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Probabilistic, entropy-maximizing control of large-scale neural synchronization

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Abstract

Oscillatory neural activity is dynamically controlled to coordinate perceptual, attentional and cognitive processes. On the macroscopic scale, this control is reflected in the U-shaped deviations of EEG spectral-power dynamics from stochastic dynamics, characterized by disproportionately elevated occurrences of the lowest and highest ranges of power. To understand the mechanisms that generate these low- and high-power states, we fit a simple mathematical model of synchronization of oscillatory activity to human EEG data. The results consistently indicated that the majority (~95%) of synchronization dynamics is controlled by slowly adjusting the probability of synchronization while maintaining maximum entropy within the timescale of a few seconds. This strategy appears to be universal as the results generalized across oscillation frequencies, EEG current sources, and participants (N = 52) whether they rested with their eyes closed, rested with their eyes open in a darkened room, or viewed a silent nature video. Given that precisely coordinated behavior requires tightly controlled oscillatory dynamics, the current results suggest that the large-scale spatial synchronization of oscillatory activity is controlled by the relatively slow, entropy-maximizing adjustments of synchronization probability (demonstrated here) in combination with temporally precise phase adjustments (e.g., phase resetting generated by sensorimotor interactions). Interestingly, we observed a modest but consistent spatial pattern of deviations from the maximum-entropy rule, potentially suggesting that the mid-central-posterior region serves as an "entropy dump" to facilitate the temporally precise control of spectral-power dynamics in the surrounding regions.

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Introduction

A great deal of evidence suggests that the coordination of oscillatory activity contributes to controlling neural communications that are necessary for effective operations of perception, attention, memory, and cognition (e.g., Fries, 2005; Palva et al., 2005; Palva & Palva, 2007; 2012; Busch & VanRullen, 2010; Engel & Fries, 2010; Duzel et al., 2010; Mathewson et al., 2011; 2010; Voytek et al., 2010; Donner & Siegel, 2011; Arnal & Giraud, 2012; Hipp et al., 2012; Klimesch, 2012; Spaak et al., 2012; Bonnefond & Jensen, 2015; Michalareas et al., 2016; Müller et al., 2016). While fine-tuned coordination likely involves controlling the phases of oscillatory activity across frequency bands, the impact of oscillatory activity also depends on the size of synchronized neural population. For large-scale neural activity detected by scalp-recorded electroencephalography (EEG), the spectral power obtained at an EEG current source reflects the size of synchronously oscillating population within its spatial resolution. EEG spectral power fluctuates at each current source reflecting the dynamic changes in large-scale synchronization of oscillatory activity. The goal of the current study was to elucidate the mechanisms that control these large-scale synchronization dynamics.

One way to investigate dynamic control is to compare spectral-power dynamics between EEG and their phase-scrambled controls. Phase scrambling randomizes cross-frequency phase relations, thus destroying temporal structures that depend on cross-frequency phase alignment, rendering spectral-power dynamics stochastic (memory free) while preserving time-averaged power spectra. Because stochastic dynamics reflect a Poisson process (see below), phase-scrambled spectral-power dynamics are characterized by exponential power distributions. Real EEG spectral-power dynamics deviate from exponential profiles in a characteristic U-shaped manner with disproportionately elevated occurrences of the lowest and highest ranges of power (see below). This indicates that real spectral-power dynamics exhibit intermittent bursts of extensive oscillatory synchronization separated by periods of sparse synchronization (compared with stochastic dynamics). How are these periods of extensive and sparse synchronization generated?

On the one hand, the brain neural network may actively boost large-scale synchronization or inhibit it in precise temporal coordination with behavioral demands. On the other hand, the network may indirectly influence large-scale synchronization by increasing or decreasing the probability of synchronization on a relatively slow timescale while generally maintaining maximum entropy for energy efficiency. As described in the results section, these possibilities can be evaluated in a relatively simple manner.

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Methods

1. Participants

Fifty-two Northwestern University students (35 women, 1 non-binary; ages 18 to 29 years, M = 20.75, SD = 2.52) gave informed consent to participate for monetary compensation (\$10/hr). All were right-handed, had normal hearing and normal or corrected-to-normal vision, and had no history of concussion. They were tested individually in a dimly lit or dark room. The study protocol was approved by the Northwestern University Institutional Review Board. Participants p1-p7 and p12-p28 (N = 24) participated in the rest-with-the-eyes-closed condition where their EEG was recorded for ~5 min while they rested with their eyes closed and freely engaged in spontaneous thoughts. This condition was always run first for those who also participated in the nature-video condition. Participants p8-p28 (N = 21) also participated in the nature-video condition where their EEG was recorded for ~5 min while they viewed a silent nature video. To evaluate the test-retest reliability, the nature-video condition was run twice (20-30 min apart), labeled as earlier viewing and later viewing in the analyses. A generic nature video was presented on a 13-inch, 2017 MacBook Pro, 2880(H)-by-1800(V)-pixel-resolution screen with normal brightness and contrast settings, placed 100 cm away from participants, subtending $16^{\circ}(H)$ -by- $10^{\circ}(V)$ of visual angle. Participants p29-p52 (N = 24) participated in the replication of the rest-with-the-eyes-closed condition and the rest-with-the-eyes-open-in-dark condition which was the same as the former except that the room was darkened and participants kept their eyes open while blinking naturally.

2. EEG recording and pre-processing

While participants rested with their eyes closed, rested with their eyes open in dark, or viewed a silent nature video for approximately 5 min, EEG was recorded from 60 scalp electrodes (although we used a 64-electrode montage, we excluded signals from noise-prone electrodes, *Fpz*, *Iz*, *T*9, and *T10*, from analyses) at a sampling rate of 512 Hz using a BioSemi ActiveTwo system (see www.biosemi.com for details). Electrooculographic (EOG) activity was monitored using four face electrodes, one placed lateral to each eye and one placed beneath each eye. Two additional electrodes were placed on the left and right mastoid area. The EEG data were preprocessed using EEGLAB and ERPLAB toolboxes for MATLAB (Delorme & Makeig, 2004; Lopez-Calderon & Luck, 2014). The data were re-referenced offline to the average of the two mastoid electrodes, bandpass-filtered at 0.01 Hz-80 Hz, and notch-filtered at

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60 Hz (to remove power-line noise that affected the EEG signals from some participants). For the EEG signals recorded while participants rested with the eyes open in dark or while they viewed a silent nature video, an Independent Component Analysis (ICA) was conducted using EEGLABs' *runica* function (Makeig et al., 1996; 2000) to remove components reflecting blinks. To reduce effects of volume conduction (to within adjacent sites; e.g., Cohen, 2014), to virtually eliminate the effects of reference electrode choices, as well as to facilitate data-driven determinations of EEG current sources, we applied the surface-Laplacian transform to all EEG data (Hjorth, 1980; Kayser and Tenke, 2006; Tenke and Kayser, 2012), using the Perrin and colleagues' method (e.g., Perrin et al., 1987; Perrin et al., 1989a; 1989b) with a typical set of parameter values (Cohen, 2014). We refer to the surface-Laplacian transformed EEG signals that represent the current sources under the 60 scalp sites simply as EEG signals.

3. EEG analysis

3.1. EEG temporal derivative. An example 1 sec EEG waveform at a central site FCz from one participant (at rest with the eyes closed) is shown in Figure 1A (black curve). The mean spectral-amplitude profile of the full length (~5 min) version of the same data, with the fast Fourier transform (FFT) computed on each consecutive 5 sec waveform and then averaged, is shown in Figure 1B (black curve; the shaded area represents ±1 standard error of the mean). The general linear decrease in the spectral amplitude for higher frequencies with a slope of approximately 1 (in log-log scale) reflects the 1/f background profile largely explained by the neuronal Ornstein-Uhlenbeck process that exhibits a random-walk type behavior (e.g., Koch, 1999; Mazzoni et al., 2008; see He, 2014 for a review of the various factors that contribute to the $1/f^{\beta}$ spectral background; see Gao et al., 2017 for contributions of the excitatory and inhibitory dynamics). The spectral "bumps" seen around 10 Hz, 20 Hz, and 30 Hz indicate the characteristic bands of oscillation frequencies that the neural population reflected at this site for this person may utilize for communication and/or information processing. Taking the temporal derivative of EEG ($\frac{\Delta EEG}{\Delta t}$, where Δt is the temporal resolution, i.e., 1/512 sec) (see the black curve in Figure 1C) largely removes the 1/f background (due to trigonometric properties) to highlight the oscillatory activity (see the black curve in Figure 1D). While Figure 1D shows an example at one site from one participant, we confirmed that taking the temporal derivative generally flattened the background spectral-amplitude profiles across sites and participants. Thus, to highlight the dynamics of oscillatory activity (over and above the general 1/f spectral background) with a simple continuous mathematical operation that macroscopically estimates

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the underlying electrical currents (without having to estimate $1/f^{\beta}$ in sliding time windows to discount their influences), we used the EEG temporal derivative, which we call EEGd.

3.2. Computing spectral power as a function of time. The spectral-amplitude profiles shown in Figure 1B and 1D are time-averaged (standard fast Fourier transforms). To investigate how spectral power (amplitude squared) fluctuated, we used a Morlet wavelet-convolution method (e.g., Cohen, 2014) to extract spectral amplitudes as a function of time. Each EEGd waveform was decomposed into a time series of spectral power using Morlet wavelets with twenty center frequencies f_c 's between 6 Hz and 50 Hz, encompassing the θ , α , β , and γ bands. The f_c 's were logarithmically spaced as neural temporal-frequency tunings tend to be approximately logarithmically scaled (e.g., Hess & Snowden, 1992; Lui et al., 2007). The accompanying *n* factors (roughly the number of cycles per wavelet, $n = 2\pi f \cdot SD$, where *SD* is the wavelet standard deviation) were also logarithmically spaced between 4.4 and 14.5, yielding the temporal resolutions ranging from *SD* = 117 ms (at 6 Hz) to *SD* = 46 ms (at 50 Hz) and spectral resolutions ranging from *FWHM* (full width at half maximum) = 3.2 Hz (at 3 Hz) to *FWHM* = 8.2 Hz (at 50 Hz). These values strike a good balance for the temporal/spectral-resolution trade-off, and are typically used in the literature (e.g., Cohen, 2014).

3.3. Generating phase-scrambled controls. We generated phase-scrambled control data whose spectral power fluctuated stochastically (i.e., unpredictably in a memory free manner) while maintaining the time-averaged spectral-amplitude profiles of the real EEG data. While phase-scrambling can be performed using several different methods, we chose discrete cosine transform, DCT (e.g., Kiya et al., 2010). In short, we transformed each 5 min EEG waveform with type-2 DCT, randomly shuffled the signs of the coefficients, and then inverse-transformed it with type-3 DCT (the "inverse DCT"), which yielded a phase-scrambled version. DCT phase-scrambling is similar to DFT (discrete Fourier transform) phase-scrambling except that it is less susceptible to edge effects. We verified that DCT phase-scrambling yielded a desired outcome, generating waveforms whose spectral-power fluctuations conformed to exponential distributions (see below) indicative of a Poisson point process (a stochastic process), with virtually no distortions to the time-averaged spectral-amplitude profiles of EEG or EEGd (e.g., the blue curves overlap the black curves in Figure 1B and 1D).

3.4. Computing entropy per interval, d. We computed entropy for non-overlapping intervals of duration d (sec). For each interval, we divided spectral power values into N_{bins} bins using the Freedman-Diaconis method (Freedman & Diaconis, 1981),

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$$N_{bins} = ceil\left(\frac{max}{2 \cdot iqr \cdot [d \cdot 512]^{-1/3}}\right),$$

where *max* is the maximum spectral-power value corresponding to the highest bin, *iqr* is the interquartile range, *d*•512 is the number of spectral-power values available within each *d* (sec) interval sampled at 512 Hz, and *ceil* takes the nearest larger integer. The value of *max* was chosen such that the highest bin reached the 99.9th percentile or higher spectral-power value for each frequency and behavioral condition (because spectral-power values varied primarily as a function of frequency and condition). The *iqr* values were computed per frequency per condition, averaged across frequencies, then averaged within the same behavioral condition, yielding three values, one for the rest-with-the-eyes-closed condition (averaged across the original and replication condition), one for the rest-with-the-eyes-open-in-dark conditions). Thus, *N_{bins}* was optimized for the condition-specific *iqr* and the number of data points within the *d* (sec) interval while the same range [0, *Max*] was used in all cases. Using these spectral-power bins, we generated the probability distribution of spectral power values for each *d* (sec) interval (per frequency per site per participant per condition), and computed the corresponding entropy as,

$$S = \sum_{i=1}^{i=N_{bins}} p_i \cdot ln(p_i),$$

where S is entropy, p_i is the proportion of spectral-power values within the *i*th bin, and N_{bins} is the number of bins.

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Fig 1. The use of the temporal derivative of EEG (EEGd) and DCT-phase-scrambled controls for investigating the dynamic control of synchronization of oscillatory activity. A. An example of 1 sec EEG waveform (black) and its DCT-phase-scrambled control (blue) at *FCz* from one participant. **B.** The mean spectral-amplitude profiles of the full length (~5 min) versions of the same EEG data (black) and its DCT-phasescrambled control (blue), with the fast Fourier transform (FFT) computed on each consecutive 5 sec waveform and then averaged, plotted in a log-log format. **C.** The temporal derivatives, which we call EEGd, of the example EEG waveform (black) and its DCT-phase-scrambled control (blue) shown in A. **D.** The mean spectralamplitude profiles of the full length (~5 min) versions of the same EEGd data (black) and its DCT-phasescrambled control (blue), with the fast Fourier transform (FFT) computed on each consecutive 5 sec waveform and then averaged, plotted in a spectralamplitude profiles of the full length (~5 min) versions of the same EEGd data (black) and its DCT-phasescrambled control (blue), with the fast Fourier transform (FFT) computed on each consecutive 5 sec waveform and then averaged, plotted in a semi-log format. For B and D, the shaded areas represent ±1 standard error of the mean based on the FFTs computed on multiple 5 sec waveforms. The units are arbitrary (a.u.).

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Results

We started with a simple stochastic model of neural synchronization. As most neural connections are short-range (e.g., Buzsaki, 2006), we postulated that, at each moment, synchronization would sequentially spread from neural-unit to neural-unit with the probability p_{term} that the rapid spreading would terminate at any given unit. It is reasonable to assume that the number of neural units is large and p_{term} is small. Thus, the probability that the size of synchronized neural population, *N* units, is larger than *n* units at a given timepoint, is provided by the Poisson equation,

$$P(N > n) = \frac{(p_{term} \cdot n)^0}{0!} e^{-p_{term} \cdot n} = e^{-p_{term} \cdot n} - Eq. 1.$$

Then, the probability density function f(n) for the occurrence of a synchronized population of size *n* can be obtained by solving,

$$\int_{n}^{\infty} f(n) dn = e^{-p_{term} \cdot n} - Eq. 2$$

yielding,

$$f(n) = p_{term} e^{-p_{term} \cdot n} - Eq. 3.$$

The exponential form of f(n) indicates that the model yields maximum entropy for fluctuations in n for a given temporal average $\langle n \rangle$. Thus, our model (*Eq.* 1) describes a simple macroscopic mechanism that generates synchronization dynamics that maximize entropy for a given temporal average (*Eq.* 3). We note that any reasonable model that leads to an exponential probability density function for n with its parameters related to the probability of synchronization would be just as appropriate for our discussion.

It is reasonable to assume that EEG spectral power at a given site is proportional to the size of the synchronously oscillating neural population *n* within the accessible current sources. Then, *Eq.* 3 predicts an exponential distribution for the fluctuations of spectral power for phase-scrambled EEG (which are rendered stochastic). Our data confirmed this prediction (the thinner horizontal lines in Figure 2). Our goal was to elucidate the mechanisms that make the real EEG spectral-power dynamics deviate from stochastic (exponential) dynamics in the characteristic U-shaped manner (the thicker curves in Figure 2). To this end, we considered the relationship between average spectral power and entropy.

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Fig 2. Probability distributions of spectral power as residual deviations (in ratio) from exponential fits to phase-scrambled controls. The distributions are shown for representative frequency bands, θ (6.0 Hz), α (10.5 Hz), β (14.6 Hz and 20.6 Hz) and γ (31.9 Hz, 40.0 Hz, and 50.1 Hz), color-coded from cooler to warmer. The five panels show the residual distributions for the five conditions: 5-min rest with the eyes closed (Rest EC), its replication (Rest EC rep), 5-min rest with the eyes open in dark (Rest EO DK), and the earlier and later 5-min viewing of a silent nature video (Nature video). Timepoint-by-timepoint spectral-power values obtained in each condition were normalized to the median power per frequency per site per participant, then averaged across sites (*x*-axis). Note that all distributions for the phase-scrambled controls tightly conform to the exponential form (the thinner horizontal lines at *y* = 1), whereas the distributions for the real EEG data (the thicker curves) deviate from the exponential form in a characteristic U-shaped manner with elevated occurrences of the lowest and highest ranges of power. The shaded areas represent ±1 standard error of the mean with participants as the random effect.

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Stochastic dynamics such as the spectral-power dynamics of the phase-scrambled controls (the thinner horizontal lines in Figure 2) are well fit by Eq. 3 with a constant p_{term} . Nevertheless, the effective value of p_{term} within an interval of duration d, which we call $p_{term.d}$, stochastically fluctuates with the variance given by,

$$Var(p_{term.d}) \propto p_{term} \cdot (1 - p_{term})/d - Eq. 4.$$

Note that this is analogous to the familiar coin-tossing example. While the probability of getting heads is stationary with $p_{heads} = \frac{1}{2}$ for each (fair) coin toss, the effective probability of getting heads, that is, the actual proportion of heads obtained for a given set of *N* trials $p_{heads.N}$ is variable, with its set-to-set variance given by, $Var(p_{heads.N}) = p_{heads} \cdot (1 - p_{heads})/N$.

For stochastic dynamics of synchronization, Eq. 3 holds within an interval of any duration (given that it includes sufficient data points to reliability evaluate the probability distribution of *n*), so that the average size of a synchronized population <n> and entropy *S* within any interval of duration *d* are given by,

$$\langle n \rangle = \int_0^\infty f(n) \cdot n \, dn = \frac{1}{p_{term.d}} - Eq. 5, \text{ and}$$
$$S = -\int_0^\infty f(n) \cdot \ln\{f(n)\} \, dn = 1 - \ln(p_{term.d}) - Eq. 6,$$

where f(n) is given by Eq. 3 with $p_{term.d}$ substituted for p_{term} . Note that even if $p_{term.d}$ varied beyond the level of stochastic fluctuations (Eq. 4), Eqs. 5 and 6 would still hold if $p_{term.d}$ remained effectively constant on the timescale of d.

Taking the natural log of Eq. 5, we get, $ln(\langle n \rangle) = -ln(p_{term.d})$. Substituting this into Eq. 6 yields a linear relationship between entropy, *S*, and the log average size of the synchronized population, $ln(\langle n \rangle)$,

 $S = ln(\langle n \rangle) + 1 - Eq. 7.$

As we assume that the observed spectral power *SP* at each site is proportional to the size of the synchronized neural population *n*, we have

$$SP = k \cdot n - Eq. 8,$$

where k is the constant of proportionality. Taking the temporal average yields,

$$\langle SP \rangle = k \cdot \langle n \rangle - Eq. 9.$$

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Taking the natural log of Eq. 9, $ln(\langle SP \rangle) = ln(\langle n \rangle) + ln(k)$, and solving for $ln(\langle n \rangle)$, we get,

$$ln(\langle n \rangle) = ln(\langle SP \rangle) - ln(k) - Eq. 10.$$

Substituting Eq. 10 into Eq. 7 yields,

$$S = ln(\langle SP \rangle) + 1 - ln(k) - Eq. 11.$$

Note that any attenuation of *SP* due to the use of scalp-recorded EEG to compute spectral power is absorbed in k (*Eq.* 8). The computation of entropy, *S*, requires binning of spectral-power values (see Methods 3.4) to generate their probability distribution per d (sec) interval, which tends to underestimate entropy. We accommodated this underestimation of the true entropy, *S*, by the observed entropy, *S*_{obs}, by introducing a scaling factor a and an additive term b,

$$S_{obs} = aS + b$$
 — Eq. 12,

where $0 \le a \le 1$; *a* approaches 1 and *b* approaches 0 with a larger number of data points and finer spectral-power bins per interval. Substituting *Eq.* 12 into *Eq.* 11 yields,

 $S_{obs} = a \cdot [ln(\langle SP \rangle) + 1 - ln(k)] - Eq. 13.$

The parameter b has been absorbed in k because linear fitting cannot distinguish between b and k. As such, the observed value of k would be difficult to interpret.

Phase-scrambled spectral-power dynamics (which we have confirmed to obey *Eq.* 3; Figure 2) should obey *Eq.* 13 for intervals of any duration *d* (given that it includes sufficient data points to reliability evaluate the probability distribution of *SP*). To confirm this prediction, we divided each ~5 min EEG recording period into non-overlapping *d* (sec) intervals and computed average spectral power *<SP>* and entropy S_{obs} for each interval. The use of a longer interval, providing a larger number of *SP* values per interval, would make the relationship between In(*<SP>*) and S_{obs} tighter by increasing the accuracy of estimating *<SP>* and S_{obs} . However, it would reduce the variability in *<SP>* and S_{obs} across intervals (*Eq.* 4) and also reduce the number of In(*<SP>* $)-S_{obs}$ pairs to evaluate their relationship over time. We present our primary analyses with *d* = 3 sec; the choice of this particular duration will be justified below.

Each upper-left panel in Figure 3 shows, for the phase-scrambled controls, the 2D-density plot of $ln(\langle SP \rangle)$ - S_{obs} pairs for all d = 3 sec intervals for all frequencies, sites, and participants for a specific condition. Density is color-coded as percentile so that confidence intervals can be inferred. As predicted by *Eq.* 13, the relationship between $ln(\langle SP \rangle)$ and S_{obs} for the phase-

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scrambled controls was linear for all conditions, rest with the eyes closed, its replication, rest with the eyes open in dark, and the earlier and later viewing of a silent nature video.

We note that minor deviations from linearity occurred in the extreme ranges of spectral power for technical reasons. First, the binning of spectral-power values necessary to compute entropy per time interval (see Methods 3.4) caused an underestimation of entropy, generating the slight upward curvature in the lowest spectral-power range especially for ln[<SP>] < 0 (see the lower-left portions of the left panels in Figure 3) due to the floor effect (entropy > 0). Second, the use of a fixed maximum spectral-power bin (necessary to compute entropy over the same range of spectral-power bins in all cases) prevented extremely high-power values (though up to at least 99.9th percentile of the values were retained; see Methods 3.4) from contributing to the computation of entropy, causing an underestimation of entropy in the highest spectral-power range especially for ln[<SP>] > 5.5 (see the upper-right portions of the left panels in Figure 3). These extreme ranges of ln[<SP>] were excluded from the subsequent analyses (also from the computation of the linear fits shown in Figure 3).

Notwithstanding these minor deviations for the extreme values of $ln(\langle SP \rangle)$, the crucial observation is that the relationship between $ln(\langle SP \rangle)$ and S_{obs} were consistently linear for all conditions for the phase-scrambled controls, obeying *Eq.* 13. Because *Eq.* 13 derives from *Eq.* 3 (describing a maximum-entropy distribution), the linear relationships defined by the phase-scrambled controls indicate the *line of maximum entropy*.

Remarkably, the relationship between $ln(\langle SP \rangle)$ and S_{obs} for the real EEG data tightly clustered along the line of maximum entropy (the lower-left panels in Figure 3). Because average spectral power considerably varied depending on frequency, sites, and participants, the range of temporal variations in $ln(\langle SP \rangle)$ and S_{obs} are obscured when all $ln(\langle SP \rangle)$ - S_{obs} pairs are simply plotted together. To focus on the temporal variation in $ln(\langle SP \rangle)$ and S_{obs} , we aligned the 2D-density plot for the phase-scrambled control for each frequency, site, and participant at its center at (0,0) and equivalently translated the density plots for the corresponding real EEG data.

The centered relationships between $ln(\langle SP \rangle)$ and S_{obs} are shown in the main panels in Figure 3. The 2D-densoty plots for the phase-scrambled controls are shifted upward to avoid overlaps with those for the real EEG data, with the parallel gray dashed oblique lines indicating the line of maximum entropy. It is clear that the ranges of average spectral power $\langle SP \rangle$ (for d =3 sec intervals) were substantially extended in the real EEG data relative to their phasescrambled controls while consistently following the line of maximum entropy. This pattern appears to be universal, observed in all conditions (the main panels in Figure 3), all representative frequencies per condition (Figure 4), and all participants (Figures S1-S2). These

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results suggest that spectral-power dynamics maintain maximum entropy on the timescale of a few seconds while generating large power variations (relative to phase-scrambled controls) by changing the probability of synchronization on slower timescales (*Eq*s. 5-6).



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Fig 3. Relationship between log average spectral power In(<SP>) and entropy S_{obs} for d = 3 sec intervals. Each set of three panels shows the data for a specific condition: 5-min rest with the eyes closed (Rest EC), its replication (Rest EC rep), 5-min rest with the eyes open in dark (Rest EO DK), and the earlier and later 5-min viewing of a silent nature video (Nature video). For each condition, the $ln(\langle SP \rangle)$ - S_{obs} pair was computed for each non-overlapping d = 3sec interval per frequency per site per participant. Upper-left panels. 2D-density plots of all In(<SP>)-S_{abs} pairs for the phase-scrambled controls. The linear fits (gray dashed oblique lines) indicate the line of maximum entropy indicative of spectral power fluctuations that maximize entropy for a given value of average spectral power (see text). Lower-left **panels.** 2D-density plots of all $In(\langle SP \rangle)$ - S_{obs} pairs for the real EEG data. Note that the distributions follow the line of maximum entropy (the gray dashed oblique lines) defined by the phase-scrambled controls. Main panels. Re-plotting of the 2D-density plots for both the phase-scrambled controls and the real EEG data after aligning the phasescrambled 2D-density plot for each frequency, site, and participant at its center at (0,0) and equivalently translating the corresponding real-data density plots. The 2D-density plots for the phase-scrambled controls are shifted upward to avoid overlaps with those for the real EEG data. The centering shows that the dynamic ranges of average spectral power (per d = 3 sec interval) were substantially extended along the line of maximum entropy (the gray dashed oblique lines) for the real EEG data relative to their phase-scrambled controls in all conditions. This pattern was observed for all representative frequencies (Figure 4) and participants (Figures S1-S2). Thus, on the timescale of up to about 3 sec, spectral power appears to be controlled in such a way that the dynamic ranges are substantially extended (relative to stochastic dynamics) while tightly conforming to the line of maximum entropy. All panels. Density is color-coded as percentile so that confidence intervals can be inferred. The extreme ranges of spectral power, $ln(\langle SP \rangle) < 0$ and $ln(\langle SP \rangle) > 5.5$ were excluded from the computations of the line of maximum entropy and the centered 2D-density plots (the main panels) to avoid the binning-related distortions (see text).

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Fig 4. The same as the main panels in Figure 3, but the centered relationships are shown separately for the representative frequencies (rows) and conditions (columns). The phase-scrambled distributions are shifted upward to avoid overlaps with the real-data distributions, and the gray dashed oblique lines indicate the line of maximum entropy. Note that the dynamic ranges of average spectral power were substantially extended along the line of maximum entropy for the real EEG data relative to their phase-scrambled controls for all representative frequencies in all conditions.

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To examine how closely the spectral-power dynamics followed the line of maximum entropy we computed the probability distributions of entropy around the line of maximum entropy for the real EEG data and their phase-scrambled controls. While the line of maximum entropy was virtually identical for all participants (e.g., Figures S1 and S2), here we computed it separately for each participant to increase the accuracy in estimating entropy distributions around it. The bins to compute the distributions were determined by the maximum (*max*), minimum (*min*), and inter-quartile range (*iqr*) of the entropy distributions for the relevant real EEG data and their phase-scrambled controls (per participant), with the number of bins given by, $N_{bins} = (max, min)$

 $ceil\left(\frac{max-min}{2 \cdot iqr \cdot N_{intervals}}\right)$ (Freedman & Diaconis, 1981), where $N_{intervals}$ is the number of d (sec) intervals for which $In(\langle SP \rangle)$ - S_{obs} pairs were computed.

These probability distributions are plotted in Figure 5A for the five conditions for representative interval durations, d = 1, 3, 10, 20, 40, and 90 sec. The negative and positive values on the *x*-axis indicate the negative and positive deviations from the line of maximum entropy, *x*-scale is normalized so that all distributions for the phase-scrambled controls have the same standard deviation, and the *y*-axis indicates probability density. The shaded areas represent the distributions for the phase-scrambled controls (symmetric about the line of maximum entropy regardless of *d* as expected), the solid curves represent the distributions for the real data, and the solid areas represent the regions where the probability density was higher for the real data than for their phase-scrambled controls.

The real and phase-scrambled distributions were virtually indistinguishable for d = 1 sec and d = 3 sec for all five conditions (the two left columns in Figure 5A), indicating that the real data tightly followed the line of maximum entropy up to d = 3 sec. For longer interval durations, the real distributions progressively extended in the lower-entropy direction, seen as the solid-colored negative tails increasing in the third through the last column in Figure 5A. We quantified these lower-entropy tails for the real EEG data by computing the *proportions of lower-entropy intervals* (**PrLEI**) for the real data relative to their phase-scrambled controls. Specifically, for each distribution we computed the real minus phase-scrambled probability density wherever the density was higher for the real data than for the phase-scrambled controls, and summed those differences (multiplied by the bin width to convert to proportions) separately on the negative and positive sides, then subtracted the sum on the positive side from the sum on the negative side. This algorithm essentially yielded the proportion of the real-data distribution while compensating for any changes in distribution widths (approximately corresponding to the solid-colored negative tails of the real data in Figure 5A). For example, PrLEI = 0.2 would indicate

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that for a given interval duration d, the occurrences of lower-entropy intervals for the real data were 20% more frequent than for their phase-scrambled control.

We computed PrLEI values per participant per condition and plotted them as a function of interval duration *d* in Figure 5B. The circular symbols connected with thick lines indicate the median PrLEI values with the thin dotted lines showing the values for the individual participants. While PrLEI became large for longer interval durations (note the *y*-axis is reversed), the median PrLEI values remained small (< 5%) and condition independent up to about *d* = 3 sec. This indicates that up to the timescale of a few seconds, only up to about 5% of intervals of the real EEG data (on average) more negatively deviated from the line of maximum entropy than their phase-scrambled controls. That is, on average, greater than 95% of spectral-power dynamics followed the line of maximum entropy on the timescale of a few seconds. Even at the level of individual participants, only a few (out of 52), yielded PrLEI values greater than 10% for *d* = 3 sec (the dotted lines in Figure 5B).

The PrLEI values (for d = 3 sec) were consistently low for all frequencies for all conditions (Figure 6A) and globally low at all sites for all conditions (Figure 6B). Nevertheless, the data potentially suggest an interesting spatial pattern. We *z*-transformed the PrLEI values across sites per participant to quantify the consistency of regional deviations in PrLEI from the spatial average as *t*-values (with |t|>3.95 for Bonferroni-corrected 2-tailed significance at $\alpha = 0.05$) (Figure 6C). Cooler colors indicate regions with lower-than-average PrLEI values while warmer colors indicate regions with higher-than-average PrLEI values. In the eyes-open conditions, entropy was near maximal (very low PrLEI values) in the mid-central-posterior region (the dark blue regions highlighted with dotted circles in the lower three rows of Figure 6B and 6C). At the same time, consistent elevations in the PrIEI values (though still low with the means of less than 8.7% for all sites for all conditions) were observed in areas surrounding the mid-central-posterior region (Figure 6C). In particular, in the eyes-closed conditions the PrLEI values were focally elevated in the right-lateral region (the upper two rows in Figure 6C).



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Fig 5. Probability distributions of entropy S_{obs} relative to the line of maximum entropy for the real EEG data and their phase-scrambled controls as a function of interval duration d. A. Probability distributions for the phase-scrambled controls (shaded areas) and real EEG data (solid outlines), with the line of maximum entropy labeled as 0 on the x-axis and the negative tails of the real-data distributions toward lower entropy shown as solid areas. The rows correspond to the five conditions and the columns correspond to the representative interval durations d (sec). The probability distributions have been normalized so that the standard deviations are equalized for all phase-scrambled-control distributions. Note that up to about d = 3 sec, the distributions for the real EEG data and their phase-scrambled controls virtually overlap. B. Proportions of lower-entropy intervals (PrLEI) for the real EEG data relative to their phase-scrambled controls (approximately the solid-colored negative tails shown in A) as a function of interval duration d (sec). This measure indicates the proportions of d (sec) intervals for which the real EEG data yielded lower entropy than predicted by the line of maximum entropy. The circular symbols connected with thick lines indicate the median PrLEI values with the five conditions color-coded as in A (the black dotted lines indicating the replication of the rest-with-the-eyes-closed condition and the blue dotted lines indicating the later viewing of the nature-video condition) with the thin dotted lines showing the PrLEI values for the individual participants. Note that for the interval durations up to about d = 3 sec the real EEG data closely followed the line of maximum entropy with less than ~5% deviations (in median PrLEI values) across all conditions, suggesting that neural dynamics on the spatial-scale of EEG current sources generally maintain maximum entropy up to the timescale of a few seconds (see text).

Interval duration d (sec)

0.8

0.9

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Fig 6. Proportions of lower-entropy intervals (PrLEI) for d = 3 sec for the real EEG data relative to their phasescrambled controls as a function of frequency and site. PrLEI indicates the proportion of d (sec) intervals for which the real EEG data had lower entropy than predicted by the line of maximum entropy. **A.** PrLEI as a function of frequency. The thick lines indicate the median PrLEI values with the thin lines showing the values for the individual participants. The rows correspond to the five conditions. Note that the median PrLEI values were low regardless of frequency or condition. **B.** PrLEI as a function of site. The rows correspond to the five conditions. The mean PrLEI values were globally low across all sites and conditions. The mid-central-posterior region (highlighted with dotted circles) yielded particularly low PrLEI values in the eyes-open conditions (rest-with-the-eyes-open-in-dark and nature-video conditions) (the lower three topoplots). **C.** Same as B, but the data from each participant were z-transformed across sites to quantify the consistency of regional deviations of PrLEI values from the spatial average as t values (with |t|>3.95 for Bonferroni-corrected 2-tailed significance at $\alpha = 0.05$). Cooler colors indicate regions with lower-than-average PrLEI values while warmer colors indicate regions with higher-than-average PrLEI values. The t-values confirm that the PrLEI values were consistently low in the mid-central-posterior region in the eyes-open conditions (see B). Further, consistent elevations in the PrLEI values (though still low with the means of less than 8.7% for all sites and conditions) occurred in areas surrounding the midcentral-posterior region, particularly in the right-lateral region in the eyes-closed conditions (the upper two topoplots).

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Discussion

The dynamics of EEG spectral power deviate from stochastic dynamics in a U-shaped manner, such that the occurrences of the lowest and highest ranges of power are elevated (Figure 2). We used a simple mathematical model of synchronization dynamics to investigate the mechanisms that generate these characteristic deviations.

We modeled synchronization dynamics as simple chain reactions, where synchronization sequentially spreads from neural-unit to neural-unit at each moment with the probability p_{term} for the rapid spreading to terminate (*Eq.* 1). Although one may question the physiological relevance of postulating synchronization to independently spread at each time moment, the model (*Eq.* 1) is useful in the sense that it provides a simple computational mechanism that generates synchronization dynamics that maximize entropy for a given temporal average (*Eq.* 3). Note that the inferences that we have drawn are valid irrespective of the physiological plausibility of the specific model because they are based on how EEG spectral-power dynamics obeyed the rule of maximum entropy (i.e., *Eq.* 13 derived from *Eq.* 3). If the simple chain-reaction model of synchronization dynamics (*Eq.* 1) were physiological relevant, the parameter p_{term} could be interpreted as the probability of termination of the sequential spreading of synchronization. If not, p_{term} directly related to the temporal average of the size of synchronized population (*Eq.* 5).

We assumed that EEG spectral power was proportional to the size of the synchronously oscillating neural population accessible at each site. For a constant p_{term} , the model predicted stochastic dynamics (*Eq.* 3) with the temporal variation of spectral power exponentially distributed. The model further predicted that if p_{term} remained constant the fluctuations of log average spectral power and entropy should be associated along the line of maximum entropy (*Eq.* 13 derived from Eq. 3) on any timescale. These predictions were confirmed for the phase-scrambled controls (Figure 5A). *Eq.* 13 further predicted that even if p_{term} substantially varied as a function of time, if it remained relatively constant up to some timescale *d* (sec) the fluctuations of log average spectral power and entropy should still be associated along the line of maximum entropy up to that timescale (*Eqs.* 5, 6).

The results clearly showed that up to a few seconds (d = 3 sec), the dynamics of the EEG data closely followed the line of maximum entropy (Figure 3) for all representative frequencies (Figure 4) and all participants (Figures S1-S2) whether they rested with their eyes closed, rested with their eyes open in a darkened room, or viewed a silent nature video. The EEG dynamics

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were nearly as tightly distributed around the line of maximum entropy as were their phasescrambled controls (see the d = 3 sec column highlighted with the rectangle in Figure 5A). Any systematic deviations from the line of maximum entropy were small up to d = 3 sec with the median PrLEI values remaining low for all frequencies and conditions, especially for the eyesopen conditions where the median values were consistently below ~5% (Figure 6A). These results suggest that the majority (~95%) of the low- and high-spectral-power states that deviated from stochastic dynamics (Figure 2) were generated by relatively slow mechanisms that generally maintain maximum entropy within the timescale of a few seconds while changing the probability of synchronization on slower timescales to substantially extend the dynamic range of spectral power along the line of maximum entropy (Figure 3 [the main panels], Figure 4, and Figures S1 and S2).

This conclusion may seem counterintuitive because precisely coordinated actions and mental operations require tight controls of oscillatory neural dynamics. One possibility is that the large-scale spatial synchronization of oscillatory activity may be controlled by a combination of the relatively slow, entropy-maximizing adjustments of synchronization probability and the temporally precise adjustments of phase such as phase-resetting generated by sensorimotor interactions. For example, the inter-regional and cross-frequency coordination of large-scale oscillatory activity may be generally controlled by slowly co-varying the probabilities of synchronization across the relevant regions and frequency bands while maintaining maximum entropy on the timescale of a few seconds for energy efficiency. At the same time, the precisely timed coordination of inter-regional and cross-frequency synchronization of oscillatory activity may utilize phase-resetting initiated by punctate sensorimotor signals such as those generated by multisensory environmental stimuli as well as goal-directed and routine sensorimotor events such as blinks, saccades, and active touch (e.g., Rajkai et al., 2008; Fiebelkorn et al., 2011; Mercier et al., 2013; Thorne & Debener, 2014; Wutz et al., 2014; Sugiyama et al., 2019; see Ding & Simon, 2013 and Voloh & Womelsdorf, 2016 for reviews).

While the deviations from the line of maximum entropy were globally low on the timescale of a few seconds at all sites for all conditions (Figure 6B), we observed some notable spatial patterns. In particular, in the eyes-open conditions entropy was near maximal in the mid-central-posterior region (highlighted with dotted circles in the lower three rows in Figure 6B and 6C). This potentially suggests that, especially in the presence of substantial sensory input (the eyes-open conditions here), the mid-central-posterior region plays the role of an "entropy dump" to facilitate the temporally precise control of spectral-power dynamics in the surrounding regions. In the eyes-closed conditions, we observed focal PrLEI elevations in the right-lateral region (the

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upper two rows in Figure 6C), potentially suggesting that this cortical region plays a role in controlling synchronization dynamics for constructing spontaneous imagery and thoughts (which were encouraged in the rest-with-the-eyes-closed condition). While these interpretations are highly speculative, analyzing deviations of spectral-power dynamics from the line of maximum entropy may provide a useful method for tracking the spatiotemporal occurrences of temporally precise controls of large-scale spectral dynamics.

In summary, we used a simple mathematical model of synchronization to investigate the mechanisms that make EEG spectral-power dynamics deviate from stochastic dynamics in a characteristic U-shaped manner (Figure 2). The results have clearly shown that the majority (~95%) of this control is universally (across frequencies, sites, and behavioral conditions) accomplished by slowly changing the probability of synchronization while maintaining maximum entropy on the timescale of a few seconds. The results may further suggest that the mid-central-posterior region potentially serves as an entropy dump to facilitate the generation of precisely controlled spectral-power dynamics in the surrounding regions.

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References

Arnal, L.H., Giraud, A.L., 2012. Cortical oscillations and sensory predictions. *Trends in Cognitive Sciences*, *16* (7), 390-398.

Bonnefond, M, & Jensen, O. (2015). Gamma activity coupled to alpha phase as a mechanism for top-down controlled gating. *PLoS ONE, 10(6)*: e0128667, 1-11.

Busch, N. A., & VanRullen, R. (2010). Spontaneous EEG oscillations reveal periodic sampling of visual attention. *Proceedings of the National Academy of Sciences, 107(37)*, 16048-16053.

Buzsaki, G. (2006). Rhythms of the Brain, Oxford University Press.

Cohen, M. X. (2014). Analyzing Neural Time Series Data: Theory and Practice, MIT Press.

Delorme, A., & Makeig, S. (2004). EEGLAB: an open source toolbox for analysis of single-trial EEG dynamics including independent component analysis. *Journal of neuroscience methods*, *134*(1), 9-21.

Ding, N., & Sinom, J. J. (2013). Power and phase properties of oscillatory neural responses in the presence of background activity. *Journal of Computational Neuroscience, 3*4, 337-343.

Donner, T. H., & Siegel, M. (2011). A framework for local cortical oscillation patterns. *Trends in Cognitive Sciences*, *15*(5), 191-199.

Duzel, E., Penny, W. D., & Burgess, N. (2010). Brain oscillations and memory. *Current Opinion in Neurobiology, 20*, 143-149.

Engel, A. K., & Fries, P. (2010). Beta-band oscillations—signaling the status quo? *Current Opinion in Neurobiology, 20*, 156-165.

Fiebelkorn, I. C., Foxe, J. J., Butler, J. S., Mercier, M. R., Snyder, A. C., & Molholm, S. (2011). Ready, set, reset: stimulus-locked periodicity in behavioral performance demonstrates the consequences of cross-sensory phase reset. *Journal of Neuroscience*, *31*(*27*), 9971–9981.

Menceloglu et al. 25

Freedman, D. and Diaconis, P. (1981). On the histogram as a density estimator: L2 theory. *Probability Theory and Related Fields, 57 (4)*, 453-476.

Fries, P. (2005). A mechanism for cognitive dynamics: neural communication through neural coherence. *Trends in Cognitive Sciences*, *9(10)*, 474-480.

Gao, R., Peterson, E. J., & Voytek, B. (2017). Inferring synaptic excitation/inhibition balance from field potentials. *Neuroimage*, *158*, 70-78.

He, B. J. (2014). Scale-free brain activity: past, present, and future. *Trends in Cognitive Sciences*, *18*(9), 480-487.

Hess, R. F. & Snowden, R. J., (1992) Temporal properties of human visual filters: Number, shapes and spatial covariance, *Vision Research*, *32*, 47-60.

Hipp, J. F., Hawellek, D. J., Corbetta, M., Siegel, M., & Engel, A. K. (2012). Large-scale cortical correlation structure of spontaneous oscillatory activity. *Nature Neuroscience*, *15(6)*, 884-890.

Hjorth, B. (1980). Source derivation simplifies topographical EEG interpretation. *American Journal of EEG Technology 20*, 121-132.

Kayser, J., Tenke, C. E. (2006). Principal components analysis of Laplacian waveforms as a generic method for identifying ERP generator patterns: I. Evaluation with auditory oddball tasks. *Clinical Neurophysiology*, *117*, 348-368.

Kiya, H., Ito, I., & Miron, S. (2010). "Phase scrambling for image matching in the scrambled domain" in Signal Processing, *InTech*, 97-414.

Klimesch, W. (2012). Alpha-band oscillations, attention, and controlled access to stored information. *Trends in Cognitive Sciences*, *16(12)*, 606-617.

Koch, C. (1999). Biophysics of Computation: Information Processing in Single Neurons. Oxford.

Menceloglu et al. 26

Lopez-Calderon, J., & Luck, S. J. (2014). ERPLAB: an open-source toolbox for the analysis of event-related potentials. *Frontiers in human neuroscience*, *8*, 213.

Lui, L. L., Bourne, J. A., Rosa, M. G. P. (2007). Spatial and temporal frequency selectivity of neurons in the middle temporal visual area of new world monkeys (Callithrix jacchus). *European Journal of Neuroscience*, *25*, 1780-1792.

Mathewson, K. E., Gratton, G., Fabiani, M., Beck, D. M., & Ro, T. (2010). To see or not to see: prestimulus alpha phase predicts visual awareness. *Journal of Neuroscience, 29*, 2725-2732.

Mathewson, K. E., Lleras, A., Beck, D. M., Fabiani, M., Ro, T., Gratton, G. (2011). Pulsed out of awareness. EEG alpha oscillations represent a pulsed-inhibition of ongoing cortical processing. *Frontiers in Psychology, 2, A99*, 1-15.

Mazzoni, A., Panzeri, S., Logothetis, N. K., & Brunel, N. (2008). Encoding of naturalistic stimuli by local field potential spectra in networks of excitatory and inhibitory neurons. *PLoS Computational Biology*, *4*(*12*): e1000239. doi:10.1371/journal.pcbi.1000239

Makeig, S., Bell, A. J., Jung, T.-P. and Sejnowski, T. J. (1996). "Independent component analysis of electroencephalographic data," In: D. Touretzky, M. Mozer and M. Hasselmo (Eds). Advances in Neural Information Processing Systems 8:145-151, MIT Press, Cambridge, MA.

Makeig, Scott et al. "EEGLAB: ICA Toolbox for Psychophysiological Research". WWW Site, Swartz Center for Computational Neuroscience, Institute of Neural Computation, University of San Diego California <www.sccn.ucsd.edu/eeglab/>, 2000. [World Wide Web Publication].

Menceloglu, M., Grabowecky, M., & Suzuki, S. (under review). Spectral-power associations reflect amplitude modulation and within-frequency interactions on the sub-second timescale and cross-frequency interactions on the seconds timescale.

https://biorxiv.org/cgi/content/short/2020.01.15.908103v1

Menceloglu et al. 27

Mercier, M. R., Foxe, J. J., Fiebelkorn, I. C., Butler, J. S., Schwartz, T. H., & Molholm, S. (2013). Auditory-driven phase reset in visual cortex: human electrocorticography reveals mechanisms of early multisensory integration. *NeuroImage*, *79*, 19–29.

Michalareas, G., Vezoli, J., van Pelt, S., Schoffelen, J.-M., Kennedy, H., & Fries, P. (2016). Alpha-beta and gamma rhythms subserve feedback and feedforward influences among human visual cortical areas. *Neuron, 89*, 384-397.

Müller, V., Perdikis, D., Oertzen, T., Sleimen-Malkoun, R, Jirsa, V., & Lindenberger, U. (2016). Structure and topology dynamics of hyper-frequency networks during rest and auditory oddball performance. *Frontiers in Computational Neuroscience, 10, Article 108*, 1-25.

Palva, S., & Palva, M. (2007). New vistas for a-frequency band oscillations. *Trends in Neurosciences*, *30*(*4*), 150-158

Palva, S., & Palva, J. M. (2012). Discovering oscillatory interaction networks with M/EEG: challenges and breakthroughs. *Trends in Cognitive Sciences*, *16*(*4*), 219-230.

Palva, J. M., Palva, S., & Kaila, K. (2005). Phase synchrony among neuronal oscillations in the human cortex. *Journal of Neuroscience*, *25(15)*, 3962-3972.

Perrin, F., Pernier, J., Bertrand, O., & Echallier, J. F. (1989a). Corrigenda EEG 02274. *Electroencephalography and Clinical Neurophysiology* 76, 565.

Perrin, F., Pernier, J., Bertrand, O., & Echallier, J. F. (1989b). Spherical Splines for Scalp Potential and Current Density Mapping. *Electroencephalography and Clinical Neurophysiology 72(2)*: 184-187.

Perrin, F., Pernier, J., Bertrand, O., Giard, M. H., & Echallier, J. F. (1987). Mapping of Scalp Potentials by Surface Spline Interpolation. *Electroencephalography and Clinical Neurophysiology*, *66 (1)*, 75-81.

Rajkai C., Lakatos P., Chen C. M., Pincze Z., Karmos G., Schroeder C. E. (2008). Transient cortical excitation at the onset of visual fixation. *Cerebral Cortex*, *18(1)*, 200–209.

Menceloglu et al. 28

Spaak, E., Bonnefond, M., Maier, A., Leopold, D. A., & Jensen, O. (2012). Layer-specific entrainment of gamma-band neural activity by the alpha rhythm in monkey visual cortex. *Current Biology*, *22*(*24*), 2313–2318.

Tenke, C. E., & Kayser, J. (2012). Generator localization by current source density (CSD): Implications of volume conduction and field closure at intracranial and scalp resolutions. *Clinical Neurophysiology*, *123*, 2328–2345.

Sugiyama, S., Kinukawa, T., Takeuchi, N., Nishihara, M., Shioiri, T., & Inui, K. (2019). Tactile cross-modal acceleration effects on auditory steady-state response. *Frontiers in Integrative Neuroscience*, 13:72. <u>doi.org/10.3389/fnint.2019.00072</u>

Thorne, J. D., & Debener, S. (2014). Look now and hear what's coming: On the functional role of cross-modal phase reset. *Hearing Research*, *307*, 144–152.

Voloh, B., & Womelsdorf, T. (2016). A role of phase-resetting in coordinating large scale neural networks during attention and goal-directed behavior. *Frontiers in Systems Neuroscience*, 10:18. doi: 10.3389/fnsys.2016.00018

Voytek, B., Canolty, R. T., Shestyuk, A., Crone, N. E., Parvizi, J., & Knight, R. T. (2010). Shifts in gamma phase-amplitude coupling frequency theta to alpha over posterior cortex during visual tasks. *Frontiers in Human Neuroscience, 4, A191*, 1-9.

Wutz, A., Weisz, N., Braun, C., & Melcher, D. (2014). Temporal windows in visual processing: "prestimulus brain state" and "poststimulus phase peset" segregate visual transients on different temporal scales. *Journal of Neuroscience*, *34*(4), 1554–1565.

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Supplementary figures

Figure S1. Individual participants' data for the main panels in Figure 3 for participants p1p28 who participated in the rest-with-the-eyes-closed condition, the nature-video condition, or both.

Figure S2. Individual participants' data for the main panels in Figure 3 for participants p29-p52 who participated in the replication of the rest-with-the-eyes-closed condition and the rest-with-the-eyes-open-in-dark condition.

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Fig S1. Individual participants' data for the main panels in Figure 3 for participants p1-p28 who participated in the restwith-the-eyes-closed condition, the nature-video condition, or both. All participants who participated in the nature-video condition provided data for both earlier and later viewings. The dynamic ranges of average spectral power (for d = 3 sec intervals) were moderately to substantially extended along the line of maximum entropy (the gray dashed oblique lines) for the real EEG data relative to their phase-scrambled controls for all participants for all conditions. Note that the degree of extension of spectral-power dynamic range does not appear to be a trait-like property as it substantially varied for some participants between conditions.



Fig S2. Individual participants' data for the main panels in Figure 3 for participants p29-p52 who participated in the replication of the rest-with-the-eyes-closed condition and the rest-with-the-eyes-open-in-dark condition. The dynamic ranges of average spectral power (for d = 3 sec intervals) were moderately to substantially extended along the line of maximum entropy (the gray dashed oblique lines) for the real EEG data relative to their phase-scrambled controls for all participants for all conditions. Note that the degree of extension of spectral-power dynamic range does not appear to be a trait-like property as it substantially varied for some participants between the two similar conditions.