

The Neuroanatomy of Speech Processing: A Large-scale Lesion Study

Corianne Rogalsky¹, Alexandra Basilakos², Chris Rorden², Sara Pillay³, Arianna N. LaCroix¹, Lynsey Keator², Soren Mickelsen¹, Steven W. Anderson⁴, Tracy Love⁵, Julius Fridriksson², Jeffrey Binder³, and Gregory Hickok⁶

Abstract

■ The neural basis of language has been studied for centuries, yet the networks critically involved in simply identifying or understanding a spoken word remain elusive. Several functional–anatomical models of critical neural substrates of receptive speech have been proposed, including (1) auditory-related regions in the left mid-posterior superior temporal lobe, (2) motor-related regions in the left frontal lobe (in normal and/or noisy conditions), (3) the left anterior superior temporal lobe, or (4) bilateral mid-posterior superior temporal areas. One difficulty in comparing these models is that they often focus on different aspects of the sound-to-meaning pathway and are supported by different types of stimuli and tasks. Two auditory tasks that are typically used in separate studies—syllable discrimination and word comprehension—often yield different conclusions. We assessed syllable discrimination (words and nonwords)

INTRODUCTION

The neural basis of speech perception-defined here as the processes up to and including the successful activation of a phonological word form-has been a contentious topic for decades, if not centuries, and remains so today. Hickok and Poeppel (2000, 2004, 2007) have reviewed the earlier literature and argued that task effects are a significant source of the lack of agreement: Depending on what task is used, researchers can get different answers. In particular, they pointed out that syllable discrimination, judging whether two syllables are the same or different, can yield misleading results because they do not necessarily tap into all of the same processes involved in processing speech sounds under more normal, ecologically valid listening conditions, such as during comprehension. Specifically, it was argued that discrimination tasks recruit not only perceptual systems in the superior temporal gyrus (STG) but also more dorsal, motor speech-related pathways. Comprehension tasks implicate perceptual systems

and word comprehension (clear speech and with a noise masker) in 158 individuals with focal brain damage: left (n = 113) or right (n = 19) hemisphere stroke, left (n = 18) or right (n = 8) anterior temporal lobectomy, and 26 neurologically intact controls. Discrimination and comprehension tasks are doubly dissociable both behaviorally and neurologically. In support of a bilateral model, clear speech comprehension was near ceiling in 95% of left stroke cases and right temporal damage impaired syllable discrimination. Lesion-symptom mapping analyses for the syllable discrimination and noisy word comprehension tasks each implicated most of the left superior temporal gyrus. Comprehension but not discrimination tasks also implicated the left posterior middle temporal gyrus, whereas discrimination but not comprehension tasks also implicated more dorsal sensorimotor regions in posterior perisylvian cortex.

in the STG as well but do not engage the more dorsal stream and instead engage more ventral middle temporal regions. Despite highlighting these potential problems, discrimination tasks are still regularly employed to assess the neural basis of speech perception, which may perpetuate confusion and contraction in the field (Stokes, Venezia, & Hickok, 2019; Skipper, Devlin, & Lametti, 2017; Venezia, Saberi, Chubb, & Hickok, 2012). Aiming to test the task-dependence claim and to map the neural correlates of speech perception, we compared stimulusmatched word-picture matching comprehension tasks with same-different auditory syllable discrimination tasks in a large sample of unilateral stroke survivors. In what follows, we provide a brief summary of the major hypotheses regarding the neural basis of speech perception and then describe the present experiment. It is important to note that we do not include in our summary an important body of literature on more fine-grained aspects of speech processing in aphasia that has contributed knowledge of the psycholinguistic subtleties of disordered speech abilities and their implications for neurolinguistic models of speech processing. We exclude this work here because, unfortunately, neural correlates of the behavioral observations are not available beyond aphasia syndrome-level

¹Arizona State University, Tempe, ²University of South Carolina, Columbia, ³Medical College of Wisconsin, Milwaukee, ⁴University of Iowa, Iowa City, ⁵San Diego State University, California, ⁶University of California, Irvine

classification of cases. We encourage the reader to consult Blumstein (1998) for a thorough review of this early literature.

Carl Wernicke (Wernicke, 1874) proposed that the left superior temporal lobe serves as the substrate for what we today generally call speech perception. The proposal is still viable today (Albouy, Benjamin, Morillon, & Zatorre, 2020) and is intuitively appealing in that the region is in auditory-related cortex and in the language-dominant hemisphere. However, the claim has been challenged from several angles, leading to at least three current competitors to Wernicke's hypothesis.

The first challenge, based mostly on studies of phoneme perception using tasks such as syllable discrimination, disputes the view that sensory cortex alone is sufficient for perceptual processes, as putatively perceptual speech tasks have been found to invoke the involvement of various levels and subsystems of motor cortex (Pulvermüller et al., 2006; Pulvermüller, Hauk, Nikulin, & Ilmoniemi, 2005; Wilson, Saygin, Sereno, & Iacoboni, 2004; Watkins, Strafella, & Paus, 2003; Rizzolatti & Arbib, 1998). This view was first proposed in the form of the motor theory of speech perception (Liberman, Cooper, Shankweiler, & Studdert-Kennedy, 1967; Liberman, 1957) and indirectly linked the process to motor cortex. It was subsequently challenged by several examples of normal speech perception ability in the face of impaired or nonexistent motor speech ability (Bishop, Brown, & Robson, 1990; Kuhl & Miller, 1975; Eimas, Siqueland, Jusczyk, & Vigorito, 1971), but the motor theory claim was revived with the discovery of mirror neurons in macaque monkeys (Gallese, Fadiga, Fogassi, & Rizzolatti, 1996). Consistent with this motorcentric view of speech perception, modern neuroimaging studies have shown that (i) motor speech regions indeed activate during listening to phonemes and syllables (Pulvermüller et al., 2006; Wilson et al., 2004; Hickok, Buchsbaum, Humphries, & Muftuler, 2003; Buchsbaum, Hickok, & Humphries, 2001) and (ii) transcranial stimulation (TMS) studies have reported modest performance declines on syllable identification or discrimination tasks (D'Ausilio et al., 2009; Meister, Wilson, Deblieck, Wu, & Iacoboni, 2007; see Stokes et al., 2019, for a detailed evaluation of effect size). However, strong versions of the motor centric hypothesis remain improbable for reasons given above (Hickok, 2014). A more moderate proposal in this vein is that the motor system augments speech perception only under noisy listening conditions, as broadly suggested by evidence that people with Broca's aphasia have difficulty processing speech under degraded listening conditions (Wilson, 2009; Moineau, Dronkers, & Bates, 2005; Utman, Blumstein, & Sullivan, 2001). However, this hypothesis is complicated by the fact that lesions associated with Broca's aphasia also often involve damage to the superior temporal lobe (Fridriksson, Fillmore, Guo, & Rorden, 2015).

A second challenge to Wernicke's model came from neuroimaging studies at the beginning of this century showing a left dominant and specifically anterior superior temporal region that appeared to respond selectively to intelligible speech (Scott, Blank, Rosen, & Wise, 2000). Although this anterior superior temporal region has been proposed to be critical for speech-specific processing more broadly, the stimuli in these studies have typically been sentences (Evans et al., 2014; Narain et al., 2003; Scott et al., 2000), raising the possibility that the activation in these regions is at a postphonological stage of processing (Brennan & Pylkkänen, 2012; Rogalsky, Love, Driscoll, Anderson, & Hickok, 2011). Furthermore, subsequent higher-powered replications of the original report show a bilateral pattern of activation involving both anterior and posterior temporal areas (McGettigan et al., 2012; Okada et al., 2010). Nonetheless, this work highlights the potential role of the anterior temporal lobe (ATL) in addition to more traditional posterior temporal areas.

Finally, a third challenge came from the study of people with language disorders (aphasia) following brain injury: It was found that left hemisphere damage generally produced only mild deficits in speech perception (as measured by word to picture matching tasks with phonological foils), whereas bilateral superior temporal lesions could produce profound deficits even when pure-tone hearing thresholds are normal, that is, "word deafness" (Poeppel, 2001; Buchman, Garron, Trost-Cardamone, Wichter, & Schwartz, 1986; Miceli, Gainotti, Caltagirone, & Masullo, 1980; Henschen, 1926). In addition, studies of the isolated right hemisphere in callosotomy patients or during Wada procedures found that the right hemisphere exhibited good comprehension of most auditory words and simple phrases (Hickok et al., 2008; Zaidel, 1985; Gazzaniga & Sperry, 1967). These findings led to the proposal that speech perception can be mediated in the superior mid to posterior temporal lobes bilaterally (Hickok et al., 2008; Hickok & Poeppel, 2000, 2004, 2007; Goodglass, 1993; Bachman & Albert, 1988; Buchman et al., 1986; Zaidel, 1985; Gazzaniga & Sperry, 1967), which would explain the substantial effects of bilateral damage, whereas individuals with unilateral damage (to either hemisphere) are typically only mildly impaired on auditory word comprehension tasks. Of course, the bilateral claim is consistent with a large body of functional activation studies showing bilateral superior temporal lobe involvement in speech perception (DeWitt & Rauschecker, 2012, 2016; Leonard, Baud, Sjerps, & Chang, 2016; Moses, Mesgarani, Leonard, & Chang, 2016; Overath, McDermott, Zarate, & Poeppel, 2015; Okada et al., 2010; Hickok & Poeppel, 2007; Myers, 2007; Okada & Hickok, 2006; Binder et al., 2000).

Much of the modern research on this topic has been carried out using neuroimaging methods, which cannot assay the causal involvement of activated networks. There have been attempts to assess the competing proposals more directly using neuropsychological (Bak & Hodges, 2004) or, as noted above, neural interference methods (TMS; D'Ausilio et al., 2009; Mottonen & Watkins, 2009; Meister et al., 2007). However, questions have been raised about the generalizability of these results to natural receptive speech processing, particularly for the TMS studies, because the reported effects on perceptual accuracy are small (estimated to amount to a decrement of only about 1–2 dB; Stokes et al., 2019), are only found for syllable discrimination or identification tasks not comprehension (Schomers, Kirilina, Weigand, Bajbouj, & Pulvermüller, 2015) and are evident only in near-threshold listening conditions (see Stokes et al., 2019; Hickok, 2014, for discussion). Thus, despite more than a century of study, the neural basis of speech perception remains controversial.

To summarize, we have outlined the following four hypotheses regarding the neural basis of speech perception, again, defined as the processes up to and including the successful activation of a phonological word form:

- 1. Classical model—speech perception is uniquely supported by the left mid-to-posterior superior temporal lobe.
- 2. Motor model—speech perception is dependent on (strong version) or augmented by (weak version) one or more motor subsystems involved in speech production. Augmentation for speech perception under the weak model is typically assumed to hold for noisy or otherwise effortful listening conditions.
- 3. Left anterior temporal model—speech perception is dependent on the left anterior superior temporal lobe.
- 4. Bilateral temporal model—speech perception is bilaterally organized in the mid-to-posterior superior temporal lobe.

We have also outlined an additional claim that discrimination and comprehension tasks implicate shared networks in the STG but then engage different dorsal (discrimination) versus ventral (comprehension) pathways beyond this region. Thus,

5. Task-dependent claim—speech processing as measured by discrimination tasks will implicate the mid-to-posterior STG and more dorsal regions, while speech processing as measured by comprehension will implicate the mid-to-posterior STG and more ventral regions.

To test these hypotheses, we conducted a large-scale lesion study employing two auditory word comprehension tasks and two syllable discrimination tasks. Both of the auditory word comprehension tasks involved matching a spoken word to a picture in an array of four where one picture was the target and the other three were phonological, semantic, or unrelated distractors. In one version of the task, speech was presented in a quiet background (clear speech), and in the other version, speech was presented in white noise. The speech-innoise task served two purposes, one to test claims about the role of the motor system in processing noisy speech and, second, to ensure that the distribution of performance on word comprehension is below ceiling, thus enabling lesion mapping.

The syllable discrimination tasks involved the presentation of pairs of monosyllabic speech stimuli with minimally contrasting onsets and a 1-sec interstimulus interval. The listener was asked to decide whether the two stimuli were the same or different. One version of the task involved real words and the other involved nonwords. Despite the intuitive assumption that both discrimination and auditory word comprehension tasks measure an ability to extract sound-based phonological information from a speech signal, these two tasks are reported to be doubly dissociable (Hickok & Poeppel, 2004; Miceli et al., 1980). Particularly paradoxical are cases in which the ability to discriminate pairs of syllables (e.g., da from ba) is impaired while the ability to differentiate words with these same sounds (e.g., doll versus ball) is spared in comprehension tasks. This has confounded models of the neural basis of speech perception and recognition for decades because one can reach different conclusions depending on which task is used (Hickok, 2009, 2014; Hickok & Poeppel, 2000). We sought to resolve this issue in this study by employing both tasks in a large sample. We studied people with either left or right hemisphere chronic stroke, left or right anterior temporal lobectomy, and neurologically intact controls. For our largest group (the left hemisphere stroke group), we use lesion-behavior mapping to identify brain regions responsible for receptive speech deficits.

Predictions

Based on their evaluation of previously published studies, Hickok and Poeppel (2000, 2004, 2007) predict that word comprehension and syllable discrimination tasks are doubly dissociable both behaviorally and neurally, even when the syllables in the discrimination task are words. This would argue strongly that discrimination tasks add some process to speech perception that is not necessarily used during normal word comprehension (see the work of Burton, Small, & Blumstein, 2000). Assuming that researchers are aiming to understand how speech is processed "in the wild" (this is indeed our own goal) and assuming that the word comprehension task is a closer approximation to ecological validity, such a result would seriously question the validity of discrimination tasks in studies of speech perception. Turning to the neuroanatomical models, the classical model predicts that damage to the posterior superior temporal lobe should yield substantial deficits on both tasks. The motor models predict that damage to frontal, motor-related regions—which may include primary motor cortex (D'Ausilio et al., 2009), premotor cortex (Meister et al., 2007), inferior frontal gyrus (Mottonen & Watkins, 2009; Burton et al., 2000), or subparts thereof-should cause speech perception deficits. In this study, we sample all of these regions and so will be in position to test each of these possibilities. The strong variant predicts deficits on all tasks, whereas the weak variant predicts impairment on tasks involving speech in noise. The left anterior temporal model predicts that unilateral mid-to-anterior superior temporal lobe damage should substantially interfere with word comprehension via disruption to systems critical for recovering intelligibility. It is less clear what predictions this model has for nonword discrimination. The bilateral temporal model generally predicts minimal impairment on speech perception tasks, perhaps of any type, following unilateral damage. The task-dependent claim predicts overlapping lesion correlates for syllable discrimination and word comprehension tasks in the STG, with differences emerging in dorsal regions (discrimination only) and ventral regions (comprehension only).

METHODS

Participants

Consent from all participants (described below) was obtained according to the Declaration of Helsinki, and the experiments were approved by the institutional review boards of all the institutions at which the work was performed.

Lesion Participants

The following groups of participants were examined: 113 individuals who had previously experienced a left hemisphere stroke (Figure 1), 19 individuals who had previously experienced a right hemisphere stroke (Figure 2A, B), 18 individuals who had previously undergone a unilateral left temporal lobectomy, and 8 individuals who had previously undergone a unilateral right temporal lobectomy (Figure 2C, D). See Table 1 for demographic information of each group. The left hemisphere stroke participants were tested at six institutions: University of Iowa (n = 40), University of South Carolina (n = 32), Medical College of Wisconsin (n = 24), San Diego State University (n = 9), Arizona State University (n = 7), and University of California, Irvine (n = 1). The right hemisphere stroke participants were tested at University of Iowa (n = 17), University of South Carolina (n = 1), and Arizona State University (n = 1). All lobectomy participants were tested at the University of Iowa. Participants were included in this study based on the following criteria: (i) a chronic focal (6 months or more postonset) lesion because of a single stroke in either the left or right cerebral hemisphere or a unilateral temporal lobectomy, (ii) no significant anatomical abnormalities other than the signature lesion of their vascular event (or evidence of surgery for the lobectomy participants) nor signs of multiple strokes, (iii) an absence of a history of psychological or neurological disease other than stroke (or seizure disorder for the lobectomy participants), (iv) native English speaker, (v) right-handed prestroke, and (v)



Figure 1. Overlap map of the areas of damage in the 113 participants with a left hemisphere stroke.

ability to follow task instructions. The temporal lobectomy patients all had a seizure disorder that required lobectomy surgery for the treatment of their seizures.¹ All stroke participants had ischemic strokes, with the exception of two participants in the left hemisphere group with hemorrhagic strokes.

Aphasia type was assessed in the stroke and lobectomy groups via site-specific protocols that included the Western Aphasia Battery, Boston Diagnostic Aphasia Examination, and clinical ratings. The left hemisphere stroke group consisted of the following aphasia diagnoses: Broca's (n =30), anomic (n = 15), conduction (n = 9), residual aphasia (n = 8), Wernicke's (n = 7), transcortical motor (n = 5), mixed nonfluent (n = 5), global (n = 3), mild Broca's/anomic (n = 2), and no aphasia (n = 29). None of the participants in the lobectomy groups or in the right hemisphere stroke group were diagnosed with aphasia.

Given the proposed role of the right mid-posterior STG in speech perception, the right hemisphere stroke patients were divided into two groups: those with right mid-posterior STG damage (n = 11) and those with no mid-posterior STG damage (n = 8; max overlap is 3 of 8 participants in portions of the inferior frontal gyrus, precentral gyrus, and insula; Figure 2A, B). Mid-posterior STG was defined as any gray or white damage in the STG adjacent or posterior to Heschl's gyrus as determined by visual inspection by one of the authors (Rogalsky) with extensive training in cortical neuroanatomy. The area of maximum lesion overlap for the right



Figure 2. (A, B) Overlap map of the areas of damage in the participants with a right hemisphere stroke (A) without and (B) with mid-posterior STG damage. (C, D) Overlap map of the areas resected in the participants who underwent a (C) left or a (D) right temporal resection.

	п	Age (Years)	Sex (M/F)	Education (Years)	Time Post (Years)
Left Hemi. Stroke	113	61 (31–86)	67/46	15 (8–20)	6.1 (0.5–32)
Right Hemi. Stroke	19	52 (25–75)	13/6	15 (10-20)	11 (0.5–35)
Left Lobectomy	18	48 (29-62)	8/10	13.5 (10–16)	4 (0.7–14)
Right Lobectomy	8	41 (23-60)	3/5	14 (12–18)	9 (4–18)
Control	26	59 (12–16)	5/21	15 (12–16)	n/a

Table 1. Demographic Information for Each Participant Group

Note that "time post" refers to the number of years since a stroke for the left hemisphere and right hemisphere stroke groups, and refers to the number of years postresection for the left lobectomy and right lobectomy groups.

hemisphere (RH) stroke participants with mid-posterior STG damage is located in white matter underlying the posterior temporal lobe more generally.

Control Participants

Twenty-six adults were recruited and tested at San Diego State University. All control participants were native English speakers, right-handed, and self-reported no history of psychological or neurological disease. See Table 1 for demographic information.

Informed consent was obtained from each participant prior to participation in the study, and all procedures were approved by the institutional review boards of UC Irvine, University of South Carolina, San Diego State University, Arizona State University, Medical College of Wisconsin, and University of Iowa.

Standard hearing and vision screenings were performed during the recruitment of all participants to ensure adequate hearing and vision abilities to complete the experimental tasks and aphasia assessments. The exact hearing and vision screenings varied across sites, but also all participants were provided with several practice trials for each task during which presentation volume could be adjusted to each participant's preferences.

Materials

This study focuses on two types of tasks: auditory word comprehension and syllable discrimination. The tasks were administered as part of an extensive psycholinguistic test battery to assess receptive and productive speech abilities. Within the battery, individual tests themselves were presented in a nonfixed pseudorandom order; the order and proximity within the test battery of the tasks described below was random across participants. Items within each test described below were presented in a fixed random order at all testing sites, except for Medical College Wisconsin (n = 24), where items were presented in a random order for each participant.

Auditory Comprehension Measures

Auditory word comprehension in a clear background. A four-alternative forced-choice word-to-picture matching task previously used by Rogalsky, Rong, Saberi, and Hickok (2011) and similar to Baker, Blumstein, and Goodglass' (1981; Blumstein, Baker, & Goodglass, 1977) paradigm was administered. The task contained 20 trials. In each trial, an auditory word was presented 1 sec after the appearance of a picture array depicting four objects on a computer screen. The words were recorded by a male native speaker of American English. The participant was instructed to point to the picture that matched the auditorily presented word. The picture array contained the target and three distractors. Ten of the 20 trials included a one-syllable target word and its matching picture, as well as a picture representing the phonemic distractor that was a minimal pair to the target (4 place of articulation contrasts, 4 voicing contrasts, and 2 manner contrasts), a semantic distractor, and an unrelated distractor, which was semantically related to the phonological distractor. For example, the target goat was presented with distractor images of a coat, a pig, and a shoe. The auditory stimuli were counterbalanced such that each picture in each array was presented once as the target. All of the words presented have an estimated average age of acquisition under 6 years old (range: 2.6-5.7 years old; Kuperman, Stadthagen-Gonzalez, & Brysbaert, 2012) and ranged in frequency from 1.33 to 86.86 per million words (Brysbaert & New, 2009). The frequency of each error type was calculated, as well as proportion correct for each participant's overall performance in each task.

Auditory word comprehension in noise. The auditory comprehension in noise task contains the same word and picture array pairings as the auditory comprehension word–picture matching task described above, except that the auditory stimuli were presented in 14-dB Gaussian white noise. The noise began 1 sec before the onset of the word and ended 1 sec after the end of the word.

Syllable Discrimination Measures

We employed two syllable discrimination measures: a real word discrimination task and a nonword discrimination task. These two syllable discrimination tasks were modeled on previous discrimination tasks used with individuals with aphasia (Caplan & Waters, 1995; Baker et al., 1981; Blumstein et al., 1977). Each of these discrimination tasks contained 40 trials. The real word discrimination task contained pairs of one-syllable real words, and the nonword discrimination task contained one-syllable nonwords, with neighborhood density and phonotactic probability between the real words and nonwords matched. The Irvine Phonotactic Online Dictionary (Vaden, Halpin, & Hickok, 2009) was used to select the real words and nonwords, and to calculate neighborhood density and phonotactic probability for each word and nonword. Each word and its corresponding nonword have the same onset consonant phoneme. Paired-samples t tests comparing each word with its matched nonword indicate that there is no significant difference between the two stimulus types for density, t(39) = 0.14, p = .88, or phonotactic probability, t(39) = 0.18, p = .86. The words used in the real word discrimination task do not overlap with the words in the auditory comprehension tasks described above, but the minimal pairs in the syllable discrimination tasks are the same minimal pairs present in the auditory word comprehension tasks.

The real word and nonword discrimination tasks both were structured in the following manner: There were 40 trials in each task, and in each trial, a pair of onesyllable words (or nonwords) were presented via headphones while a fixation cross was presented on the computer screen. Each trial presented the two words (or nonwords) in one of four arrangements: A-B, B-A, A-A, and B-B such that, for half of the trials, the correct answer was "same" and, for the other half, the correct answer was "different." In the "same" trials, two different tokens of the same syllable were presented. The same male native English speaker of the words in the comprehension task described above recorded all of the stimuli. There was a 1-sec interval between each word (or nonword) in a pair, and 5 sec between each pair (more time between pairs was given, if necessary, for the patient's comfort). Patients were instructed to determine if the two words (or nonwords) presented were the same or different. "Different" trials contained two syllables that differed by one feature of the onset consonant (e.g., "puff" vs. "cuff" in the real word discrimination task, or "pag" vs. "kag" in the nonword discrimination task). Participants made their response either by speaking their answer or by pointing to the correct answer written in large print on a paper adjacent to the computer screen. Signal detection methods were used to determine how well participants could discriminate between the same and different pairs by calculating the measure d' for the real word and nonword discrimination tasks, respectively (Swets, 1964).

Neuroimaging

All stroke and lobectomy participants (n = 158) underwent MRI scanning using a 3-T or 1.5-T MRI system at the respective testing site. T1-MRIs and T2-MRIs with 1-mm³ resolution were collected and used to demarcate the lesion; the lesion demarcation was conducted manually by welltrained and experienced individuals in lesion-mapping studies. The lesion maps were smoothed with a 3-mm full-width half-maximum Gaussian kernel to remove jagged edges associated with manual drawing. Enantiomorphic normalization (Nachev, Coulthard, Jäger, Kennard, & Husain, 2008) was conducted using SPM12 and in-house MATLAB scripts in the following way: A mirrored image of the T1 image (reflected around the midline) was coregistered to the native T1 image. Then, we created a chimeric image based on the native T1 image with the lesioned tissue replaced by tissue from the mirrored image (using the smoothed lesion map to modulate this blending, feathering the lesion edge). SPM12's unified segmentationnormalization (Ashburner & Friston, 2005) was used to warp this chimeric image to standard space, with the resulting spatial transform applied to the actual T1 image as well as the lesion map. The normalized lesion map was then binarized, using a 50% probability threshold.

Lesion-symptom Mapping

All of the lesion-symptom mapping (LSM) routines used here are integrated into the NiiStat toolbox for MATLAB (https://www.nitrc.org/projects/niistat). The following routines were performed: Univariate analyses fitting a general linear model (GLM) were completed to identify anatomical regions in which the percentage of lesioned voxels was associated with performance on the auditory comprehension in noise, word discrimination, or nonword discrimination tasks. A GLM was fit for performance on each task within each of the 94 left hemisphere ROIs defined by the John's Hopkins University (JHU) atlas (den Ouden et al., 2019; Faria et al., 2012; Mori et al., 2008). An ROI approach, compared with voxel-based statistics, increases statistical power by reducing the severity of multiple comparison correction (Rorden, Fridriksson, & Karnath, 2009). Univariate analyses were conducted for the following variables: auditory word comprehension in noise (proportion correct), word discrimination (d'), and nonword discrimination (d'). Clear speech auditory word comprehension was not used for LSM because of a ceiling effect with insufficient variability; we discuss this further below. The behavioral data were significantly skewed (i.e., Z-skew >1.96); thus, deskewing was performed in NiiStat prior input into the GLM by computing the standard error of the skew and applying a square root transform to the data.

Permutation thresholding included 4000 permutations to correct for multiple comparisons (p < .05 controlled for familywise error). As ROIs that are infrequently damaged will have low statistical power while increasing the number of comparisons, only ROIs where at least 11 participants (i.e., approximately 10% of the largest left hemisphere patient sample size in a task) had damage were included in the analyses.

Variability because of overall lesion size was regressed out of each LSM. Data collection site also was used as a covariate to remove variance accounted for by collection site, including potential variance because of site-specific differences in scanner properties and protocols. All reported coordinates are in Montreal Neurological Institute (MNI) space. Figure 3 displays the extent of coverage of the ROIs included in the LSM analyses; the coverage does not vary between the analyses.

Post Hoc Comparisons of LSMs

Based on the close proximity and partial overlap of the significant ROIs identified by the LSMs described above, we decided to use conjunction and independence tests to further investigate the relationships between (1) the brain regions supporting real word discrimination versus those regions supporting nonword discrimination, and (2) the brain regions supporting nonword discrimination versus auditory comprehension in noise. The conjunction tests were computed in each ROI using an in-house MATLAB script following the valid conjunction test procedure recommended by Nichols, Brett, Andersson, Wager, and Poline (2005). Code for the conjunction analysis can be found here: https://raw.githubusercontent.com /rordenlab/spmScripts/master/nii_thresh_conjunction.m. This conjunction analysis procedure generates, in each Figure 3. Map of the extent of coverage by the ROIs from the JHU atlas included in the LSM analyses of the left hemisphere group. Note the ample coverage of the superior and middle portions of the temporal lobe, as well as premotor, inferior frontal, and primary motor cortex.



ROI, the *p* value of the *z* values under the conjunction null, that is, the union of the null hypotheses for the two tasks' *z* scores. ROIs with *p* < .05 indicate that the damage in that ROI is associated with lower performance in both tasks. The independence tests were computed in each ROI using a custom contrast in the nii_stat toolbox (e.g., 1 – 1) to determine if damage in each ROI is significantly more related to one task than the other; in other words, if an ROI's *z* score for one task is significantly different than the *z* score for another task (*p* < .05, Freedman-Lane permutation as described by Winkler, Ridgway, Webster, Smith, & Nichols, 2014).

Because of a combination of time limitations and site-specific testing protocols, not all participants completed all speech perception tasks. Table 1 indicates the sample sizes for each task within each participant group. Within each lesion group, there are no qualitative differences in the lesion distributions or lesion sizes of the participants who did and did not complete each task.

RESULTS

Overall Behavioral Results

A summary of the descriptive statistics for each group on each of the tasks is presented in Table 2.

Auditory Word Comprehension Tasks

All groups were at ceiling on the clear auditory word comprehension task. In the left hemisphere (LH) stroke group, the mean performance was 97% correct (Figure 4) with both the left (L) and right (R) anterior temporal lobectomy groups scoring at 100% correct. More variance in performance was found on the auditory word comprehension in noise task, enabling statistical analyses. A one-way ANOVA across the five groups (control, LH stroke, RH stroke, L lobectomy, R lobectomy) revealed a significant Group effect in performance on the auditory word comprehension in noise task, F(4, 170) = 10.6, p < .001, with the LH stroke group differing from all other groups ($ps \le$.002, Bonferroni correction of $\alpha = .05/10 = .005$) and no other group differences (Figure 5, left panel).

Syllable Discrimination Tasks

A 5 × 2 mixed ANOVA was computed for the performance on the two syllable discrimination tasks (i.e., word discrimination and nonword discrimination) in each of the five groups revealing significant main effects of Task, F(1, 138) = 17.80, p < .001, with higher scores on the word than nonword discrimination, and Group, F(4, 138) =9.20, p < .001. The interaction was not significant, F(4, 138) =0.38, p = .83. The LH stroke group performed

	Control	LH Stroke	RH Stroke	Left ATL	Right ATL
Clear Auditory	Word Comprehension				
n	21	109	19	18	8
Mean	1	0.97	0.99	1	1
Median	1	1	1	1	1
SD	0	0.08	0.03	0	0
Range	1–1	0.4–1	0.9–1	1–1	1–1
Auditory Word	l Comprebension in Noi	ise			
n	22	109	19	17	8
Mean	0.91	0.78	0.88	0.87	0.91
Median	0.90	0.80	0.90	0.85	0.90
SD	0.05	0.14	0.05	0.07	0.05
Range	0.85-1	0.3–1	0.8–0.95	0.75–1	0.85–1
Word Discrim	ination				
n	25	78	18	17	8
Mean	5.05	4.20	4.72	5.02	5.15
Median	5.15	4.22	5.15	5.15	5.15
SD	0.29	1.06	0.78	0.38	0
Range	4.22–5.15	1.71–5.15	2.68-5.15	3.86–5.15	5.15-5.15
Nonword Disc	rimination				
n	23	77	18	17	8
Mean	4.79	3.75	4.29	4.58	4.76
Median	5.15	4.22	4.22	5.15	5.15
SD	0.57	1.27	1.02	0.86	0.56
Range	3.61-5.15	0-5.15	2.07-5.15	2.68-5.15	3.86-5.15

Table 2. Descriptive Statistics for the Clear Auditory Word Comprehension (Proportion Correct), Auditory Word Comprehension in Noise (Proportion Correct), Word Discrimination (*d*-prime Values), and Nonword Discrimination (*d*-prime Values) Tasks

significantly worse than the control, L lobectomy, and R lobectomy groups ($ps \le .002$, Bonferroni correction of $\alpha = .05/10 = .005$) with no other significant group differences. The LH stroke group numerically performed worse than the RH stroke group across both syllable discrimination tasks, but the comparison did not withstand multiple comparison correction (p = .015; Figure 5, right panel).

As noted in the Introduction, previous observations indicated that performance on auditory word comprehension and syllable discrimination tasks are doubly dissociable. We assessed this in our sample by categorizing performance as impaired or spared on the auditory word comprehension in the noise task and the word discrimination task. "Impaired" was defined as performance greater than 2 *SD*s below the control mean; "spared" was defined as performance that is within or above .5 *SD*s of the control mean. Seventy-five LH stroke participants completed both of these tasks. Table 3 presents participant counts that fall into the four possible spared-impaired categories on the two tasks. It is apparent that these two tasks are dissociable, with 13 cases (17.3%) exhibiting impaired performance on the auditory word comprehension task but spared performance on the word discrimination task, and 10 cases (13.3%) showing the reverse pattern. There are fewer cases with spared auditory word comprehension in noise in the face of impaired word discrimination but recall that performance on clear word comprehension is at ceiling for the majority of LH stroke participants. There



Figure 4. (A) Distribution of performance of the left hemisphere stroke group on the clear auditory comprehension task. Numbers in parentheses indicate the actual number of left hemisphere stroke participants who performed at that accuracy level (out of a total 109 who completed the task). Note that chance performance would be .25 proportion correct (not shown). (B) Distribution of performance of the left hemisphere stroke group on the clear auditory comprehension task as a function of word discrimination (top) and nonword discrimination (bottom) task performance.

are 59 such participants who were both at ceiling on the clear auditory word comprehension task and completed the word discrimination task; 30 (50.8%) were impaired on the latter, which was also presented as clear speech. This provides substantial evidence that these tasks are dissociable.

Within the LH stroke group, correlation analyses performed between the auditory word comprehension in noise and word discrimination tasks revealed a statistically significant relationship, r(73) = .39, $r^2 = .15$, p < .001. However, it is evident from the scatterplot in Figure 6 that, consistent with the categorization-based analysis of task dissociability, continuous variation in performance on one task is a rather poor predictor of variation on the other, accounting for only 15% of the variance. A partial correlation between the auditory word comprehension in noise and word discrimination tasks controlling for lesion size was also significant, r(72) = .0.32, p = .005, suggesting that overall lesion size is not accounting for the relationship between the two tasks.

Lesion-Behavior Mapping Results

LSMs were computed within the left hemisphere participants for auditory word comprehension in noise, word discrimination, and nonword discrimination. The auditory word comprehension in noise task identified four adjacent ROIs covering much of the superior temporal lobe (Figure 7, Table 4). These ROIs included the superior and middle temporal gyri and underlying white matter, as well as extending into the inferior portion of the supramarginal gyrus; Heschl's gyrus also was implicated as it is part of the anterior superior temporal ROI (JHU atlas label = "superior temporal gyrus," z = 3.08, ROI center of mass = MNI coordinates -51 - 13 1; Figure 8A). The LSM of word discrimination identified two significant ROIs (p < .05, corrected; Table 4) adjacent to one another (JHU atlas labels "superior temporal gyrus" and "posterior temporal gyrus"; Figure 7, 8B; Table 4), implicating most of the STG, as well as Heschl's gyrus and the planum temporale, as well as an additional significant ROI, consisting of a portion of the superior longitudinal fasciculus (SLF) underlying the supramarginal gyrus (z =3.46; ROI center of mass = -35 - 24 29 MNI). The LSM of nonword discrimination identified the three ROIs implicated in real word discrimination (i.e., the two STG ROIs, and the SLF ROI), as well as an additional ROI in the supramarginal gyrus extending into the postcentral gyrus (z = 2.87, ROI center of mass = -51 - 2933 MNI), which is dorsal to the supramarginal gyrus region implicated in the auditory word comprehension in noise LSM (Figure 7, 8B). All three LSMs overlap in the STG ROI that includes mid-temporal regions including Heschl's gyrus and extends into the more posterior STG (orange, Figure 8C; Table 4).

The results for the post hoc conjunction and independence tests of the LSMs are as follows: For the word discrimination versus the nonword discrimination contrast (LSMs shown in Figures 7 and 8B), the conjunction test identified that damage in the left STG (z = 3.12), the left posterior STG (z = 3.04), and the left superior longitudinal fasciculus (z = 2.99) ROIs (i.e., all three ROIs that were separately significant for each task; Table 4) were each related to significantly lower performance in both tasks (p < .05); the independence test did not identify any ROIs as being significantly more implicated in the nonword than the word discrimination task, or vice versa (i.e., z < 2.96, p > .05 in all ROIs). For the nonword discrimination versus auditory word comprehension in noise LSMs (LSMs shown in Figure 8C), the conjunction test identified damage in the left STG ROI (z = 3.21) as being significantly related to lower performance in both tasks (p < .05); the independence test identified damage in the left supramarginal gyrus (z = 3.21), left posterior STG (z = 3.20),



Figure 5. Performance on the auditory word comprehension in noise (left) and syllable discrimination tasks (right) in each participant group. Error bars represent standard error. Overlapping points in the discrimination task graph have been offset slightly to make them visible. Ctrl = control group; ATL = anterior temporal lobectomy; Str = stroke.

	Word Discrimination		
Auditory Word Comprehension in Noise	Impaired	Spared	
Impaired	26	13	
Spared	10	12	

Impaired is defined as performance that is > 2 SDs below the control mean; spared is defined as performance that is within or above .5 SDs of the control mean.



Figure 6. Relation between the auditory word comprehension in noise and word discrimination tasks within the LH stroke group. Overlapping points have been offset slightly to make them visible.

and left superior longitudinal fasciculus ROIs (z = 3.57) as being significantly more related to lower performance for nonword discrimination than for auditory comprehension in noise, and damage in the posterior middle temporal gyrus (z = 3.84) and sagittal striatum (z = 3.91) ROIs as being significantly more related to lower performance for auditory comprehension in noise than for nonword discrimination (p < .05). In short, the findings from the post hoc conjunction and independence analysis statistically confirm the map shown in Figure 8C.

Right Hemisphere Results

To test for effects of right hemisphere damage on speech perception, we partitioned the RH stroke group into those cases with ("Temporal +") or without ("Temporal –") damage (Figure 2A, B) to the posterior superior temporal lobe and compared them to controls on the auditory word comprehension in noise task and on each syllable discrimination task. The one-way ANOVA across the three groups (RH Temporal +, RH Temporal –, and control) on the auditory word comprehension in noise task yielded a main effect of Group that did not pass the $\alpha > .05$ threshold but approached significance, F(2, 40) = 3.03, p = .061, with the superior temporal lobe damaged group performing numerically worse than those without damage there (Figure 8, left panel). A mixed ANOVA on the two syllable discrimination tasks across the same three groups



Figure 7. LSM results in the left hemisphere stroke group. Significant ROIs are displayed (p < .05, permutation-based family-wise error corrected) on sagittal slices through the left hemisphere for each of the three tasks for which LSM was computed.

Table 4. *z* Scores of Each Significant ROI in the Auditory Word Comprehension in Noise Task and the Syllable Discrimination Tasks (i.e., Word Discrimination and Nonword Discrimination)

JHU Atlas ROI Description (ROI Number and Label)	Auditory Word Comprebension in Noise	Word Discrimination	Nonword Discrimination
Left supramarginal gyrus (29 SMG_L)	_	_	2.87
Left STG (35 STG_L)	3.04	3.27	3.49
Left rentrolenticular part of internal capsule (135 RLIC_L)	3.08	—	—
Left Sagittal stratum ^a (151 SS_L)	3.67	_	_
Left superior longitudinal fasciculus ^b (155 SLF_L)	_	3.46	3.30
Left posterior STG (184 PSTG_L)	_	3.67	3.60
Left posterior middle temporal gyrus (186 PSMG_L)	3.29	—	—

^a The JHU ROI atlas notes that the sagittal stratum includes inferior longitudinal fasciculus and inferior fronto-occipital fasciculus.

^b The JHU ROI atlas denotes this ROI as being part of the SLF, but it is noteworthy that it is difficult to distinguish between SLF and arcuate fasciculus fibers in this region.

revealed a significant effect of Task, F(1, 38) = 7.4, p = .01, with nonword discrimination performance lower than word discrimination, and a significant effect of Group, F(2, 38) = 7.5, p = .002, with the RH temporal + group performing significantly worse than the controls and the

RH temporal – group (ps < .002, Bonferroni correction of $\alpha = .05/3 = .017$); there was no significant difference between the control group and the RH temporal – group (Figure 9, bottom). The Task × Group interaction was not significant, F(2, 38) = 1.5, p = .24.



Figure 8. Lateral view summary of brain regions implicated in each task in the left hemisphere stroke group, p < .05, permutation-based family-wise error corrected. (A) Left lateral view of the regions significantly implicated in auditory word comprehension in noise (B) Left lateral view of the regions significantly implicated in word and nonword discrimination. Note that the word discrimination regions are a subset of the regions implicated in nonword discrimination in noise and nonword discrimination indicated in orange.



Figure 9. Right hemisphere auditory word comprehension in noise (top) and syllable discrimination performance (bottom). Error bars represent standard error. Overlapping points in the syllable discrimination graph (bottom) have been offset slightly to make them visible.

DISCUSSION

This study sought to test four functional anatomic models of speech perception:

- 1. Classical model—speech perception is uniquely supported by the left mid-to-posterior superior temporal lobe.
- 2. Motor model—speech perception is dependent on (strong version) or augmented by (weak version) frontal motor systems involved in speech production, particularly under noisy listening conditions.
- 3. Left anterior temporal model—speech perception is dependent on the left anterior superior temporal lobe.
- 4. Bilateral temporal model—speech perception is bilaterally organized in the mid-to-posterior superior temporal lobe.

We also sought to evaluate potential behavioral and neural differences on two tasks that are often used to assess speech perception:

5. Task dependent claim—discrimination and comprehension tasks are dissociable both behaviorally and neurally.

To this end, we examined syllable discrimination and auditory word comprehension in chronic stroke, anterior temporal lobectomy, and unimpaired control participants. Consistent with prior work, we found that performance on auditory word comprehension and syllable discrimination tasks are dissociable behaviorally.² Performance on the clear auditory word comprehension task was at ceiling for the vast majority of the left hemisphere stroke group, precluding lesion analysis on that task. LSM using the auditory word comprehension in noise task as well as both syllable discrimination tasks revealed an association between each of these tasks and the left mid-to-posterior STG. Task differences were also observed, however. The syllable discrimination tasks also uniquely implicated regions in the inferior parietal lobe and planum temporale, whereas the auditory word comprehension in noise task uniquely implicated a more ventral region in the middle temporal gyrus. Lesions involving the ATL or frontal, motor-related regions were not associated with poorer performance on any of the tasks, although the superior longitudinal fasciculus, which terminates in several areas of the frontal cortex including premotor regions, was implicated in both syllable discrimination tasks. Individuals with right hemisphere temporal lobe damage exhibited significant deficits on the syllable discrimination tasks and approached significance on the auditory word comprehension in noise task. In the following sections, we discuss the implications of these findings for the specific hypotheses we aimed to test, starting with the task effects.

Task Dependence

Noting early reports of the paradoxical dissociability of auditory word comprehension tasks from syllable discrimination tasks (Miceli et al., 1980; Basso, Casati, & Vignolo, 1977; Blumstein et al., 1977), Hickok and Poeppel argued that the two tasks involve shared resources in the STG including what they termed "spectrotemporal analysis" in and around primary auditory cortex and extending into the ventral extent of the gyrus in the superior temporal sulcus, but then tapped into divergent processing streams beyond that (Hickok & Poeppel, 2007). They argued that discrimination tasks, in addition, relied on working memory systems in the motor-related dorsal stream (inferior parietal lobe, posterior planum temporale, and posterior frontal cortex; Buchsbaum & D'Esposito, 2019; Schomers, Garagnani, & Pulvermüller, 2017; Buchsbaum et al., 2011; McGettigan et al., 2011; Hickok et al., 2003), whereas auditory word comprehension tasks also relied on more ventral lexical access systems in the middle temporal gyrus region (Pillay, Binder, Humphries, Gross, & Book, 2017; Indefrey, 2011; Lau, Phillips, & Poeppel, 2008; Liebenthal, Binder, Spitzer, Possing, & Medler, 2005; Indefrey & Levelt, 2004).

Our findings are highly consistent with this hypothesis. Our study replicates previous work demonstrating the dissociability of syllable discrimination and auditory word comprehension tasks and identifies both shared and distinct regions implicated in the two tasks. The shared region involves the mid-to-posterior STG, which was implicated in both word and nonword discrimination tasks, as well as the auditory word comprehension in noise task. The two discrimination tasks, but not the comprehension in noise task, additionally implicated the planum temporal region (part of the posterior STG ROI), which includes area Spt, thought to serve as an auditory– motor interface for speech production (Hickok, 2012; Buchsbaum et al., 2011; Hickok, Okada, & Serences, 2009; Hickok et al., 2003). The nonword discrimination task uniquely implicated additional inferior parietal cortex in the supramarginal gyrus. In contrast, the auditory word comprehension in noise task, but not the discrimination tasks, implicated the posterior middle temporal gyrus, thought to be important for lexical-semantic access (Indefrey, 2011; Lau et al., 2008; Hickok & Poeppel, 2004, 2007; Indefrey & Levelt, 2004; de Zubicaray, Wilson, McMahon, & Muthiah, 2001).

Given that functional imaging and TMS have implicated frontal, motor-related regions in the performance of syllable discrimination and related tasks (Lee, Turkeltaub, Granger, & Raizada, 2012; D'Ausilio et al., 2009; Meister et al., 2007; Blumstein, Myers, & Rissman, 2005; Watkins & Paus, 2004; Jancke, Wustenberg, Scheich, & Heinze, 2002; Burton et al., 2000; Zatorre, Evans, Meyer, & Gjedde, 1992), we expected to see some frontal ROIs correlated with deficits on our discrimination task. Yet, we did not. This result does not necessarily mean that motor speech systems play no causal role at all in speech processing using some tasks and under some conditions. The TMS studies suggest otherwise as does the present finding that temporal-parietal dorsal stream regions that play an important role in speech production are implicated in syllable discrimination. Presumably, these temporal-parietal areas are interfering with discrimination task performance as a function of their network connectivity with frontal, motor-related areas. Our finding of damage to the superior longitudinal fasciculus being associated with poorer performance on both syllable discrimination tasks, but not the comprehension task, also supports view. Yet for reasons that are not entirely clear, disrupting posterior nodes in the dorsal stream sensorimotor network seems to have the dominant effect on syllable discrimination relative to damage in frontal motor speech areas (Caplan, Gow, & Makris, 1995).

What is clear is that none of these dorsal stream regions, when damaged, impair the ability to comprehend words. Further additional support for this conclusion comes from a recent study of two cases of acquired opercular syndrome, both with complete anarthria secondary to bilateral removal of most of the inferior, posterior frontal cortex. Despite being completely unable to voluntarily control their vocal tracts, both individuals were able to comprehend words at ceiling levels (Walker, Rollo, Tandon, & Hickok, 2021).

Our neuroanatomical findings for these tasks explain the behavioral patterns of association and dissociation. Overall, across the LH stroke group, word discrimination and word comprehension (in noise) are weakly but significantly correlated (Figure 6). This covariation is presumably related to shared involvement of auditory and phonological systems in the STG. The dissociability of the two tasks (Figure 6 and Table 3) is explained on the basis of the distinct pathways recruited by the two tasks beyond the STG. Damage to dorsal stream systems can impact discrimination without affecting comprehension and damage to ventral temporal lobe systems can impact comprehension without affecting discrimination.

The Classical Model of Speech Perception

The classical model of speech perception, originally proposed by Wernicke, was not supported by the present data. This model, which holds that the left STG is the primary substrate for speech perception, predicts that damage to this region should cause substantial deficits on all receptive speech tasks. This was not the case as the vast majority of individuals in our LH stroke group were at ceiling on the clear auditory word comprehension task although the task required the ability to resolve minimal phonemic pair differences. This finding not only shows that damage to classical Wernicke's area is not sufficient for producing substantial deficits in word comprehension, but more generally that damage anywhere in the left hemisphere is not sufficient. This, in turn, indicates that receptive speech up to the word level is not strongly lateralized and is more consistent with a bilateral model, which we discuss below.

The Motor Models of Speech Perception

Our study provided little support for the strong motor model, which predicts that damage to motor speech areas should cause deficits in the ability to perceive speech sounds and therefore affect both syllable discrimination and auditory word comprehension. As noted, clear auditory word comprehension was largely unaffected by unilateral stroke even with a substantial number of individuals in the LH stroke group exhibiting clinically significant motor speech deficits (e.g., Broca's aphasia). Furthermore, damage to frontal motor-speech areas was not significantly correlated with deficits on any of our tasks. As pointed out above, the motor model does gain partial support in that damage to temporal-parietal dorsal stream regions, which are part of the speech production network, are implicated in performance on discrimination tasks. However, these effects do not generalize to the more ecologically valid comprehension tasks.

Weaker versions of the motor model also did not gain support from our study. These models argue that motor speech systems play an augmentative role in speech perception under noisy listening conditions (Wilson, 2009; Moineau et al., 2005). We studied auditory word comprehension in noise and found that damage to superior temporal and middle temporal regions was correlated with deficits; dorsal stream regions were not implicated. This finding contradicts a previous, smaller-scale study that reported that word comprehension suffered with degraded speech in people with Broca's aphasia, who typically have damage to frontal, motor-related speech areas (Moineau et al., 2005). There are two difficulties interpreting this previous study, however. One is that Broca's aphasia is not caused solely by damage to frontal cortex and indeed regularly also involves damage to classical Wernicke's area in the mid and posterior temporal lobe (Fridriksson et al., 2015), that is, regions that our study linked to deficits on the auditory word comprehension in noise task. Thus, it could be that it was damage to Wernicke's area rather than motor speech areas that was the source of the effect reported by Moineau et al. The second difficulty is that the behavioral analysis did not take response bias into account. The task was to decide whether a word-picture pair matched or did not. Performance on such a task is affected not only by perceptual ability but also response bias. Previous work using fMRI (Venezia et al., 2012) and lesion (Rogalsky et al., 2018) methods have linked motor speech areas to changes in response bias. Moineau et al. report significant response bias effects in their data, yet report only percent correct measures rather than *d*-prime statistics, which would correct for bias (Green & Swets, 1966). Thus, even if the lesion distribution in Broca's aphasia was not a confound, it is unclear if Moineau et al.'s observed effects are because of perceptual effects or changes in response bias. The data from the present, larger-scale study that includes a direct analysis of lesion patterns suggest that one or both of the confounds in the Moineau et al. study impacted their findings.

One could argue that our study did not include the full range of tasks that have been shown to implicate frontal motor-related systems in speech processing. For example, Burton et al. (2000) showed that discriminating the onset phoneme in syllable pairs such as *dip-tip* did not consistently yield frontal activation, whereas in pairs such as *dip-ten*, it did, which they argued was because of explicit phoneme segmentation (similar to a phonological awareness task) or working memory. Our task used stimuli more similar to *dip-tip*, leaving open the possibility that a different task requiring explicit segmentation or working memory may recruit frontal motor-related systems. However, such a possibility does not weaken our claim that speech perception as it is employed under more natural conditions does not substantively involve the frontal motor speech system. Rather, it would strengthen the view that motor involvement is an artifact of tasks that require motor-related processes, such as working memory (Hickok & Poeppel, 2000, 2004, 2007).

The Anterior Temporal Model of Speech Perception

The anterior temporal model predicts that damage to the left ATL should have a substantial impact on the ability to comprehend speech. Our findings provide partial support for this model in that, for all tasks, lesions implicated midposterior as well as mid-anterior STG regions. The left superior temporal gyrus ROI that was implicated in all tasks includes mid-anterior STG and extends anteriorly such that it nearly included the coordinates for fMRI



Figure 10. Red: Significant ROIs identified by the LSM analyses for the auditory word comprehension in noise task; green: The left lobectomy group's lesion maps (areas shown indicate lesion in at least 2 participants); yellow: overlap between the auditory word comprehension in noise ROIs and the left lobectomy lesion maps. Blue circles indicate the five anterior temporal coordinates identified by Scott et al. (2000; MNI: -546-16, -66-12-12) or Evans et al. (2014; MNI: -582-16, -60-2-12, -584-20) as exhibiting main effects of speech intelligibility or a similar contrast.

activations in the ATL that have been claimed to be the foci involved in intelligible speech processing (Figure 10), as revealed in studies of intelligible versus unintelligible sentences (Evans et al., 2014; Narain et al., 2003; Scott et al., 2000).

These fMRI activation foci largely fall within the region impacted by this study's left hemisphere ATL resection cases, which as a group scored lower than controls and similar to the group average for individuals with right posterior temporal strokes, but quite a bit better than the left hemisphere stroke group. This suggests a role for the left ATL in speech perception, but does not support the claim that the network is strongly left dominant (Rauschecker & Scott, 2009; Scott et al., 2000) as more severe deficits should have been apparent with left hemisphere damage, even on the clear auditory word comprehension task if the network was strongly left dominant. Furthermore, in the context of the ATL model, it has been claimed that posterior superior temporal regions primarily support auditorymotor functions, not intelligibility (Rauschecker & Scott, 2009), thus predicting that posterior temporal damage should not be implicated in word comprehension. Our findings suggest otherwise and argue that the "pathway for intelligible speech" involves much of the STG and extends into the posterior middle temporal gyrus consistent with higher-powered functional imaging studies on intelligible speech (Evans et al., 2014; Okada et al., 2010).

The Bilateral Temporal Model of Speech Perception

The bilateral temporal model predicts that speech perception as measured by auditory word comprehension will not be substantially impaired by unilateral damage to either hemisphere and that any mild deficits will be associated with superior temporal lobe damage. This is precisely the result we found. Clear speech word comprehension was at or near ceiling in the vast majority of all people studied, and poorer performance on the auditory word comprehension in noise task was associated with damage to the STG and posterior middle temporal gyrus. This was true in the left hemisphere where we had a sufficient sample size to perform LSM; A nonsignificant trend (p = .06) in the same direction was found as well in the 11 participants with right hemisphere mid-posterior superior temporal lobe lesions.

Previous authors have argued for a bilateral capacity for receptive speech at the word level based on data from split brain studies, Wada procedures, and stroke (Hickok et al., 2008; Hickok & Poeppel, 2000, 2004, 2007; Goodglass, 1993; Bachman & Albert, 1988; Buchman et al., 1986; Zaidel, 1985; Gazzaniga & Sperry, 1967). Until recently, however, direct evidence has been lacking for an association between speech perception impairment and unilateral right hemisphere damage, although higher-level language deficits have been documented (Gajardo-Vidal et al., 2018). This study provides such evidence from stroke and joins a recent TMS study reporting similar effects when left or right superior temporal regions were interrupted in healthy participants (Kennedy-Higgins, Devlin, Nuttall, & Adank, 2020).

A recent large-scale lesion study using a clinical assessment tool (Western Aphasia Battery-Revised) reported that unilateral left hemisphere stroke can cause single-word comprehension deficits (in clear speech), which map onto left temporal lobe regions (Bonilha et al., 2017). However, scores on this task include subtests that assess aspects of reading (matching spoken words to printed letters and numbers) as well as shapes, colors, and body parts in addition to more common objects. In light of the high level of performance on this study 's clear auditory word comprehension task, it is likely that clinical assessment tools tap into a broader range of abilities that go beyond basic phonemic perception and lexical-semantic access, which is what our tasks were designed to test.

Although group performance was exceptionally high on the clear auditory word comprehension task (mean = 97%) with 95.4% of our sample performing at or better than 90% correct, there were a handful of cases (\sim 5%) who were substantially impaired. How should these cases be interpreted with respect to claims about the lateralization pattern for auditory word comprehension? We suggest that these cases likely reflect anomalous dominance, that is, cases where auditory word comprehension is atypically dependent on the left hemisphere. In general, the incidence of anomalous cerebral lateralization for language in strong right handers is in that same range with estimates at approximately 5% (Knecht et al., 2000) and functional brain imaging work suggests the incidence of almost complete left dominance is itself atypical and with a frequency of approximately 5% (Springer et al., 1999). One might counter that because aphasia was not an inclusion criterion at all of our testing sites, the actual incidence of word comprehension impairments following a left hemisphere stroke might be underestimated in our sample. Yet even at sites that recruited only people with aphasia, only 4 of 63 (6.3%) performed worse than 90% correct on the clear word comprehension test. Moreover, the reverse sampling bias argument can be made: Recruiting participants on the basis of the presence of aphasia following unilateral stroke likely overestimates left hemisphere dominance for language because people with naturally more bilateral organization will be less likely to present with aphasia following unilateral stroke.

A final potential concern regarding the present data for the bilateral model is whether plastic reorganization is responsible for the well-preserved word comprehension ability. That is, perhaps performance is at ceiling because the right hemisphere is taking on speech recognition function after left hemisphere insult. However, data from a large sample of acute stroke (Rogalsky, Pitz, Hillis, & Hickok, 2008) as well as a smaller sample of acute left hemisphere deactivation in a Wada study (Hickok et al., 2008) show similar effects.

Although our findings strongly support the bilateral model, this does not necessarily imply symmetry of function (Morillon et al., 2010; Hickok & Poeppel, 2007; Poeppel, 2003; Zatorre, Belin, & Penhune, 2002). Indeed, we found greater impairment following left compared with right hemisphere damage on our auditory word comprehension in noise task and our discrimination tasks. This could reflect left–right computational differences within the speech perception systems (Albouy et al., 2020; Zatorre et al., 2002; Poeppel, 2001) or their interaction with other more lateralized networks—lexical-semantic, sensorimotor, attention—that may augment the efficiency of the left hemisphere for processing speech for some tasks.

Summary

In summary, the findings from this study support a neuroanatomical model of basic auditory word identification and comprehension in which the critical regions include much of the superior temporal lobe, bilaterally: a hybrid of the anterior temporal and bilateral models discussed above. This work also confirmed the dissociability, both behaviorally and anatomically, of comprehension- and discrimination-based auditory speech tasks. Given that a large of number of studies of "speech perception" use discrimination-type tasks, particularly those that implicate dorsal, motor-related networks (D'Ausilio et al., 2009; Mottonen & Watkins, 2009; Meister et al., 2007), the present finding calls for a critical re-evaluation of that body of work.

Reprint requests should be sent to Corianne Rogalsky, Department of Speech and Hearing Science, Arizona State University, P.O. Box 870102, Tempe, AZ 85287-0102, or via e-mail: Corianne.rogalsky@asu.edu.

Funding Information

Gregory Hickok, National Institute on Deafness and Other Communication Disorders (https://dx.doi.org/10.13039 /100000055), grant number: R01 DC009659. Alexandra Basilakos, National Institute on Deafness and Other Communication Disorders (https://dx.doi.org/10.13039 /100000055), grant number: T32 DC0014435.

Diversity in Citation Practices

Retrospective analysis of the citations in every article published in this journal from 2010 to 2021 reveals a persistent pattern of gender imbalance: Although the proportions of authorship teams (categorized by estimated gender identification of first author/last author) publishing in the *Journal of Cognitive Neuroscience (JoCN)* during this period were M(an)/M = .407, W(oman)/M = .32, M/W = .115, and W/W = .159, the comparable proportions for the articles that these authorship teams cited were M/M = .549, W/M = .257, M/W = .109, and W/W = .085 (Postle and Fulvio, *JoCN*, 34:1, pp. 1–3). Consequently, *JoCN* encourages all authors to consider gender balance explicitly when selecting which articles to cite and gives them the opportunity to report their article's gender citation balance.

Notes

1. The participants with temporal lobectomies have lesions localized in the anterior temporal lobe (Figure 2), a region that is often left intact by all but the most severe middle cerebral artery strokes (Holland & Lambdon Ralph, 2010) and therefore of particular value to this study. However, there is evidence that epileptic brains have variable functional organization, including laterality of speech processing (Swanson, Sabsevitz, Hammeke, & Binder, 2007; Hertz-Pannier et al., 2002; Swanson et al. 2002). Thus, findings in this population should be interpreted carefully.

2. One possible explanation of these tasks being dissociable might be general task difficulty differences, but there are two strong pieces of evidence against a general task difficulty effect and indicating a speech perception-related effect: (1) The ROIs identified are in auditory-related areas and not frontal regions typically implicated in task demands, and (2) the vast majority of the errors induced by adding noise were phonemic errors (i.e., selecting the phonemic foil picture).

REFERENCES

- Albouy, P., Benjamin, L., Morillon, B., & Zatorre, R. J. (2020). Distinct sensitivity to spectrotemporal modulation supports brain asymmetry for speech and melody. *Science*, *3*67, 1043–1047. https://doi.org/10.1126/science.aaz3468, PubMed: 32108113
- Ashburner, J., & Friston, K. J. (2005). Unified segmentation. *Neuroimage*, *26*, 839–851. https://doi.org/10.1016/j .neuroimage.2005.02.018, PubMed: 15955494
- Bachman, D. L., & Albert, M. L. (1988). Auditory comprehension in aphasia. In F. Boller & J. Grafman (Eds.), *Handbook of neuropsychology* (Vol. 1, pp. 281–306). New York: Elsevier.
- Bak, T. H., & Hodges, J. R. (2004). The effects of motor neurone disease on language: Further evidence. *Brain and Language*, 89, 354–361. https://doi.org/10.1016/S0093-934X(03)00357-2, PubMed: 15068918
- Baker, E., Blumstein, S. E., & Goodglass, H. (1981). Interaction between phonological and semantic factors in auditory comprehension. *Neuropsychologia*, 19, 1–15. https://doi.org /10.1016/0028-3932(81)90039-7, PubMed: 7231654
- Basso, A., Casati, G., & Vignolo, L. A. (1977). Phonemic identification defects in aphasia. *Cortex*, *13*, 85–95. https://doi .org/10.1016/S0010-9452(77)80057-9, PubMed: 844311
- Binder, J. R., Frost, J. A., Hammeke, T. A., Bellgowan, P. S., Springer, J. A., Kaufman, J. N., et al. (2000). Human temporal lobe activation by speech and nonspeech sounds. *Cerebral Cortex*, 10, 512–528. https://doi.org/10.1093/cercor/10.5.512, PubMed: 10847601
- Bishop, D. V., Brown, B. B., & Robson, J. (1990). The relationship between phoneme discrimination, speech production, and language comprehension in cerebral-palsied individuals. *Journal of Speech and Hearing Research*, 33, 210–219. https://doi.org/10.1044/jshr.3302.210, PubMed: 2359262
- Blumstein, S. E. (1998). Phonological aspects of aphasia. In M. T. Sarno (Ed.), *Acquired aphasia* (3rd ed., pp. 157–185). San Diego: Academic Press. https://doi.org/10.1016/B978 -012619322-0/50008-7
- Blumstein, S. E., Baker, E., & Goodglass, H. (1977). Phonological factors in auditory comprehension in aphasia. *Neuropsychologia*, 15, 19–30. https://doi.org/10.1016/0028 -3932(77)90111-7, PubMed: 831150
- Blumstein, S. E., Myers, E. B., & Rissman, J. (2005). The perception of voice onset time: An fMRI investigation of phonetic category structure. *Journal of Cognitive Neuroscience*, 17, 1353–1366. https://doi.org/10.1162 /0898929054985473, PubMed: 16197689
- Bonilha, L., Hillis, A. E., Hickok, G., den Ouden, D.-B., Rorden, C., & Fridriksson, J. (2017). Temporal lobe networks supporting the comprehension of spoken words. *Brain*, 140, 2370–2380. https://doi.org/10.1093/brain/awx169, PubMed: 29050387
- Brennan, J., & Pylkkänen, L. (2012). The time-course and spatial distribution of brain activity associated with sentence processing. *Neuroimage*, 60, 1139–1148. https://doi.org/10.1016/j.neuroimage.2012.01.030, PubMed: 22248581
- Brysbaert, M., & New, B. (2009). Moving beyond Kučera and Francis: A critical evaluation of current word frequency norms and the introduction of a new and improved word frequency measure for American English. *Behavior Research Methods*, 41, 977–990. https://doi.org/10.3758/BRM.41.4.977, PubMed: 19897807
- Buchman, A. S., Garron, D. C., Trost-Cardamone, J. E., Wichter, M. D., & Schwartz, M. (1986). Word deafness: One hundred years later. *Journal of Neurology, Neurosurgery, and*

Psychiatry, 49, 489–499. https://doi.org/10.1136/jnnp.49.5 .489, PubMed: 2423648

- Buchsbaum, B. R., Baldo, J., Okada, K., Berman, K. F., Dronkers, N., D'Esposito, M., et al. (2011). Conduction aphasia, sensory-motor integration, and phonological short-term memory—An aggregate analysis of lesion and fMRI data. *Brain and Language*, *119*, 119–128. https://doi.org/10.1016/j .bandl.2010.12.001, PubMed: 21256582
- Buchsbaum, B. R., & D'Esposito, M. (2019). A sensorimotor view of verbal working memory. *Cortex*, 112, 134–148. https://doi.org/10.1016/j.cortex.2018.11.010, PubMed: 30639088
- Buchsbaum, B. R., Hickok, G., & Humphries, C. (2001). Role of left posterior superior temporal gyrus in phonological processing for speech perception and production. *Cognitive Science*, 25, 663–678. https://doi.org/10.1207 /s15516709cog2505_2
- Burton, M. W., Small, S., & Blumstein, S. E. (2000). The role of segmentation in phonological processing: An fMRI investigation. *Journal of Cognitive Neuroscience*, 12, 679–690. https://doi.org/10.1162/089892900562309, PubMed: 10936919
- Caplan, D., Gow, D., & Makris, N. (1995). Analysis of lesions by MRI in stroke patients with acoustic-phonetic processing deficits. *Neurology*, 45, 293–298. https://doi.org/10.1212 /WNL.45.2.293, PubMed: 7854528
- Caplan, D., & Waters, G. S. (1995). On the nature of the phonological output planning processes involved in verbal rehearsal: Evidence from aphasia. *Brain and Language*, 48, 191–220. https://doi.org/0.1006/brln.1995.1009, PubMed: 7728516
- D'Ausilio, A., Pulvermuller, F., Salmas, P., Bufalari, I., Begliomini, C., & Fadiga, L. (2009). The motor somatotopy of speech perception. *Current Biology*, *19*, 381–385. https://doi .org/10.1016/j.cub.2009.01.017, PubMed: 19217297
- den Ouden, D.-B., Malyutina, S., Basilaskos, A., Bonilha, L., Gleichgerrcht, E., Yourganov, G., et al. (2019). Cortical and structural-connectivity damage correlated with impaired syntactic processing in aphasia. *Human Brain Mapping*, 40, 2153–2173. https://doi.org/10.1002/hbm.24514, PubMed: 30666767
- DeWitt, I., & Rauschecker, J. P. (2012). Phoneme and word recognition in the auditory ventral stream. *Proceedings* of the National Academy of Sciences, U.S.A., 109, E505–E514. https://doi.org/10.1073/pnas.1113427109, PubMed: 22308358
- DeWitt, I., & Rauschecker, J. P. (2016). Convergent evidence for the causal involvement of anterior superior temporal gyrus in auditory single-word comprehension. *Cortex*, 77, 164–166. https://doi.org/10.1016/j.cortex.2015.08.016, PubMed: 26387007
- de Zubicaray, G. I., Wilson, S. J., McMahon, K. L., & Muthiah, S. (2001). The semantic interference effect in the picture–word paradigm: An event-related fMRI study employing overt responses. *Human Brain Mapping*, *14*, 218–227. https://doi.org/10.1002/hbm.1054, PubMed: 11668653
- Eimas, P. D., Siqueland, E. R., Jusczyk, P., & Vigorito, J. (1971). Speech perception in infants. *Science*, 171, 303–306. https:// doi.org/10.1126/science.171.3968.303, PubMed: 5538846
- Evans, S., Kyong, J. S., Rosen, S., Golestani, N., Warren, J. E., McGettigan, C., et al. (2014). The pathways for intelligible speech: Multivariate and univariate perspectives. *Cerebral Cortex*, 24, 2350–2361. https://doi.org/10.1093/cercor/bht083, PubMed: 23585519
- Faria, A. V., Joel, S. E., Zhang, Y., Oishi, K., van Zjil, P. C. M., Miller, M. I., et al. (2012). Atlas-based analysis of resting-state functional connectivity: Evaluation for reproducibility and multi-modal anatomy-function correlation studies.

Neuroimage, *61*, *613–621*. https://doi.org/10.1016/j .neuroimage.2012.03.078, PubMed: 22498656

Fridriksson, J., Fillmore, P., Guo, D., & Rorden, C. (2015). Chronic Broca's aphasia is caused by damage to Broca's and Wernicke's areas. *Cerebral Cortex*, 25, 4689–4696. https://doi .org/10.1093/cercor/bhu152, PubMed: 25016386

Gajardo-Vidal, A., Lorca-Puls, D. L., Hope, T. M. H., Parker Jones, O., Seghier, M. L., Prejawa, S., et al. (2018). How right hemisphere damage after stroke can impair speech comprehension. *Brain*, *141*, 3389–3404. https://doi.org/10 .1093/brain/awy270, PubMed: 30418586

Gallese, V., Fadiga, L., Fogassi, L., & Rizzolatti, G. (1996). Action recognition in the premotor cortex. *Brain*, *119*, 593–609. https://doi.org/10.1093/brain/119.2.593, PubMed: 8800951

Gazzaniga, M. S., & Sperry, R. W. (1967). Language after section of the cerebral commissures. *Brain*, *90*, 131–148. https://doi .org/10.1093/brain/90.1.131, PubMed: 6023071

Goodglass, H. (1993). Understanding aphasia. San Diego: Academic Press.

Green, D. M., & Swets, J. A. (1966). *Signal detection theory and psychophysics*. New York: Wiley.

Henschen, S. E. (1926). On the function of the right hemisphere of the brain in relation to the left in speech, music and calculation. *Brain*, 49, 110–123. https://doi.org/10 .1093/brain/49.1.110

Hertz-Pannier, L., Chiron, C., Jambaqué, I., Renaux-Kieffer, V., Van de Moortele, P.-F., Delalande, O., et al. (2002). Late plasticity for language in a child's non-dominant hemisphere: A pre- and post-surgery fMRI study. *Brain*, *125*, 361–372. https://doi.org/10.1093/brain/awf020, PubMed: 11844736

Hickok, G. (2009). The functional neuroanatomy of language. *Physics of Life Reviews*, 6, 121–143. https://doi.org/10.1016/j .plrev.2009.06.001, PubMed: 20161054

Hickok, G. (2012). Computational neuroanatomy of speech production. *Nature Reviews Neuroscience*, 13, 135–145. https://doi.org/10.1038/nrn3158. PubMed: 22218206

Hickok, G. (2014). *The myth of mirror neurons: The real neuroscience of communication and cognition*. New York: W. W. Norton & Company.

Hickok, G., Buchsbaum, B., Humphries, C., & Muftuler, T. (2003). Auditory–motor interaction revealed by fMRI: Speech, music, and working memory in area Spt. *Journal of Cognitive Neuroscience*, *15*, 673–682. https://doi.org/10.1162/089892903322307393, PubMed: 12965041

Hickok, G., Okada, K., Barr, W., Pa, J., Rogalsky, C., Donnelly, K., et al. (2008). Bilateral capacity for speech sound processing in auditory comprehension: Evidence from Wada procedures. *Brain and Language*, 107, 179–184. https://doi .org/10.1016/j.bandl.2008.09.006, PubMed: 18976806

Hickok, G., Okada, K., & Serences, J. T. (2009). Area Spt in the human planum temporale supports sensory-motor integration for speech processing. *Journal of Neurophysiology*, *101*, 2725–2732. https://doi.org/10.1152/jn .91099.2008, PubMed: 19225172

Hickok, G., & Poeppel, D. (2000). Towards a functional neuroanatomy of speech perception. *Trends in Cognitive Sciences*, 4, 131–138. https://doi.org/10.1016/S1364-6613(00) 01463-7, PubMed: 10740277

Hickok, G., & Poeppel, D. (2004). Dorsal and ventral streams: A framework for understanding aspects of the functional anatomy of language. *Cognition*, *92*, 67–99. https://doi.org/10 .1016/j.cognition.2003.10.011, PubMed: 15037127

Hickok, G., & Poeppel, D. (2007). The cortical organization of speech processing. *Nature Reviews Neuroscience*, 8, 393–402. https://doi.org/10.1038/nrn2113, PubMed: 17431404

Holland, R., & Lambon Ralph, M. A. (2010). The anterior temporal lobe semantic hub is a part of the language neural network: Selective disruption of irregular past tense verbs by rTMS. Cerebral Cortex, 20, 2771–2775. https://doi.org/10 .1093/cercor/bhq020, PubMed: 20223808

Indefrey, P. (2011). The spatial and temporal signatures of word production components: A critical update. *Frontiers in Psychology*, 2, 255. https://doi.org/10.3389/fpsyg.2011.00255, PubMed: 22016740

Indefrey, P., & Levelt, W. J. (2004). The spatial and temporal signatures of word production components. *Cognition*, 92, 101–144. https://doi.org/10.1016/j.cognition.2002.06.001, PubMed: 15037128

Jancke, L., Wustenberg, T., Scheich, H., & Heinze, H. J. (2002). Phonetic perception and the temporal cortex. *Neuroimage*, *15*, 733–746. https://doi.org/10.1006/nimg.2001.1027, PubMed: 11906217

Kennedy-Higgins, D., Devlin, J. T., Nuttall, H. E., & Adank, P. (2020). The causal role of left and right superior temporal gyri in speech perception in noise: A transcranial magnetic stimulation study. *Journal of Cognitive Neuroscience*, *32*, 1092–1103. https://doi.org/10.1162/jocn_a_01521, PubMed: 31933438

Knecht, S., Dräger, B., Deppe, M., Bobe, L., Lohmann, H., Flöel, A., et al. (2000). Handedness and hemispheric language dominance in healthy humans. *Brain*, *123*, 2512–2518. https://doi.org/10.1093/brain/123.12.2512, PubMed: 11099452

Kuhl, P. K., & Miller, J. D. (1975). Speech perception by the chinchilla: Voiced–voiceless distinction in alveolar plosive consonants. *Science*, *190*, 69–72. https://doi.org/10.1126 /science.1166301, PubMed: 1166301

Kuperman, V., Stadthagen-Gonzalez, H., & Brysbaert, M. (2012). Age-of-acquisition ratings for 30,000 English words. *Behavior Research Methods*, 44, 978–990. https://doi.org/10.3758 /s13428-012-0210-4, PubMed: 22581493

Lau, E. F., Phillips, C., & Poeppel, D. (2008). A cortical network for semantics: (De)constructing the N400. *Nature Reviews Neuroscience*, 9, 920–933. https://doi.org/10.1038/nrn2532, PubMed: 19020511

Lee, Y. S., Turkeltaub, P., Granger, R., & Raizada, R. D. (2012). Categorical speech processing in Broca's area: An fMRI study using multivariate pattern-based analysis. *Journal of Neuroscience*, *32*, 3942–3948. https://doi.org/10.1523 /JNEUROSCI.3814-11.2012, PubMed: 22423114

Leonard, M. K., Baud, M. O., Sjerps, M. J., & Chang, E. F. (2016). Perceptual restoration of masked speech in human cortex. *Nature Communications*, 7, 13619. https://doi.org/10.1038 /ncomms13619, PubMed: 27996973

Liberman, A. M. (1957). Some results of research on speech perception. *Journal of the Acoustical Society of America*, *29*, 117–123. https://doi.org/10.1121/1.1908635

Liberman, A. M., Cooper, F. S., Shankweiler, D. P., & Studdert-Kennedy, M. (1967). Perception of the speech code. *Psychological Review*, 74, 431–461. https://doi.org/10.1037 /h0020279, PubMed: 4170865

Liebenthal, E., Binder, J. R., Spitzer, S. M., Possing, E. T., & Medler, D. A. (2005). Neural substrates of phonemic perception. *Cerebral Cortex*, 15, 1621–1631. https://doi.org /10.1093/cercor/bhi040, PubMed: 15703256

McGettigan, C., Evans, S., Rosen, S., Agnew, Z. K., Shah, P., & Scott, S. K. (2012). An application of univariate and multivariate approaches in fMRI to quantifying the hemispheric lateralization of acoustic and linguistic processes. *Journal of Cognitive Neuroscience*, 24, 636–652. https://doi.org/10.1162/jocn_a_00161, PubMed: 22066589

McGettigan, C., Warren, J. E., Eisner, F., Marshall, C. R., Shanmugalingam, P., & Scott, S. K. (2011). Neural correlates of sublexical processing in phonological working memory. *Journal of Cognitive Neuroscience*, 23, 961–977. https://doi .org/10.1162/jocn.2010.21491, PubMed: 20350182 Meister, I. G., Wilson, S. M., Deblieck, C., Wu, A. D., & Iacoboni, M. (2007). The essential role of premotor cortex in speech perception. *Current Biology*, *17*, 1692–1696. https://doi.org /10.1016/j.cub.2007.08.064, PubMed: 17900904

Miceli, G., Gainotti, G., Caltagirone, C., & Masullo, C. (1980). Some aspects of phonological impairment in aphasia. *Brain* and Language, 11, 159–169. https://doi.org/10.1016/0093 -934X(80)90117-0, PubMed: 7427715

Moineau, S., Dronkers, N. F., & Bates, E. (2005). Exploring the processing continuum of single-word comprehension in aphasia. *Journal of Speech, Language, and Hearing Research, 48*, 884–896. https://doi.org/10.1044/1092-4388 (2005/061), PubMed: 16378480

Mori, S., Oishi, K., Jiang, H., Jiang, L., Li, X., Akhter, K., et al. (2008). Stereotaxic white matter atlas based on diffusion tensor imaging in an ICBM template. *Neuroimage*, 40, 570–582. https://doi.org/10.1016/j.neuroimage.2007.12.035, PubMed: 18255316

Morillon, B., Lehongre, K., Frackowiak, R. S. J., Ducorps, A., Kleinschmidt, A., Poeppel, D., et al. (2010). Neurophysiological origin of human brain asymmetry for speech and language. *Proceedings of the National Academy* of Sciences, U.S.A., 107, 18688–18693. https://doi.org/10.1073 /pnas.1007189107, PubMed: 20956297

Moses, D. A., Mesgarani, N., Leonard, M. K., & Chang, E. F. (2016). Neural speech recognition: Continuous phoneme decoding using spatiotemporal representations of human cortical activity. *Journal of Neural Engineering*, *13*, 056004. https://doi.org/10.1088/1741-2560/13/5/056004, PubMed: 27484713

Mottonen, R., & Watkins, K. E. (2009). Motor representations of articulators contribute to categorical perception of speech sounds. *Journal of Neuroscience*, 29, 9819–9825. https://doi.org/10.1523/JNEUROSCI.6018-08.2009, PubMed: 19657034

Myers, E. B. (2007). Dissociable effects of phonetic competition and category typicality in a phonetic categorization task: An fMRI investigation. *Neuropsychologia*, *45*, 1463–1473. https:// doi.org/10.1016/j.neuropsychologia.2006.11.005, PubMed: 17178420

Nachev, P., Coulthard, E., Jäger, H. R., Kennard, C., & Husain, M. (2008). Enantiomorphic normalization of focally lesioned brains. *Neuroimage*, *39*, 1215–1226. https://doi.org/10.1016/j .neuroimage.2007.10.002, PubMed: 18023365

Narain, C., Scott, S. K., Wise, R. J. S., Rosen, S., Leff, A., Iversen, S. D., et al. (2003). Defining a left-lateralized response specific to intelligible speech using fMRI. *Cerebral Cortex*, *13*, 1362–1368. https://doi.org/10.1093/cercor/bhg083, PubMed: 14615301

Nichols, T., Brett, M., Andersson, J., Wager, T., & Poline, J.-B. (2005). Valid conjunction inference with the minimum statistic. *Neuroimage*, *25*, 653–660. https://doi.org/10.1016/j .neuroimage.2004.12.005, PubMed: 15808966

Okada, K., & Hickok, G. (2006). Identification of lexical-phonological networks in the superior temporal sulcus using fMRI. *NeuroReport*, *17*, 1293–1296. https://doi.org/10 .1097/01.wnr.0000233091.82536.b2, PubMed: 16951572

Okada, K., Rong, F., Venezia, J., Matchin, W., Hsieh, I. H., Saberi, K., et al. (2010). Hierarchical organization of human auditory cortex: Evidence from acoustic invariance in the response to intelligible speech. *Cerebral Cortex*, 20, 2486–2495. https://doi.org/10.1093/cercor/bhp318, PubMed: 20100898

Overath, T., McDermott, J. H., Zarate, J. M., & Poeppel, D. (2015). The cortical analysis of speech-specific temporal structure revealed by responses to sound quilts. *Nature Neuroscience*, 18, 903–911. https://doi.org/10.1038/nn.4021, PubMed: 25984889 Pillay, S. B., Binder, J. R., Humphries, C., Gross, W. L., & Book, D. S. (2017). Lesion localization of speech comprehension deficits in chronic aphasia. *Neurology*, 88, 970–975. https:// doi.org/10.1212/WNL.00000000003683, PubMed: 28179469

Poeppel, D. (2001). Pure word deafness and the bilateral processing of the speech code. *Cognitive Science*, *25*, 679–693. https://doi.org/10.1207/s15516709cog2505_3

Poeppel, D. (2003). The analysis of speech in different temporal integration windows: Cerebral lateralization as "asymmetric sampling in time.". *Speech Communication*, 41, 245–255. https://doi.org/10.1016/S0167-6393(02)00107-3

Pulvermüller, F., Hauk, O., Nikulin, V. V., & Ilmoniemi, R. J. (2005). Functional links between motor and language systems. *European Journal of Neuroscience*, *21*, 793–797. https://doi.org/10.1111/j.1460-9568.2005.03900.x, PubMed: 15733097

Pulvermüller, F., Huss, M., Kherif, F., Moscoso del Prado Martin, F., Hauk, O., & Shtyrov, Y. (2006). Motor cortex maps articulatory features of speech sounds. *Proceedings of the National Academy of Sciences*, U.S.A., 103, 7865–7870. https://doi.org/10.1073/pnas.0509989103, PubMed: 16682637

Rauschecker, J. P., & Scott, S. K. (2009). Maps and streams in the auditory cortex: Nonhuman primates illuminate human speech processing. *Nature Neuroscience*, *12*, 718–724. https://doi.org/10.1038/nn.2331, PubMed: 19471271

Rizzolatti, G., & Arbib, M. A. (1998). Language within our grasp. *Trends in Neurosciences*, 21, 188–194. https://doi.org/10 .1016/S0166-2236(98)01260-0

Rogalsky, C., LaCroix, A. N., Chen, K. H., Anderson, S. W., Damasio, H., Love, T., et al. (2018). The neurobiology of agrammatic sentence comprehension: A lesion study. *Journal of Cognitive Neuroscience*, 30, 234–255. https://doi .org/10.1162/jocn a 01200, PubMed: 29064339

Rogalsky, C., Love, T., Driscoll, D., Anderson, S. W., & Hickok, G. (2011). Are mirror neurons the basis of speech perception? Evidence from five cases with damage to the purported human mirror system. *Neurocase*, *17*, 178–187. https://doi.org/10.1080/13554794.2010.509318, PubMed: 21207313

Rogalsky, C., Pitz, E., Hillis, A. E., & Hickok, G. (2008). Auditory word comprehension impairment in acute stroke: Relative contribution of phonemic versus semantic factors. *Brain and Language*, *107*, 167–169. https://doi.org/10.1016/j.bandl.2008 .08.003, PubMed: 18823655

Rogalsky, C., Rong, F., Saberi, K., & Hickok, G. (2011). Functional anatomy of language and music perception: Temporal and structural factors investigated using functional magnetic resonance imaging. *Journal of Neuroscience*, *31*, 3843–3852. https://doi.org/10.1523/JNEUROSCI.4515-10 .2011, PubMed: 21389239

Rorden, C., Fridriksson, J., & Karnath, H.-O. (2009). An evaluation of traditional and novel tools for lesion behavior mapping. *Neuroimage*, 44, 1355–1362. https://doi.org/10 .1016/j.neuroimage.2008.09.031, PubMed: 18950719

Schomers, M. R., Garagnani, M., & Pulvermüller, F. (2017). Neurocomputational consequences of evolutionary connectivity changes in perisylvian language cortex. *Journal* of Neuroscience, 37, 3045–3055. https://doi.org/10.1523 /JNEUROSCI.2693-16.2017, PubMed: 28193685

Schomers, M. R., Kirilina, E., Weigand, A., Bajbouj, M., & Pulvermüller, F. (2015). Causal influence of articulatory motor cortex on comprehending single spoken words: TMS evidence. *Cerebral Cortex*, 25, 3894–3902. https://doi.org/10 .1093/cercor/bhu274, PubMed: 25452575

Scott, S. K., Blank, C. C., Rosen, S., & Wise, R. J. S. (2000). Identification of a pathway for intelligible speech in the left temporal lobe. *Brain*, *123*, 2400–2406. https://doi.org/10 .1093/brain/123.12.2400, PubMed: 11099443 Skipper, J. I., Devlin, J. T., & Lametti, D. R. (2017). The hearing ear is always found close to the speaking tongue: Review of the role of the motor system in speech perception. *Brain* and Language, 164, 77–105. https://doi.org/10.1016/j.bandl .2016.10.004, PubMed: 27821280

Springer, J. A., Binder, J. R., Hammeke, T. A., Swanson, S. J., Frost, J. A., Bellgowan, P. S., et al. (1999). Language dominance in neurologically normal and epilepsy subjects: A functional MRI study. *Brain*, *122*, 2033–2046. https://doi.org /10.1093/brain/122.11.2033, PubMed: 10545389

Stokes, R. C., Venezia, J. H., & Hickok, G. (2019). The motor system's [modest] contribution to speech perception. *Psychonomic Bulletin & Review*, 26, 1354–1366. https://doi .org/10.3758/s13423-019-01580-2, PubMed: 30945170

Swanson, S. J., Sabsevitz, D. S., Hammeke, T. A., & Binder, J. R. (2007). Functional magnetic resonance imaging of language in epilepsy. *Neuropsychology Review*, 17, 491–504. https:// doi.org/10.1007/s11065-007-9050-x, PubMed: 18058239

Swanson, S. J., Binder, J. R., Possing, E. T., Hammeke, T. A., Sabsevitz, D. S., Spanaki, M., et al. (2002). fMRI language laterality during a semantic decision task: Age of onset and side of seizure focus effects. Paper presented at the Annual Meeting of the International Neuropsychological Society. Toronto, Canada.

Swets, J. A. (1964). *Signal detection and recognition by human observers*. Los Altos: Peninsula Publishing.

Utman, J. A., Blumstein, S. E., & Sullivan, K. (2001). Mapping from sound to meaning: Reduced lexical activation in Broca's aphasics. *Brain and Language*, 79, 444–472. https://doi.org /10.1006/brln.2001.2500, PubMed: 11781053

Vaden, K. I., Halpin, H. R., & Hickok, G. S. (2009). Irvine phonotactic online dictionary, Version 1.4. [Data file]. Available from https://www.iphod.com.

Venezia, J. H., Saberi, K., Chubb, C., & Hickok, G. (2012). Response bias modulates the speech motor system during syllable discrimination. *Frontiers in Psychology*, *3*, 157. https://doi.org/10.3389/fpsyg.2012.00157, PubMed: 22723787

Walker, G. M., Rollo, P. S., Tandon, N., & Hickok, G. (2021). Effect of bilateral opercular syndrome on speech perception. *Neurobiology of Language*, *2*, 335–353. https://doi.org/10 .1162/nol_a_00037

Watkins, K. E., & Paus, T. (2004). Modulation of motor excitability during speech perception: The role of Broca's area. *Journal of Cognitive Neuroscience*, 16, 978–987. https://doi.org/10.1162/0898929041502616, PubMed: 15298785

Watkins, K. E., Strafella, A. P., & Paus, T. (2003). Seeing and hearing speech excites the motor system involved in speech production. *Neuropsychologia*, 41, 989–994. https://doi.org/10.1016/S0028-3932(02)00316-0, PubMed: 12667534

Wernicke, C. (1874). In G. H. Eggert (Ed.), *Wernicke's works* on aphasia: A sourcebook and review (pp. 91–145). The Hague: Mouton.

Wilson, S. M. (2009). Speech perception when the motor system is compromised. *Trends in Cognitive Sciences*, 13, 329–330. https://doi.org/10.1016/j.tics.2009.06.001, PubMed: 19646917

Wilson, S. M., Saygin, A. P., Sereno, M. I., & Iacoboni, M. (2004). Listening to speech activates motor areas involved in speech production. *Nature Neuroscience*, 7, 701–702. https://doi.org /10.1038/nn1263, PubMed: 15184903

Winkler, A. M., Ridgway, G. R., Webster, M. A., Smith, S. M., & Nichols, T. E. (2014). Permutation inference for the general linear model. *Neuroimage*, 92, 381–397. https://doi.org/10.1016/j.neuroimage.2014.01.060, PubMed: 24530839

Zaidel, E. (1985). Language in the right hemisphere. In D. F. Benson & E. Zaidel (Eds.), *The dual brain: Hemispheric specialization in humans* (pp. 205–231). Guilford Press.

Zatorre, R. J., Belin, P., & Penhune, V. B. (2002). Structure and function of auditory cortex: Music and speech. *Trends in Cognitive Sciences*, 6, 37–46. https://doi.org/10.1016/S1364 -6613(00)01816-7, PubMed: 11849614

Zatorre, R. J., Evans, A. C., Meyer, E., & Gjedde, A. (1992). Lateralization of phonetic and pitch discrimination in speech processing. *Science*, 256, 846–849. https://doi.org/10.1126 /science.256.5058.846, PubMed: 1589767