INTRODUCTION

The use of spontaneous resting electroencephalographic (EEG) activity as an objective measure of individual differences in psychological functioning has a long history (Davidson, 1984; Klimesch, 1999; Knyazev, 2007; Schutter & Knyazev, 2012). Resting EEG has typically been divided into standardized bands based on canonical (i.e., fixed)
spectral power bands, from the slowest frequencies in delta (0–4 Hz), through theta (4–7 Hz), alpha (7–13 Hz), beta (13–30 Hz), to the fastest frequencies in gamma (30+ Hz), but are also sometimes defined on an individual basis from the alpha peak frequency (Babiloni et al., 2020). A persistent but growing focus is on the use of these resting EEG measures (usually quantified from canonical bands) as biomarkers for optimal and suboptimal executive function, particularly in the context of identifying healthy versus unhealthy executive function development and decline with aging (Arns et al., 2013; Babiloni et al., 2006). Recent research suggests that individual and intraindividual differences in executive function broadly, and attentional control specifically, are associated with the ratio between fronto-central standard theta to beta oscillations, purported to represent differences in periodic activity within these frequency bands (Angelidis et al., 2016).

1.1 | Theta-beta ratios

The ratio between fronto-central theta and beta oscillations has been proposed as a marker of executive [dys]function associated with Attention-Deficit Hyperactive Disorder (ADHD; Arns et al., 2013) where higher ratios—indicating relatively greater fronto-central theta than fronto-central beta—are characteristic of ADHD. In nonclinical populations, theta-beta ratio has been found to be strongly negatively correlated with self-report and behavioral measures of executive function, while theta and beta alone are not (Perone et al., 2018), and smaller theta-beta ratios are indicative of better cognitive control, executive control, and increased vigilance (Angelidis et al., 2016, 2018; Putman et al., 2010, 2014; van Son, Schalbroeck, et al., 2018). Several studies have shown that beyond being a marker of attentional control, theta-beta ratios are negatively correlated with response-inhibition by threatening stimuli (Putman et al., 2010) and are positively correlated with attentional capture by mildly threatening stimuli relative to highly threatening stimuli (van Son, Angelidis, et al., 2018). Theta-beta ratios have also been found to negatively correlate with advantageous decision-making in reinforcement learning paradigms (Massar et al., 2014; Schutter & Van Honk, 2005). Other studies have reported that theta-beta ratios are positively associated with risk taking behavior, and that theta and beta alone do not (Massar et al., 2012). Recent studies have also shown that theta-beta ratios increase during mind-wandering (van Son, De Blasio, et al., 2019; van Son, de Rover, et al., 2019). Together, these studies suggest that theta-beta ratios are related to attentional control broadly, as well as to more specific emotional and rewarding contexts, such that lower theta-beta ratios are reflective of more control or focus.

Theta-beta ratios are argued to reflect the reciprocal regulation of bottom-up subcortical processes by top-down cortical processes (Knyazev, 2007; Schutter & Knyazev, 2012). Although much of the research supporting this subcortical–cortical model of theta-beta ratios and executive function has been indirect, a recent study has provided support for involvement of cortical networks (van Son, de Rover, et al., 2019). Specifically, van Son, de Rover, et al. (2019) showed that not only are theta-beta ratios lower when participants exert attentional control compared to when they engage in mind-wandering, but that these changes are associated with decreased functional connectivity between dorsolateral prefrontal cortex (DLPFC) and the dorsal anterior cingulate cortex (ACC)—regions which have been associated with executive function in multiple domains (Seeley et al., 2007).

Given age-related decline in executive function (Buckner, 2004; Lustig & Jantz, 2015) and in cortical integrity (Fjell et al., 2017; Madden et al., 2009, 2012), one would assume a straightforward relationship between theta-beta ratio and age. The mapping between age and the theta-beta ratio is more complicated, however. First, theta has been observed to increase with age, potentially through a migration of alpha activity to the upper frequencies of theta, discussed in more detail below (Klimesch, 1999). Second, resting theta recorded from the same fronto-central scalp locations used in theta-beta ratio research has been positively correlated with cognitive function in older adults (73 adults ages 56–70, Finnigan & Robertson, 2011; 53 adults ages 18–89, Vlahou et al., 2014). Third, in child and young adult samples, theta-beta ratios have been reported to be negatively correlated with age (41 young adults ages 18–31 years, Angelidis et al., 2016; 41 children ages 8–12 years, Clarke et al., 2001; 101 children ages 7–16 years, Ogrim et al., 2012; 162 children ages 3–9 years, Perone et al., 2018; 28 young adults ages 19–28 years, Putman et al., 2010, but see Putman et al., 2014 for a non-replication in 77 young adults with a mean age of 19.9 years), and with cognitive function (Angelidis et al., 2016, 2018; Putman et al., 2010, 2014; 128 young adults with a mean age of 22.3 years, Schutte et al., 2017). Thus, it remains unclear what drives any association between age and theta-beta ratio, complicated in part by the reliance on canonical frequency bands for calculating theta-beta ratio in populations with shifts in individual alpha peak frequencies in the existing literature.

1.2 | Individual alpha peak frequencies

The frequency at which power in the alpha band (7–13 Hz) peaks, known as individual alpha peak frequency, has
been found to be negatively correlated with age in adulthood (Clark et al., 2004; Klimesch, 1997), and is reduced in individuals with Alzheimer’s disease (Klimesch, 1997). Higher individual alpha peak frequency across adulthood is associated with better working memory, better reading comprehension, and a larger general intelligence factor (Angelakis et al., 2004; Clark et al., 2004; Grandy et al., 2013; Klimesch, 1997), suggesting it is an indicator of cognitive capacity or preparedness.

Greater alpha power has been associated with reductions in blood flow across wide areas of the frontal and parietal cortex (Jensen & Mazaheri, 2010; Laufs et al., 2003). During working memory tasks, greater local alpha power during a trial was associated with better memory performance (Jensen & Mazaheri, 2010), and decreases in BOLD activation, particularly in areas of the default mode network (Anticevic et al., 2010; Daselaar et al., 2004). Together, these findings suggest that alpha power indexes the ability to inhibit task-irrelevant regions while performing cognitive tasks (Jensen & Mazaheri, 2010), processes which tend to decline with advancing age.

With regards to theta-beta ratios, because the peak alpha frequency is found in lower frequencies with age, some of the EEG power associated with the alpha band (commonly defined as 7–13 Hz) may be mistakenly attributed to power in the canonical theta band (commonly defined as 4–7 Hz) in older adults, driving increases in canonical theta band power with age and therefore changes in theta-beta ratios. While some work suggests that relative canonical theta in older adults may be positively correlated with measures of memory, attention, and executive functioning, the potential role of alpha leaking into the canonical theta band, as indexed by individual peak alpha frequency, remains unclear (Finnigan & Robertson, 2011). Therefore, understanding how resting canonical theta band power, canonical beta band power, theta-beta ratios calculated from canonical theta and beta band power, and individual peak alpha frequencies are cross-sectionally interrelated across the adult age-span is a key to beginning to understand how these EEG metrics may relate to healthy aging, particularly since the existing literature on theta-beta ratios in age relies on canonical power band definitions.

1.3 Aperiodic and periodic neural activity

The association between traditional EEG metrics and healthy aging is further complicated by recent observations that frequency band measures of periodic activity are influenced by aperiodic activity present across all frequencies (Donoghue, Dominguez, & Voytek, 2020; Donoghue, Haller, et al., 2020; Keil et al., 2022; Voytek et al., 2015). In initial conceptualizations of aperiodic activity, steeper spectra were interpreted as indicating greater synchronization; flatter spectra were interpreted as indicating reduced synchronization (i.e., greater neural noise; Voytek & Knight, 2015). More recent data suggest that the slope of the EEG spectra is related to the ratio of excitatory to inhibitory neural activity, while the height of the spectra is related to neural spiking rates (Donoghue, Haller, et al., 2020; Waschke et al., 2021). Greater excitatory to inhibitory activity is reflected in flatter spectra (Gao et al., 2017), and greater spiking activity is reflected in greater overall spectral power (Manning et al., 2009; Miller et al., 2012). Aperiodic activity—in particular the slope of the spectra—has been associated with age, mediates cross-sectional associations between age and cognitive function (Voytek et al., 2015), and is associated with physiological markers of cognitive decline (Tran et al., 2020) and processing speed (Ouyang et al., 2020).

The impact of aperiodic activity on EEG metrics is particularly pronounced for theta-beta ratios. For example, the association between the exponent of aperiodic activity (i.e., the gradient of the spectra slope) and the theta-beta ratio has been found to be significantly stronger than the association between the periodic measures of both theta and beta (Donoghue, Dominguez, & Voytek, 2020). Beyond suggesting that measures of both theta and beta are severely confounded by aperiodic activity, the strong association between theta-beta ratios and aperiodic activity found in prior work implies that the individual differences in the theta-beta ratio may primarily reflect individual differences in excitatory to inhibitory neural activity (Donoghue, Dominguez, & Voytek, 2020). Furthermore, in the case of the lack of definable peaks within a given power band, it is ambiguous whether group or individual differences are due to changes in periodic power or instead the aperiodic component.

1.4 The present study

Theta-beta ratios have been found to be negatively associated with age, such that larger ratios—indicating relatively greater theta power than beta power—are observed in samples of younger participants compared to older participants. However, child and young adult samples with restricted age ranges (e.g., children 3–9 years old in Perone et al., 2018; young adults 19–28 years old in Putman et al., 2010) that rely upon canonical band definitions, currently predominate the studies of age-related differences in theta-beta ratios. In the present study, we extend the research on fronto-central theta-beta ratios (calculated from canonical bands) by examining whether the negative association between age and
canonical fronto-central theta-beta ratios is observed in a large sample featuring a wide adult age range (from 36 to 84 years). Additionally, we examine to what extent any associations between canonical fronto-central theta-beta ratios and age are accounted for by age-related differences in fronto-central individual alpha peak frequency and in the fronto-central aperiodic (1/f-like) component of the neural power spectrum. Finally, we examine the unique associations between age and canonical fronto-central theta band power and canonical fronto-central beta band power individually and controlling for fronto-central individual alpha peak frequency and the fronto-central aperiodic component.

We conducted a partially preregistered secondary analysis of data from the Midlife in the US Study’s Neuroscience Project (see http://midus.wisc.edu/ and Ryff et al. 2021 for additional details), examining whether the correlation between canonical fronto-central theta-beta ratios and age is due to variation in associations between age and canonical fronto-central theta, age and canonical fronto-central beta, age-related decreases in individual peak alpha frequencies, or age-related flattening of the aperiodic component. Based on previous studies, we developed and tested two preregistered hypotheses. First, we tested whether the negative association between canonical fronto-central theta-beta ratios and age is replicated in a large sample of older adults ranging in age from 36 to 84 years old, hypothesizing that greater age will be associated with lower canonical fronto-central theta-beta ratios (Table 1, confirmatory hypothesis 1). Second, we used the RestingIAF package (https://github.com/corcoran/restingIAF; Corcoran et al., 2018) to test whether there would be a pattern of fronto-central alpha peak frequency “slowing” with age, predicting that older age would be associated with lower fronto-central individual alpha peak frequencies (Table 1, confirmatory hypothesis 2). We examined whether the association between canonical fronto-central theta-beta ratios and age was preserved when statistically adjusting for individual differences in fronto-central individual alpha peak frequencies, and examined if the relationships held controlling for gender and race. We explored the relationship between the aperiodic 1/f-like component of the power spectrum and age, and if the association between canonical fronto-central theta-beta ratios and age was preserved when statistically adjusting for changes in the aperiodic component. Additionally, we examined if fronto-central individual alpha peak frequencies or fronto-central aperiodic component could mediate the relationship between canonical fronto-central theta-beta ratios and age. Finally, we examined the extent to which canonical fronto-central theta power and canonical fronto-central beta power are uniquely associated with age at time of recording (Table 1, additional hypothesis E1). Additional analyses are included in the supplemental materials to ensure the findings are not specific to analytical choices or specific EEG metric quantification methods described in the main manuscript.

Analyses and hypotheses regarding canonical fronto-central theta-beta ratio, fronto-central individual alpha peak frequency, and age (including the specific fronto-central ROI) were preregistered prior to the extraction of new EEG frequency metrics and their statistical analysis at https://osf.io/n57au. Additionally, the new EEG reprocessing pipeline for the extraction of canonical fronto-central theta power, canonical fronto-central beta power, and individual alpha peak frequencies was registered separately at https://osf.io/wfkmn. See Table 1 for a summary of the preregistration status of all analyses and exclusion criteria.

2  |  METHOD

2.1  |  Sensitivity power analysis

We used G*Power 3.1 (Faul et al., 2009) to conduct sensitivity power analysis prior to data processing for a sample of 300 participants as an estimate for the final usable sample size after applying our criteria for usable EEG data. This analysis indicated that we would have 95% power to detect a Pearson’s correlation of 0.20, and 95% power to detect a small to medium sized effect in regression analyses ($f^2 = 0.06$).

2.2  |  Participants

The present study used data collected during the second wave of Midlife in the US (MIDUS) in the Neuroscience Project (2004–2009), consisting of 331 participants from the main MIDUS cohort. These respondents included three distinct subsamples: the Main Longitudinal (n = 135), Twin (n = 88), and Milwaukee (n = 108) subsamples (see http://midus.wisc.edu/midus2/project5/ for additional details about sampling strategies within these subsamples). The Main Longitudinal and Twin subsamples...
Table 1  Summary of preregistration status of analyses (from https://osf.io/n57au) and exclusion criteria

| Analysis | Preregistration status |
|----------|------------------------|
| Section 3.1: Pearson’s pairwise correlations | **Preregistered:** Age with canonical fronto-central theta-beta ratio, with and without controlling for gender Age with RestingIAF defined fronto-central individual alpha peak frequencies, with and without controlling for gender **Post hoc, determined to be a useful additional analytic strategy:** Partial correlations controlling for gender and race Report all other additional pairwise correlation combinations and descriptive statistics **Post hoc, reviewer suggested:** Age with FOOOF defined fronto-central aperiodic exponent and offset Canonical fronto-central theta-beta ratio with FOOOF defined fronto-central aperiodic exponent and offset RestingIAF defined fronto-central individual alpha peak frequencies with FOOOF defined fronto-central aperiodic exponent and offset |
| Section 3.2: Partial correlations | **Implied in background of preregistration:** Due to author oversight, analyses regarding controlling for RestingIAF defined fronto-central individual alpha peak frequencies are implied in the preregistration introduction but not explicitly outlined **Post hoc, reviewer suggested:** Age with canonical fronto-central theta-beta ratio, controlling separately for fronto-central aperiodic exponent and offset |
| Section 3.3: Mediational analyses | **Post hoc, determined to be a useful additional analytic strategy:** Relationship between age and canonical fronto-central theta-beta ratio mediated by RestingIAF defined fronto-central individual alpha peak, FOOOF defined aperiodic offset, and/or FOOOF defined aperiodic exponent |
| Section 3.4: Hierarchical regression analyses | **Preregistered:** Step 1 (canonical fronto-central theta and canonical fronto-central beta regressed on age) and Step 2 (RestingIAF defined fronto-central individual peak alpha added) **Post hoc, reviewer suggested:** Step 3 (FOOOF defined aperiodic exponent) |
| Supplemental material, S2: General estimating equation analysis | **Preregistered:** GEE analyses repeating main correlational analyses, controlling for genetic dependencies within family |
| Supplemental material, S3: Explore nonlinear age and theta-beta ratio relationships | **Preregistered:** Quadratic regression of canonical fronto-central theta-beta ratio with age |
| Supplemental material, sections S4-S10 | **Post hoc, reviewer suggested:** Repeat analyses on eyes closed only, with different theta-beta ratio and individual alpha peak specification methods |

Preregistered exclusion criteria (reproduced from https://osf.io/n57au and https://osf.io/wfkmn)

1. 50% epochs retained for spectral power density metrics
2. 50% of channels resulting in definable alpha peaks

Post hoc exclusion criteria

1. Poor FOOOF model fit, defined as less than three standard deviations below the mean in $R^2$ model fit for the fronto-central composite

Table 1 contains individuals who participated in the initial wave of MIDUS data collection approximately 10 years prior. The Milwaukee subsample contained individuals who participated in the baseline MIDUS Milwaukee study initiated in 2005. Demographic information is presented in Table 2.

All data collection procedures were approved by the UW-Madison Institutional Review Board, and informed consent was obtained for all participants. Participants with unusable resting spectral power EEG data ($n = 12, 3.6\%$), without identifiable alpha peak frequencies ($n = 48, 14.5\%$), and/or without adequate FOOOF model fit ($n = 9$, 3.0\%) were excluded.
2.3% were excluded from analyses, yielding a final sample of \( n = 268 \) participants.

### Materials

#### Demographics

Demographic variables are publicly available via Colectica ([http://midus.colectica.org/](http://midus.colectica.org/)) and the Inter-university Consortium for Political and Social Research (ICPSR; [https://www.icpsr.umich.edu/web/ICPSR/series/203](https://www.icpsr.umich.edu/web/ICPSR/series/203)). From the MIDUS 2 Neuroscience Project data set, we used age at time of EEG data collection, gender, race (dichotomized as White/Black, Indigenous, and People of Color [BIPOC] for analyses), and Family ID. Family ID was used to account for genetic dependencies in follow-up analyses in the supplemental materials. See Table 2 for a breakdown of demographics.

#### Procedure

### EEG recording

EEG data were collected using a 128 channel geodesic net of Ag/AgCl electrodes in the GSN200 montage (see pre-processing preregistration figure, [https://osf.io/wfknn](https://osf.io/wfknn); Electrical Geodesics, Inc, 2007) encased in saline dampened sponges (Electrical Geodesics, Inc [EGI], Eugene, OR) with impedances reduced to less than 100 KΩ while ensuring that electrolyte “bridges” (see Greischar et al., 2004) had not formed. After the net was placed, participants were escorted into a soundproof booth where
they were seated in front of a computer screen. A computer located outside the booth recorded the data. Signals were amplified and sampled at 500 Hz with an online bandpass filter from 0.1 to 100 Hz at 16-bit precision using an online vertex (Cz) reference. The participant was instructed to rest for six 1-min periods. During three of the 1-minute periods they were asked to keep their eyes open; for the remaining three 1-min periods they were asked to keep their eyes closed. The order of the eyes open/eyes closed was pseudorandomized, with two fixed orders counterbalanced across participants. Participants then completed an emotional picture viewing task (data not presented here), followed by another baseline resting recording for six 1-min periods. Prior data processing was restricted to alpha asymmetry variables from the first baseline recording, collapsed across participants. Participants then completed an emotional picture viewing task (data not presented here), followed by another baseline resting recording for six 1-min periods. Prior data processing was restricted to alpha asymmetry variables from the first baseline recording, collapsed across the entire 6-minute period (e.g., Hostinar et al., 2017). The current analyses focus on metrics extracted from this first resting recording, collapsed across eyes open, and eyes closed periods. Additional analyses examining eyes closed only epochs are available in the supplemental materials and do not change the interpretations of the analyses. Raw continuous data as well as the summary metrics described below in sections 2.5.2, 2.5.3, and 2.5.4 are available upon request. See https://osf.io/fgnxnt/ for information about how to access the data used in the following analyses.

2.5 | Data reduction

2.5.1 | EEG preprocessing

Offline the EEG data were filtered (60 Hz notch, 0.5 Hz high-pass), bad channels identified and removed, and bad sections of data identified and removed. EEGLab6 was originally used to conduct a PCA/ICA to identify 20 components (such that PCA was first applied for the reduction constrained to 20 components, followed by an ICA for the differentiation of components), which were visually inspected to identify components to remove obvious blink, eye movement, and other artifacts. No further PCA or ICA dimension reduction was conducted after artifactual components were removed. Bad channels were replaced using a spherical spline interpolation. These are the original preprocessing steps from the initial alpha asymmetry pipeline that were preserved in the reprocessing pipeline, detailed in Ryff et al. (2021). Data from the eyes open and eyes closed conditions were collapsed.5 The fronto-central ROI was preregistered to comprise of the average composite of the F3/Fz/F4 analog channels.6

2.5.2 | Spectral power for canonical fronto-central theta-beta ratio

Data processing for spectral power for canonical fronto-central theta-beta ratios was completed using EEGLab 2019.1 scripts implemented in MATLAB 2019b. Data were re-referenced to the average reference and Cz was imputed. Continuous resting data were epoched into 2 s segments with 50% overlap, and bad segments were rejected if there was a voltage deviation on any channel of ±100 μV. As preregistered, participants with more than 50% of the total number of epochs rejected were excluded from analyses in a list-wise fashion (n = 12). EEG spectral power at each predefined canonical spectral band (theta: 4–7 Hz; beta: 13–30 Hz) was extracted using a 2 s Hamming window padded by a factor of 2 with 50% overlap. Spectral power was extracted individually for each channel, then averaged over the fronto-central composite ROI and were transformed to a theta-beta ratio by dividing the former by the latter and subsequently log-normalized.7

2.5.3 | EEG reprocessing: Individual alpha peak frequency

Fronto-central individual alpha peak frequency from the initial baseline recording was extracted using the RestingIAF package (https://github.com/corcorana/restingIAF; Corcoran et al., 2018), using adjustments to the parameters based on our sample of older adults as recommended by Corcoran et al. (2018). The RestingIAF package algorithmically identifies the peak activity within the alpha band using the Savitzky–Golay filter (SGF), a nonparametric curve fitting technique, whereby the PSD estimates are smoothed using the SGF before estimating the first and second order derivatives. These derivatives are then used to identify a spectral peak, and the first

5Alternative analyses on just the eyes closed conditions are available in the supplemental materials and do not change interpretations.

6As shown in the reprocessing pipeline registration (https://osf.io/wkmmn), the fronto-central composite of F3/Fz/F4 was comprised of the EGI GSN200 electrode montage (Electrical Geodesics, Inc, 2007) sensors 12, 20, 21, 25, 29 (comprising the analog for F3), sensors 4, 5, 118, 119, 124 (comprising the analog for F4), and sensor 11 (comprising the analog for Fz), and was selected based on existing theta-beta ratio literature. See Keil et al. (2022) for a discussion of the importance of preregistering ROIs in frequency-based EEG analyses.

7Alternative analyses using individually defined theta and beta bands based on individual alpha peak frequency to create the theta-beta ratio are available in the supplemental materials and do not change interpretations.
derivative is additionally used to identify the individual alpha band windows based upon where the “shoulders” of the alpha peak are located (see Corcoran et al., 2018 for additional information). We used a 2 s Hamming window with 50% overlap, as well as the following RestingIAF algorithm settings: $F_w = 11$ (SGF frame width, with larger numbers indicating more smoothing, results in a frequency span of $\sim 2.69$ Hz); $k = 4$ (SGF polynomial degree, higher values result in less smoothing and less peak height attenuation); $W_a = [6, 14]$ Hz (the frequency domain within which evidence for peak activity was searched); $f_{Range} = [1, 40]$ Hz (range of frequencies used to fit the algorithm), $mpow = 0.6$ (the minimum power value a local maximum needed to exceed to qualify as a peak candidate), $pDiff = 0.20$ (the minimum proportion of peak height by which the highest peak candidate had to exceed other peaks within the search window $W_a$ to be assigned as the alpha peak frequency), $cMin = 3$ (minimum number of channel estimates necessary for returning results). Estimates were extracted individually for each channel, then averaged over the fronto-central composite ROI. As preregistered, individuals who did not exhibit a definable fronto-central individual alpha peak value in 50% of the sensors used for the composites or 50% of the overall scalp were excluded from analyses in a list-wise fashion ($n = 55$). Fronto-central individual alpha peak frequency for the current study was quantified as the average composite of the F3/Fz/F4 analog channels to assess the impact of age-related differences in individual peak alpha at fronto-central sites on canonical theta-beta ratio measured at the same fronto-central sites.

### 2.5.4 Modeling periodic and aperiodic power spectrum components

Spectral power density was extracted individually for each channel using a 2 s Hamming window padded by a factor of 2 with 50% overlap using EEGLab 2019.1 scripts implemented in MATLAB 2019b from 0 to 250 Hz in 0.25 Hz increments for all sensors then analyzed using FOOOF 1.0.0 (Donoghue, Haller, et al., 2020; https://foof-tools.github.io/) in Python (version 3.9) to fit aperiodic and periodic components from 2 to 40 Hz.

FOOOF algorithmically fits a model to estimate both the aperiodic (1/f-like) component of EEG spectral power density as well as overlying periodic oscillatory “peaks”, by first fitting an aperiodic component (modeled after a Lorentzian function) with a specific offset value (corresponding to the y-intercept of the aperiodic component) and exponent (corresponding to the “flatness” of the 1/f curve, equivalent to the sign-flipped slope of a linear fit in log-log space), which is then regressed out of the PSD, leaving behind periodic peaks. These peaks are then iteratively modeled by fitting a Gaussian around the central frequency of each peak, until the maximum number of peaks fit is reached or no more peaks meeting the algorithm’s criteria are available (see Donoghue, Haller, et al., 2020 for additional details). Periodic and aperiodic components were estimated from the PSD ranging from 2 to 40 Hz, without a knee, with peaks limited in width from 1–6 Hz, a minimum peak height of 0.05, a relative peak threshold of 1.5 standard deviations, and a maximum number of six peaks fit. The resulting models were defined as having poor fit if they were less than three standard deviations below the mean in $R^2$ model fit for the fronto-central composite, resulting in $n = 9$ failing to meet the $R^2 = 0.862$ threshold and were excluded in a list-wise fashion. Finally, the aperiodic offset and exponent values for the frontal F3/Fz/F4 composite were extracted. Details regarding additional alternative EEG metrics are discussed in the supplemental materials.

### 3 RESULTS

As our preregistered analyses focused on metrics extracted from the resting data collapsed across eyes open and eyes closed periods, we report all analyses below on metrics extracted from the combined recordings. Parallel analyses were conducted on alternative EEG metrics are reported in the supplemental materials (see Supplemental Materials section S3). Overall, the pattern of results remained the same regardless of the choice of EEG metric quantification (e.g., canonical and individual band power, metrics extracted from eyes closed only data, etc.). All statistical analyses were performed in R (version 4.1.2). RMarkdown scripts and output for all analyses reported below and in the supplemental materials are available at https://osf.io/hdrax/. See Table 1 for a breakdown of which analyses were preregistered. We first report pairwise correlational analyses, including our two preregistered analyses regarding our hypotheses that: (1) resting canonical fronto-central theta-beta ratios will be inversely associated with age, and (2) fronto-central individual alpha peak frequencies will be inversely associated with age. Next, we report

As detailed in the supplemental materials, use of the FOOOF defined individual alpha peak frequency instead of the RestingIAF defined individual alpha peak frequency did not change the analyses. Additionally, it was not possible to create a metric of aperiodic adjusted canonical theta-beta ratio using FOOOF defined aperiodic adjusted canonical fronto-central theta power and aperiodic adjusted canonical fronto-central beta power, as only $n = 42$ participants had a definable aperiodic adjusted canonical fronto-central theta peak. This resulted in a severe floor effect, with $n = 229$ participants with a FOOOF-derived aperiodic adjusted canonical fronto-central theta-beta ratio of zero.
our exploratory analyses examining partial correlations between canonical fronto-central theta-beta ratio and age controlling for fronto-central individual alpha peak frequency and the fronto-central aperiodic 1/f component. Then, we explore the extent to which fronto-central individual alpha peak frequencies and the fronto-central aperiodic component metrics mediate the relationship between canonical fronto-central theta-beta ratio and age. Finally, we explore the unique associations between canonical fronto-central theta and canonical fronto-central beta with age, with and without controlling for fronto-central individual alpha peak frequencies and the fronto-central aperiodic exponent. We report false discovery rate (FDR) corrected $p$ values for pairwise and partial correlations, because FDR corrections have been shown to have increased power over other correction methods, particularly in cases with many comparisons and when the number of non-null hypotheses increase (Benjamini & Hochberg, 1995). Given the expectation that EEG metrics would be significantly intercorrelated, we opted for FDR to preserve as much statistical power as was feasible while controlling for false discoveries.

### 3.1 Pairwise Pearson’s correlation analyses

Our first confirmatory hypothesis was initially tested using Pearson’s correlations between log-normalized theta-beta ratios and age. As shown in Table 3 and in Figure 1, resting canonical fronto-central theta-beta ratios were negatively correlated with age ($r = -0.24$, 95% CI $[-0.35, -0.12]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$), such that the ratio of slow-wave to fast-wave activity was lower for older participants. Consistent with our second confirmatory hypothesis, as well as consistent with a prior unpublished analysis of this data set and previous independent studies in adults (Clark et al., 2004; Klimesch, 1999), we also observed a significant negative correlation between individual alpha peak frequency and age ($r = -0.17$, 95% CI $[-0.28, -0.05]$, $p_{uncorrected} = 0.006$, $p_{fdr} = 0.008$), such that fronto-central peak alpha frequencies were lower in older participants (Table 2, Figure 1). Additionally, consistent with prior work (Voytek et al., 2015), the fronto-central aperiodic exponent was negatively correlated with age ($r = -0.24$, 95% CI $[-0.35, -0.13]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$), consistent with a “flattening” of the aperiodic component with age. Finally, we replicated prior work (Donoghue, Dominguez, & Voytek, 2020) by finding that canonical fronto-central theta-beta ratio is more strongly correlated with the fronto-central aperiodic exponent ($r = 0.71$, 95% CI $[0.64, 0.76]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$) than with canonical fronto-central beta ($r = -0.28$, 95% CI $[-0.38, -0.16]$, $p_{uncorrected} < 0.001$, $p_{fdr} < .001$), Fisher’s z $= 6.90$, $p < .001$, or canonical fronto-central theta ($r = 0.50$, 95% CI $[0.41, 0.59]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$), Fisher’s z $= 3.89$, $p < 0.001$, suggesting that canonical theta-beta ratios are highly confounded with the aperiodic exponent.

Generalized Estimating Equations (GEE) confirmed these relationships held when adjusting for genetic dependencies between twin and sibling participants in the sample ($n_{twin/sibling} = 71$; Supplemental Table S6 for details of the GEE analyses). We also examined the partial correlations controlling for gender and race, and still observed a significant negative correlation between age and canonical fronto-central theta-beta ratio ($r = -0.23$, 95% CI $[-0.34, -0.11]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$), a significant negative correlation between age and fronto-central individual alpha peak frequency ($r = -0.18$, 95% CI $[-0.30, -0.06]$, $p_{uncorrected} = 0.003$, $p_{fdr} = 0.003$), and a significant negative correlation between age and fronto-central aperiodic exponent ($r = -0.23$, 95% CI $[-0.34, -0.11]$, $p_{uncorrected} < 0.001$, $p_{fdr} < 0.001$), while the relationship between age and fronto-central aperiodic exponent remained nonsignificant ($r = -0.09$, 95% CI $[-0.21, 0.03]$, $p_{uncorrected} = 0.155$, $p_{fdr} < 0.181$).

### 3.2 Partial correlation analyses

Next, we examined the partial correlations between age and canonical fronto-central theta-beta ratio, controlling separately for fronto-central individual alpha peak, fronto-central aperiodic offset, and fronto-central aperiodic exponent. As shown in Table 4 and Figure 2, the partial correlation between canonical fronto-central theta-beta ratio and age becomes nonsignificant only when controlling for the fronto-central aperiodic exponent, $r_{partial} = -0.10$, 95% CI $[-0.21, 0.02]$, $p_{uncorrected} = .110$, $p_{fdr} = 0.110$. This suggests that in adults, the flattening of the aperiodic curve with age, as denoted by the aperiodic exponent, may be largely driving the relationship between canonical theta-beta ratio and age.

### 3.3 Mediation analyses

To further understand the relationship between age and canonical fronto-central theta-beta ratio, we conducted a series of exploratory mediational analyses to see if fronto-central individual alpha peak frequency, fronto-central aperiodic offset, or fronto-central aperiodic exponent would fully mediate the relationship between age
| TABLE 3 Correlations and descriptive statistics between age and EEG metrics, collapsed across eyes open and eyes closed (n = 268) |
| --- |
|   | Mean (SD) | 1. Age | 2. Canonical Fronto-Central Theta | 3. Canonical Fronto-Central Beta | 4. Canonical Fronto-Central Theta-Beta Ratio | 5. Fronto-Central Individual Alpha Peak Frequency | 6. Fronto-Central Aperiodic Exponent |
| 1. Age | 55.8 (11.0) | — | — | — | — | — | — |
| 2. Canonical Fronto-Central Theta | 0.77 (1.17) | 0.01 [−0.11, 0.13] | — | — | — | — | — |
| 3. Canonical Fronto-Central Beta | 0.22 (0.17) | 0.42 [0.32, 0.52] | — | — | — | — | — |
| 4. Canonical Fronto-Central Theta-Beta Ratio | 1.09 (0.68) | 0.50 [0.41, 0.59] | — | — | — | — | — |
| 5. Fronto-Central Individual Alpha Peak Frequency | 9.31 (0.98) | −0.57 [−0.80, −0.32] | — | — | — | — | — |
| 6. Fronto-Central Aperiodic Exponent | 1.23 (0.26) | 0.44 [0.34, 0.53] | — | — | — | — | — |
| 7. Fronto-Central Aperiodic Offset | 0.43 (0.44) | 0.65 [0.58, 0.72] | — | — | — | — | — |

Note: 95% confidence intervals for pairwise correlations displayed in brackets followed by uncorrected and FDR corrected p values.
and canonical fronto-central theta-beta ratio. Mediation analyses were conducted using the processR package in R (Moon, 2021), with maximum likelihood estimation and 10,000 bootstrap estimates of standard error. See Figure 3.

As shown in Table 5, only the fronto-central aperiodic exponent fully mediated the relationship between age and canonical fronto-central theta-beta ratio, such that the direct effect ($c'$) between age and canonical fronto-central theta-beta ratio was nonsignificant ($c' = -0.004, 95\% CI [-0.010, 0.001], p = .121$). This suggests that the relationship between canonical fronto-central theta-beta ratio and age is driven primarily by the aperiodic exponent, not periodic activity in the canonical theta and beta bands or leakage of alpha into the canonical theta band with age. However, we wanted to determine the extent to which canonical theta and beta have unique associations with age, if any, apart from the theta-beta ratio. Therefore, we conducted a hierarchical regression, regressing age on canonical theta and beta, adding individual alpha peak frequency in the second block and adding the aperiodic exponent in the third block. The first two blocks of the analysis were preregistered as exploratory analysis E1 in the preregistration (https://osf.io/n57au), and the third block was added as an exploratory step to include the aperiodic component. Because controlling for aperiodic offset did not substantially change the relationship between canonical theta-beta ratio and age or mediate the relationship between age and canonical theta-beta ratio, and because the aperiodic exponent and offset are highly intercorrelated ($r = 0.75, 95\% CI [0.70, 0.80], p_{uncorr} < 0.001, p_{dfr} < 0.001$), we conducted the stepwise

3.4 | Hierarchical regression analyses

Taken together, the correlational and meditational analyses suggest that the relationship between canonical fronto-central theta-beta ratio and age in older adults is due to the underlying aperiodic exponent, not periodic activity in the canonical theta and beta bands or leakage of alpha into the canonical theta band with age. However, we wanted to determine the extent to which canonical theta and beta have unique associations with age, if any, apart from the theta-beta ratio. Therefore, we conducted a hierarchical regression, regressing age on canonical theta and beta, adding individual alpha peak frequency in the second block and adding the aperiodic exponent in the third block. The first two blocks of the analysis were preregistered as exploratory analysis E1 in the preregistration (https://osf.io/n57au), and the third block was added as an exploratory step to include the aperiodic component. Because controlling for aperiodic offset did not substantially change the relationship between canonical theta-beta ratio and age or mediate the relationship between age and canonical theta-beta ratio, and because the aperiodic exponent and offset are highly intercorrelated ($r = 0.75, 95\% CI [0.70, 0.80], p_{uncorr} < 0.001, p_{dfr} < 0.001$), we conducted the stepwise
analyses with only the aperiodic exponent to avoid issues of multicollinearity.

As shown in Table 6, in Block 2 canonical fronto-central theta was significantly associated with age when controlling for fronto-central individual alpha peak frequency, $b = -1.86, t(264) = 2.84, p = .005$. However, canonical fronto-central theta was nonsignificantly associated with age when controlling for the fronto-central aperiodic exponent in block 3, $b = -0.53, t(264) = 0.75, p = .456$. In Block 3, there were significant relationships between canonical fronto-central beta and age, $b = -2.71, t(264) = 3.96, p < .001$, and the fronto-central aperiodic exponent and age, $b = -1.18, t(264) = 4.32, p < .001$, suggesting that there is a significant increase in periodic activity in the canonical beta band with age, as well as the age-related flattening of the aperiodic component and “slowing” of the individual alpha peak frequency. The lack of unique variance associated with canonical theta power over and above the aperiodic component is consistent with the lack of definable peaks (with the FOOOF package) within the canonical theta band, as described in the supplemental materials.

4 | DISCUSSION

In the preregistered portion of the current study, we aimed to replicate and extend previous observations that canonical fronto-central theta-beta ratios and fronto-central individual alpha peak frequency are associated with age in a large sample of 268 adults featuring a wide age range (36–84 years). Consistent with preregistered predictions and previous studies, we found that both canonical fronto-central theta-beta ratios and fronto-central individual alpha peak frequencies were negatively correlated with age. Exploratory analyses indicated that the association between canonical fronto-central theta-beta ratios and age remained when controlling for fronto-central individual alpha peak frequencies, demonstrating that age-related decreases in canonical fronto-central theta-beta ratios are not due to age-related decreases in fronto-central individual alpha peak frequencies. Instead, the relationship between canonical fronto-central theta-beta ratios and age were reduced when controlling for the fronto-central aperiodic exponent. Additionally, mediation analyses found that only the fronto-central aperiodic exponent fully mediated the relationship between age and canonical fronto-central theta-beta ratios. Furthermore, this effect appears to be robust against multiple ways of defining theta-beta ratios and individual alpha peaks, and consistent across eyes closed only recordings, as described in the supplemental materials.
4.1 | Understanding how aperiodic components, canonical theta-beta ratios and individual alpha peak frequencies change over the lifespan

Our results also reveal a complex pattern of associations between canonical fronto-central theta-beta ratios, fronto-central individual alpha peak frequency, fronto-central aperiodic activity, and age. Consistent with previous studies (Voytek et al., 2015), we observed that the aperiodic exponent was negatively associated with age, suggesting relatively synchronized aperiodic firing in younger versus older adults. However, the age-related differences in aperiodic offset reported by prior research (Voytek et al., 2015) were not significant in our sample ($p_{uncorr} = 0.071$, $p_{FDR} = 0.083$). We observed that the age-related differences in the aperiodic exponent are preserved into older adulthood, and are not limited only to the younger (e.g., <44 years of age) populations reported on in previous studies (Donoghue, Dominguez, & Voytek, 2020), or the relatively small samples used in others (Voytek et al., 2015). We also observed that the association between age and canonical fronto-central theta-beta ratios is reduced when statistically adjusting for the fronto-central aperiodic exponent, consistent with the observation that individual differences in ratio metrics are likely confounded with individual differences in aperiodic activity, especially when there is no clear peak within the particular power band. Critically, the association between age and canonical fronto-central theta-beta ratio is fully mediated by the fronto-central aperiodic exponent.

4.2 | Limitations of the current study

The current study has some methodological limitations, particularly regarding the preregistered decision
to examine the data combined across eyes open and eyes closed periods and calculate individual alpha peak frequency from fronto-central ROI. Combining eyes open and eyes closed data results in unequal number of epochs between the two conditions. Additionally, alpha power is known to be strongest during eyes closed recordings from posterior sites, which may have impeded our ability to detect individual alpha peak frequency. However, additional analyses reported in the supplemental materials on only the eyes closed data, as well as from individual alpha peak frequency calculated from across all sensors, neither substantially increased the number of RestingIAF package definable peaks, nor changed the interpretation of the analyses. Using the FOOOF package to define individual alpha peak frequency did increase the number of definable individual alpha peaks to \( n = 302 \), but the results do not change with this alternative method of defining individual alpha peaks (see Supplemental Materials for full details). Additionally, the decision to use visual artifactual screening makes the preprocessing stream non-reproducible without getting a list of artifactual components. However, we decided to keep the original data preprocessing pipeline from the initial MIDUS 2 EEG data release the same to increase consistency with the publicly available MIDUS 2 alpha asymmetry metrics (http://midus.colectica.org/; https://www.icpsr.umich.edu/web/ICPSR/series/203). The current study is also limited by examining these relationships cross-sectionally across age. Additional longitudinal work is needed to tease apart the unique developmental trajectories of theta-beta ratio and individual peak alpha frequency.

### 4.3 | Implications for fronto-central aperiodic activity, canonical fronto-central theta-beta ratio, and fronto-central individual alpha peak frequency as markers of executive function and healthy aging

Taken together, our findings complicate the interpretation of fronto-central theta-beta ratio as a marker of executive function. In adolescents and young adults, higher theta-beta ratios are associated with more executive dysfunction and related to ADHD (Arns et al., 2013), and lower theta-beta ratios are associated with better attentional control (Perone et al., 2018). Considering older age-related decline in executive function (Buckner, 2004; Lustig & Jantz, 2015), fronto-central theta-beta ratios may exhibit a curvilinear relationship with executive functioning, such that better executive functioning is related to a moderate level of fronto-central theta-beta ratio. Additionally, it may be that adolescence and younger adults are more
prone to disruptions related to elevated theta-beta ratios and older adults are more prone to reductions in theta-beta ratios potentially driven by normative aging processes. The moderate level of theta-beta ratio may reflect an optimal balance in the bidirectional regulation of bottom-up subcortical processes by top-down cortical processes that theta-beta ratio is putatively suggested to index (Knyazev, 2007; Schutter & Knyazev, 2012).

However, considering recent data regarding the physiological mechanisms and functions of neural noise, the theta-beta ratio model advanced in previous studies is increasingly difficult to support. As Donoghue, Dominguez, and Voytek (2020) observed and we have replicated, the association between theta-beta ratios and age is confounded by age-related differences in aperiodic activity. Inter- and intraindividual differences in aperiodic activity have also been strongly and consistently associated with variation in cognitive function (Tran et al., 2020; Voytek et al., 2015), and provide a parsimonious and physiologically plausible mechanism for variation in cognitive function across the lifespan relating to the ratio of excitatory to inhibitory activity (Donoghue, Haller, et al., 2020; Gao et al., 2017; Waschke et al., 2021).

Given the relationships between aperiodic activity, individual alpha peak frequency, and theta-beta ratio with age, as well as existing research linking aperiodic activity to cognitive function (Tran et al., 2020; Voytek et al., 2015), individual alpha peak frequency with memory-related aspects of executive functioning (i.e., Clark et al., 2004) and theta-beta ratio with attention-related aspects of executive functioning (Angelidis et al., 2016), these markers appear to be promising, but potentially overlapping and redundant measures of healthy aging. Further research is needed to confirm the unique associations of aperiodic activity, individual alpha peak frequency and theta-beta ratio with memory, executive functioning, and measures of healthy and pathological aging.

## 5 Conclusion

Overall, we found that both fronto-central theta-beta ratios and fronto-central individual alpha peak frequencies were cross-sectionally negatively associated with age, and that age-related decreases in fronto-central theta-beta ratios are not due to age-related decreases in fronto-central individual alpha peak frequencies. This suggests that changes in both theta-beta ratios and individual alpha peak frequencies may index differential components of healthy aging. Critically, our findings highlight confounds between theta-beta ratio and the aperiodic exponent, suggesting that both metrics should be considered in understanding power-based EEG metrics and aging. Future research should explicitly examine multiple facets of executive function (including working memory, attention control, and response inhibition) to determine how theta-beta ratios, aperiodic components, and individual alpha peak frequencies at rest relate to cognitive functioning in older adulthood, and if these measures are suitable as biomarkers for healthy and pathological aging. Additionally, we are limited by the cross-sectional nature of the study from determining if these cross-sectional relationships between age and resting EEG metrics reflect an underlying developmental trajectory in aging. Future longitudinal research is needed to trace the developmental trajectory of theta-beta ratios, aperiodic components, and individual alpha peak across the lifespan.
AUTHOR CONTRIBUTIONS
Anna J. Finley: Conceptualization; data curation; formal analysis; investigation; supervision; visualization; writing – original draft; writing – review & editing.
Douglas J. Angus: Conceptualization; formal analysis; writing – original draft; writing – review and editing.
Carien Van Reekum: Data curation; formal analysis; investigation; methodology; writing – review and editing.
Richard J. Davidson: Funding acquisition; methodology; project administration; resources; writing – review and editing.
Stacey M. Schaefer: Data curation; investigation; methodology; project administration; resources; supervision; writing – review and editing.

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ORCID
Anna J. Finley © https://orcid.org/0000-0002-0881-9147
Douglas J. Angus © https://orcid.org/0000-0001-9722-2475

REFERENCES
Angelakis, E., Lubar, J. F., Stathopoulos, S., & Kounios, J. (2004). Peak alpha frequency: An electroencephalographic measure of cognitive preparedness. *Clinical Neurophysiology*, 115(4), 887–897. https://doi.org/10.1016/j.clinph.2003.11.034
Angelidis, A., Hagenaars, M., van Son, D., van der Does, W., & Putman, P. (2018). Do not look away! Spontaneous frontal EEG theta/beta ratio as a marker for cognitive control over attention to mild and high threat. *Biological Psychology*, 135, 8–17. https://doi.org/10.1016/j.biopsycho.2018.03.002
Angelidis, A., van der Does, W., Schakel, L., & Putman, P. (2016). Frontal EEG theta/beta ratio as an electrophysiological marker for attention control and its test-retest reliability. *Biological Psychology*, 121, 49–52. https://doi.org/10.1016/j.biopsycho.2016.09.008
Anticevic, A., Repovs, G., Shulman, G. L., & Barch, D. M. (2010). When less is more: TFP and default network deactivation during encoding predicts working memory performance. *NeuroImage*, 49(3), 2638–2648. https://doi.org/10.1016/j.neuroimage.2009.11.008
Arns, M., Connors, C. K., & Kraemer, H. C. (2013). A decade of EEG theta/beta ratio in ADHD: A meta-analysis. *Journal of Attention Disorders*, 17(5), 374–383. https://doi.org/10.1177/1087054712460087
Babiloni, C., Barry, R. J., Başar, E., Blinowska, K. J., Cichocki, A., Drinkenburg, W. H. I. M., Klimesch, W., Knight, R. T., Lopes da Silva, F., Nunez, P., Oostenveld, R., Jeong, J., Pascual-Marqui, R., Valdes-Sosa, P., & Hallett, M. (2020). International federation of clinical neurophysiology (IFCN)–EEG research working group: Recommendations on frequency and topographic analysis of resting state EEG rhythms. Part 1: Applications in clinical research studies. *Clinical Neurophysiology*, 131(1), 285–307. https://doi.org/10.1016/j.clinph.2019.06.234
Babiloni, C., Binetti, G., Cassarino, A., Forno, G. D., Percio, C. D., Ferrari, F., Ferri, R., Frisoni, G., Galderisi, S., Hirata, K., Lanuzza, B., Miniusi, C., Mucci, A., Nobili, F., Rodriguez, G., Romani, G. L., & Rossini, P. M. (2006). Sources of cortical rhythms in adults during physiological aging: A multicentric EEG study. *Human Brain Mapping*, 27(2), 162–172. https://doi.org/10.1002/hbm.20175
Benjamini, Y., & Hochberg, Y. (1995). Controlling the false discovery rate: A practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society: Series B (Methodological)*, 57(1), 289–300. https://doi.org/10.1111/1467-9868.00203
Buckner, R. L. (2004). Memory and executive function in aging and AD: Multiple factors that cause decline and reserve factors that compensate. *Neuron*, 44(1), 195–208. https://doi.org/10.1016/j.neuron.2004.09.006
Clark, C. R., Veltmeyer, M. D., Hamilton, R. J., Simms, E., Paul, R., Hermens, D., & Gordon, E. (2004). Spontaneous alpha peak frequency predicts working memory performance across the age span. *International Journal of Psychophysiology*, 53(1), 1–9. https://doi.org/10.1016/j.ijpsycho.2003.12.011
Clarke, A. R., Barry, R. J., McCarthy, R., & Selikowitz, M. (2001). Age and sex effects in the EEG: Differences in two subtypes of attention-deficit/hyperactivity disorder. *Clinical Neurophysiology*, 112(5), 815–826. https://doi.org/10.1016/S1388-2457(01)00487-4
Corcoran, A. W., Alday, P. M., Schlesewsky, M., & Bornkessel-Schlesewsky, I. (2018). Toward a reliable, automated method of individual alpha frequency (IAF) quantification. *Psychophysiology*, 55(7), e13064. https://doi.org/10.1111/psyp.13064
Daselaar, S. M., Prince, S. E., & Cabeza, R. (2004). When less means more: Deactivations during encoding that predict subsequent memory. *NeuroImage*, 23(3), 921–927. https://doi.org/10.1016/j.neuroimage.2004.07.031
Davidson, R. J. (1984). Hemispheric asymmetry and emotion. In Approaches to Emotion (pp. 39–57). Psychology Press.
Donoghue, T., Dominguez, J., & Voytek, B. (2020). Electrophysiological frequency band ratio measures conflate periodic and aperiodic neural activity. *Neuron*, 27(3), 921–927. https://doi.org/10.1016/j.neuron.2019.06.028
Donoghue, T., Haller, M., Peterson, E. J., Varma, P., Sebastian, P., Gao, R., Noto, T., Lara, A. H., Wallis, J. D., Knight, R. T., Shetystuk, A., & Voytek, B. (2020). Parameterizing neural power spectra into periodic and aperiodic components. *Nature Neuroscience*, 23(12), 1655–1665. https://doi.org/10.1038/s41593-020-00744-x
Electrical Geodesics, Inc. (2007, January 31). Electrical Geodesics, Incorporated.
SUPPORTING INFORMATION
Additional supporting information may be found in the online version of the article at the publisher’s website.

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