

Negative BOLD Signals During Speech Comprehension.

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Abstract

Speech comprehension studies have generally focused on the isolation and function of regions with positive blood oxygen level-dependent (BOLD) signals with respect to a resting baseline. Although regions with negative BOLD signals in comparison to a resting baseline have been reported in language-related tasks, their relationship to regions of positive signals is not fully appreciated. Based on the emerging notion that the negative signals may represent an active function in language tasks, we test the hypothesis that negative BOLD signals during receptive language are more associated with comprehension than content-free versions of the same stimuli. Regions associated with comprehension of speech were isolated by comparing responses to passive listening to natural speech to two incomprehensible versions of the same speech: one that was digitally time-reversed and one that was muffled by removal of high frequencies. The signal polarity was determined by comparing the BOLD signal during each speech condition to the BOLD signal during a resting baseline. As expected, stimulation-induced positive signals relative to resting baseline were observed in the canonical language areas with varying signal amplitudes for each condition. Negative BOLD responses relative to resting baseline were observed primarily in fronto-parietal regions and were specific to the natural speech condition. However, BOLD signal remained indistinguishable from baseline for the unintelligible speech conditions. Variations in connectivity between brain regions with positive and negative signals were also specifically related to the comprehension of natural speech. These observations of anticorrelated signals related to speech comprehension are consistent with emerging models of cooperative roles represented by BOLD signals of opposite polarity.

Introduction

Language reception and production are the two primary components of human communication. Spoken language is fundamental to human social behavior and language learning starts early in life. Comprehension of speech involves many levels of input processing, from the simplest sound processing, to recognizing speech components, assigning meaning to words, and finally integrating the words into sentences and the sentences within a context of the narrative for the understanding of the speech. Integration of all these processes involves multiple associated brain regions. The neural correlates of these tasks have been extensively studied and canonical language areas are well known in the temporal and frontal cortices. Blood oxygenation-level-dependent (BOLD) signals acquired by functional MRI increase with respect to a resting baseline indicating stimulation-induced activation of these regions.

On the other hand, BOLD signals in areas associated with semantic processing have been widely observed as stimulation-induced deactivations with respect to a resting baseline during a wide range of non-semantic tasks. These regions, collectively designated as a default mode network (DMN), have been also reported in meta-analyses of spontaneous cognition (Biswal et al, 1995; Shulman et al, 1997; Gusnard et al, 2001; Mazoyer et al, 2001; Fox et al, 2007). However, semantic tasks have not typically been associated with deactivations within the DMN regions, and Binder (Binder et al, 1999) proposed that during conscious passive states like rest, subjects engaged in task-unrelated thoughts that were essentially semantic and therefore semantic tasks do not show deactivation of the DMN as other tasks do. In agreement with this idea, non-semantic tasks used in language studies (perceptual matching, non-word reading) generally produced deactivations that overlap with

the DMN regions (Mechelli et al, 2003; Rissman et al, 2003; Binder et al, 2005; Xiao et al, 2005). Stimulation-induced deactivations observed in the DMN regions would then represent a direct competition for attention and executive resources where regions that carry out effortful tasks (guided by exogenous signals) suppress regions that carry out task-unrelated thoughts (guided by endogenous signals).

These observations have shifted the focus of functional imaging studies initially from task-induced activations to task-induced deactivations (Buckner et al, 1996; McKiernan et al, 2003). When task-induced activations and deactivations are observed in comparison to a resting baseline, they reflect both positive and negative BOLD signals (above or below the signal of a resting baseline). This is distinguished from signals that are compared to another task and are positive or negative relative to activity in a related contrast. Further, multiple cognitive tasks have been shown to engage anti-correlated positive and negative signals (Fox et al, 2005; Buckner et al, 2008; Vincent et al, 2008; Anticevic et al, 2010; Anderson et al, 2011). Although the roles of brain regions with positive BOLD signals are well studied in receptive language processes (Zekveld et al, 2006; Ackermann et al, 2010; Price, 2012), the putative roles of brain regions with negative BOLD signals relative to a resting baseline and the interaction between these regions are not well appreciated.

So-called deactivations observed in many language studies are the result of comparisons between two conditions. The absence of a resting baseline precludes the distinction of deactivations with negative BOLD signal from those due to different task demands across conditions. For example, to compare semantic processing of content and function words, Diaz et al. used a working memory task in which non-words have to be remembered or retrieved, and interspersed with words that were task-irrelevant. This design,

where the baseline condition is a simple task rather than a resting baseline, is optimal to differentiate areas more responsive to function words or content words, but poor at determining which areas elicit a positive or negative BOLD response relative to a baseline. The observed deactivation of the dorsolateral prefrontal cortex for the word condition with respect to the non-word baseline cannot be assumed to correspond to a negative BOLD signal (Diaz et al, 2009).

Language studies where deactivations correspond to negative BOLD signals with respect to a resting baseline are few (Yoncheva et al, 2010) and usually focused on regions of the default mode network (DMN) (Schafer et al, 2009; Seghier et al, 2009; Snijders et al, 2009). The activation or deactivation of DMN nodes has been shown to vary with task: language tasks that engage episodic memory retrieval activate posterior nodes of the DMN, and deactivate the anterior node in middle prefrontal cortex. Furthermore, language tasks have been shown to recruit nodes in regions of the parietal lobule (Sestieri et al, 2011) or subregions of left angular gyrus (Seghier et al, 2010) that are anatomically separate from the DMN. The interrelation between activations and deactivations, in summary, has been shown to depend on the task and the specific demands, but remains poorly understood. In this study we investigate the relationship between stimulation-induced activations and deactivations during levels of speech comprehension. We specifically ask if the intelligibility of a stimulus induces variable signal deactivations. If so, would those deactivations be only located within the DMN system?

Methods

Subjects

Fifteen healthy right-handed subjects, 6 male, ages 30.5 ± 7.5 years/old were recruited for this study. Signed consent from the subjects was obtained prior to participation in the study in accordance with established institutional guidelines at Columbia University, Cornell University, and at Yale School of Medicine. Data from one subject was discarded due to uncorrectable movement artifacts.

Imaging

All scans were collected on a 1.5-tesla General Electric (Milwaukee, WI) magnetic resonance scanner with a standard head coil. We used foam to cushion the subjects' head and paper tape across the forehead to assist subjects to remain stable during the scanning sessions. Functional scans consisted of axial single shot EPI time series (128x128 mm matrix, 21 contiguous slices, field of view = 190 mm, $1.5 \times 1.5 \times 5$ mm voxels, TR = 3000 ms, TE = 43 ms, flip angle = 60°) obtained parallel to the AC/PC line. A total of 120 images were acquired in each run. Conventional high-resolution (T1-weighted) images were also acquired along sagittal planes and at the same axial plane locations as the T2*-weighted images.

Study Paradigm

The activation paradigm consisted of two separate runs with alternating blocks of stimulation and rest. The auditory stimuli were personal narratives that were pre-recorded by a friend or family member. Sentences that narrated an event of the subject's life were employed to enhance engagement during passive listening and simulate natural language. Three segments of the narratives, each lasting 18 seconds, were selected for each run. These

segments were modified to create three auditory conditions: i) natural speech condition: the segment was played as recorded, the content was comprehensible and sounded like natural speech; ii) reversed speech condition: the segment was digitally time-reversed so meaning and prosody were removed, and iii) muffled speech condition: frequency components higher than 400Hz and lower than 80Hz were removed by a band-pass filter in order to preserve fundamental frequency variability, lexical tone, prosody, and speech rate but rendered the content unintelligible. The stimuli were adjusted to equate for perceived loudness. The nine auditory segments were interleaved with resting baseline (periods of no stimulation), each lasting 18 seconds, the first block in a run being a rest, and the order of the conditions was counterbalanced across runs. Subjects wore headphones designed to reduce background scanner noise (Resonance Technology, Inc.). All subjects were instructed to keep their eyes closed and “listen to the narratives interspersed between periods of rest”. These instructions were consistent for all conditions to assure that, over all, rest periods were comparable across conditions. Before the acquisition of the data, 6-10 images were acquired during a typical speech segment to confirm that the subject was able to understand the language over the scanner noise. Subjects were instructed to listen to the stimuli in all cases and were informed that in some cases they might not understand the content. All subjects were queried following each run regarding their experience. In all cases subjects reported hearing all stimuli, understanding the natural speech, and not understanding the content of the reversed or the muffled speech.

Data Analysis

Preprocessing. The first five images from each run were removed and runs were concatenated for preprocessing and analysis. Initial analysis of image quality to discard image artifacts and excessive motion was performed with the CANlab software diagnostic tools (Cognitive and Affective Neuroscience Lab, University of Colorado at Boulder). Image processing and analysis was performed with Statistical Parametric Mapping (SPM)8 software (Wellcome Department of Imaging Neuroscience, University College, London, UK). Preprocessing steps included normalization to standard MNI space, motion correction using a sinc interpolation algorithm, and smoothing with a 6 mm full-width-half-maximum Gaussian kernel (FWHM). The BOLD response was modeled by boxcar functions for each condition (natural, reversed and muffled speech) that were convolved with the canonical hemodynamic response function (hrf). A 160 seconds high-pass filter was used to remove low frequency confounds and a low-pass filter shaped to match the canonical hemodynamic response function was used to remove unknown temporal autocorrelations. Motion parameters and the global value for each frame and detected spikes using the diagnostic tools were then used as nuisance regressors for subsequent analysis.

BOLD Activation Analysis. Statistical analysis of each subject's data was based on the General Linear Model (GLM) with three task regressors (natural, reversed, and muffled conditions), 6 motion parameters and a session regressor on each voxel in the entire brain volume. The single-subject contrast images were used for a second-level random effects group analysis. SPM[T] volumes were generated to investigate the effect of listening to all conditions (natural, reversed, and muffled speech) in comparison with the resting baseline as well as the contrast of natural speech greater than reversed and muffled speech to isolate regions associated with processing of intelligible speech. Individual voxel thresholds were

set at $p < 0.001$ ($t \geq 3.79$). To protect against false-positive results an effective $p \leq 0.02$ corrected threshold was satisfied by using only clusters of 40 or more voxels as determined by 10000 Monte Carlo simulations of whole brain fMRI data with the above parameters using AlphaSim in AFNI (v2009). Visualization and assignments of anatomic labels and MNI coordinates for active regions was performed using xjview toolbox, (<http://www.alivelearn.net/xjview>).

Block Signal Averaging. The average of the block signal were calculated for the clusters associated to speech comprehension (natural speech versus reversed and muffled speech). The time courses for all the voxels in each cluster were used. Each time course was divided into 36-seconds segments consisting of 18-second condition block and its following 18-second resting block. The resulting (speech + rest) blocks were grouped according to condition (natural, reversed and muffled speech). The average time course for all the 36-second blocks for each condition was calculated using matlab.

Psycho-physiological Interaction (PPI) Analysis. Functional connectivity was assessed using PPI analysis. This analysis identifies contributions of one brain region to another brain region that change with the experimental condition. Regions with responses during natural speech that were greater than muffled and reversed speech conditions for the group analysis were combined as a “comprehension seed” for the PPI (Uddin et al, 2009). Time-course series were extracted from the composite comprehension seed (network seed) for each participant and summarized as the first principal component across all voxels. The time series were adjusted with respect to the main effect of the stimulation to partial out confounds. The resulting residual signals were convolved with the psychological factor (intelligibility of the speech given by the comparison of natural vs. muffled and reversed

speech) to create the PPI regressors (Friston et al, 1997). A general linear model was applied with the PPI regressor, the network seed time-series, the three experimental task regressors and the 6 motion parameters as effects of no interest. SPM t-maps were thresholded at $P < 0.001$ ($t \geq 3 > 79$, uncorrected at the voxel level) with clusters size of 40 voxels.

Results

Positive and negative signals associated with all listening conditions

The conjunction of all speech conditions (natural, reversed and muffled) compared to resting baseline engaged regions with both positive BOLD signals (Figure 1 red clusters and Table 1.a) and with negative BOLD signals (Figure 1, blue clusters and Table 1.b) with respect to the resting baseline. The positive signals extended over bilateral temporal regions (middle, superior and transverse gyri), left frontal regions (inferior and precentral gyri), left hippocampal gyrus, and right putamen. As expected, the positive signals included traditional areas for auditory and language processing. The negative signals associated with all speech conditions constituted clusters over bilateral medial, middle and superior frontal, precuneus in the inferior parietal cortex, anterior, middle and posterior cingulate gyrus, middle occipital gyrus, left hippocampal gyrus, bilateral parahippocampal gyri and left thalamus. Most of these clusters have generally been associated with the DMN. As expected for a language task, the lateral temporal cortex, which is usually observed as a negative signal region included in the DMN, was observed here as a positive signal region. Notably, the posterior cingulate cortex engaged with the DMN in previous studies was not observed in this contrast.

===== Figure 1, Table 1=====

Positive and negative signals associated with speech comprehension

Speech comprehension, as observed during the natural speech compared to muffled and reversed speech conditions (meaningful speech with respect to incomprehensible speech), engaged activated regions (Figure 2, brain slices in the left, red clusters and Table 2.a) and deactivated regions (Figure 2 left panels, blue clusters and Table 2.b) for the comparison of speech conditions. Activated regions included bilateral temporal regions (middle temporal extending to temporo-parietal junction), frontal regions (left inferior and bilateral superior), bilateral parahippocampal gyrus, right hippocampal gyrus, caudate, and thalamus.

The time-course and magnitude of the block-averaged BOLD signals of bilateral temporal cortices and left inferior frontal regions are shown in the top row of Figure 2. These graphs confirm the positive BOLD signal (above the resting baseline) for the natural speech condition on the temporal and frontal cortices. The BOLD signal is also positive for the reversed and muffled speech conditions. The amplitude of the positive BOLD signal varies with condition and is higher for the natural speech than for the reversed and muffled speech, and amplitude for reversed speech exceeds that of muffled speech. In addition, the posterior cingulate gyrus and neighboring calcarine extending to precuneus and cuneus were also observed. The time course for this cluster (Figure 2, third row, right panel) shows that the BOLD signal is positive (above the resting baseline) for the natural speech condition (red), close to resting baseline for the reversed speech condition (blue) and negative (below the resting baseline) for the muffled condition (green) during the first six initial images corresponding to the task epoch (0-18 seconds).

===== Figure 2 =====

The deactivated regions (Figure 2, brain slices, left column, blue clusters and Table 2.b) associated with the natural speech in comparison to reversed and muffled speech included frontal regions (bilateral middle and left inferior gyri), parietal regions (bilateral superior, inferior, precuneus and postcentral gyri), left occipital regions (superior and cuneus) and right insula (see Table 2b for complete list with MNI coordinates). On the other hand, the frontal eye field areas usually observed in visual attention tasks were not observed here in the passive auditory task.

The time-course and magnitude of the BOLD signals of representative regions are shown in the second to bottom rows of Figure 2. Importantly, the graphs show the negative BOLD signals (below the resting baseline) in these regions for the natural speech condition, and illustrate that the amplitudes for the negative signals are smaller than those of the positive BOLD signals as expected (Kannurpatti et al, 2004). Therefore the results can be interpreted as true deactivations of these regions during the natural speech condition rather than a relative difference of natural speech condition in comparison to the reversed and muffled speech conditions. In addition, the BOLD signals during muffled and reversed conditions in these regions are close to baseline levels confirming that these regions are not engaged during the reversed or muffled conditions. Although the graphs shown in Figure 2 correspond to each condition and its following resting baseline for the given region of interest, each region was obtained by the SPM contrast and therefore the signal was assumed to reflect a significant change above the average of combined resting baselines.

Summarizing, the global deactivated fronto-parietal signals associated with intelligible speech were functionally anticorrelated with the traditional language system.

Functional connectivity of negative signals during language comprehension

To understand the dynamics of these negative signals during processing of intelligible speech we looked at the functional connectivity of all regions associated with language comprehension. We used the comprehensive set of fronto-parietal deactivated regions observed for the contrast of natural speech against reversed and muffled speech as a seed for PPI (clusters of negative BOLD signal seen in blue in Figure 2). These deactivated brain regions increased connectivity with supplementary motor area (SMA) (Table 3A.a and Figure 3.A, red clusters) and decreased connectivity with the canonical language regions in the middle and superior temporal gyri, and inferior frontal gyrus (Table 3A.b and Figure 3.A, blue clusters) during speech comprehension. These brain regions were observed in the contrast of all speech conditions against resting baseline (Figure 1). Thus, the global deactivated fronto-parietal signals associated with intelligible speech processing were functionally less connected with the traditional language system.

===== Figure 3, Table 2 =====

Functional connectivity of positive signals during language comprehension

Similarly, we used the comprehensive group of activated regions previously observed for the contrast of natural speech greater than reversed and muffled speech as a seed for PPI (clusters of positive BOLD signal seen in red in Figure 2A). and observed that the activated temporal and inferior frontal regions increased connectivity with the bilateral temporal cortices (middle, superior and transverse temporal cortex), and with left supramarginal gyrus, left hippocampal gyrus, and right postcentral (Table 3B.a and Figure

3.B, red clusters). On the other hand, these frontal and temporal activated regions decreased their connectivity to frontal gyrus (medial, middle and superior frontal gyrus), precuneus in the inferior parietal lobule, medial structures (anterior and posterior cingulate gyri), and thalamus (Table 3B.b and Figure 3.B, blue clusters). Most of these areas were observed as deactivations in Figure 1A (ventro-medial prefrontal cortex, posterior cingulate cortex (PCC) and posterior inferior parietal lobule) and are usually observed as part of the DMN. Thus, the activated regions associated with speech comprehension appear more connected to areas responsible for low-level auditory processing and less connected to the medial regions when the stimuli are meaningful.

Summarizing, we find that processing of natural, meaningful speech involves a constellation of brain regions with negative and positive BOLD signals relative to a resting baseline. While the amplitude of the positive BOLD signal was graded for the three conditions, the amplitude of the negative BOLD signal was not graded and only present during the intelligible speech condition. Regions of positive and negative signals were collectively distinguished by their functional connectivity. When the speech was meaningful the areas of positive BOLD signal decreased their connectivity to the DMN, and the areas of negative BOLD signal decreased their connectivity to the traditional language areas in frontal and temporal cortices.

Discussion

Our results indicate that engagement of particular activated and/or deactivated regions is related to the nature of the language processes, with language comprehension of natural intelligible speech engaging a large-scale frontal-parietal deactivated signal not

observed when the stimuli are unintelligible. The contrast of natural versus muffled and reversed speech demonstrated clusters of positive BOLD signal with respect to the resting baseline within the temporal cortex and frontal cortex that correspond well with current accounts of large-scale networks associated with auditory language comprehension of sentences and syntactic processing located in temporal and frontal areas (Fiebach et al, 2004; Hunter et al, 2006; Humphries et al, 2007; Saur et al, 2008; Saur et al, 2010; Seghier et al, 2010; Snijders et al, 2010; Papoutsi et al, 2011; Tyler et al, 2011). Current models associate language processing with larger networks with components in temporal, frontal gyrus, and subcortical regions (Hirsch et al, 2000; Catani et al, 2005; Saur et al, 2008; Friederici, 2009; Kelly et al, 2010; Xiang et al, 2010; Papoutsi et al, 2011; Barbas et al, 2013). Neuroimaging studies confirm that mapping phonological representations onto lexical conceptual representations involves the anterior superior temporal, middle temporal and inferior temporal sulcus that connect to premotor areas (BA 44 and 6) (Warren et al, 2009). In addition, semantic processing engages the superior temporal sulcus and the superior temporal gyrus as well as the middle temporal gyrus connecting to the ventrolateral prefrontal cortex (BA 45 and 47) (Vigneau et al, 2006; Saur et al, 2010; Weiller et al, 2011).

We also observed strong positive activation of hippocampal and parahippocampal gyrus during language comprehension that is paired with activation of the precuneus/posterior cingulate cortex suggesting the engagement of memory functions as well. The posterior inferior parietal lobule and the precuneus have been associated with episodic memory retrieval (Shannon et al, 2004; Huijbers et al, 2012). The positive time course for meaningful speech and negative signal for muffled speech in the posterior cingulate gyrus and precuneus is in agreement with the role of this region in semantic

processing observed by Seghier and Price (Seghier et al, 2012). A meta-analysis of 120 studies points to the role of precuneus/posterior cingulate cortex as an interface between the semantic network and the hippocampal memory system to encode meaningful events into episodic memory (Binder et al, 2011). On the other hand, the hippocampus and adjacent areas in the medial temporal lobe that are associated with episodic memory function are also part of the DMN (Greicius et al, 2004). In fact, hippocampal formation shows spontaneous correlations with many of the major regions of the DMN (Vincent et al, 2008). In our study, hippocampal gyrus (slice -20) is active in two contrasts: all speech conditions vs. resting baseline (Figure 1), and meaningful speech (forward speech vs. reversed and muffled speech, slice -20). These results indicate that the signal amplitude for forward speech is greater than for the other speech conditions, although all conditions have a positive signal in this region. Therefore, hippocampal activation may be related to recollection associated with intelligibility of speech that brings back memories, but it could also be related to recollection associated to other attributes of the stimuli (i.e., recognition of a familiar voice, emotional recollection associated to it). On the other hand, parahippocampal gyrus and a region of the hippocampus are also deactivated for all speech conditions vs. resting baseline (Figure 1, slices -8 and 0) suggesting that the signal is below the resting baseline for all the conditions to some degree. Therefore, these regions do not decrease activation with the intelligibility of the speech as the areas shown in Figure 2. The results show a heterogeneity of sub-regions of the hippocampal gyrus regarding activation or deactivation during language comprehension. Although our study was not designed to disentangle the precise roles of these structures for language comprehension, the different patterns of activation suggest a different contribution of these sub-regions of the hippocampal system to episodic memory

associated with language processing.

The contrast of natural versus muffled and reversed speech also demonstrated clusters of negative BOLD signal with respect to the resting baseline. Notably, negative BOLD responses with respect to the resting baseline have not been consistently reported in language studies possibly because this signal is smaller than the positive BOLD signal for a similar change of neural activity (Shmuel et al, 2006). The negative BOLD signals with respect to the resting baseline observed in our study encompass the intraparietal sulcus, parietal and the insular regions; areas that have been described as a dorsal attention network activated during externally oriented tasks (Corbetta, 1998; Fox et al, 2005; Gazzaley et al, 2007), and have also been associated with a control network that responds to cues that indicate task onset (Dosenbach et al, 2008). Dosenbach has proposed that this network encompasses regions that initiate attentional control triggered by the cue and regions that process trial-by-trial performance feedback to adjust control. Based on these studies we expected the frontal parietal network to be silent in the absence of an explicit task. Furthermore, we expected this network to be active during the presentation of the muffled and reversed speech due to the greater effort of processing these unusual stimuli. However the deactivation of these regions during passive listening to natural speech suggests a different engagement. This result could be interpreted as a suppression of the attention network but further investigation is necessary.

It has been widely shown that cerebral activity is coupled with the positive BOLD responses by localized changes in blood oxygenation that produce a mismatch of changes in cerebral blood flow (CBF), cerebral blood volume (CBV) and cerebral metabolic rate of oxygen consumption (CMRO₂) (Kim et al, 2012). Positive BOLD responses correlate

with increases in CBV and CBF. However, the neuronal origin of the negative BOLD signal appears to be less understood. The coupling with decreases of CBF and CMRO₂ is supported by many studies (Shmuel et al, 2002; Blumenfeld et al, 2004; Englot et al, 2009; Mullinger et al, 2014) suggesting that the intrinsic dynamics serve cognitive and perceptual processes (Shmuel et al, 2002; Shmuel et al, 2006). However, negative BOLD signals do not necessarily reflect decreased CBF associated with neural activity. The signal polarity can also be explained by a decrease of CBF due to a redistribution of CBF into neighboring regions that causes a local decrease of CBF (known as vascular-steal effect) (Harel et al, 2002; Kannurpatti et al, 2004). Other possible mechanisms include a large increase in CMRO₂ without an adequate CBF increase in areas with low vascular reactivity (Schridde et al, 2008), or an increase of dopamine release that causes an increased activity of subcortical structures and vasoconstriction of neighboring microvessels (Shih et al, 2009). Therefore, the hemodynamic mechanism may be different for positive and negative BOLD signals. The techniques employed in this study do not differentiate between these possible underlying mechanisms. Finally, negative BOLD signals have been observed in caudate and putamen despite their increased neural activity during spontaneous seizures (Mishra et al, 2011). Therefore, caution must be used in the interpretation of the negative BOLD signal for these structures and in cases of altered hemodynamics in brain pathologies.

The observed negative and positive signals observed during language comprehension might be interpreted as cooperative and separate processes. In this frame, the negative signals in the absence of goal-directed tasks might allow for a more efficient processing of the auditory information, and to a greater extent when the speech is meaningful. We observe that regions specific for processing natural intelligible language exert both an enhancement

of functional connectivity on traditional language regions and a weakening of functional connectivity to the deactivated areas during speech comprehension. This may serve to reduce interference with other processes.

Alternatively, the meaningful auditory stimuli, selectively engaging posterior tempo-parietal cortex and posterior cingulate gyrus, could amplify neural representations of task-relevant information while disengaging from representations of task-irrelevant information in fronto-parietal signals. Amplification of the brain response to task-relevant information has been previously shown in cognitive control paradigms (Egner et al, 2005; Chadick et al, 2011). These effects could be applied directly from the posterior temporal or the posterior cingulate cortex or mediated by subcortical areas such as thalamus or the insular cortex.

Binder suggested that if the observed deactivations relative to the resting baseline were due to relocation of resources, their amplitudes would be expected to vary with task difficulty (Binder et al, 1999). In this study we used natural speech and two unintelligible versions of the same auditory stimuli to isolate brain regions that process meaning, and observed a variation of the positive BOLD signal relative to the resting baseline for the three speech conditions but without a similar variation in the negative signal. However, a further variation in the level of speech comprehension may be necessary to assess the responsiveness of the BOLD signal and could be the target of future studies to investigate factors that affect brain regions with negative BOLD signals outside the DMN.

The observation of both positive and negative signals relative to the resting baseline only for natural speech suggests a possible interaction between positive and negative signals during high-level (meaningful) language processing. Functional connectivity using an inclusive seed composed of all the regions of positive BOLD signals, mainly composed by

canonical language areas in the superior/middle temporal and inferior frontal gyri, showed increased connectivity with auditory and language regions of the temporal cortex. These findings are in agreement with reported increased connectivity between the left posterior temporal cortex and left inferior frontal gyrus, bilateral middle temporal, left posterior inferior temporal gyrus and right occipital gyrus comparing ambiguous and unambiguous sentences (Snijders et al, 2010). Papoutsi and colleagues (Papoutsi et al, 2011) found only negative PPI effects for ambiguous compared to unambiguous sentences. Coupling was observed between left posterior middle temporal and bilateral ACC/GF_d, precuneus and left middle occipital gyrus. Similarly, we found strong decreases in connectivity from the canonical language areas to medial regions of the DMN. On the other hand, functional connectivity using an inclusive seed composed of all the regions of negative BOLD signal, mainly composed of fronto-parietal areas in the middle frontal gyrus and superior and inferior parietal areas, showed decreased connectivity to temporal and frontal speech processing regions and increased connectivity only to the SMA. The SMA was deactivated during the contrast of all conditions against baseline.

These global connectivity patterns suggest that regions with negative and positive BOLD signal relative to the resting baseline dynamically decrease their connectivity during speech comprehension, coupled with functional changes in connectivity to deactivated regions that are considered part of the DMN. Thus, the neural dynamics of language comprehension include both strengthening connectivity between positive BOLD regions and temporal regions and decreasing connectivity between positive BOLD regions and DMN regions and between negative fronto-parietal network and traditional language areas.

Conclusion

We test the hypothesis that large-scale networks of negative signals relative to a resting baseline are a fundamental component of the neural underpinnings of speech comprehension, and that the neural dynamics of language comprehension involve interactions between large-scale networks including both positive and negative anti-correlated signals. In our study, fMRI was employed to acquire neural responses during listening to meaningful natural speech in comparison to incomprehensible versions of the same speech. An extensive temporal-frontal language network of areas with positive signals relative to the resting baseline was observed as expected. Further, negative signals relative to the resting baseline were observed in fronto-parietal regions associated specifically with speech comprehension. These findings demonstrate that receptive-language induced negative BOLD signals are anticorrelated with the receptive-language induced positive BOLD signals during comprehension of spoken language. The anti-correlation between positive and negative BOLD signals relative to the resting baseline is observed during the intelligible natural speech and not during the unintelligible speech. Thus, both distinct positive and negative signal responses are observed in the contrast of speech against baseline and in the contrast of natural speech against muffled and reversed speech. These results suggest that the differences between processing meaningful intelligible speech and unintelligible speech are not only shown by patterns of positive signals, but that the patterns of negative signals reflect processes also contributing to the processing of language stimuli. While the language regions with positive BOLD signal relative to the resting baseline increased connectivity to adjacent temporal areas and decreased connectivity to the medial areas typically associated to the DMN, the fronto-parietal regions with negative BOLD

signal relative to the resting baseline decreased their connectivity to temporal and frontal language areas. These findings suggest that the neural dynamics of processing natural speech include specific strengthening and weakening of connectivity between networks of positive and negative BOLD signals, consistent with a model of large-scale anticorrelated processes in language comprehension.

Understanding the functional relationships between positive and negative BOLD signals relative to the resting baseline could further our understanding of language deficits of neurological origin such as autism and acquired language deficits such as disorders of consciousness (Rodriguez Moreno et al, 2010). The passive task employed in this study allows natural listening without imposing laboratory-constrained conditions. The personalized natural narrative segments used in our paradigm motivates future applications to clinical situations where the ability to respond is impaired. Subjects were asked to attend to the narratives with their eyes closed, and comprehension of the natural narratives was confirmed for all imaging runs and failure to comprehend was confirmed for the muffled and reversed conditions. Language conditions that impose minimal attention requirements such as the passive listening have been used for patients that have limited cognitive or motor abilities, and this utility provides an additional rationale for studying fundamental properties of language comprehension using a passive listening paradigm.

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Author Disclosure Statement

No competing financial interests exist.

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Figure Captions

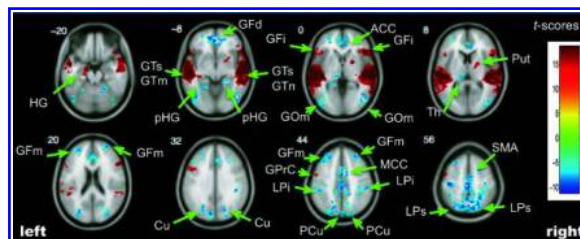


FIG. 1. Main effect of listening to all speech conditions (natural, reversed and muffled) in comparison to resting baseline (effective corrected threshold $p \leq 0.02$ for cluster size 40, color bar= t-scores). Red: positive signals, cyan: negative signals.

ACC: anterior cingulate cortex, Cu: cuneus, GF_i: inferior frontal gyrus, GF_d: medial frontal gyrus, GF_m: middle frontal gyrus, GH: hippocampal gyrus; GOM: middle occipital gyrus, GP_rC: precentral gyrus, GT_m: middle temporal gyrus, GT_s: superior temporal gyrus, LP_i: inferior parietal lobule, LP_s: superior parietal lobule, MCC: middle cingulate cortex, PCu: precuneus, pHG: parahippocampal gyrus, Put: putamen, SMA: supplementary motor area, Th: thalamus.

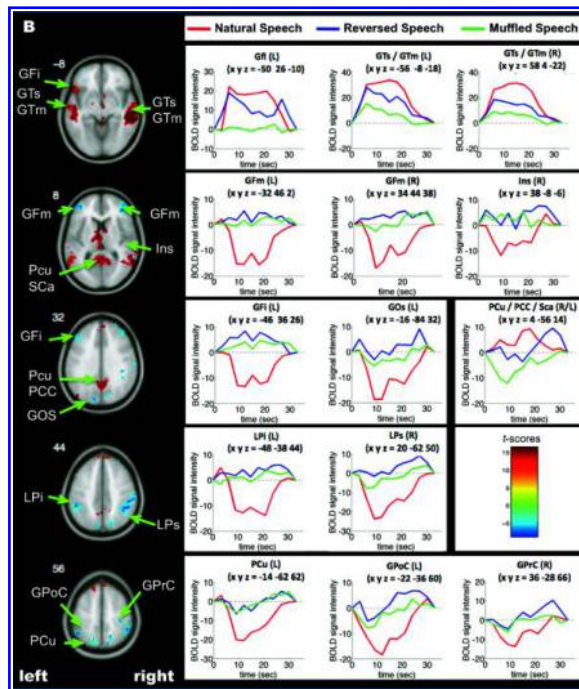


FIG. 2. Main effect of language comprehension (natural speech in comparison to muffled and reversed speech) (effective corrected threshold $p \leq 0.02$ for cluster size 40, color bar= t-scores). Representative brain slices are shown in the left panels. Red: positive signals, cyan: negative signals. The time course for the block-average BOLD signals are shown in the right panels. Signals above the zero baseline correspond to positive BOLD signal relative to a resting baseline. Signals below the zero baseline correspond to negative BOLD signals relative to a resting baseline. Red: natural speech, blue: reversed speech and green: muffled speech.

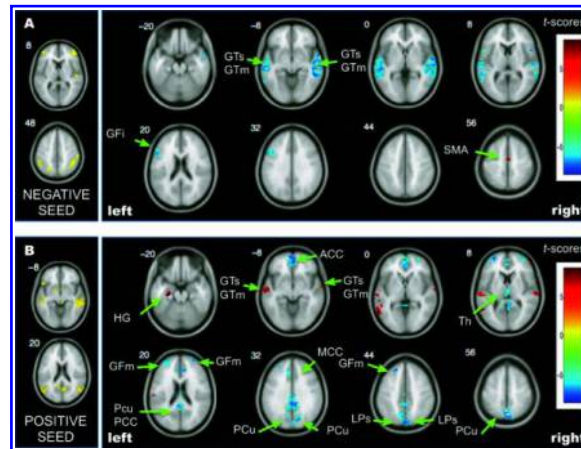


FIG. 3. PPI results of functional connectivity for the networks associated with language comprehension. Red represent increases in connectivity with the seed network and blue clusters represent decreases in connectivity (effective corrected threshold $p \leq 0.02$ for cluster size 40). A. Changes in connectivity with the negative fronto-parietal network seed (shown in yellow in the left panel) for the natural speech in comparison to the muffled and reversed speech. B. Changes in connectivity with the positive regions' network seed (shown in yellow in the left panel) for the natural speech compared to the muffled and reversed speech.

TABLE 1. MAIN EFFECT OF LISTENING: PASSIVE LISTENING TO ALL SPEECH CONDITIONS IN COMPARISON TO RESTING BASELINE (EFFECTIVE CORRECTED THRESHOLD $P \leq 0.02$ FOR CLUSTER SIZE 40).

<i>Brodmann's</i>				
<i>Region</i>	<i>Anatomical Area</i>	<i>Area</i>	<i>T max</i>	<i>x,y,z (mm)</i>
<i>a. Listening > Baseline (Positive BOLD network)</i>				
Temporal	Superior (L)	22	10.69	-42 -24 8

Temporal	Superior (L)	22	10.52	-48 -30 4
Temporal	Superior (L)	42	11.19	-62 -18 10
Temporal	Superior (R)	22	13.42	52 -14 -6
Temporal	Superior (R)	22	12.71	46 -24 0
Temporal	Superior (R)	22	12.3	62 -22 -2
Frontal	Inferior Operculum (L)	45	5.43	-52 18 12
Frontal	Inferior Orbitalis (L)	47	6.5	-52 20 -8
Frontal	Precentral (L)	6	5.75	-50 -4 54
Frontal	Precentral (L)	6	5.52	-54 0 48
Frontal	Precentral (L)	6	4.21	-46 -4 46
Frontal	Precentral (L)	9	5.61	-48 12 32
Limbic	Hippocampus (L)	34	6.58	-16 -6 -20
Limbic	Hippocampus (L)	35	4.49	-20 -20 -18
Limbic	Hippocampus (L)		5.27	-24 -12 -16
Sublobar	Putamen (R)		9.78	20 0 4
Sublobar	Putamen (R)		4.22	30 10 -6

b. Listening < Baseline (Negative BOLD network)

Temporal	Middle (R)	37	11.67	44 -66 0
Frontal	Medial Orbital (R)	10	9.53	2 52 -8
Frontal	Middle (L)	8	11.32	-28 22 54
Frontal	Middle (R)	10	5.84	28 58 22
Frontal	Middle (L)	10	6.95	-30 38 14
Frontal	Middle (L)	11	5.46	-26 46 2

Frontal	Middle (R)	10	4.68	26 30 38
Frontal	Middle (R)	8	6.29	26 36 50
Frontal	Middle (R)	9	6.09	34 34 38
Frontal	Superior (L)	6	5.62	-16 4 62
Frontal	Superior (L)	6	5.48	-20 0 70
Frontal	Superior (L)	6	3.94	-18 14 64
Frontal	Superior (R)	6	7.54	22 10 58
Frontal	Superior Orbital (L)	10	6.72	-26 54 -6
Limbic	Anterior Cingulate (R)	32	10.19	8 44 -10
Limbic	Hippocampus (L)		9.12	-22 -40 0
Limbic	Parahippocampal Gyrus (L)	19	9.98	-24 -50 -10
Limbic	Parahippocampal Gyrus (R)	30	8.87	26 -42 -10
Occipital	Fusiform (L)	19	9.01	38 -68 -12
Occipital	Fusiform (L)	19	6.65	-44 -68 -20
Parietal	Precuneus (L)	7	11.1	-4 -62 58
Parietal	Precuneus (R)	7	11.99	2 -18 74
Sublobar	Thalamus Pulvinar (L)		8.31	-12 -26 2
Sublobar	Thalamus Pulvinar (L)		4.15	-16 -32 6
Sublobar	Thalamus (L)		4.42	-8 -22 16
Cerebellum	(R)	4, 5	8.68	18 -52 -20

x, y, z = MNI coordinates for activation peak of significant clusters.

TABLE 2. MAIN EFFECT OF LISTENING TO INTELLIGIBLE SPEECH: PASSIVE LISTENING TO NATURAL SPEECH IN COMPARISON TO PASSIVE LISTENING TO MUFFLED AND REVERSED SPEECH (EFFECTIVE CORRECTED THRESHOLD $P \leq 0.02$ FOR CLUSTER SIZE 40).

<i>Brodmann's</i>				
<i>Region</i>	<i>Anatomical Area</i>	<i>Area</i>	<i>T max</i>	<i>x,y,z (mm)</i>
<i>a. Listening to Natural Speech > Reversed and Muffled Speech (Positive BOLD signal)</i>				
Temporal	Middle (L)	21	8.53	-56 -8 -18
Temporal	Middle (L)	21	7.64	-48 -14 -10
Temporal	Middle (L)	22	7.39	-54 -26 -2

Temporal	Middle (R)	21	11.97	58 4 -22
Temporal	Middle (R)	21	9.78	64 -34 -8
Temporal	Middle (R)	21	9.21	56 -36 -10
Frontal	Inferior Orbital (L)	47	5.32	-50 26 -10
Frontal	Inferior Orbital (L)	47	4.27	-40 22 -14
Frontal	Inferior Triangular (L)	47	5.18	-44 22 -2
Frontal	Superior (L)	8	5.47	-14 36 52
Frontal	Superior (L)	8	4.8	-12 30 58
Frontal	Superior Medial (L)	9	6.13	-4 56 38
Frontal	Superior Medial (L)	9	4.7	-2 46 38
Frontal	Superior Medial (R)	8	5.73	6 36 60
Frontal	Superior Medial (R)	9	4.89	4 52 40
Limbic	Parahippocampal (L)	28	6.12	-20 -20 -20
Limbic	Parahippocampal (R)	25	5.97	24 -24 -22
Limbic	Hippocampal (R)	35	5.82	32 -18 -26
Limbic	Hippocampal (R)	35	5.66	30 -16 -16
Occipital	Calcarine/Precuneus (R)	23	8.56	4 -56 14
Sublobar	Amygdala (L)		7.14	-16 -2 -14
Sublobar	Caudate Body (R)		8.44	8 8 10
Sublobar	Thalamus Ventral Ant. (L)		8.05	-6 -6 8
Cerebellum	Declive (R)		4.64	6 -82 -26
Cerebellum	Declive of Vermis (L)		5.06	-2 -84 -28

b. Listening to Natural Speech < Reversed and Muffled Speech (negative BOLD signal)

Frontal	Inferior Triangular (L)	46	6.14	-46 36 26
Frontal	Middle (L)	10	7.09	-32 46 2
Frontal	Middle (L)	46	7.08	-42 48 10
Frontal	Middle (R)	9	7.72	34 44 38
Frontal	Middle (R)	10	5.98	38 42 8
Frontal	Middle (R)	10	5.89	48 46 22
Frontal	Paracentral (L)	5	5.8	-12 -36 62
Frontal	Precentral (R)	4	5.51	36 -28 66
Occipital	Cuneus (L)	19	5.84	-10 -88 26
Occipital	Superior (L)	19	7	-16 -84 32
Parietal	Inferior (L)	40	6.37	-48 -38 44
Parietal	Inferior (L)	40	6.16	-44 -52 54
Parietal	Inferior (L)	40	5.65	-52 -52 50
Parietal	Postcentral (L)	3	6.37	-22 -36 60
Parietal	Postcentral (R)	4	6.58	36 -30 56
Parietal	Postcentral (R)	40	7.51	60 -22 46
Parietal	Precuneus (L)	7	5.68	-12 -48 60
Parietal	Precuneus (L)	7	5.2	-14 -62 62
Parietal	Precuneus (L)	31	6.25	-24 -76 16
Parietal	Superior (L)	7	4.86	-14 -68 54
Parietal	Superior (L)	7	4.23	-22 -70 58
Parietal	Superior (R)	7	6.27	20 -62 50
Parietal	Superior (R)	7	5.85	14 -74 52

Parietal	Superior (R)	7	5.34	16 -58 56
Parietal	Superior (R)	40	8.37	52 -42 60
Sublobar	Insula (R)	13	6.05	38 -8 -6
Sublobar	Insula (R)	13	5.33	42 -2 4
Sublobar	Insula (R)	13	4.1	38 -10 6

TABLE 3B. CHANGES IN FUNCTIONAL CONNECTIVITY FOR THE LANGUAGE COMPREHENSION REGIONS WITH POSITIVE BOLD SIGNAL ASSOCIATED WITH SPEECH DURING THE NATURAL SPEECH IN COMPARISON TO THE REVERSED AND MUFFLED SPEECH (EFFECTIVE CORRECTED THRESHOLD $P \leq 0.02$ FOR CLUSTER SIZE 40)

<i>Brodmann's</i>				
<i>Region</i>	<i>Anatomical Area</i>	<i>Area</i>	<i>T max</i>	<i>x,y,z (mm)</i>
<i>a. Increased Connectivity during Listening to Natural Speech</i>				
Temporal	Middle (L)	21	7.51	-60 -58 0
Temporal	Middle (L)	21	6.77	-56 -46 4
Temporal	Middle (L)	21	6.29	-58 -10 -8
Temporal	Middle (L)	21	5.29	-66 -34 -6

Temporal	Middle (L)	21	5.12	-58 -32 -4
Temporal	Middle (L)	39	3.93	-52 -56 10
Temporal	Middle (R)	21	5.35	64 -4 -10
Temporal	Superior (R)	22	4.94	66 -10 0
Temporal	Transverse (L)	41	5.8	-44 -24 12
Parietal	Postcentral (R)	40	5.83	64 -20 14
Parietal	Supramarginal (L)	40	6.72	-54 -24 14
Limbic	Hippocampal (L)	35	8.47	-28 -20 -20

b. Decreased Connectivity during Listening to Natural Speech

Frontal	Medial Orbital (R)	10	-7.13	8 62 -6
Frontal	Medial Orbital (R)	32	-8.33	4 42 -4
Frontal	Middle (L)	10	-7.33	-40 50 8
Frontal	Middle (L)	10	-5.19	-32 46 16
Frontal	Middle (L)	10	-7.2	-24 34 46
Frontal	Middle (L)	10	-5.82	-32 56 4
Frontal	Middle (R)	10	-5.89	30 58 20
Frontal	Middle Orbital (R)	10	-6.36	34 60 -8
Frontal	Superior (L)	8	-4.08	-12 36 38
Frontal	Superior (L)	10	-7.17	-26 46 28
Frontal	Superior (L)	10	-6.4	-24 56 14
Frontal	Superior (R)	10	-6.03	32 62 12
Frontal	Superior medial (L)	9	-7.76	-4 44 24
Limbic	Posterior Cingulate (R)	30	-8.01	2 -52 12

Limbic	Posterior Cingulate (R)	31	-7.9	0 -38 30
Limbic	Posterior Cingulate (R)	31	-7.88	4 -44 10
Occipital	Cuneus (R)	7	-9.89	20 -70 28
Parietal	Precuneus (L)	7	-7.69	-2 -62 56
Parietal	Precunues (L)	7	-5.25	-12 -64 30
Parietal	Precunues (R)	7	-11.37	4 -60 62
Sublobar	Thalamus Medial Dorsal (R)		-4.71	2 -12 6
Sublobar	Thalamus (R)		-8.54	6 -16 16

TABLE 3A. CHANGES IN CONNECTIVITY FOR THE FRONTO-PARIETAL REGIONS WITH NEGATIVE BOLD SIGNAL ASSOCIATED WITH SPEECH DURING THE NATURAL SPEECH IN COMPARISON TO THE REVERSED AND MUFFLED SPEECH (EFFECTIVE CORRECTED THRESHOLD $P \leq 0.02$ FOR CLUSTER SIZE 40)

<i>Brodmann's</i>				
<i>Region</i>	<i>Anatomical Area</i>	<i>Area</i>	<i>T max</i>	<i>x,y,z (mm)</i>
<i>a. Increased Connectivity during Listening to Natural Speech</i>				
Frontal	Supplementary Motor Area (R)	6	6.35	0 -12 52
<i>b. Decreased Connectivity during Listening to Natural Speech</i>				
Temporal	Superior (L)	21	-7.15	-56 -26 -8
Temporal	Superior (L)	21	-6.56	-50 -40 2
Temporal	Superior (L)	22	-6.09	-58 -30 0

Temporal	Superior (R)	21	-8.31	64 -16 -6
Temporal	Superior (R)	21	-8.21	58 -6 -4
Temporal	Superior (R)	22	-7.71	64 -2 -8
Frontal	Inferior Operculum (R)	44	-5.75	52 10 8
Frontal	Inferior Triangularis (L)	9	-6.68	-46 16 28
Frontal	Inferior Triangularis (L)	45	-6.49	-54 16 24