Abstract

How do animals distinguish between sensations coming from external sources and those resulting from their own actions? A corollary discharge system has evolved that involves the transmission of a copy of motor commands to sensory cortex, where the expected sensation is generated. Through this mechanism, sensations are tagged as coming from self, and responsiveness to them is minimized. The present study investigated whether neural phase synchrony between motor command and auditory cortical areas is related to the suppression of the auditory cortical response. We recorded electrocorticograms from the human brain during a vocalizing/listening task. Neural phase synchrony between Broca’s area and auditory cortex in the gamma band (35 to ∼50 Hz) in the 50-msec time window preceding speech onset was greater during vocalizing than during listening to a playback of the same spoken sounds. Because prespeech neural synchrony was correlated ($r = -0.83$, $p = .006$), with the subsequent suppression of the auditory cortical response to the spoken sound, we hypothesize that phase synchrony in the gamma band between Broca’s area and auditory cortex is the neural instantiation of the transmission of a copy of motor commands. We suggest that neural phase synchrony of gamma frequencies may contribute to transmission of corollary discharges in humans.

INTRODUCTION

There is rapidly growing evidence that synchrony among large populations of neurons is essential for cognitive functions (Engel, Fries, & Singer, 2001; Von Stein, Chiang, & Konig, 2000) and that gamma band activity correlates with various cognitive processes (Pesaran, Pezaris, Sahani, Mitra, & Andersen, 2002; Fries, Reynolds, Rorie, & Desimone, 2001; Varela, Lachaux, Rodriguez, & Martinerie, 2001; Singer, 1999a). It has been hypothesized that cognitive functions require flexibility in the routing of neuronal signals through anatomically connected structures; specifically, communication between two neuronal groups (either within a region or between regions) mechanistically depends on coherence (or phase locking or synchrony) of neural activity between them (Fries, 2005). Synchronous oscillations in gamma-frequency band (e.g., 30–80 Hz) have been implicated in the mechanisms that integrate spatially distributed processing and signal the relatedness of neurons coding for different features of the same object (Varela et al., 2001; Singer, 1999a). Long-range neural integration may be mediated by neuronal groups oscillating together, precisely phase-locking neural activity in the gamma-frequency bands over short periods (Womelsdorf et al., 2007; Fell et al., 2001; Lachaux, Rodriguez, Martinerie, & Varela, 1999; Miltner, Braun, Arnold, Witte, & Taub, 1999; Rodriguez et al., 1999).

This mechanism provides an explanation for integration of cortical information processing between primary and association cortical areas (e.g., primary visual and middle temporal cortical areas; Chen et al., 2007) and different sensory cortical areas (e.g., somatosensory and auditory cortical areas; Lakatos, Chen, O’Connell, Mills, & Schroeder, 2007). Thus, neuronal communication between cortical areas via gamma-band coherence may subserve the integration of cognitive processes (Fries, 2005; Varela et al., 2001; Singer, 1999b).

The experimental evidence mentioned above supports the proposal that synchronous neuronal oscillation in gamma-frequency band is the “functional building blocks” of information processing (Basar-Eroglu, Struber, Schurmann, Stadler, & Basar, 1996), supporting “binding” of brain functions across cortical networks (Phillips & Singer, 1997). Recent studies show that gamma-band synchrony is used for long-range communication across distant brain areas in monkeys, for example, between FEFs and area V4 (Gregoriou, Gotts, Zhou, & Desmone, 2009) and between lateral intraparietal area and medial-temporal area (Saalmann, Pigarev, & Vidyasagar, 2007). One neural mechanism that may rely on cortical communication via gamma-band phase synchrony is the “forward model” (Webb, 2004), involving the transmission of a copy of motor commands from cortical areas involved in planning and executing a motor act to sensory areas involved in processing the resulting sensations. Because the forward model mechanism involves long-range coordinated...
motor–sensory communication, enhancement of neural synchrony might be evident before the execution of motor acts. In fact, local field potential recordings from somatosensory barrel cortex of rats show neural synchrony preceding exploratory whisking in the 25- to 45-Hz band, perhaps triggered by the transfer of a copy of motor commands to somatosensory cortex (Hamada, Miyashita, & Tanaka, 1999). Additional support for a forward model using gamma-band phase synchrony comes from data showing coherent 25 to 35 Hz oscillations in the motor and somatosensory cortices of awake behaving rhesus monkeys (Murthy & Fetz, 1992) and gamma-band coherence between frontal and temporal electrodes in humans (Ford, Gray, Faustman, Heinks, & Mathalon, 2005).

An increasingly recognized forward model is the “corollary discharge.” It has been suggested that the corollary discharge mechanism suppresses sensations that match anticipated sensory consequences of self-generated motor acts (Crapse & Sommer, 2008; Poulet & Hedwig, 2006; Sperry, 1950; Von Holst & Mittelstaedt, 1950; Helmholz, 1867, 1925). The theoretical concept of a forward model has been shared and studied by multiple disciplines in a broad variety of scientific fields such as physiology (Poulet & Hedwig, 2002; Sommer & Wurtz, 2002; Bell, 1981; Davis, Siegler, & Mpitsos, 1973), psychology (Grusser, 1986), psychiatry (Shergill, Samson, Bays, Frith, & Wolkert, 2005; Ford, Mathalon, Heinks, Kalba, & Roth, 2001; Feinberg, 1978), and cybernetics (Guenther, Hampson, & Johnson, 1998; Wolkert, Ghahramani, & Jordan, 1995; Jordan & Rumelhart, 1992). Taking the singing male cricket (Poulet & Hedwig, 2006) as an example, crickets hear through their ears located on their forelegs very close to where 100-dB sound-pressure level chirps are generated. However, singing crickets respond to other cricket’s songs; therefore, the cricket’s CNS must have a way to deal with self-generated intensive sounds during chirping. To avoid being overwhelmed by its own chirps, there is tight coordination between wing motor and sensory systems (Poulet & Hedwig, 2006) using mechanisms of corollary discharge (Crapse & Sommer, 2008). In addition, these concepts apply to cybernetics, as autonomous robots need to distinguish between the consequences of their own actions and the actions of others moving toward them (McKinstry, Seth, Edelman, & Krichmar, 2008).

A forward model of motor–sensory modulation has been proposed to explain how people tell the difference between their own inner verbal experiences or thoughts and “voices” or auditory verbal hallucinations that sound as if they are coming from external sources (Ford, Roach, Faustman, & Mathalon, 2007). This model proposes that a copy of motor commands, transferred from speech production regions in the frontal lobes to the auditory cortex, produces an expectancy of consequences in the auditory cortex to prepare it for an imminent self-generated speech sound. This minimizes the auditory cortical response to these self-generated sounds and provides a mechanism for recognizing these sounds as their own. Support for this mechanism in the human auditory system comes from both intracortical and noninvasive studies. Intracortically, single unit recordings were made during a presurgical planning procedure from the exposed surface of the right and left temporal cortex; while patients talked, auditory cortical responses were reduced compared with when the patients listened to others talking (Creutzfeldt, Ojemann, & Lettich, 1989). Findings from noninvasive, human electrophysiological studies, using EEG and magneto-encephalography synchronized to the onset of vocalization, show similar auditory cortical suppression (i.e., N100 or M100 suppression) during vocalizing (Ford, Roach, et al., 2007; Heinks-Maldonado, Mathalon, Gray, & Ford, 2005; Houde, Nagarajan, Sekihara, & Merzenich, 2002; Ford et al., 2001; Curio, Neuloh, Numminen, Jousmaki, & Hari, 2000). In these studies, the N100 of the EEG-based ERP or the M100 of the magneto-encephalography-based response reveals that activity in auditory cortex, 100 msec after speech onset, is reduced during vocalizing compared with when the same speech sound is recorded and played back. The corollary discharge also works during self-stimulation paradigms; when a tone burst (Martikainen, Kaneko, & Hari, 2005; Schafer & Marcus, 1973) or a speech sound (Ford, Gray, Faustman, Roach, & Mathalon, 2007) results from indirect motor acts of pushing a button, it elicits a smaller N100 than when that exact sound is passively heard. Data from humans are consistent with data from nonhuman primates (Eliades & Wang, 2003; Muller-Preuss & Ploog, 1981) and crickets (Poulet & Hedwig, 2002, 2006). Eliades and Wang (2003) recorded from single units in primary auditory cortex of marmoset monkeys during vocalizing. Like Creutzfeldt et al. (1989), they reported vocalization-induced suppression beginning before vocalization, with both inhibition and excitation of units beginning after vocal onset (Eliades & Wang, 2003, 2005, 2008). They suggested that the origin of the inhibition was from the speech production areas and excitation from the re-afferent of the self-produced vocalization.

A recent diffusion tensor imaging study (Frey, Campbell, Pike, & Petrides, 2008) suggested that Brodmann’s area 45 (Broca’s area) connects with the superior temporal gyrus via the extreme capsule fiber system, providing a plausible route by which a copy of motor commands from Broca’s area travels to auditory cortex. In the present study, we studied forward model mechanisms linking Broca’s area and auditory cortex in patients implanted with cortical electrodes for neurosurgical evaluation while they spoke (vocalizing condition) and then listened to a playback of these recorded sounds (listening condition). We used the amplitude of the N100 component of the auditory ERP to index the cortical response to the self-produced speech sounds during both vocalizing and listening. The N100 has a complex set of subcomponents with both tonotopic and nontonotopic generators (Woods, 1995). Although one of its subcomponents may emanate from primary auditory cortex (Näätänen & Picton, 1987), it peaks too late to reflect the initial cortical volley, which is seen
on the medial aspect of Heschl’s gyrus (Liegeois-Chauvel, Musolino, & Chauvel, 1991; Celesia, Broughton, Rasmussen, & Branch, 1968) and occurs within the first 15 msec after stimulus onset. Determined by clinical necessity, our intracranial electrode grid coverage of the temporal lobe did not allow us to record these early components but only the activity in the more lateral aspects of the supratemporal plane. The high signal-to-noise ratio of these intracranial recordings allowed us to investigate the mechanisms of long-range neural oscillations in a wide frequency range as well as to localize where signals reflecting the copy of motor commands start and where the electrophysiological effects of its expected sensory consequence are seen. Although miniature saccades have been shown to affect the gamma-band response recorded with scalp electrodes during the period from ~200 to 300 msec after stimulus onset (Yuval-Greenberg, Tomer, Keren, Nelken, & Deouell, 2008), intracranial recordings provide a sensitive measure of cortically generated gamma activity. On the basis of our previous findings (Ford, Gray, et al., 2007; Ford, Roach, et al., 2007; Ford et al., 2005), we hypothesized that there would be greater neuronal phase synchrony in the gamma-frequency range (Ford et al., 2005) between Broca’s area and auditory cortex during vocalizing than during listening. We further hypothesized that this neural phase synchrony during vocalizing would reflect the copy of motor commands, as evidenced by a correlation between this synchrony and the auditory cortical suppression of responsiveness during vocalizing (Ford, Gray, et al., 2007; Ford, Roach, et al., 2007).

METHODS
Participants
Participants were three male patients with treatment-resistant epilepsy being evaluated for surgical resection of seizure foci. The experiments were undertaken with the understanding and written consent of each participant. The use of human participants is in conformity with the “Guiding Principles for Research Involving Animals and Human Beings,” and data are HIPAA-compliant. Protocols were approved in advance by the Yale Human Investigation Committee. These patients underwent a phased evaluation that included MRI, interictal single photon emission CT, interictal PET, neuropsychological assessment, video/EEG with ictal and interictal scalp recording in Phase 1, and intracerebral sodium amobarbital testing in Phase 2. Only patients in whom the epileptic area(s) was not determined and/or in whom there was significant discordant data were offered Phase 3 evaluation (i.e., intracranial EEG monitoring). This monitoring was done to localize (1) the seizure focus for later resection and (2) critical language and motor areas to be avoided during surgery.

Patient A was a 45-year-old right-handed white man with medication-resistant complex partial seizures. MRI showed bilateral hippocampal atrophy and increased sig-

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tal change on T2-weighted images. He had resection of right anterior temporal pole and anteromedial hippocampectomy. Pathology showed diffuse glial proliferation of white matter in the right temporal lobe and neuronal loss and gliosis of the hippocampus consistent with hippocampal sclerosis. Patient B was a 37-year-old right-handed white man with medication-resistant complex partial seizures. MRI showed small T2 change in subcortical/cortical lesion in the left frontal lobe, suggesting cortical dysplasia and some T2 signal changes in the right parietal lobe, which were nonspecific. He had resections of posterior superior temporal gyrus and middle temporal gyrus with multiple subpial transections of inferior frontal and parietal regions. Pathology showed fragments of superficial neocortex with artifactualy altered neurons. Patient C was a 52-year-old right-handed white man with medication-resistant complex partial seizures. MRI showed bilateral hippocampal atrophy, and the amygdala was larger on the right than left. He had trans-sylvian amygdalohippocampectomy with preservation of lateral temporal cortex. Pathology showed that the left anterior hippocampus had partial loss of neurons and gliosis consistent with hippocampal sclerosis.

Grid placement was determined entirely by clinical necessity. None of the reported electrode sites in this study was identified as epileptogenic or within the resected area at surgery. We were limited to the patient population presenting for evaluation of their seizure focus for treatment of medically intractable seizures and did not have any influence over the selection and demographic makeup of the patient population. There was no bias in the surgical program regarding patients’ gender, socioeconomic status, or ability to pay for clinical care.

Electrode Localization
For each patient, preoperative MRI scans were acquired, and after surgical installation of the electrodes, both postoperative anatomical MRI scans (with a slice thickness of 1.6 mm) and CT images (with a slice thickness of 1.1 mm) were also acquired. The electrodes were interactively identified in the CT images and then were localized by using the bioimage suite software package developed at Yale University (www.bioimagesuite.org). To determine the location of electrodes included in the analyses, individual electrode locations were marked on postoperative CT images. Briefly, the metal contacts in each electrode are extremely bright in the postoperative CT images. They can be visualized using simple thresholding, which yields clusters of bright voxels. Then interactively, these clusters are labeled by a trained technologist, and the centroid of these clusters is used to define the electrode position. Marked CT images were registered to postoperative MRI scans using a six-parameter rigid transformation, and postoperative MRI scans were then registered with preoperative MRI scans using a nonlinear grid-based transformation (Papademetris, Jackowski, Schultz, Staib, & Duncan, 2004) to account for...
the distortion of the brain that occurs as a result of the craniotomy. Before the later registration, the skulls are stripped from both MRI scans to enhance convergence.

**Experimental Paradigm and Instrumentation**

Participants were tested in a vocalizing/listening experiment using Presentation software (Neurobehavioral Systems Inc., Albany, CA). The experimental paradigm consisted of blocks where the word “Ready” was first presented on the screen for 1 sec, followed by an exclamation point for 1 sec and then a plus sign for 7 sec. Participants were instructed to prepare to say “ah” when the “Ready” message was presented, and then when the plus sign appeared (and remained on the screen for 7 sec), they said “ah” roughly every 1 to 2 sec until the plus sign disappeared. Finally, the word “Rest” would appear on the screen for 5 sec, indicating that participants could stop vocalizing and prepare for the next trial. This vocalizing condition block was repeated 15 times, with an extended break in the middle of the whole recording. During listening condition blocks, recorded sounds from the vocalizing condition were played back, and participants were instructed simply to listen. The participants were trained to produce uniform, brisk utterances with minimal tongue, jaw, and throat movements. Before data acquisition, the participants uttered “ah” several times to facilitate sound system calibration and acclimation to the environment.

Presentation software was used to run both conditions from a laptop running Windows XP and a Sound Blaster Audigy 2 ZS PCMCIA sound card (Creative Technology Ltd., Jurong East, Singapore). The sound card was used to monitor and record the participant and the onset of sound recording or playback. This difference is under half of a millisecond. It is worth noting that although the same sequence of sounds is heard during vocalizing and listening, each individual sound is unique. All patients heard speech sounds binaurally through Etymotic ear insert headphones (Etymotic Research, Inc., Elk Grove Village, IL).

**Data Collection and Event-related Potential Processing**

Electrocorticography (ECoG) data were recorded from a selection of 128 subdural strip and grid electrodes implanted in the neurosurgery patients. The impedance of each electrode was between 5 and 10 kΩ. Signals were amplified using alternating current amplifiers (SA Instrumentation Co., San Diego, CA) before analog band-pass filtering from 0.1 to 100 Hz. The software for collecting ECoG data is homemade (Yale New Haven Hospital, New Haven, CT). Signals were digitized at 14-bit resolution (DAP 4200a; Microstar Laboratories, Bellevue, WA) and sampled at either 1000 Hz (Patients A and B) or 2000 Hz (Patient C). ECoG signals were referenced to a peg electrode. The peg electrode was designed so that the contact lay within the diploic space. The location of the peg electrode was individualized in that the surgeon tried to place it as far as possible from the intracranial electrodes used for the patient. The ground electrode is a four-contact strip placed facing the dura. For off-line processing, the ECoG data were epoched to onsets of participant-generated speech sounds and were baseline corrected. Then, a 60-Hz second-order infinite impulse response notch filter was used.

**Statistical and Data Analysis**

Our analysis methods for Figures 1 to 3 are described in detail here. Supplementary Figure 1a is a schematic illustration of sequences of permutation. The classical Broca’s area (i.e., yellow-circled electrode in Figure 1A; blue-circled electrodes in Figure 3A–C) was determined by intraoperative language mapping for each patient by interruption of speech production (complete speech arrest) with direct electrical stimulation at each electrode. Auditory cortex (i.e., red-circled electrode in Figure 1A; yellow-circled electrodes in Figure 3A–C) was not determined by functional mapping but instead by gyral anatomy and auditory N/M100 component (Reite et al., 1994). Because the interelectrode spacing was approximately 1 cm and primary auditory areas extended for an average of 24 mm.
(Artacho-Perula et al., 2004), we decided to select three electrodes for representing auditory cortex in each patient. None of the recorded electrode sites was identified as epileptogenic or within the resected area at surgery. No data from tissue involved in seizure activity are reported here.

Measuring Phase Synchrony between Electrodes: Phase-Locking Value

Intracranial electrode placements of interest of Patient A are shown in Figure 1A. The ECoG data were epoched to onsets of participant-generated speech sounds, and the averaged ERPs to speech onset of electrodes A12 and G8 are shown in Figure 1B and C, respectively. The red and blue traces are the ERPs for the vocalizing and the listening conditions, respectively. The most N100 suppression of Patient A during vocalizing compared with listening was observed and illustrated in Figure 1C at the electrode contact G8, suggesting that G8 is in or adjacent to primary auditory cortex (Eliades & Wang, 2005, 2008). Eliades and Wang (2005) suggested that the origin of the suppression was from the speech production areas, so we selected the electrode contact A12 in Figure 1A located in Broca’s area.

The neuroelectric signals at the two circled electrode contacts (i.e., A12 and G8) were used to measure phase synchrony between those two contacts by phase-locking values (PLVs) for vocalizing and listening (Figure 1D and E, respectively). These PLVs represent the intertrial variability of the phase differences between two electrode contacts (Fell et al., 2001; Lachaux et al., 1999; Rodriguez et al., 1999). If the phase difference between two electrode contacts varies little across trials, PLV is close to 1 (perfect phase synchrony). Phase synchrony of neuronal oscillation in the gamma-band frequencies has been proposed and shown as an integrative mechanism that brings long-range sets of neurons together into coherently oscillating neuronal groups that can interact effectively.

To generate Figure 1D and E, first, continuous recordings were epoched off-line from −1000 to 1000 msec to avoid edge effects of the wavelet transformation for time window −300 to 300 msec. Seventy-six epoched trials were then spectrally decomposed by a Morlet wavelet transform in the gamma-frequency range from 25 to 50 Hz in 1-Hz step using the freqanalysis_wltconvol.m from FieldTrip toolbox (Radboud University, Nijmegen, the Netherlands) in

![Figure 1](https://example.com/figure1.png)
MATLAB (The MathWorks, Inc., Natick, MA). The wavelet family we used is defined by a constant ratio, $f_0 / \sigma_f = 7$, where $f_0$ is the center frequency and $\sigma_f = 1 / (2\pi f_0)$. At 25 Hz, this leads to a wavelet duration $(2\sigma_f)$ of 89.1 msec and to a spectral bandwidth $(2\sigma_f)$ of 7.1 Hz and at 40 Hz to a duration of 55.7 msec and a bandwidth of 11.4 Hz. Second, for each pair of electrodes, instantaneous phase was extracted by the wavelet decomposition for each time point and frequency on the data from each trial. The result of the wavelet decomposition is a matrix with complex elements. Using the function angle.m in MATLAB, we extracted the phase information from the complex matrix. Third, for each time point of each trial, a phase difference (\(\Delta \theta_{\text{time,trial}}\)) between two selected electrode contacts (e.g., A12-and-G8-paired electrodes in Figure 1A) was determined. Using phase differences of $N$ number of trials, PLV (see Equation 1), ranging from 0 (no synchronization) to 1 (perfect synchronization), was calculated at each time point (Lachaux et al., 1999). Lastly, the third step was repeated for all selected frequencies.

$$PLV_t = \frac{1}{N} \left| \sum_{n=1}^{N} e^{i(\Delta \theta_{t,n})} \right|$$ (1)

**Permutation Procedure for Testing Differences between Two Conditions**

To demonstrate how to differentiate significant PLVs from background fluctuations in each of different experimental conditions, one could implement certain techniques of randomization as described by Lachaux et al. (1999). However, because the results of those randomization techniques would only reveal the significant PLVs within a condition, they would not be able to determine whether the differences of PLVs between two conditions (e.g., Figure 1F) are significant. On the basis of the principles of permutation tests (Maris, Schoffelen, & Fries, 2007; Wilcox, 2004), we developed a new permutation procedure specifically designed for testing the significances of the differences of PLVs between two conditions (e.g., the differences of PLVs between vocalizing and listening conditions in Figure 1F). The details of applied permutation procedure were described and illustrated in Supplementary Figure 1a. The significant differences of PLVs between two conditions in Figure 1F are shown in Supplementary Figure 1b.

**Percentage Changes of Significant Differences of PLVs between Conditions**

Percentage changes of significant differences of PLVs between conditions are shown in Figure 2, and the formula is ($t$ is the time point in msec and $f$ is the frequency in Hz)

$$\text{Percentage change}_{(t,f)\text{significant}} = \frac{(\text{PLV}_{(t,f)\text{Vocalizing}} - \text{PLV}_{(t,f)\text{Listening}})_{\text{significant}}}{\text{PLV}_{(t,f)\text{Listening}}} \times 100\%$$ (2)

Because it was based on only the “significant” differences of PLVs between vocalizing and listening conditions (i.e., $[\text{PLV}_{\text{Vocalizing}} - \text{PLV}_{\text{Listening}}]_{\text{significant}}$, Supplementary Figure 1b), red regions would represent significant enhanced changes of PLVs during vocalizing compared with the listening condition, blue regions would represent significant suppressed changes of PLVs during vocalizing compared with the listening condition, and green regions would represent nonsignificant changes of PLVs between two conditions. One of the advantages of percentage changes (or modulation) is that enhanced/suppressed phase synchronies across different patients can be compared.

**Relationship between N100 Suppression and PLVs of Gamma Frequencies**

**PLV between Broca’s area and auditory cortex.** One Broca’s area electrode and three auditory cortex electrodes were selected for each patient, making three pairs of electrodes for each patient (i.e., A12-G8, A12-G9, and A12-G10 for Patient A in Figure 3A; G3-G50, G3-G42, and G3-G34 for Patient B in Figure 3B; G6-G45, G6-G57, and G6-G29 for Patient C in Figure 3C).

For each pair of electrodes, we first calculated the percent change between conditions for all PLVs that were significantly different during vocalizing and listening (i.e., the method for generating Figure 2). Next, we averaged these values between the $-50$- and the $0$-msec time window.
avoiding possible event-related changes in coherence that might cause significant differences of PLVs. These mean values represented the degree of synchrony for each electrode pair for each participant, and higher gamma phase synchrony is reflected in larger mean values in the x-axis of Figure 3D. These mean values were used to assess the degree of association between interregion synchrony during vocalizing and subsequent suppression of auditory cortical activity to the spoken sound.

Relating N100 suppression to gamma phase synchrony during vocalizing. N100 SUPPRESSION. The averaged ERPs to speech onset of auditory cortex electrodes (three for each patient) are yellow-circled and shown in Supplementary Figure 2a–c. The averaged ERPs to speech onset of green-rectangle-selected electrodes are shown in the bottom-left corner of Figure 3A–C (magenta trace is vocalizing and blue trace is listening; also shown in the Supplementary materials). From the averaged ERP, the N100 was measured as the most negative peak between 50 and 150 msec, area ±25 msec around the peak were calculated (i.e., the hollow circles and the dashed line; $r = - .825$, $p = .006$). In addition, a mixed model ANCOVA was applied to those hollow-circled values, $F(1, 5) = 9.058$, $p = .03$. The other two-tailed Pearson’s $r$ between the best N100 suppression (from the green-rectangle-selected electrode) and the grand mean of averaged percentage changes of significant differences of PLVs was computed (i.e., the solid circles and line; $r = - .998$, $p = .044$).

Figure 3. Relationship between N100 suppression and averaged percent change of significant differences of PLVs. Intracranial electrode placements of Patients A, B, and C are shown in panels A, B, and C, respectively. The neuroelectric signals of circled electrode contacts (i.e., Broca’s area circled in blue and auditory cortex circled in yellow) were used for calculating PLVs and the averaged percent change of significant differences of PLVs (see details in the Methods section). The averaged ERPs to speech onset of green-rectangle-selected electrodes are shown in the bottom-left corner of panels A, B, and C (magenta trace is vocalizing and blue trace is listening). (D) The averaged percent change of significant differences of PLVs is on the x-axis, and N100 suppression is shown on the y-axis (see details in the Methods section). Two-tailed Pearson’s correlation coefficients ($r$) and ANCOVA results are shown in the upper-right corner. A two-tailed Pearson’s correlation coefficient ($r$) between the yellow-circled N100 suppression and its corresponding averaged percentage change of significant differences of PLVs was calculated (i.e., the hollow circles and the dashed line; $r = - .825$, $p = .006$). In addition, a mixed model ANCOVA was applied to those hollow-circled values, $F(1, 5) = 9.058$, $p = .03$. The other two-tailed Pearson’s $r$ between the best N100 suppression (from the green-rectangle-selected electrode) and the grand mean of averaged percentage changes of significant differences of PLVs was computed (i.e., the solid circles and line; $r = - .998$, $p = .044$).
N100 was seen best at electrodes over primary auditory cortex (i.e., the green rectangle within the yellow circle shown in Figure 3A and C) in Patients A and C. However, in Patient B, N100 was best seen at an electrode well above the electrodes placed over primary auditory cortex (i.e., the green rectangle shown in Figure 3B and Supplementary Figure 2b). Because each participant’s anatomy is different and because electrodes are placed for clinical necessity not for scientific convenience, it is likely that the electrode placement over the temporal lobe in Patient B was not ideal for recording N100. We suggest that in Patients A and C, N100 was generated in superior temporal gyrus, directly under the electrodes shown in the green rectangle, with the dipole generating N100 being perpendicular to the electrode. In Patient B, we suggest that N100 was generated deep in the STS, at an orientation tangential to the electrodes on the gyrus, making it electrically invisible at those sites. Instead, the dipole generating N100 was likely perpendicular to electrodes well above the generator but visible to them. Depending on the orientation of the dipole plane generating the auditory evoked potentials, one might expect to observe evoked potentials generated in auditory cortex volume conducted to a distant site. Unfortunately, we do not have the necessary coverage over the area to confirm this suggestion with a dipole source analysis.

**RESULTS**

**PLVs between Broca’s and Auditory Cortical Areas**

The PLVs between electrodes over Broca’s area and auditory cortex are greater during vocalizing (Figure 1D) than during listening (Figure 1E), which suggests that there is more phase synchrony between those two electrode sites during vocalizing than listening. Figure 1F shows the difference between PLVs during vocalizing and listening. We found a similar vocalizing-related correlation in the scalp-recorded EEG over Broca’s and Wernicke’s areas in the gamma band (Ford et al., 2005), without the spatial precision afforded by this study.

On the basis of the results of our permutation test (see details in the Methods section and Supplementary Figure 1A and B), Figure 2 shows the percent changes in the significant differences of PLVs between vocalizing and listening conditions with listening condition as the baseline. Between −50 and 100 msec, phase synchronies between Broca’s and auditory cortical areas during vocalizing were significantly greater (warm colors in Figure 2) than those during listening for two frequency ranges within the 25- to 50-Hz frequency window (i.e., 25 to ∼33 Hz and 35 to ∼50 Hz). In the period of −50 to 0 msec, before the onset of speech, there are more significant gamma band phase synchronies between Broca’s and auditory cortical areas during vocalizing in the frequency range from 35 to ∼50 Hz. In the period of 40 to 120 msec, after the onset speech, there are more significant phase synchronies between those two areas during vocalizing in frequency ranges from 25 to ∼33 Hz and also from ∼38 to 50 Hz.

**Cortical Interaction and N100 Suppression**

As can be seen in Figure 3D, higher gamma phase synchrony is reflected in larger values on the x-axis, and more
N100 suppression is seen in smaller numbers on the y-axis. A significant negative correlation (indicated by the dashed line in Figure 3D; \( r = -0.825, p = 0.06 \)) means that higher gamma phase synchronies between Broca’s and auditory cortical areas correlate with greater N100 suppression in auditory cortices. To avoid problems associated with non-independence of data points (three pairs of electrodes from each participant), we also calculated the relationship between N100 suppression and gamma phase synchrony using a mixed model ANCOVA, with the “patient” variable included as a random factor, gamma phase synchrony as a covariate, and N100 suppression as a dependent variable. Patient did not affect N100 suppression, but phase synchrony did, \( F(1, 5) = 9.058, p = 0.03 \), consistent with the Pearson’s correlation analysis.

Although there were only three points (one for each patient) in our second correlational analysis, the correlation coefficient showed a significant trend that higher gamma phase synchronies between Broca’s and auditory cortical areas correlated with greater N100 suppression (i.e., the solid line in Figure 3D; \( r = -0.998, p = 0.044 \)). In both correlation analyses, it is clear that Patient B had the lowest PLVs and the least suppression of N100.

As described previously in our methods, all the ECoG data that were used for calculating Figure 3D preceded the onset of vocalization (i.e., between -50 and 0 msec; 0 msec is the onset of vocalization). Because of the fact that movement artifacts are much less prominent in ECoG than scalp-recorded EEG and that data during vocalization were excluded from the analyses for Figure 3D, it is unlikely that this finding could result from movement in the vocalizing condition. Lastly, we illustrated that the percentage changes of significant differences of PLVs were not widespread and not due to artifacts recorded from the same electrode grid in the Supplementary Figure 3A and B.

**DISCUSSION**

Helmholtz (1867) described a forward model mechanism that allows us to discriminate between real moving objects and movements on the retina resulting from eye movements. This can be demonstrated by comparing the visual image during a saccade with the visual image experienced when the eye is externally moved by tapping the corner of the eye. In the former case, but not in the latter, visual perception is suppressed, a phenomenon known as saccadic suppression (Thiele, Henning, Kubischik, & Hoffmann, 2002). After Helmholtz (1867, 1925), others suggested that this suppression was accomplished through the action of an efference copy (Von Holst & Mittelstaedt, 1950) or a corollary discharge (Sperry, 1950). This mechanism is responsible for our critical ability to unconsciously recognize and disregard sensations resulting from our own actions (Von Holst & Mittelstaedt, 1950).

Although the action of this mechanism is undeniable and essential to negotiating and sensing our complex environment, until recently its neurophysiological basis was unknown. It is now being studied in vertebrates and invertebrates with electrophysiological methods similar to those used with humans, making findings cross-translational. The concept of a forward model is increasingly recognized as an important modulator of motor control, sensory processing, and cognition (Crapse & Sommer, 2008). It is argued that the forward model is an internal loop that uses the copy of motor commands to predict the expected sensation and to modulate its sensory processing. To this end, there should be a specific output stream to sensory input areas from areas generating this output. This kind of stream or pathway has been identified in animals with “simple” nervous systems, such as crickets, by analyses of synaptic connectivity (Poulet & Hedwig, 2006).

In the present study, we investigated neuronal communication between Broca’s area and auditory cortex. Our data suggest that neural phase synchrony of gamma frequencies (e.g., 35–50 Hz) between Broca’s area and auditory cortex may at least partially explain the neural mechanism of corollary discharge in humans. Gamma-band phase synchrony in the 50-msec period immediately preceding speech onset may be the instantiation of a copy of motor commands being sent from Broca’s area to auditory cortex. Although we cannot argue causality from correlation, in that it is correlated with the suppression of auditory cortical responses, 100 msec after speech onset supports this interpretation. Furthermore, the gamma-band phase synchrony precedes speech onset; therefore, these synchronies were not due to the auditory responses of the uttered speech.

Although the functional meaning of specific frequencies is still unclear, synchronous oscillations at specific frequencies could identify even distant neuronal populations as belonging to the same functional network of spatially distributed neuronal assemblies (Bastiaansen & Hagoort, 2006). Possibly, only a portion of the gamma band is used for modulating sensory processing in the output stream from Broca’s area to auditory cortex in our proposed forward model, and different gamma frequencies may participate in distinct neurophysiological mechanisms and reflect different aspects of cortical activation during information processing (Roopun et al., 2008). However, the range of frequencies greater than 80 Hz remains largely unexplored, and the function of high gamma oscillation is unknown.

To rule out the simple effects of agency and predictability on our findings, we studied the effects of agency (subjects pressed a button to deliver an “ah”) and predictability (a visual warning stimulus gave precise information about when an “ah” would be presented); in neither case was cortical suppression of the sensation as great as when subjects said “ah” (Ford, Gray, Faustman, Roach, & Mathalon, 2007). It is also worth mentioning that when we shifted the pitch of the “ah” during talking, there is still suppression, but less (Heinks-Maldonado et al., 2005).

We are becoming increasingly aware that locally specialized functions in the brain must be coordinated with each
other. In our vocalizing/listening paradigm, the data show that participants with higher gamma phase synchrony between Broca’s area and auditory cortex (before the onset of speech) have greater N100 suppression to the spoken sound. We speculate that (1) the transmission of a copy of motor commands to appropriate sensory cortex may be an emergent property of a self-organizing system, (2) communication among neuronal populations requires precise matching of the relative phase of distinct frequencies, and (3) integration of those mechanisms is responsible for our critical capability to unconsciously recognize and disregard sensations from our own actions. This type of ongoing motor–sensory interaction illustrates how important the neural system’s constant-changing “context” is in processing new external sensory “content” (Arieli, Sterkin, Grinvald, & Aertsen, 1996).

We recognize that intracranial data sets are traditionally small and the conclusions drawn from them may not be generalized to a large population; however, the results were significant in both correlation analyses and suggest a very promising possibility that should be replicated in another sample. In addition, the data are rare and valuable, and the findings are consistent with previous findings and models described below.

The pioneering work by Creutzfeldt et al. (1989), the only other such study using intracortical recordings from the human auditory cortex during speech, showed that while patients talked, auditory cortical responses were reduced compared with when the patients listened to others talking. However, this work is not directly comparable with ours because they used tape-recorded words and sentences of different lengths, spoken by different speakers and not by the patients themselves. Also, they used single unit recordings and their recordings were restricted to regions to be resected, which excluded recordings from language areas. Nevertheless, their findings also suggested possible mechanisms for a forward model that could explain how people tell the difference between external and self-generated sounds.

Our results are consistent with the findings in auditory–visual (AV) speech studies (Kauramaki et al., 2010; Skipper, van Wassenhove, Nusbaum, & Small, 2007; van Wassenhove, Grant, & Poeppel, 2005). Van Wassenhove et al. (2005) suggested that the first computational stage of AV speech integration is a feature-extraction stage where visual information enables the prediction of the auditory input, and this early visual processing can be used in a forward model, that is, “analysis-by-synthesis” model, for computing the residual error between the following auditory input and the internal prediction. Later, Skipper et al. (2007) illustrated this predictive mechanism using fMRI and showed that AV and visual speech perception yielded more robust activities in the motor system associated with speech production, supporting their hypothesis that motor commands can be used to predict the sensory consequences of those corresponding movements through the transmission of a copy of those motor commands. Kauramaki et al. (2010) suggest that the origin of the top–down inputs that cause N100 or M100 suppression during lipreading is from the speech production system that acts like mirror neurons (Rizzolatti & Craighero, 2004). Specifically, it was hypothesized that during lipreading the observers do not vocalize themselves, but their speech production cortical areas are activated through “mirroring” of observed vocal actions; a copy of this mirroring response could then be sent to auditory cortex and enhance auditory processing (Kauramaki et al., 2010). In the present study, our findings provide new information for explaining the possible underlying mechanisms of a forward model or corollary discharge/efference copy in AV speech research. In addition, our results support the consensus that interacting cortical areas are associated with the production and coordination of speech, illustrated and investigated by a comprehensive speech acquisition and production model, the Directions Into Velocities of Articulators (DIVA) model (Guenther, Ghosh, & Tourville, 2006; Guenther et al., 1998). According to the DIVA model, the expected auditory consequences of self-generated speech are encoded in the forward model projections from the “speech sound map” (hypothesized to lie in the left posterior inferior frontal gyrus and ventral premotor cortex) to the “auditory target map” (hypothesized to lie in bilateral planum temporale and superior temporal gyrus); if the incoming auditory signals are within the predicted range, there is an inhibition that cancels the excitatory effects of those incoming auditory signals. Converging with the hypothesis of auditory expectations projected from the speech sound map to the auditory target map in the DIVA model, our findings may suggest how multiple cortical areas interact with one another. Furthermore, the present study may contribute potential mechanisms to be used in EEG-based brain–machine interfaces for providing communication capabilities to profoundly paralyzed patients (Guenther et al., 2010).

In addition to dampening irrelevant sensations resulting from our own actions, the forward model provides a mechanism for automatic distinction between internally and externally generated percepts across sensory modalities and may even operate in the realm of covert thoughts, which have been viewed as our most complex motor act (Feinberg, 1978). An extension of the forward model to the process of thinking would be that the covert rehearsal of a word or phrase requires activation of the auditory cortical areas involved in speech perception, and in this case, the generation of corollary discharge is essential to distinguishing between self- and externally generated sensations (Shergill et al., 2002). The translational potential for the present study and other studies of the corollary discharge mechanism is illustrated by Feinberg (1978), who suggested that patients with schizophrenia may have a deficit in this basic neural mechanism, making it difficult for them to distinguish their own inner thoughts and memories from external voices, resulting in auditory hallucinations. Specifically, if a copy of a memory, a thought, or
REFERENCES


