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### A possible functional localiser for identifying brain regions sensitive to sentence-level prosody

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Investigations of how we produce and perceive prosodic patterns are not only interesting in their own right but can inform fundamental questions in language research. We here argue that functional magnetic resonance imaging (fMRI) in general – and the functional localisation approach in particular – has the potential to help address open research questions in prosody research and at the intersection of prosody and other domains. Critically, this approach can go beyond questions like 'where in the brain does mental process *x* produce activation' and towards questions that probe the nature of the representations and computations that subserve different mental abilities. We describe one way to functionally define regions sensitive to sentence-level prosody in individual subjects. This or similar 'localiser' contrasts can be used in future studies to test the hypotheses about the precise contributions of prosody-sensitive brain regions to prosodic processing and cognition more broadly

### Keywords: fMRI; prosody; intonation

### Introduction

Patterns of pitch and loudness in speech, as well as the ways in which words are grouped temporally, provide an important source of information in both language acquisition (e.g., Gleitman & Wanner, 1982; Jusczyk, 1997; Jusczyk, Cutler, & Redanz, 1993; Mattys, Jusczyk, Luce, & Morgan, 1999) and language processing (e.g., Bader, 1998; Marslen-Wilson, Tyler, Warren, Grenier, & Lee, 1992). Investigating how people produce and/or perceive certain aspects of prosody is not only interesting in its own right, but can also inform broader issues in the architecture of human language. For example, investigations of prosody have been used to ask about the basic meaning units of language (e.g., Selkirk, 1984; cf. Watson & Gibson, 2004), or about whether we produce language with a comprehender in mind (e.g., Albritton, McKoon, & Ratcliff, 1996; Breen, Fedorenko, Wagner, & Gibson, 2010; Snedeker & Trueswell, 2003; cf. Kraljic & Brennan, 2005; Schafer, Speer, Warren, & White, 2000). Although we have learned a tremendous amount over the last several decades, some key questions about the mechanisms that support prosodic processing remain unanswered (or at least are still actively debated). For example, are the same or distinct mechanisms used for processing prominence patterns vs. temporal grouping information in the speech signal? Does extracting prosodic information from speech rely on specialised cognitive and neural machinery, or is it instead supported by some of the mechanisms that are engaged by other mental processes, like musical processing or social cognition? The answers to these

questions would importantly constrain the possibilities for the kinds of representations that mediate prosodic processing, which might in turn allow us to tackle even more challenging questions. For example, how do prosodic representations interact with syntactic/semantic representations in both constructing utterances in the course of production and extracting meaning from the linguistic signal in the course of comprehension? In the current paper, we will (1) argue that functional magnetic resonance imaging (fMRI) is a powerful – and currently underused in the domain of prosody – tool that can help address some of these open questions, and (2) describe an fMRI approach that we think is promising for doing so.

The rest of this paper is organised as follows: We begin with a brief summary of previous investigations of the brain basis of prosodic processing. We then outline an fMRI approach, which has been successful in other domains but has not yet been applied in the domain of prosody. In particular, we argue for the importance of defining regions of interest (ROI) functionally in individual subjects (e.g., Fedorenko, Hsieh, Nieto-Castanon, Whitfield-Gabrieli, & Kanwisher, 2010; Saxe, Brett, & Kanwisher, 2006). We then motivate and present the current study, which provides one possible way to identify brain regions sensitive to sentence-level prosody. We conclude with a discussion of how this (or similar) 'functional localisers' can be used in future work to tackle theoretically important questions in the domain of prosody and beyond.

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## Previous investigations of the brain basis of prosodic processing

As in other cognitive domains, two primary sources of evidence have informed our understanding of how prosodic processing is instantiated in the brain: (1) studies of patients with brain damage, and (2) neuroimaging (PET, fMRI) investigations. Both kinds of studies have implicated a large number of cortical regions in the temporal, frontal and parietal lobes, as well as some subcortical structures (for reviews see e.g., Baum & Pell, 1999; Friederici & Alter, 2004; Kotz, Meyer, & Paulmann, 2006; Van Lancker & Breitenstein, 2000; Van Lancker Sidtis, Pachana, Cummings, & Sidtis, 2006; Wildgruber, Ackermann, Kreifelts, & Ethofer, 2006; Wong, 2002).

A question that has perhaps received the most attention in the literature is that of lateralisation of prosodic processing. Some early patient findings have suggested that deficits in prosody perception and production – especially affective prosody - typically arise after damage to the right hemisphere (e.g., Bowers, Coslett, Bauer, Speedie, & Heilman, 1987; Bradvik et al., 1991; Bryan, 1989; Darby, 1993; Dykstra, Gandour, & Stark, 1995; Heilman, Scholes, & Watson, 1975; Ross, 1981; Ross & Mesulam, 1979; Ross, Thompson, & Yenkosky, 1997; Schmitt, Hartje, & Willmes, 1997; Starkstein, Federoff, Price, Leiguarda, & Robinson, 1994; Tucker, Watson, & Heilman, 1977; Weintraub, Mesulam, & Kramer, 1981; see also Blumstein & Cooper, 1974; Herrero & Hillix, 1990; Ley & Bryden, 1982, for behavioural evidence of right hemisphere superiority for prosodic processing from healthy individuals). The right hemisphere superiority for processing affective prosody fits nicely with work on emotional processing in other domains (e.g., Sackeim, Gur, & Saucy, 1978; Strauss & Moscovitch, 1981; cf. Caltagirone et al., 1989; Kowner, 1995). However, other studies have shown that the left hemisphere also plays an important role, especially for linguistic prosody, leading to the argument that both hemispheres may be important (e. g., Baum, Daniloff, Daniloff, Lewis, 1982; Behrens, 1988; Blumstein & Goodglass, 1972; Emmorey, 1987; Goodglass & Calderon, 1977; Heilman, Bowers, Speedie, & Coslett, 1984; Pell & Baum, 1997; Schlanger, Schlanger, & Gerstman, 1976; Seron, Van der Kaa, Vanderlinden, Remits, & Feyereisen, 1982; Shapiro & Danly, 1985; Speedie, Coslett, & Heilman, 1984; Van Lancker, 1980; Van Lancker & Sidtis, 1992; Wertz, Henschel, Auther, Ashford, & Kirshner, 1998; Zurif & Mendelsohn, 1972). In a recent review and meta-analysis of neuropsychological investigations, Witteman, van IJzendoorn, van de Velde, van Heuven, and Schiller (2011; see also Kotz et al., 2006) conclude that both hemispheres are necessary for both emotional and linguistic prosodic perceptions, but the right hemisphere plays a relatively greater role in processing emotional prosody. In particular, the right hemisphere damage leads to (1) greater deficits in emotional compared to linguistic prosody, and (2) greater deficits in emotional prosody compared to left hemisphere damage. Consistent with this review, neuroimaging studies often find bilateral cortical activations for prosodic (including affective prosodic) manipulations, as well as some additional subcortical structures (e.g., George et al., 1996; Grandjean et al., 2005; Imaizumi et al., 1997; Kotz et al., 2003; Phillips et al., 1998).

In summary, although many cortical and subcortical brain regions in both hemispheres have been implicated in prosodic processing and a number of proposals have been advanced for the functions of these regions (e.g., Ethofer, Anders, Erb, Herbert et al., 2006; Friederici & Alter, 2004; Kotz & Schwartze, 2010; Van Lancker Sidtis et al., 2006; Wildgruber, Ethofer, Kreifelts, & Grandjean, 2009), most researchers agree that more work is needed in order to understand the precise contributions of these different regions to perceiving and producing prosody.

# Functional localisation as an alternative to the traditional fMRI approach

Many previous neuroimaging studies have focused on the question of which hemisphere or which particular brain region is engaged by some aspect(s) of prosodic processing. This kind of a question is a necessary starting point, but the ultimate goal of cognitive science and cognitive neuroscience is to understand the function(s) of each relevant component of the mind/brain. In particular, for any given brain region, we would like to know what kinds of knowledge representations it stores and works with, and/or what computations it performs on particular stimuli. To be able to answer – or at least begin to answer – these questions, multiple hypotheses need to be evaluated about each key brain region. As a result, no single study will be sufficient. In order to accumulate knowledge across studies and labs, it is important to be able to refer to the 'same' region from one brain to the next. We have recently been arguing that the traditional fMRI approach is not well suited for comparing results across studies as needed for accumulating knowledge (e.g., Fedorenko et al., 2010; Fedorenko & Kanwisher, 2009; Fedorenko, Nieto-Castañón, & Kanwisher, 2012). In particular, in the traditional group-based approach, brains are aligned in the common stereotaxic space and activation overlap is examined across individual brains. However, because of anatomical variability (e.g., Amunts et al., 1999; Brodmann, 1909; Geschwind & Levitsky, 1968; Juch, Zimine, Seghier, Lazeyras, & Fasel, 2005; Miller et al., 2002; Ono, Kubik, & Abernathey, 1990; Tomaiuolo et al., 1999; Wohlschläger et al., 2005; Zilles et al., 1997), individual activations do not line up well across brains, especially in the frontal and temporal lobes (e.g., Frost & Goebel, 2012; Tahmasebi et al., 2012). Consequently, locations (e.g., sets of  $\{x,y,z\}$  coordinates) in the stereotaxic space are not optimally suited for comparing results across individuals and studies.

For example, imagine a scenario where one study reports activation in or around some anatomical location (e.g., superior temporal gyrus, STG) for a manipulation of affective prosody, and another study reports a nearby location (also within the STG) for a manipulation of linguistic prosody. Based on this pattern, one could arrive at two opposite conclusions about the relationship between affective and linguistic prosody. On the one hand, it could be argued that the two locations are close enough to each other (falling within the same broad anatomical region) to count them as the 'same' region, which would imply that affective and linguistic prosody rely on the same mechanisms. On the other hand, it could be argued that because the activations do not fall in exactly the same coordinates in the stereotaxic space, they are two nearby but distinct regions, which would imply that affective and linguistic prosody are supported by different mechanisms. We know that in many parts of the brain small but functionally distinct regions lie side by side (e.g., the fusiform face area (FFA) and the fusiform body area - Schwarzlose, Baker, & Kanwisher, 2005; or different regions within the left inferior frontal gyrus -Fedorenko, Duncan, & Kanwisher, 2012). Consequently, without comparing the two manipulations to each other in the same individual, it is impossible to determine which interpretation is correct.

An approach that has been proposed as an alternative to the traditional fMRI approach involves (1) identifying ROIs functionally in each individual brain (i.e., regions that exhibit a particular functional signature), and then (2) probing the functional profiles of those regions in additional studies in an effort to narrow down the range of possible hypotheses about their function(s). For example, using a contrast between faces and objects, Kanwisher, McDermott, and Chun (1997) identified a region in the fusiform gyrus that responds more strongly during the processing of faces than during the processing of objects. This region can be robustly found in any individual's brain in just a few minutes of scanning. Then, across many subsequent studies, the responses of this region were examined to many new stimuli and tasks to try to understand what drives the stronger response to faces (e. g., Downing, Chan, Peelen, Dodds, & Kanwisher, 2006; Kanwisher, Tong, & Nakayama, 1998). Because the same 'localiser' task (the faces > objects contrast in this example) is used across studies, the results can be straightforwardly compared. This 'functional localisation' approach is the standard approach in the field of vision research, and it has recently been successfully extended to other domains (e.g., social cognition - Saxe & Kanwisher, 2003; speech perception - Belin et al., 2000; Hickok, Okada, & Serences, 2009; language -Fedorenko et al.,

2010; January et al., 2009; Pinel et al., 2007). In addition to facilitating knowledge accumulation, the functional localisation approach yields higher sensitivity and functional resolution, i.e., it is more likely to detect an effect when it is present, and it is better at distinguishing between nearby functionally different regions, which is especially important for addressing questions of functional specificity (e.g., Nieto-Castañon & Fedorenko, 2012).

In summary, the functional localisation approach is more conducive to asking deep questions about the nature of the representations and computations underlying a particular mental process, compared to the traditional fMRI approach. We therefore advocate the adoption of this approach for investigating prosodic processing. A prerequisite for this approach is a 'localiser' task that can identify domain- or process-relevant regions at the level of individual subjects. We here propose one possible 'localiser' for brain regions sensitive to sentence-level prosody.

### **Experiment**

It is not obvious what experimental contrast(s) are best suited for discovering brain regions sensitive to prosodic processing. Previous studies have used several approaches: (1) stimulus manipulations where different kinds of prosodic contours are compared to one another (e.g., Doherty, West, Dilley, Shattuck-Hufnagel, & Caplan, 2004; Grandjean et al., 2005; Kotz et al., 2003); (2) stimulus manipulations where a prosodic contour is compared to some control condition(s) where some aspects of prosody are degraded (e.g., Humphries, Love, Swinney, & Hickok, 2005; Newman, Supalla, Hauser, Newport, & Bayelier, 2010; Wiethoff et al., 2008; Zhao et al., 2008); and (3) task manipulations where participants perform either a task that draws attention to prosody (e.g., classifying the emotion that the intonation is conveying) vs. some control task on the same stimuli (e.g., Buchanan et al., 2000; Ethofer, Anders, Erb, Droll et al., 2006; Ethofer, Anders, Erb, Herbert et al., 2006; Gandour et al., 2003; George et al., 1996; Mitchell, Elliott, Barry, Cruttenden, & Woodruff, 2003). The first approach only makes sense if we assume that the neural representations/ processes of different prosodic contours (either different affective contours, like happy vs. sad prosody, or different linguistic prosodic contours, like statements vs. questions) are spatially distinct. It is not clear that such an assumption is warranted. And if the same patch of cortex processes different kinds of contours, then any contrast of this sort would not reveal those regions. For example, the FFA (Kanwisher et al., 1997) responds to a wide range of face stimuli, and contrasting, for example, sad vs. neutral faces or male vs. female faces would miss this region.

The second and third approaches seem more promising. Any brain region engaged in prosodic processing should respond more when the signal contains a prosodic contour, compared to when some features of this contour are present to a lesser extent or absent. Similarly, brain regions engaged in prosodic processing may be expected to respond more when the task requires paying attention to the prosodic features of the stimulus compared to some other features, although this approach may not work well if perceiving prosody is highly automatic. In that case, prosody-sensitive regions would be engaged to a similar extent regardless of the specific task and may thus be subtracted out in a task-based contrast. Consistent with this notion, some studies that have used task manipulations report activations in what appear to be the highly domain-general regions of the fronto-parietal network (e. g., Buchanan et al., 2000), which respond across a wide range of cognitive demands and which are generally sensitive to salient stimuli (e.g., Duncan & Owen, 2000; Fedorenko, Duncan, & Kanwisher, 2013; see Duncan, 2010, for a recent review of this brain system). As a result of these potential concerns with the stimulus manipulations that compare different prosodic contours, and task manipulations, in the current study we chose to use a stimulus manipulation that compares stimuli that have sentence prosody to those that do not.

The design used here has not been previously used to the best of our knowledge. We used 'structured' linguistic stimuli (sentences and Jabberwocky sentences, which obey the rules of English syntax but use pseudowords instead of the content words) and 'unstructured' linguistic stimuli (lists of words and lists of pseudowords; see (2) below for sample stimuli).2 These linguistic materials were presented visually and auditorily. The contrast between structured and unstructured linguistic stimuli has some features that are the same regardless of the modality of presentation. For example, structured stimuli contain syntactic information and compositional semantic information, and that holds for both visually and auditorily presented materials. Importantly, however, auditorily presented structured stimuli - read naturally (cf. Humphries et al., 2005) – involve sentence-level prosodic contours, which are not present, or present to a lesser degree, in the unstructured stimuli (see Methods), as shown schematically in (1). Thus, subtracting the unstructured conditions from the structured conditions in the auditory materials should activate brain regions sensitive to sentence-level prosodic contours. But this subtraction also includes whatever makes the structured materials structured (i.e., syntax and compositional semantics). In order to isolate the prosody-relevant component of auditory structured stimuli, we contrasted the 'structured' > unstructured' comparison for the auditorily presented stimuli with the same comparison for the visually presented stimuli.

(1) A schematic illustration of the logic of the experimental design (structured stimuli: sentences, Jabberwocky sentences; Unstructured stimuli: word lists, pseudoword lists):

|                         | Structured > | Structured > |
|-------------------------|--------------|--------------|
|                         | Unstructured | Unstructured |
|                         | VISUAL       | AUDITORY     |
|                         | presentation | presentation |
| Syntax                  | +            | +            |
| Compositional semantics | +            | +            |
| Sentence-level prosody  | _            | +            |

Although silent reading of sentences has been argued to activate prosodic representations (Fodor, 1998; see e.g., Bader, 1998; Hirose, 2003; Swets, Desmet, Hambrick, & Ferreira, 2007, for experimental evidence), previous work on visual imagery has established that to the extent that mental simulations of a particular sensory experience activate the corresponding sensory cortices, these activations are not nearly as robust as those elicited by actual sensory stimulation (e.g., O'Craven & Kanwisher, 2000). With respect to our design, this finding suggests that the sentence-level prosodic response to visually presented sentences and Jabberwocky sentences will be much weaker than to the same materials presented auditorily. Consequently, we reasoned that we can target brain regions that are sensitive to sentence-level prosodic contours with a conjunction of two contrasts: (1) a greater response to structured than unstructured auditory stimuli, and (2) no difference between structured and unstructured visual stimuli.<sup>3</sup> In other words, this conjunction of contrasts is aimed at brain regions that selectively respond to the presence of structure in the auditory stimuli.

To discover these prosody-sensitive<sup>4</sup> regions, we use a method that was recently developed for investigating high-level language regions (Fedorenko et al., 2010) and subsequently validated on the well-known ventral visual stream regions (Julian, Fedorenko, Webster, & Kanwisher, 2012; see also Fedorenko, McDermott, Norman-Haignere, & Kanwisher, 2012, for the application of this method to musical processing). This method - the group-constrained subject-specific (GSS) method – is an alternative to the traditional random-effects analysis, where individual activation maps are aligned in the common space and a t-test is performed in every voxel. The GSS analysis discovers patterns that are spatially systematic across subjects without requiring voxel-level overlap, thus accommodating inter-subject variability in the anatomical location of functional regions. This method thus improves sensitivity and functional resolution in cases where the effects are anatomically variable (Nieto-Castañon & Fedorenko, 2012).

### Methods

### **Participants**

Twelve participants (seven females) between the ages of 18 and 30 – students at MIT and members of the surrounding community – were paid for their participation.<sup>5</sup> Participants were right-handed native speakers of English. All participants had normal or corrected-to-normal vision and were naïve to the purposes of the study. All participants gave informed consent in accordance with the requirements of the Internal Review Board at MIT.

### Design, materials and procedure

Each participant performed several runs of the visual version of the experiment and several runs of the auditory version of the experiment. The entire scanning session lasted between 1.5 and 2 hours.

#### Materials

There were four types of stimuli: sentences, word lists, Jabberwocky sentences and pseudoword lists. One hundred and sixty items were constructed for each condition, and each item was eight words/pseudowords long. Sample items for each type of stimulus are shown in (2) below. For details of how the materials were created, see Fedorenko et al. (2010; Experiments 2-3). All the materials are available from: http://web.mit.edu/evelina9/ www/funcloc.html. After each stimulus, a word (for Sentences and Word-lists conditions) or a pseudoword (for Jabberwocky and Pseudoword-lists conditions) appeared and participants were asked to decide whether this word/pseudoword appeared in the immediately preceding stimulus. Participants were instructed to press one of two buttons to respond. [In previous work, we have established that activations in the language-sensitive brain regions are similar regardless of whether this memory probe task is included or whether participants are simply reading the materials with no task (Fedorenko, submitted; Fedorenko et al., 2010)].

(2) Sample items:

Sentences: THE DOG CHASED THE CAT ALL DAY LONG

A RUSTY LOCK WAS FOUND IN THE DRAWER Word lists: BECKY STOP HE THE LEAVES BED LIVE MAXIME'S

FOR THE JUICE UP AROUND GARDEN LILY TRIES Jabberwocky: THE GOU TWUPED THE VAG ALL LUS RALL

A CLISY NYTH WAS SMASP IN THE VIGREE Pseudoword lists: BOKER DESH HE THE DRILES LER CICE FRISTY'S

FOR THE GRART UP AROUND MEENEL LALY SMEBS

One hundred and twenty-eight of the 160 items were used for the auditory versions of the materials (fewer materials

were needed because four, not five, items constituted a block of the same length as the visual presentation, because visual presentation is typically faster than the average speaking rate). The materials were recorded by a native speaker using the Audacity software, freely available at http://audacity.sourceforge.net/. The speaker was instructed to produce the sentences and the Jabberwocky sentences with a somewhat exaggerated prosody. These two conditions were recorded in parallel in order to make the prosodic contours across each pair of a regular sentence and a Jabberwocky sentence as similar as possible. The speaker was further instructed to produce the Word-lists and Pseudoword-lists conditions not with a list intonation, where each word/pseudoword is a separate intonational phrase (e.g., Schubiger, 1958; Couper-Kuhlen, 1986), but in a way that would make them sound more like a continuous stream of speech. This was done to make the low-level acoustic properties (e.g., frequency of boundaries in a stimulus) more similar between the structured and the unstructured stimuli (see Discussion and Appendix C for the results of the acoustic analyses of the materials, and for sample pitch tracks). We reasoned that with this recording strategy the auditory structured stimuli would only differ from the auditory unstructured stimuli in the presence of intonation patterns typical of English sentences (present in the structured stimuli vs. absent - or present to a lesser extent - in the unstructured stimuli).

### Visual presentation

Words/pseudowords were presented in the centre of the screen one at a time in all capital letters. No punctuation was included in the Sentences and Jabberwocky conditions, in order to minimise differences between the Sentences and Jabberwocky conditions on the one hand, and the Word-lists and Pseudoword-lists conditions on the other. Each trial lasted 4800 ms, which included (1) a string of eight words/pseudowords each presented for 350 ms, (2) a 300 ms fixation, (3) a memory probe appearing on the screen for 350 ms, (4) a period of 1000 ms during which participants were instructed to press one of two buttons, and (5) a 350 ms fixation. Participants could respond any time after the memory probe appeared on the screen. There were five trials in each block.

### Auditory presentation

Stimuli were presented over scanner-safe earphones. Each trial lasted 6000 ms, which included (1) the stimulus (whose total duration varied between 3300 ms and 4300 ms), (2) a 100 ms beep tone indicating the end of the sentence, (3) a memory probe presented auditorily (maximum duration 1000 ms), (4) a period (lasting until the end of the trial) during which participants were instructed to press one of two buttons. Participants could respond

any time after the onset of the memory probe. There were four trials in each block.

Each run lasted for 464 sec (7 min 44 sec) and consisted of 16 24-sec blocks, grouped into four sets of four blocks with 16-sec fixation periods at the beginning of the run and after each set of blocks. Condition order was counterbalanced across runs and participants, and auditory and visual runs were alternated. Each item was presented in either visual or auditory modality, but not both; which items were presented in which modality varied across participants. Each participant, except for one, completed four visual and four auditory runs. The remaining participant completed two visual and seven auditory runs.

### fMRI data acquisition

Structural and functional data were collected on the whole-body 3 Tesla Siemens Trio scanner with a 12channel head coil at the Athinoula A. Martinos Imaging Center at the McGovern Institute for Brain Research at MIT. T1-weighted structural images were collected in 128 axial slices with 1.33 mm isotropic voxels (TR = 2000 ms, TE = 3.39 ms). Functional, blood oxygenation leveldependent (BOLD) data were acquired using an EPI sequence (with a 90° flip angle), with the following acquisition parameters: 32 4-mm thick near-axial slices acquired in the interleaved order (with 10% distance factor), 3.1 × 3.1 mm in-plane resolution, FoV in the phase encoding (A >> P) direction 200 mm and matrix size  $96 \times 96$  mm, TR = 2000 ms and TE = 30 ms. The first 8 sec of each run were excluded to allow for steady state magnetisation.

### fMRI data analyses

MRI data were analysed using SPM5 (http://www.fil.ion.ucl.ac.uk/spm) and custom matlab scripts (available from http://web.mit.edu/evelina9/www/funcloc.html). Each subject's data were motion corrected and then normalised into a common brain space (the Montreal Neurological Institute, MNI template) and resampled into 2-mm isotropic voxels. Data were smoothed using a 4-mm Gaussian filter, and high-pass filtered (at 200 sec).

To identify brain regions sensitive to sentence-level prosodic contours, we performed a GSS analysis (Fedorenko et al., 2010; Julian et al., 2012). To do so, we first created – for each individual subject – a map containing voxels that satisfy the following two criteria: (1) a significant effect of structure, i.e., Sentences+Jabberwocky > Word-lists+Pseudoword-lists (at p < .01 uncorrected level) for the auditory materials, and (2) no significant effect of structure (at p < .1) for the visual materials. We then overlaid these maps to create a probabilistic overlap map, and then divided this map into

'parcels' using the watershed image parcellation algorithm. This algorithm discovers key topographical features (i.e., the main 'hills') of the activation landscape (see Fedorenko et al., 2010, for details). We then identified parcels which – when intersected with the individual maps - contained responses in at least 9/12 individual subjects (i.e.,  $\sim$  75%). Twelve parcels satisfied this criterion. Finally, because there is currently less agreement about how to interpret deactivations in fMRI, we selected a subset of these 12 parcels that responded to each of the (1) auditory Sentences, and (2) auditory Jabberwocky sentences conditions reliably above the fixation baseline. Four parcels remained. [See Appendix A for figures showing all 12 parcels and their responses to the eight experimental conditions. Note that some of these parcels – not discussed below - look spatially similar to regions implicated in some previous studies of prosodic processing (e.g., medial frontal regions; e.g., Alba-Ferrara, Hausmann, Mitchell, & Weis, 2011; Heilman, Leon, & Rosenbek, 2004) and may thus require further investigation.]

For each of the four key parcels, we estimated the response magnitude to each condition in individual subjects using an *n*-fold cross-validation procedure, so that the data used to define the functional ROI (fROIs) and to estimate the responses were independent (e.g., Kriegeskorte, Simmons, Bellgowan, & Baker, 2009; Vul & Kanwisher, 2010). Each parcel from the GSS analysis (Figure 1) was intersected with each subject's activation map containing voxels that satisfy the criteria described above (i.e., an effect of structure for auditory materials, and a lack of such an effect for visual materials) for all but one run of the data. All the voxels that fell within the boundaries of the parcel were taken as that subject's fROI (see Appendix B for figures showing sample individual fROIs). This procedure was iterated across all possible partitions of the data, and the responses were then averaged across the left-out runs to derive a single response magnitude per subject per region per condition (see Appendix A for responses to individual conditions). Statistical tests were performed on these values. Two contrasts were examined to test for sensitivity to structure in the visual and auditory conditions, respectively: (1) Sentences + Jabberwocky > Word-lists + Pseudoword-lists for visual materials; and (2) Sentences+Jabberwocky > Word-lists + Pseudoword-lists for auditory materials.

### Results

We discovered four regions in which the majority of subjects showed a selective response to structure in the auditory materials (i.e., a greater response to sentences and Jabberwocky sentences than to word and pseudoword lists for auditory, but not for visual, materials). These regions (parcels) are shown in Figure 1 (see Table 1 for a

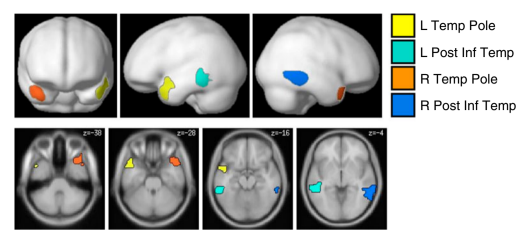


Figure 1. Top: Prosody-sensitive parcels projected onto the cortical surface. The parcels show the locations where most subjects showed activation for the relevant contrasts (i.e., an effect of structure for the auditory, but not for the visual, conditions; see Methods for details). Bottom: Parcels projected onto axial slices (colour assignments are the same in both views). [These parcels are available from the first author upon request and will soon be made available at: http://web.mit.edu/evelina9/www/funcloc.html].

summary of key properties) and include bilateral regions in the superior temporal poles, and bilateral regions in the posterior inferior temporal lobes. (Note that the activations of any individual subject within a parcel typically constitute only a small fraction of the parcel; see Table 1 for average sizes of the individual fROIs; see also Appendix B.) Each of the four regions was present in at least 9/12 subjects; the left temporal pole region was present in 10/12 subjects.<sup>6</sup>

In Figure 2, we present the responses of these prosody-sensitive regions to the presence of structure in the visual and auditory conditions (estimated using cross-validation, as described in Methods). The sensitivity to structure in the auditory conditions is highly robust in each of these regions (p's < .001). However, none of the regions show an effect of structure in the visual conditions (t's < 1.24; see Table 2 for the details of the statistics).

### Discussion

Four brain regions in the temporal cortices were found to be sensitive to sentence-level prosodic contours, as evidenced by a stronger response to structured (sentences, Jabberwocky sentences) compared to unstructured (word lists, pseudoword lists) conditions for the auditory, but not visual, presentation. These regions include bilateral superior temporal pole regions, and bilateral regions in the posterior inferior temporal lobes. We now discuss a few theoretical and methodological points that the current study raises.

## The prosody-sensitive regions discovered in the current study

What can we currently say about the four regions discovered in the current experiment? We begin by ruling out a few possibilities. *First*, these regions are not likely engaged in low-level auditory processing given that (1) both structured and unstructured auditory stimuli require basic auditory analysis and (2) they are not located in or around primary auditory cortex (e.g., Morosan et al., 2001).

Second, by design, we know that these four regions are not part of the 'language network' (e.g., Binder et al., 1997; Fedorenko et al., 2010), whose regions respond to linguistic stimuli similarly across modalities (see also Braze et al., 2011). For example, in Figure 3 below we show sensitivity to structure in the visual vs. auditory conditions in the language regions (defined by a greater response to sentences than to lists of pseudowords, as

Table 1. Basic information on the prosody-sensitive brain regions. The units for the parcel sizes and the average individual fROI sizes are 2 mm isotropic voxels.

|                                 | Left hemisphere |               | Right hemisphere |               |
|---------------------------------|-----------------|---------------|------------------|---------------|
|                                 | Temp pole       | Post Inf Temp | Temp pole        | Post Inf Temp |
| Present in n/12 subjs           | 10              | 9             | 9                | 9             |
| Parcel size                     | 723             | 889           | 765              | 885           |
| Average size of individual fROI | 41              | 46            | 36               | 47            |

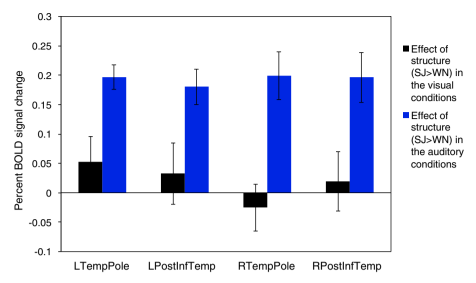


Figure 2. Sensitivity to structure in the visual vs. auditory conditions in individually defined prosody-sensitive regions. (The responses are estimated using n-fold cross-validation, as discussed in Methods, so that data to define the fROIs and estimate the responses are independent).

described in Fedorenko et al., 2010). As can be clearly seen, all of the regions of the 'language network' show highly robust sensitivity to structure in both modalities (all p's < .001). Note that this is in spite of the fact that language activations encompass extended portions of the temporal cortices, including anterior temporal regions, especially in the left hemisphere. It appears then that language and prosody-sensitive regions are located near each other but are nevertheless functionally distinct. This result is consistent with the findings from the neuropsychological literature that at least some aphasic individuals have intact prosodic abilities (e.g., Hughlings-Jackson, 1931; Schmitt et al., 1997).

It is worth noting, however, that one possibility that needs to be tested in future work is that the regions identified in the current study are, in fact, part of the language network, but are less sensitive to structure (i.e., syntax and combinatorial semantics) than the 'core' language regions (e.g., Figure 3 below). Sentence-level prosodic contours (present in the auditory materials) may

make the structural information more salient thus recruiting these possibly less sensitive-to-structure brain regions. To evaluate this interpretation, the response in the regions reported here needs to be tested to stimuli that contain sentence-level prosodic contours but do not contain any linguistic information (e.g., using degraded auditory materials like those used in Meyer, Alter, Friederici, Lohmann, & von Cramon, 2002). A robust response to such stimuli would suggest sensitivity to prosodic contours rather than reinforcement of the structural information in linguistic stimuli.

And *third*, we know that these regions are not part of the domain-general fronto-parietal 'multiple-demand' network (e.g., Duncan, 2010) whose regions respond across a wide range of demanding cognitive tasks: regions of the multiple-demand network respond more to unstructured compared to structured stimuli (Fedorenko, Duncan et al., 2012; Fedorenko et al., 2013). Again, this is in spite of the fact that demanding cognitive tasks frequently activate posterior inferior temporal cortices (e.g., Duncan, 2006).

Table 2. Effects of sensitivity to structure in the visual and auditory conditions. We report uncorrected p values, but the effects of structure in the auditory conditions would remain significant after a Bonferroni correction for the number of regions (even including all 12 regions discovered by the GSS method).

|                                                   | Left hemisphere               |                              | Right hemisphere             |                             |
|---------------------------------------------------|-------------------------------|------------------------------|------------------------------|-----------------------------|
|                                                   | TempPole                      | PostInfTemp                  | TempPole                     | PostInfTemp                 |
| df                                                | 9                             | 8                            | 8                            | 8                           |
| 1. Effect of structure in the visual conditions   | t < 1.24;                     | <i>t</i> <1;                 | <i>t</i> <1;                 | <i>t</i> <1;                |
| 2. Effect of structure in the auditory conditions | n.s. $t = 9.42$ ; $p < .0001$ | n.s. $t = 5.98;$ $p < .0005$ | n.s. $t = 4.85$ ; $p < .001$ | n.s. $t = 4.63;$ $p < .001$ |

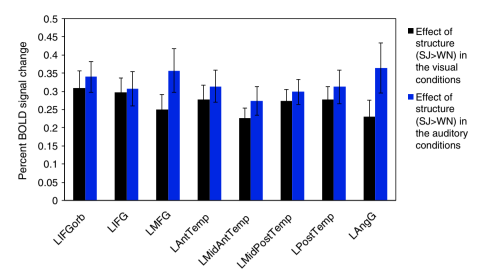


Figure 3. Sensitivity to structure in the visual vs. auditory conditions in brain regions sensitive to high-level linguistic processing (defined in individual subjects using the sentences > pseudoword lists contrast; Fedorenko et al., 2010). (The responses are estimated using n-fold cross-validation, so that data to define the fROIs and estimate the responses are independent).

So what do we think the prosody-sensitive regions discovered here might be doing? We hypothesise that at least some of the regions we report here may store typical prosodic contours. Their response may thus be driven by how well the prosodic contour of an incoming stimulus matches these stored prosodic 'templates'. This proposal is similar to Peretz et al.'s proposal for some of the music-sensitive brain regions, which are proposed to store pitch / rhythm patterns characteristic of the music that the person has been exposed to (e.g., Peretz & Coltheart, 2003; also Fedorenko, McDermott et al., 2012).

In order to better understand the acoustic features that may be associated with sentence-level prosodic contours, we performed a series of acoustic analyses on the auditory materials used in the current study. In particular, we identified word/pseudoword boundaries in each audio file using the Prosodylab-Aligner tool (Gorman, Howell, & Wagner, 2011), extracted a set of acoustic features from each word/pseudoword using Praat (Boersma & Weenink, 2005), and then analysed those features statistically. The procedure and the results are described in detail in Appendix C, but we here highlight the key results.

Although, as intended in creating these materials, the structured and unstructured stimuli turned out to be quite well matched on a number of acoustic dimensions (e.g., mean pitch, a falling pitch and intensity pattern across the stimulus, etc.), we did observe some differences. First, the maximum pitch was higher, the minimum pitch was lower, and the power was lower in the structured compared to the unstructured conditions. And second, there was greater variability across the stimulus in the structured conditions than in the unstructured conditions with respect to duration, maximum pitch, and centre pitch, and lower variability with respect to power.

These observed acoustic differences between structured and unstructured conditions provide some preliminary hints about the relevant features of the sentence-level prosodic contours that may be contributing to or driving the fMRI effects. As discussed above, in order to understand how exactly these regions contribute to prosodic processing, many studies are needed that would examine the responses of these functionally defined regions to a variety of new stimuli and tasks. For example, by manipulating different acoustic features above separately in a controlled fashion we would be able to narrow in on the ones that contribute the most to the observed effects.

In terms of relating the current findings to previous work on prosody, the superior temporal pole regions look similar to regions that have been reported in previous neuroimaging studies (e.g., Humphries et al., 2005; Mayer et al., 2002). For example, Humphries et al. (2005) reported some regions in the vicinity of superior temporal pole that respond more to stimuli with sentence-like prosody (compared to those with list prosody) even in cases where the stimuli were not linguistically meaningful. That would be consistent with the possible interpretation above, that these regions store typical prosodic 'templates' and respond when there is a match between a stimulus and these stored representations. The regions in the posterior inferior temporal lobes have not been previously reported in investigations of prosody to the best of our knowledge, except for one unpublished study (Davis et al., 2010).

A number of previous studies have reported activations in and around the regions we observed in the current study for linguistic contrasts that do not appear to have anything to do with prosodic processing. For example, activations in or near the superior temporal pole regions

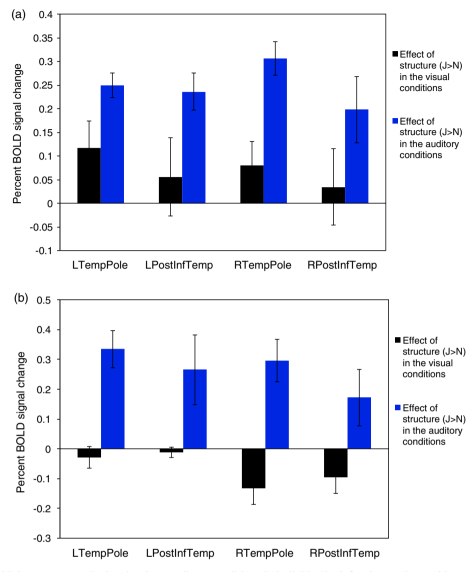


Figure 4. (a): Sensitivity to structure in the visual vs. auditory conditions in individually defined prosody-sensitive regions, for the subset of conditions consisting of real words: sentences and word lists). The fROIs are defined by a conjunction of two contrasts: (1) a greater response to sentences than word lists in the auditory conditions, and (2) no difference between sentences and word lists in the visual conditions. (Here, as in all the other analyses, the responses are estimated using n-fold cross-validation, as discussed in Methods, so that data to define the fROIs and estimate the responses are independent.) The Sentences > Word lists contrast is significant for the auditory conditions in all four ROIs (ps<.0005, except for RPostInfTemp whose p<.05). The Sentences > Word lists contrast is not significant for the visual conditions (except for LTempPole where it reaches significance at p<.05). (b): Sensitivity to structure in the visual vs. auditory conditions in individually defined prosody-sensitive regions, for the subset of conditions consisting of pseudowords: Jabberwocky sentences and pseudoword lists). The fROIs are defined by a conjunction of two contrasts: (1) a greater response to Jabberwocky sentences than pseudoword lists in the auditory conditions, and (2) no difference between Jabberwocky sentences and pseudoword lists in the visual conditions. The Jabberwocky > Pseudoword-lists contrast is significant for the auditory conditions in three of the four ROIs (ps<.05) and marginal in RPostInfTemp (p=.062). The Jabberwocky > Pseudoword-lists contrast is not significant for the visual conditions in any of the ROIs.

have been reported for violations of syntactic structure (e.g., Friederici, Rüschemeyer, Hahne, & Fiebach, 2003) and effortful speech perception (e.g., Adank, 2012). And parts of the inferior posterior temporal cortex have been implicated in visual word recognition (e.g., Baker et al., 2007; Cohen et al., 2000; Petersen, Fox, Snyder, &

Raichle, 1990; Polk & Farah, 1998; for a review see e.g., McCandliss, Cohen, & Dehaene, 2003), recalling wordforms in non-alphabetical languages (e.g., Kawahata, Nagata, & Shishido, 1988; Nakamura et al., 2000; Soma, Sugishita, Kitamura, Maruyama, & Imanaga, 1989) and spelling (e.g., Rapcsak & Beeson, 2004). In examining

these studies, we should keep in mind that there is no way to determine with certainty whether or not activated regions are the 'same' as the regions we report here, as discussed in the Introduction. To find that out, one would need to examine activations for the different contrasts within the same individuals. Nevertheless, examining activations observed in and around the regions discussed here may be important in generating hypotheses to be evaluated in future work.

## How good is the current 'localiser' as a localiser for prosody-sensitive regions?

We want to stress that the 'localiser' task proposed here is certainly not the only way to find prosody-sensitive brain regions, and we are not even arguing that this is necessarily the *best* contrast to use. The goal of the current study is to provide a proof of concept. In particular, we show that it is possible to define prosody-sensitive regions (with stable functional profiles, as indicated by the replicability of the effects with acrossruns cross-validation) in individual subjects. This result suggests that the functional localisation approach is feasible in prosody research.

As with any localiser, before investigating the responses of the functional ROIs reported here to new conditions, it is important to first establish the robustness of this localiser contrast. In particular, a good localiser should not be sensitive to changes in the specific materials used, in the particular speaker used or in the specific task. Furthermore, only a subset of the conditions from the current study may suffice for fROI definition in future investigations. For example, in Figure 4, we show that qualitatively and quantitatively similar profiles obtain when only four of the eight conditions are used (i.e., only meaningful materials, i.e., sentences and word lists visually and auditorily presented - Figure 4a, or only meaningless materials, i.e., Jabberwocky sentences and pseudoword lists visually and auditorily presented -Figure 4b).

Other localiser contrasts might work just as well or better. There are different approaches to developing a functional localiser for a particular mental process / set of mental processes. One can imagine initially 'casting a wide net' with a broad functional contrast. In the extreme, you can imagine starting with something like 'listening to sentences > fixation'. This contrast activates a wide range of brain regions including those in the primary and secondary auditory cortices, the whole language network (e.g., Fedorenko et al., 2010) and some of the regions in the 'multiple-demand' network (e.g., Duncan, 2010). Across many studies, one could then try to narrow in on the parts of this extended set of brain regions that may be more specifically engaged in dealing with prosody by examining the responses of these various regions to new

conditions. In the current study, we chose a narrower starting point. As discussed in the beginning of the paper, this contrast may not (and almost certainly does not) include all of the brain regions important for prosodic processing, which may or may not be a problem, depending on the goals of a particular research study / programme. For example, brain regions that respond to implicit prosody would not be included in the current set given the design we chose.

As with any experimental approach, many possibilities are perfectly valid in using the functional localisation approach, as long as the researcher is (1) clear about what mental process(es) the localiser contrast targets and (2) careful in interpreting the results in line with all the possibilities for what the regions could be responding to (see Fedorenko et al., 2010; Saxe et al., 2006, for discussions). For example, Wiethoff et al. (2008) have reported stronger neural responses in the right middle STG for emotional compared to neutral prosodic stimuli. However, they then demonstrated that this greater response can be explained by a combination of arousal and acoustic parameters like mean intensity, mean fundamental frequency and variability of fundamental frequency. As a result, the region in the STG is engaged during the processing of emotional prosody but only by virtue of the acoustic characteristics of those stimuli. For any candidate prosody-sensitive region – including those reported here - it is important to consider all of the possible alternatives for what could be driving the response to prosody. The functional localisation approach is perfectly suited for doing so, allowing for easy comparisons across studies and labs.

Some general guidelines for creating powerful yet quick 'localisers' include the following (for general advice on fMRI designs, see e.g., Huettel, Song, & McCarthy, 2008):

- (1) Use a blocked, not event-related, design (blocked designs are much more powerful due to the additive nature of the BOLD signal; e.g., Birn et al., 2002; Friston, Zarahn, Josephs, Henson, & Dale, 1999).
- (2) Use as few conditions as possible (from that perspective, the localiser used in the current experiment is not ideal; this is because this study was originally designed to address different research questions than the one asked here).
- (3) Given that the recommended block length is between ~10 and ~40 sec (e.g., Friston et al., 1999), use blocks that are as short as possible within this range. However, this choice also depends on how long individual trials are, because it is also advisable to include as many trials in a block as possible. Typical blocks are between 16 and 24 sec in duration.

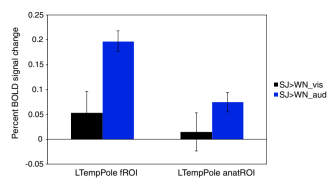


Figure 5. A comparison of the effects in the left temporal pole for individually defined functional ROIs vs. for the subject-independent anatomical ROI.

- (4) Unless the manipulation you are examining is very subtle (which is probably not a good idea for a localiser contrast anyway), 10–12 blocks per condition is generally sufficient to obtain a robust effect.
- (5) It is generally advisable to distribute the blocks across two or more 'runs' so that data can be easily split up for cross-validation (i.e., using some portion of the data to define the regions of interest, and the other portion to estimate the responses; see also Coutanche & Thompson-Schill, 2012).

Keeping in mind (1) the guidelines above, and (2) the discussion at the beginning of the Experiment section, the best contrast for identifying prosody-responsive brain regions in future work may be a two-condition contrast between prosodically 'intact' and prosodically degraded stimuli (e.g., Humphries et al., 2005; Newman et al., 2010; Wiethoff et al., 2008; Zhao et al., 2008). Our recommendation is also to remove lexical content and syntactic structure from the contrast by using either speech in an unfamiliar language or speech that has been filtered so as to remove linguistic information.

### Do we really need individual functional ROIs?

To illustrate the importance of using individual fROIs, consider Figure 5 where we show the response obtained with individually defined fROIs for the region in the left superior temporal pole (same data as in Figure 2) vs. an anatomical region in a similar location where we simply extract the response from all the voxels in that ROI in each subject (see Fedorenko et al., 2010; Fedorenko, Nieto-Castañón et al., 2012; Saxe et al., 2006, for similar demonstrations). As can be clearly seen, although the general patterns of response are similar, the effects are much larger and more statistically robust for individually defined fROIs. Given that individual fROIs constitute

only a small portion of the relevant parcel (see Table 1), this is not surprising: in the case of the subject-independent anatomical ROI many voxels that are included in the analysis do not have the right functional properties and thus 'dilute' the effects (see Nieto-Castañon & Fedorenko, 2012).

### Future research

The ability to define prosody-sensitive regions functionally in individual subjects opens the door to a research programme investigating the functional profiles of these (or other functionally defined prosody-sensitive) regions in an effort to understand their contributions to prosodic processing. For example, it is important to discover the necessary and sufficient features that a stimulus must possess in order to elicit a response in these regions. This question applies both to linguistic stimuli (e.g., different types of prosodic contours) and non-linguistic stimuli. For example, if some of these regions indeed store prosodic 'templates' (i.e., commonly encountered prosodic patterns), we can probe the nature of these representations. We can ask how long these prosodic chunks have to be to elicit a response in these regions, by presenting foreign or low-pass filtered speech split up into prosodic patterns of various durations. Or we can ask how abstract these representations are, by examining adaptation effects in these regions to the same/similar prosodic patterns produced by speakers with different voices or presented at different speeds.

Cognitive tasks across several domains have been shown to activate regions in and around temporal poles (see e.g., Olson, Plotzker, & Ezzyat, 2007 for a review), including music (e.g., Fedorenko, McDermott et al., 2012; Peretz & Zatorre, 2005), social cognition (e.g., Fletcher et al., 1995) and abstract semantics (e.g., Patterson, Nestor, & Rogers, 2007). It is possible to construct multiple hypotheses about possibly shared computations between prosody and each of these other domains, especially music and social cognition (see some of the reviews cited in the introduction for discussions of some ideas along these lines). Examining the response of the regions reported here to musical stimuli and to non-prosodic socially-salient stimuli is necessary for evaluating such hypotheses.

Once we make progress in functionally characterising these regions, we can start investigating the relationship between these regions and other regions / networks in the brain, by examining anatomical connectivity (e.g., using DTI) and functional resting-state correlations (e.g., Fox & Raichle, 2007). In particular, we can examine the relationship between prosody-sensitive regions and primary auditory regions (e.g., Morosan et al., 2001), regions that have been implicated in pitch processing (Patterson, Uppenkamp, Johnsrude, & Griffiths, 2002), language regions (Fedorenko et al., 2010), multiple-demand regions

(Duncan, 2010) and regions that support social cognition (Saxe & Kanwisher, 2003), among others.

Multi-voxel pattern analyses (e.g., Norman, Polyn, Detre, & Haxby, 2006) might also prove valuable in studying prosody-sensitive regions. In particular, as discussed in the introduction, there is no reason to necessarily expect differences in the mean BOLD response for different types of prosodic contours. However, a region that is important in prosodic processing may be able to distinguish among these various prosodic contours in its fine-grained pattern of spatial activity, in spite of showing similar BOLD responses. This method can thus be used to ask which distinctions are represented in each of these regions.

Furthermore, once we have sufficiently narrowed down the range of hypotheses about the functions of these regions, we can use these regions as targets for transcranial magnetic stimulation (TMS) to investigate their causal role in various computations.

#### Conclusion

We have here argued that fMRI in general, and the functional localisation approach in particular, holds promise for asking and answering theoretically important questions in the domain of prosody. We have presented one possible functional localiser for identifying prosodysensitive brain regions in individual subjects, demonstrating the feasibility of this method for investigating prosodic processing. We hope that this approach can help shift the field from asking questions about *where* something happens in the brain to *how* it happens.

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### Notes

- In principle, studies that ask questions like 'does a particular manipulation activate brain region x?' could also inform deep issues in cognitive science, but this is only possible in cases where region x is characterised sufficiently well to serve as a neural 'marker' of a particular mental process. With a few exceptions, most brain regions lack such detailed functional characterisation and thus are not suitable for use as markers of particular mental processes (see e.g., Poldrack, 2006).
- 2. Note that this experiment was not originally designed to study prosodic processing. Hence the inclusion of both meaningful (Sentences, Word lists) and meaningless (Jabberwocky, Pseudoword lists) conditions may seem unmotivated. However, we think it ends up being a *strength* of this experiment to be able to generalise across the presence of meaning in the stimuli: as we will show in the Results, similar patterns hold for meaningful and meaningless conditions when examined separately.
- One important caveat to keep in mind is that individuals may differ with respect to how strongly they activate prosodic representations during silent reading. In the extreme case, if an individual activated prosodic representations during silent reading to the same degree as during auditory linguistic processing, then the contrast proposed here would fail to identify any prosody-sensitive regions in that individual. As will be shown below, the proposed contrast successfully identifies regions with the specified functional properties in the majority of individuals, suggesting that a substantial proportion of individuals have brain regions that respond more to the presence of structure in the auditory stimuli than in the visual stimuli (which we argue plausibly reflects sensitivity to sentence-level prosody). Once these prosody-sensitive brain regions are established as robust to irrelevant differences in the materials, task, etc., investigating individual differences in their response profiles and relating them to behaviour will be a fruitful avenue for future research.
- Although in the remainder of the paper we will use the term 'prosody-sensitive', the reader should keep in mind that we are referring to brain regions that are sensitive to sentencelevel prosodic contours.
- 5. The dataset used for the current study is the same dataset as that used in Experiment 3 in Fedorenko et al. (2010).
- Not being able to define a fROI in every single subject using the fixed-threshold approach - i.e., when parcels are intersected with thresholded individual activation maps is not uncommon. For example, when developing a localiser for high-level language regions, Fedorenko et al. (2010) considered a region meaningful if it could be defined in 80% or more of individual participants (see also Julian et al., 2012, where 60% or more of individual participants is used as a criterion for selecting meaningful high-level visual regions). An alternative that would enable one to define a fROI in every single subject would be to move away from the fixed-threshold approach. In particular, once a region has 'established itself' (i.e., once we know that it emerges consistently across people, is stable within individuals, and is robust to various properties of the localiser contrast), we can simply take the top – with respect to the t-values for the relevant functional contrast - 5% or 10% of voxels within some spatial constraint (defined anatomically or with the use of functional parcels obtained from a GSS analysis). This approach ensures that (1) a fROI is defined in every individual (and thus the results are generalisable to the

whole population, as opposed to the proportion of the population for whom the fROIs could be defined), and (2) fROIs are of the same size across individuals (see Nieto-Castañon & Fedorenko, 2012, for a discussion). The reason that we used the fixed-threshold approach in the current paper is that it is mathematically not trivial to use the top-n-voxels approach for the conjunction of multiple contrasts, which is what we use in the current study.

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### Appendix A

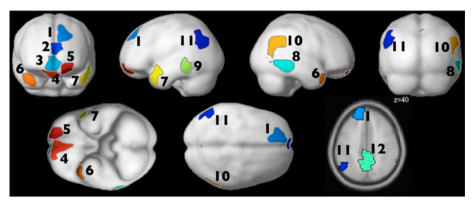


Figure A1. Twelve parcels that satisfied the functional criteria (i.e., a selective response to structure in the auditory stimuli; see Methods for details) and were present in at least 9/12 subjects. (Regions #6-9 are the regions discussed in the main body of the paper and shown in Figures 1 and 2.)

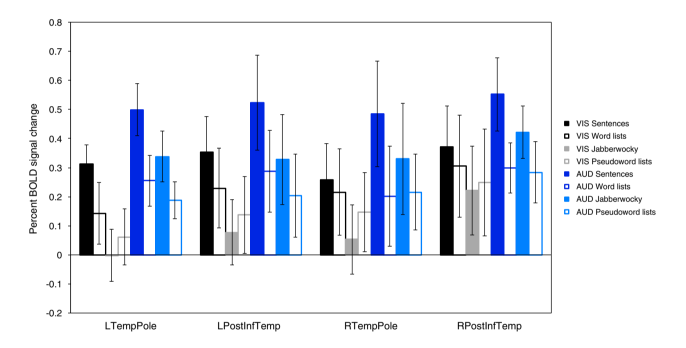


Figure A2. Responses of the four regions shown in Figure 2 to the individual experimental conditions.

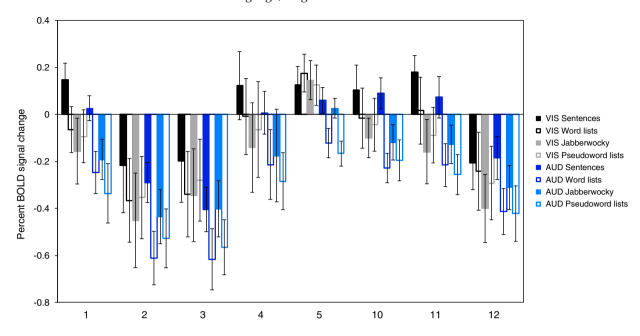


Figure A3. Responses of the regions not shown in Figure 2 (i.e., regions #1-5, 10–12 in Figure A1) to the individual experimental conditions. Region numbers correspond to the numbers of the regions in Figure A1.

### Appendix B

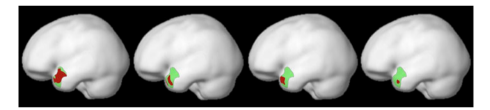


Figure B1: Sample individual fROIs for the Left Temporal Pole region. The parcel is shown in green, and the individual activations are shown in red.

Appendix C

Pitch tracks for the four conditions of two sample items

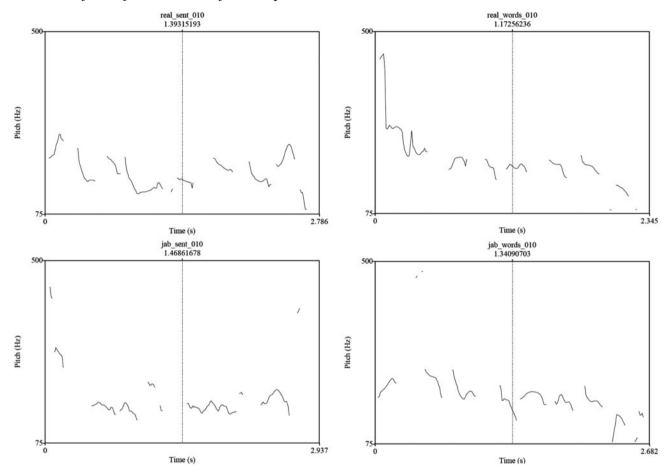


Figure C1a: Pitch tracks for Item #010 (real\_sent = sentence; real\_words = word list; jab\_sent = Jabberwocky sentence; and jab\_words = pseudoword list).

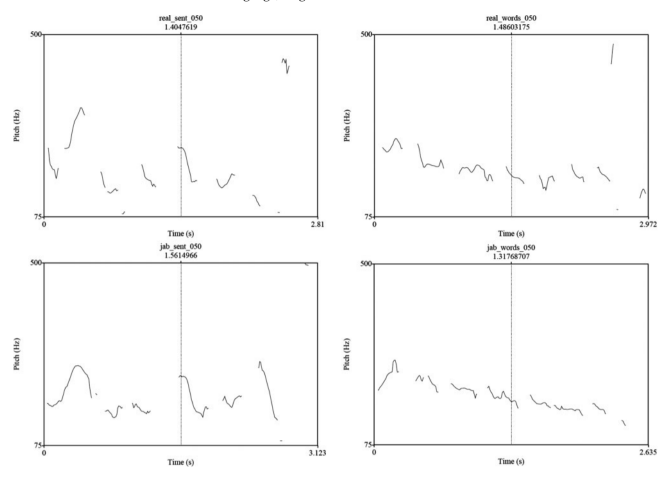


Figure C1b: Pitch tracks for Item #050 (real\_sent = sentence; real\_words = word list; jab\_sent = Jabberwocky sentence; and jab\_words = pseudoword list).

### Acoustic analyses of the experimental materials

### Methods

The analysis was performed in three steps:

1. The audio files were aligned with the corresponding text files to identify the word/pseudoword boundaries in each audio file using the Prosodylab Aligner tool (Gorman et al., 2011).

The alignment was performed following the instructions provided at http://prosodylab.org/tools/aligner/. The aligner uses the CMU pronouncing dictionary (available at http://www.speech.cs.cmu.edu/cgi-bin/cmudict). A few words from the Sentences (n = 4) and Word-lists (n = 6) conditions were not in the dictionary. And, as expected, a large number of pseudowords in the Jabberwocky sentences and Pseudoword-lists conditions were not in the dictionary (544 and 547, respectively). Consequently, a total of 686 unique words were added to the dictionary. Their pronunciation was generated by the tools available at http://www.speech.cs.cmu.edu/tools/lextool.html.

To align the audio files to the text files, the aligner takes as input a \*.wav file and a \*.lab file (i.e., the text file with the orthographic words that are matched to the items in the dictionary), and it outputs a \*.TextGrid file, which has the forced time divisions of the \*.wav file for each word (and phoneme) in the \*.lab file.

The aligner was unsuccessful in forcing the alignment for a small number of items in each condition (1 in the Sentences condition, 1 in the Word-lists condition, 18 in the Jabberwocky sentences condition, and 17 in the Pseudoword-lists condition). A plausible reason for this is that the acoustic pattern has to be fairly close to the phonetic form of the word in the dictionary, and some of the pseudowords may have been pronounced quite differently from the pronunciation generated by the dictionary. All the analyses were thus performed on 127 sentences, 127 word lists, 110 Jabberwocky sentences, and 111 pseudoword lists.

2. A set of acoustic features was extracted from each word/pseudoword in each aligned audio file using Praat (Boersma & Weenink, 2005).

The extraction was performed with a Praat script written by Michael Wagner. This script takes as input a \*.wav file and a \*.TextGrid file and outputs a set of pre-specified acoustic measures.

We extracted the following features for each word/pseudoword:

### Duration measure:

• duration (s) [word/pseudoword duration]

### Pitch measures:

- mean pitch (Hz) [mean fundamental frequency (F0) of the word/pseudoword]
- maximum pitch (Hz) [highest F0 observed in the word/pseudoword]
- minimum pitch (Hz) [lowest F0 observed in the word/pseudoword]
- initial pitch (Hz) [mean F0 of the first 5% of the word/pseudoword]
- centre pitch (Hz) [mean F0 of the 5% of the word/pseudoword centred on the midpoint of the word/pseudoword]
- final pitch (Hz) [mean F0 of the last 5% of the word/pseudoword]

### Other measures:

- mean intensity (dB) [mean loudness level in the word/pseudoword]
- maximum intensity (dB) [highest dB level observed in the word/pseudoword]
- minimum intensity (dB) [lowest dB level observed in the word/pseudoword]
- power (Pascal<sup>2</sup>) [power across the entire word/pseudoword]
- 3. Several statistical tests were performed on the extracted values.

Analyses reported here were conducted with the lme4 package (Bates, Maechler, & Dai, 2008) for the statistical language R (R Development Core Team, 2008).

We asked the following two questions:

- i. Averaging across the eight words/pseudowords in a sentence/word list, do conditions differ with respect to any of the acoustic features above? In particular, is there a difference between structured conditions (Sentences + Jabberwocky sentences) and unstructured conditions (Word lists + Pseudoword lists)? For exploratory purposes, we also examined the changes in the acoustic features over the course of the sentence / word list.
- ii. Do conditions differ with respect to how much they vary over the course of the sentence / word list in any of the acoustic features above?

### Results

The figures that summarise the results of the analyses appear below.

First, for each acoustic feature we show the mean value (averaged across words/pseudowords within a stimulus, and then averaged across trials within each condition) in a bar graph on the left, and the word-by-word values in a line graph on the right in Figures C2a-i. To quantify the differences, we conducted a series of linear mixed-effects regressions with the two experimental factors (structure: sentences vs. word lists; and lexical content: real words vs. pseudowords) as fixed effects, and items as random effects (we included both an intercept and a slope for each item following Barr et al., 2013). Of most interest to the current investigation are potential differences between structured and unstructured conditions. (We find a few differences between real words and pseudowords, but we don't discuss these here given that all of the key fMRI analyses focus on the structured vs. unstructured contrast collapsing across real words and pseudowords as in Figure 2, or treating them separately as in Figure 4a and b.)

It is worth noting that the creation of the materials was successful in that the structured and unstructured conditions are quite well matched in a number of ways. In particular, they do not differ in terms of mean pitch, initial, centre or final pitch, and maximum or minimum intensity (all ts < 1, except for centre pitch where t = |1.73|). Furthermore, the changes in various features over the course of the stimulus (see line graphs in the figures below) also look similar across structured and unstructured conditions, at least in their general trends. For example, the pitch and intensity values decrease over the course of the sentence / word list and the duration values increase at the end of the sentence / word list.

However, we do find a few differences between structured and unstructured conditions, and these differences plausibly contribute to the effects we observe in fMRI. Because we are examining 11 acoustic features, a correction for multiple comparisons is needed. We chose the most stringent, Bonferroni, correction: with 11 tests, the p value has to be smaller than 0.05/11 = 0.0045. With respect to pitch, we find that words/pseudowords in the structured conditions are produced with higher maximum pitch (t = |5.71|) and lower minimum pitch (t = |6.32|). Furthermore, words/pseudowords in the structured conditions are produced with less power than in the unstructured conditions (t = |3.88|). With respect to duration, we find that on average, words/pseudowords in the structured conditions are longer, but the effect doesn't reach significance at the corrected p level (p < 0.02), and in any case, as can be seen in Figure C2a, structured-ness appears to interact with real word / pseudoword dimension, so this difference is unlikely to contribute to the observed differences in fMRI.

In all the graphs below, the error bars represent standard errors of the mean over items. For the line graphs, the error bars were computed as follows: the standard deviation for all the structured (or unstructured, for the unstructured conditions) items was divided by the square root of the number of structured (or unstructured) items. Note also that in all the line graphs below, we averaged condition means for the real words and pseudoword conditions to get each of structured and unstructured values. The reason for doing that (instead of averaging across items) is that the number of items differs slightly between conditions – see Methods part 1 above.

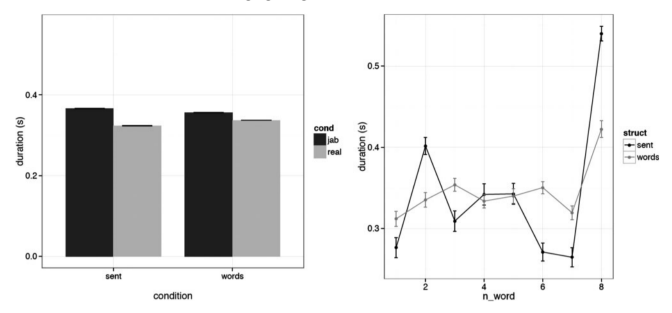


Figure C2a: Left: Average *duration* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *duration* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

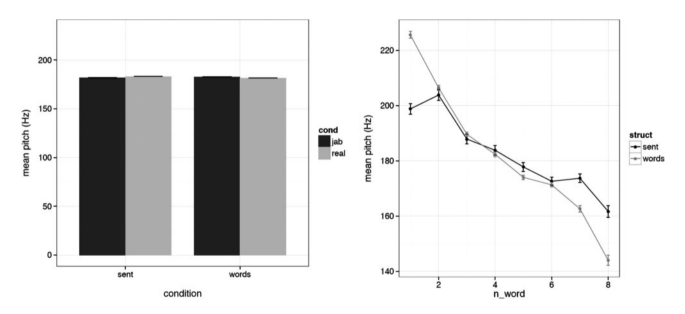


Figure C2b: Left: Average *mean pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *mean pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

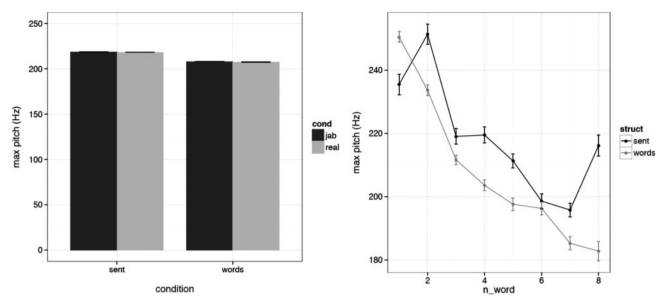


Figure C2c: Left: Average *maximum pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *maximum pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

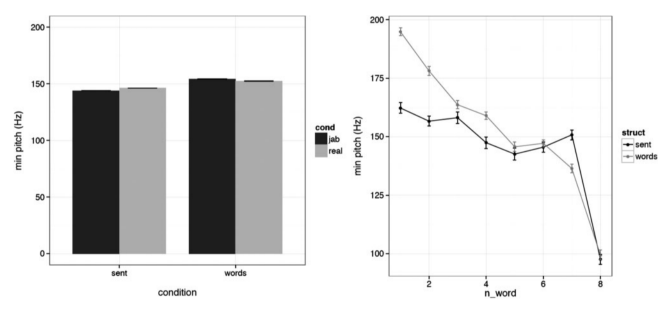


Figure C2d: Left: Average *minimum pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *minimum pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

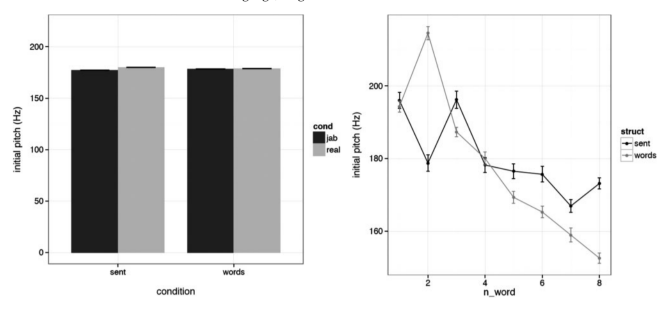


Figure C2e: Left: Average *initial pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *initial pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

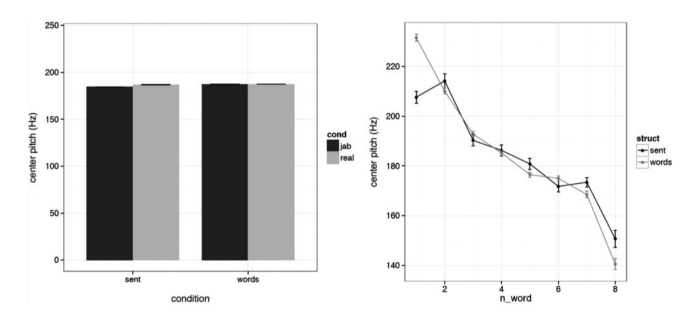


Figure C2f: Left: Average *centre pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *centre pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

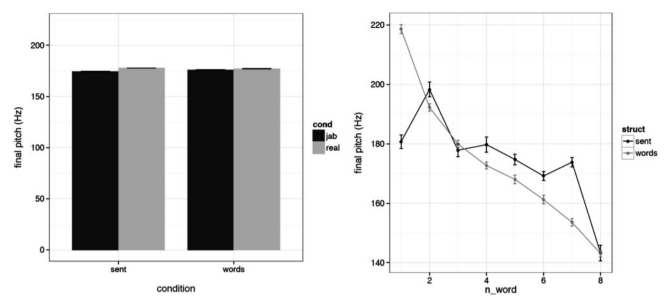


Figure C2g: Left: Average *final pitch* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *final pitch* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

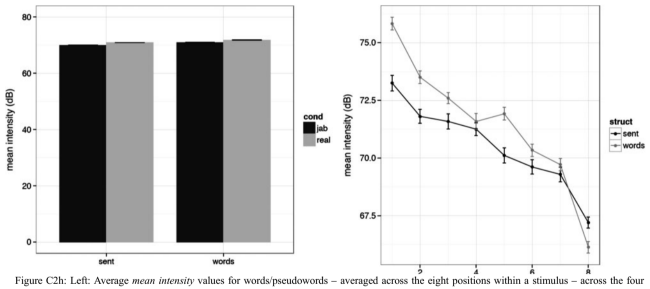


Figure C2h: Left: Average *mean intensity* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *mean intensity* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

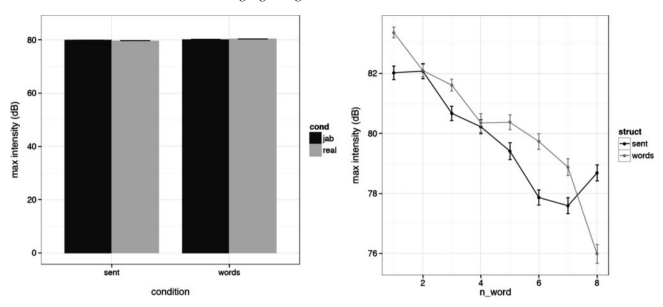


Figure C2i: Left: Average *maximum intensity* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *maximum intensity* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

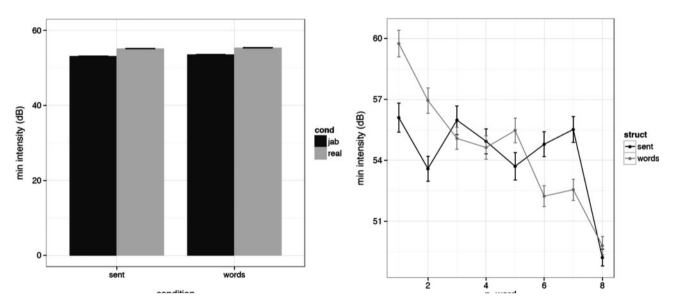


Figure C2j: Left: Average *minimum intensity* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *minimum intensity* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

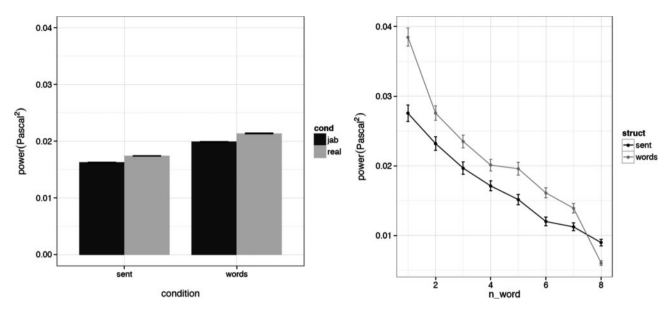


Figure C2k: Left: Average *power* values for words/pseudowords – averaged across the eight positions within a stimulus – across the four conditions (sentences with real words, Jabberwocky sentences, lists of real words, lists of pseudowords). Right: Average *power* values for words/pseudowords at each of the eight positions for the structured (black) vs. unstructured (grey) conditions.

And second, we also find some differences between structured and unstructured conditions in terms of the amount of variability across the eight words/ pseudowords. As in the first analysis, because we are examining 11 acoustic features, we are applying a Bonferroni correction, and the p value has to be smaller than 0.05/11 = 0.0045. With respect to duration, we find that words/pseudowords in the structured conditions are more variable over the course of the stimulus (t = |10.33|). With respect to pitch, the structured and unstructured conditions do not differ significantly with respect to variability in mean pitch, minimum pitch, initial pitch, or final pitch (all ts < 1.5). However, the words/pseudowords in the structured conditions are more variable in terms of maximum pitch (t = |4.28|) and centre pitch (t = |5.41|). None of the three intensity measures reveal significant differences between structured and unstructured conditions, although there is a trend for more variability in minimum intensity in the structured conditions. Finally, we find that words/pseudowords in the structured conditions are less variable over the course of the stimulus in terms of power (t = |4.29|).