

Review

Broadband changes in the cortical surface potential track activation of functionally diverse neuronal populations[☆]

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ABSTRACT

We illustrate a general principal of electrical potential measurements from the surface of the cerebral cortex, by revisiting and reanalyzing experimental work from the visual, language and motor systems. A naive decomposition technique of electrocorticographic power spectral measurements reveals that broadband spectral changes reliably track task engagement. These broadband changes are shown to be a generic correlate of local cortical function across a variety of brain areas and behavioral tasks. Furthermore, they fit a power-law form that is consistent with simple models of the dendritic integration of asynchronous local population firing. Because broadband spectral changes covary with diverse perceptual and behavioral states on the timescale of 20–50 ms, they provide a powerful and widely applicable experimental tool.

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Contents

Introduction	712
Methods	712
Human subjects	712
Electrocorticographic recording	714
Cortical rendering and electrode localization	714
Spectral analysis and decomposition (Fig. 1)	714
Tasks	714
Face-house picture task (Figs. 1 and 2)	714
Visual search task (Fig. 3)	714
Verb generation task (Fig. 4)	715
Finger movement task (Fig. 5)	715
Results and discussion	715
Decoupling the cortical power spectrum to reveal broadband spectral changes	715
Linking behavior to asynchronous broadband activity across tasks and brain regions	715
The significance of the power law form of the broadband spectral change	717
Examples of genuinely rhythmic high-frequency activity	718
How broadband changes have gained attention	718
Conclusion	719
References	719

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Introduction

When examining the extracellular voltage power spectrum of occipital or Rolandic cortex, the shift from rest to task is strongly associated with an increase in power at high frequencies (greater than ~50 Hz), and often accompanied by a decrease in power at low frequencies (less than ~30 Hz) (Aoki et al., 1999; Crone et al., 1998a,b, 2001; Miller et al., 2007, 2010b). When we first investigated these signals, our interpretation was that there would be different high-frequency oscillations specific to each cortical region and function. However, in each case that we examined the high-frequency spectral changes lacked a definitive upper bound, and appeared to extend to the highest frequencies resolvable with our sampling rate and noise floor.

This presented a pair of questions. First, might the high-frequency spectral changes reflect a broadband change in the power spectrum? Second, might this broadband change track asynchronous processes in the underlying neuronal dynamics? In this paper, we review and reanalyze data supporting the claim that broadband shifts are indeed a generic marker of circuit activation in the neocortex (Manning et al., 2009; Miller, 2010; Miller et al., 2009b,d; Ray and Maunsell, 2011; Whittingstall and Logothetis, 2009). Moreover, we discuss how broadband spectral change can be obscured by rhythmic phenomena at lower (e.g. α , β) frequencies, and we illustrate how the changes in high frequencies are different in kind from changes in the canonical lower-frequency rhythms.

Before describing the details of the broadband power shifts, it is important to distinguish rhythmic and non-rhythmic neural processes and their manifestations in the field potential. Rhythmic processes (such as the occipital α and peri-central motor β rhythms) exhibit band-limited spectral peaks, reflective of a characteristic oscillatory time-scale (Bates, 1951; Crone et al., 1998b; Jasper, 1941; Jasper and Andrews, 1938; Jasper and Penfield, 1949; Miller et al., 1940, 2007; Penfield, 1954; Pfurtscheller, 1999). Non-rhythmic processes do not have an oscillatory timescale and do not produce isolated peaks in the power spectrum. Nonetheless, a non-rhythmic process may exhibit a distinctive spectral profile, and changes in its amplitude will produce changes in the spectrum of the electrical field (Bullock et al., 2003).

We have proposed (Miller et al., 2009b) that the broadband “1/f” background, ubiquitous in neocortical field spectra, reflects a non-rhythmic underlying process with an intuitive connection to neuronal population activity. In particular, the approximately power-law profile of the broadband component can be understood as the signature of the summation of asynchronously arriving synaptic inputs in the dendrites of pyramidal neurons (Bedard et al., 2006; Miller et al., 2009b). The dendritic summation model predicts a particular form of power-law in the spectrum above 80 Hz, and this prediction matches the empirical spectral profile to >500 Hz (once one corrects for gain-modulation in amplifier hardware).

Regardless of the details of the biophysics that produce the broadband 1/f component, it is clear that (i) the underlying process is not rhythmic and (ii) the process is strongly associated with the activation of local circuits. Behavior-locked increases in the broadband component have been demonstrated in peri-Rolandic cortex (during finger tapping) and in the occipital lobe (during visual search) via intracranial recordings

from the surface of the human neocortex (Miller et al., 2009c, 2010a). More generally, task-locked power increases in the “high gamma” band of the power spectrum (~80–200 Hz) are widely observed in mammalian cortical electrophysiology (Canolty et al., 2007; Crone et al., 1998a, 2001; Edwards et al., 2010; Foster et al., 2012; Gunduz et al., 2011; Hermes et al., 2012; Jacobs and Kahana, 2009; Mesgarani and Chang, 2012; Miller et al., 2007; Pei et al., 2011; Sederberg et al., 2003; Swann et al., 2009; Vidal et al., 2010). In many cases, these wide-band power shifts are probably reflective of an underlying broadband shift, especially when there is no band-limited peak observed above 80 Hz.

Direct support for the broadband model was provided by simultaneous measurements of unit activity and field potentials. Based on 2030 identified units across numerous human neocortical sites, Manning et al. (2009) demonstrated a positive relationship between the average firing rate of the units and the amplitude of the broadband component. Recent work in macaque visual cortex found a robust relationship between broadband high-frequency power increases and single unit activity, and was also able to separate the (non-rhythmic) high-frequency broadband effect from those attributable to (rhythmic) visual gamma (Ray and Maunsell, 2011).

Because rhythmic (band-limited) and non-rhythmic (broadband) components of the power spectrum have distinct frequency profiles and exhibit distinct task-related changes, they can be decoupled from one another (Miller et al., 2009d) (Fig. 1). Once decoupling is performed, the spectrum of the electrical field can be expressed as a composite of rhythmic and non-rhythmic components (Figs. 1–5). A broadband shift can then be understood as a change in the amplitude of a non-rhythmic, stochastic, process, which reflects the dendritic integration of asynchronous inputs.

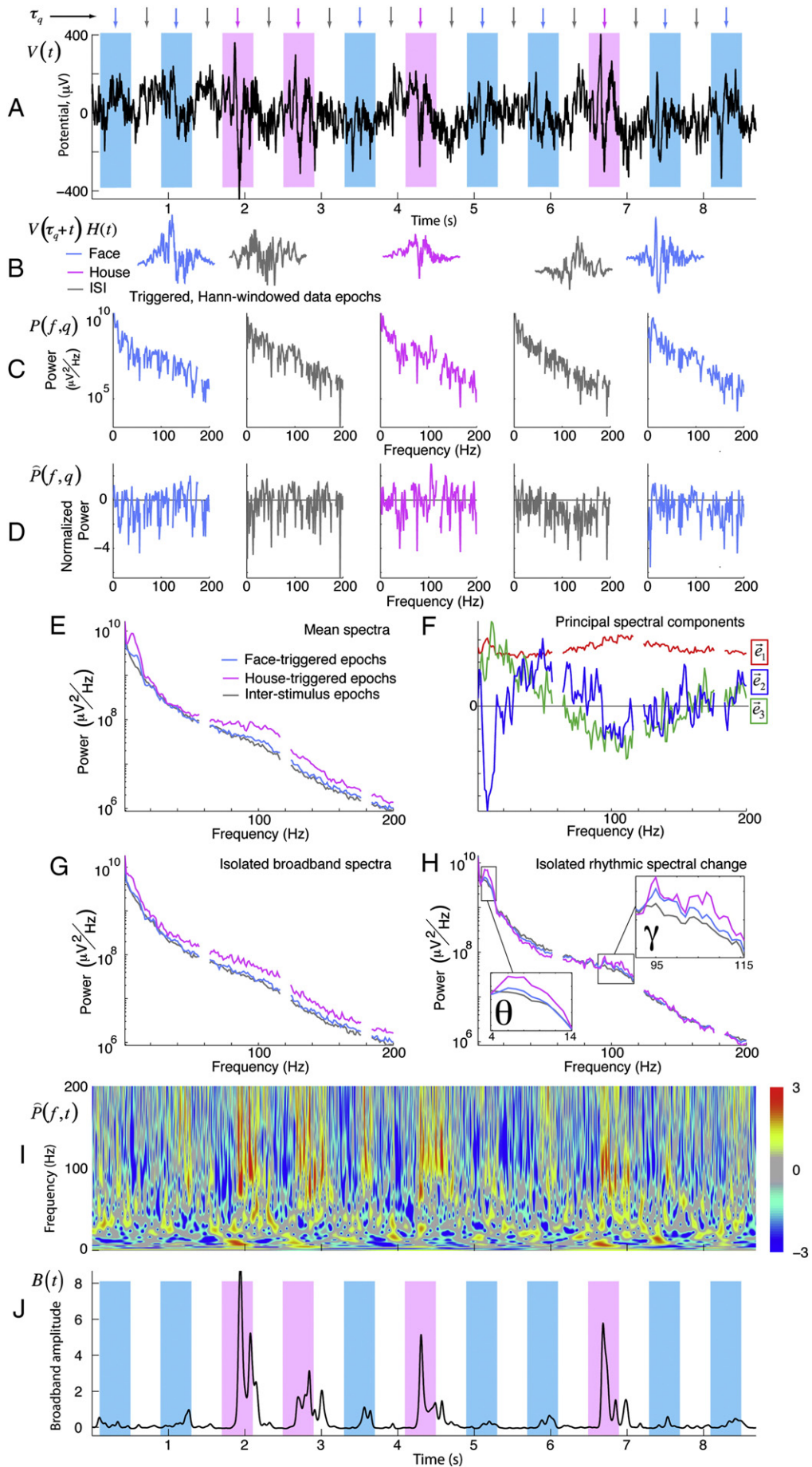
In this review, we begin by demonstrating that broadband shifts are a ubiquitous marker of local cortical activity; we present data from four different settings: face and house detection (inferotemporal cortex, Fig. 2), visual search (peri-Calcarine cortex, Fig. 3), speech production (peri-Sylvian and peri-Rolandic cortex, Fig. 4), and finger tapping (peri-Rolandic cortex, Fig. 5). In each setting, spatially focal shifts in broadband power are observed in a task-selective manner. We then illustrate how broadband power shifts may appear to be band-limited in certain contexts due to contamination by amplifier noise at high frequencies and masking due to the general anticorrelation of alpha and beta rhythm power at low frequencies (Fig. 6).

Methods

Human subjects

Five patients participated in the study, one patient each is reflected in Figs. 1–5. All were patients at Harborview Hospital in Seattle, WA, USA, with sub-dural electrocorticographic (ECoG) grids placed for extended clinical monitoring and localization of seizure foci, in the course of the treatment for medically-refractory epilepsy. All patients participated in a purely voluntary manner, after providing informed written consent, under a protocol approved by the Institutional Review Board of the University of Washington.

Fig. 1. Decoupling the cortical spectrum to reveal broadband spectral change. Simple black-and-white pictures of faces and houses were presented to a patient while recording from an ECoG electrode on the parahippocampal gyrus. (A) Raw potential, V , with time triggers, τ_q , in the middle of each stimulus (q) and ISI period (noted with arrows, 400 ms each). Times of face presentation are shown with pink background, with house presentations in blue. The gray triggers are the middle of ISI periods. (B) 1-s epochs of raw potential. Centered at each time τ_q , are extracted and Hann-windowed: $V(\tau_q + t)H(t)$. (C) The power spectral density (PSD) of each of these epochs is calculated: $P(f, q)$. (D) Each spectrum is normalized with respect to the mean spectrum across trials of all types, and the log is taken, to produce the transformed PSD: $P(f, q)$. (E) The average power spectrum is shown for all face/house/ISI presentations in pink/blue/gray. (F) Principal spectral components, \bar{e}_1 , \bar{e}_2 , \bar{e}_3 (in red, blue, and green; “PSCs”) of the normalized spectra (as in D) are identified, and reveal motifs of change in the power spectrum. The first is primarily broadband increase across all frequencies, whereas the 2nd to 4th typically capture changes in rhythmic, peaked phenomena of power spectral change. (G) Isolated broadband spectra for each stimulus type — the 2nd to 4th PSCs have been removed and the power spectra are reconstructed, and broadband increase across all frequencies is revealed during presentation of house stimuli. (H) Reconstructed power spectra of the 2nd to 4th PSCs only. Selective increase during house stimuli in the theta and high gamma range are extracted from the raw spectra and observed here. (I) The dynamic power spectral density after dividing by the average across all time, then taking the log: similar to D, but where the power spectrum is estimated at each point in time, using 7-cycle Morlet wavelets: $P(f, t)$. (J) The dynamic spectrum, $P(f, t)$, is projected into the 1st PSC, \bar{e}_1 , smoothed, z-scored, and exponentiated to obtain the timecourse of broadband spectral change, $B(t)$. Comparison with stimuli reveals that the cortex beneath this electrode is involved in visual processing of both face and house stimuli, but house stimuli more dramatically than face stimuli.



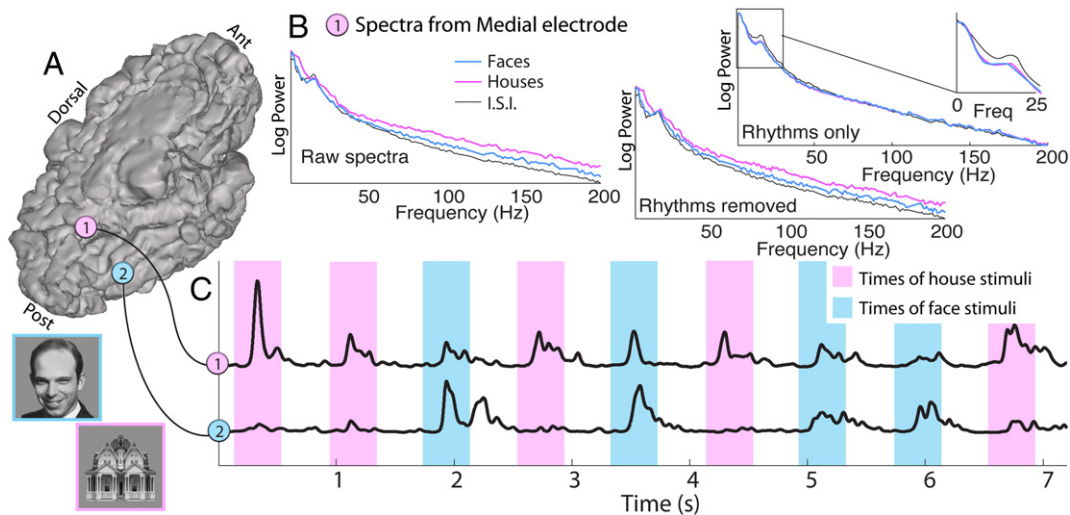


Fig. 2. Inferotemporal category-specific response. (A) A medial (“1”) and a lateral (“2”) inferotemporal electrode are shown in situ. (B) The raw power spectral density (“raw spectra”) from site “1” in panel (A) is decomposed as illustrated in Fig. 1. (C) The broadband time series from each site. Note that the lateral electrode shows selective increase during face pictures, and the medial electrode shows significant response during both face and house stimuli, but with larger response for house picture stimuli. Of the 300 presentations of face and house pictures, all 300 were correctly identified from the spontaneous stream of data (with 22 false positives, (Miller et al., 2009a)). The correct face or house label could be identified with 95% accuracy, with 23 ms precision (23 ms standard deviation).

Electrocorticographic recording

The platinum electrode arrays (Ad-Tech, Racine, WI) were configured as combinations of “grid” (4×8 , 8×8) and strip arrays. The electrode pads had 4 mm diameter (2.3 mm exposed), 1 cm inter-electrode distance, and were embedded in silastic. These arrays were surgically placed on the sub-dural brain surface during the treatment for epilepsy. ECoG signals were split into two identical sets. One set was fed into the clinical EEG system (XLTEK, Oakville, Ontario, Canada) and the other set was recorded with Synamps2 (Neuroscan, El Paso, TX) biosignal amplifiers at 1 kHz with an instrumental bandpass-filter from 0.3 Hz to 200 Hz. ECoG signals were acquired from the experimental amplifiers using the general-purpose BCI2000 software (Schalk et al., 2004), which was also used for visual stimulus presentation. Audio was recorded using a Logitech USB desktop microphone at 11 kHz, and synchronized with a trigger in the BCI2000 program (Fig. 4). Finger position was recorded using a sensor data glove (5DT, Irvine, CA, Fig. 5).

Cortical rendering and electrode localization

The relationship between electrode position and gyral anatomy was determined by first rendering the cortical surface from a pre-operative MRI, using either the freesurfer (Dale et al., 1999) or spm5 (Ashburner and Friston, 2005; Friston et al., 1995) environment. Then, electrode positions were calculated with respect to this pre-operative MRI from post-operative computed tomography (CT) using the CTMR package of Hermes et al. (2010), demonstrated to accurately localize the electrode positions within an error of ~4 mm (the same as the size of the electrodes).

Spectral analysis and decomposition (Fig. 1)

After rejection of artifactual or epileptiform electrodes, scalp-referenced ECoG potentials were re-referenced with respect to the common average reference across all remaining electrodes. From each electrode, samples of power spectral density (PSD; $P(f, \tau_q)$) as a function of frequency (f) were calculated from 1 second epochs centered at the midpoint of each “events” (different for each task), τ_q , where a Hann taper was applied, the Fourier transform was implemented, and the

resulting quantity was multiplied by its complex conjugate. These samples of power spectral density were then decomposed using a principal-component type approach to obtain “principal spectral components” (PSCs), as illustrated in Fig. 1 and described in the associated caption. Reconstruction of the PSDs from a subset of all of the PSCs differentiates underlying motifs, and the 2nd–4th PSCs were first removed, and then examined in isolation in Figs. 1–5.

Continuous time-frequency approximations (dynamic spectra) were calculated using a wavelet approach, and projected onto the 1st PSC. After smoothing (by filtering with a Gaussian window of width $\sigma = 50$ ms), z-scoring, and exponentiating, a “broadband time series”, $B(t)$, is obtained. There is a power law in the cortical PSD of the form $P(f, t) \sim A(t)f^{-\chi}$, which is revealed by broadband fluctuations in the PSD across all frequencies (separate from the classic θ , α , β , and γ rhythmic motifs) (Miller et al., 2009b,d), and appears to directly correlate with local neuronal population firing rate (Manning et al., 2009; Miller, 2010). The exponent χ is addressed in the discussion. We have proposed that $B(t)$ approximates multiplicative scaling in the timecourse of $A(t)$, and may be directly correlated with multiplicative factor in the average neuronal firing rate of neuronal populations. This manuscript demonstrates that $B(t)$ is highly correlated to task-specific dynamics at very short timescales, and is a general property across many brain areas.

Tasks

Face-house picture task (Figs. 1 and 2)

Pictures of simple, luminance and contrast matched, grayscale faces and houses (10 cm by 10 cm at 1 m distance) were displayed in random order for 400 ms each, with 400 ms inter-stimulus interval between. Patients were asked to report a simple target (a single upside-down house). “Event times,” τ_q , were denoted at midpoint of picture presentation or each blank screen in between stimuli. Fig. 2 results also are quantified in Miller et al. (2009a).

Visual search task (Fig. 3)

The patient surveyed an array of colored squares (free to saccade, without a fixation cross), on each trial stating the color of a box that was in a particular direction adjacent to a starred box. “Event times,”

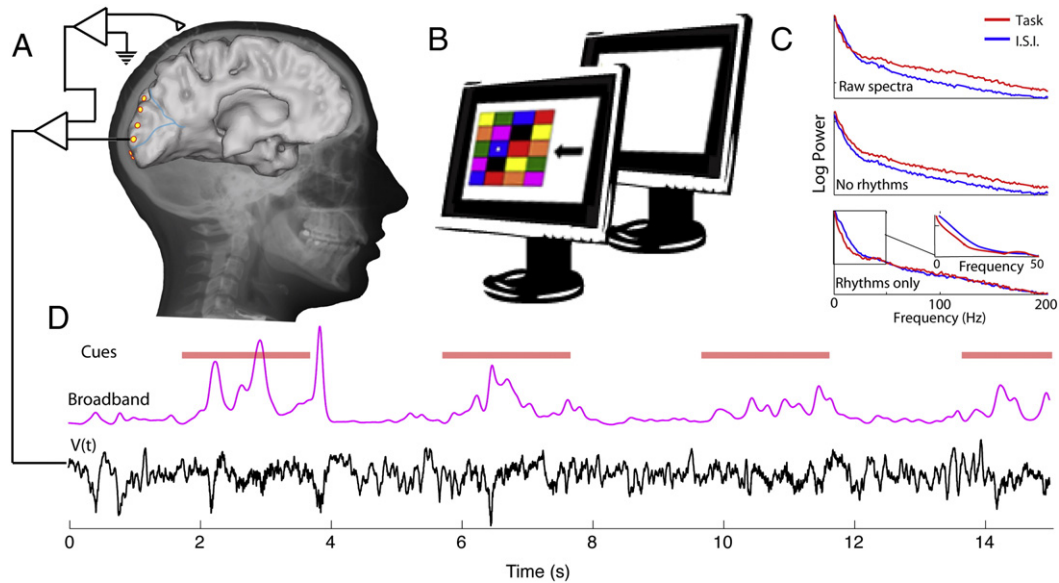


Fig. 3. Peri-calcarine broadband response during a visual search task. (A) The ECoG potential is measured from occipital lobe brain surface sites during participation in a visual search task, and decomposed as in Fig. 1. (B) The visual search task consists of a 4-by-5 array of colored squares. Each cue consists of a star in one of the boxes (in this case a blue box), and an arrow. The patient states the color of the box in the direction of the arrow from the star (“green” would be the appropriate response in this case). Visual search cues are shown for 2 s, with blank inter-stimulus-interval (ISI) cues shown for intervening 2 s. (C) As illustrated previously, power spectral changes during different portions of the experiment (top panel) may be appreciated and naively decoupled into broadband changes (middle panel, 2nd–4th PSCs omitted) and changes in brain rhythms (bottom panel, 2nd–4th PSCs only, with prominent theta and alpha range changes). (D) Broadband increases are observed during the various saccades required for processing of each cue. Figure modified from Miller et al. (2010a).

τ_q , were denoted at midpoint of box array presentation or blank screen in between. For a full description, see Miller et al. (2010a).

Verb generation task (Fig. 4)

Patients stated verbs that were connected to written nouns on a screen. “Event times,” τ_q , were denoted at times of speech onset or silent periods in between. For a full description of the task, see Miller et al. (2011). The data have been analyzed differently in this manuscript from the original.

Finger movement task (Fig. 5)

Patients performed self-paced movements of individual fingers in response to simple visual cues. “Event times,” τ_q , were denoted at times of peak finger displacements or points of rest. For a full description, see Miller et al. (2009d, 2012).

Results and discussion

Decoupling the cortical power spectrum to reveal broadband spectral changes

The decoupling approach illustrated in Fig. 1 is used to identify which spectral components covary as the PSD changes over time. The covarying frequency components are referred to as motifs. The strength of the linear spectral decomposition is that it makes few assumptions regarding the underlying structure of the motifs, except that they are orthogonal. The most robust phenomenon in the PSD, seen in every motif decomposition and across four different tasks, is a broadband increase that is roughly even in magnitude across frequencies when plotted in log space (Figs. 1–5). In addition, there are task-related changes in the amplitude of low frequency rhythms (usually but not always decreases), with a center frequency that varies across cortical locations.

Because the low frequency rhythms do not completely decohere during tasks, there is typically residual amplitude in the peaked portions of the PSD during task engagement. The naive mechanism used in this manuscript does not assume the form for the underlying spectral motifs,

and the underlying rhythms are frequently present in the background shape of the “broadband” decoupled.

We include an illustration and description of the decoupling technique so that the reader may gain an intuitive understanding for how a broadband motif in spectral changes emerges naively. This method is important from a conceptual viewpoint because it illustrates that a ‘band-independent’ motif across many frequencies is specific for the dynamics of functionally diverse neuronal populations in different brain areas and across different tasks. From a practical viewpoint, filtering for a “high frequency band” (HFB) may capture a majority of the variance in broadband spectral change. Although the extracted broadband was demonstrated to vary with behavior more closely than with HFB changes (Miller et al., 2009c; Supplemental material), the decoupling process is more analytically burdensome. For practical purposes, isolating HFB changes is a reasonable and appropriate approximation of the broadband change. For example, HFB power in the 76–100 Hz range approximated the broadband component of intracranial peri-Rolandic recordings (Miller et al., 2009d; Supplemental Figure 15). However, we emphasize that the specific HFB frequency range will vary depending on sites and recording systems, and should always be selected after inspecting the power spectrum for peaked rhythms and signal-to-noise characteristics. In conclusion, when interpreting HFB changes, it is essential to make the distinction between asynchronous broadband change (reflecting a population-averaged stochastic process) and synchronized rhythmic change (reflecting some coherent property of cortical microcircuitry).

Linking behavior to asynchronous broadband activity across tasks and brain regions

As shown in Figs. 1–5, broadband spectral change reflects selective activation of local cortical circuits across many tasks and brain areas. In Figs. 1 and 2 we see that, broadband responses within the fusiform and parahippocampal gyri track the presentation of visual stimuli (such as faces and houses) on a single stimulus basis with high temporal fidelity. Furthermore, the category-specific nature of these inferotemporal loci is sufficiently robust that the stimulus category

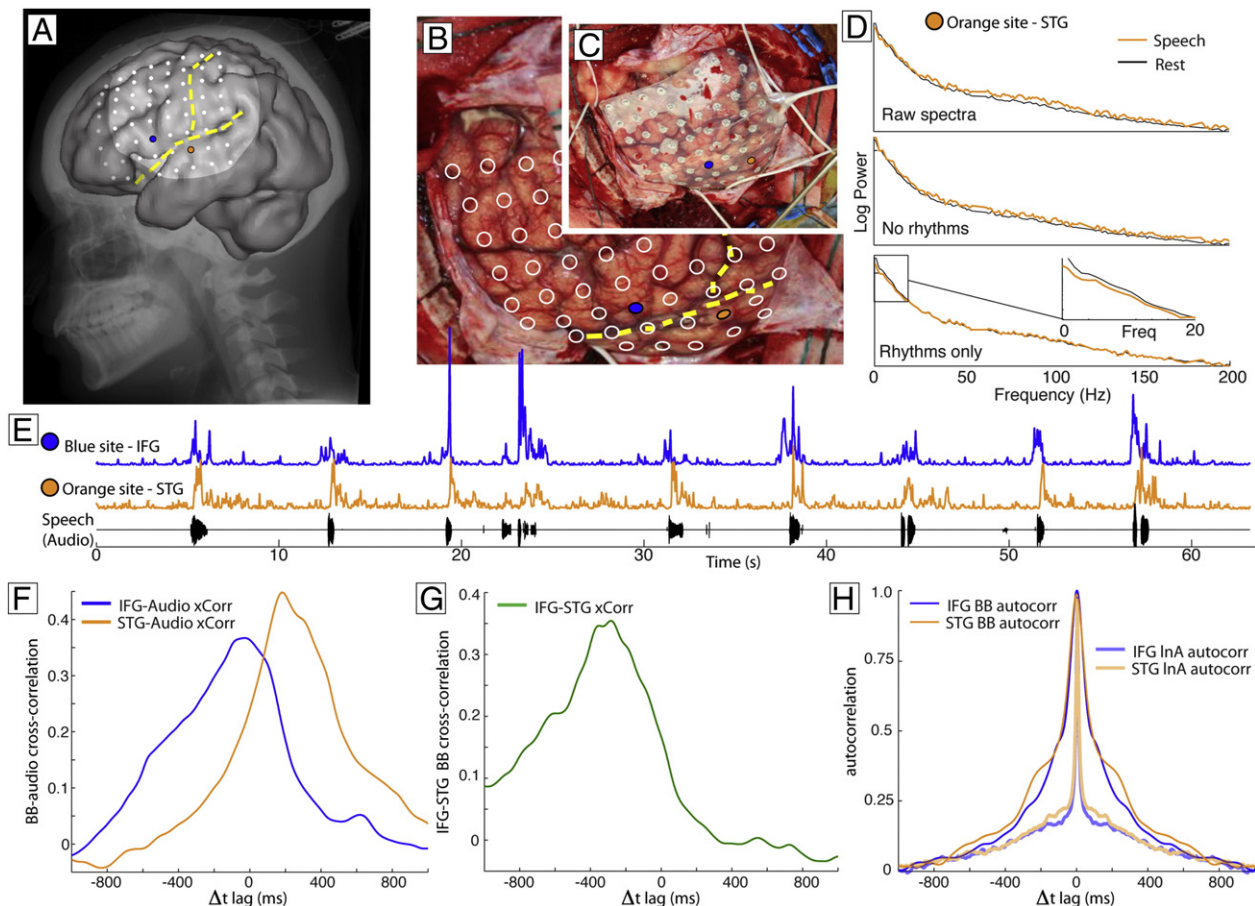


Fig. 4. Language and auditory area activity during verb generation. (A) The cortical anatomy is shown, with the non-shaded region representing the area of the exposed craniotomy, as seen in panels B and C. Yellow lines indicate the Sylvian fissure and the central sulcus. The blue dot is an electrode site in the inferior frontal gyrus (IFG) – Broca area (Brodmann area 44). The orange dot is an electrode site on the superior temporal gyrus (STG – Brodmann area 22). The white dots are the locations of the remainder of the electrode sites. (B) The locations of the electrodes are shown on the exposed brain surface, and the craniotomy with the grid in situ is also shown (C). (D) As illustrated previously, power spectral changes during different portions of the experiment (top panel) may be appreciated and naively decoupled into broadband (BB) changes (middle panel, 2nd–4th PSCs omitted) and changes in brain rhythms (bottom panel, 2nd–4th PSCs only). (E) BB timecourses from IFG (blue trace) and STG (orange trace). Simultaneous audio recording is also shown (black trace). (F) The lagged cross-correlation between BB from the two ECoG channels and the envelope of the audio trace (absolute value of the Hilbert transformed signal) shows that IFG activity (blue site) precedes speech by 30 ms, while STG activity (orange site) follows speech by 180 ms. (G) Similar comparison of cross-correlation between the IFG and STG sites shows that IFG activity leads STG activity by 280 ms. (H) The fidelity and discriminability of the time-varying BB can be estimated by examining the auto-correlations of the BB from the two sites. As would be expected, the unsmoothed InA autocorrelation falls off much faster. Panels A–C modified from Miller et al. (2011).

can be classified on individual trials by simple inspection of the response trace. Of the 300 presentations of face and house pictures for the patient in Fig. 2, all 300 were correctly identified from the spontaneous stream of data (with 22 false positives, Miller et al., 2009a). The correct face or house label could be identified with 95% accuracy, with 23 ms precision (23 ms standard deviation).

In peri-calcarine cortex (Fig. 3) the broadband time series reveals an activated dynamical state during visual search, but not while resting or staring at a blank screen. Within each cue period, there are broadband fluctuations that presumably reflect individual saccades associated with gaze shifts required for the individual elements of task performance – finding the target box, making a saccade to the arrow, returning to the target, and then identifying the appropriate color.

Broadband changes also reveal the fine temporal dynamics of speech and language in frontal and temporal areas (Fig. 4). An intracranial recording from Broca's area shows broadband activity that immediately precedes speech during verb generation. Conversely, recording from a superior temporal gyral auditory site shows broadband increase following the onset of speech production. Examining the temporal cross-correlations of broadband spectral change and the speech envelope, we see that inferior frontal gyrus (IFG) activity precedes speech by 30 ms, while superior temporal gyral (STG) activity follows speech by

180 ms. Cross-correlation between activity from the IFG and STG sites shows that IFG leads STG activity by 280 ms. The fact that $30 + 180 < 280$ demonstrates that simplified cross-correlations cannot take into account complex temporal dynamics, and serve only as a first pass for interaction times between cortical areas and behavioral parameters. The cortical activity during speech at each of these cortical sites is sustained for multiple seconds (panel 4E), this 70 ms discrepancy is likely explained by more complex within-trial dynamics. The falloff of the autocorrelation functions in Fig. 4H suggest that meaningful broadband signal change happens at well under 100 ms.

Finally, as seen in the hand region of dorsal pre-central cortex, adjacent sites are highly specific for the movements of individual fingers, with minimal overlap, and robustly capture the dynamics of each individual finger flexion (Figure 5, from a larger study, detailed in Miller et al., 2012). Close examination of the timing between motor cortex activity and movement onset is approximately 85 ms (± 39 ms SD) (Miller et al., 2009d).

These data demonstrate, across four behavioral task and multiple cortical regions, that broadband power spectral change can be used to track local cortical activity, with meaningful resolution, at approximately 20–50 ms timescale. This suggests that broadband power changes in the ECoG potential are a generic marker of activation for populations of

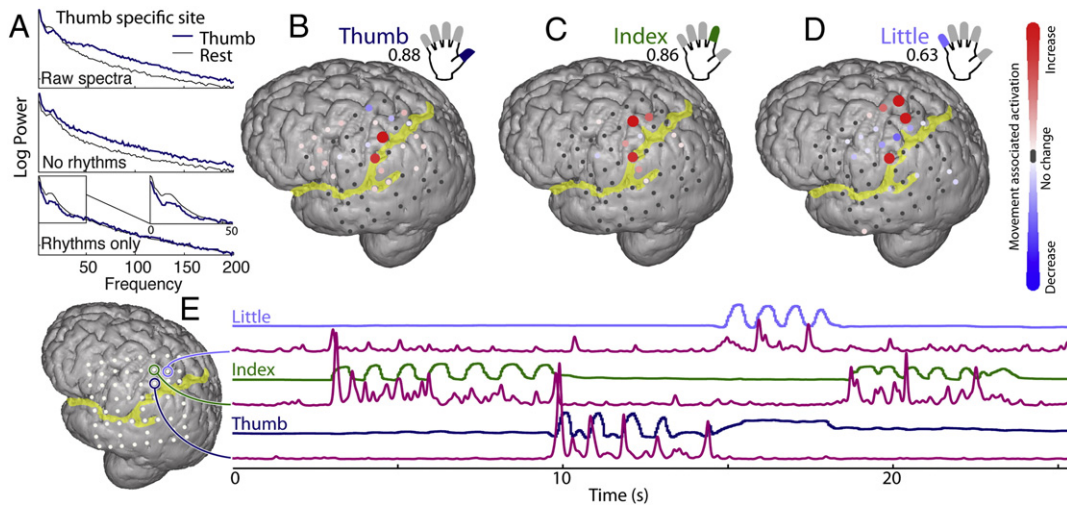


Fig. 5. ECoG broadband power resolves somatotopic representation of fingers. (A) Power spectral changes, raw and decoupled, during thumb movement and rest. (B–D) Changes in broadband at different cortical sites for movement of thumb, index, and little finger, are each compared with rest. Colors denote a signed r^2 measurement of increases and decreases in power with movement relative to rest (individually scaled with maximum noted below each label). (E) Traces of thumb (dark blue), index (green), and little finger (light blue) flexion, with corresponding timecourse of broadband spectral change (pink) for 3 pre-central cortical sites. Note that broadband timecourse from each site is remarkably specific for the flexion of only one finger. Figure partially modified from the supplement of Miller et al. (2012).

cortical neurons. Future studies will be required to demonstrate that broadband shifts are directly related to changes in population firing rate throughout the cerebral cortex. However, in light of the conserved geometry of superficial pyramidal neurons, this relationship is likely to extend well beyond the occipital (Ray and Maunsell, 2011), temporal (Manning et al., 2009), and somatomotor (Mollazadeh et al., 2009) areas where it has been directly confirmed.

The significance of the power law form of the broadband spectral change

As demonstrated previously, the ECoG cortical power spectral density has a power law form at high frequency, fitting tightly to $P(f) \sim 1/f^\chi$, with $\chi = 4$ (Miller et al., 2009b). If low frequency oscillations are avoided, either by selecting cortical sites where they are absent, or removing them using a principle component approach, a two-Lorentzian form is revealed. A single Lorentzian form indicates the presence of stochastic activity with a particular biophysical timescale (Miller et al., 2009b) while a double Lorentzian indicates the presence of two biophysical timescales within the circuit that generates the cortical surface potential. With behavior, there is a proportional increase at each frequency, e.g. $P(f,t) \sim A(t)P(f)$. This structure is important, because these two-Lorentzian forms can emerge generically from noise-like processes that have two simple correlation times (Sigeti and Horsthemke, 1987). Transmitted action potentials between neurons are approximately Poisson-distributed in time, and appear as a pure noise-like, Lorentzian, process when averaged over many neurons (Dayan and Abbott, 2001). Laminar recordings, measured alongside the surface potential, suggest that the current source densities (CSD) in different cortical lamina are due to synaptic currents and ensuing current dipoles in the dendritic tree (Mitzdorf, 1985). In contrast, propagating action potentials in axons and axon terminals do not contribute strongly to the CSD at spatial scales of ~ 50 – 300 μm , the scale where CSD varies, setting up the dipole currents which the LFPs (and by extension ECoG potentials) measure. The overall magnitude, $A(t)$, of the measured power spectrum results from the averaging of many input action potentials to the population of neurons in the vicinity of a recording electrode. The shape of the power spectrum results from the combination of two simple known neuronal processes (each with a characteristic correlation time and resulting Lorentzian form), such as: temporal integration in dendrites or soma, exponentially decaying membrane currents, low-pass RC

filtering by tissue, or local network connectivity which, when modeled, produces precisely the measured form. The integer exponent of 2 in each $1/f^2$ term indicates an underlying stochastic process, suggesting that the ECoG signal does not resolve self-organized critical processes (which would be associated with non-integer exponents).

A simple single-neuron model for the generation of the cortical potential may provide insight into the two-Lorentzian form (Bedard et al., 2006; Miller, 2010; Miller et al., 2009b). Two biophysical processes determine the spectral form: one process is the exponentially decaying post-synaptic current of fast-synapses (decay timescale 2.3 ms Sabatini and Regehr, 1996), and the other process is the temporal integration and leakage of polysynaptic input in the dendritic tree (leakage timescale 100 ms (Koch et al., 1996)). This is only one of many models that are consistent with the measured form, but it is parsimonious and empirically grounded. Our model suggests that broadband spectral shifts generically reflect changes in the input firing rate to a local neuronal population. It may be that inputs to more superficial lamina dominate, or that pyramidal neurons have a larger contribution than interneurons, or that, in some contexts, specific frequency ranges dominate because of peculiarities of local circuitry (Buzsaki and Draguhn, 2004; Cunningham et al., 2003; Rotstein et al., 2005; Sohal et al., 2009; Steriade, 2006; Tiesinga et al., 2008). Additionally, inter-spike temporal correlations will be important at smaller spatial scales. However, the internal correlations between neuronal events are likely to be lost by averaging over large spatial areas, and the spectral changes that we measure using subdural macro-electrodes generically inform us about the overall number of events taking place in the neuronal population.

This hypothesis was tested explicitly by comparing spiking rate to both band-limited and broadband power spectral changes in human microelectrode recordings (Manning et al., 2009). The best predictor of firing rate was the broadband feature of the PSD. This relation was robust, significant, and reproduced across a large number of individuals and brain sites. Several years later, this was reproduced in non-human primates, with similar results (Ray and Maunsell, 2011), and in a setting where the broadband component was concretely distinguished from the narrow-band occipital gamma rhythm.

When extracted, this broadband spectral change then likely reflects mean firing rate, spatially averaged over the neuronal population beneath. If the size of the electrode is made smaller, then fewer neurons

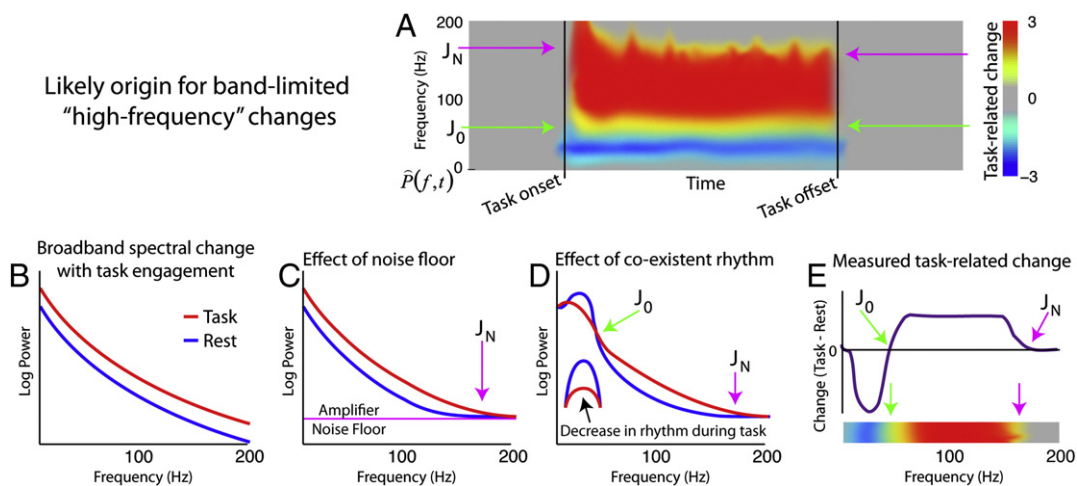


Fig. 6. Why “band-limited, high frequency” features may be seen in experimental data. (A) Stereotypical time–frequency plot, with task-associated power increases observed in a wide frequency range, beginning at $f = J_0$, extending to $f = J_N$. Panel B shows idealized time-averaged broadband power-law spectra seen during task. (C) At high frequencies, the size of the amplifier noise floor drowns out the power in the electric potential due to cortical activity (we denote the frequency where this becomes relevant as J_N). (D) At lower frequencies, the presence of co-existent synchronous rhythms obscures changes in the asynchronous broadband, and there is a crossover in power between the task and rest spectra (J_0). (E) When the difference between task and rest spectra shown in panel D are plotted, it becomes clear why spectrograms have band-limited, task-associated, high frequency power changes: They are bound on the low end by J_0 and the high end by J_N .

are being averaged over. The resolution of cortical representation is directly tied to the electrode size and density of spacing. By intrinsically averaging over many stochastic synaptic inputs, broadband spectral change reveals a robust correlate of local cortical activity on the single trial basis, with high temporal fidelity.

Examples of genuinely rhythmic high-frequency activity

We have emphasized the ubiquity and significance of broadband power shifts, and have suggested that (apparently) band-limited spectral changes in the high-gamma range may in many cases actually reflect a broadband shift. However, there are also functionally critical processes, such as hippocampal sharp wave ripples (Buzsaki et al., 1983; Okeefe and Nadel, 1979; Ylinen et al., 1995) that have genuine high-frequency oscillatory signatures. Other examples include the gamma rhythm in the occipital lobe, which can be observed at frequencies as high as 90 Hz (Fig. 1H) (Ray and Maunsell, 2011; Womelsdorf et al., 2006) and which appear to be related to the intrinsic timescales of interneuronal activity (Traub et al., 2001).

Rhythmic and non-rhythmic processes in the high-frequency range may co-occur, if, for example, synchronization of one subpopulation co-occurs with asynchronous firing in another subpopulation of neurons. Therefore the development of tools for separating synchronous and asynchronous processes is crucial. A separation method based on power spectral covariance was illustrated in Fig. 1, and other tools based on signal coarse-graining (Yamamoto and Hughson, 1993) have also been employed. It is crucial to test the performance of these methods in cases where the asynchronous broadband component deviates from its expected form (e.g. if it does not precisely follow a power law).

The implication of broadband power shifts will also vary as a function of the spatial geometry of the electrode. When recording with penetrating electrodes, sharp edges in the field potential (e.g. due to individual action potentials) may introduce wide-band energy into the spectrum. Depending on their spectral profile, these edge-induced bursts may be difficult to distinguish from the asynchronous broadband shifts that reflect population activity (Scheffer-Teixeira et al., 2013). However, when recording with surface electrodes as in the examples shown here (Figs. 1–5), the possibility of spike contamination is dramatically reduced, as the signal is aggregated across hundreds of

thousands of neurons, and is dominated by currents in proximal apical dendrites (Mitzdorf, 1985).

It has been shown in a variety of behavioral settings that the amplitude of this broadband spectral change is modulated by the phase of low-frequency rhythms — a specific instance of “phase amplitude coupling” (PAC) (Miller et al., 2010a, 2012). As such, this modulation likely reflects a macroscopic index of well-known spike-field interaction (Buzsaki and Draguhn, 2004; Mollazadeh et al., 2009; Okun et al., 2009; Traub et al., 2001), and provides evidence that averaged spiking activity in widespread cortical circuits, reflected by this broadband, can be entrained with the phase of underlying rhythms.

How broadband changes have gained attention

More than 40 years ago, Brindley and Craggs observed that the power in the 80–250 Hz frequency range of the electric potential time series from motor cortex was dynamically and somatotopically increased in adjacent motor areas during movement (Brindley and Craggs, 1972). This finding lay obscure and dormant for more than 25 years, until Nathan Crone and colleagues independently re-discovered that the high-frequency (~75–100 Hz) portion of the human ECoG potential was highly specific for the dynamics of different motor functions (Crone et al., 1998a). Although both groups proposed that this high-frequency power was a correlate of specific cortical activity, the nature and relationship to underlying physiology was unknown. Crone's group has more clearly postulated that very local populations of neurons were selectively synchronized with one another at multiple fast timescales, in a spatially-overlapping tableau of microdomains, with the spatial scale of coherence falling off with increasing frequency, producing a $1/f$ shape in the PSD (Crone et al., 2011).

However, the lack of a clear upper bound in the difference between active and inactive spectra in motor cortex led to the hypothesis (Miller et al., 2007), and later demonstration (Miller et al., 2009b,d) that some behavior-related changes in the cortical spectrum do not correspond to synchronized populations of neurons, but are rather asynchronous and broadband in nature. As illustrated in Fig. 6, these broadband spectral changes are often mistaken for band-limited changes. The broadband is obscured at lower frequencies by intersection with classic area-specific oscillations such as the α and β rhythms in visual and motor cortices (J_0 in Figure 6 Miller et al., 2008). At high frequencies, the power in the cortically generated electrical potential falls off rapidly as

$P \sim 1/f^4$. Because all amplifiers have an intrinsic noise floor, the brain signal becomes lost in the instrument noise at an upper bound (J_N in Figure 8 Miller et al., 2009b). When researchers observing only the $1/f$ correlate of local brain activity between J_0 and J_N , they may mistakenly refer to it as a band-limited “high-gamma power”.

Focusing on changes in particular frequency bands is practical and computationally expedient, but we suggest caution in interpreting spectral changes (especially those above 60 Hz) as changes in cortical oscillations. The oscillatory interpretation is most plausible when the high-frequency spectral shift is large and tightly band-limited, conditions which are rarely met in extra-cortical recordings. We propose that the first interpretation of a high-frequency change in the spectrum should be in terms of broadband spectral shifts. These broadband shifts, which can be characterized using the methods described above, are the signature of changes in asynchronous firing rate within local population, and they are therefore both expected and observed (Figs. 1–5) in wide-ranging experimental settings. An advantage of this approach is that once a broadband shift is isolated, concurrent changes in the θ , α , β , and γ rhythms can be more readily and precisely observed.

Conclusion

Having surveyed a variety of experimental settings, we reiterate the claim that the popular “high-gamma” range of the cortical electrical potential is widely misunderstood. In most cases, changes in spectral power in this band do not reflect changes in synchronous, rhythmic, action potentials in underlying cortex. We propose that, especially for recordings outside the cortex, spectral changes in this band most often reflect changes in asynchronous activity, which has a power-law spectral signature. This power-law process, observable via broadband spectral changes during behavior, likely correlates with the mean neuronal population activity. The data indicate that this broadband spectral signature is robustly linked to function across cortical areas and behaviors, thereby generically tracking the activation of cortical populations with high temporal fidelity.

References

- Aoki, F., Fetz, E.E., Shupe, L., Lettich, E., Ojemann, G.A., 1999. Increased gamma-range activity in human sensorimotor cortex during performance of visuomotor tasks. *Clin. Neurophysiol.* 110, 524–537.
- Ashburner, J., Friston, K.J., 2005. Unified segmentation. *NeuroImage* 26, 839–851.
- Bates, J.A., 1951. Electrical activity of the cortex accompanying movement. *J. Physiol.* 113, 240–257.
- Bedard, C., Kroeger, H., Destexhe, A., 2006. Does the $1/f$ frequency scaling of brain signals reflect self-organized critical states? *Phys. Rev. Lett.* 97, 118102.
- Brindley, G.S., Craggs, M.D., 1972. The electrical activity in the motor cortex that accompanies voluntary movement. *Proc. Physiol. Soc.* 28P.
- Bullock, T.H., McClune, M.C., Enright, J.T., 2003. Are the electroencephalograms mainly rhythmic? Assessment of periodicity in wide-band time series. *Neuroscience* 121, 233–252.
- Buzsaki, G., Draguhn, A., 2004. Neuronal oscillations in cortical networks. *Science* 304, 1926–1929.
- Buzsaki, G., Leung, L.W., Vanderwolf, C.H., 1983. Cellular bases of hippocampal EEG in the behaving rat. *Brain Res.* 287, 139–171.
- Canolty, R.T., Soltani, M., Dalal, S.S., Edwards, E., Dronkers, N.F., Nagarajan, S.S., Kirsch, H.E., Barbaro, N.M., Knight, R.T., 2007. Spatiotemporal dynamics of word processing in the human brain. *Front. Neurosci.* 1, 185–196.
- Crone, N.E., Miglioretti, D.L., Gordon, B., Lesser, R.P., 1998a. Functional mapping of human sensorimotor cortex with electrocorticographic spectral analysis. II. Event-related synchronization in the gamma band. *Brain* 121 (Pt 12), 2301–2315.
- Crone, N.E., Miglioretti, D.L., Gordon, B., Sieracki, J.M., Wilson, M.T., Uematsu, S., Lesser, R.P., 1998b. Functional mapping of human sensorimotor cortex with electrocorticographic spectral analysis. I. Alpha and beta event-related desynchronization. *Brain* 121 (Pt 12), 2271–2299.
- Crone, N.E., Hao, L., Hart Jr., J., Boatman, D., Lesser, R.P., Irizarry, R., Gordon, B., 2001. Electrocorticographic gamma activity during word production in spoken and sign language. *Neurology* 57, 2045.
- Crone, N.E., Korzeniewska, A., Franaszczuk, P.J., 2011. Cortical gamma responses: searching high and low. *Int. J. Psychophysiol.* 79, 9–15.
- Cunningham, M.O., Davies, C.H., Buhl, E.H., Kopell, N., Whittington, M.A., 2003. Gamma oscillations induced by kainate receptor activation in the entorhinal cortex in vitro. *J. Neurosci.* 23, 9761–9769.
- Dale, A.M., Fischl, B., Sereno, M.I., 1999. Cortical surface-based analysis. I. Segmentation and surface reconstruction. *NeuroImage* 9, 179–194.
- Dayan, P., Abbott, L.F., 2001. *Theoretical Neuroscience: Computational and Mathematical Modeling of Neural Systems*. Massachusetts Institute of Technology Press, Cambridge, Mass.
- Edwards, E., Nagarajan, S.S., Dalal, S.S., Canolty, R.T., Kirsch, H.E., Barbaro, N.M., Knight, R.T., 2010. Spatiotemporal imaging of cortical activation during verb generation and picture naming. *NeuroImage* 50, 291–301.
- Foster, B.L., Dastjerdi, M., Parvizi, J., 2012. Neural populations in human posteromedial cortex display opposing responses during memory and numerical processing. *Proc. Natl. Acad. Sci. U. S. A.* 109, 15514–15519.
- Friston, K.J., Holmes, A.P., Worsley, K.J., Poline, J.B., Frith, C.D., Frackowiak, R.S.J., 1995. Statistical parametric maps in functional imaging: a general linear approach. *Hum. Brain Mapp.* 2, 189–210.
- Gunduz, A., Brunner, P., Daich, A., Leuthardt, E.C., Ritaccio, A.L., Pesaran, B., Schalk, G., 2011. Neural correlates of visual–spatial attention in electrocorticographic signals in humans. *Front. Hum. Neurosci.* 5, 89.
- Hermes, D., Miller, K.J., Noordmans, H.J., Vansteensel, M.J., Ramsey, N.F., 2010. Automated electrocorticographic electrode localization on individually rendered brain surfaces. *J. Neurosci. Methods* 185, 293–298.
- Hermes, D., Miller, K.J., Vansteensel, M.J., Aarnoutse, E.J., Leijten, F.S., Ramsey, N.F., 2012. Neurophysiologic correlates of fMRI in human motor cortex. *Hum. Brain Mapp.* 33, 1689–1699.
- Jacobs, J., Kahana, M.J., 2009. Neural representations of individual stimuli in humans revealed by gamma-band electrocorticographic activity. *J. Neurosci.* 29, 10203–10214.
- Jasper, H.H., 1941. Electrical activity of the brain. *Annu. Rev. Physiol.* 3, 377–398.
- Jasper, H.H., Andrews, H.L., 1938. Brain potentials and voluntary muscle activity in man. *J. Neurophysiol.* 1, 87.
- Jasper, H., Penfield, W., 1949. Electrocorticograms in man: effect of voluntary movement upon the electrical activity of the precentral gyrus. *Eur. Arch. Psychiatry Clin. Neurosci.* 183, 163–174.
- Koch, C., Rapp, M., Segev, I., 1996. A brief history of time (constants). *Cereb. Cortex* 6, 93–101.
- Manning, J.R., Jacobs, J., Fried, I., Kahana, M.J., 2009. Broadband shifts in local field potential power spectra are correlated with single-neuron spiking in humans. *J. Neurosci.* 29, 13613.
- Mesgarani, N., Chang, E.F., 2012. Selective cortical representation of attended speaker in multi-talker speech perception. *Nature* 485, 233–236.
- Miller, K.J., 2010. Broadband spectral change: evidence for a macroscale correlate of population firing rate? *J. Neurosci.* 30, 6477–6479.
- Miller, F.R., Stavaky, G.W., Woonton, G.A., 1940. Effects of eserine, acetylcholine and atropine on the electrocorticogram. *J. Neurophysiol.* 3, 131.
- Miller, K.J., Leuthardt, E.C., Schalk, G., Rao, R.P., Anderson, N.R., Moran, D.W., Miller, J.W., Ojemann, J.G., 2007. Spectral changes in cortical surface potentials during motor movement. *J. Neurosci.* 27, 2424–2432.
- Miller, K.J., Shenoy, P., den Nijs, M., Sorensen, L.B., Rao, R.N., Ojemann, J.G., 2008. Beyond the gamma band: the role of high-frequency features in movement classification. *IEEE Trans. Biomed. Eng.* 55, 1634.
- Miller, K.J., Hermes, D., Schalk, G., Ramsey, N.F., Jagadeesh, B., den Nijs, M., Ojemann, J.G., Rao, R.P., 2009a. Detection of spontaneous class-specific visual stimuli with high temporal accuracy in human electrocorticography. *IEEE Eng. Med. Biol. Soc.* 2009, 6465–6468.
- Miller, K.J., Sorensen, L.B., Ojemann, J.G., den Nijs, M., 2009b. Power-law scaling in the brain surface electric potential. *PLoS Comput. Biol.* 5, e1000609.
- Miller, K.J., Weaver, K.E., Ojemann, J.G., 2009c. Direct electrophysiological measurement of human default network areas. *Proc. Natl. Acad. Sci.* 106, 12174.
- Miller, K.J., Zanos, S., Fetz, E.E., den Nijs, M., Ojemann, J.G., 2009d. Decoupling the cortical power spectrum reveals real-time representation of individual finger movements in humans. *J. Neurosci.* 29, 3132.
- Miller, K.J., Hermes, D., Honey, C.J., Sharma, M., Rao, R.P., den Nijs, M., Fetz, E.E., Sejnowski, T.J., Hebb, A.O., Ojemann, J.G., Makeig, S., Leuthardt, E.C., 2010a. Dynamic modulation of local population activity by rhythm phase in human occipital cortex during a visual search task. *Front. Hum. Neurosci.* 4, 197.
- Miller, K.J., Schalk, G., Fetz, E.E., den Nijs, M., Ojemann, J.G., Rao, R.P.N., 2010b. Cortical activity during motor execution, motor imagery, and imagery-based online feedback. *Proc. Natl. Acad. Sci.* 108, 4430–4435.
- Miller, K.J., Abel, T.J., Hebb, A.O., Ojemann, J.G., 2011. Rapid online language mapping with electrocorticography. *J. Neurosurg. Pediatr.* 7.
- Miller, K.J., Hermes, D., Honey, C.J., Hebb, A.O., Ramsey, N.F., Knight, R.T., Ojemann, J.G., Fetz, E.E., 2012. Human motor cortical activity is selectively phase-entrained on underlying rhythms. *PLoS Comput. Biol.* 8, e1002655.
- Mitzdorf, U., 1985. Current source-density method and application in cat cerebral cortex – investigation of evoked-potentials and EEG phenomena. *Physiol. Rev.* 65, 37–100.
- Mollazadeh, M., Aggarwal, V., Thakor, N.V., Law, A.J., Davidson, A., Schieber, M.H., 2009. Coherency between spike and LFP activity in M1 during hand movements. *Neural Engineering*, 2009. NER '09. 4th International IEEE/EMBS Conference on, pp. 506–509.
- Okeefe, J., Nadel, L., 1979. The hippocampus as a cognitive map. *Behav. Brain Sci.* 2, 520–528.
- Okun, M., Naim, A., Lampl, I., 2009. Intracellular Recordings in Awake Rodent Unveil the Relation Between Local Field Potential and Neuronal Firing. *Society for Neuroscience*, Chicago, IL, USA.
- Pei, X., Barbour, D.L., Leuthardt, E.C., Schalk, G., 2011. Decoding vowels and consonants in spoken and imagined words using electrocorticographic signals in humans. *J. Neural Eng.* 8, 046028.
- Penfield, W., 1954. Mechanisms of voluntary movement. *Brain* 77, 1.
- Pfurtscheller, G., 1999. Event-Related Desynchronization (ERD) and Event Related Synchronization (ERS). *Williams and Wilkins*, Baltimore.

- Ray, S., Maunsell, J.H., 2011. Different origins of gamma rhythm and high-gamma activity in macaque visual cortex. *PLoS Biol.* 9, e1000610.
- Rotstein, H.G., Pervouchine, D.D., Acker, C.D., Gillies, M.J., White, J.A., Buhl, E.H., Whittington, M.A., Kopell, N., 2005. Slow and fast inhibition and an H-current interact to create a theta rhythm in a model of CA1 interneuron network. *J. Neurophysiol.* 94, 1509–1518.
- Sabatini, B.L., Regehr, W.G., 1996. Timing of neurotransmission at fast synapses in the mammalian brain. *Nature* 384, 170–172.
- Schalk, G., McFarland, D.J., Hinterberger, T., Birbaumer, N., Wolpaw, J.R., 2004. BCI2000: a general-purpose brain–computer interface (BCI) system. *IEEE Trans. Biomed. Eng.* 51, 1034–1043.
- Scheffer-Teixeira, R., Belchior, H., Leao, R.N., Ribeiro, S., Tort, A.B., 2013. On high-frequency field oscillations (>100 Hz) and the spectral leakage of spiking activity. *J. Neurosci.* 33, 1535–1539.
- Sederberg, P.B., Kahana, M.J., Howard, M.W., Donner, E.J., Madsen, J.R., 2003. Theta and gamma oscillations during encoding predict subsequent recall. *J. Neurosci.* 23, 10809–10814.
- Sigeti, D., Horsthemke, W., 1987. High-frequency power spectra for systems subject to noise. *Phys. Rev. A* 35, 2276–2282.
- Sohal, V.S., Zhang, F., Yizhar, O., Deisseroth, K., 2009. Parvalbumin neurons and gamma rhythms enhance cortical circuit performance. *Nature* 459, 698–702.
- Steriade, M., 2006. Grouping of brain rhythms in corticothalamic systems. *Neuroscience* 137, 1087–1106.
- Swann, N., Tandon, N., Canolty, R., Ellmore, T.M., McEvoy, L.K., Dreyer, S., DiSano, M., Aron, A.R., 2009. Intracranial EEG reveals a time- and frequency-specific role for the right inferior frontal gyrus and primary motor cortex in stopping initiated responses. *J. Neurosci.* 29, 12675–12685.
- Tiesinga, P., Fellous, J.M., Sejnowski, T.J., 2008. Regulation of spike timing in visual cortical circuits. *Nat. Rev. Neurosci.* 9, 97–107.
- Traub, R.D., Kopell, N., Bibbig, A., Buhl, E.H., LeBeau, F.E.N., Whittington, M.A., 2001. Gap junctions between interneuron dendrites can enhance synchrony of gamma oscillations in distributed networks. *J. Neurosci.* 21, 9478–9486.
- Vidal, J.R., Ossandon, T., Jerbi, K., Dalal, S.S., Minotti, L., Ryvlin, P., Kahane, P., Lachaux, J.P., 2010. Category-specific visual responses: an intracranial study comparing gamma, beta, alpha, and ERP response selectivity. *Front. Hum. Neurosci.* 4, 195.
- Whittingstall, K., Logothetis, N.K., 2009. Frequency-band coupling in surface EEG reflects spiking activity in monkey visual cortex. *Neuron* 64, 281–289.
- Womelsdorf, T., Fries, P., Mitra, P.P., Desimone, R., 2006. Gamma-band synchronization in visual cortex predicts speed of change detection. *Nature* 439, 733–736.
- Yamamoto, Y., Hughson, R.L., 1993. Extracting fractal components from time-series. *Physica D* 68, 250–264.
- Ylinen, A., Bragin, A., Nadasdy, Z., Jando, G., Szabo, I., Sik, A., Buzsaki, G., 1995. Sharp wave-associated high-frequency oscillation (200 Hz) in the intact hippocampus: network and intracellular mechanisms. *J. Neurosci.* 15, 30–46.