Check for updates

scientific reports

REGISTERED REPORT

OPEN Variations of autonomic arousal mediate the reportability of mind blanking occurrences

Paradeisios Alexandros Boulakis^{1,2}, Nicholas John Simos¹, Stefania Zoi¹, Sepehr Mortaheb^{1,2}, Christina Schmidt^{2,3}, Federico Raimondo^{4,5,7} & Athena Demertzi^{1,2,6,7}

Mind blanking (MB) is the inability to report mental events during unconstraint thinking. Previous work shows that MB is linked to decreased levels of cortical arousal, indicating dominance of cerebral mechanisms when reporting mental states. What remains inconclusive is whether MB can also ensue from autonomic arousal manipulations, pointing to the implication of peripheral physiology to mental events. Using experience sampling, neural, and physiological measurements in 26 participants, we first show that MB was reported more frequently in low arousal conditions, elicited by sleep deprivation. Also, there was partial evidence for a higher occurence of MB reports in high arousal conditions, elicited by intense physical exercise. Transition probabilities revealed that, after sleep deprivation, mind wandering was more likely to be followed by MB and less likely to be followed by more mind wandering reports. Using classification schemes, we found higher performance of a balanced random forest classifier trained on both neural and physiological markers in comparison to performance when solely neural or physiological were used. Collectively, we show that both cortical and autonomic arousal affect MB report occurrences. Our results establish that MB is supported by combined brainbody configurations, and, by linking mental and physiological states, they pave the way for novel embodied accounts of spontaneous thinking.

'The stage 1 protocol for this Registered Report was accepted in principle on 02/01/23. The protocol, as accepted by the journal, can be found at: 10.17605/OSF.IO/SH2YE' Techniques: Life sciences techniques, Biophysical methods [Electrocardiography - EKG]; Life sciences techniques, Biophysical methods [Electroencephalography - EEG]; CTS received date: 27.11.2024.

Keywords Mind blanking, Experience sampling, Brain-body interactions, Machine learning, Spontaneous thinking

During ongoing mentation, our mind constantly shifts across different mental states. These mental states typically bear some content ("what we think about") and indicate a relationship towards that content (i.e., perceiving, fearing, hoping, remembering)¹. As we move through the environment, our thoughts fluctuate between the external and internal milieu^{2,3}, resulting in a fluid stream of consciousness⁴. External content is tightly coupled to the processing of environmental stimuli and task-demanding conditions. Internal content is more associated with self-referential processing and internal dialogue, widely referred to as "mind wandering" (MW)⁴. Inclusive as this external-internal dipole may seem, it does not capture the full scope of the "aboutness" of mental content. Recent work has highlighted another mental state, where people report that they are "thinking of nothing" or "their mind just went away", a phenomenological experience termed "mind blanking" (MB)⁵. As MB is relatively new in the landscape of ongoing cognition, the extent of MB episodes in daily and clinical settings remains widely uncharacterized. For example, a recent study found that MB might be miscategorized as MW in ADHD symptom evaluation⁶. Therefore, the experience of MB occurrences poses a challenge to our everyday functioning and our understanding of the continuous nature of the stream of consciousness.

¹Physiology of Cognition Lab, GIGA-CRC Human Imaging Unit, GIGA Research, University of Liège, Liège, Belgium. ²Fund for Scientific Research FNRS, Brussels, Belgium. ³Sleep & Chronobiology Lab, GIGA-CRC Human Imaging Unit, GIGA Research, University of Liège, Liège, Belgium. ⁴Institute of Neuroscience and Medicine, Research Centre Jülich, Brain & Behaviour (INM-7), Jülich, Germany. ⁵Institute of Systems Neuroscience, Medical Faculty, Heinrich Heine University Düsseldorf, Düsseldorf, Germany. ⁶Psychology and Neuroscience of Cognition Research Unit, University of Liège, Liège, Belgium. ⁷Federico Raimondo and Athena Demertzi contributed equally to this work. [™]email: a.demertzi@uliege.be

Currently, there is no clear answer as to how MB reports are generated. So far, behavioral studies show that MB can arise after conscious mental effort to empty our mind⁷⁻⁹, is usually unintentional^{5,10,11} and gets reported less frequently during unconstrained thinking compared to MW and sensory/perceptual mental states^{5,11-13}. At the brain level, the inability to report mental events after the prompt to "empty the mind" has been associated with activation of the anterior cingulate/medial prefrontal cortex, and deactivation of inferior frontal gyrus/ Broca's areas and the hippocampus, which the authors interpreted as the inability to verbalize internal mentation (inner speech)⁸. Recently, we found that the functional connectome of fMRI volumes around MB reports was similar to a unique brain pattern of overall positive inter-areal connectivity¹² which was also characterized by increased amplitude of fMRI global signal (i.e. averaged connectivity across all grey matter voxels), an implicit indicator of low arousal¹⁴⁻¹⁶. For example, the amplitude of the global signal correlated negatively with EEG vigilance markers (alpha, beta EEG frequency bands), while increases in EEG vigilance due to caffeine ingestion were associated with reduced global signal amplitude¹⁴. Our findings corroborate recent EEG-related evidence supporting the possibility of "local sleeps" during MB reportability^{10,17}. "Local sleeps" refer to the scalp distribution of EEG potentials during wakefulness, in the form of high-intensity, slow oscillatory activity in the theta/delta band, which could differentiate between MB and MW, with more frontocentral potentials tied to MW and parietal to MB¹⁰. Together, the presence of slow waves preceding MB reports and the high fMRI global signal hint toward the role of arousal in mental content reportability. Starting from this line of evidence, we infer that arousal fluctuations drive MB reportability.

Arousal is a multidimensional construct generally referring to the behavioral state of being awake and alert, supporting wakefulness, responsiveness to environmental stimuli, and attentiveness^{18,19}. Anatomically, arousal is supported by the ascending arousal system, the autonomic nervous system, and the endocrine system¹⁸. Early on, Lacey viewed arousal in terms of behavioral arousal (indicated by a responding organism, like restlessness and crying), cortical arousal (evidenced by desynchronized fast oscillatory activity), and autonomic arousal (indicated by changes in bodily functions)²⁰. Cortical arousal is self-generated through the reticulate formation and propagated through dorsal, thalamic, and ventral subthalamic pathways²¹, and can be indexed by the alpha, theta, and delta EEG bands during wakefulness^{22,23}. Lower levels of cortical arousal in the form of slow waves have been associated with an increased number of missed stimuli in behavioral tasks^{11,23} and decreased thought intensity²⁴. Also, lower levels of arousal indexed by pupil size have been correlated with a higher probability of MB reports in sustained attention tasks^{11,25,26}.

Much as it may have been done in terms of cortical arousal, the present study will focus on how autonomic arousal influences MB reportability, which is widely understudied. Our choice is justified by the theoretical assumption that mental function is tightly linked to peripheral body functions, as expressed by the embodied cognition stance²⁷. Briefly, embodiment holds that cognition is bound to a living body interacting with a dynamic environment, and conceptualizes cognition as the result of brain-body interactions during dynamic contexts. From that perspective, modifications in autonomic arousal are expected to lead to differential reportability of mental states. Autonomic arousal links the body and the brain through spinal cord projections from peripheral organs to the brainstem and can be indexed by physiological signals reflecting sympathetic/parasympathetic balance, such as heart rate, galvanic skin response, and fluctuations in pupil size²⁸. Converging evidence suggests that afferent physiological signals and biological rhythms, such as the cardiac or the respiratory phase, play a modulatory role in conscious perception^{29,30}, metacognition³¹, affective salience of information³², and perceptual confidence of sensory sampling³³, both during task performance and in-silico simulations³⁴. Alterations in autonomic arousal were also found to influence brain activity in that fMRI volumes characterized by lower arousal levels (indexed by decreased pupil size) showed reduced in-between network integration and inter-subject variability in comparison to scans characterized by high arousal levels (indexed by increased pupil size)³⁵.

Taken together, we here advocated for a link between autonomic arousal and thought reportability. Firstly, we examined how MB report distribution shifted across different autonomic arousal conditions. To this end, we used experience sampling under differently elicited arousal conditions. Experience sampling is a though-sampling methodology, where people are probed to report their mental state at random intervals, probed by an external cue⁴. We employed this task at three distinct arousal conditions: *Baseline, High* (post-workout), and *Low* (post-sleep deprivation). Our operational hypothesis was that optimal levels of autonomic arousal (fixed variable) are necessary for optimal mental state reportability (dependent variable). We expected that deviations from optimal levels, such as after sleep deprivation or intense physical exercise, would alter our stream of thought and promote more frequent MB reports (Supplementary Table S1 for the full scope of our hypotheses). Secondly, we opted to identify specific brain-body interaction patterns that would promote MB reportability. To this end, we utilized multimodal neurophysiological recordings and a machine-learning approach to decode MB reports from arousal measurements.

Methods Design

The study included healthy volunteers recruited after campus poster advertisements, intranet electronic invitations, and through the ULiège "petites annonces" e-campus platform. Inclusion criteria were: (a) right-handedness, (b) age>18 years, (c) minimal exercise background (<2h per week), (d) good subjective sleep quality (Pittsburgh Sleep Quality Index [PSQI] $\leq 5^{36}$), (e) habitual sleep duration of 8 ± 1 hours. Exclusion criteria were: (a) history of developmental, psychiatric, or neurological illness resulting in documented functional disability, (b) severe anomalies in pupil shape or inability to open both eyes preventing pupil measurement³⁷, (c) analgesic medication which may affect physiological arousal, (d) history of psychiatric illness pertaining to anxiety disorders or scores < 9 in the General Anxiety Disorder-7 (GAD-7 scale)³⁸ as anxious participants may

experience biased perceptions of their bodily states³⁹, (e) extreme chronotypes, (f) shift work or traveling over time zones in the past 3 months.

Experience sampling was utilized in a within-participants repeated-measures design. During an experience sampling session, participants laid restfully and were directed to let their minds wander, without any specific instructions towards internal (daydreaming, memories, prospective events) or external thoughts (body sensations, sensory stimuli in their immediate environment). Auditory probes (total n=40, 500 Hz simple tones) invited participants to report what they were thinking at the moment just preceding the probe. The inter-probe interval was sampled from a uniform distribution between 110 and 120 seconds. Report times were monitored online to examine if participants missed the probe or fell asleep due to our experimental manipulation. In case of a report time > 6 s, participants were reminded to report their mental state as soon as they heard the probe and indicate they were awake via button press. In case of unresponsiveness, the experimenters manually awakened the participant. Depending on the probes' trigger times and participants' reaction times, a recording lasted on average 70-90 minutes. We chose to present 40 probes (overall length approximately 1 hour and 15 minutes) to avoid fatigue/drowsiness and the possibility of participants returning to baseline arousal after the experimental manipulations. Also, the relatively large experience sampling interval, compared to previous studies, was used to record enough samples to accurately estimate physiological markers from slow oscillatory signals, such as heart-rate variability. Upon the probe, participants had to choose among four distinct choices describing their mental state: mind blanking (MB), mind wandering (MW), perceptual sensations (SENS), or sleep (SLEEP). These response options were chosen to minimize assumptions about what the actual partition of mental states might be. For example, debates about what can be classified as MW⁴⁰ refer to whether MW is a coherent cluster of events^{1,41} and how it is separated from awareness and processing of environmental stimuli^{40,42}. We believe that our division respects the literature on internal/external thought-orientation brain networks^{3,43,44} while introducing minimum assumptions as to the actual content of each state. The introduction of the sleep option facilitated the identification of trials where participants fell asleep due to the reduced vigilance. Participants indicated their responses via button press from a response keyboard placed under their dominant hand. We repeated the experience sampling task on three distinct days, over the span of two weeks under three conditions: (a) experience sampling during spontaneous thinking without arousal modulations (Baseline), (b) experience sampling elicited through short, high-intensity interval training (High Arousal), (c) experience sampling after total sleep deprivation (Low Arousal) (Fig. 1). The goal of both arousal manipulations was to promote distinct changes in physiological and cortical markers associated with arousal mechanisms (Supplementary Table S2). Monitoring of arousal changes was done with physiological and cortical measurements. In case when participants did not show distinct cortical and physiological changes after our arousal manipulations, they were excluded



Fig. 1. Experimental protocol. *Top* The experience sampling task invited participants to sit idly and relax, letting their minds wander. Every 110–120 s, a 500 Hz auditory cue probed participants to report what they were thinking at that moment. Participants were able to choose from 4 presented responses: Mind blanking (MB), Mind wandering (MW), Perceptual Sensations (SENS), and Sleep (SLEEP). *Bottom* Repeated-measures autonomic arousal recordings. To test how spontaneous thoughts unfold over time across different arousal conditions, we first invited people for baseline assessments on Day 1 (*Baseline* condition). On Day 2 participants underwent a 15-minute high-intensity exercise (*High Arousal* condition) and on Day 3 they participated in a total sleep deprivation protocol (*Low Arousal* condition). The *High* and *Low Arousal* conditions were counter-balanced across participants. Multimodal physiological recordings were used to monitor arousal manipulations. The dataset was constituted of EEG, pupillometry, ECG, EDA, and respiratory data; the arrows indicate the hypothesized directions of the derived metrics.

from further analysis. Effect monitoring was done by examining the heart rate in *High Arousal* as well as the EEG spectra in both *High* and *Low Arousal*.

In *High Arousal*, participants first performed high-intensity interval activity in the form of static cycling. They started with a warm-up training session of 3 minutes to avoid potential muscle trauma and then cycled for 45 s as fast as possible. A resting period of 15 s followed. A total number of 10 workout cycles was administered. The choice of this timing protocol rested on previous studies indicating that similar exercise routines produce distinct and sustained sympathetic activity^{45,46} and cortical excitation⁴⁶, which can last between 30 and 90 minutes after exercise cessation⁴⁷.

In Low Arousal, participants performed the experience sampling task after one night of total sleep deprivation. Sleep deprivation leads to an arousal state that is behaviourally distinct from typical wakefulness^{48,49}, promotes specific neuronal signatures ("local sleeps" in the delta band)¹¹, and has a distinct physiological expression. Critically, we do not wish to claim that sleep states are identical to "local sleeps", nor do we suggest an overlap between low arousal due to sleep deprivation and unconsciousness during sleep. To acquire estimates of their mean sleep schedule, participants wore an actimeter for one week before the total sleep deprivation protocol (Supplementary Fig. S1; available for 24/26 subjects due to data corruption). The total sleep deprivation protocol was as follows: A week prior to sleep deprivation, participants were provided with an actimetry device to track wake-sleep schedule, and were instructed to follow a consistent 8 h sleep schedule. On the deprivation day, participants arrived at the lab one hour before their normal sleep time to extract their actimetry baseline data, estimate the optimal sleep deprivation window, and to provide baseline vigilance, drowsiness, and sleepiness measurements. After a total sleep deprivation of 26 h (16 h of typical wakefulness, 8 h of sleep deprivation, and a 2h post-sleep deprivation period) participants began the post-sleep deprivation, experience sampling session. As an example, a participant who typically slept at 12 am would arrive at the lab at 11 pm, start sleep deprivation at 12 am, finish sleep deprivation at 8 am, and perform the experience sampling task at 10 am. Should slow-wave activity during wakefulness follow the same circadian modulation it follows during sleep⁵⁰, a potential confound that could have lowered the power of our analysis is the time window of the experience sampling task. However, as suggested in⁵⁰, the relative time-window we selected did not fall under a critical point of large reductions in the amplitude of the slow-waves. The 2-hour, post-deprivation waiting window allowed us to match the time of the experience sampling across the 3 conditions, avoiding potential circadian confounds on experience sampling, as we could easier match sleep-wake cycles and the time of the experience sampling within each participant. We chose this sleep manipulation as similar manipulations have been previously used to examine the effects of sleep pressure^{51,52}, and have been shown to elicit distinct low-arousal cortical profiles^{53,54}, as well as changes in the sympathetic/parasympathetic balance⁵⁵.

Sleep deprivation was controlled with regard to light influence (illuminance = 15 lux), caloric intake (standardized meals every 4 h), and body posture (semi-recumbent position during scheduled wakefulness) to minimize potential masking effects on the sleep-wake regulatory system. Participants were not allowed to stand up except for regularly scheduled bathroom visits and did not have any indications of the time of the day. The experimenters continually monitored participants to keep them awake. In case of a sleep event, the experimenters first tried to awaken the participant through an intercom, and in case of failure, they manually awakened the participant. We also monitored for sleep lapses through the experience sampling tasks. In case participants closed their eyes for a time period of > 30 seconds, they were probed by a tone to wake up. If they did not, the experimenter in the room would awaken the participant.

An one-week interval took place between sleep deprivation and further recordings in order to minimize potential carry-over effects of sleep deprivation on our follow-up conditions. In that way, the participants' sleep schedules would also reset to their respective normal cycles. The order of the three arousal conditions was randomized. As a post-registration note, we randomized only the order between sleep deprivation and post-exercise, to add a training session before the baseline that allowed participants to get acquainted with the protocol, without external task impositions, that might have confounded the protocol understanding.

Sampling plan

We used a Neyman-Pearson frequentist approach to balance false-negative and false-positive rates by setting power to 95% and establishing a Type I error rate (alpha) of 5%. To estimate the desired sample size, a simulation approach was utilized: data were generated consistent with a latent binomial regression model, in which one categorical predictor with 3 levels (Base, High, Low) predicted a binary outcome Y (occurrence of MB or not). An original probability $p_{MB} = 0.1$ was specified as the underlying generative probability in the baseline model based on previous research^{5,11,12}. We allowed the random intercepts and slopes to vary freely vary around a normal distribution with a standard deviation of s.d. = 0.1. Given that no previous study to our knowledge has provided evidence for the distribution of the effect sizes of arousal on mental reports, and to account for possible reverse effects (such as decreased MB report probability), we reasoned that a meaningful yet conservative effect for the *Low Arousal* condition would be an odds ratio of 1.6 and an odds ratio of 0.55 for the *High Arousal* condition. Since our initial hypothesized distribution is expected to yield ~3–5 MB reports per session^{11,12}, this effectively translates to a small effect size of interest of at least 3 more reports across conditions.

Considering these parameters, for each population sample, ranging from 5 to 50 participants, we sampled 500 datasets, and fit a binomial model with the participant ID as a random factor, keeping the regression coefficients for the levels of the predictor constant. Based on the simulation analysis, using a false positive threshold of 0.05, we required a sample size of 26 participants to achieve a power of 0.95 (Supplementary Fig. S2).

Data analysis Behavioral data

Statistical analysis was performed using generalized linear mixed-effects models. To address whether arousal affects MB occurrence, we used a binomial, linear model with arousal as a categorical independent variable, and the proportion of mental reports across a sampling period (40 trials) as our dependent variable. Data were binary coded (occurence or not of MB report) and fit into the model using a "logit" link. Given that the underlying distribution was unknown, a Bernoulli distribution minimized the assumptions about the model. In order to examine whether the multinomial distribution of mental reports itself changes across different arousal conditions, we used the generalized estimating equations (GEE) approach, an extension of generalized mixedeffects models that can account for correlated, repeated-measures count data from multinomial distributions^{56,57}. Mental reports were aggregated as counts across participants and conditions, and we examined shifts in reaction time distribution using the three experimental arousal conditions as predictors. We considered as reaction times the intervals between the response probe and the participant's report. To examine reaction times as a function of mental states, we specified a generalized linear mixed-effect model with mental reports and arousal conditions as categorical variables and used a gamma distribution with an "inverse" link function. As reaction times are usually an indicator of arousal effects on the task performance, an effect of arousal condition as a covariate might be informative about a potential shift of the overall slower mental report times distribution and about the arousal condition of the subject itself. This choice of distribution and link minimizes assumptions about the model, respects the positive, skewed distribution of reaction times, and was previously found to provide a better fit compared to other link functions⁵⁸. To examine whether arousal shifts the dynamics of mental reports, i.e. one state might be more likely to be followed by MB in one of the arousal states compared to Baseline, we estimated dynamical transition probabilities across different mental states using Markov models. The transition probabilities of MB were then compared using a linear model with an identity link, with the transition probabilities as the dependent variable and the arousal condition as the categorical, independent variable.

All specified models were compared against null models using likelihood ratio tests. We introduced the participant's ID as an a-priori random factor, i.e., we allowed the model's intercept to vary. In case we contrasted multiple models, *p*-values were corrected using Bonferroni correction. In case of significance of a fixed predictor, we used corrected pairwise comparisons to examine the marginal means of the predictors.

Brain-based measures

Physiological and cortical timeseries were segmented based on the response probe time. We considered the 110-second period before the response probe as a meaningful analysis epoch, representing the neuronal and physiological dynamics that result in a specific mental state. This period was used in subsequent analyses.

We recorded EEG with an EasyCap (64 active electrodes) connected to an actiCHamp system (Brain Products GmbH) using the 10–20 standard configuration. A ground electrode was placed frontally (Fpz in the 10–20 system). Online, we referenced the electrodes to a frontal electrode. Impedance was kept below 20 k Ω . As a post-registration note, we originally registered to keep impedance below 10 k Ω . However, we decided to leverage the strength of active electrodes to follow the research standard of 20 k Ω . To minimize impedance, we used conductive gel. Data were sampled at a sampling frequency of 500 Hz. Preprocessing included band-pass filtering (0.1–45 Hz, FIR filter), notch filtering (50 Hz), and epoch definition (t_start = 110 s preceding the probe, t_max= probe). As a post-registration note, during EEG preprocessing we observed low-frequency (<1 Hz) artifacts, such as sweat during the post-exercise session, that contaminated the quality of the signal. Therefore, we decided to reanalyze our data using a 1 Hz high-pass filter to minimize the presence of those artifacts. By visual inspection, we checked and removed noisy electrodes and epochs. In case of discarding more than 50% of the total epochs for a single participant, that participant was discarded from future analysis. We then used ICA decomposition to remove non-neuronal components such as blinks, heartbeats, muscle artifacts, etc. Finally, channels removed due to rejection were interpolated using neighboring channels, and all channels were referenced to the average.

Based on EEG recordings, we estimated three classes of measures: (1) measures estimating spectral power raw and normalized power spectra, median spectral frequency (MSF), spectral edge 90 (SEF90), and spectral edge 95 (SEF95), (2) measures estimating information content—spectral entropy, Kolmogorov-Chaitin complexity (K) and permutation entropy, and (3) measures estimating functional connectivity—symbolic mutualiInformation (SMI) and weighted symbolic mutual information (wSMI). Power spectrum density (PSD) was computed over the delta (1–4 Hz), theta (4–8 Hz) alpha (8–12 Hz), beta (12–30 Hz), gamma (30–45 Hz) spectral bands, using the Welch spectrum approximation (segments = 512 ms, overlap = 400ms). Segment rejections were windowed using a Hanning window and zero-padded to 4096 samples. Kolmogorov-Chaitin complexity was computed by compressing a discretization of the signal using a histogram approach with 32 bins. Permutation entropy was obtained by computing the entropy of a symbolic transformation of the signals, within the alpha, delta, and theta bands. SMI and wSMI were then computed from the same symbolic transformation, but data was first filtered using current source density estimates to diminish the volume conduction. SMI and wSMI were computed in theta, delta, and alpha bands⁵⁹. From the available connectivity metrics, we chose to use only wSMI as it is the only one that can detect purely nonlinear interaction dynamics and can be computed for each epoch⁶⁰.

Physiological measures

Electrocardiogram (ECG) data were acquired using the BIOPAC MP160 system (BIOPAC SYSTEMS Inc.) and the BIOPAC ECG100C amplifier. The data were sampled at a sampling frequency of 2 kHz and recorded using the AcqKnowledge v4.4 software. ECG disposable adhesive skin electrodes were used in a bipolar arrangement

of two electrodes and ground. The positive electrode was at the non-dominant wrist of the participant and the negative was on the contralateral ankle. The ground electrode was placed on the ipsilateral ankle.

ECG data were filtered with a notch filter (0.05 Hz) to remove baseline wander artifacts. A Butterworth highpass filter was applied (0.5 Hz) to attenuate linear drifts and physiological artifacts. Powerline interference was attenuated with a notch filter (50 Hz). Finally, the data were smoothed with a 3rd-order polynomial Savitzky-Golay filter. Peaks were detected using the native Neurokit2 algorithm. Finally, data were epoched based on the partition scheme in the EEG preprocessing section.

ECG metrics were grouped into three domains: time, spectral power, and information content. Time-domain metrics were (a) heart rate (HR), (b) standard deviation of the RR intervals (SDNN), and (c) root mean square of successive differences (RMSSD). Spectral power features were (a) low frequency of the heart rate variability (LF-HRV), (b) high frequency of the heart rate variability (HF-HRV), and (c) LF/HF HRV ratio. Information content metrics were (a) approximate entropy (AE), (b) sample entropy (SE), and (c) multiscale entropy (MSE). Initially, we used the native Neurokit2 algorithm to extract the peaks of the QRS complex. RR intervals were estimated as the sequential difference of the peak times. We estimated the time domain features based on the RR timeseries. For the spectral power metrics, the RR was evenly resampled at 4 Hz. Power spectra were computed over the LF-HRV (0.04–0.15 Hz) and the HF-HRV (0.15–0.4 Hz) frequency bands. The power spectrums were estimated using the Welch procedure.

Respiration. Respiratory data was acquired using a respiratory belt and amplified through the BIOPAC DA100C amplifier. Data were sampled at a sampling frequency of 2 kHz and recorded using the AcqKnowledge v4.4 software.

Respiratory metrics were grouped in the time and information content domain. Time-domain metrics were (a) respiration rate and (b) respiration rate variability. Information content was estimated based on multiscale entropy.

Pupillometry. Eye movements and pupil size in both eyes were recorded using oculometric glasses (Drowsimeter R100; Phasya, S.A) with a sampling frequency of 120 Hz. The eye tracker was calibrated at the start of each recording. Data was epoched based on the epoching scheme in the EEG preprocessing section. We identified 100 ms blink periods around blinks and removed the whole segment, as pre- and post-blink periods can introduce pupil dilation artifacts while the eye is recovering to its standard size. We interpolated segments using 3rd-degree cubic interpolation. Dilation speed outliers were calculated by estimating the median absolute deviation (MAD) of each value. Samples exceeding the deviation threshold were removed. Pupil dilation was smoothed using a moving average filter and baseline-corrected with a 100 ms period 2 s after the probe.

Pupil metrics were grouped in the same three domains: time, spectral power, and information content. Timedomain metrics were: (1) blink rate, (2) pupil size, and (3) pupil size variability. Spectral power metrics were: (1) low frequency pupil component (LFC), (2) high-frequency pupil component (HFC). The information content metric is multiscale entropy. The power spectra were estimated using the Welch procedure. As a post-registration note, we encountered issues extracting pupil metrics at the *Low Arousal* condition, as participants tended to have their eyes closed or partially closed for most of the trials. As our device was not sensitive to capture dilation in this setting, we additionally estimated (a) blink rate, (b) blink duration, (c) blink rate variability, (d) mean eye openness, (e) eye openness variability, (f) percentage of 70% eye closure and (g) percentage of 80% eye closure. As stated below, our registered plan was to reliably estimate all time, frequency, and complexity metrics that can be of use to our classifiers. Therefore, while we do not deviate from our original registered protocol, it is of note that these features could not be estimated reliably.

Electrodermal activity (EDA) data was acquired through skin electrodes on the index and middle finger and amplified through the BIOPAC EDA100C amplifier. Data was sampled at a sampling frequency of 2k Hz and recorded using the AcqKnowledge v4.4 software. All EDA metrics originated from the time domain: (a) galvanic skin response (GSR), (b) tonic EDA, and (c) phasic EDA. Extraction of the phasic and tonic components of the EDA was conducted with deconvolution of the EDA signal with a biologically plausible impulse response function with initially fixed parameters that are iteratively optimized per participant⁶¹.

Pattern recognition

To examine the physiological counterpart of the behavioral shifts in MB reports, we employed a supervised decoding approach. Using the multimodal neurophysiological measurements during the three experience sampling sessions, we trained multiple classifiers to discriminate across MB, MW, and SENS reports and identify whether MB is supported by a unique brain-body interaction pattern. This approach allowed us to extract meaningful brain-body interactions from the proposed arousal metrics without being conservative about the nature of the multiple comparisons between the various brain and body metrics.

As features, we opted to collect meaningful data in the time, frequency, information, and connectivity domain, unless such measurements could not be reliably estimated within our selected time window. The goal of the multiple selected metrics was to capture potential diverse spatiotemporal relationships (low-high frequency interactions, phase-amplitude interactions) that might extend across different recording modalities. Overall, we computed 57 features.

As targets, we used the participants' mental states (MB, MW, and SENS). Since this creates a multiclass classification problem, we focused on the binary classification of MB vs other reports. We expected to acquire 40 samples per participant and condition (i.e. baseline and arousal states), giving a total of 1040 (26*40) samples per condition. We expected that 5% of the samples to correspond to the target report (MB), yielding an imbalanced problem with only 52 target samples per condition.

As learning algorithms, we tested parametric and non-parametric models, such as Support Vector Machines (SVM), Random Forests (RF), and Extremely Randomized Trees (ET). SVM is a classification technique that aims to separate labeled inputs by creating a hyperplane that maximizes the distance of their

features. Given a set of n-labeled inputs, SVM provides a hyperplane in an n-dimensional space that maximally separates the differently labeled groups. An RF classifier is a meta-estimator. Various classifiers ("decision trees") are trained in different parts of the input dataset, and each classifier uses only that part of the dataset to predict the label of the input. Then, the predictions of each classifier are pooled ("bagged") together, and an optimal decision is chosen based on the label with the most predictions ("votes"). Finally, an ET classifier is a meta-estimator that employs a similar voting scheme. However, in the case of the ET classifier, trees are trained on all the features and the cutoff point of the trees (how the various metric nodes are arranged to reach a decision) is randomized. Since our problem is highly imbalanced, we also tested outlier detection algorithms (i.e. one-class classifiers), aiming to isolate MB from the other reports by considering MB as either an inlier or outlier. Therefore, we tested the one-class counterparts of the SVM (One-class SVM) and RF (Isolation Forests) algorithms.

For model selection and performance estimation, we employed two different cross-validation approaches. First, we used a 5-fold stratified cross-validation scheme trained with all the samples. This provided us with performance estimates of classifiers aimed at obtaining patterns of brain and body function that can predict MB reports in known participants. As a second approach, we used a 5-fold group stratified cross-validation scheme, using participants as groups. In this scenario, each participant was either on the train or on the test set. Thus, it aimed at learning general patterns of brain and body function that could predict the report of MB in unseen participants. In other terms, the first approach aimed at learning patterns that could discriminate MB from other reports while accounting for each participant's variance, while the second strengthened the claim, aiming to learn general patterns that could be found in unseen participants.

As performance metrics, we report a) recall, b) precision, c) F1-score, d) area under the ROC curve (AUC), and e) balanced accuracy. Recall is the ratio of how often an item was classified correctly as a positive outcome (True Positive / True Positive + False Negative). Similarly, precision is the ability of the model to return only the data points in the relevant class (True Positive / True Positive + False Positive). F1-score is the harmonic mean of precision and recall. The AUC curve is another evaluation metric that summarizes how well the classifier predicts a class based on different thresholds of true positive and false positive ratios. Finally, balanced accuracy is an evaluation metric suitable for imbalanced datasets, where one class appears at significantly different frequencies than the others. Balanced accuracy is useful because it is estimated as the average of specificity and sensitivity, simultaneously controlling for very high precision due to classifying nothing as the infrequent class and very high recall due to classifying everything as the infrequent class.

We selected each model's hyperparameters using nested cross-validation (same scheme as the outer cross-validation), using the F1-score as our optimization metric.

To evaluate the variance in the classifier performance and compare it to chance level, we performed repeated cross-validation (10 times), while training also a "dummy" classifier to obtain the empirical chance level of the training samples distribution. This type of classifier generates predictions based on the distribution of training samples for each class without accounting for the features.

The decoding analysis was implemented in Python using Julearn⁶² and Scikit-Learn⁶³. Metrics were estimated from existing Python libraries: MNE⁶⁴, NICE⁶⁵, Neurokit⁶⁶, and custom in-lab Python functions.

Results

Participants

To achieve a power of 0.95 at an alpha threshold of 0.05, we acquired 3 sessions of 40 trials per session from 26 participants (mean age= 26.38, sd= 4.53, min= 20, max= 40; female= 11). As a post-registration note, in case participants could not adhere to the strict 3-week protocol (30% total sessions), they were rescheduled to a later date that respected their sleep schedules to avoid time windows with potential extreme slow-wave activity⁵⁰. Due to data corruption, one participant had 30 trials in one of the three sessions, and one participant had 33 trials in one of the three sessions. The remaining two sessions were completed for both participants.

Behavioral data

Occurrences of mental state reports alter across arousal conditions

We found a main effect for mental states, with MB being reported at significantly lower rates (Mean proportions±SD: MW=0.56±0.21, SENS=0.2±.14, MB= 0.12±0.13; Kruskal H= 124.07, p= 1.2e-27, eta²= 0.53) compared to MW (Dunn's test= -10.75, p_{FDR} = 1.8e-26) and to SENS (Dunn's test= -2.85, p_{FDR} = 4.3e-03). Additionally, MW was reported significantly more frequently compared to SENS (Dunn's test= 7.9, p_{FDR} = 4.3e-15; Fig. 2). As the study was focused on wakeful mental states, "SLEEP" reports were not included in the analysis (Mean proportions ±SD: *Baseline*= 0.03±.05, *High Arousal*= 0.05±.07, *Low Arousal*= 0.26±.21, Total= 0.1±.17).

We found that a model including all conditions outperformed a null model with only an intercept (Full_{LogLik} = -1021, Null_{LogLik} = -1046.83, χ^2 = 51.57, df = 2, p_{Bonf} = 6.1e-12): MB was reported significantly more frequently in *Low Arousal* compared to *Baseline* (Marginal Mean= -0.79, SE= 0.14, CL= [-1.16, -0.43], p_{FDR} = 1.8e-08) and to *High Arousal* (Marginal Mean= -0.97, SE= 0.15, CL= [-1.35, -0.59], p_{FDR} = 7.9e-11) (Fig. 3a). However, MB reports during *Baseline* and *High Arousal* were comparable (Marginal Mean= 0.17, SE= 0.15, CL= [-0.21, 0.56], p_{FDR} = 2.4e-01). A visual inspection of the individual marginal means showed that this effect was mostly consistent across participants and was not driven by extreme cases (Fig. 3b-d).

Additionally, generalized estimating equations (GEE) showed a significant interaction for MW between *Low Arousal—Baseline* (beta= 6, SE= 1.5, CL= [3.06, 8.94], p_{FDR} = 6.4e-05) and *Low—High Arousal* (beta= 8.23, SE=1.6, CL= [5.1, 11.36], p_{FDR} = 2.6e-07). We also found significant interactions in SENS reports, such that SENS tended to be higher in *Baseline* compared to *High* (*SENS Baseline—SENS High*: beta= 2.54, SE= 0.81, CL= [0.96, 4.12], p_{FDR} = 1.7e-3) and *Low Arousal* (*SENS Baseline—SENS Low*: beta= 2.46, SE= 0.77, CL= [0.96, 3.97], p_{FDR} = 1.3e-3). It is of note that this analysis yielded no significant results for MB, but the overall trend of the beta estimates was consistent with our positive results of the logit model above (Supplementary Fig. S3).



Fig. 2. Mind blanking (MB) was reported significantly less frequently compared to mind wandering (MW) and perceptual sensations (SENS) across all arousal conditions, validating what is generally reported in the literature. Density kernels show overall data dispersion and clustering trends. Point plots are individual subject estimates. Box plots show medians and interquartile ranges, while whiskers indicate extreme values and diamonds indicate outliers.



Fig. 3. The frequency of mind blanking (MB) reports altered across the three arousal conditions. (**a**) MB report probability increased in *Low Arousal* (after sleep deprivation) compared to *High Arousal* (after intense exercise) and *Baseline*. Density kernels indicate overall data dispersion and clustering trends. Point plots represent participants' MB report probabilities. Box plots indicate medians and interquartile ranges, whiskers indicate extreme values, and diamonds indicate data outliers. (**b**–**d**) Bar plots denote single-subject marginal means, comparing MB reports across arousal conditions. Compared to *Baseline*, there was no significant change during *High Arousal* (**b**). However, there was a visible trend favoring an increased probability of MB reports in the *Low Arousal* condition compared to baseline and *High Arousal*, signifying that the effect was present in most participants (**c**–**d**).

.....

MB was characterized by higher reaction times

There was a main effect of arousal conditions, with reports during *Baseline* being reported the fastest and during *Low Arousal* the slowest (Fig. 4a). Also, there was a main effect of mental states, with MW reports being reported the fastest and MB reports the slowest (Fig. 4b). A significant interaction between MW and arousal showed that MW was reported the slowest in *Low Arousal* (Fig. 4c). A significant interaction between MB and arousal condition showed that MB was reported the slowest in *High Arousal* and *Low Arousal* (Fig. 4e). A model including both arousal and reaction times outperformed simplified models including only null or main effect terms (Full_{LogLik} = 2889.76, χ^2 = 47.1, df= 4, p_{Bonf}= 1.5e-09; Fig. 4c). For a detailed overview of main effects and interactions, see Supplementary Table S3.

Transition probabilities showed reduced probability to transition to MW in Low arousal

Markov transition probabilities indicated significant differences only between *High* and *Low Arousal* conditions (Fig. 5), such that MW was more likely to be followed by MB (t= 3.26, CI= [0.03, .15], p_{FDR} = 9.7e-03, Cohen's D= 0.74). Also in *Low Arousal*, both MW (t= -3.79, CI= [-0.31, -0.9], p_{FDR} = 7.6e-03, Cohen's D= -0.86) and SENS



Fig. 4. Mental states had different reaction times depending on arousal conditions. (**a**) Reaction times at *Baseline* arousal were reported the fastestlowest, followed by *High* (after exercise) and *Low Arousal* (after sleep deprivation), collapsed across all arousal levels. Point plots show individual subject estimates. Box plots show medians and interquartile ranges, while whiskers show extreme values. (**b**) Mind wandering (MW) was reported the fastest, followed by Sensations (SENS) and mind blanking (MB), collapsed across all mental states. Point plots show individual subject estimates. Boxplots show medians and interquartile ranges, while whiskers show extreme values. (**c**-**e**) Interaction between arousal condition and mental state reaction times: MW was reported the slowest in *Low Arousal* compared to *Baseline* and *High Arousal*, while MB was reported the slowest in the *Low Arousal* and *High Arousal* conditions compared to *Baseline*.

(t= -3.43, CI= [0.37, -.009], p_{FDR} = 9.5e-03, Cohen's D= -0.77) were less likely to be followed by MW (Fig. 5; Supplementary Fig. S4).

Exploratory analysis 1: MB frequency did not correlate with SLEEP frequency

As we wanted to avoid participants confounding MB and SLEEP reports, we opted for a paradigm that allowed participants to report both. Spearman correlations on each condition examined whether these two states were correlated. We did not find any significant correlation between MB and SLEEP report probabilities across any arousal condition. (*Baseline*: r = 0.13, p = 5.3e-01, *High Arousal*: r = 0.31, p = 1.3e-01, *Low Arousal*: r = -0.05, p = 8.2e-01) (Supplementary Fig. S5). To strengthen the claim that MB and SLEEP reports do not covary, we additionally ran separate equivalence tests on each correlation. No test was able to reject an equivalence claim (*Baseline*: z = -0.34, p = 3.7e-01, *High Arousal*: z = 0.54, p = 7e-01, *Low Arousal*: z = 0.72, p = 2.3e-01). Therefore, these results remain indeterminate.

Exploratory analysis 2: High arousal: MB reports increased at the start, but not the end, of the experience sampling session

While we found that MB reports were more frequently in *Low Arousal*, we did not find any significant effect of *High Arousal*. In our original hypothesis (Supplementary Table S1), we registered a potential alternative explanation for the absence of an effect of high arousal in mental state report frequency. High arousal, as elicited

Low – High Transition Probabilities



Fig. 5. After sleep deprivation (*Low Arousal*), participants were more likely to transition from mind wandering (MW) to mind blanking (MB) compared to the condition of physical exercise (*High Arousal*). Additionally, participants were less likely to transition to MW, either when departing from reports about sensory perception (SENS) or from MW itself. Arrows indicate the direction of the mental state transition. Bold font indicates statistical significance (FDR corrected).

by high-intensity exercise, might not last for the full session, and our session would represent a gradual return to *Baseline Arousal*. To test for potential effects of more frequent MB reports only at the start of the experience sampling we split the *High Arousal* session in two parts and compared the count of MB reports across the start and the end of the experiment. Using a chi-squared test we found a significant effect, with MB reports being more frequent (divergence= 4.08, p= 3.2e-02) during the first half of the *High Arousal* condition compared to the second half (MB_{start} = 93, MB_{end} = 66). We additionally attempted to validate this hypothesis by splitting the session into 4 and 6 discrete segments of 10 and 7 trials each and replicated the same analysis. However, this analysis did not reach significance. Finally, to provide further evidence for reduced occurrences of MB across time, we considered only the first and last 10 trials. We found a significant effect of more frequent MB occurrences (divergence= 7.39, p= 6.6e-03), with the first 10 trials of the *High Arousal* condition inducing more MB compared to the second half (MB_{start} = 51, MB_{end} = 27).

Classification of MB reports was outperformed by classification containing both BRAIN-BODY markers

We trained a cohort of different classification algorithms and evaluated their capacity to classify MB reports from mental states with content (MW, SENS) based on 26 BRAIN (EEG) and 31 BODY features (12 ECG, 4 EDA, 8 RSP, 7 EYE), spanning time, frequency, information, and connectivity domains for each mental state report. In our original report, we registered that these features would be estimated across the 110 s pre-probe window, with bad epochs being dropped. However, across an 110 s epoch, even a nonlinearity of 1 s would result in epoch removal, leaving a total clean sample of 25 / 78 sessions (29.4%), and a total of 1060/3120 (33.3%) clean epochs. Therefore, to preserve datapoints and data quality and minimize data discarding due to brief non-linearities, we opted for an extra step in bad epoch removal. After the initial epoch definition of 110 s, we followed it up by partitioning that epoch into 5 s sub-epochs, resulting in 22 sub-epochs per epoch. We then proceeded to do bad epoch removal and EEG marker estimation on those sub-epochs. If an epoch consisted of more than 50% bad sub-epochs, it was discarded. Then, we averaged across within each epoch, resulting in no lost sessions, and a total of 2734 / 3120 (87.6%) total sample size.

Having a final 2734 reports x 57 features matrix per report, we trained multiple classifiers on the total dataset, to examine whether a specific brain-body profile would outperform chance level classification of MB reports (Table 1).

Due to the unbalanced nature of our dataset, we evaluated classifier performance based on balanced accuracy, as it avoids inflated performance estimates on unbalanced datasets. Overall, we found that a balanced random forest (a random forest that undersamples the majority class in each bootstrap to equate class count) had above-chance performance and outperformed all other examined classifiers (Fig. 6a). We additionally examined whether we could predict unknown subjects, by leaving a subset of subjects out on each iteration. Due to the high degree of per-fold variance, we do not consider any classifier as meaningfully performing above chance level (Fig. 6b). Importantly, these results were replicated when we trained the classifiers in the 1 Hz filtered data (Supplementary Fig. S6a,b; Supplementary Table S4).

Having established that MB reports can be predicted from known subjects, we then examined whether a brain-body data pattern would outperform classifiers trained solely on either BRAIN or BODY features. To this end, we fit and optimized a separate balanced random forest classifier on discrete feature combinations of our dataset. For a full report of the performance on different features, see Table 2 and Supplementary Table S5.

Overall, we found that a classifier trained on both BRAIN and BODY markers numerically outperformed classifiers trained solely on BRAIN or BODY features across all our performance metrics (Fig. 7a,c; Supplementary Fig. S7a,c; Table 2; Supplementary Table S5). To evaluate the impact of the number of features on the capacity of

Examined	Classifier	Recall	Precision	F1	ROC AUC	Balanced accuracy
Known subjects	Balanced RF	0.62, [0.6, 0.64]	0.26, [0.26, 0.27]	0.37, [0.36, 0.37]	0.71, [0.7, 0.72]	0.66, [0.65, 0.67]
	SVM	0.29, [0.28, 0.31]	0.28, [0.27, 0.29]	0.29, [0.27, 0.3]	0.62, [0.61, 0.63]	0.58, [0.58, 0.59]
	ET	0.16, [0.15, 0.17]	0.61, [0.58, 0.64]	0.25, [0.23, 0.26]	0.73, [0.72, 0.74]	0.57, [0.56, 0.58]
	RF	0.14, [0.13, 0.15]	0.57, [0.53, 0.6]	0.22, [0.21, 0.23]	0.71, [0.7, 0.72]	0.56, [0.56, 0.56]
	IF	0.14, [0.13, 0.16]	0.2, [0.19, 0.22]	0.17, [0.15, 0.18]	0.52, [0.52, 0.53]	0.52, [0.52, 0.53]
	OC SVM	0.89, [0.86, 0.92]	0.15, [0.14, 0.15]	0.25, [0.25, 0.25]	0.51, [0.5, 0.51]	0.51, [0.5, 0.51]
	DUMMY	0.14, [0.13, 0.15]	0.14, [0.13, 0.15]	0.14, [0.13, 0.15]	0.5, [0.49, 0.5]	0.5, [0.49, 0.5]
Unknown Subjects	Balanced RF	0.46, [0.41, 0.51]	0.18, [0.16, 0.2]	0.25, [0.23, 0.27]	0.55, [0.53, 0.57]	0.54, [0.53, 0.56]
	IF	0.23, [0.19, 0.27]	0.18, [0.16, 0.2]	0.19, [0.17, 0.22]	0.53, [0.51, 0.54]	0.53, [0.51, 0.54]
	RF	0.05, [0.04, 0.06]	0.36, [0.29, 0.44]	0.08, [0.06, 0.09]	0.54, [0.52, 0.55]	0.51, [0.51, 0.52]
	OC SVM	0.87, [0.82, 0.92]	0.14, [0.13, 0.15]	0.24, [0.22, 0.26]	0.51, [0.5, 0.52]	0.51, [0.5, 0.52]
	ET	0.03, [0.02, 0.03]	0.36, [0.26, 0.45]	0.05, [0.04, 0.06]	0.53, [0.52, 0.55]	0.51, [0.5, 0.51]
	DUMMY	0.15, [0.14, 0.16]	0.15, [0.13, 0.17]	0.14, [0.13, 0.16]	0.5, [0.49, 0.51]	0.5, [0.5, 0.51]
	SVM	0.2, [0.17, 0.22]	0.16, [0.14, 0.17]	0.16, [0.15, 0.17]	0.49, [0.47, 0.5]	0.5, [0.49, 0.51]

Table 1. A balanced random forest classifier outperformed all classifiers when compared across balancedaccuracy. Cells indicate mean and 95% CI. RF = Random Forest; SVM = Support Vector Machine; ET =Extreme Trees; IF = Isolation Forest; OC SVM = One-Class Support Vector Machine



Fig. 6. Classification performance was above chance level when mind blanking (MB) reports were pooled across subjects, but not after training on a subset of participants and classifying the remaining subset. (**a**) A balanced random forest classifier provided the highest classification performance across all examined classifiers including known subjects. (**b**) A balanced random forest classifier provided the highest classifier provided the highest classification performance across all examined classifiers on unknown samples. However, due to the high variance, we could not consider it meaningful. Individual points indicate performance on the folds of the repeated cross-validation. Results are ordered based on descending order of performance. Chance-level performance is indicated by the Dummy classifier. RF = random forest; SVM = support vector machine; ET = extreme trees; IF = isolation forest; OC SVM = one-class support vector machine.

the learning algorithm to extract relevant information, we also trained the balanced random forest model using randomly shuffled bodily features. EEG features were not altered. The model with the shuffled values showed a decline in classification performance, providing evidence that, when classifying mental states, a model trained on both brain and body data learns unique information from both domains (Fig. 7d; Supplementary Fig. 7d). For feature importance, we calculated Shapley Additive Explanations (SHAP) values for each feature in our dataset. SHAP values estimate the marginal contribution of each feature, averaged across every potential feature combination. In this manner, each value represents how much this feature contributes to the classification, after controlling for the impact of other features on this feature's importance. We found that the model relied mostly on EEG and EYE openness features to discriminate MB reports when pooling MB occurrences across all three conditions. (Fig. 7b; For an extensive list of all SHAP values, see Supplementary Fig. S8). Importantly, feature importance did not substantially change when filtering the data with a 1 Hz filter (Supplementary Fig. S7b; For an extensive list of all SHAP values, S9). Overall, the comparable performance of the

Classifier	Recall	Precision	F1	ROC AUC	Balanced Accuracy
BRAIN + BODY	0.62, [0.6, 0.64]	0.26, [0.26, 0.27]	0.37, [0.36, 0.37]	0.71, [0.7, 0.72]	0.66, [0.65, 0.67]
BRAIN	0.61, [0.59, 0.62]	0.24, [0.24, 0.25]	0.35, [0.34, 0.36]	0.7, [0.69, 0.71]	0.65, [0.64, 0.65]
BODY	0.59, [0.58, 0.6]	0.22, [0.21, 0.22]	0.32, [0.31, 0.32]	0.66, [0.66, 0.67]	0.61, [0.61, 0.62]
EYE	0.57, [0.55, 0.59]	0.21, [0.21, 0.22]	0.31, [0.3, 0.32]	0.64, [0.63, 0.65]	0.61, [0.6, 0.62]
ECG	0.55, [0.54, 0.57]	0.18, [0.17, 0.18]	0.27, [0.26, 0.27]	0.58, [0.57, 0.59]	0.56, [0.55, 0.57]
EDA	0.6, [0.57, 0.63]	0.17, [0.17, 0.17]	0.26, [0.26, 0.27]	0.57, [0.56, 0.58]	0.55, [0.54, 0.56]
RSP	0.52, [0.5, 0.54]	0.15, [0.15, 0.16]	0.24, [0.23, 0.24]	0.53, [0.52, 0.54]	0.52, [0.51, 0.53]

 Table 2. A classifier trained on a combination of BRAIN and BODY features outperformed classifiers trained solely on BRAIN or BODY features, when evaluated with balanced accuracy. Cells indicate mean and 95% CI.



Fig. 7. Classification of MB improves when considering both BRAIN and BODY. (**a**) Balanced random forest classifier trained on a combination of BRAIN and BODY features outperformed classifiers trained solely on BRAIN or BODY features when evaluated with balanced accuracy. Individual points indicate performance on the folds of the repeated cross-validation. (**b**) Subset of the 10 features with the highest mean of the absolute SHAP values obtained from the balanced random forest classifier. (**c**) The per-fold differences between the classifier trained on both BRAIN and BODY features and the one trained only on BRAIN data suggest that incorporating both feature domains provides a slight performance improvement over using BRAIN data alone. The shaded region indicates better performance for the classifier trained on both feature domains. The star indicates the mean difference. The solid, horizontal line represents the 95% highest-density intervals of the distribution. Red dots indicate per-fold differences. (**d**) The per-fold differences between the classifier trained on both BRAIN and BODY features and the one trained on BRAIN asymptotic trained on both BRAIN and BODY features and the one trained on BRAIN and shuffled BODY data suggest that the model with both BRAIN and BODY data does not consider the body markers as noise.

.....

models, and the high degree of overlap in the ranking of the feature importance point to the robustness of the models.

Exploratory analysis 3: Feature importance altered across arousal conditions. The decoding analysis in known samples showed that we can predict MB instances from the combination of brainbody markers with adequate accuracy when MB instances were aggregated across different arousal conditions.



mean(|SHAP value|) (average impact on model output magnitude)

Fig. 8. Ranking of features by mean absolute SHAP value extracted from the balanced random forest classifier varied across different arousal conditions. (**a**) Magnitude of SHAP values for a balanced random forest classifier trained on MB reports collected during the *Baseline Arousal condition*. The model relied mostly on features from the EEG frequency domain. (**b**) Magnitude for SHAP values for a classifier trained on MB reports collected during the *Low Arousal* condition (after sleep deprivation). The model mostly used spectral power in the EEG delta band. (**c**) Magnitude for SHAP values for a classifier trained on MB reports collected during the *High Arousal condition* (after intense exercise). The model relied mostly on features from eye openness, EDA, and the EEG frequency domain

We were further interested in whether this classification was achieved based on a universal mechanism, or whether we could detect arousal-dependent brain-body configurations that predict MB. To this end, we trained a balanced random forest classifier solely on data acquired from *Baseline*, from *High*, and from *Low Arousal*. We found that *Baseline* had the best performance (0.67, [0.65, 0.68]), followed by *Low Arousal* (0.64, [0.63, 0.65]), and finally *High Arousal* (0.61, [0.6, 0.63]). We retained comparable performance when examining the arousal partitions of the 1 Hz filtered dataset (Supplementary Table S6-7). Examining the SHAP values for each arousal state, we saw that the models relied on distinct feature domains. During *Baseline*, the model relied on markers from the frequency domain of EEG (Fig. 8a). During *Low Arousal*, MB classification was obtained using the delta band power, by far the most dominant marker (Fig. 8b). Finally, in *High Arousal*, the model did not rely on a single feature, rather in a combination of eye openness, GSR, and the frequency domain of EEG (Fig. 8c). Similar feature importances were observed in the 1 Hz filtered dataset (Supplementary Fig. S10). However, in the 1 Hz filtered dataset, we observed that ECG features tended to rank higher (Supplementary Fig. S11–16).

Exploratory analysis 4: Feature importance altered based on the pre-probe analysis window

A potential caveat of utilizing the full pre-probe period of 110 s before a report is that we might capture multiple mental states, and the actual statistical regularities might be weakened when averaged across. With this consideration, we examined whether we could improve classification performance when classifying MB from the last 10 s before a report. We defined a secondary brain-body data matrix, with BODY features that could be estimated from 10 s of body activity. Across both 0.1 and 1 Hz filters we retained comparable performance in the classifiers trained on both EEG and bodily markers, as well as solely EEG or BODY markers (Supplementary Fig. S17–20; Supplementary Table S8 and 9). However, we observed decreased performance in the classifier trained solely in the eye openness data (Supplementary Table S8 and 9). An examination of feature importance showed that the beta, delta, and theta bands of the EEG frequency domains remained the most important EEG features, but there was a reduction in the importance of the EYE features and an increase in the importance of EDA (Supplementary Fig. S17b, 18, 19b, 20). Importantly, our results were not affected by the choice of filtering parameters, indicating robustness of our results to preprocessing parameters.

Discussion

We used experience sampling combined with EEG and peripheral physiological recordings under different autonomic arousal conditions to determine whether MB reports in neurotypical individuals were supported by distinct brain-body configurations compared to mental states with reportable content. Overall, our results show that MB is a mental state that becomes more prevalent in low and partially in high arousal states, and that MB is driven by both brain and body processes, providing evidence for an embodied account of MB.

Behaviorally, we found that MB was reported at significantly lower rates compared to sensory experiences or MW, irrespective of the arousal condition. This finding is in line with past research showing that MB rates vary between 5 and 10% of total probe instances, across both uninterrupted thinking¹² and task engagement¹¹. We also show that sleep deprivation significantly increased the frequency of MB occurrences. Sleep deprivation

has been shown to induce a low arousal state during which cognitive performance declines⁶⁷, metabolic and physiological processes change⁶⁸, and unique neuronal markers, such as slow-wave activity, emerge⁶⁹. After sleep deprivation, participants also tend to perform worse in sustained attention tasks⁷⁰, with results suggesting a true effect of sleep deprivation on more "misses" (no response when necessary) compared to "false alarms" (response when unnecessary)⁷¹, a finding that was recently shown as a behavioral correlate of MB¹¹. Additionally, sleep deprivation and mounting sleep pressure have been positively correlated with more MW instances^{72,73}, suggesting an overall mode shift from task engagement to MW⁷⁴. Our results challenge these past findings by showing that participants were more likely to experience an MB event rather than MW after sleep deprivation. We also found that MW was in fact more likely to decrease after sleep deprivation. This is further supported by the results of the transition analysis, where MW reports were less likely to be followed by another MW report, and more likely to be followed by MB. Such discrepancies in the reportability of MW after sleep deprivation could be possibly explained by the explicit inclusion of MB as a reportable mental state in the experience sampling that our design opted for. In other words, it might be that the observed MW occurrence increases after sleep deprivation can be accounted for by MB reports, once participants have the chance to opt between these two mental states in a more fine-grained way.

In terms of high arousal induced by high-intensity exercise, our analysis did not reveal any significant effects on MB occurrences. As per the provided registered protocol alternative explanation (Supplementary Table 1), we hypothesized that this arousal manipulation might not have been overall effective as it could not have produced effects that would last across the whole experience sampling session. To test whether MB frequency reports would differ between the beginning and at the end of the session, we split the dataset into two parts. When split, we indeed found a significant difference between the frequency of MB reports. This result was replicated when considering only the first and last 10 trials per subject, which maximized the distance between initial and final physiological arousal within the session. However, we were not able to find any differences when the data were split into smaller bins. Together, we consider that these results provide partial evidence for our registered hypothesis, showing that residual high arousal effects after intense exercise can increase the frequency of MB reports.

In addition to the frequency of mental states across arousal conditions, we also examined whether rreaction times differ across arousal conditions and mental states. In general, reports in low arousal tended to be the slowest, consistent with a wide range of attention tasks that show slower reaction times in sleep deprivation compared to baseline arousal⁷⁵. We consider these findings as additional evidence that the arousal manipulation was effective in that it lowered overall vigilance levels of the . We also observed a main effect of mental states, such that MB tended to be reported significantly slower compared to MW and sensations. Contrary to our current results, we recently found that MB was reported faster when compared to other mental states when content had to be evaluated¹². This apparent mismatch in results can be explained when considering that MB can be a state devoid of content, and therefore, there is the binary consideration of "yes/no" when evaluating thought content, which might be a relatively fast decision. This can be different, for example, from the evaluation of content-full mental states, which demand a sequential evaluation of both content presence ("yes/no") and content evaluation ("what is the content about?"). This way, the difference in results can be explained by the imposition of an additional cognitive evaluation. Overall, we suggest that these results might reflect a gradient of vigilance, with participants being the most alert at baseline arousal, and progressively declining during high and low arousal conditions, as well as more vigilant when reporting mental states with content compared to MB. Of note, we observed two interesting interactions between mental states and arousal conditions. MW tended to be reported slower in low arousal compared to baseline and high, which is consistent with our interpretation of reaction times as marking vigilant states. However, as we also observed that MB reports tended to be reported slower in both high and low arousal conditions, we speculate that this might be preliminary evidence that arousal modulates how engaged participants are with their current mental states. In this sense, exercise fatigue can lead to an MB state that takes longer to recover from when probed for a report.

A final explanatory analysis revolved around the relationship between sleep and MB. We recently posited that MB is a distinct mental state characterized by a unique phenomenological profile of no content⁷⁶, and unique neuronal markers, characterized by high cortical integration and low cortical segregation¹². This neuronal configuration is atypical of wakefulness⁷⁷, and is more closely reminiscent of brain configurations during deep sleep⁷⁸. In conjunction with the presence of slow wave intrusions during wakefulness as a marker of MB reports¹¹, a classic marker of NREM sleep, an emerging issue is whether MB is a misrepresented instance of sleep. This issue is further complicated by the postulation that in MB there is no content⁷⁶, and thus does not functionally represent a wakeful state where people can report content. To avoid this pitfall, we introduced sleep as a potential report during experience sampling. We found that people discretely reported MB and sleep, providing evidence that when provided with such options, people can differentiate between these two experiences. Additionally, we did not find that MB and sleep tended to covary. To strengthen this claim, we ran equivalence tests for each correlation across arousal conditions. However, no test showed a positive result for equivalence. Therefore, these results remain indeterminate, with a trend for no relationship between MB and sleep.

Having established that MB occurrence varied across different physiological arousal conditions, we then examined whether MB could be decoded by brain and body markers. With the aim of showing single trial prediction, we trained different models on EEG and physiological markers, spanning time, spectral, complexity, and connectivity domains. Overall, we were able to achieve above-chance-level classification, showing that there exist unique brain-body patterns that can discriminate MB reports from mental states with content. However, we were not able to show above-chance-level classification when training classifiers on unknown subjects. Therefore, our results are not generalizable to novel populations due to the high amount of variance between subjects. Of importance is the result that a combination of EEG and physiological markers marginally, but consistently, outperformed both EEG and physiological markers. Overall, we observed an improvement of 2–5

% in classification performance in balanced accuracy. This improvement can be attributed to unique information inherent in body signals, as evidenced by the comparison of the classifier trained on both brain and body data compared to classifiers trained solely on brain data or brain and shuffled body data. The classifier trained on both brain and body data does not consider body features as noise or redundant. Overall, while our results suggest a high degree of overlap between brain and body information in MB, they indicate that information about MB extracted from the body is partially independent of the EEG features. Feature importance ranking derived from the classification model indicates that the low and mid frequencies of the EEG power spectrum and metrics of eye openness are useful predictors of MB. This finding was consistent across analysis windows and preprocessing parameters. Importantly, all classifiers trained on body markers had above chance performance with variant degrees of variability, with the highest performing being the EYE (eye openness) and the ECG (heart-rate variability), providing evidence that MB can be decoded solely from bodily signals.

To further validate our protocol, we ran two exploratory analyses, with the aim to examine whether classification performance varies based on the analyzed pre-probe window and whether feature importance alters across arousal conditions (For a full Discussion, see the Supplementary Discussion on Methodology). Overall, when examining a classifier trained on a brief 10 s window before MB reports, we found similar performance compared to the full 110 s classifier. What was interesting was that, while EEG performance remained the same, performance on classifiers trained solely on body features decreased. As brain-physiology coupling occurs at varying time delays across cardiac⁷⁹ and respiratory domains⁸⁰, we interpret these results as evidence that bodily contributions on MB are based on slow, oscillatory processes that might not be captured from examining short pre-probe periods. At the same time, our classification analysis on separate arousal conditions showed distinct brain-body configurations that can predict MB reports. As our decoding approach did not permit any inference of the directionality effect, or decomposing interactions within and across physiology modalities, at this stage we claim that our results point to discrete physiological pathways that elicit MB reports. Overall, we show that our enhanced classification is retained across different analysis windows and different arousal conditions.

Similarly, enhanced classification when considering a brain-heart matrix compared to solely brain markers was also shown for patients with disorders of consciousness, where the inclusion of cardiac features outperformed classification based solely on EEG markers⁸¹. To our knowledge, our results are the first to extend multivariate decoding past the brain-heart axis and consider the inclusion of multiple unique bodily afferent sources in classifying mental states. The overall success of the brain-body decoding paradigm in classifying consciousness levels and mental states provides evidence that bodily information is not redundant and is not necessarily fully represented within brain dynamics. Instead, an embodied approach, stressing bidirectional information routes between brain and body can provide better predictive power and assist in more comprehensive, generative models of experience^{34,82}.

A neurobiological explanation of our results comes from an integrative model of content, task engagement, and arousal which suggests that the relationship between thought and arousal can be conceptualized as an inverted u-curve. This means that an optimal arousal level modeled by the locus coerulius-norepinephrine (LC-NE) release is necessary to actively engage and control our thoughts, either during task engagement or MW⁸³. This stance treats thought as an active task, where engagement is necessary for clear content and control of thought dynamics. As arousal tapers off to non-optimal levels of the inverted U-curve, we experience concurrent, opposing thoughts that serve exploratory purposes for optimal performance, such as exploring different strategies. This necessitates flexibility and malleability of content. We here suggest that our results supplement this model by providing an account of the extremities of the optimal U-curve. As the model suggests degradation of thought clarity when we move closer to arousal extremities, we consider MB reports as instances where no content can be clear or present, extending this unifying framework to the entire arousal U-curve. Neurophysiologically, this model has translated to investigations of pupil dilation as a function of mental state and task engagement with pupil size yielding both positive^{26,84} and null results¹¹ in discriminating on-task vs off-task mental states, as well as contrasting MB and MW. Part of the ascending arousal network, the LC modulates cardiac, galvanic, respiratory, and pupillary activity^{28,85}. In addition, the LC innervates projections responsible for eyelid openness⁸⁶. The combinatorial high performance of different body markers in classifying MB reports, and the evidence that altered levels of arousal increase MB occurrences provide further support for the modulatory role of the ascending arousal system in mental states and thought reportability.

From a theoretic perspective, our study challenges the conception that brain information is uniquely suitable to understand thought reportability and provides support for an embodied account of the mind. Embodiment moves the seat of mental events away from the brain and reformulates cognition as resulting from brain-body interactions. An extensive literature has shown how cataloged cardiac, respiratory, gut, and pupillary effects on perception³⁰, action⁸⁷, metacognition³¹ and consciousness⁸¹, while the collective interplay of peripheral systems has discriminatory power for clinical⁸⁸ and consciousness classification⁸⁹. We show here that within embodiment, the body is not only facilitatory, but also might impede access to our mental lives. Under specific brain-body configurations, we are not able to clearly formulate mental content.

Some limitations pertain to our study. First, the nature of experience sampling discretizes the continuous nature of ongoing thinking. As there is no consensus on how long a mental state might last, or whether all mental states last the same length, results might average across different mental states. While we attempted to circumvent this problem by analyzing different pre-probe windows, it remains unclear whether all mental states last the same, and what is their actual duration. Secondly, the post-exercise setup might be suboptimal in examining the effects of high arousal on ongoing cognition. Neuronal and electrophysiological recordings have shown that the duration of the effects of exercise on ongoing brain and physiological activity^{45–47} is highly variant. In addition, it is unclear whether brain and body recover to baseline states at the same rates, potentially confounding the post-exercise importance of cortical and physiological markers in cognition. Experience sampling with online probes during exercise could overcome this challenge.

In conclusion, our study suggests that MB is an arousal-modulated mental state, with a unique cortical and physiological profile. We think that our results pave a new paradigm for an embodied account of mental states, where the phenomenology of our mental lives is expressed based on both our body and our brain state. At the same time, our results challenge the neurocentric approach to mental state research, putting emphasis on the constant brain-body interactions that shape our cognition. As MB research continues to evolve, we consider our findings elaborative for clinical and experimental accounts of spontaneous thinking, where we move towards a complex and dynamic conception of our mind.

Data availability

The aggregated raw data in a BIDS format, the trained machine-learning models, experimental and analysis logs, and result dataframes can be found at https://doi.org/https://doi.org/10.58119/ULG/174Q6G.

Code availability

All codes to replicate the power analysis, the experience sampling paradigm, and the present analysis can be found at https://gitlab.uliege.be/Paradeisios.Boulakis/mind_blanking_arousal. An archived version of the code at the time of submission can be found at https://doi.org/https://doi.org/10.58119/ULG/174Q6G.

Received: 11 August 2022; Accepted: 25 November 2024 Published online: 10 February 2025

References

- Christoff, K., Irving, Z. C., Fox, K. C. R., Spreng, R. N. & Andrews-Hanna, J. R. Mind-wandering as spontaneous thought: A dynamic framework. Nat. Rev. Neurosci. 17, 718–731 (2016).
- Van Calster, L., D'Argembeau, A., Salmon, E., Peters, F. & Majerus, S. Fluctuations of attentional networks and default mode network during the resting state reflect variations in cognitive states: Evidence from a novel resting-state experience sampling method. J. Cogn. Neurosci. 29, 95–113 (2017).
- 3. Vanhaudenhuyse, A. et al. Two distinct neuronal networks mediate the awareness of environment and of self. J. Cogn. Neurosci. 23, 570–578 (2011).
- Smallwood, J. & Schooler, J. W. The science of mind wandering: Empirically navigating the stream of consciousness. Annu. Rev. Psychol. 66, 487–518 (2015).
- 5. Ward, A. & Wegner, D. Mind-blanking: When the mind goes away. Front. Psychol. 4, 650 (2013).
- Van den Driessche, C. et al. Attentional lapses in attention-deficit/hyperactivity disorder: Blank rather than wandering thoughts. Psychol. Sci. 28, 1375–1386 (2017).
- Kawagoe, T., Onoda, K. & Yamaguchi, S. Different pre-scanning instructions induce distinct psychological and resting brain states during functional magnetic resonance imaging. *Eur. J. Neurosci.* 47, 77–82 (2018).
- Kawagoe, T., Onoda, K. & Yamaguchi, S. The neural correlates of 'mind blanking': When the mind goes away. *Hum. Brain Mapp.* 40, 4934–4940 (2019).
- 9. Winter, U. et al. Content-free awareness: EEG-fcMRI correlates of consciousness as such in an expert meditator. *Front. Psychol.* **10**, 3064 (2020).
- 10. Andrillon, T. et al. Does the mind wander when the brain takes a break? local sleep in wakefulness, attentional lapses and mindwandering. *Front. Neurosci.* **13**, 949 (2019).
- Andrillon, T., Burns, A., Mackay, T., Windt, J. & Tsuchiya, N. Predicting lapses of attention with sleep-like slow waves. *Nat. Commun.* 12, 3657 (2021).
- 12. Mortaheb, S. et al. Mind blanking is a distinct mental state linked to a recurrent brain profile of globally positive connectivity during ongoing mentation. *Proc. Natl. Acad. Sci. U.S.A* **119**, e2200511119 (2022).
- Schooler, J. W., Reichle, E. D. & Halpern, D. V. Zoning out while reading: evidence for dissociations between experience and metaconsciousness. in *Thinking and seeing: visual metacognition in adults and children* 203–226 (MIT Press, 2004).
- Wong, C. W., Olafsson, V., Tal, O. & Liu, T. T. The amplitude of the resting-state fMRI global signal is related to EEG vigilance measures. *Neuroimage* 83, 983–990 (2013).
- 15. Chang, C. et al. Tracking brain arousal fluctuations with fMRI. Proc. Natl. Acad. Sci. U.S.A 113, 4518–4523 (2016).
- 16. Wong, C. W., Olafsson, V., Tal, O. & Liu, T. T. Anti-correlated networks, global signal regression, and the effects of caffeine in resting-state functional MRI. *Neuroimage* **63**, 356–364 (2012).
- 17. Vyazovskiy, V. V. et al. Local sleep in awake rats. Nature 472, 443-447 (2011).
- 18. Beri, D. & K, J. R. Physiological correlates of arousal: A metaanalytic review. J. Neurol. & Neurosci. 10, (2019).
- 19. Ogilvie, R. D. The process of falling asleep. Sleep Med. Rev. 5, 247-270 (2001).
- Lacey, J. L. Somatic response patterning and stress: some revisions of activation theory. in *Psychological stress* (eds. Appley, M. H. & Trumbell, R.) 14–37 (Appleton-Century-Crofts, 1967).
- 21. Jones, B. E. Arousal systems. Front. Biosci. 8, s438-451 (2003).
- Cantero, J. L., Atienza, M., Salas, R. M. & Gómez, C. M. Alpha EEG coherence in different brain states: An electrophysiological index of the arousal level in human subjects. *Neurosci. Lett.* 271, 167–170 (1999).
- Pinggal, E., Dockree, P. M., O'Connell, R. G., Bellgrove, M. A. & Andrillon, T. Pharmacological manipulations of physiological arousal and sleep-like slow waves modulate sustained attention. J. Neurosci. 42, 8113–8124 (2022).
- 24. Perogamvros, L. et al. The phenomenal contents and neural correlates of spontaneous thoughts across wakefulness, nrem sleep, and rem sleep, *J. Cogn. Neurosci.* 29, 1766–1777 (2017).
- Stawarczyk, D., François, C., Wertz, J. & D'Argembeau, A. Drowsiness or mind-wandering? Fluctuations in ocular parameters during attentional lapses. *Biol. Psychol.* 156, 107950 (2020).
- Unsworth, N. & Robison, M. K. Tracking arousal state and mind wandering with pupillometry. Cogn. Affect. Behav. Neurosci. 18, 638–664 (2018).
- 27. Varela, F. J., Thompson, E. & Rosch, E. The embodied mind: cognitive science and human experience. (MIT Press, 2017).
- Wang, C.-A. *et al.* Arousal effects on pupil size, heart rate, and skin conductance in an emotional face task. *Front. Neurol.* 9, (2018).
 Al, E. et al. Heart-brain interactions shape somatosensory perception and evoked potentials. *Proc. Natl. Acad. Sci. U.S.A* 117, 10575–10584 (2020).
- Park, H.-D., Correia, S., Ducorps, A. & Tallon-Baudry, C. Spontaneous fluctuations in neural responses to heartbeats predict visual detection. Nat. Neurosci. 17, 612–618 (2014).
- Garfinkel, S. N. et al. What the heart forgets: cardiac timing influences memory for words and is modulated by metacognition and interoceptive sensitivity. *Psychophysiol.* 50, 505–512 (2013).

- 32. Garfinkel, S. N. et al. Fear from the heart: sensitivity to fear stimuli depends on individual heartbeats. J. Neurosci. 34, 6573–6582 (2014).
- 33. Allen, M. et al. Unexpected arousal modulates the influence of sensory noise on confidence. eLife 5, e18103 (2016).
- Allen, M., Levy, A., Parr, T. & Friston, K. J. In the body's eye: the computational anatomy of interoceptive inference. *PLoS Comput. Biol.* 18, e1010490 (2022).
- 35. Lee, K. et al. Arousal impacts distributed hubs modulating the integration of brain functional connectivity. *Neuroimage* 258, 119364 (2022).
- Buysse, D. J., Reynolds, C. F., Monk, T. H., Berman, S. R. & Kupfer, D. J. The pittsburgh sleep quality Index: A new instrument for psychiatric practice and research. *Psychiatry Res.* 28, 193–213 (1989).
- Koenig, J., Jarczok, M. N., Fischer, J. É. & Thayer, J. F. The association of (effective and ineffective) analgesic intake, pain interference and heart rate variability in a cross-sectional occupational sample. *Pain Med.* 16, 2261–2270 (2015).
- Spitzer, R. L., Kroenke, K., Williams, J. B. W. & Löwe, B. A brief measure for assessing generalized anxiety disorder: The GAD-7. Arch. Intern. Med. 166, 1092–1097 (2006).
- Siess, J., Blechert, J. & Schmitz, J. Psychophysiological arousal and biased perception of bodily anxiety symptoms in socially anxious children and adolescents: A systematic review. Eur. Child Adolesc. Psychiatry 23, 127–142 (2014).
- 40. Seli, P., Risko, E. F., Smilek, D. & Schacter, D. L. Mind-wandering with and without intention. Trends Cogn. Sci. 20, 605-617 (2016).
- 41. Seli, P. et al. Mind-wandering as a natural kind: A family-resemblances view. Trends Cogn. Sci. 22, 479-490 (2018).
- 42. Smallwood, J. Distinguishing how from why the mind wanders: A process-occurrence framework for self-generated mental activity. *Psychol. Bull.* 139, 519–535 (2013).
- Fox, K. C. R., Spreng, R. N., Ellamil, M., Andrews-Hanna, J. R. & Christoff, K. The wandering brain: Meta-analysis of functional neuroimaging studies of mind-wandering and related spontaneous thought processes. *Neuroimage* 111, 611–621 (2015).
- 44. Smallwood, J. et al. The neural correlates of ongoing conscious thought. iScience 24, 102132 (2021).
- 45. Gourine, A. V. & Ackland, G. L. Cardiac vagus and exercise. Physiology 34, 71-80 (2019).
- Lin, M.-A. et al. Resistance-induced brain activity changes during cycle ergometer exercises. BMC Sports Sci. Med. Rehabil. 13, 27 (2021).
- Basso, J. C. & Suzuki, W. A. The effects of acute exercise on mood, cognition, neurophysiology, and neurochemical pathways: a review. *Brain Plast.* 2, 127–152 (2017).
- Gorgoni, M. et al. Topographic electroencephalogram changes associated with psychomotor vigilance task performance after sleep deprivation. Sleep Med. 15, 1132–1139 (2014).
- 49. Borbély, A. A., Daan, S., Wirz-Justice, A. & Deboer, T. The two-process model of sleep regulation: a reappraisal. J. Sleep Res. 25, 131–143 (2016).
- Lazar, A. S., Lazar, Z. I. & Dijk, D.-J. Circadian regulation of slow waves in human sleep: Topographical aspects. Neuroimage 116, 123–134 (2015).
- 51. Birchler-Pedross, A. et al. Subjective well-being is modulated by circadian phase, sleep pressure, age, and gender. J. Biol. Rhythms 24, 232–242 (2009).
- 52. Sagaspe, P. et al. Influence of age, circadian and homeostatic processes on inhibitory motor control: A go/nogo task study. *PLOS ONE* 7, e39410 (2012).
- Cajochen, C. et al. Dynamics of frontal low EEG-activity and subjective sleepiness under high and low sleep pressure. Sleep 24, A77–A77 (2001).
- 54. Wu, J. et al. Decreased resting-state alpha-band activation and functional connectivity after sleep deprivation. Sci. Rep. 11, 484 (2021).
- Glos, M., Fietze, I., Blau, A., Baumann, G. & Penzel, T. Cardiac autonomic modulation and sleepiness: physiological consequences of sleep deprivation due to 40h of prolonged wakefulness. *Physiol. Behav.* 125, 45–53 (2014).
- 56. Liang, K.-Y. & Zeger, S. L. Longitudinal data analysis using generalized linear models. Biometrika 73, 13-22 (1986).
- 57. Fitzmaurice, G. M., Laird, N. M. & Ware, J. H. Applied longitudinal analysis. (Wiley, 2011).
- Lo, S. & Andrews, S. To transform or not to transform: Using generalized linear mixed models to analyse reaction time data. Front. Psychol. 6, 1171 (2015).
- King, J.-R. et al. Information sharing in the brain indexes consciousness in noncommunicative patients. *Curr. Biol.* 23, 1914–1919 (2013).
- 60. Imperatori, L. S. et al. EEG functional connectivity metrics wPLI and wSMI account for distinct types of brain functional interactions. Sci. Rep. 9, 8894 (2019).
- 61. Benedek, M. & Kaernbach, C. A continuous measure of phasic electrodermal activity. J. Neurosci. Methods 190, 80-91 (2010).
- 62. Hamdan, S. et al. Julearn: An easy-to-use library for leakage-free evaluation and inspection of ML models. *GigaByte* 2024, gigabyte113 (2024).
- 63. Pedregosa, F. et al. Scikit-learn: Machine learning in python. J Mach. Learn. Res. 12, 2825–2830 (2011).
- 64. Gramfort, A. et al. MEG and EEG data analysis with MNE-Python. Front. Neurosci. 7, (2013).
- Engemann, D. A. et al. Robust EEG-based cross-site and cross-protocol classification of states of consciousness. *Brain* 141, 3179– 3192 (2018).
- 66. Makowski, D. et al. NeuroKit2: A python toolbox for neurophysiological signal processing. *Behav. Res. Methods* 53, 1689–1696 (2021).
- Hudson, A. N., Van Dongen, H. P. A. & Honn, K. A. Sleep deprivation, vigilant attention, and brain function: A review. Neuropsychopharmacol. 45, 21–30 (2020).
- 68. Csipo, T. et al. Sleep deprivation impairs cognitive performance, alters task-associated cerebral blood flow and decreases cortical neurovascular coupling-related hemodynamic responses. *Sci. Rep.* **11**, 20994 (2021).
- 69. Krause, A. J. et al. The sleep-deprived human brain. Nat. Rev. Neurosci. 18, 404-418 (2017).
- 70. Qi, J.-L. et al. The effects of 43 hours of sleep deprivation on executive control functions: Event-related potentials in a visual go/no go task. *Soc. Behav. Pers.* **38**, 29–42 (2010).
- Gosselin, A., De Koninck, J. & Campbell, K. B. Disentangling specific inhibitory versus general decision-making processes during sleep deprivation using a go/nogo erp paradigm. *Int. J. Psychophysiol.* 141, 18–27 (2019).
- Poh, J.-H., Chong, P. L. H. & Chee, M. W. L. Sleepless night, restless mind: Effects of sleep deprivation on mind wandering. J. Exp. Psychol. Gen. 145, 1312–1318 (2016).
- 73. Carciofo, R., Du, F., Song, N. & Zhang, K. Mind wandering, sleep quality, affect and chronotype: An exploratory study. *PLOS ONE* 9, e91285 (2014).
- Jubera-Garcia, E., Gevers, W. & Van Opstal, F. Local build-up of sleep pressure could trigger mind wandering: Evidence from sleep, circadian and mind wandering research. *Biochem. Pharmacol.* 191, 114478 (2021).
- 75. Lim, J. & Dinges, D. F. Sleep deprivation and vigilant attention. Ann. N. Y. Acad. Sci. 1129, 305-322 (2008).
- Boulakis, P. A., Mortaheb, S., van Calster, L., Majerus, S. & Demertzi, A. Whole-brain deactivations precede uninduced mindblanking reports. J. Neurosci. 43, 6807–6815 (2023).
- 77. Demertzi, A. et al. Human consciousness is supported by dynamic complex patterns of brain signal coordination. *Sci. Adv.* **5**, 7603 (2019).
- 78. El-Baba, M. et al. Functional connectivity dynamics slow with descent from wakefulness to sleep. PLOS ONE 14, e0224669 (2019).

- 79. Lin, A., Liu, K. K. L., Bartsch, R. P. & Ivanov, P. C. Delay-correlation landscape reveals characteristic time delays of brain rhythms and heart interactions. *Philos. Trans. A. Math. Phys. Eng. Sci.* **374**, 20150182 (2016).
- Bartsch, R. P., Liu, K. K., Ma, Q. D. & Ivanov, P. C. Three independent forms of cardio-respiratory coupling: Transitions across sleep stages. Comput. Cardiol. 41, 781–784 (2014).
- Raimondo, F. et al. Brain-heart interactions reveal consciousness in noncommunicating patients. *Ann. Neurol.* 82, 578–591 (2017).
 Candia-Rivera, D., Catrambone, V., Barbieri, R. & Valenza, G. Functional assessment of bidirectional cortical and peripheral
- neural control on heartbeat dynamics: A brain-heart study on thermal stress. *Neuroimage* **251**, 119023 (2022). 83. Mittner, M., Hawkins, G. E., Boekel, W. & Forstmann, B. U. A neural model of mind wandering. *Trends. Cogn. Sci.* **20**, 570–578
- (2016). 84. Unsworth, N. & Robison, M. K. Pupillary correlates of lapses of sustained attention. *Cogn. Affect. Behav. Neurosci.* 16, 601–615
- (2016).85. Iwamoto, M. et al. Respiratory entrainment of the locus coeruleus modulates arousal level to avoid physical risks from external vibration. *Sci. Rep.* 13, 7069 (2023).
- Bauvergne, C. et al. Are locus coeruleus neurons involved in blinking?. *Neurosci. Res.* 61, 182–191 (2008).
- 87. Park, H.-D. et al. Breathing is coupled with voluntary initiation of mental imagery. Neuroimage 264, 119685 (2022).
- Tian, Y. E. et al. Evaluation of brain-body health in individuals with common neuropsychiatric disorders. JAMA Psychiatry 80, 567-576 (2023).
- Bashan, A., Bartsch, R. P., Kantelhardt, J. W., Havlin, S. & Ivanov, P. C. Network physiology reveals relations between network topology and physiological function. *Nat. Commun.* 3, 702 (2012).

Acknowledgements

The experimental work was conducted at the GIGA-CRC-Human Imaging platform of ULiège, Belgium.

Author contributions

Paradeisios Alexandros Boulakis: conceptualization; data curation; formal analysis; investigation; methodology; project administration; software; visualization; writing—original draft preparation. Nicholas John Simos: investigation; software; validation; writing—review and editing. Zoi Stefania: investigation; project administration; writing—review and editing. Sepehr Mortaheb: formal analysis; software; writing—review and editing. Christina Schmidt: methodology; writing—review and editing. Federico Raimondo: formal analysis; methodology; software; validation; supervision; writing—review and editing. Athena Demertzi: conceptualization; methodology; supervision; writing—original draft preparation.

Funding

At the time of the research, PAB and SM were FNRS Research Fellows. AD and CS are FNRS Research Associates. This work was supported by the Belgian Fund for Scientific Research (FRS-FNRS), the European Union's Horizon 2020 Research and Innovation Marie Skłodowska-Curie RISE programme NeuronsXnets (Grant agreement 101007926), the European Cooperation in Science and Technology COST Action (CA18106), the Léon Fredericq Foundation, and the University and of University Hospital of Liège. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Declarations

Competing interests

The authors declare no competing interests.

Ethical approval

The experimental procedure has been approved by the CHU Liège local ethics committee and conforms with the Declaration of Helsinki and the European General Data Protection Regulation (GDPR). Before the onset of the protocol, participants provided informed consent for their participation in the study. Participants also received monetary compensation for their participation in the study.

Protocol registration

The stage 1 accepted-in-principle protocol can be found at https://osf.io/sh2ye. The authors confirm that no data for the pre-registered study was collected prior to the date of AIP.

Additional information

Supplementary Information The online version contains supplementary material available at https://doi.org/1 0.1038/s41598-024-81618-1.

Correspondence and requests for materials should be addressed to A.D.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License, which permits any non-commercial use, sharing, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if you modified the licensed material. You do not have permission under this licence to share adapted material derived from this article or parts of it. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by-nc-nd/4.0/.

© The Author(s) 2025