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Four-dimensional functional cortical maps of visual and auditory language: intracranial recording

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SUMMARY

Objective: The strength of presurgical language mapping using electrocorticography (ECoG) is its outstanding signal fidelity and temporal resolution, but the weakness includes limited spatial sampling at an individual patient level. By averaging naming-related high-gamma activity at non-epileptic regions across a large number of patients, we provided the functional cortical atlases animating the neural dynamics supporting visual-object and auditory-description naming at the whole brain level.

Methods: We studied 79 patients who underwent extraoperative ECoG recording as epilepsy presurgical evaluation, and generated time-frequency plots and animation videos delineating the dynamics of naming-related high-gamma activity at 70–110 Hz.

Results: Naming task performance elicited high-gamma augmentation in domain-specific lower-order sensory areas and inferior-precentral gyri immediately after stimulus onset. High-gamma augmentation subsequently involved widespread neocortical networks with left hemispheric

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DISCLOSURE

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dominance. Left posterior temporal high-gamma augmentation at several hundred milliseconds before response onset exhibited a double dissociation; picture naming elicited high-gamma augmentation preferentially in regions medial to the inferior-temporal gyrus, whereas auditory naming elicited high-gamma augmentation more laterally. The left lateral prefrontal regions including Broca's area initially exhibited high-gamma suppression subsequently followed by high-gamma augmentation at several hundred milliseconds before response onset during both naming tasks. Early high-gamma suppression within Broca's area was more intense during picture compared to auditory naming. Subsequent lateral-prefrontal high-gamma augmentation was more intense during auditory compared to picture naming.

Significance: This study revealed contrasting characteristics in the spatiotemporal dynamics of naming-related neural modulations between tasks. The dynamic atlases of visual and auditory language might be useful for planning of epilepsy surgery. Differential neural activation well explains some of the previously reported observations of domain-specific language impairments following resective epilepsy surgery. Video materials might be beneficial for the education of lay people about how the brain functions differentially during visual and auditory naming.

Keywords

Physiological high-frequency oscillations (HFOs); Four dimension (4D); Intracranial electroencephalography (EEG) recording; Speech; Event-related high-gamma modulation

INTRODUCTION

You might say 'Cat', 'Kitty' or 'Kitten', if you witness one walking in the street. You might answer 'Bird' or 'Plane', if you are verbally asked: 'What flies in the sky?' Word finding abilities in visual or auditory domain are crucial for humans. Visual object naming (referred to as 'picture naming' below) and auditory description naming ('auditory naming') are commonly used for evaluation of the baseline language competence as well as for localization of language-related eloquent areas in patients scheduled to undergo epilepsy surgery¹. Transient language impairment elicited by electrical stimulation and task-related neural activation on electrocorticography (ECoG) or functional MRI (fMRI) are useful biomarkers to localize eloquent areas in clinical practice²⁻⁴. The strength of ECoG- over fMRI-based language mapping is its outstanding signal fidelity and temporal resolution⁵. Augmentation of high-gamma ECoG activity is an excellent summary measure of cortical activation, since it is tightly associated with increased neuronal firing rate⁶, hemodynamic activation⁷, glucose metabolism⁸, probability of stimulation-induced language impairment⁹, and risk of post-resection language impairment^{10,11}. In contrast, it remains less certain if task-related attenuation of alpha/beta activities is linearly correlated to the underlying cortical activation^{12,13}. Though ECoG can sample only selected regions for each patient, averaging of task-related high-gamma activity_{70-110 Hz} at non-epileptic regions across a large number of patients has effectively yielded a movie animating the dynamics of auditory naming-related neural modulation at the whole-brain level¹². In this study of 79 patients with focal epilepsy (Table 1), we evaluated the spatiotemporal dynamics of picture and auditory naming-related high-gamma activity at 7,348 electrode sites across 66 regions of interest (ROIs). We believe the resulting dynamic atlases of visual and auditory language

would be useful for planning of epilepsy surgery. Previous behavioral studies of patients undergoing epilepsy surgery indicated that focal resection of a region outside the lower-order sensory areas (i.e.: primary visual/auditory areas in the present study) may impair naming function of a certain domain but preserve others¹⁴. Therefore, we hypothesized that domain-specific naming-related high-gamma modulation would be evident not only in the primary visual/auditory areas but also in association areas. We expected that this study would clarify the *timing* of given naming-related neural modulations including *activation* and *suppression* at an order of tens of milliseconds, whereas our previous studies of smaller samples focused on neural activation alone at limited numbers of ROIs^{11,15}. Evaluation of task-related neural suppression is important as well, based on the theory of antagonism suggesting that effective task completion is facilitated by transient suppression of undesirable cortical function at given moments of performance^{16,17}.

METHODS

Patients

The inclusion criteria consisted of: (i) extraoperative ECoG recording at Children's Hospital of Michigan or Harper University Hospital in Detroit between January 2008 and July 2016, (ii) age of four years and above, and (iii) brain mapping with measurement of naming-related high-gamma activity in visual and auditory domains (Figure 1). The exclusion criteria consisted of: (i) brain malformations confounding the anatomical landmarks for the central, calcarine, or lateral sulcus, (ii) severe cognitive dysfunction (anticipation of inability to complete naming tasks or verbal IQ<70 when available), (iii) inability to complete either naming task, (iv) primary spoken language other than English, (v) previous epilepsy surgery, (vi) visual field deficit on confrontation, and (vii) right-hemispheric language dominance as suggested by either Wada test or left-handedness associated with left-hemispheric congenital neocortical lesions¹¹. We determined handedness based on the observation of preferred hand for writing tasks. No formal testing of handedness (e.g. Edinburgh handedness inventory) was performed in this study. The present study was approved by the Institutional Review Board at Wayne State University, and written informed consent was obtained from the patients or guardians of patients.

Acquisition of ECoG and 3D surface MR images

The detailed methods were previously described^{12,18}. Chronically-implanted subdural electrodes (10 mm center-to-center distance) recorded ECoG signals at a sampling frequency of 1,000 Hz. Preoperative 3T MRI, including fluid-attenuated inversion recovery (FLAIR) images, was visually assessed by an experienced pediatric neuroradiologist who was blinded to scalp video-EEG data. Electrode sites classified as seizure onset zone (SOZ) or those affected by any structural lesion were excluded from further analysis. Those showing interictal spikes or artifacts during either task were also excluded from analysis. In general, we do not treat sharply contoured waves as interictal epileptiform discharges, if they occur only rarely during the entire recording or they can be attributed to some forms of physiological waves such as background fluctuations, lambda waves, mu rhythm, and sleep spindles^{19,20}. Thus, a total of 7,348 electrode sites were available for further analysis.

Using preoperative volumetric MR image, a 3D surface image was created with the location of electrodes defined on the surface¹². The spatial normalization of electrode sites was performed using FreeSurfer scripts (<http://surfer.nmr.mgh.harvard.edu>). All electrode sites were registered on the averaged FreeSurfer surface image. Each recording site was assigned an anatomical label based on the Desikan-Killiany atlas²¹ (Figure 2). The number of electrode sites eligible for analysis is indicated in Table 2. The statistical power to determine the significance in the degree of high-gamma modulation was dependent on the number of electrode sites at each ROI. For example, the number of electrode sites was 345 and 39 in the left inferior precentral and posterior cingulate regions, respectively; thus, the former ROI had 3.0 times greater statistical power than the latter. Lack of significant high-gamma modulation should be interpreted as failure to demonstrate significant modulation at ROIs where small numbers of electrode sites were available.

Picture and auditory naming tasks

During the picture naming task¹⁵, patients were instructed to overtly name an object presented on an LCD monitor (Experiment Center®, SensoMotoric Instruments, Boston, MA). Stimuli consisted of common gray-scale line-drawn objects (up to 60 objects; such as ‘cat’ and ‘chair’)²². The timings of stimulus presentations, patient behaviors, and ECoG signals were synchronized using a photosensor and microphone¹⁸. ECoG traces were aligned to ‘stimulus onset’ and ‘response onset’ (Figure 1A). Trials not accompanied by overt correct answers were excluded from analysis. For example, when patients were shown a picture of skunk, incorrect answers would include: ‘*I don’t know*’, ‘*What is this?*’, or a bad smell gesture. Conversely, correct answers would include ‘*Skunk*’, ‘*Squirrel*’, and so on. A gray-scale picture of skunk could be perceived as a squirrel for some individuals. Nouns relevant based on each patient’s criteria were treated as correct answers.

During the auditory naming task¹², patients were instructed to overtly verbalize an answer for a series of auditory sentence questions (such as ‘*What flies in the sky?*’). Up to 100 questions were presented with the Presentation® (Neurobehavioral Systems, Albany, CA). ECoG traces were then aligned to ‘stimulus onset’, ‘stimulus offset’, and ‘response onset’ (Figure 1B). Likewise, trials not accompanied by overt, correct, noun answers were excluded from analysis.

Time-frequency ECoG analysis

We determined the spatiotemporal dynamics of naming-related high-gamma modulation using the method identical to that previously reported¹². ECoG time-voltage signals were transformed into the time-frequency domain, in steps of 10 ms and 5 Hz, using a complex-demodulation time-frequency transformation^{23,24}. At each 10-ms epoch at each recording site, we measured the percent change in high-gamma_{70–110} Hz amplitude (a measure proportional to the square root of power) relative to the mean amplitude during a resting period between 600 to 200 ms prior to stimulus onset (Figure 1). Time-frequency analysis was repeated with a time-lock to ‘stimulus onset’ and ‘response onset’ for picture naming, whereas ‘stimulus onset’, ‘stimulus offset’, and ‘response onset’ for auditory naming. We plotted the mean and standard error (SE) of high-gamma amplitude change across all available channels at each ROI as a function of time. We determined the timing of

statistically-significant augmentation (or suppression) of high-gamma activity at each ROI in each hemisphere during each naming task, using studentized bootstrap statistics followed by correction for multiple comparisons across time windows¹². Specifically, if high-gamma amplitude was beyond or below the 99.99% confidence interval (i.e.: mean \pm 3.89 \times standard error) of that during the resting period, the change in high-gamma amplitude was treated as significant at a given ROI commonly in picture and auditory naming tasks. The aforementioned procedures effectively determined at what ROIs common and differential high-gamma modulations were noted during picture and auditory naming tasks. It should be noted that there was no statistical difference between ‘the mean across all electrode sites within each ROI’ and ‘the grand-mean across individual means within each ROI’, as shown in Supplementary Figure S1; therefore, the risk of a single patient’s data severely deviating the calculated mean from the true mean was small in the present study. This analytic approach benefited from a maintained statistical power.

High-gamma amplitude measures were spatially presented with a Gaussian half-width at half maximum of 7.5 mm, and sequentially animated on the average FreeSurfer pial surface image as a function of time. Grand-averaging of all available patients’ data finally yielded the maps showing the dynamics of picture and auditory naming-related high-gamma modulations at the whole-brain level (**Supplementary Videos S1 and S2**).

RESULTS

Behavioral results

We studied 79 patients who satisfied both inclusion and exclusion criteria (Table 1). The mean number of included trials was 55.3 per patient (SE: 0.7) during the picture naming task and 83.2 per patient (SE: 2.3) during the auditory naming task. Thus, time-frequency analysis of high-gamma activity during the auditory naming task benefited from an approximately 1.2 times greater signal-to-noise ratio, compared to that during picture naming, because the number of trials was larger during the auditory naming task. The mean response time was 1,481 ms (SE: 63 ms) during picture naming and 1,407 ms (SE: 65 ms) during auditory naming. Younger age was associated with a longer response time somewhat in picture naming ($p=0.07$ on linear regression analysis; regression coefficient: -15 [95%CI: -31 to $+1$]) and auditory naming ($p=0.02$; regression coefficient: -19 [95%CI: -35 to -3]).

High-gamma modulation during picture naming

Supplementary Video S1 best visualizes the dynamics of naming-related high-gamma modulations in both visual and auditory domains and at the whole brain level. **Figures 3 and 4** present the snapshots of picture and auditory naming-related high-gamma modulations. Sustaining high-gamma augmentation took place in the bilateral occipital lobes within 100 ms after stimulus onset (Figures 5A and 5B). High-gamma augmentation subsequently involved the bilateral ventral occipito-temporal pathways (Figures 5E and 5F) as well as inferior-precentral regions (Figure 5D) at 100–300 ms after stimulus onset. Conversely, high-gamma suppression took place in the left inferior-frontal gyrus including pars orbitalis, triangularis, and opercularis at 300–600 ms after stimulus onset (Figures 5H–5J). At 600 ms prior to response onset and after, high-gamma augmentation involved the pars opercularis of

the left inferior-frontal gyrus (Figure 5H). Prior to and during the response phase, high-gamma augmentation involved the bilateral posterior superior-frontal region as well as, pre- and post-central gyri (Figures 3D and 5D). High-gamma augmentation of the left superior-temporal gyrus was minimal prior to response onset and evident only at response onset and after (Figure 5C).

High-gamma modulation during auditory naming

Sustained high-gamma augmentation took place in the bilateral superior-temporal gyri within 100 ms after stimulus onset (Figure 5C). High-gamma augmentation subsequently involved the inferior-precentral regions at 100–200 ms after stimulus onset (Figure 5D). Conversely, high-gamma suppression took place in the pars orbitalis of the left inferior-frontal gyrus at 300–600 ms after stimulus onset (Figures 5H–5J). Between stimulus offset and response onset, high-gamma augmentation involved widespread regions of the left hemisphere, whereas high-gamma suppression involved portions of the right lateral prefrontal regions (Figures 4B and 4C). Prior to and during the response phase, high-gamma augmentation involved the bilateral posterior superior-frontal regions as well as, pre- and post-central gyri (Figures 4D and 5D). High-gamma augmentation of the left superior-temporal gyrus was minimal within 300 ms prior to response onset and evident at response onset and after (Figure 5C).

Spatial-temporal characteristics of domain-specific high-gamma modulation

Domain-specific high-gamma augmentation was most evident in the primary visual/auditory areas. Specifically, picture and auditory naming tasks elicited high-gamma augmentation in the bilateral occipital and superior-temporal regions within 100 ms, respectively (Figures 5A–5C). The posterior temporal association neocortex showed domain-specific high-gamma augmentation. Picture naming task resulted in a *medial-to-lateral* gradient in the magnitude of high-gamma augmentation at 200–800 ms prior to response onset, which was greater in the posterior-fusiform and smaller in the posterior middle-temporal region (Figures 5E–5G). Conversely, the auditory naming task resulted in a *lateral-to-medial* gradient in the magnitude of high-gamma augmentation at the same time window.

Picture naming, compared to auditory naming, was associated with a smaller high-gamma activity in the left lateral prefrontal regions including the inferior-frontal gyrus (Figures 5H–5K). The degree of high-gamma suppression at 328 electrode sites within the left inferior-frontal gyrus at 300–600 ms following stimulus onset was 2.2 times more intense during picture compared to auditory naming ($p=0.01$ on studentized bootstrapping test; -0.029 [i.e.: -2.9%] during picture naming vs -0.013 [-1.3%] during auditory naming; Figure 5H–5J). Conversely, the degree of high-gamma suppression at 304 right inferior-frontal electrode sites at the same time window was not different between tasks ($p=0.97$; -1.0% during picture naming vs -0.9% during auditory naming).

During both naming tasks, lateral prefrontal regions showed high-gamma augmentation with left dominance at 200–800 ms prior to response onset (Supplementary Video S1). Such left-hemispheric dominant high-gamma augmentation was more intense in amplitude and more extensive in space and time during auditory compared to picture naming. Auditory naming

showed left hemisphere-specific high-gamma augmentation during this time window at 12 ROIs involving many in the left lateral prefrontal regions as well as inferior-parietal and posterior middle-temporal regions (Figures 5F-5K; Supplementary Video S2). Conversely, picture naming showed left-hemisphere specific high-gamma augmentation in the pars opercularis of inferior-frontal gyrus alone. The degree of high-gamma augmentation at 328 left inferior-frontal electrode sites during this time window was much more intense in auditory compared to picture naming ($p < 0.001$; +10.6% during auditory naming vs +0.8% during picture naming). The degree of high-gamma suppression at 304 right inferior-frontal electrode sites during this time window was not different between tasks ($p = 0.28$; -0.2% during auditory naming vs -0.8% during auditory naming).

Spatial-temporal characteristics of domain-common high-gamma modulation

Commonly during picture and auditory naming tasks, the inferior-precentral regions showed high-gamma augmentation initiated at 100–300 ms following stimulus onset, and sustained this until the overt response (Figure 5D). Likewise, the inferior-postcentral regions showed high-gamma augmentation initiated at 500–900 ms prior to response onset (Supplementary Video S2), and the posterior superior-temporal gyri showed high-gamma augmentation immediately following response onset (Figure 5C).

DISCUSSION

Significance of domain-specific high-gamma modulation

Our novel ECoG-based dynamic atlases revealed contrasting characteristics in the spatiotemporal profiles of naming-related neural modulations between task domains at the whole brain level (Supplementary Videos S1 and S2). Picture naming elicited early high-gamma augmentation in the occipital lobes, whereas auditory naming elicited early high-gamma augmentation in the superior-temporal gyrus over both hemispheres. High-gamma augmentation in these regions likely reflects, at least in part, lower-order perceptual processing in the visual²⁵ or auditory domain^{1,26}. Previous behavioral studies suggested that stroke in the left occipital lobe can cause difficulty in reading words or sentences partly because of the visual field deficit with letter naming ability being reasonably well maintained²⁷. The left posterior superior-temporal gyrus has been suggested to be involved in both lower-order acoustic processing as well as critical phonological processing ultimately necessary for reproduction of speech sounds²⁸. Our previous study demonstrated that stimulation at the left superior-temporal gyrus resulted in either auditory hallucination or naming error¹².

The left posterior temporal lobe neocortex showed a differential spatial gradient in the degree of high-gamma augmentation between task domains (Figures 5E-5G). More medial structures including the fusiform gyrus showed high-gamma augmentation relatively greater during picture naming. Conversely, more lateral structures including the posterior middle-temporal gyrus showed high-gamma augmentation relatively greater during auditory naming. These observations are consistent with previous electrical stimulation, fMRI, and ECoG studies^{10,29,30}. The observed spatial gradient provides a potential explanation for postoperative decline in visual more than auditory naming skills, as reported in a systematic

review of 21 behavioral studies of patients undergoing left anterior-medial temporal lobe resection¹⁴. It is plausible to speculate that extensive resection of the medial temporal lobe structures, as commonly performed in left anterior-medial temporal lobe resection³¹, may be associated with collateral damage of the neural pathways preferentially supporting picture naming function.

Contrasting spatiotemporal characteristics of naming-related high-gamma augmentation noted in the left posterior temporal neocortex between tasks support the notion that the exact boundary of Wernicke's region may be difficult to define universally using the anatomical landmarks alone and that the definition has largely varied across investigators³². While the left posterior superior-temporal gyrus, among perisylvian structures, was most commonly treated as a part of Wernicke's region in previous reports^{32,33}, picture naming-related high-gamma augmentation in the left superior-temporal gyrus was minimal prior to response onset but became significant during response. Commonly during picture and auditory naming tasks, high-gamma augmentation involved the left posterior inferior-temporal gyrus prior to response onset (Figure 5F). Thus, our ECoG study supports the previous fMRI and lesion-deficit studies reporting that the left posterior inferior-temporal gyrus supports semantic processing in both visual and auditory domains^{30,34}.

The left inferior-frontal gyrus showed early high-gamma suppression subsequently followed by high-gamma augmentation prior to the response phase during both naming tasks. Early high-gamma suppression was more intense (picture: -2.9% vs auditory: -1.3%) and subsequent augmentation less intense ($+0.8\%$ vs $+10.6\%$) in picture naming as compared to auditory naming. Our study is consistent with previous studies reporting that hemodynamic and neural activation in the left frontal lobe was greater during auditory naming compared to during picture naming^{10,30}. Collective evidence indicates that picture naming task can be completed with less activation of the left lateral prefrontal regions including the left inferior-frontal gyrus, known to include Broca's area. Less extensive activation of the left inferior-frontal gyrus during picture naming can be in part attributed to the experimental paradigms employed. Picture naming does not require syntactic processing as required by auditory naming, and the left inferior-frontal gyrus is suggested to support syntactic function^{35,36}. Recent ECoG studies showed that inferior-frontal high-gamma augmentation was fairly symmetric during an auditory verbal working memory task which requires neither syntactic nor semantic processing³⁷, but highly left-hemispheric dominant in an auditory descriptive naming task¹² such as the one employed herein. Other cognitive processes might also account for a greater extent of high-gamma augmentation in the left lateral prefrontal regions during auditory naming. A previous ECoG study reported that high-gamma augmentation within the left lateral prefrontal regions including the inferior-frontal gyrus was more intense during naming of ambiguous images compared to that of unambiguous common objects³⁸. The former naming task additionally requires 'estimation' and 'selection', whereas the latter naming task can be rapidly completed in an automatic manner³⁸.

The left inferior precentral gyrus commonly showed high-gamma augmentation within 100–300 ms following stimulus onset, earliest among all frontal lobe ROIs (Figure 5). At the same time window, high-gamma activity was rather suppressed in the left inferior-frontal

gyrus during picture and auditory naming tasks. These observations do not support the classic language model, as illustrated in the previous research^{32,39}, that internal word image generated in the left temporo-parietal neocortex is transferred initially and solely to the left inferior-frontal gyrus and subsequently to the left inferior precentral gyrus in a rostral-to-caudal direction. Based on the theory of antagonism^{16,17}, one can hypothesize that early suppression of the unnecessary function in the left inferior-frontal gyrus may facilitate effective completion of the naming task or that energy resources may be optimally distributed to the regions actively processing stimuli. Our recent ECoG study utilizing an auditory verbal working memory task suggested that the left inferior-precentral gyrus supports initial maintenance of a mental representation of memory items, whereas the left inferior-frontal gyrus contributes to later scanning function to determine a match among previously encountered items³⁷. Studies utilizing cortico-cortical evoked potentials and diffusion tensor tractography demonstrated that the left inferior-precentral gyrus is effectively connected structurally and functionally to both superior-temporal and inferior-frontal gyri^{40,41}.

Methodological considerations

We cannot rule out the possibility that the epileptogenic zone may have affected the spatiotemporal dynamics of naming-related high-gamma activity. Indeed, we previously reported that SOZ, compared to non-SOZ, has a smaller chance to generate significant naming-related high-gamma augmentation¹¹. Thus, we have excluded both SOZ and sites affected by a structural lesion from further analysis. Our observation of high-gamma augmentation spatially different across auditory and visual domains cannot be explained by the effect of SOZ localization, because ECoG signals between tasks were compared within the same patient cohort. Furthermore, interictal spike discharges *per se* are accompanied by a pathological increase in high-gamma power^{42,43}. Thus, the present study also excluded electrode sites showing interictal epileptiform discharges during the task, as it is a common practice in ECoG-based cognitive neuroscience research³. Our previous ECoG study of four patients with left frontal lobe epilepsy reported that slow waves immediately following interictal spikes may transiently inhibit naming-related cortical activity. However, this appeared to be restricted to regions involved in spike-and-wave discharge and did not appear to affect sites not involved in spike-and-wave discharges⁴⁴. Further analysis of a large number of patients at the whole brain level is necessary to better determine the transient effects of focal spike-and-wave discharges on the language network, taking into account that interictal focal spike discharges are suggested to have an impact on widespread networks beyond the regions from which spikes are generated⁴⁵.

Subdural disk electrodes used in the present study do not effectively sample ECoG signals from the cortex facing a sulcus. Further studies using depth electrodes may be warranted to determine if high-gamma augmentation in the superior-temporal gyri is attributed to activation of the immediately underlying cortex or that in the planum temporale.

Our patient cohort included those of age ranging from 5 to 44 years. Thus, great care needs to be taken in interpreting the results of time-frequency analysis. The spatial-temporal dynamics of averaged naming-related high-gamma activity, as presented in Supplementary

Video S1, likely reflect the neural dynamics exerted commonly by patients across a wide range of ages. In other words, neural dynamics exerted by a specific age group alone may have been underestimated. Development of the language systems is expected to be incomplete in younger children⁴⁶. Our previous ECoG studies of auditory naming-related neural dynamics reported that older age was associated with a greater degree of inferior-precentral high-gamma augmentation during stimulus listening as well as a greater degree of left rostral middle-frontal high-gamma augmentation immediately following stimulus offset^{12,47}. In the present study, younger patients had a longer response time compared to older ones. The effect of variance in response time across age is expected to be more prominent at the time windows long after stimulus onset as well as long before response onset. In order to distinguish high-gamma modulations related to perceptual and motor processing, we have provided the time-frequency plots time-locked to stimulus onset and response onset, respectively.

We employed a multivariate linear regression analysis to determine whether the response accuracy was independently associated with any of the age, age of epilepsy onset, history of generalized tonic-clonic seizures (GTCs), or each of the eight lobes affected by seizure onset zone (i.e.: left frontal, left temporal, left parietal, left occipital, right frontal, right temporal, right parietal, and right occipital lobes; Table 1). This *post-hoc* analysis failed to demonstrate the association between the response accuracy and any of the aforementioned 11 variables during either naming task (uncorrected $p > 0.05$).

Future directions

While the present study demonstrated that auditory naming, compared to picture naming, elicited more extensive high-gamma augmentation which involved the left lateral prefrontal and posterior cingulate regions, we still do not know whether auditory naming is more useful than picture naming for localization of the critical language areas or prediction of language outcome following cortical resection. Correlation between naming-related high-gamma activity and postoperative neuropsychological assessment in large populations will be ultimately necessary to address this question. A recent systematic review infers that the extent of cortex showing auditory or picture naming-related high-gamma augmentation was more extensive than the critical language area defined by electrical stimulation mapping in pediatric populations⁹. Conversely, resection of naming-related high-gamma sites can result in postoperative language impairment even though stimulation-defined critical language areas were preserved^{10,11}.

The strength of ECoG-based language mapping includes its more robust signal fidelity against overt speech-related artifacts and its capability to determine the detailed temporal dynamics of task-related neural modulations. Thus, inclusion of larger numbers of patients with a wide range of age in future collaborative works are expected to determine the effect of age on naming-related high-gamma modulations at given time windows at the whole brain level. We plan to test our hypothesis that efficient naming is supported by selective elaboration of new network connectivity throughout development. We are willing to share our dataset with investigators who express an interest in studying the neurobiology of language or the clinical utility of naming-related high-gamma activity.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Key Points:

- We provided the cortical atlases animating the neural dynamics supporting visual and auditory language function at the whole brain level.
- Visual and auditory naming tasks elicited early high-gamma augmentation in lower-order visual and auditory areas, respectively.
- Visual and auditory naming tasks commonly elicited early high-gamma augmentation in inferior-precentral gyri.
- Visual and auditory naming tasks elicited domain-specific high-gamma modulations in the left posterior temporal areas.
- Visual naming task, compared to auditory, elicited smaller high-gamma augmentation in the left prefrontal areas.

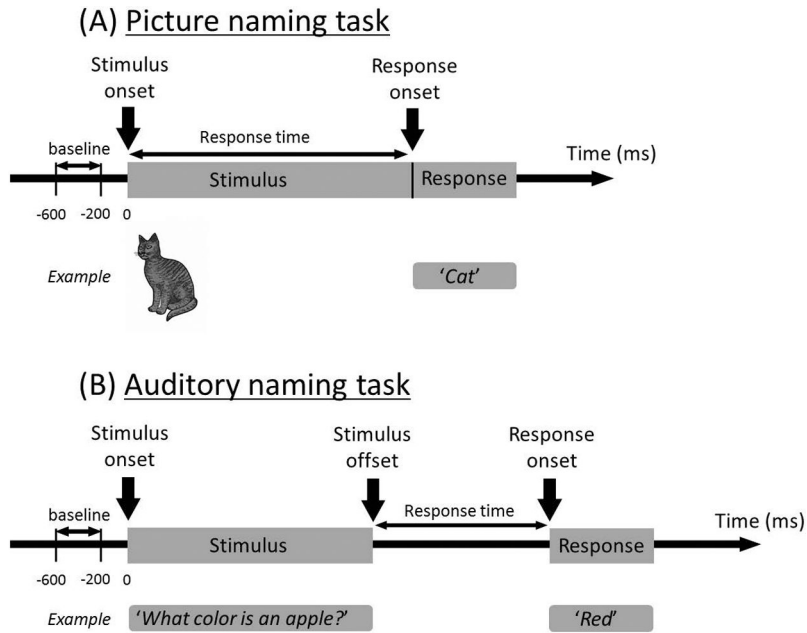


Figure 1. Picture and auditory naming tasks.

(A) Picture naming task: Patients were asked to look at and overtly name each object during extraoperative ECoG recording. At the end of each overt response, the examiner pressed the button to present the next stimulus following a cross in the center of the screen presented for 2 or 2.5 seconds. We then measured the percent change of high-gamma activity compared to that during the baseline period. The response time was defined as the period between stimulus onset and response onset. (B) Auditory naming task: Patients were asked to listen to a series of questions and to overtly verbalize a relevant answer for each question. The median duration across each question was 1,800 ms. At the end of each overt response, the examiner pressed the button to present the next stimulus following a silence of 2 or 2.5 seconds. The response time was defined as the period between stimulus offset and response onset.

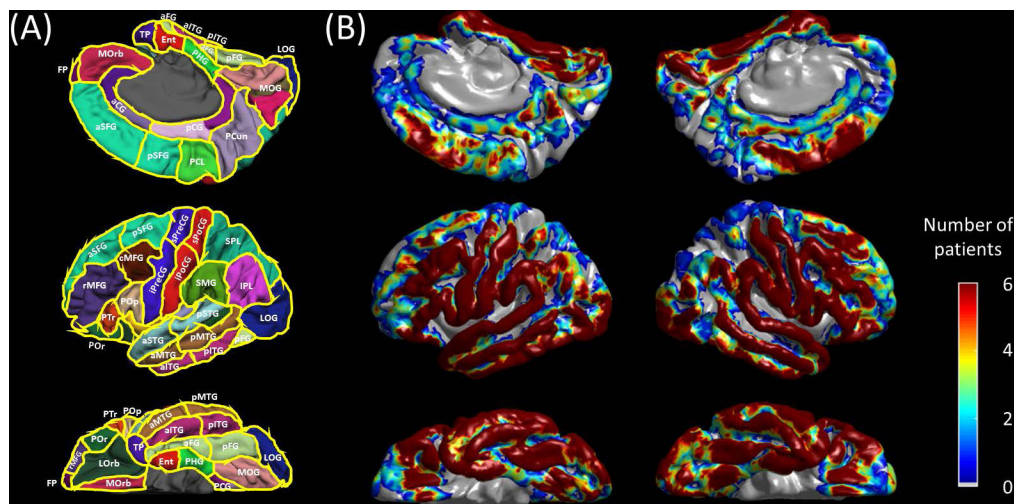


Figure 2. Regions of interest (ROIs) and distribution of subdural electrodes included in the analysis.

(A) aCG = anterior cingulate gyrus; aFG = anterior fusiform gyrus; aITG = anterior inferior temporal gyrus; aMTG = anterior middle temporal gyrus; aSFG = anterior superior frontal gyrus; aSTG = anterior superior temporal gyrus; cMFG = caudal middle frontal gyrus; Ent = entorhinal gyrus; FP = frontal pole; IPL = inferior parietal lobule; iPoCG = inferior postcentral gyrus; iPreCG = inferior precentral gyrus; LG = lingual gyrus; LOrb = lateral orbitofrontal gyrus; LOG = lateral occipital gyrus; MOG = medial occipital gyrus; MOrb = medial orbitofrontal gyrus; pCG = posterior cingulate gyrus; PCL = paracentral lobule; PCun = precuneus gyrus; pFG = posterior fusiform gyrus; PHG = parahippocampal gyrus; pITG = posterior inferior temporal gyrus; pMTG = posterior middle temporal gyrus; POp/PTr/POR = pars opercularis/pars triangularis/pars orbitalis within the inferior frontal gyrus. pSFG = posterior superior frontal gyrus; pSTG = posterior superior temporal gyrus; rMFG = rostral middle frontal gyrus; SMG = supramarginal gyrus; SPL = superior parietal lobule; sPoCG = superior postcentral gyrus; sPreCG = superior precentral gyrus; TP = temporal pole. (B) The numbers of patients whose ECoG data at a given site contributed to further analysis are presented.

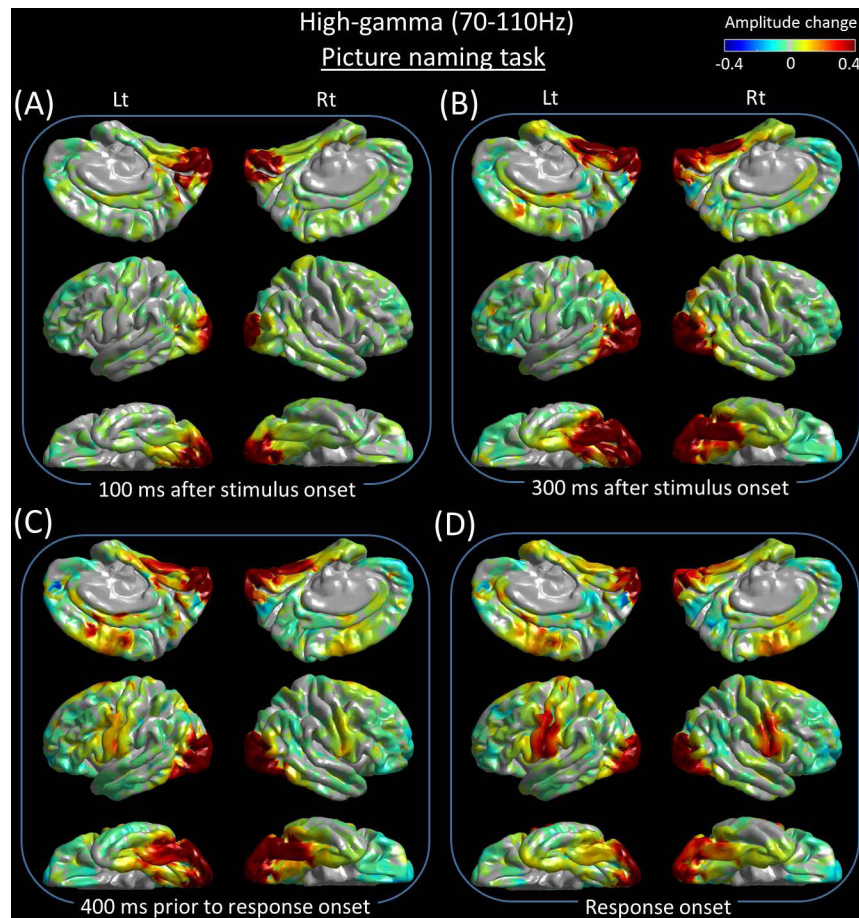


Figure 3. Snapshots of picture naming-related high-gamma activity.

Event-related amplitude augmentation is reflected by red, whereas suppression by blue. '+0.4' indicates that the amplitude was augmented by 40% compared to the mean during the resting period between -600 and -200 ms relative to stimulus onset. (A) 100 ms after stimulus onset. (B) 300 ms after stimulus onset. (C) 400 ms prior to response onset. (D) Response onset.

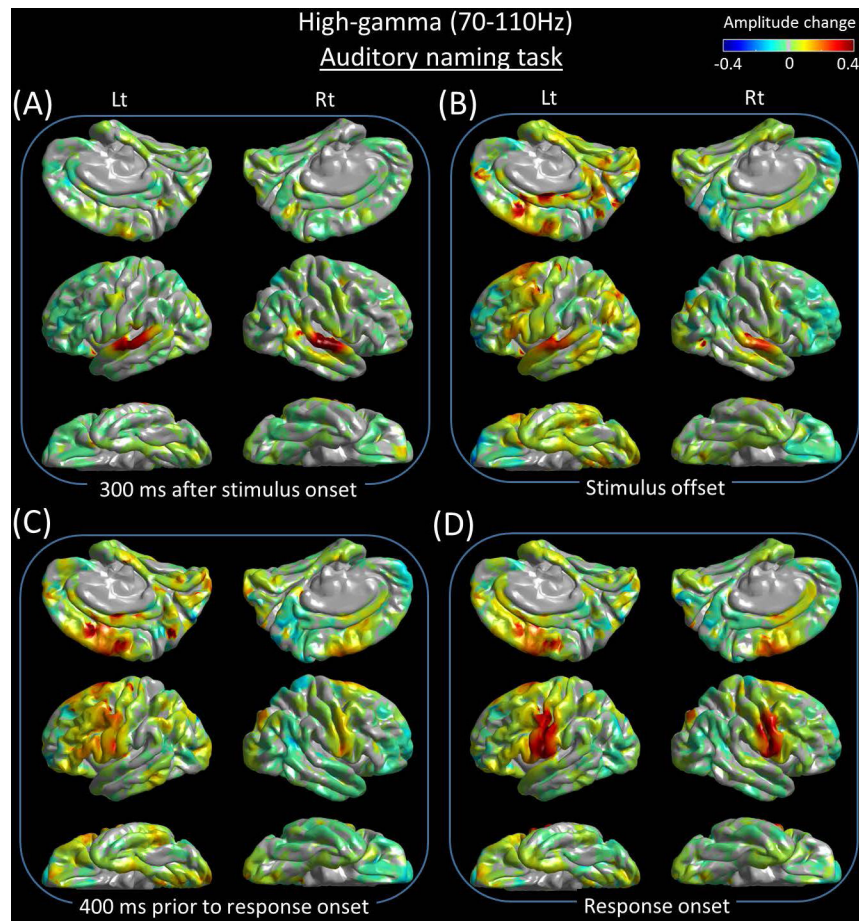
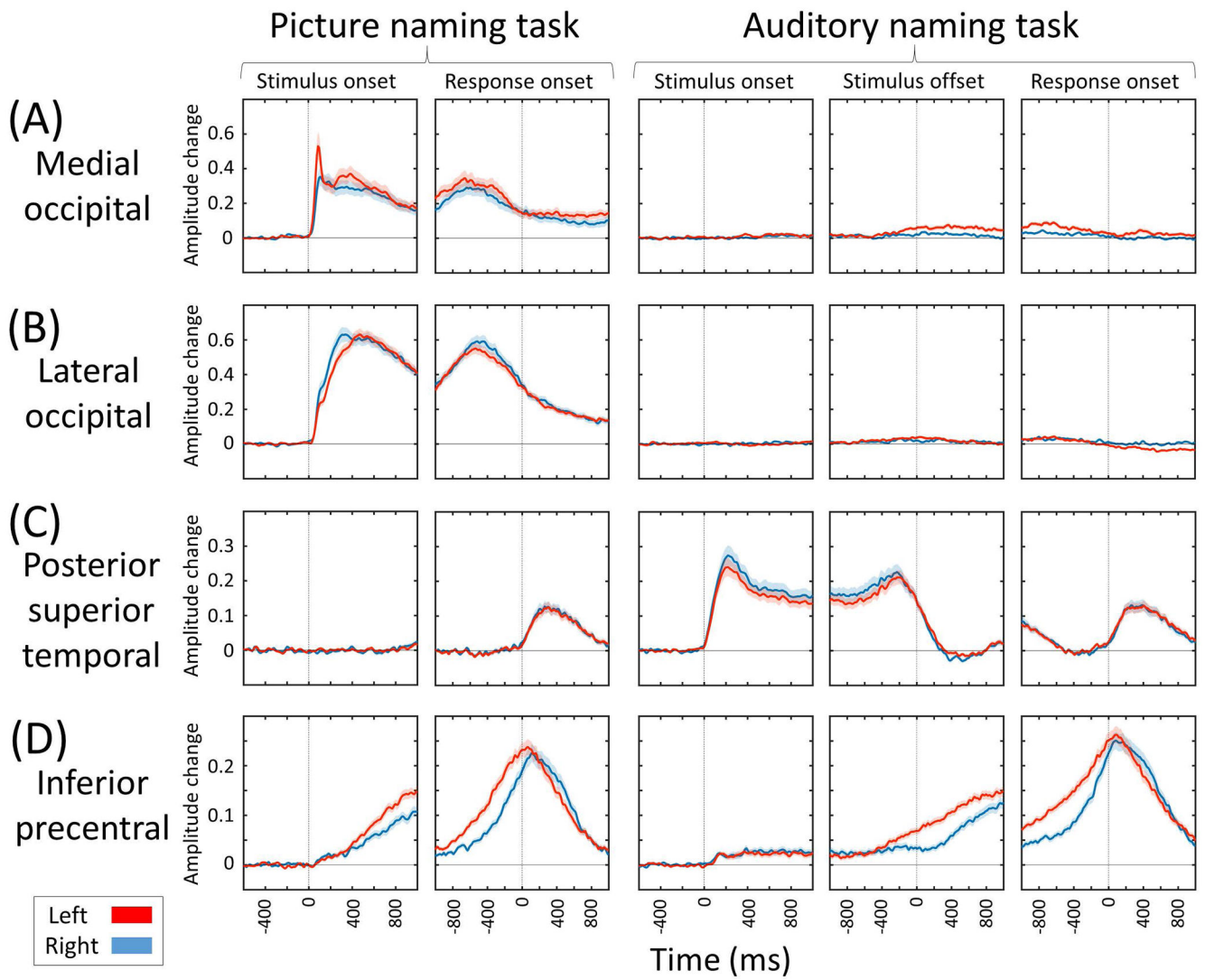
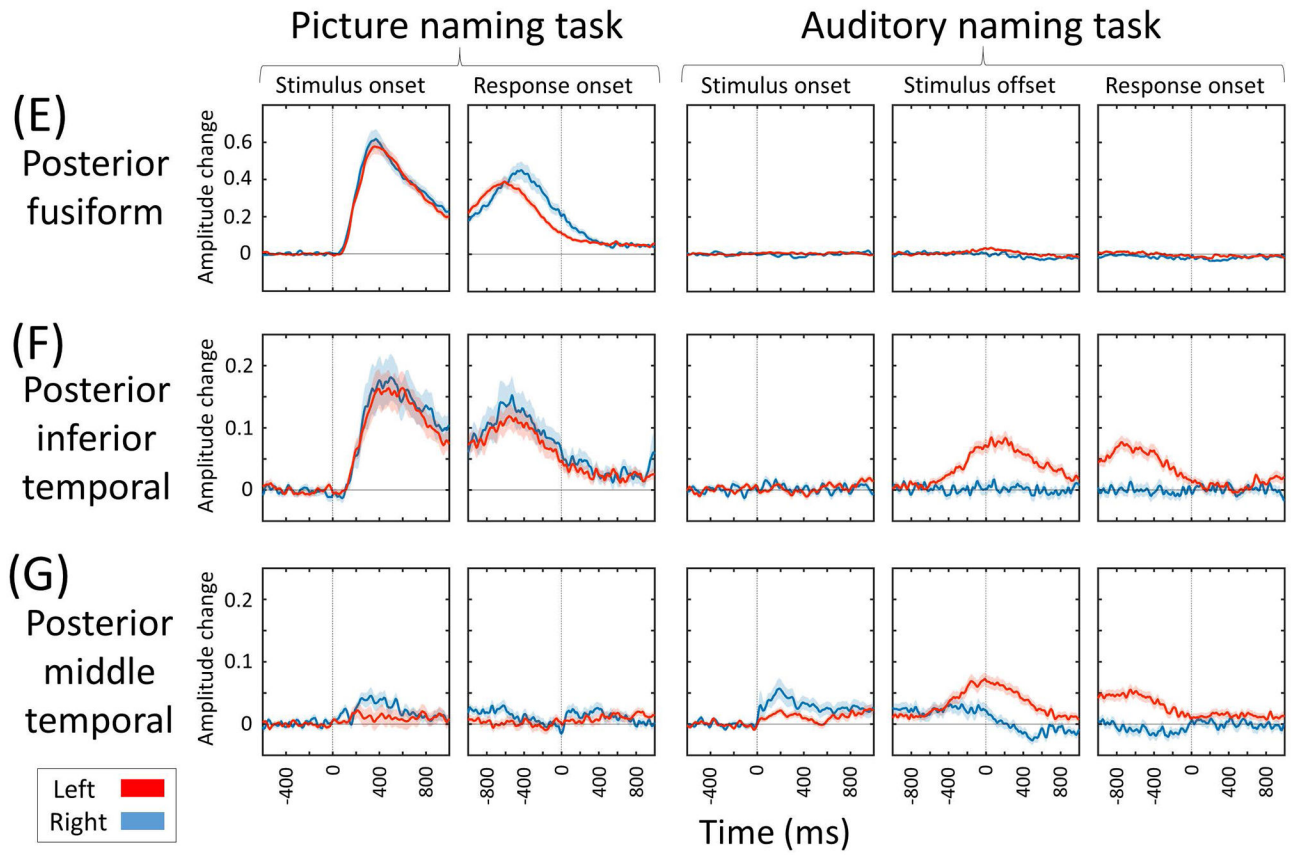


Figure 4. Snapshots of auditory naming-related high-gamma activity.
 (A) 300 ms after stimulus onset. (B) Stimulus offset. (C) 400 ms prior to response onset. (D) Response onset.





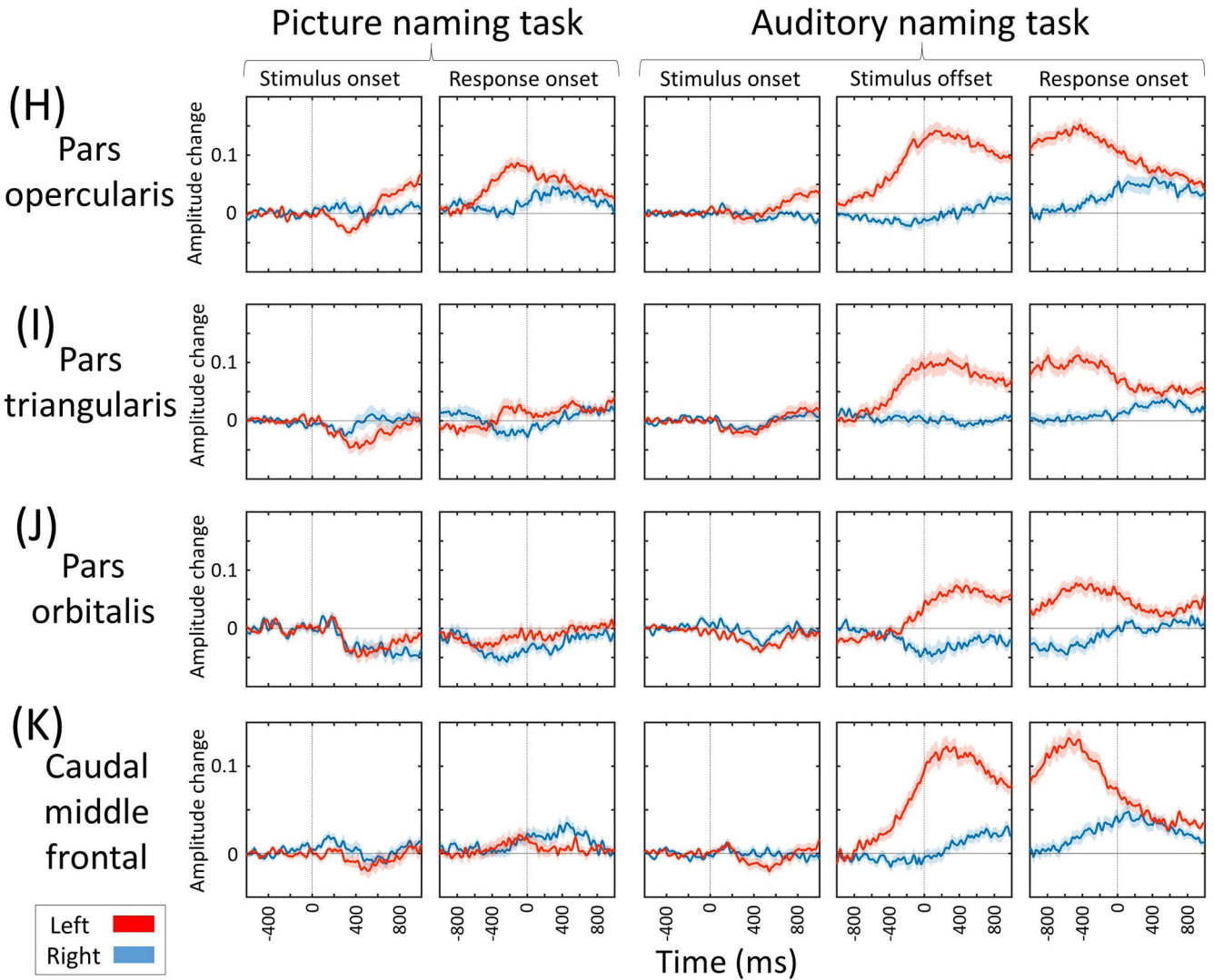


Figure 5. Region of interest analysis of naming-related high-gamma activity.

(A) Medial-occipital region. (B) Lateral-occipital region. (C) Posterior-superior temporal region. (D) Inferior-precentral region. Supplementary Video S2 shows the time-frequency plots of naming-related high-gamma activity at the whole brain level. (E) Posterior fusiform region. (F) Posterior inferior-temporal region. (G) Posterior middle-temporal region. (H) Pars opercularis. (I) Pars triangularis. (J) Pars orbitalis of the inferior-frontal gyrus. (K) Caudal middle-frontal gyrus.

Table 1.

Patient profile.

Mean age (year old)	15.2
Median age (year old)	14
Range of age (year old)	5 – 44
Range of age of epilepsy onset (year old)	0 – 34.8
Proportion of patients with history of GTCs (%)	57.0
Proportion of male (%)	53.2
Proportion of left handedness (%)	10.1
Proportion of sampled hemisphere (%)	
Left	33 (41.8)
Right	28 (35.4)
Both	18 (22.8)
Seizure onset zone [*] . Number of patients (%)	
Left frontal	11 (13.9)
Left temporal	26 (32.9)
Left parietal	8 (10.1)
Left lateral occipital	1 (1.3)
Right frontal	11 (13.9)
Right temporal	19 (24.1)
Right parietal	14 (17.7)
Right lateral occipital	4 (5.1)
Mean number of antiepileptic drugs	1.9
Median number of antiepileptic drugs	2
Range of number of antiepileptic drugs	1 – 4
Antiepileptic drugs ^{**} . Number of patients (%)	
Carbamazepine	6 (7.6)
Oxcarbazepine	33 (41.8)
Lacosamide	26 (32.9)
Zonisamide	5 (6.3)
Phenytoin	6 (7.6)
Lamotrigine	20 (25.3)
Levetiracetam	32 (40.5)
Valproate	10 (12.7)
Clobazam	7 (8.9)
Topiramate	4 (5.1)
Etiology. Number of patients (%)	
Tumor	16 (20.3)
Dysplasia	22 (27.8)

Hippocampal sclerosis	8 (10.1)
Dysplasia + Hippocampal sclerosis	1 (1.3)
No definitive lesion other than gliosis	31 (39.2)
Arteriovenous malformation	1 (1.3)
Mean response accuracy (%)	91.3

Eighty-four percent of patients were 19 years old or younger.

*. Sixteen patients had seizure onset zone involving at least two lobes.

**.. Fifty five patients took at least two antiepileptic drugs.

Response accuracy was defined as the number of correct trials divided by the number of all trials.

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Table 2.

The number of electrode sites at regions of interest (ROIs).

ROIs	Left	Right
aSFG: anterior superior-frontal gyrus.	80	118
pSFG: posterior superior-frontal gyrus.	69	95
rMFG: rostral middle-frontal gyrus.	249	302
cMFG: caudal middle-frontal gyrus.	200	175
POR: Pars orbitalis of the inferior-frontal gyrus.	90	77
PTr: Pars triangularis of the inferior-frontal gyrus.	100	137
POp: Pars opercularis of the inferior-frontal gyrus.	138	90
FP: frontal pole.	1	8
MOrb: medial orbitofrontal gyrus.	21	26
LOrb: lateral orbitofrontal gyrus.	120	92
PCL: paracentral gyrus.	29	44
sPreCG: superior precentral gyrus.	52	64
iPreCG: inferior precentral gyrus.	345	296
sPoCG: superior postcentral gyrus.	73	44
iPoCG: inferior postcentral gyrus.	270	242
PCun: Precuneus.	50	49
SPL: superior parietal lobule.	46	22
SMG: supramarginal gyrus.	291	226
IPL: inferior parietal lobule.	84	116
aSTG: anterior superior-temporal gyrus.	149	134
pSTG: posterior superior-temporal gyrus.	190	135
aMTG: anterior middle-temporal gyrus.	111	86
pMTG: posterior middle-temporal gyrus.	183	87
aITG: anterior inferior-temporal gyrus.	103	91
pITG: posterior inferior-temporal gyrus.	114	58
aFG: anterior fusiform gyrus.	85	102
pFG: posterior fusiform gyrus.	139	75
TP: temporal pole.	39	50
Ent: entorhinal gyrus.	43	64
PHG: parahippocampal gyrus.	22	20
LOG: lateral occipital gyrus.	207	153
MOG: Medial occipital gyrus.	127	137
aCG: anterior cingulate gyrus.	14	19
pCG: posterior cingulate gyrus.	39	41