

All spectral frequencies of neural activity reveal semantic representation in the human anterior ventral temporal cortex

Saskia L. Frisby¹, Ajay D. Halai¹, Christopher R. Cox², Alex Clarke³, Akihiro Shimotake^{4,5}, Takayuki Kikuchi⁶, Takeharu Kuneida^{6,7}, Yoshiki Arakawa⁶, Ryosuke Takahashi⁴, Akio Ikeda⁸, Riki Matsumoto^{4,9}, Timothy T. Rogers¹⁰, & Matthew A. Lambon Ralph¹

¹ MRC Cognition and Brain Sciences Unit, University of Cambridge, Cambridge, UK

² Department of Psychology, Louisiana State University, Baton Rouge, USA

³ Department of Psychology, University of Warwick, Warwick, UK

⁴ Department of Neurology, Kyoto University Graduate School of Medicine, Kyoto, Japan

⁵ Clinical Research Center, National Hospital Organization Utano Hospital, Kyoto, Japan

⁶ Department of Neurosurgery, Kyoto University Graduate School of Medicine, Kyoto, Japan

⁷ Department of Neurosurgery, Ehime University Graduate School of Medicine, Ehime, Japan

⁸ Department of Epilepsy, Movement Disorders and Physiology; Kyoto University Graduate School of Medicine; Kyoto; Japan

⁹ Division of Neurology, Kobe University Graduate School of Medicine, Kusunoki-cho, Kobe, Japan

¹⁰ Department of Psychology, University of Wisconsin–Madison, Madison, USA

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Abstract:

Intracranial electrophysiology offers a unique insight into the nature of information representation in the brain – it can be used to disentangle information encoded in local neuronal activity (gamma and high gamma frequencies) from information encoded via long-range interactions (lower frequencies). We used regularised logistic regression to decode animacy from time-frequency power and phase extracted from electrocorticography (ECoG) grid electrode data recorded on the surface of human vATL. Power in gamma (30 – 60 Hz) and high gamma (60 – 200 Hz) produced reliable decoding, indicating that semantic information is indeed expressed by local populations in vATL. However, power from a wide range of frequencies (4 – 200 Hz) produced significantly higher decoding accuracy and also exhibited the same rapidly-changing dynamic code previously observed when decoding voltage. These findings support the theory that semantic information is encoded by a local vATL “hub” that interacts with distributed cortical “spokes”.

1. Introduction

Semantic cognition is our ability to understand and respond in a meaningful way to objects that we see, language that we hear, events that we experience, and any other stimuli in our environment. The ventral anterior temporal lobes (vATLs) play a critical role in such abilities (Lambon Ralph et al., 2017; Patterson et al., 2007; Rogers et al., 2004), but it remains unclear *how* neural activity in these regions encodes semantic information. Electrocorticography (ECoG) grid electrode data are a rare and informative resource, sensitive to activity at a wide range of frequencies including very high-frequency activity thought to indicate local neuronal activity (Başar-Eroglu et al., 1996; Bressler, 1990; Merker, 2013; Nir et al., 2007). Although prior work has shown that surface voltages express information about stimulus animacy (is it an animal or not?; Y. Chen et al., 2016; Rogers et al., 2021), voltages are a conflation of high- and lower-frequency activity; therefore, apparent representation of semantic information by the vATL may actually reflect incoming signals from more remote areas. We address these questions by applying regularised logistic regression — a multivariate decoding procedure that makes minimal assumptions about the nature of neural representations (Cox & Rogers, 2021; Frisby et al., 2023) — to time-frequency power and phase extracted from a large sample of ECoG data ($n = 18$).

Extensive prior work suggests that the vATLs serve as a representational “hub”, interacting with multiple “spokes” that each encode modality-specific information about the environment. By virtue of these interactions, and over the course of learning and development, the vATLs come to express transmodal conceptual similarity structure and enable generalisation of knowledge across items and contexts (Lambon Ralph et al., 2010, 2017; Patterson et al., 2007; Rogers et al., 2004). The “hub-and-spoke” model is supported by convergent evidence from neuropsychology (Hodges & Patterson, 2007), positron emission tomography (PET; Devlin et al., 2000), distortion-corrected functional magnetic resonance imaging (fMRI; Binney et al., 2010; Halai et al., 2014; Visser et al., 2010), magnetoencephalography (MEG; Mollo et al., 2017), neuro-computational simulations (L. Chen et al., 2017; Rogers et al., 2004, 2021), transcranial magnetic stimulation (TMS; Pobric et al., 2007, 2010a, 2010b), invasive cortical stimulation mapping (Lüders et al., 1991; Matoba et al., 2024), and ECoG (Halgren et al., 2006; Nobre & McCarthy, 1995; Sato et al., 2021; Shimotake et al., 2015).

Multivariate decoding approaches have only recently been applied to ECoG data in an attempt to characterise, not simply where and when task-related changes in neuronal activity occur, but what information those changes represent. Most of these analyses have focused on voltage (Y. Chen et al., 2016; Cox et al., 2024; Liu et al., 2009; Nagata et al., 2022; Rogers et al., 2021, cf. Clarke, 2020; Rupp et al., 2017; Wang et al., 2011). However, voltage reflects a blend of activity at different frequencies and, while gamma (> 30 Hz) and high gamma ($> \sim 60$ Hz) activity is thought to reflect local neuronal activity (Başar-Eroglu et al., 1996; Bressler, 1990; Merker, 2013; Nir et al., 2007), activity at lower frequencies may reflect transmission of information over varying cortical distances (Canolty & Knight, 2010; Lachaux et al., 2012), which may be important for assembling the spatially-distributed representations posited by the hub-and-spoke model.

The first aim of this paper was to clarify the role of time-frequency power and phase in semantic representation in the vATL and to relate this to voltage findings (Y. Chen et al., 2016; Cox et al., 2024; Liu et al., 2009; Nagata et al., 2022; Rogers et al., 2021) and to other sources of evidence (Lambon Ralph et al., 2017). The dataset consisted of ECoG data collected from a large cohort of patients ($n = 18$) while they named line drawings of animate and inanimate objects. ECoG offers excellent spatial and temporal resolution and the use of grid electrodes on the cortical surface offers much greater coverage than can be obtained with cortical depth electrodes — important for detecting semantic representations that may be distributed across space (Merker, 2013; Rogers et al., 2021). Crucially for this study, intracranial methods are necessary for revealing high gamma activity that is blocked by the skull and scalp when using noninvasive methods such as EEG or MEG (Lachaux et al., 2012; Parvizi & Kastner, 2018).

We used regularised logistic regression to decode information from time-frequency power and phase — a method that, unlike decoding methods that have previously been applied to ECoG time-frequency data (Clarke, 2020; Rupp et al., 2017; Wang et al., 2011) makes very few assumptions about the properties of the underlying code (Frisby et al., 2023). Regularised regression can detect representations whether individual populations or frequencies encode pieces of information separately, or whether the relationship between information and neural activity can be understood only by considering multiple populations and/or frequencies simultaneously (Frisby et al., 2023). Since it relies on parameter fitting

rather than simple correlations, it is resistant to false positives and false negatives associated with purely correlational approaches (Cox et al., 2024).

We tested whether animacy (a) can be decoded from time-frequency power and/or phase; (b) if so, from which frequency ranges and at what timepoints; and (c) how these results relate to decoding of voltage. These analyses therefore allowed us to determine whether time-frequency- and voltage-based decoding offer similar or different conclusions about the timecourse of semantic representation in the vATL. Decoding accuracy in gamma and high gamma ranges (independently of other ranges) allowed us to determine whether semantic information is encoded by local neural activity within the vATL.

The second aim of this paper was to assess whether the dynamic and nonlinear changes to semantic representation observed in voltage decoding (Rogers et al., 2021) also arise within or across frequency ranges. Dynamic change in representation can be measured via the *temporal generalisation* of a decoding model (Carlson et al., 2013; Cichy et al., 2014; Contini et al., 2017; King & Dehaene, 2014; Rogers et al., 2021). The model is fitted to neural data at a given timepoint, then is assessed at other timepoints. If a classifier shows above-chance hold-out accuracy when fitted to timepoint A, and also shows above-chance accuracy when used to decode at timepoint B, we can conclude that information is encoded similarly at both timepoints. Conversely, if a classifier trained and succeeding in timepoint A then fails when tested on timepoint B, and if the reverse pattern also holds (a model trained on B succeeds on B but fails on A), this suggests that the target information is encoded by different neural patterns at different timepoints. That is, such a pattern suggests dynamic change in neural representation of the target information.

When applied to ECoG voltage data, this technique revealed four interesting characteristics of the semantic code in the vATL (Rogers et al., 2021):

1. Classifiers fitted and tested at overlapping 50 ms windows showed reliable decoding from about 100ms post stimulus onset through the remainder of the processing window (1650ms), suggesting that animacy is *constantly decodable* from the moment stimulus-driven activity reaches the vATL until the response.

2. Classifiers trained at a given window generalised to temporal neighbours (both preceding and succeeding), but not to temporally remote windows; that is, voltage-based neural decoding showed *local temporal generalisation*, suggesting a dynamically-changing neural code for animacy.

134 3. The width of the temporal generalisation window for a given classifier was quite
135 narrow early in stimulus processing but widened over time, indicating that dynamic change
136 to the voltage-based code is initially quite rapid but slows with further processing.

137 4. The direction of the neural code — whether an increase in voltage at a particular
138 electrode signified increased or decreased probability that the stimulus is animate —
139 changed over the course of processing, so that voltage deflections might signify different
140 things at different timepoints at the same electrode.

141 Together these properties suggest that, while information about animacy is
142 expressed in voltage throughout stimulus processing, the precise way that this information
143 is encoded changes rapidly and nonlinearly, with change slowing as the system settles into a
144 steady state. Moreover, these patterns mirrored those observed in a recurrent deep neural
145 network model of semantic processing, suggesting that such models may offer a useful way
146 of thinking about representational dynamics in real neural systems (Rogers et al., 2021).

147 The second goal of this paper was to assess whether these properties (constant
148 decodability, local temporal generalisation, widening generalisation window, and change in
149 code direction) are observed when decoding time-frequency power or phase information,
150 across ranges or within each range considered independently. Answers to these questions
151 could clarify whether these dynamic characteristics of representation arise from rapid local
152 activity (i.e., driven by gamma/high gamma) and/or from more remote input (driven by
153 lower frequencies).

156 1. Results

158 1.1. Relative decoding accuracy using spectral frequency power or phase vs. voltage

160 The ECoG data were collected from 6 - 52 subdural grid electrodes implanted over
161 the ventral anterior temporal lobe (vATL) in 18 patients. The patients named line drawings
162 of animals and inanimate objects that were used in prior work and matched for age of
163 acquisition, concreteness, word frequency, and familiarity (Dilkina & Lambon Ralph, 2012).
164 Time-frequency power and phase were extracted using complex Morlet wavelet convolution
165 (see Methods). Classifiers were fitted using LASSO regularisation (Tibshirani, 1996) and

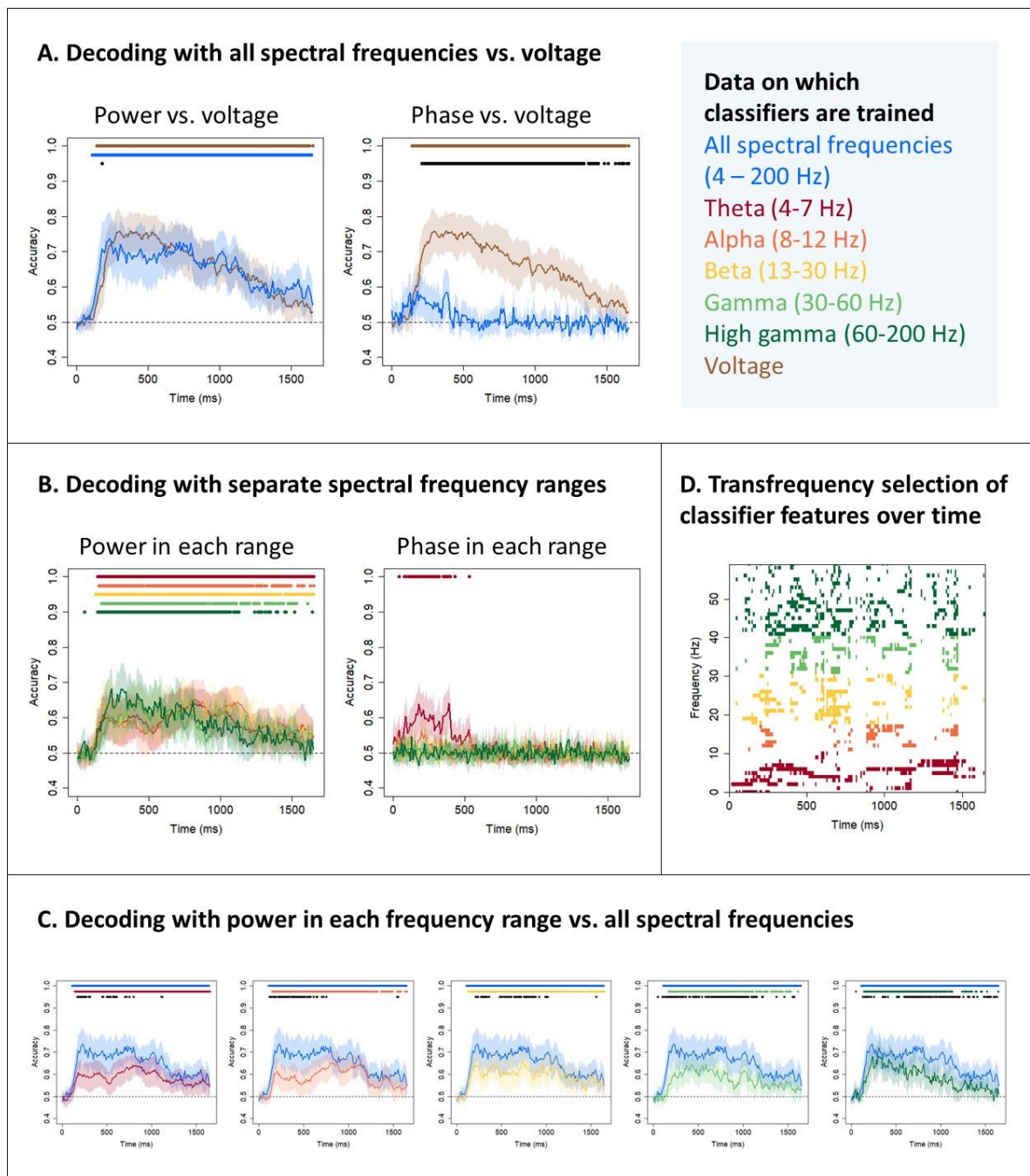
assessed using ten-fold cross-validation. The classifiers were trained either on frequency features (vectors of power or phase values extracted from multiple frequencies from multiple electrodes at a single timepoint) or on voltage features (vectors of voltage values extracted from multiple electrodes from a 50 ms window centred on the timepoint of interest). Separate classifiers were trained for timepoints between 0 and 1650 ms in 10 ms time steps (0 ms, 10 ms, 20 ms, ...). Each classifier was tested on every possible timepoint.

We first tested whether it was possible to decode the animacy of the stimuli using time-frequency power and/or phase and whether decoding performance was comparable to voltage (Rogers et al., 2021). We compared classifiers trained on frequency features of power or phase at 60 frequencies, logarithmically spaced between 4 and 200 Hz, to classifiers trained on voltage features. Figure 1A shows the results. Time-frequency power showed an almost identical decoding profile to voltage – hold-out accuracy rose to around 0.7 at 200 ms post stimulus onset and remained significantly above chance (0.5) throughout the time window. At no point was it possible to decode animacy from classifiers trained on phase.

Next, to disentangle the role of local neural activity versus distal input, we trained classifiers on frequency features composed only of power or phase values from one frequency range - theta (4 – 7 Hz), alpha (8 – 12 Hz), beta (13 – 30 Hz), gamma (30 – 60 Hz) and high gamma (60 – 200 Hz). Figure 1B shows the decoding profile in each range. For power, the patterns were almost indistinguishable, showing the same rise to about 0.6 and remaining above chance for most of the time window of interest. On the assumption that gamma and high gamma frequencies represent local neural activity, these results confirmed that local neural activity in the vATL expresses semantic information. However, we found that power in every range was sufficient, and no range was necessary, for above-chance decoding. For phase, decoding was rarely above chance, with the exception of weak decoding using theta data in the window 0 - 500 ms. The remaining analyses therefore focused on characterising semantic information encoded in time-frequency power.

Figure 1C shows hold-out accuracy for classifiers trained on power within a single range compared to accuracy for classifiers trained on power at all 60 frequencies. For each range there was a time window during which decoding on all frequencies reached its peak and decoding on a single range performed significantly less well. This indicated that no individual range contains sufficient information to enable decoding accuracy comparable to

198 accuracy based on all frequencies. To investigate which combination of frequencies enabled
199 such successful decoding, we inspected the coefficients on each frequency. Figure 1D shows
200 these averaged across electrodes for one example patient (similar plots for all patients are
201 shown in Supplementary Materials S1, Figure S1; plots showing coefficients on individual
202 electrodes for the one sample patient are shown in Supplementary Materials S1, Figure S2).
203 Throughout the time window, frequencies across the whole range were selected by the
204 classifier and thus contributed to decoding performance. This is evidence that the vATL
205 represents semantic information via a *transfrequency* code, by which we mean that
206 frequencies within the theta, alpha, beta, gamma and high gamma ranges all contribute to
207 information representation.



208

209 Figure 1: Semantic decoding with time-frequency power and phase. (A) Mean and 95%
 210 confidence interval of the hold-out accuracy for classifiers trained on power or phase
 211 frequency features for all frequencies between 4 and 200 Hz (blue) or on voltage features
 212 (brown). Coloured dots indicate a significant difference between classifier accuracy and
 213 chance (0.5, one-sample t-test with probabilities adjusted to control the false-discovery rate
 214 at $\alpha = 0.05$). Black dots indicate a significant difference between accuracies at a given
 215 timepoint (paired t-tests with probabilities adjusted to control the false-discovery rate at $\alpha =$
 216 0.05). (B) Mean and 95% confidence interval of the hold-out accuracy for classifiers trained
 217 on frequency features composed only of power or phase values from a given range - theta
 218 (4 – 7 Hz, red), alpha (8 – 12 Hz, orange), beta (13 – 30 Hz, yellow), gamma (30 – 60 Hz, light
 219 green) and high gamma (60 – 200 Hz, dark green). Coloured dots indicate a significant

difference between classifier accuracy and chance (0.5, one-sample t-test with probabilities adjusted to control the false-discovery rate at $\alpha = 0.05$). (C) Mean and 95% confidence interval of the hold-out accuracy for classifiers trained on frequency features composed only of power or phase values from a given range compared to classifiers trained on frequency features including all frequencies between 4 and 200 Hz. Coloured dots indicate a significant difference between classifier accuracy and chance; black dots indicate a significant difference between accuracies. (D) Feature selection of each frequency at each timepoint (averaged across electrodes) for classifiers trained on frequency feature vectors including all frequencies between 4 and 200 Hz for a single patient. Features with a nonzero coefficient are shown in colour.

1.2. Does the time-frequency semantic code change dynamically?

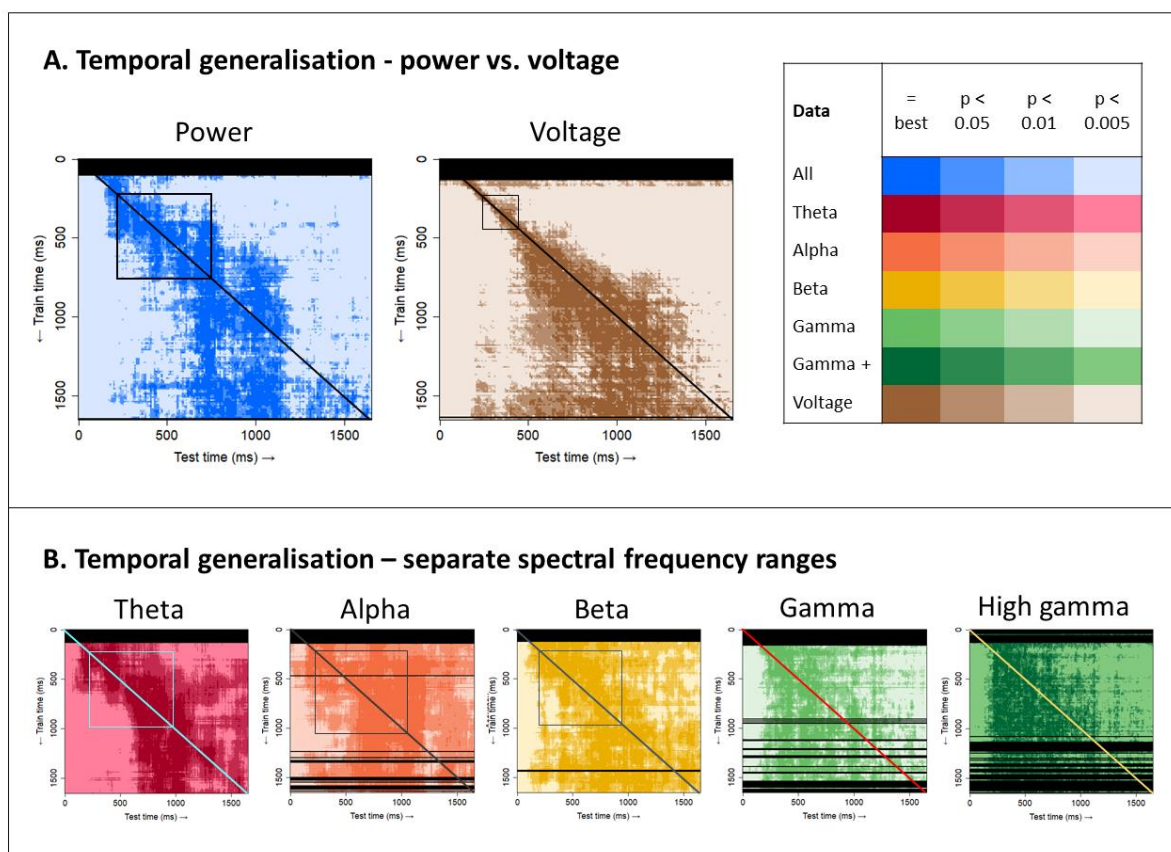
Our second question was whether the time-frequency code changes dynamically over stimulus processing in a manner similar to the voltage code. The preceding results show that time-frequency power exhibits one property also observed in voltage, namely constant decodability - animacy can be decoded from whole-spectrum power, and from power within each frequency range independently, from about 100-200ms post onset until about the mean onset of naming (1190ms in the subset of patients analysed by Rogers et al., 2021). Next, we applied the method of temporal generalisation to investigate whether time-frequency power, like voltage, shows (a) local temporal generalisation, indicating a changing neural code, and (b) a widening generalisation window, indicating rapid initial change that slows over time (Rogers et al., 2021).

Figure 2A shows the generalisation profile of classifiers trained on power at all frequencies (left) and those trained on voltage (right). The diagonal of each matrix indicates the hold-out accuracy of classifiers trained and tested at the same timepoint. Points closer to the diagonal indicate how well a classifier generalises to neighbouring timepoints, while points far from the diagonal indicate how well it generalises to timepoints further away. Local temporal generalisation is signified by better decoding close to the diagonal. In these plots, the most saturated colours indicate classifier accuracies statistically indistinguishable from the best-performing classifier at the same timepoint. Increasingly desaturated colours indicate accuracies that are reliably worse than the best-performing classifier at increasingly strict significance thresholds. Horizontal black bars indicate classifiers that did not exceed chance performance even when evaluated on held-out stimuli from the training timepoint.

The superimposed squares on each plot provide illustrative referents for evaluating whether the temporal generalisation pattern signifies a dynamically changing code. If classification accuracy is significantly higher for the two on-diagonal corners than the two off-diagonal corners, this signifies that (a) the neural signal reliably encodes semantic information at the two timepoints but (b) the signal changes in nature – whatever features of the signal support the good performance at the training timepoint are not present at the more remote testing points. This pattern is clearly observed for both voltage and whole-spectrum power (Figure 2A), indicating that the transfrequency semantic code exhibits dynamic properties similar to voltage.

When decoding from each range separately, however, the local generalisation pattern was less clear overall, and arguably present only in lower frequency ranges (theta, alpha, and beta; Figure 2B). Gamma and high gamma showed a different pattern: classifiers fitted at a given timepoint generalised approximately equally well to all prior timepoints back to the first timepoint where decoding was reliable (about 100-200ms post stimulus onset). That is, classifiers trained on gamma and high gamma showed excellent *retrograde* generalisation. This pattern yields the vertical “edge” clearly evident in the high gamma matrix, and blurrier but still present in the gamma matrix. The classifiers also generalised to future timepoints, but with performance often tailing off for more remote future timepoints — that is, they showed diminishing *anterograde* generalisation.

Together these observations suggest two interesting things about the semantic code as expressed within and across different frequency ranges. First, whatever information a classifier trained on gamma or high gamma power exploits at its own training timepoint must also be present in the signal at all earlier time points, all the way back to the first moment semantic information appears (producing good retrograde generalisation). However, information exploited by a classifier at its training timepoint may *not* be present in the signal at future timepoints, suggesting that these features “drop out” over the course of stimulus processing, producing a graded drop-off in anterograde generalisation. Second, the dynamically-changing pattern observed in the transfrequency code may be carried predominantly by lower frequency information.



284

285 Figure 2: Temporal generalisation of classifiers. For each test time, the train time (vertical
 286 dimension) of the best classifier, plus the train times of other classifiers that perform
 287 statistically indistinguishably from the best ($p > 0.05$, paired t-test with probabilities
 288 adjusted to control the false-discovery rate at $\alpha = 0.05$), are shown in the most saturated
 289 colour. Progressively desaturated colours show the train time of classifiers exhibiting a
 290 significant difference from the best classifier ($p < 0.05$, $p < 0.01$, $p < 0.005$). Timepoints
 291 where classifiers trained at that timepoint do not perform significantly better than chance
 292 even when tested at the training timepoint (0.5, one-sample t-test with probabilities
 293 adjusted to control the false-discovery rate at $\alpha = 0.05$) are shown in black. Local temporal
 294 generalisation is indicated by significantly better performance on timepoints at or near the
 295 train time (the diagonal shown on each plot) than at more remote timepoints. The squares
 296 highlight cases where hold-out accuracies at two training timepoints (the two corners on the
 297 diagonal) are both significantly better than when the same classifiers are used to decode
 298 more remote timepoints (the two off-diagonal corners). (A) Results for classifiers trained on
 299 power frequency features for all frequencies between 4 and 200 Hz (blue) or on voltage
 300 features (brown). (B) Results for classifiers trained on power frequency features from a
 301 single range - theta (4 – 7 Hz, red), alpha (8 – 12 Hz, orange), beta (13 – 30 Hz, yellow),
 302 gamma (30 – 60 Hz, light green) and high gamma (60 – 200 Hz, dark green).

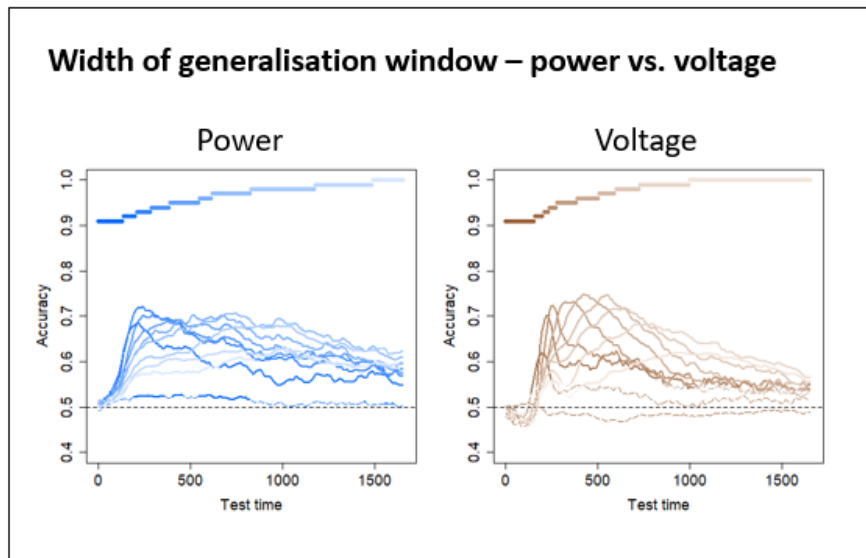


Figure 3: Width of generalisation window. Classifiers trained on power frequency features for all frequencies between 4 and 200 Hz (shades of blue) or on voltage features (shades of brown) were grouped into ten clusters via agglomerative hierarchical clustering (see Methods). The “timecourses” show the mean hold-out accuracy for classifiers within each cluster at each timepoint. Lines are solid where there was a significant difference between classifier accuracy and chance (0.5, one-sample t-test with probabilities adjusted to control the false-discovery rate at $\alpha = 0.05$) and dashed where there is no significant difference. Coloured bars show the grouped timepoints in each cluster. Statistics supporting this pattern are reported in Supplementary Materials 1, Figure S4).

We decided to focus subsequent analyses on frequency ranges in which a dynamic code was most clearly observed, namely voltage and whole-spectrum power. We next assessed whether the window of above-chance temporal generalisation widened over time, consistent with an overall slowing of representational change. To this end, we grouped the classifiers into ten clusters based on similarity in their temporal decoding profiles (see Methods). That is, classifiers that show similar patterns of prediction accuracy across all test timepoints were clustered together, and we computed the mean accuracy profile over time for each cluster. Figure 3 shows the result. Classifiers trained earlier showed a sharper rise and fall in accuracy, whereas classifiers trained later exhibited a wider window of generalisation (similar plots for classifiers trained on separate frequency ranges are shown in Supplementary Materials 1, Figure S3; statistics supporting this claim are reported in Supplementary Materials 1, Figure S4). Thus both voltage and tranfrequency power show a slowing of dynamic representational change over the course of stimulus processing.

Finally, we assessed whether the observed dynamic change was accompanied by change in the apparent code direction—that is, whether an increase in voltage or in power predicted a greater likelihood that the stimulus is an animal at some points in time, and a greater likelihood that it is inanimate at others. Change in code direction is another hallmark of nonlinear representational change (Rogers et al., 2021). To see this, we aggregated the classifier coefficients across patients for each timepoint and plotted these on the cortical surface. Specifically, at each cortical location, we first calculated the number of electrodes across patients that had a non-zero coefficient, then computed the proportion of these having a negative coefficient (since animals were coded as 0 and inanimate objects as 1, a negative coefficient means that an increase in voltage or power is associated with increased probability that the stimulus is an animal). Figure 4 shows these proportions at 4 different timepoints post stimulus onset, for classifiers trained on whole-spectrum power (top) or voltage (bottom). Warm colours indicate areas and timepoints where most selected electrodes have negative coefficients, while cool colours indicate areas and timepoints where most selected electrodes have positive coefficients. Green shades indicate areas and timepoints where selected electrodes have a mixture of positive and negative coefficients. We also animated the coefficient change over time, with videos for classifiers trained on all frequencies, individual frequency ranges, and voltage available at https://github.com/slfrisby/ECoG_LASSO/.

For both voltage and power, changes in code direction were observed. For voltage, for instance, the anterior-most part of the ventral temporal lobe was uniformly positive at 200ms and uniformly negative at 600ms. By 1000ms, midway between the anterior and posterior extent of coverage, coefficients showed a medial-to-lateral change in direction, suggesting that voltage increases signify animals in the more medial portion and non-animals in the more lateral portion. Whole-spectrum power also showed changes in code direction over time; for instance, in Figure 4, the medial anterior portion of ventral temporal cortex showed a negative relationship between power and animacy at 400ms, but a positive relationship by 1000ms. At 1000ms, the transfrequency coefficients also showed a medial-to-lateral change, with increased power signifying animals all along the medial extent of the coverage area, but showing a more mixed pattern across the more lateral extent.

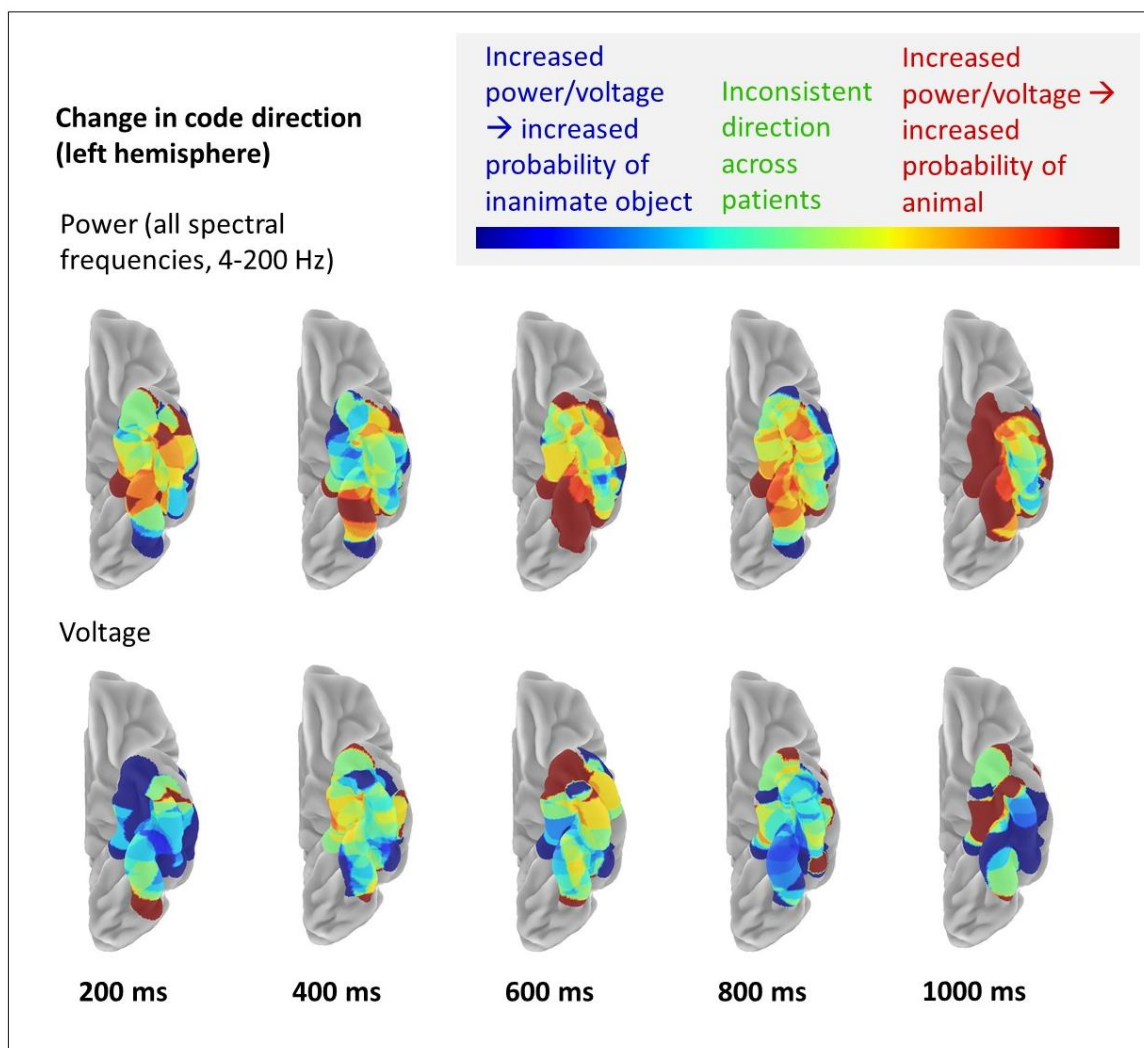


Figure 4: Change in code direction. Coefficients for classifiers trained on power frequency features for all frequencies between 4 and 200 Hz, or on voltage features, are projected onto the ventral surface of the left hemisphere. Time is shown relative to stimulus onset. Colours indicate the proportion of classifier coefficients at that location with the same sign (see Methods). Warm colours indicate that coefficients are mostly negative across patients (since animals were coded as 0 and inanimate objects as 1, a negative coefficient indicates that an increase in power is associated with increased probability that the stimulus is animate); cool colours indicate that coefficients are mostly positive across patients (an increase in power is associated with increased probability that the stimulus is inanimate); green shades indicate that the region was selected in multiple patients but that the sign of the coefficient is not consistent across patients.

2. Discussion

While it has long been clear that the vATLs play an important role in human semantic cognition (Lambon Ralph et al., 2017; Patterson et al., 2007; Rogers et al., 2004), precisely *how* neural populations encode such information remains uncertain. Recent

studies applying multivariate decoding to voltages recorded from the cortical surface suggest that semantic information is expressed in a rapidly-changing code (Rogers et al., 2021), but, with respect to uncovering local neural representations, such studies remain limited, since voltages can be influenced by a variety of sources both local and remote. The current work instead applied multivariate decoding to a time-frequency decomposition of ECoG data, with two results that advance our understanding of the vATL semantic code in important ways. First, information about animacy is expressed by power in all frequencies simultaneously - it appears within each range considered independently, and jointly across all ranges, for most of the duration of stimulus processing. Moreover, decoding accuracy was reliably better when models were fitted to all frequencies simultaneously compared to each individual range; and classifiers fitted to the whole spectrum always selected features from all frequency ranges no matter which timepoint they were trained on. This result strongly suggests conjoint encoding of semantic information across the power spectrum (Frisby et al., 2023). Second, decoding from both power and voltage revealed characteristics of dynamic representational change: constant decodability, local temporal generalisation, a widening generalisation window, and changes in code direction. This is likely to be driven by low frequencies – gamma and high gamma classifiers showed a contrasting pattern of excellent retrograde generalisation back to the earliest successful decoding timepoint, but anterograde generalisation that diminished for more remote timepoints.

Assuming that the frequency of activity is inversely related to the distance over which information is being transmitted (Canolty & Knight, 2010; Lachaux et al., 2012), our results are consistent with the claim that both local neuronal activity within the vATL (reflected in gamma and/or high gamma power) and distal inputs to the region (reflected in theta, alpha and beta) encode semantic information. These findings therefore agree with multiple univariate studies that variously associate theta (Hermes et al., 2014), alpha (Clarke et al., 2018), beta (Abel et al., 2015; Sato et al., 2021), gamma and high gamma (Arya, 2019; Cervenka et al., 2013; Chan et al., 2011; Crone et al., 2001; Crone & Hao, 2002; Edwards et al., 2010; Forseth et al., 2018; Kojima et al., 2013; Nakai et al., 2017, 2019; Snyder et al., 2023; Tanji, 2005; Wang et al., 2011) with semantic tasks. One interpretation of our results is that information is represented redundantly in multiple frequencies. A second possibility is that, since the brain exhibits properties of a dynamical system (Jones et al., 2015; Yuste, 2015), rhythmic patterns may be an epiphenomenon of normal functioning or even of

epileptiform activity (Drane et al., 2008; Hamberger, 2007; Henry et al., 1998). Therefore, some frequency ranges may represent information not in a “strong” sense (meaning that the brain uses time-frequency power to represent information for downstream computation) but in a “weak” sense (meaning that time-frequency power is decodable but is not used as the principal means of representation; Watrous et al., 2015).

However, the finding that classifiers trained on whole-spectrum power outperform those trained on a single frequency range and select features from across the frequency spectrum (Figures 1C and 1D) supports a more nuanced conclusion. Classifiers trained with LASSO regularisation prefer sparse solutions, in which only a small subset of features receive nonzero coefficients. If the information in the different frequency ranges were redundant, a classifier would not assign nonzero coefficients to many features that spanned the frequency range. Instead, these results support the notion of *transfrequency* representation – information is encoded *across* multiple frequency ranges (sometimes called frequency “multiplexing”, Watrous et al., 2015). Some previous univariate studies have proposed specific roles that different frequency ranges may play - abstraction from multiple episodic experiences (Sauseng & Klimesch, 2008), binding multiple features of an item (Merker, 2013; Sauseng & Klimesch, 2008; Stryker, 1989; Tallon-Baudry & Bertrand, 1999; von der Malsburg, 1995), encoding semantic similarity (Solomon et al., 2019; Spiers, 2020), and enabling retrieval independent of sensory modality (Sauseng & Klimesch, 2008) in a way that is appropriate for the context (Halgren et al., 2015; Jackson, 2021; Klimesch, 2012; Marko et al., 2019). However, on the simple assumption that the frequency of activity is inversely related to transmission distance, the notion of transfrequency representation is highly consistent with the hub-and-spoke model (Lambon Ralph et al., 2017; Patterson et al., 2007; Rogers et al., 2004). According to this model, neither the vATL “hub” nor the modality-specific “spokes” are individually sufficient for full semantic cognition – rather, information is encoded conjointly in multiple regions and in the interactions between them (Frisby et al., 2023).

This study was the first to investigate whether activity in different frequency ranges exhibits the same pattern of dynamic representational change seen in voltage (Rogers et al., 2021). Although dynamic change was clearly evident when decoding whole-spectrum power, gamma and high gamma showed a different pattern – excellent retrograde generalisation and diminishing anterograde generalisation. The good retrograde

generalisation indicates that features exploited by a classifier at its training timepoint must also be present, and must signify the same information in the same way, for all preceding timepoints (up to the earliest successful timepoint). That is, this part of the code must be relatively *static* over processing from the earliest arrival of perceptual signals up to the time where the classifier is fitted. Conversely, the diminishing anterograde generalisation indicates that features exploited by a classifier at its training timepoint eventually fall away at future timepoints.

Figure 5 illustrates in schematic form what such a decoding pattern might suggest about how animacy is encoded in high-frequency activity (and thus, by inference, in local neural activity). Each curve represents a hypothetical response to animate stimuli in the gamma or high gamma range for one electrode over time. Every electrode shows increased gamma power for animate stimuli from early in processing, but the timecourse of this change differs between electrodes. Where gamma increases most strongly early in processing, it also diminishes most rapidly; where gamma increases are smaller initially, they persist longer. Since the LASSO penalty prefers to assign features the smallest coefficients possible, models fitted with LASSO regularisation will select whichever electrodes show the strongest response at the training timepoint. Thus models fitted to the earliest timepoints will select the red electrodes, those fitted slightly later will select the orange ones, etc. Generalisation to future timepoints will diminish as the gamma power decreases on the selected electrodes, producing diminishing anterograde generalisation. Because all electrodes carry useful signal from the onset, however, each classifier generalises successfully to all prior time points, producing good retrograde generalisation. The generalisation pattern is thus consistent with the view that animacy is encoded in the local activity of many neural populations, some responding strongly but then quickly diminishing, and other initially responding less strongly but persisting for a longer duration.

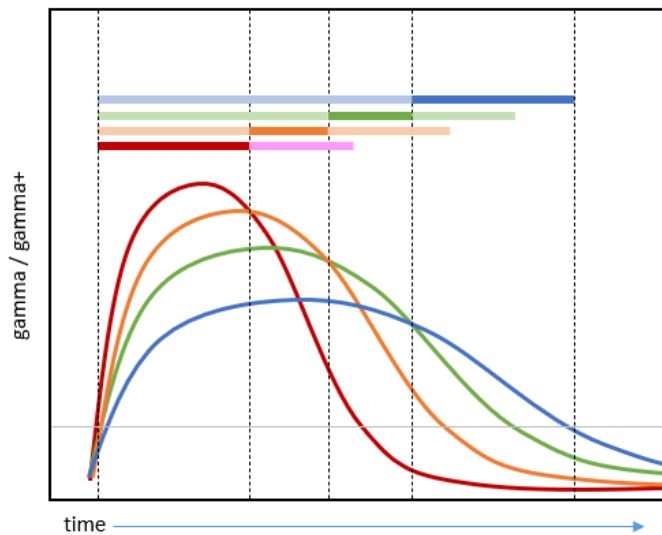


Figure 5: Hypothesised semantic code in gamma and/or high gamma producing full retrograde generalisation and limited anterograde generalisation. Each curve represents the hypothesised gamma/high gamma response to animate stimuli by one electrode. Where curves are above the grey horizontal line, they encode information about animacy. All electrodes encode animacy from early in processing, but models regularised with the LASSO penalty will place coefficients on whichever electrode has the strongest response at a given timepoint. The saturated parts of the coloured horizontal bars thus indicate which electrode is selected by a classifier at each timepoint. The desaturated parts indicate the generalisation profile of the resulting classifier. Electrodes that show the strongest responses early on diminish in power most rapidly, limiting anterograde generalisation. Because all electrodes carry information about animacy from the onset of the signal, however, all classifiers generalise well to the beginning of the processing window.

Finally, the current work carries important implications for interpreting other sources of information about semantic representation. Noninvasive methods such as EEG and MEG cannot discern activity within high gamma frequencies because it is blocked by the skull and the scalp. The current work suggests that dynamic representational change is mainly observed in lower frequency ranges that are more readily measurable via these other techniques—thus the comparatively static code revealed in high frequency ranges may be obscured in other imaging modalities. Additionally, the current work highlights circumstances in which representations in neural activity differ in their nature from representations observed in neuro-computational models (which exhibit the dynamic properties observed in voltage and transfrequency power, not the comparatively static code observed in high frequencies). Further work should explore the circumstances under which representational properties in brains and machines converge and diverge (Cox et al., 2015).

2.1. Conclusion

To summarise, we have shown that it is possible to decode semantic information from every frequency range between theta and high gamma. Although individual frequency ranges contained enough information to drive decoding above chance, decoding from the whole frequency spectrum at once produced the highest decoding accuracy, equivalent to using voltage. Decoding from the whole frequency spectrum also revealed the same set of dynamic properties observed in voltage; however decoding of high-frequency ranges alone showed a qualitatively different pattern, suggesting that local firing activity expresses a more static semantic code. Together these results provide an important step toward understanding how neural systems encode semantic information.

3. Methods

3.1. Patients

19 patients participated in the study (labelled 01-22 to enable comparison with previous work; Y. Chen et al., 2016; Cox et al., 2024; Rogers et al., 2021; Shimotake et al., 2015). Of these 19 patients, one was later excluded because too many trials were contaminated with artefacts (see Preprocessing – ECoG), leaving 18 patients for decoding. For one further patient (Patient 20), no exact MNI coordinates were available for their electrodes. This patient was included in the analyses, but excluded from visualisation (see Experimental Questions). All patients were native speakers of Japanese. Information about patients' age, sex, handedness, and clinical presentation is summarised in Table 1.

Each patient was implanted with subdural electrodes arranged in grids or strips for presurgical monitoring (mean 91 electrodes, range 56 – 128 electrodes per patient). 16 patients had electrodes implanted in the left hemisphere, of which 10 – 52 electrodes (mean 27 electrodes) were within grids or strips that covered the vATL (where grids or strips included both electrodes that covered ventral anterior temporal cortex and electrodes that covered lateral or posterior temporal cortex, the whole grid or strip was included in the analysis). In the remaining 3 patients, with electrodes implanted in the right hemisphere, 6 –

28 electrodes (mean 20 electrodes) were within grids or strips that covered the vATL. Grids in which no electrodes covered the vATL were excluded from analysis. Electrodes that were included in the analysis are shown in Figure 6. The electrodes were platinum, with a recording diameter of 2.3 mm and an inter-electrode distance of 1 cm (ADTECH, WI). Three patients were also implanted with depth electrodes (sEEG), but these were not included in the analysis.

All patients gave written informed consent and the study was approved by the ethics committee of the Kyoto University Graduate School of Medicine (#C533).

	Patient 01	Patient 02	Patient 03	Patient 04
Age, sex, handedness	22, M, R	29, M, R&L	17, F, R	38, F, R
WAIS-R/WAIS-III* (VIQ, PIQ, TIQ)	70, 78, 69	72, 78, 72	67, 76, 69	84, 97, 89
WMS-R (verbal, visual, general, attention, delayed recall)	99, 64, 87, 91, 82	99, 92, 97, 87, 83	51, <50, <50, 81, 56	75, 111, 83, 62, 53
WAB	95.6	96	97.2	98.5
WADA	Left	Bilateral	Left	Left
Age of seizure onset	16	10	12	29
Seizure type	FAS→FIAS, FBTCs	FAS→FIAS	FAS→FIAS	FAS→FIAS
Ictal ECoG onset	aMTG	PHG	PHG	PHG
MRI	L basal frontal cortical dysplasia, L anterior temporal arachnoid cyst	L posterior temporal cortical atrophy	L temporal tip arachnoid cyst	L HS/HA
Pathology	FCD type I	FCD type IIIa	Palmini FCD type IB	FCD type IIa
	Patient 05	Patient 06	Patient 07	Patient 08
Age, sex, handedness	55, M, R	34, M, L	41, F, R	27, F, R
WAIS-R/WAIS-III* (VIQ, PIQ, TIQ)	105, 99, 103	55, **, 44	72, 83, 75	106, **, 105
WMS-R (verbal, visual, general, attention, delayed recall)	71, 117, 84, 109, 72	52, <50, <50, 55, <50	83, 111, 89, 94, 82	112, 114, 114, 81, 100
WAB	98	88	97.3	99.6
WADA	Left	Left	Right	Left
Age of seizure onset	55	12	19	16
Seizure type	FIAS (once)	FAS→FIAS	FAS→FIAS	FAS→FIAS
Ictal ECoG onset	None	R parietal lobe/pMTG	PHG	ventral anterior temporal
MRI	L medial temporal lobe low-grade glioma	R parietal cerebral atrophy & contusion, R hippocampal sclerosis/atrophy	L HS/HA, L parieto-occipital perinatal infarction	R medial temporal cyst
Pathology	Diffuse astrocytoma	Post-traumatic change(parietal)/scar(temporal)/ HS	FCD type I	FCD type I
	Patient 09	Patient 10	Patient 11	Patient 12
Age, sex, handedness	51, M, R	38, F, R	29, F, R	40, M, L

WAIS-R/WAIS-III* (VIQ, PIQ, TIQ)	73, 97, 83	109, 115, 112	62, 80, 67	93, 105, 98
WMS-R (verbal, visual, general, attention, delayed recall)	80, 101, 85, 91, 91	71, 79, 70, 90, 58	64, 94, 68, 79, 79	74, 94, 77, 110, 96
WAB	89.6	96.9	95.8	99
WADA	Left	Left	Left	Right
Age of seizure onset	43	28	12	6
Seizure type	FIAS	FAS→FIAS	FAS→FIAS	FAS→FIAS
Ictal ECoG onset	mITG	SMG	PHG	PHG
MRI	L temporal cavernoma	L parietal operculum tumour	L HS	R HS/HA
Pathology	Arteriovenous malformation	Oligoastrocytoma	non-neoplastic brain tissue	FCD type IIIa
	Patient 13	Patient 14	Patient 15	Patient 20
Age, sex, handedness	22, M, R	42, M, R	35, M, R	23, F, R
WAIS-R/WAIS-III* (VIQ, PIQ, TIQ)	86, 79, 81	96, 84, 90	82, 86, 82	67, 82, 71
WMS-R (verbal, visual, general, attention, delayed recall)	55, 79, 53, 90, 54	73, 85, 73, 103, 76	75, 89, 75, 92, 81	78, 119, 86, 82, 64
WAB	97.4	99.9	99.2	94.2
WADA	Left	Left	Left	Bilateral
Age of seizure onset	14	27	20	15
Seizure type	FAS→FIAS	FAS→FIAS	FIAS	FIAS
Ictal ECoG onset	PHG	PHG	PHG	IPL/SMG
MRI	L HS/HA	L HS	L HS/HA	No apparent lesion
Pathology	FCD type IIIa	FCD type IIIa	FCD type IIIa	FCD type I
	Patient 21	Patient 22		
Age, sex, handedness	40, M, R	28, F, R		
WAIS-R/WAIS-III* (VIQ, PIQ, TIQ)	86, 97, 90	80, 69, 72		
WMS-R (verbal, visual, general, attention, delayed recall)	111, 113, 113, 87, 109	77, 89, 77, 121, 73		
WAB	92.2	100		
WADA	Left	Left		
Age of seizure onset	30	12		
Seizure type	FIAS	FIAS		
Ictal ECoG onset	aMTG	PHG		
MRI	No apparent lesion	L HS/HA		
Pathology	FCD type I	FCD type IIIa		

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535 Table 1: Patient characteristics. Abbreviations: WAIS-R – Wechsler Adult Intelligence Scale
536 (1991), WAIS-III - Wechsler Adult Intelligence Scale (1997), VIQ – Verbal IQ, PIQ –
537 Performance IQ, TIQ – full-scale IQ, WMS-R – Wechsler Memory Scale (1987), WAB –
538 Western Aphasia Battery, FAS – focal aware seizure, FIAS – focal impaired awareness
539 seizure, FBTCS – focal to bilateral tonic-clonic seizure, aMTG – anterior middle temporal
540 gyrus, PHG – parahippocampal gyrus, pMTG – posterior middle temporal gyrus, mITG –
541 medial inferior temporal gyrus, SMG - supramarginal gyrus, ITG – inferior temporal gyrus,
542 IPL – intraparietal lobule, HS – hippocampal sclerosis, HA – hippocampal atrophy, FCD –
543 focal cortical dysplasia. * - The WAIS-R was used to test patients 01-06 and the WAIS-III was
544 used to test other patients. ** - missing score.

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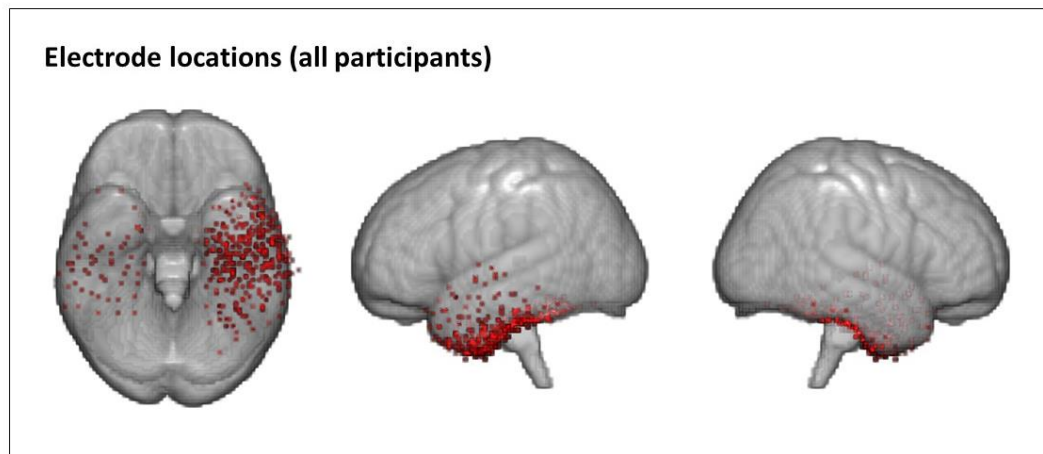


Figure 6: Electrode locations for all patients. Electrodes are overlaid on an MNI template (MNI152NLin2009cAsym).

3.2. Stimuli, task, and acquisition

Stimuli were the same 100 line drawings used previously (Y. Chen et al., 2016; Rogers et al., 2021; Shimotake et al., 2015) – 50 animals and 50 nonliving items including buildings, tools, musical instruments and other household objects (Morrison et al., 1997; https://github.com/slfrisby/ECOG_LASSO/tree/main/data_info/stimuli/). There were no significant differences between the categories with respect to age of acquisition, visual complexity, concreteness, familiarity, word frequency, name agreement and non-semantic visual structure (Barry et al., 1997; Rogers et al., 2021). MATLAB r2010a was used to display stimuli on a PC screen.

There were four runs per patient, collected in a single session. Within each run, each stimulus was presented once in a random order. Stimuli were presented for 5 seconds each, with no break between stimuli.

Patients were instructed to name each picture as quickly and accurately as possible. Data for nine patients were recorded at 2000 Hz (with a low-pass filter of 600 Hz) and data for ten patients were recorded at 1000 Hz (with a low-pass filter of 300 Hz). Time of naming onset was measured. Responses and eye fixation were monitored via video.

3.3. Data analysis

3.3.1. Preprocessing – structural MRI

A clinical MPRAGE T₁-weighted anatomical scan was acquired before and after electrode implantation. The location of each electrode was identified on each 2D slice of the post-surgical scan. *fnirt* in FSL (<https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/>; Jenkinson et al., 2012; Smith et al., 2004) was used to coregister the electrode positions to the pre-surgical scan and then to MNI space (MNI152NLin2009cAsym). The position of each electrode was then manually adjusted to the surface.

3.3.2. Preprocessing – ECoG

Preprocessing was performed using in-house MATLAB (r2023b) code, composed of functions from EEGLAB (2023.1; <https://eeglab.org/>; Delorme & Makeig, 2004) and available at https://github.com/slfrisby/ECoG_LASSO/. First, the CleanLine EEGlab plugin (v2.0; <https://github.com/scn/cleanline/>; Delorme, 2023; Mitra & Bokil, 2007) was used to remove line noise at 60 Hz and the harmonics 120 and 180 Hz, without scanning for exact line noise frequencies and with a sliding window step size of 2 (all other parameters were set as defaults). CleanLine uses a multitaper fast Fourier transform to identify and remove sinusoidal noise and, though it removes noise less completely than traditional notch-filtering, is less disruptive to the overall frequency structure of the data. Data were then filtered using *eegfilt* (EEGLAB). The low cutoff was set to 0.5 Hz (to remove slow drifts; Delorme, 2023) and the high cutoff set to 300 Hz (for consistency across patients - some data were recorded at 1000 Hz using equipment that imposed a low-pass filter at 300 Hz). Channels that lay below the seizure onset zone (identified in the clinical examinations) or with poor contact (identified by visual inspection) were removed. Data were epoched between -1000 and 3000 ms relative to stimulus onset and baseline-corrected using the mean response across trials between -200 and -1 ms. Data from the nine patients recorded at 2000 Hz was then down-sampled to match the ten patients recorded at 1000 Hz by boxcar averaging pairs of neighbouring timepoints. A common average reference was applied using *reref* (EEGLAB).

Addressing artefacts in the data was a five-step process. First, the mean voltage for each channel was calculated over timepoints and trials. Any trial that contained a value more extreme than 10 standard deviations from the mean was rejected. Second, all trials were inspected individually and any that appeared to contain obvious interictal epileptiform

activity, muscle activity, “electrode pop” or other artefacts were rejected. Since repeated presentations of the same stimulus were to be averaged it was important to check whether there was at least one good trial for each stimulus. One patient lacked any good trials for 18 stimuli (7 animate, 11 inanimate) and so was excluded from further analysis (and does not appear in Table 1). Another lacked any good trials for 3 stimuli (1 animate, 2 inanimate) – because of the small and relatively balanced number of missing stimuli, we included this patient. For the next steps an independent component analysis (ICA) was conducted using *runica* (EEGLAB) extracting n components, where n was 75% of the total number of good electrodes (Clarke, 2020). For the third artefact rejection step, the SASICA EEGLAB plugin (<https://github.com/dnacombo/SASICA/>; Chaumon et al., 2015) was used to identify ICA components with weak autocorrelation, which is characteristic of muscle activity. Components identified by SASICA were inspected and, in order to minimise rejection of signal-carrying data, rejected if the channel appeared to contain only autocorrelated noise and no neural activity. Fourth, SASICA was used to highlight components with focal trial activity (i.e., activity in only a few trials, characteristic of muscle activity), “electrode pop” or other artefacts. Rather than rejecting the component across all trials, the atypical trials were identified via visual inspection and only those trials were removed. The ICA was then re-run and the process repeated until no components contained significant focal trial activity. Fifth, the possibility of microsaccade artefacts in the data was assessed (Clarke, 2020; Jerbi et al., 2009; Kovach et al., 2011; Yuval-Greenberg et al., 2008). The independent component activations were filtered for activity in the frequency range associated with microsaccades (20 – 190 Hz). The activations were then convolved with a saccade-related potential template and the number of microsaccades per second was calculated (Craddock et al., 2016). Since every component contained very few microsaccades (< 0.000012 microsaccades per second) and no component obviously contained more microsaccades than any other, no components were rejected based on this assessment.

Time-frequency power and phase were extracted using complex Morlet wavelet convolution (Bertrand et al., 1994), implemented with *timefreq* (EEGLAB). Power values were extracted from every trial between 0 and 1650 ms in 10 ms time steps and between 4 and 200 Hz in 60 logarithmically-spaced frequency steps. A 5-cycle wavelet was used at 4 Hz, increasing to a 15-cycle wavelet at 200 Hz (Clarke, 2020). Power values were then averaged across repeated presentations of the same stimulus and any missing trials were interpolated

with the median power value across all trials. Decibel normalisation was performed using the mean power across trials between -300 and -100 ms (calculated separately for each electrode and frequency) – a gap before stimulus onset was used to mitigate the effect of temporal leakage of trial activity into the baseline during wavelet convolution. Since phase values cannot be averaged (Cohen, 2014), preprocessed trials were averaged across repeated presentations of the same stimulus before phase values were extracted using the same time and frequency steps used for the extraction of power. As a comparison, preprocessed voltage was averaged over repeated presentations of the same stimulus.

The impact of preprocessing decisions on decoding results is explored in Supplementary Materials 2. In summary, decoding of semantic structure (animacy) with spectral frequency features remains unchanged after each step in this extensive and careful cleaning.

3.3.3. Multivariate classification

3.3.3.1. Decoding approach

Logistic regression classifiers were trained to discriminate animate from inanimate stimuli using *glmnet* in R (Friedman et al., 2010). For power and phase, frequency feature vectors were created for each stimulus at each timepoint (0 ms, 10 ms, 20 ms, ...) by concatenating power or phase values for all electrodes for all frequencies in a range of interest. Voltage feature vectors were created for each stimulus at each timepoint by concatenating voltage values for all electrodes in a 50 ms window centred on the timepoint of interest. The voltage feature vectors therefore reflected activity around the timepoint of interest – but note that so did frequency feature vectors, since Morlet wavelet convolution takes activity at neighbouring timepoints into account. These vectors were provided as input to the classifiers. Each classifier was trained on vectors generated at a single timepoint for a single patient.

The classifiers used LASSO (L1) regularisation (Tibshirani, 1996), which applies a penalty that scales with the sum of the absolute values of the coefficients and thus produces solutions in which many features receive coefficients of zero. This approach was selected because it can be used to assess whether the same electrodes or frequencies are used to represent information at different timepoints. If the information used by a classifier trained at timepoint A is present at timepoint B, the classifier will perform well at timepoint

B; other units, that may be in different states at the two timepoints, will receive coefficients of zero and therefore will not affect classifier performance (Rogers et al., 2021).

Classifier accuracy was assessed using 10-fold cross-validation. In each outer loop, 10 out of 100 stimuli (5 animate, 5 inanimate) were held out. The remaining 90 stimuli were used to search a range of 100 values (0.2 – 0.002, logarithmically spaced) for the regularisation parameter that resulted in the smallest mean squared error. This process was implemented using *cv.glmnet* (glmnet) with parallel model fitting implemented with foreach (<https://cran.r-project.org/web/packages/foreach/index.html>; all other parameters were set as defaults). A model with the best regularisation parameters was tested on the 10 stimuli in the outer loop hold-out set. The same model was tested on the same 10 stimuli at all other timepoints. The process was then repeated 10 times with different final hold-out sets. This process yielded both a main timecourse of decoding accuracy (i.e., mean accuracy, over folds, of classifiers tested on the timepoint at which they were trained) and a generalisation matrix in which the y-coordinate of a cell is the time at which the classifier was trained and the x-coordinate is the time at which the classifier was tested. Finally, a single classifier was trained at every timepoint using all 100 stimuli for training. The accuracy of these classifiers was not evaluated – they were used only for inspection of coefficients.

To assess group-level performance, the timecourses of decoding accuracy and the generalisation matrices were averaged across patients.

3.3.3.2. Experimental questions

We first assessed whether power, phase and voltage each contained sufficient information to enable decoding. We therefore created both voltage feature vectors and frequency feature vectors of power and phase data (separately) using all 60 frequencies between 4 and 200 Hz and used these as input to classifiers. We compared each group-average timecourse to chance (0.5) using one-tailed, one-sample t-tests. We compared power and voltage, and phase and voltage, using paired t-tests. Probabilities were adjusted to control the false-discovery rate at $\alpha = 0.05$ (Benjamini & Hochberg, 1995). We confirmed that near-identical patterns of results were obtained in the 8 patients with left-hemisphere electrodes previously analysed by (Rogers et al., 2021), the 7 patients with left-hemisphere electrodes analysed for the first time in this work, and the 3 patients with right-hemisphere

electrodes (Supplementary Materials 1, Figure S5). Having done so, we combined all patients into a single group for all subsequent analyses.

We next investigated whether there were differences in decoding performance between power and phase within different frequency ranges. We therefore divided the 60 frequencies into theta (4 – 7 Hz, 11 frequencies), alpha (8 - 12 Hz, 7 frequencies), beta (13 – 30 Hz, 13 frequencies), gamma (30 – 60 Hz, 10 frequencies) and high gamma (60 – 200 Hz, 19 frequencies) ranges. We constructed frequency vectors of power and phase data using only power or phase at frequencies within each range and used these as input to separate classifiers. We compared each group-average timecourse to chance and then compared each range to the timecourse of decoding using all 60 frequencies.

We then tested whether, if either power or phase data contained sufficient information to enable decoding, the power or phase code exhibited the same distributed, dynamic properties identified by Rogers et al. (2021) in both the voltage data (for the first subset of patients) and the computational hub-and-spoke model. We tested for these properties using both frequency vectors constructed using all 60 frequencies and vectors constructed using only frequencies within each range. Constant decodability had already been assessed by one-sample t-tests of each group-average timecourse against chance.

To assess local temporal generalisation we inspected the group-average generalisation matrices generated by testing each classifier at every possible timepoint. To test whether the patterns observed were statistically significant, we identified the best-performing classifier at each timepoint and then conducted paired t-tests to assess whether classifiers trained at every other possible timepoint performed equally well. Probabilities were adjusted to control the false-discovery rate at $\alpha = 0.05$ (Benjamini & Hochberg, 1995).

To assess the shape of the generalisation window over time we computed the pairwise cosine distance between each row of the generalisation matrix, then applied agglomerative hierarchical clustering using *hclust* (native R) and cut the tree to create 10 clusters. We selected this cluster number to make these results comparable to those of Rogers et al. (2021). We averaged the timecourses of classifiers within each cluster and inspected performance over time (again, probabilities were adjusted to control the false-discovery rate). To quantify the patterns observed we took classifiers trained every 50 ms (i.e., 0 ms, 50 ms, 100 ms ...; this ensured that, for classifiers trained on voltage, consecutive 50 ms windows did not overlap and were therefore independent). We calculated the area

under the curve between each timecourse and a horizontal line at chance (0.5) and fitted a piecewise linear regression to the area values using the *segmented* library in R, using *segmented* with the Bayesian Information Criterion to determine the number and location of breakpoints to produce the best model given the number of free parameters.

To assess and visualise changes in code direction we first took the classifiers trained on all data at each timepoint and then (separately for every electrode, every patient and every timepoint) calculated the mean classifier coefficient within each frequency range (or over all 60 frequencies, or over the 50 ms within each window in the case of voltage). These mean classifier coefficients were converted to one NIFTI volume per patient and per timepoint, at 2.5 mm isotropic resolution in MNI space (MNI152NLin2009cAsym), using SPM12 (<https://www.fil.ion.ucl.ac.uk/spm/>) implemented in MATLAB r2023b. One patient (Patient 20) was excluded from this analysis because no MNI coordinates were available for their electrodes. Other patients were missing MNI coordinates from only a subset of electrodes; we included those patients in the analysis but ignored coefficients on the electrodes with missing coordinates. We projected coefficients in each volume to the pial surface (fsaverage template; Dale et al., 1999; Fischl et al., 1999) using *-volume-to-surface-mapping* with trilinear interpolation and smoothed them on the surface at 6 mm FWHM using *-metric-smoothing* in Connectome Workbench 1.5.0. We calculated the proportion of negative coefficients that each vertex received (since animals were coded as 0 and inanimate objects as 1, a negative coefficient indicates that an increase in power is associated with increased probability that the stimulus is animate). We plotted these proportions on the pial surface using *plot_surf_stat_map* in *nilearn* 0.10.3 implemented in Python 3.9. Finally, we animated the results – ffmpeg 5.1.6 (<https://ffmpeg.org/>) was used to concatenate plots of coefficients at successive timepoints with a frame rate of 10 (10 times slower than real time) and motion interpolation.

Data and code availability: We are unable to share raw data for this study because the patients did not provide informed consent to do so. However, matrices containing power, phase and voltage features (columns) for each stimulus (rows) will be made available upon peer review and acceptance. Code is available at https://github.com/slfrisby/ECoG_LASSO/.

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References

- Abel, T. J., Rhone, A. E., Nourski, K. V., Kawasaki, H., Oya, H., Griffiths, T. D., Howard, M. A., & Tranel, D. (2015). Direct physiologic evidence of a heteromodal convergence region for proper naming in human left anterior temporal lobe. *Journal of Neuroscience*, 35(4), 1513–1520. <https://doi.org/10.1523/JNEUROSCI.3387-14.2015>
- Arya, R. (2019). Similarity of spatiotemporal dynamics of language-related ECoG high-gamma modulation in Japanese and English speakers. *Clinical Neurophysiology*:

- 794 *Official Journal of the International Federation of Clinical Neurophysiology*, 130(8),
795 1403–1404. <https://doi.org/10.1016/j.clinph.2019.05.006>
- 796 Barry, C., Morrison, C. M., & Ellis, A. W. (1997). Naming the Snodgrass and Vanderwart
797 pictures: Effects of age of acquisition, frequency, and name agreement. *The*
798 *Quarterly Journal of Experimental Psychology Section A*, 50(3), 560–585.
799 <https://doi.org/10.1080/783663595>
- 800 Başar-Eroglu, C., Strüber, D., Schürmann, M., Stadler, M., & Başar, E. (1996). Gamma-band
801 responses in the brain: A short review of psychophysiological correlates and
802 functional significance. *International Journal of Psychophysiology*, 24(1–2), 101–112.
803 [https://doi.org/10.1016/S0167-8760\(96\)00051-7](https://doi.org/10.1016/S0167-8760(96)00051-7)
- 804 Benjamini, Y., & Hochberg, Y. (1995). Controlling the false discovery rate: A practical and
805 powerful approach to multiple testing. *Journal of the Royal Statistical Society, Series*
806 *B (Statistical Methodology)*, 57(1), 289–300.
- 807 Bertrand, O., Bohorquez, J., & Pernier, J. (1994). Time-frequency digital filtering based on an
808 invertible wavelet transform: An application to evoked potentials. *IEEE Transactions*
809 *on Biomedical Engineering*, 41(1), 77–88. *IEEE Transactions on Biomedical*
810 *Engineering*. <https://doi.org/10.1109/10.277274>
- 811 Binney, R. J., Embleton, K. V., Jefferies, E., Parker, G. J. M., & Lambon Ralph, M. A. (2010).
812 The ventral and inferolateral aspects of the anterior temporal lobe are crucial in
813 semantic memory: Evidence from a novel direct comparison of distortion-corrected
814 fMRI, rTMS, and semantic dementia. *Cerebral Cortex*, 20(11), 2728–2738.
815 <https://doi.org/10.1093/cercor/bhq019>
- 816 Bressler, S. L. (1990). The gamma wave: A cortical information carrier? *Trends in*
817 *Neurosciences*, 13(5), 161–162. [https://doi.org/10.1016/0166-2236\(90\)90039-D](https://doi.org/10.1016/0166-2236(90)90039-D)
- 818 Canolty, R. T., & Knight, R. T. (2010). The functional role of cross-frequency coupling. *Trends*
819 *in Cognitive Sciences*, 14(11), 506–515. <https://doi.org/10.1016/j.tics.2010.09.001>
- 820 Carlson, T., Tovar, D. A., Alink, A., & Kriegeskorte, N. (2013). Representational dynamics of
821 object vision: The first 1000 ms. *Journal of Vision*, 13(10), 1–1.
822 <https://doi.org/10.1167/13.10.1>
- 823 Cervenka, M. C., Corines, J., Boatman-Reich, D. F., Eloyan, A., Sheng, X., Franaszczuk, P. J., &
824 Crone, N. E. (2013). Electrocorticographic functional mapping identifies human
825 cortex critical for auditory and visual naming. *NeuroImage*, 69, 267–276.
826 <https://doi.org/10.1016/j.neuroimage.2012.12.037>
- 827 Chan, A. M., Baker, J. M., Eskandar, E., Schomer, D., Ulbert, I., Marinkovic, K., Cash, S. S., &
828 Halgren, E. (2011). First-pass selectivity for semantic categories in human
829 anteroventral temporal lobe. *Journal of Neuroscience*, 31(49), 18119–18129.
830 <https://doi.org/10.1523/JNEUROSCI.3122-11.2011>
- 831 Chaumon, M., Bishop, D. V. M., & Busch, N. A. (2015). A practical guide to the selection of
832 independent components of the electroencephalogram for artifact correction.
833 *Journal of Neuroscience Methods*, 250, 47–63.
834 <https://doi.org/10.1016/j.jneumeth.2015.02.025>
- 835 Chen, L., Lambon Ralph, M. A., & Rogers, T. T. (2017). A unified model of human semantic
836 knowledge and its disorders. *Nature Human Behaviour*, 1(3), 0039.
837 <https://doi.org/10.1038/s41562-016-0039>

- Chen, Y., Shimotake, A., Matsumoto, R., Kunieda, T., Kikuchi, T., Miyamoto, S., Fukuyama, H., Takahashi, R., Ikeda, A., & Lambon Ralph, M. A. (2016). The 'when' and 'where' of semantic coding in the anterior temporal lobe: Temporal representational similarity analysis of electrocorticogram data. *Cortex*, 79, 1–13. <https://doi.org/10.1016/j.cortex.2016.02.015>
- Cichy, R. M., Pantazis, D., & Oliva, A. (2014). Resolving human object recognition in space and time. *Nature Neuroscience*, 17(3), 455–462. <https://doi.org/10.1038/nn.3635>
- Clarke, A. (2020). Dynamic activity patterns in the anterior temporal lobe represents object semantics. *Cognitive Neuroscience*, 11(3), 111–121. <https://doi.org/10.1080/17588928.2020.1742678>
- Clarke, A., Devereux, B. J., & Tyler, L. K. (2018). Oscillatory dynamics of perceptual to conceptual transformations in the ventral visual pathway. *Journal of Cognitive Neuroscience*, 30(11), 1590–1605. https://doi.org/10.1162/jocn_a_01325
- Cohen, M. X. (2014). *Analyzing Neural Time Series Data: Theory and Practice*. MIT Press.
- Contini, E. W., Wardle, S. G., & Carlson, T. A. (2017). Decoding the time-course of object recognition in the human brain: From visual features to categorical decisions. *Neuropsychologia*, 105, 165–176. <https://doi.org/10.1016/j.neuropsychologia.2017.02.013>
- Cox, C. R., & Rogers, T. T. (2021). Finding distributed needles in neural haystacks. *Journal of Neuroscience*, 41(5), 1019–1032. <https://doi.org/10.1523/JNEUROSCI.0904-20.2020>
- Cox, C. R., Rogers, T. T., Shimotake, A., Kikuchi, T., Kunieda, T., Miyamoto, S., Takahashi, R., Matsumoto, R., Ikeda, A., & Lambon Ralph, M. A. (2024). Representational similarity learning reveals a graded multidimensional semantic space in the human anterior temporal cortex. *Imaging Neuroscience*, 2, 1–22. https://doi.org/10.1162/imag_a_00093
- Cox, C. R., Seidenberg, M. S., & Rogers, T. T. (2015). Connecting functional brain imaging and Parallel Distributed Processing. *Language, Cognition and Neuroscience*, 30(4), 380–394. <https://doi.org/10.1080/23273798.2014.994010>
- Craddock, M., Martinovic, J., & Müller, M. M. (2016). Accounting for microsaccadic artifacts in the EEG using independent component analysis and beamforming. *Psychophysiology*, 53(4), 553–565. <https://doi.org/10.1111/psyp.12593>
- Crone, N. E., Boatman, D., Gordon, B., & Hao, L. (2001). Induced electrocorticographic gamma activity during auditory perception. *Clinical Neurophysiology*, 112(4), 565–582. [https://doi.org/10.1016/S1388-2457\(00\)00545-9](https://doi.org/10.1016/S1388-2457(00)00545-9)
- Crone, N. E., & Hao, L. (2002). Functional dynamics of spoken and signed word production: A case study using electrocorticographic spectral analysis. *Aphasiology*, 16(9), 903–927. <https://doi.org/10.1080/02687030244000383>
- Dale, A. M., Fischl, B., & Sereno, M. I. (1999). Cortical surface-based analysis: I. Segmentation and surface reconstruction. *NeuroImage*, 9(2), 179–194. <https://doi.org/10.1006/nimg.1998.0395>
- Delorme, A. (2023). EEG is better left alone. *Scientific Reports*, 13(1), 2372. <https://doi.org/10.1038/s41598-023-27528-0>

- 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. <https://doi.org/10.1016/j.jneumeth.2003.10.009>
- Devlin, J. T., Russell, R. P., Davis, M. H., Price, C. J., Wilson, J., Moss, H. E., Matthews, P. M., & Tyler, L. K. (2000). Susceptibility-induced loss of signal: Comparing PET and fMRI on a semantic task. *NeuroImage*, 11(6), 589–600. <https://doi.org/10.1006/nimg.2000.0595>
- Dilkina, K., & Lambon Ralph, M. A. (2012). Conceptual structure within and between modalities. *Frontiers in Human Neuroscience*, 6. <https://doi.org/10.3389/fnhum.2012.00333>
- Drane, D. L., Ojemann, G. A., Aylward, E., Ojemann, J. G., Johnson, L. C., Silbergeld, D. L., Miller, J. W., & Tranel, D. (2008). Category-specific naming and recognition deficits in temporal lobe epilepsy surgical patients. *Neuropsychologia*, 46(5), 1242–1255. <https://doi.org/10.1016/j.neuropsychologia.2007.11.034>
- 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(1), 291–301. <https://doi.org/10.1016/j.neuroimage.2009.12.035>
- Fischl, B., Sereno, M. I., & Dale, A. M. (1999). Cortical surface-based analysis: II. Inflation, flattening, and a surface-based coordinate system. *NeuroImage*, 9(2), 195–207. <https://doi.org/10.1006/nimg.1998.0396>
- Forseth, K. J., Kadipasaoglu, C. M., Conner, C. R., Hickok, G., Knight, R. T., & Tandon, N. (2018). A lexical semantic hub for heteromodal naming in middle fusiform gyrus. *Brain*, 141(7), 2112–2126. <https://doi.org/10.1093/brain/awy120>
- Friedman, J., Hastie, T., & Tibshirani, R. (2010). Regularization paths for generalized linear models via coordinate descent. *Journal of Statistical Software*, 33(1). <https://doi.org/10.18637/jss.v033.i01>
- Frisby, S. L., Halai, A. D., Cox, C. R., Lambon Ralph, M. A., & Rogers, T. T. (2023). Decoding semantic representations in mind and brain. *Trends in Cognitive Sciences*, 27(3), 258–281. <https://doi.org/10.1016/j.tics.2022.12.006>
- Halai, A. D., Welbourne, S. R., Embleton, K., & Parkes, L. M. (2014). A comparison of dual gradient-echo and spin-echo fMRI of the inferior temporal lobe. *Human Brain Mapping*, 35(8), 4118–4128. <https://doi.org/10.1002/hbm.22463>
- Halgren, E., Kaestner, E., Marinkovic, K., Cash, S. S., Wang, C., Schomer, D. L., Madsen, J. R., & Ulbert, I. (2015). Laminar profile of spontaneous and evoked theta: Rhythmic modulation of cortical processing during word integration. *Neuropsychologia*, 76, 108–124. <https://doi.org/10.1016/j.neuropsychologia.2015.03.021>
- Halgren, E., Wang, C., Schomer, D. L., Knake, S., Marinkovic, K., Wu, J., & Ulbert, I. (2006). Processing stages underlying word recognition in the anteroventral temporal lobe. *NeuroImage*, 30(4), 1401–1413. <https://doi.org/10.1016/j.neuroimage.2005.10.053>
- Hamberger, M. J. (2007). Cortical language mapping in epilepsy: A critical review. *Neuropsychology Review*, 17(4), 477–489. <https://doi.org/10.1007/s11065-007-9046-6>

- Henry, T. R., Buchtel, H. A., Koepp, R. A., Pennell, P. B., Kluin, K. J., & Minoshima, S. (1998). Absence of normal activation of the left anterior fusiform gyrus during naming in left temporal lobe epilepsy. *Neurology*, 50(3), 787–790. <https://doi.org/10.1212/WNL.50.3.787>
- Hermes, D., Miller, K. J., Vansteensel, M. J., Edwards, E., Ferrier, C. H., Bleichner, M. G., van Rijen, P. C., Aarnoutse, E. J., & Ramsey, N. F. (2014). Cortical theta wanes for language. *NeuroImage*, 85, 738–748. <https://doi.org/10.1016/j.neuroimage.2013.07.029>
- Hodges, J. R., & Patterson, K. (2007). Semantic dementia: A unique clinicopathological syndrome. *The Lancet Neurology*, 6(11), 1004–1014. [https://doi.org/10.1016/S1474-4422\(07\)70266-1](https://doi.org/10.1016/S1474-4422(07)70266-1)
- Jackson, R. L. (2021). The neural correlates of semantic control revisited. *NeuroImage*, 224, 117444. <https://doi.org/10.1016/j.neuroimage.2020.117444>
- Jenkinson, M., Beckmann, C. F., Behrens, T. E. J., Woolrich, M. W., & Smith, S. M. (2012). FSL. *NeuroImage*, 62(2), 782–790. <https://doi.org/10.1016/j.neuroimage.2011.09.015>
- Jerbi, K., Freyermuth, S., Dalal, S., Kahane, P., Bertrand, O., Berthoz, A., & Lachaux, J.-P. (2009). Saccade related gamma-band activity in intracerebral EEG: Dissociating neural from ocular muscle activity. *Brain Topography*, 22(1), 18–23. <https://doi.org/10.1007/s10548-009-0078-5>
- Jones, M. N., Willits, J., & Dennis, S. (2015). In J. R. Busemeyer, Z. Wang, J. T. Townsend, & A. Eidels (Eds.), *Models of Semantic Memory*. Oxford University Press. <https://doi.org/10.1093/oxfordhb/9780199957996.013.11>
- King, J.-R., & Dehaene, S. (2014). Characterizing the dynamics of mental representations: The temporal generalization method. *Trends in Cognitive Sciences*, 18(4), 203–210. <https://doi.org/10.1016/j.tics.2014.01.002>
- Klimesch, W. (2012). Alpha-band oscillations, attention, and controlled access to stored information. *Trends in Cognitive Sciences*, 16(12), 606–617. <https://doi.org/10.1016/j.tics.2012.10.007>
- Kojima, K., Brown, E. C., Matsuzaki, N., Rothermel, R., Fuerst, D., Shah, A., Mittal, S., Sood, S., & Asano, E. (2013). Gamma activity modulated by picture and auditory naming tasks: Intracranial recording in patients with focal epilepsy. *Clinical Neurophysiology*, 124(9), 1737–1744. <https://doi.org/10.1016/j.clinph.2013.01.030>
- Kovach, C. K., Tsuchiya, N., Kawasaki, H., Oya, H., Howard, M. A., & Adolphs, R. (2011). Manifestation of ocular-muscle EMG contamination in human intracranial recordings. *NeuroImage*, 54(1), 213–233. <https://doi.org/10.1016/j.neuroimage.2010.08.002>
- Lachaux, J.-P., Axmacher, N., Mormann, F., Halgren, E., & Crone, N. E. (2012). High-frequency neural activity and human cognition: Past, present and possible future of intracranial EEG research. *Progress in Neurobiology*, 98(3), 279–301. <https://doi.org/10.1016/j.pneurobio.2012.06.008>
- Lambon Ralph, M. A., Jefferies, E., Patterson, K., & Rogers, T. T. (2017). The neural and computational bases of semantic cognition. *Nature Reviews Neuroscience*, 18(1), 42–55. <https://doi.org/10.1038/nrn.2016.150>

- Lambon Ralph, M. A., Sage, K., Jones, R. W., & Mayberry, E. J. (2010). Coherent concepts are computed in the anterior temporal lobes. *Proceedings of the National Academy of Sciences*, 107(6), 2717–2722. <https://doi.org/10.1073/pnas.0907307107>
- Liu, H., Agam, Y., Madsen, J. R., & Kreiman, G. (2009). Timing, timing, timing: Fast decoding of object information from intracranial field potentials in human visual cortex. *Neuron*, 62(2), 281–290. <https://doi.org/10.1016/j.neuron.2009.02.025>
- Lüders, H., Lesser, R. P., Hahn, J., Dinner, D. S., Morris, H. H., Wylie, E., & Godoy, J. (1991). Basal temporal language area. *Brain*, 114(2), 743–754. <https://doi.org/10.1093/brain/114.2.743>
- Marko, M., Cimrová, B., & Riečanský, I. (2019). Neural theta oscillations support semantic memory retrieval. *Scientific Reports*, 9(1), 17667. <https://doi.org/10.1038/s41598-019-53813-y>
- Matoba, K., Matsumoto, R., Shimotake, A., Nakae, T., Imamura, H., Togo, M., Yamao, Y., Usami, K., Kikuchi, T., Yoshida, K., Matsushashi, M., Kunieda, T., Miyamoto, S., Takahashi, R., & Ikeda, A. (2024). Basal temporal language area revisited in Japanese language with a language function density map. *Cerebral Cortex*, 34(6), bhae218. <https://doi.org/10.1093/cercor/bhae218>
- Merker, B. (2013). Cortical gamma oscillations: The functional key is activation, not cognition. *Neuroscience & Biobehavioral Reviews*, 37(3), 401–417. <https://doi.org/10.1016/j.neubiorev.2013.01.013>
- Mitra, P., & Bokil, H. (2007). *Observed Brain Dynamics*. Oxford University Press. <https://doi.org/10.1093/acprof:oso/9780195178081.001.0001>
- Mollo, G., Cornelissen, P. L., Millman, R. E., Ellis, A. W., & Jefferies, E. (2017). Oscillatory dynamics supporting semantic cognition: MEG evidence for the contribution of the anterior temporal lobe hub and modality-specific spokes. *PLOS ONE*, 12(1), e0169269. <https://doi.org/10.1371/journal.pone.0169269>
- Morrison, C. M., Chappell, T. D., & Ellis, A. W. (1997). Age of acquisition norms for a large set of object names and their relation to adult estimates and other variables. *The Quarterly Journal of Experimental Psychology Section A*, 50(3), 528–559. <https://doi.org/10.1080/027249897392017>
- Nagata, K., Kunii, N., Shimada, S., Fujitani, S., Takasago, M., & Saito, N. (2022). Spatiotemporal target selection for intracranial neural decoding of abstract and concrete semantics. *Cerebral Cortex*, 32(24), 5544–5554. <https://doi.org/10.1093/cercor/bhac034>
- Nakai, Y., Jeong, J., Brown, E. C., Rothermel, R., Kojima, K., Kambara, T., Shah, A., Mittal, S., Sood, S., & Asano, E. (2017). Three- and four-dimensional mapping of speech and language in patients with epilepsy. *Brain*, 140(5), 1351–1370. <https://doi.org/10.1093/brain/awx051>
- Nakai, Y., Sugiura, A., Brown, E. C., Sonoda, M., Jeong, J., Rothermel, R., Luat, A. F., Sood, S., & Asano, E. (2019). Four-dimensional functional cortical maps of visual and auditory language: Intracranial recording. *Epilepsia*, 60(2), 255–267. <https://doi.org/10.1111/epi.14648>
- Nir, Y., Fisch, L., Mukamel, R., Gelbard-Sagiv, H., Arieli, A., Fried, I., & Malach, R. (2007). Coupling between Neuronal Firing Rate, Gamma LFP, and BOLD fMRI Is Related to

- Interneuronal Correlations. *Current Biology*, 17(15), 1275–1285.
<https://doi.org/10.1016/j.cub.2007.06.066>
- Nobre, A., & McCarthy, G. (1995). Language-related field potentials in the anterior-medial temporal lobe: II. Effects of word type and semantic priming. *The Journal of Neuroscience*, 15(2), 1090–1098. <https://doi.org/10.1523/JNEUROSCI.15-02-01090.1995>
- Parvizi, J., & Kastner, S. (2018). Promises and limitations of human intracranial electroencephalography. *Nature Neuroscience*, 21(4), 474–483.
<https://doi.org/10.1038/s41593-018-0108-2>
- Patterson, K., Nestor, P. J., & Rogers, T. T. (2007). Where do you know what you know? The representation of semantic knowledge in the human brain. *Nature Reviews Neuroscience*, 8(12), 976–987. <https://doi.org/10.1038/nrn2277>
- Pobric, G., Jefferies, E., & Lambon Ralph, M. A. (2007). Anterior temporal lobes mediate semantic representation: Mimicking semantic dementia by using rTMS in normal participants. *Proceedings of the National Academy of Sciences*, 104(50), 20137–20141. <https://doi.org/10.1073/pnas.0707383104>
- Pobric, G., Jefferies, E., & Lambon Ralph, M. A. (2010a). Amodal semantic representations depend on both anterior temporal lobes: Evidence from repetitive transcranial magnetic stimulation. *Neuropsychologia*, 48(5), 1336–1342.
<https://doi.org/10.1016/j.neuropsychologia.2009.12.036>
- Pobric, G., Jefferies, E., & Lambon Ralph, M. A. (2010b). Category-specific versus category-general semantic impairment induced by transcranial magnetic stimulation. *Current Biology*, 20(10), 964–968. <https://doi.org/10.1016/j.cub.2010.03.070>
- Rogers, T. T., Cox, C. R., Lu, Q., Shimotake, A., Kikuchi, T., Kunieda, T., Miyamoto, S., Takahashi, R., Ikeda, A., Matsumoto, R., & Lambon Ralph, M. A. (2021). Evidence for a deep, distributed and dynamic code for animacy in human ventral anterior temporal cortex. *eLife*, 10, e66276. <https://doi.org/10.7554/eLife.66276>
- Rogers, T. T., Lambon Ralph, M. A., Garrard, P., Bozeat, S., McClelland, J. L., Hodges, J. R., & Patterson, K. (2004). Structure and deterioration of semantic memory: A neuropsychological and computational investigation. *Psychological Review*, 111(1), 205–235. <https://doi.org/10.1037/0033-295X.111.1.205>
- Rupp, K., Roos, M., Milsap, G., Caceres, C., Ratto, C., Chevillet, M., Crone, N. E., & Wolmetz, M. (2017). Semantic attributes are encoded in human electrocorticographic signals during visual object recognition. *NeuroImage*, 148, 318–329.
<https://doi.org/10.1016/j.neuroimage.2016.12.074>
- Sato, N., Matsumoto, R., Shimotake, A., Matsushashi, M., Otani, M., Kikuchi, T., Kunieda, T., Mizuhara, H., Miyamoto, S., Takahashi, R., & Ikeda, A. (2021). Frequency-dependent cortical interactions during semantic processing: An electrocorticogram cross-spectrum analysis using a semantic space model. *Cerebral Cortex*, 31(9), 4329–4339.
<https://doi.org/10.1093/cercor/bhab089>
- Sauseng, P., & Klimesch, W. (2008). What does phase information of oscillatory brain activity tell us about cognitive processes? *Neuroscience & Biobehavioral Reviews*, 32(5), 1001–1013. <https://doi.org/10.1016/j.neubiorev.2008.03.014>

- Shimotake, A., Matsumoto, R., Ueno, T., Kunieda, T., Saito, S., Hoffman, P., Kikuchi, T., Fukuyama, H., Miyamoto, S., Takahashi, R., Ikeda, A., & Lambon Ralph, M. A. (2015). Direct exploration of the role of the ventral anterior temporal lobe in semantic memory: Cortical stimulation and local field potential evidence from subdural grid electrodes. *Cerebral Cortex*, 25(10), 3802–3817. <https://doi.org/10.1093/cercor/bhu262>
- Smith, S. M., Jenkinson, M., Woolrich, M. W., Beckmann, C. F., Behrens, T. E. J., Johansen-Berg, H., Bannister, P. R., De Luca, M., Drobnjak, I., Flitney, D. E., Niazy, R. K., Saunders, J., Vickers, J., Zhang, Y., De Stefano, N., Brady, J. M., & Matthews, P. M. (2004). Advances in functional and structural MR image analysis and implementation as FSL. *NeuroImage*, 23, S208–S219. <https://doi.org/10.1016/j.neuroimage.2004.07.051>
- Snyder, K. M., Forseth, K. J., Donos, C., Rollo, P. S., Fischer-Baum, S., Breier, J., & Tandon, N. (2023). Critical role of the ventral temporal lobe in naming. *Epilepsia*, 64(5), 1200–1213. <https://doi.org/10.1111/epi.17555>
- Solomon, E. A., Lega, B. C., Sperling, M. R., & Kahana, M. J. (2019). *Hippocampal theta codes for distances in semantic and temporal spaces*. 10.
- Spiers, H. J. (2020). The hippocampal cognitive map: One space or many? *Trends in Cognitive Sciences*, 24(3), 168–170. <https://doi.org/10.1016/j.tics.2019.12.013>
- Stryker, M. P. (1989). Is grandmother an oscillation? *Nature*, 338(6213), 297–298. <https://doi.org/10.1038/338297a0>
- Tallon-Baudry, C., & Bertrand, O. (1999). Oscillatory gamma activity in humans and its role in object representation. *Trends in Cognitive Sciences*, 3(4), 151–162. [https://doi.org/10.1016/S1364-6613\(99\)01299-1](https://doi.org/10.1016/S1364-6613(99)01299-1)
- Tanji, K. (2005). High-frequency γ -band activity in the basal temporal cortex during picture-naming and lexical-decision tasks. *Journal of Neuroscience*, 25(13), 3287–3293. <https://doi.org/10.1523/JNEUROSCI.4948-04.2005>
- Tibshirani, R. (1996). Regression shrinkage and selection via the lasso. *Journal of the Royal Statistical Society: Series B (Methodological)*, 58(1), 267–288. <https://doi.org/10.1111/j.2517-6161.1996.tb02080.x>
- Visser, M., Jefferies, E., & Lambon Ralph, M. A. (2010). Semantic processing in the anterior temporal lobes: A meta-analysis of the functional neuroimaging literature. *Journal of Cognitive Neuroscience*, 22(6), 1083–1094. <https://doi.org/10.1162/jocn.2009.21309>
- von der Malsburg, C. (1995). Binding in models of perception and brain function. *Current Opinion in Neurobiology*, 5(4), 520–526. [https://doi.org/10.1016/0959-4388\(95\)80014-X](https://doi.org/10.1016/0959-4388(95)80014-X)
- Wang, W., Degenhart, A. D., Sudre, G. P., Pomerleau, D. A., & Tyler-Kabara, E. C. (2011). Decoding semantic information from human electrocorticographic (ECoG) signals. *Annual International Conference of the IEEE Engineering in Medicine and Biology Society. IEEE Engineering in Medicine and Biology Society. Annual International Conference, 2011*, 6294–6298. <https://doi.org/10.1109/IEMBS.2011.6091553>
- Watrous, A. J., Fell, J., Ekstrom, A. D., & Axmacher, N. (2015). More than spikes: Common oscillatory mechanisms for content specific neural representations during perception

1096 and memory. *Current Opinion in Neurobiology*, 31, 33–39.
1097 <https://doi.org/10.1016/j.conb.2014.07.024>
1098 Yuste, R. (2015). From the neuron doctrine to neural networks. *Nature Reviews*
1099 *Neuroscience*, 16(8), 487–497. <https://doi.org/10.1038/nrn3962>
1100 Yuval-Greenberg, S., Tomer, O., Keren, A. S., Nelken, I., & Deouell, L. Y. (2008). Transient
1101 induced gamma-band response in eeg as a manifestation of miniature saccades.
1102 *Neuron*, 58(3), 429–441. <https://doi.org/10.1016/j.neuron.2008.03.027>
1103