze direction away from the respective location. Color code is expressed in units of effect size Cohen's d obtained after cluster-permutation test ($p < 0.025$). B- Scalp topography and power spectrum contrast between high vs. low gaze variability trials during the study phase. Color code indicates effect size Cohen's d obtained after cluster-permutation approach at $p < 0.025$. The black asterisks in the topography and the shaded areas in the powerspectrum highlight the clusters supporting the rejection of the null hypothesis of no effect between trials dominated by high vs. low gaze variability during encoding. C- Rain cloud plots (Allen et al., 2019) illustrating the distribution of remembered (left) and forgotten (right) items split by gaze variability (high vs. low) during encoding. D- The distribution of high (left) and low (right) gaze variability split by memory performance (remembered vs. forgotten).

alpha/beta power and the variation in fixation maintenance. Using this approach, we computed Spearman's rho correlations between both timeseries for the entire recording session in each participant. In addition, surrogate data was generated by circularly shifting both time series (1000 times) and computing the corresponding correlation. This analysis yielded two distributions of correlations that can be parametrically compared (Fig. 3D). For each participant we derived two correlation coefficients. These correlations obtained from the original data were significantly different from the one generated by the surrogate data ($t_{(34)} = 6.8$, CI[0.12 0.22], $p < 7.4e-08$, Cohen's $d = 1.1541$). This effect was confirmed on the individual participant level, with 33 out of 35 participants showing a significant difference (Fig. S3). We also computed the coherence between the two timeseries to infer the frequency of any phase relationship in the covariation of gaze and occipital alpha/beta power. (see Fig. S10 for coherence between alpha power and fixation density in low frequencies peaking at 0.18 Hz).

The hypothesis related to the subsequent memory effects and results illustrated in Fig. 1 are motivated by previous work. In Fig. 2, we depict the outcomes from exploratory analyses motivated by the results illustrated in Fig. 1. While SME effects are typically based on task-specific contrasts, i.e. difference between task and pre-stimulus baseline, the results reported in Fig. 3 reveal a relationship between spontaneous alpha power fluctuations and spontaneous shifts of gaze. Thus, even though it might be tempting to conclude on the basis of the results in Fig. 1 only that the gaze-alpha power relationships are task specific, the results illustrated in Fig. 3 confirms that this relationship is spontaneous and task independent yet utilized during episodic memory formation.

3. Discussion

Previous research has shown that gaze patterns as well as amplitude modulation of neural activity in posterior cortex during encoding of visual scenes strongly predicts memory performance. However, gaze patterns and amplitude modulation have thus far been studied in isolation. Our findings indicate that we might be looking at two sides of

the same coin: gaze patterns and amplitude modulations covary and jointly predict memory performance.

We simultaneously recorded MEG and eye movements in a free viewing memory paradigm. To relate alpha/beta MEG activity as well gaze variability to successful encoding, we contrasted later remembered versus later forgotten study phase items. In line with many previous studies (Vogelsang et al., 2018; Fellner et al., 2013; Khader et al., 2010; Hanslmayr et al., 2008; Klimesch et al., 1996; Sederberg et al., 2003; Hanslmayr et al., 2012; Griffiths et al., 2019, 2021a; Strunk and Duarte, 2019; Sander et al., 2020; Griffiths et al., 2021b; Hanslmayr and Staudigl, 2014), we find an alpha/beta SME, with greater decreases in 10–20 Hz power for later remembered as compared to later forgotten items. Importantly, we here demonstrate that gaze covaries with this well-established effect. During the presentation of later remembered items, gaze was directed away from the center to a greater extent than during later forgotten items. This gaze bias indicates that more extensive visual exploration correlates with a higher probability to remember the item, a finding in line with previous studies showing that the amount of saccades on complex visual stimuli predicts whether or not the stimulus will be remembered (Voss et al., 2017). Just like the alpha/beta SME, the gaze-related SME also correlated with the reported confidence judgement. Items that were later remembered with high confidence displayed stronger gaze biases and stronger alpha/beta desynchronization than those items remembered with lower confidence. Moreover, we find that splitting the encoding EEG data between high vs. low gaze variability trials closely emulates the traditional alpha/beta SME, both in frequency and topography. Trials with high gaze bias display significantly stronger alpha/beta power decreases than trials with lower gaze bias. Contrasting subsequent memory performance based on gaze bias confirmed that higher gaze bias was predictive of successful memory performance. These results indicate that gaze variability and alpha/beta activity go hand in hand in service of memory formation, a notion in line with a previous study linking saccades and the phase of alpha/beta activity successful remembering (Staudigl et al., 2017b).

Generalizing the covariation of gaze and alpha/beta activity beyond

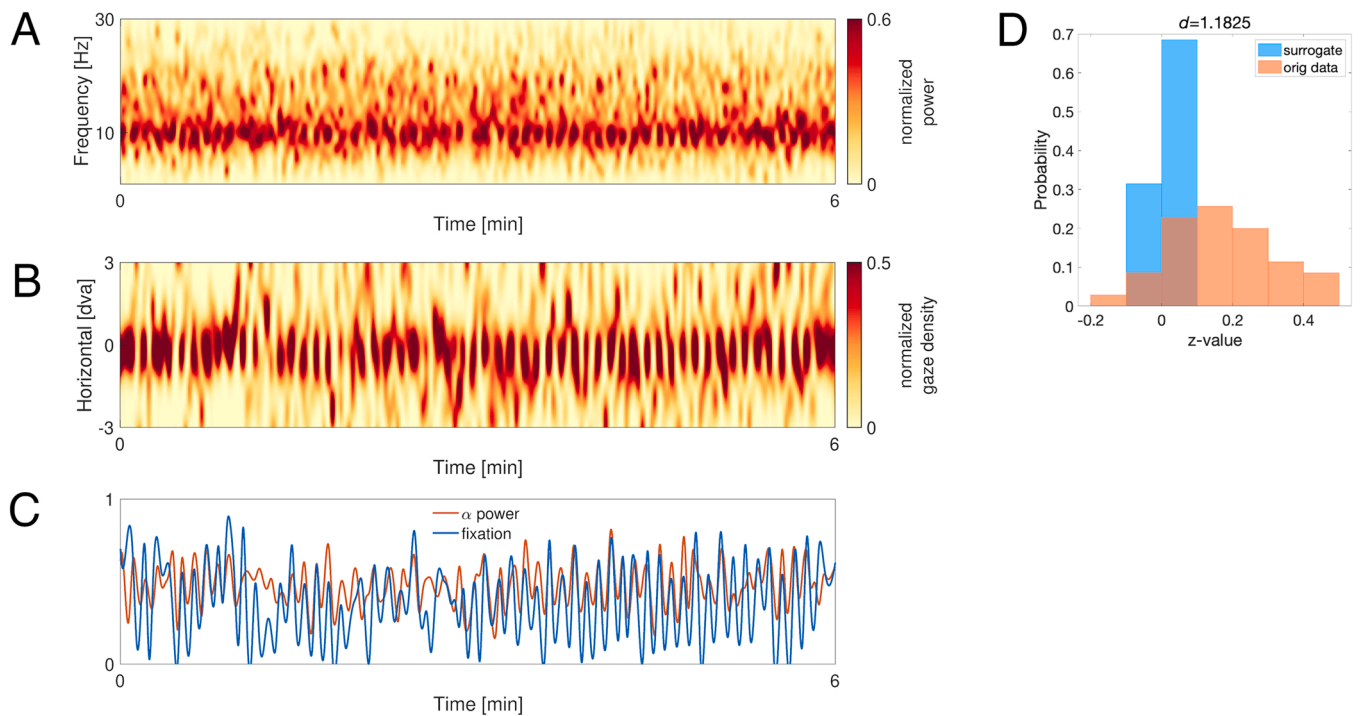


Fig. 3. Time course of the covariation of occipital alpha power and gaze direction maintenance. A-Single subject example of TFR averaged over occipital sensors, corrected for the presence of the aperiodic 1/F component. X-axis denotes time in minutes and y-axis frequency in Hz. The first 6 min of the recording are shown for illustrative purposes. Color code denotes the range corrected power $(x - \min)/(\max - \min)$ varying between 0 and 1. B- Gaze density in the horizontal dimension around the fixation location and vertical range of ± 3 dva expressed as function of time (x-axis). The middle position of the y-axis denotes the fixation location, up rightward gaze bias and down denotes leftward gaze bias. Color code illustrates the range corrected gaze density varying between 0 and 1. C- Time course of alpha power (10–14 Hz, red color) and gaze density at the fixation location (blue color) overlaid on top of each other. The x-axis is identical to A and B. D- Normalized probability histograms (sum of y-axis equals 1) of the Spearman's rho correlation coefficient between the time courses of alpha power and gaze density at fixation, converted in z-values. Red color denotes the distribution of all correlations across all participants estimated on the original time series. Blue color denotes the distribution of correlations estimated on circularly shifted time series after 1000 permutations. A significant difference was confirmed $t(34) = 6.8$, $CI[0.12 \ 0.22]$, $p < 7.4e-08$ and estimated effect size of Cohen's $d = 1.1541$.

memory encoding, we show that gaze covaries with spontaneous fluctuations of alpha/beta during the pre-stimulus baseline. Again, greater variation in gaze was correlated with larger decreases in alpha/beta power. Moving even further away from specific, task-related activity, we show that alpha/beta activity and gaze covary over the course of the whole experiment. Including an hour of simultaneous eye tracking and MEG recordings per participant, we show substantial covariation of alpha power and gaze that is task-independent. Taken together, these observations support the idea of a fundamental and domain-general link between visual alpha/beta activity and oculomotor action.

The present results are in line with recent studies indicating a strong link between eye movements and alpha/beta activity. Biases in gaze direction are associated with a topographically consistent decrease of alpha/beta power, independent of the stimulus modality (e.g. vision, audition) (Popov et al., 2021b), independent of retinal input and present during full darkness (Popov et al., 2021a). Similar associations between gaze bias and alpha/beta power modulation have recently been reported in the context of working memory (Printzlau et al., 2022; Liu et al., 2022b). Together with the here reported results, these studies make a strong case that the association between gaze and alpha/beta activity reflects a fundamental functional organization linking visual cortical signals to oculomotor action.

The present findings are also in line with previous studies showing a functionally relevant coordination of eye movements and brain activity. Eye movements affect the neural activity in- and outside visual areas (Staudigl et al., 2017b; Brunet et al., 2015; McFarland et al., 2015; Ito et al., 2011; Rajkai et al., 2008; Barczak et al., 2019; Leszczynski et al., 2021; Hamamé et al., 2014; Purpura et al., 2003), as well as in brain areas crucial for episodic memory formation: neural activity in the

primate hippocampus is sensitive to eye movements (Andrillon et al., 2015; Mao et al., 2021; Katz et al., 2022; Wagner et al., 2022; Liu et al., 2017). In particular, the phase of hippocampal low-frequency activity has been shown to be aligned to saccadic eye movements (Hoffman et al., 2013; Doucet et al., 2020), and the amount of alignment was found to be functionally relevant (Jutras et al., 2013; Staudigl et al., 2022). Such phase alignment could be a general motif in the nervous system to facilitate the organization of neural ensembles (Vолоh and Womelsdorf, 2016). The exact temporal coordination of neural activity and eye movements could be brought about by efference copies (Sommer and Wurtz, 2002). These copies of motor commands branch off corticofugal projections and could, in principle, be distributed across many brain areas. If they are capable of inducing phase alignment, efference copies could trigger the alignment of neural activity across a broad range of areas and, thereby, facilitate neural processing and communication (Buzsáki, 2010; Fries, 2015; Schneider et al., 2021).

We here demonstrate a link between eye movements and alpha/beta activity during the free viewing of visual scenes in the study phase of a memory paradigm. Alpha/beta SMEs during memory encoding are indeed a common finding, yet to what extent the present linkage between alpha/beta SME and gaze variability generalize across stimulus material such as faces and words (Fellner et al., 2019) vs. visual scenes requires further examination. Moreover, one could argue that the present findings are limited to free viewing encoding conditions, as the majority of the studies investigating SMEs ask their research participants to maintain fixation. Hence, the observation of alpha/beta variation with gaze variability might be somewhat coincidental. We hypothesize that the present findings will generalize to such conditions as well. This prediction is based on the present observation that the alpha/beta-gaze

relationship was, albeit weaker, also evident when the eyes do not move freely (e.g. maintain fixation during the baseline period, Fig. 1E,F). As fixation is not a stationary event but a process involving the control of miniature eye movements such as micro saccades (Martinez-Conde et al., 2013), a testable prediction is that independent of task instructions (fixation, free view), alpha/beta SME modulation should be also associated with gaze-related SME. An empirical question that can be examined in future studies.

So, does cortico-ocular coupling serve episodic memory encoding, as the title of this paper suggests? The answer depends on how you specify the question. Cortico-ocular coupling serves memory, because it is utilized during exploration of later remembered scenes more so than when exploring later forgotten material. However, cortico-ocular coupling is neither specific nor restricted to the memory task at hand but appears to be a more general phenomenon that might be utilized upon task demands.

In conclusion, we demonstrate that alpha/beta activity (10–20 Hz) in visual cortex and eye movements covary and jointly predict subsequent memory. Going beyond memory-related brain activity, we show that this covariation is task-independent and preserved over the course of hours. The present results thus support a complementary view on the role of alpha/beta activity, emphasizing its fundamental interrelation with oculomotor behavior.

4. Materials and methods

Part of the present dataset has been previously analyzed and documented elsewhere (Staudigl et al., 2017b).

4.1. Ethic statement

Study enrollment followed the approval of the local ethics committee-commission for human related research CMO-2014/288 region Arnhem/Nijmegen, the Netherlands. Prior to participation all volunteers were given written informed consent in accordance with the Declaration of Helsinki.

4.2. Participants

A total of 48 participants were recruited. Thirty-five were included in the present study. Thirteen participants were excluded due to: not completing the study ($N = 7$), excessive artifacts ($N = 3$) and technical difficulties during acquisition ($N = 3$). The included participants sample (24 female, age range 18–30, mean age 23.1 y) reported no history of neurological and/or psychiatric diagnosis and had normal or corrected to normal vision.

4.3. Task design and procedure

Visual scenes were presented to the participants projected onto a back-projection screen position in front of the participants inside a magnetically shielding room (MSR). The back-projection screen had the size of 39×46 cm corresponding to a visual angle of vertical $27^\circ \times 32^\circ$ horizontal dva (degrees of visual angle). Three sets of visual scenes (100 scenes each) were used. Two sets were displayed during the study (encoding) and the test phase. The scenes in the third set served as lures and were presented exclusively during the test phase. Set assignment to study and test phase was counterbalanced across participants. A training session preceded the actual memory task to ensure participants familiarity with the task procedures. Nine additional scenes not used in the main experiment were shown during the training session. All participants were made aware about the memory test prior to data acquisition.

During the study phase, visual scenes (indoor and outdoor images) were displayed for 4 s with the constrained that no more than 4 scenes from the same category could appear consecutively. Participants were instructed to report via button press whether or not the current scene

displayed was an indoor or outdoor image. The response was given during the display of a fixation cross with a variable duration between 1 and 2 s presented after each scene. During scene viewing, participants were not required to maintain fixation but were allowed to freely explore the scenes.

A short distracter session was required after each study phase. This session consisted of solving simple mathematical problems (appr. 1 min), a simple saccade task during which participants had to saccade to various locations on the screen (5 min), eyes open and eyes closed data acquisition (appr. 1 min each). The purpose of this distracter session was to prevent participants from covert rehearsing.

The test phase followed thereafter. The three sets of images (200 old items from the study phase intermixed with 100 new items) were randomly presented for viewing duration of 4 s. The randomization had the constrained that no more than 4 images of the same type (old/new) could be shown consecutively. Following each image presentation, a 6-point response scale was displayed. Participants were required to indicate whether the just presented image was “old” or “new” with the scale ranging from “very sure old”(1) to “very sure new”(6). The 6-point scale remained on the visual display until the participants response was given followed by a fixation cross of variable duration ranging between 0.75 and 1.25 s. A remembered item is defined by a correct response (old; 1, 2 or 3) that was provided in the test phase when an old item was presented; a later forgotten item is defined by an incorrect response (new; 1, 2 or 3) that was provided in the test phase when an old item was presented.

4.4. Data acquisition

Whole head magnetoencephalography (MEG) was acquired with a 275-axial gradiometer system (VSM MedTech/CTF MEG, Coquitlam, Canada) within the MSR. During data acquisition, the sampling frequency was set to 1200 Hz using a low-pass antialiasing filter at a cutoff frequency of 300 Hz. Ocular artifacts were monitored by horizontal and vertical electrooculogram using Ag/AgCl electrodes and a bipolar montage. Head movements were tracked continuously using 3 coils placed in the vicinity of the nasion and the left and right ear canals (Stolk et al., 2013). An Eyelink 1000 (SR Research) system was used to monitor horizontal and vertical eye movements of the left eye. Prior to data acquisition, eye tracker calibration was performed which involved the collection of gaze fixation samples from pre-defined positions (9 dots on a 3×3 grid) on the visual display. Raw eye tracking data was mapped to these pre-defined screen coordinates followed by a validation procedure to ensure sufficient correspondence between the position of the current gaze fixations and the one obtained during the preceding calibration. The calibration was accepted if the difference was $< 1^\circ$ dva.

4.5. Data preprocessing and analysis

4.5.1. MEG data

Offline data analyses was performed with the open source software for neuroelectric- and neuromagnetic data analysis (FieldTrip (Oostenveld et al., 2011)). The continuous data was segmented around the events of interest (scene onsets) into epochs of 8 Section (3 s baseline prior to event onset). Following a finite impulse response band-pass filter (1–40 Hz) the epochs were re-segmented to exclude potential filter artifacts resulting in time range of -2.5 – 4.5 s around the event onset. Oculo-muscular and cardiac artifacts were identified and removed from further analysis by means of independent component analysis (ICA; see Fig. S9, for a supplementary analysis without applying ICA to the data). These procedures were applied to all data epochs extracted during the study (encoding) and test (retrieval) phase. Subsequently, for each epoch, time-frequency estimates of power were computed using a sliding window of 0.5 s and a Hanning taper resulting in a frequency resolution of appr. 2 Hz. The window slid every 50 ms within the range of -2 – 4 s around the event onset. Power estimates

were averaged across epochs for each condition separately, e.g. encoding (remembered/forgotten), retrieval (remembered/forgotten). Baseline correction using the time window of -1 to -0.25 (encoding) and -0.75 to -0.25 (retrieval) was applied transforming the raw power estimates into dB change from pre-stimulus baseline. The number of trials used in the present analysis were $N_{\text{remembered}} = 145.15/30$ (M/STD) and $N_{\text{forgotten}} = 54.77/30$ (M/STD).

We followed a similar approach to split the epochs into high versus low pre-stimulus alpha power during scene encoding. Specifically, a Fast Fourier transform was applied to the data prior to stimulus onset (-1 to 0 s), including all occipital sensors (see Fig. S4). A Hanning taper was used, resulting in appr. 1 Hz frequency resolution. The mean power within the alpha/beta frequency range (10–20 Hz) was extracted and averaged over all occipital sensors. Subsequently, trials with high and low alpha power were separated by means of median split. Using this subset of high and low alpha power trials, the power estimates were re-grouped and averaged over trials. Finally, the difference between the condition's alpha power low minus alpha power high was computed.

4.5.2. Source analysis

Source reconstruction was performed using frequency domain spatial filtering algorithm (Gross et al., 2001). Using the cross-spectral density matrix constructed for a frequency of interest (e.g. alpha/beta 14 Hz \pm 4 Hz) the algorithm computes a spatial filter specific for a given brain location (i.e. voxel, 1 cm³). These filters were computed using all data during the pre and post stimulus intervals (e.g. common filter approach). Individual head models were computed using structural MR images aligned to the coordinated system of the MEG device. The details of the MR acquisition and alignment procedure are detailed in (Staudigl et al., 2017b).

4.5.3. Gaze data

The raw eye tracking data was converted from voltage to pixel coordinates following the procedures described here https://www.fieldtriptoolbox.org/getting_started/eyelink/#what-are-the-units-of-the-eye-tracker-data. Gaze density was expressed as the 2D heat map according to the procedures described here (<https://stackoverflow.com/questions/46996206/matlab-creating-a-heatmap-to-visualize-density-of-2d-point-data>). Briefly, the scatter plot of all x and y positions was converted into an image array (using *imagesc.m* in MATLAB) after a 2-D convolution (*conv2.m*) with a Gaussian filter matrix G . For the given range of x and y coordinates denoted as xG and yG and width parameter sigma (in the present case set to 2.5), $G = \exp[-xG^2/(2 * \sigma^2) - yG^2/(2 * \sigma^2)]$. After 2D conversion the eye tracking data was converted to a structure that can be read by FieldTrip akin to the one for time-frequency data. Subsequently, all approaches available for statistical treatment of time-frequency data could be applied to the gaze density data. An example illustrating the 2D heat map procedure is provided in Fig. S7.

4.5.4. Relationships between MEG and gaze data

In order to examine the relationship between the continuous MEG recording and the gaze variation as presented in Fig. 2 the following procedures were utilized. First, the continuous data from occipital sensors was segmented into trials of 2 s length. Following Fourier analysis as described above, the $1/f$ aperiodic component was removed from each trial using the *specparam* routines (Donoghue et al., 2020b) allowing the parametrization and visualization of periodic components (e.g. alpha activity) in the continuous data. Subsequently, these trials were concatenated yielding a time-frequency representation of power (e.g. Fig. 2 A). Next, for each trial, the corresponding gaze data was computed as described above, averaged along the vertical direction and concatenated, yielding a time by horizontal position spectrogram with color coded density of horizontal gaze direction (e.g. Fig. 2B). For each participant, the time course of occipital alpha power was extracted by averaging the spectral estimates across the dominant frequency band

width (e.g. 10–14 Hz) across all occipital sensors. The reason for taking 10–14 Hz rather than alpha/beta activity is that in spontaneous recordings the dominant rhythm is alpha, whereas alpha/beta decreases are scored as function of baseline or some contrast that in this case we do not have. Instead we took the spontaneous signal exhibiting strongest power mostly in the 10–14 Hz range. Similarly, the time course of gaze density averaged around fixation at 0 dva (\pm 0.5 horizontal dva) was extracted. These time courses were correlated using a bootstrapping resampling procedure with 1000 iterations yielding a distribution of correlations between the two time courses (e.g. Fig. 2D). Surrogate data was generated by circularly shifting the time series 1000 times, where at each iteration, the length of the time shift was randomly chosen ranging between 1 s and the total duration of the recording. This procedure resulted in a surrogate distribution of correlations (e.g. Fig. 2D), against which the distribution of correlations obtained from the original data was compared. All correlation coefficients were transformed into z -scores (e.g. $z = \log((1+r)/(1-r))/2$) prior to further statistical evaluation. Finally, the coherence between the MEG and gaze time series was computed. Given a sampling frequency of 0.5 Hz determined by the length of the trials used to derived these time courses (see above), the MEG and gaze data was first re-segmented into epochs of 200 s length. A multi-taper approach (Mittra and Pesaran, 1999) was used to estimate power and cross-spectral density for each trial padded with zeros at the begin and the end of each trial (500 s in total). Three tapers were used covering the frequency range from 0 to Nyquist frequency (0.25 Hz). The phase in the cross-spectra represents the phase difference between the oscillatory signals of the MEG and gaze time series, with consistent phase difference resulting in larger coherence values. Surrogate data was generated by permuting the trial order of the MEG and gaze data with respect to each other and computing coherence. This approach was repeated 1000 times and the resulting coherence values were averaged resulting in the surrogate coherence result presented in Fig. 2E.

4.5.5. Statistics

Statistical control followed the cluster-based permutation framework (Maris and Oostenveld, 2007). This approach utilizes clustering across sensors, time points, and frequency (wherever appropriate). Multiple comparisons problem was addressed by using 1000 permutations and a two-tailed alpha threshold of 0.05 ($p < 0.025$). Correlations were evaluated by utilizing Spearman's correlation coefficient ρ .

Funding

This work was supported by the European Union's Horizon 2020 research and innovation program (grant number 661373) and the European Research Council (ERC-StG 802681) awarded to TS and the Schweizerischer Nationalfonds zur Förderung der Wissenschaftlichen Forschung (SNF) Grant 105314_207580 awarded to TP.

Declaration of Competing Interest

The authors report no competing interests.

Data availability

Data will be made available on request.

Acknowledgements

The study was performed at the Donders Institute for Brain, Cognition and Behaviour. The authors would like to thank Ole Jensen and Christian F. Doeller for their involvement in conceptualizing the study. The authors would also like to thank all the participants volunteering in the experiment.

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at [doi:10.1016/j.pneurobio.2023.102476](https://doi.org/10.1016/j.pneurobio.2023.102476).

References

- Allen, M., Poggiali, D., Whitaker, K., Marshall, T.R., Kievit, R.A., 2019. Raincloud plots: a multi-platform tool for robust data visualization. *Wellcome Open Res* 4, 63. <https://doi.org/10.12688/wellcomeopenres.15191.1>.
- Andrillon, T., Nir, Y., Cirelli, C., Tononi, G., Fried, I., 2015. Single-neuron activity and eye movements during human REM sleep and awake vision. *Nat. Commun.* 6, 7884. <https://doi.org/10.1038/ncomms8884>.
- Bainbridge, W.A., Isola, P., Oliva, A., 2013. The intrinsic memorability of face photographs. *J. Exp. Psychol. Gen.* 142, 1323–1334. <https://doi.org/10.1037/a0033872>.
- Bainbridge, W.A., Dilks, D.D., Oliva, A., 2017. Memorability: A stimulus-driven perceptual neural signature distinctive from memory. *NeuroImage* 149, 141–152. <https://doi.org/10.1016/j.neuroimage.2017.01.063>.
- Barczak, A., et al., 2019. Dynamic modulation of cortical excitability during visual active sensing. *Cell Rep.* 27 (3447–3459), e3443 <https://doi.org/10.1016/j.celrep.2019.05.072>.
- Bochynska, A., Laeng, B., 2015. Tracking down the path of memory: eye scanpaths facilitate retrieval of visuospatial information. *Cogn. Process* 16 (Suppl 1), 159–163. <https://doi.org/10.1007/s10339-015-0690-0>.
- Brandt, S.A., Stark, L.W., 1997. Spontaneous eye movements during visual imagery reflect the content of the visual scene. *J. Cogn. Neurosci.* 9, 27–38. <https://doi.org/10.1162/jocn.1997.9.1.27>.
- Broers, N., Bainbridge, W.A., Michel, R., Balestrieri, E., Busch, N.A., 2022. The extent and specificity of visual exploration determines the formation of recollected memories in complex scenes. *J. Vis.* 22, 9. <https://doi.org/10.1167/jov.22.11.9>.
- Brunet, N., et al., 2015. Visual cortical gamma-band activity during free viewing of natural images. *Cereb. cortex (N. Y., N. Y.: 1991)* 25, 918–926. <https://doi.org/10.1093/cercor/bht280>.
- Buzsáki, G., 2010. Neural syntax: cell assemblies, synapses, and readers. *Neuron* 68, 362–385. <https://doi.org/10.1016/j.neuron.2010.09.023>.
- Bylinskii, Z., Isola, P., Bainbridge, C., Torralba, A., Oliva, A., 2015. Intrinsic and extrinsic effects on image memorability. *Vis. Res* 116, 165–178. <https://doi.org/10.1016/j.visres.2015.03.005>.
- Damiano, C., Walther, D.B., 2019. Distinct roles of eye movements during memory encoding and retrieval. *Cognition* 184, 119–129. <https://doi.org/10.1016/j.cognition.2018.12.014>.
- Dimigen, O., Sommer, W., Hohlfeld, A., Jacobs, A.M., Kliegl, R., 2011. Coregistration of eye movements and EEG in natural reading: analyses and review. *J. Exp. Psychol. Gen.* 140, 552–572. <https://doi.org/10.1037/a0023885>.
- Donoghue, T., et al., 2020a. Parameterizing neural power spectra into periodic and aperiodic components. *Nat. Neurosci.* 23, 1655–1665. <https://doi.org/10.1038/s41593-020-00744-x>.
- Donoghue, T., et al., 2020b. Parameterizing neural power spectra into periodic and aperiodic components. *Nat. Neurosci.* 23, 1655–1665. <https://doi.org/10.1038/s41593-020-00744-x>.
- Doucet, G., Gulli, R.A., Corrigan, B.W., Duong, L.R., Martinez-Trujillo, J.C., 2020. Modulation of local field potentials and neuronal activity in primate hippocampus during saccades. *Hippocampus* 30, 192–209. <https://doi.org/10.1002/hipo.23140>.
- Engbert, R., Mergenthaler, K., 2006. Microsaccades are triggered by low retinal image slip. *Proc. Natl. Acad. Sci. USA* 103, 7192–7197. <https://doi.org/10.1073/pnas.0509557103>.
- Fehlmann, B., et al., 2020. Visual Exploration at Higher Fixation Frequency Increases Subsequent Memory Recall. *Cereb. cortex Commun.* 1, tgaa032. <https://doi.org/10.1093/textcom/tgaa032>.
- Fellner, M.C., et al., 2019. Spectral fingerprints or spectral tilt? Evidence for distinct oscillatory signatures of memory formation. *PLoS Biol.* 17, e3000403 <https://doi.org/10.1371/journal.pbio.3000403>.
- Fellner, M.C., Bäuml, K.H., Hanslmayr, S., 2013. Brain oscillatory subsequent memory effects differ in power and long-range synchronization between semantic and survival processing. *NeuroImage* 79, 361–370. <https://doi.org/10.1016/j.neuroimage.2013.04.121>.
- Fries, P., 2015. Rhythms for cognition: communication through coherence. *Neuron* 88, 220–235. <https://doi.org/10.1016/j.neuron.2015.09.034>.
- Gbadamosi, J., Zangemeister, W.H., 2001. Visual imagery in hemianopic patients. *J. Cogn. Neurosci.* 13, 855–866. <https://doi.org/10.1162/089892901753165782>.
- Griffiths, B.J., et al., 2019. Alpha/beta power decreases track the fidelity of stimulus-specific information. *Elife* 8. <https://doi.org/10.7554/eLife.49562>.
- Griffiths, B.J., Martín-Buro, M.C., Staresina, B.P., Hanslmayr, S., 2021a. Disentangling neocortical alpha/beta and hippocampal theta/gamma oscillations in human episodic memory formation. *NeuroImage* 242, 118454. <https://doi.org/10.1016/j.neuroimage.2021.118454>.
- Griffiths, B.J., Martín-Buro, M.C., Staresina, B.P., Hanslmayr, S., 2021a. Disentangling neocortical alpha/beta and hippocampal theta/gamma oscillations in human episodic memory formation. *NeuroImage* 242, 118454. <https://doi.org/10.1016/j.neuroimage.2021.118454>.
- Griffiths, B.J., Martín-Buro, M.C., Staresina, B.P., Hanslmayr, S., Staudigl, T., 2021b. Alpha/beta power decreases during episodic memory formation predict the magnitude of alpha/beta power decreases during subsequent retrieval. *Neuropsychologia* 153, 107755. <https://doi.org/10.1016/j.neuropsychologia.2021.107755>.
- Gross, J., et al., 2001. Dynamic imaging of coherent sources: Studying neural interactions in the human brain. *Proc. Natl. Acad. Sci. USA* 98, 694–699. <https://doi.org/10.1073/pnas.98.2.694>.
- Hamamé, C.M., et al., 2014. Functional selectivity in the human occipitotemporal cortex during natural vision: evidence from combined intracranial EEG and eye-tracking. *NeuroImage* 95, 276–286. <https://doi.org/10.1016/j.neuroimage.2014.03.025>.
- Hanslmayr, S., et al., 2011. The relationship between brain oscillations and BOLD signal during memory formation: a combined EEG-fMRI study. *J. Neurosci.* 31, 15674–15680. <https://doi.org/10.1523/JNEUROSCI.3140-11.2011>.
- Hanslmayr, S., Staudigl, T., 2014. How brain oscillations form memories—a processing based perspective on oscillatory subsequent memory effects. *NeuroImage* 85 (Pt 2), 648–655. <https://doi.org/10.1016/j.neuroimage.2013.05.121>.
- Hanslmayr, S., Spitzer, B., Bäuml, K.H., 2008. Brain oscillations dissociate between semantic and nonsemantic encoding of episodic memories. *Cereb. Cortex* 19, 1631–1640. <https://doi.org/10.1093/cercor/bhn197>.
- Hanslmayr, S., Staudigl, T., Fellner, M.C., 2012. Oscillatory power decreases and long-term memory: the information via desynchronization hypothesis. *Front Hum. Neurosci.* 6, 74. <https://doi.org/10.3389/fnhum.2012.00074>.
- Hanslmayr, S., Staresina, B.P., Bowman, H., 2016. Oscillations and Episodic Memory: Addressing the Synchronization/Desynchronization Conundrum. *Trends Neurosci.* 39, 16–25. <https://doi.org/10.1016/j.tins.2015.11.004>.
- Hoffman, K.L., et al., 2013. Saccades during visual exploration align hippocampal 3–8 Hz rhythms in human and non-human primates. *Front. Syst. Neurosci.* 7, 43. <https://doi.org/10.3389/fnsys.2013.00043>.
- Ito, J., Maldonado, P., Singer, W., Grün, S., 2011. Saccade-related modulations of neuronal excitability support synchrony of visually elicited spikes. *Cereb. cortex (N. Y., N. Y.: 1991)* 21, 2482–2497. <https://doi.org/10.1093/cercor/bhr020>.
- Johansson, R., Holsanova, J., Holmqvist, K., 2006. Pictures and spoken descriptions elicit similar eye movements during mental imagery, both in light and in complete darkness. *Cogn. Sci.* 30, 1053–1079. https://doi.org/10.1207/s15516709cog0000_86.
- Johansson, R., Nystrom, M., Dewhurst, R., Johansson, M., 2022. Eye-movement replay supports episodic remembering. *Proc. Biol. Sci.* 289, 20220964. <https://doi.org/10.1098/rspb.2022.0964>.
- Jutras, M.J., Fries, P., Buffalo, E.A., 2013. Oscillatory activity in the monkey hippocampus during visual exploration and memory formation. *Proc. Natl. Acad. Sci. USA* 110, 13144–13149. <https://doi.org/10.1073/pnas.1302351110>.
- Katz, C.N., et al., 2022. A corollary discharge mediates saccade-related inhibition of single units in mnemonic structures of the human brain. *Curr. Biol.* 32 (3082–3094), e3084 <https://doi.org/10.1016/j.cub.2022.06.015>.
- Khader, P.H., Jost, K., Ranganath, C., Rosler, F., 2010. Theta and alpha oscillations during working-memory maintenance predict successful long-term memory encoding. *Neurosci. Lett.* 468, 339–343. <https://doi.org/10.1016/j.neulet.2009.11.028>.
- Klimesch, W., et al., 1996. Event-related desynchronization (ERD) and the Dm effect: does alpha desynchronization during encoding predict later recall performance. *Int. J. Psychophysiol.* 24, 47–60. [https://doi.org/10.1016/s0167-8760\(96\)00054-2](https://doi.org/10.1016/s0167-8760(96)00054-2).
- Kragel, J.E., Voss, J.L., 2022. Looking for the neural basis of memory. *Trends Cogn. Sci.* 26, 53–65. <https://doi.org/10.1016/j.tics.2021.10.010>.
- Laeng, B., Teodorescu, D.-S., 2002. Eye scanpaths during visual imagery reenact those of perception of the same visual scene. *Cogn. Sci.* 26, 207–231. https://doi.org/10.1207/s15516709cog2602_3.
- Leszczynski, M., et al., 2021. Neural activity in the human anterior thalamus during natural vision. *Sci. Rep.* 11, 17480. <https://doi.org/10.1038/s41598-021-96588-x>.
- Liu, B., Nobre, A.C., van Ede, F., 2022a. Microsaccades transiently lateralise EEG alpha activity, 2022.09.02.506318 bioRxiv. <https://doi.org/10.1101/2022.09.02.506318>.
- Liu, B., Nobre, A.C., van Ede, F., 2022b. Functional but not obligatory link between microsaccades and neural modulation by covert spatial attention. *Nat. Commun.* 13, 3503. <https://doi.org/10.1038/s41467-022-31217-3>.
- Liu, Z.X., Shen, K., Olsen, R.K., Ryan, J.D., 2017. Visual sampling predicts hippocampal activity. *J. Neurosci.* 37, 599–609. <https://doi.org/10.1523/JNEUROSCI.2610-16.2016>.
- Mao, D., et al., 2021. Spatial modulation of hippocampal activity in freely moving macaques. *Neuron* 109 (3521–3534), e3526. <https://doi.org/10.1016/j.neuron.2021.09.032>.
- Maris, E., Oostenveld, R., 2007. Nonparametric statistical testing of EEG- and MEG-data. *J. Neurosci. Methods* 164, 177–190. <https://doi.org/10.1016/j.jneumeth.2007.03.024>.
- Martinez-Conde, S., Otero-Millan, J., Macknik, S.L., 2013. The impact of microsaccades on vision: towards a unified theory of saccadic function. *Nat. Rev. Neurosci.* 14, 83–96. <https://doi.org/10.1038/nrn3405>.
- McFarland, J.M., Bondy, A.G., Saunders, R.C., Cumming, B.G., Butts, D.A., 2015. Saccadic modulation of stimulus processing in primary visual cortex. *Nat. Commun.* 6, 8110. <https://doi.org/10.1038/ncomms9110>.
- Mitra, P.P., Pesaran, B., 1999. Analysis of Dynamic Brain Imaging Data. *Biophys. J.* 76, 691–708. [https://doi.org/10.1016/S0006-3495\(99\)77236-X](https://doi.org/10.1016/S0006-3495(99)77236-X).
- Molitor, R.J., Ko, P.C., Hussey, E.P., Ally, B.A., 2014. Memory-related eye movements challenge behavioral measures of pattern completion and pattern separation. *Hippocampus* 24, 666–672. <https://doi.org/10.1002/hipo.22256>.
- Olsen, R.K., et al., 2016. The relationship between eye movements and subsequent recognition: Evidence from individual differences and amnesia. *Cortex; J. Devoted Study Nerv. Syst. Behav.* 85, 182–193. <https://doi.org/10.1016/j.cortex.2016.10.007>.

- Oostenveld, R., Fries, P., Maris, E., Schoffelen, J.M., 2011. FieldTrip: Open source software for advanced analysis of MEG, EEG, and invasive electrophysiological data. *Comput. Intell. Neurosci.* 2011, 156869 <https://doi.org/10.1155/2011/156869>.
- Otero-Millan, J., Troncoso, X.G., Macknik, S.L., Serrano-Pedraza, I., Martinez-Conde, S., 2008. Saccades and microsaccades during visual fixation, exploration, and search: foundations for a common saccadic generator, 21-18 J. Vis. 8 (21). <https://doi.org/10.1167/8.14.21>.
- Paller, K.A., Wagner, A.D., 2002. Observing the transformation of experience into memory. *Trends Cogn. Sci.* 6, 93–102. [https://doi.org/10.1016/s1364-6613\(00\)01845-3](https://doi.org/10.1016/s1364-6613(00)01845-3).
- Parish, G., Hanslmayr, S., Bowman, H., 2018. The Sync/deSync Model: How a Synchronized Hippocampus and a Desynchronized Neocortex Code Memories. *J. Neurosci.* 38, 3428–3440. <https://doi.org/10.1523/jneurosci.2561-17.2018>.
- Popov, T., Miller, G.A., Rockstroh, B., Jensen, O., Langer, N., 2021a. Alpha oscillations link action to cognition: An oculomotor account of the brain's dominant rhythm. *bioRxiv*. <https://doi.org/10.1101/2021.09.24.461634>.
- Popov, T., Langer, N., Gips, B., Weisz, N., Jensen, O., 2021b. Sound-location specific alpha power modulation in the visual cortex in absence of visual input. *bioRxiv*. <https://doi.org/10.1101/2021.03.15.435371>.
- Popov, T., Gips, B., Weisz, N., Jensen, O., 2022. Brain areas associated with visual spatial attention display topographic organization during auditory spatial attention. *Cereb. Cortex* (N. Y., N. Y.: 1991). <https://doi.org/10.1093/cercor/bhac285>.
- Printzlau, F.A.B., Myers, N.E., Manohar, S.G., Stokes, M.G., 2022. Neural Reinstatement Tracks Spread of Attention between Object Features in Working Memory. *J. Cogn. Neurosci.* 1–21. https://doi.org/10.1162/jocn_a_01879.
- Purpura, K.P., Kalik, S.F., Schiff, N.D., 2003. Analysis of perisaccadic field potentials in the occipitotemporal pathway during active vision. *J. Neurophysiol.* 90, 3455–3478. <https://doi.org/10.1152/jn.00011.2003>.
- Rajkai, C., et al., 2008. Transient cortical excitation at the onset of visual fixation. In: *Cerebral cortex* (New York, N.Y.: 1991), 18, pp. 200–209. <https://doi.org/10.1093/cercor/bhm046>.
- Sander, M.C., Fandakova, Y., Grandy, T.H., Shing, Y.L., Werkle-Bergner, M., 2020. Oscillatory Mechanisms of Successful Memory Formation in Younger and Older Adults Are Related to Structural Integrity. *Cereb. Cortex* 30, 3744–3758. <https://doi.org/10.1093/cercor/bhz339>.
- Schneider, M., et al., 2021. A mechanism for inter-areal coherence through communication based on connectivity and oscillatory power. *Neuron* 109 (4050–4067), e4012. <https://doi.org/10.1016/j.neuron.2021.09.037>.
- Sederberg, P.B., Kahana, M.J., Howard, M.W., Donner, E.J., Madsen, J.R., 2003. Theta and gamma oscillations during encoding predict subsequent recall. *J. Neurosci.* 23, 10809–10814.
- Sommer, M.A., Wurtz, R.H., 2002. A pathway in primate brain for internal monitoring of movements. *Sci. (N. Y., N. Y.)* 296, 1480–1482. <https://doi.org/10.1126/science.1069590>.
- Staudigl, T., Hartl, E., Noachtar, S., Doeller, C.F., Jensen, O., 2017a. Saccades are phase-locked to alpha oscillations in the occipital and medial temporal lobe during successful memory encoding. *PLoS Biol.* 15, e2003404.
- Staudigl, T., Hartl, E., Noachtar, S., Doeller, C.F., Jensen, O., 2017b. Saccades are phase-locked to alpha oscillations in the occipital and medial temporal lobe during successful memory encoding. *PLoS Biol.* 15, e2003404 <https://doi.org/10.1371/journal.pbio.2003404>.
- Staudigl, T., Minxha, J., Mamelak, A.N., Gothard, K.M., Rutishauser, U., 2022. Saccade-related neural communication in the human medial temporal lobe is modulated by the social relevance of stimuli. *Sci. Adv.* 8, eabl6037 <https://doi.org/10.1126/sciadv.abl6037>.
- Stolk, A., Todorovic, A., Schoffelen, J.M., Oostenveld, R., 2013. Online and offline tools for head movement compensation in MEG. *NeuroImage* 68, 39–48. <https://doi.org/10.1016/j.neuroimage.2012.11.047>.
- Strunk, J., Duarte, A., 2019. Prestimulus and poststimulus oscillatory activity predicts successful episodic encoding for both young and older adults. *Neurobiol. Aging* 77, 1–12. <https://doi.org/10.1016/j.neurobiolaging.2019.01.005>.
- Vogelsang, D.A., Gruber, M., Bergström, Z.M., Ranganath, C., Simons, J.S., 2018. Alpha Oscillations during Incidental Encoding Predict Subsequent Memory for New “Foil” Information. *J. Cogn. Neurosci.* 30, 667–679. https://doi.org/10.1162/jocn_a_01234.
- Voloh, B., Womelsdorf, T., 2016. A Role of Phase-Resetting in Coordinating Large Scale Neural Networks During Attention and Goal-Directed Behavior. *Front. Syst. Neurosci.* 10, 18. <https://doi.org/10.3389/fnsys.2016.00018>.
- Voss, J.L., Bridge, D.J., Cohen, N.J., Walker, J.A., 2017. A Closer Look at the Hippocampus and Memory. *Trends Cogn. Sci.* 21, 577–588. <https://doi.org/10.1016/j.tics.2017.05.008>.
- Wagner, I.C., Jensen, O., Doeller, C.F., Staudigl, T., 2022. Saccades are coordinated with directed circuit dynamics and stable but distinct hippocampal patterns that promote memory formation, 2022.2008.2018.504386 *bioRxiv*. <https://doi.org/10.1101/2022.08.18.504386>.
- Wynn, J.S., Shen, K., Ryan, J.D., 2019. Eye movements actively reinstate spatiotemporal mnemonic content. *Vis. (Basel)* 3 doi:10.3390/vision3020021.
- Wynn, J.S., Ryan, J.D., Buchsbaum, B.R., 2020. Eye movements support behavioral pattern completion. *Proc. Natl. Acad. Sci. USA* 117, 6246–6254. <https://doi.org/10.1073/pnas.1917586117>.