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Fronto-parietal network supports context-dependent speech comprehension

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1. Introduction

Most of us have experienced instances of joining a conversation without knowing the topic: while being able to recognise individual words and sentences, it is difficult to comprehend what is being said when lacking information about the context within which the conversation takes place. However, as soon as the topic is revealed, the subsequent as well as preceding conversation becomes clear. Previous behavioural studies have established that discourse comprehension requires knowledge of the relevant context (Bransford & Johnson, 1972; Dooling & Lachmann, 1971; Swinney, 1979). Processing of new semantic information is facilitated by predictions, based on currently available semantic representation (i.e. schema) of the narrative context. When predictions are mismatching with the incoming narrative – for example when one’s conversation partner abruptly changes the topic – they impair understanding, as the inappropriate predictions are violated by the speech stream (Wlotko & Federmeier, 2012).

Linguistic processing during context-dependent comprehension requires a number of components: mental lexicon and associated retrieval operations, selection and integration of lexical elements into larger structures (unification), as well as attention and control/error monitoring (see, e.g., the MUC model, Hagoort, 2005). During natural speech comprehension, these operations may form a loop where incoming information is integrated with previous knowledge as early as possible (Wlotko & Federmeier, 2012). But since spoken discourse unfolds over time, critical contextual information may often be available relatively late, therefore increasing the relevance of knowing context in advance. Contextual effects in natural speech comprehension thus provide a realistic model for studying the changes in brain networks subserving semantic processing. Semantic processing on local (word or sentence) and global (narrative) levels may proceed differently. Here we study the neural mechanisms underlying use of contextual information to resolve semantic ambiguity at the narrative level.
1.1. Anatomical and functional basis of the linguistic processing network

The classic linguistic processing network comprises Broca’s area in the left inferior frontal gyrus (IFG), and left inferior parietal cortex (IPC) encompassing angular (AG) and supramarginal gyri (SMG) that together with left middle (MTG) and superior temporal gyri (STG) form the classical Wernicke’s area (Binder, Desai, Graves, & Conant, 2009). Broca’s area is involved in a wide range of phonological, syntactic and semantic functions, including semantic selection (Bookheimer, 2002; Hagoort, 2005; Thompson-Schill, D’Esposito, Aguirre & Farah, 1997; Vigneau et al., 2006). Wernicke’s area, in turn, is involved in both local and global semantic processing, e.g. initial access to and retrieval of semantic representations of words (Bookheimer, 2002; Jung-Beeman, 2005; Mesulam, 1998), as well as semantic integration during discourse processing (Binder et al., 2009; Humphries, Binder, Medler, & Liebenthal, 2006; Humphries, Binder, Medler, & Liebenthal, 2007). While the left hemisphere dominates in linguistic functions, the contribution of the right hemisphere is also significant (Bookheimer, 2002; Manenti, Cappa, Rossini, & Minussi, 2008; Zempleni, Haverkort, Renken, & Stowe, 2007b; Zempleni, Renken, Hoeks, Hoogduin, & Stowe, 2007a). The exact boundaries of both Wernicke’s (Binder et al., 2009; DeWitt & Rauschecker, 2012; Mesulam, 1998) and Broca’s (Amunts et al., 1999, 2010) areas are still debated, but most researchers agree that key aspects of language processing are undertaken in this fronto-temporo-parietal language system, whose activation profile varies depending on current linguistic processing demands (Abrams et al., 2013; Tyler & Marslen-Wilson, 2008).

Several studies have confirmed the anatomical connections between Broca’s area and superior/middle temporal gyrus as well as IPC (Catanì, Jones, & Fytche, 2005; Frey, Campbell, Pike, & Petrides, 2008; Hickok & Poeppel, 2004; Saur et al., 2008; Turken & Dronkers, 2011). The dorsal pathway, which runs from premotor cortex (including BA 44 in Broca’s area) via arcuate fasciculus/ superior longitudinal fasciculus (AF/SLF) to IPC and STG, contributes to sensory-motor mapping of sounds to articulations. The ventral pathway connects prefrontal cortex and BA 45 in Broca’s area to middle temporal lobe via the extreme capsule fibre system (ECFS) and contributes to linguistic processing of sounds and mapping sounds to meanings (Friederici, 2012; Hickok & Poeppel, 2004; Rauschecker & Scott, 2009). It was also suggested that the dorsal pathway not only supports auditory-to-motor mapping, but also processing of syntactically complex sentences (Friederici, 2009, 2012; Friederici, Bahnhim, Hein, Schubotz, & Anwander, 2006) and can be segregated functionally into phonological and lexical-semantic processing systems (Friederici, 2012; Glasser & Rilling, 2008).

1.2. Broca’s area’s role in contextual understanding

Broca’s area is a good candidate hub for the hypothesised network of brain areas involved in contextual understanding. Specifically, activation of anterior portion of Broca’s area (BA 45) is increased when processing sentences with high rather than low ambiguity (Rodd, Davis, & Johnsrude, 2005), and participants with lesions spanning Broca’s area (BA 44 and 45) fail to use contextually relevant information for facilitating selection of word meaning (Bedny, Hulbert, & Thompson-Schill, 2007). Broca’s area also supports selection among semantic information held in working memory, such as decisions about task-relevant attributes of word meaning (Gabrieli, Poldrack, & Desmond, 1998; Moss et al., 2005; Thompson-Schill, D’Esposito, Aguirre & Farah, 1997). These local semantic processing functions suggest that Broca’s area could contribute to contextual understanding by guiding selection and integration of the task-relevant text or speech features (Hagoort, 2005).

Broca’s area can be further divided into BA 44 and BA 45 (Anwander, Tittgemeyer, von Cramon, Friederici, & Knösche, 2007; Poldrack et al., 1999). In the anatomical connectivity signatures of BA 44, the dorsal AF/SLF pathway is more dominant, and for BA 45 the ECFS tracts are more prominent (Anwander et al., 2007; Frey et al., 2008). Functional imaging studies also support this division (Amunts et al., 2004; Cooper, Hasson, & Small, 2011; Gough, Nobre, & Devlin, 2005; Hasson, Nusbaum, & Small, 2007; Poldrack et al., 1999). For example, Hagoort (2005) proposed that the posterior portions (BA 44) contribute to phonological, middle portions (BA 44/45) to syntactic and anterior portions (BA 45 and 47) to semantic unification. In line with this, during written word processing the anterior–ventral part of Broca’s area supports semantic processing, whereas posterior–dorsal part contributes to both semantic and phonological processing (Poldrack et al., 1999). Hemodynamic activity in BA 44 is more correlated with regions involved in phonological processing, while activity in BA 45 correlates more strongly with regions attributed to syntactic and semantic processing, and it was suggested that unification components for each linguistic modality in Broca’s area had a corresponding integration/memory component in posterior temporal cortex (Xiang, Fonteijn, Norris, & Hagoort, 2010). Another study revealed stronger functional connectivity from BA 45 to posterior occipital and temporal cortices for semantic stimuli, i.e. words than for pseudowords, letters or false fonts, while BA 44 showed similar connectivity with occipital and temporal cortices during processing of both words and phonological stimuli (pseudowords and letters). While words, pseudo-words and letters activated both subregions, the connectivity patterns indicated anterior Broca’s area involvement in assessment of semantic representations, while posterior Broca’s area connectivity profile corresponds to access to phonological representations associated with words (Bokde, Tagamets, Friedman, & Horwitz, 2001). Taken together, these findings suggest that in particular the anterior parts of Broca’s area might be critical for context-dependent processing.

1.3. Parietal and cingulate cortices in contextual understanding

The IPC and AG are heteromodal areas that have been implicated in a wide range of linguistic functions. Bilateral IPC is involved in initial access to semantic representations of words, giving them meaning (Jung-Beeman, 2005; Mesulam, 1998) and left AG is involved in global semantic processing during integration of linguistic material (Binder et al., 2009; Humphries et al., 2006, 2007). Left AG was reported in studies addressing semantic decisions and integration of prior experiences, for example, while solving problems that require extended periods of computation, or planning of future behaviour (Binder et al., 1999), memorisation of semantic information (Hasson et al., 2007) and combinatorial processes (Humphries et al., 2007). Activation of left parietal regions, including AG, has been reported while subjects make word versus pseudo-word decisions, but also during memory-intensive narrative comprehension, and top-down predictions of semantic content (for review see Price (2012)). Moreover, activation in bilateral IPC (including AG) increases as a function of complexity of linguistic stimulation from words to sentences to narratives (Xu, Kemeny, Park, Frattali, & Braun, 2005), suggesting that bilateral IPC may be involved in complex linguistic processing. Taken together, these studies suggest that left IPC is involved in contextual speech processing. For example, left IPC may function as a part of a hierarchical semantic process. As relevant contextual information facilitates top-down predictions of the semantic content, left IPC could be recruited when a matching context is
provided during narrative listening. On earlier stages of linguistic processing left IPC might contribute to access, retrieval and maintenance of semantic representations of the verbal content. On subsequent stages of linguistic processing left IPC might support integration of linguistic information.

Finally, since contextual cues provide a cognitive filter for selection of information, constant error monitoring and control for consistency between the narrative and the contextual schema is required (Hagoort, 2005). Anterior cingulate cortex (ACC) is involved in a wide range of executive functions (Abutalebi et al., 2012; Paus, 2001), such as error monitoring (Botvinick, Cohen, & Carter, 2004; Carter, Braver, Barch, Botvinick, Noll & Cohen, 1998), selective attention (Crottaz-Herbertte & Menon, 2006), and inhibition of responses (Cabeza & Nyberg, 2000). Extensive structural and functional links between ACC and prefrontal cortical suggest that the ACC could subserve general cognitive control function during narrative processing (Hagoort, 2005; Paus, 2001). Accordingly, functional connections between Broca’s region and ACC might support monitoring of consistency between narrative and the context-providing schemas.

1.4. The current study

Here we extend the previous research on contextual effects and coherence of text and context (Maguire, Frith, & Morris, 1999; Martin-Loeches, Casado, Hernández-Tamames, & Alvarez-Linera, 2008; St George, Kutas, Martinez, & Sereno, 1999) by addressing both regional responses and functional connectivity of the brain’s language network during context-dependent speech processing. To delineate the cortical network for context-dependent speech comprehension, participants listened to spoken narratives where contextual processing is manipulated using pictorial context, while their haemodynamic brain activity was measured with fMRI. We hypothesised that listening to ambiguous narratives with a semantically mismatching context would increase semantic selection demands due to lacking contextual cues, which would be reflected in increased activation in Broca’s area. In addition, we hypothesised that Broca’s area would function as the central “hub” for the fronto-temporo-parietal brain network for context-dependent speech processing. In this network, Broca’s area is interacting with IPC to perform initial access and subsequent integration of semantic representations, and with ACC to monitor the consistency of narrative in relation to context.

2. Materials and methods

2.1. Participants

Twenty right-handed healthy adults (8 males, 12 females, ages 21–47 years, mean age 26 years) with normal or corrected to normal vision and normal hearing (self-reported) volunteered for the study. Individuals with a history of neurological or psychiatric diseases, or current medication affecting the central nervous system, were excluded. All subjects were compensated for their time, and signed informed consent forms, approved by the ethics-committee of the Helsinki and Uusimaa Hospital District.

2.2. Stimuli and experimental design

Stimuli consisted of 20 ambiguous narratives spoken in Finnish, with average duration of 62±10 s (Fig. 1). Each narrative described a complex sequence of actions such as those involved in doing laundry without providing contextual cues (e.g. without mentioning items such as washing machine, laundry soap, or actions such as washing or filling a machine) that would have revealed the action being described (c.f. Bransford and Johnson, 1972; see Appendix A for example; narratives with translations are available from corresponding author). The narratives were thus difficult to comprehend without access to relevant contextual information, but readily comprehensible when appropriate contextual cues were provided. The sentences followed the rules of Finnish grammar and contained no metaphorical vocabulary; thus their lexical meaning was accessible. In a pilot experiment, 30 independent subjects read 30 narratives preceded by either mismatching or matching picture. Subsequently they evaluated the comprehensibility of each narrative, as well as congruency of each narrative-picture pair. Twenty narratives with highest comprehension and congruency ratings in matching and lowest in mismatching condition context were selected for the fMRI experiment.

The selected narratives were recorded by a male voice in an acoustically shielded room. During fMRI, the stimuli were delivered using Presentation software (Neurobehavioral Systems Inc., Albany, California, USA). The narratives were played to the subjects with an UNIDES ADU2 audio system (Unides Design, Helsinki, Finland) via plastic tubes through porous EAR-tip (Etymotic Research, ER3, IL, USA) earplugs. Sound intensity was adjusted for each subject to be loud enough to be heard over the scanner noise. Contextual cue pictures were back-projected on a semitransparent screen using a 3-micromirror data projector (Christie X3, Christie Digital Systems Ltd., Mörchengladbach, Germany), and reflected via a mirror to the subject. The viewing distance was 34 cm, and the width of the projected image was 28 cm corresponding to approximately 39.5° in the visual field.

Participants listened to the narratives in a fixed, pseudo-random order while being scanned. The picture-narrative congruency was counterbalanced so that every participant received exactly the same auditory stimulation, yet the picture context for each narrative was different across the two groups. Each narrative was preceded by either matching or mismatching picture for 11 s, followed by a grey fixation cross that was shown while the narrative was played. After the narrative, the fixation cross changed to green for 4 s to mark the start of the next trial. During the cue trial the participants were instructed to look at the picture, and during the narrative trial they were instructed to focus their eyes on the fixation cross and listen to the speech while trying to comprehend its contents. The narratives were presented in two runs both consisting of 10 trials and lasting for 12.8 min with equal number of matching and mismatching trials. Each narrative was presented to each subject only once.

2.3. fMRI acquisition and preprocessing

MRI scanning was performed with General Electric Signa 3 T MRI scanner with Excite upgrade at the Advanced Magnetic Imaging Centre at the Aalto University School of Science. Whole-brain images were acquired with T2*-weighted echo-planar imaging (EPI), sensitive to blood oxygenation level-dependent (BOLD) signal contrast, using a quadrature sixteen channel head coil. The following parameters were used: 36 axial slices, 4 mm slice thickness; TR=1800 ms; TE=30 ms; Flip angle=75°; FOV=240 mm; voxel size 3×3×4 mm³; ascending interleaved acquisition with no gaps between slices. A total of 450 volumes were acquired in both runs, and the first 6 volumes of each run were discarded. T1-weighted structural images were acquired at a resolution of 1×1×1 mm³.

fMRI data processing was carried out using FEAT (FMRI Expert Analysis Tool) Version 5.98, part of FSL (FMRIB’s Software Library, www.fmrib.ox.ac.uk/fsl). First, functional images were realigned to the middle scan by rigid-body transformations with MCFLIRT to correct for subject motion. Next, non-brain matter from functional images was removed using BET (Smith, 2002). Resulting images were high-pass filtered with Gaussian-weighted least squares fitting with a 100 s cut-off. Finally, functional images were spatially smoothed using a Gaussian kernel of FWHM 8-mm. T1-weighted structural images were cleared from non-brain tissue with BET. These images were registered to the MNI152 standard space template with 2-mm resolution by first calculating transformation parameters from structural to standard space and from functional to structural space. Then, these transformations

![Fig. 1. Sample trials with matching (top) and mismatching (bottom) contextual cues. Narratives described complex action sequences such as fishing in general terms that were ambiguous without the contextual cues. Before each narrative, participants saw a contextually matching or mismatching visual cue.](Image 318x563 to 557x735)
were concatenated and used to co-register functional datasets to the standard space. Both registration steps were performed using FLIRT (Jenkinson, Bannister, Brady, & Smith, 2002).

2.4. Comprehension and recall measurements

After the fMRI experiment, the participants were asked to rate the comprehensibility of each narrative, and to recall as many idea units as possible (see Bransford and Johnson (1972)) of each narrative as possible. Even though it is possible to memorise verbal material without understanding it, comprehension provides a significant advantage to both encoding and subsequent recall (Craik & Tulving, 1975).

2.5. Analysis of regional effects

Regional responses to narratives preceded by mismatching and matching context conditions were examined with a two-class GLM model convolved with 6 of the 12 head motion parameters. The resulting signal time courses were included in the model as effects of no interest to account for motion-related variance. After the generation of individual contrast images for the matching versus mismatching context conditions and vice versa, a second level (random effects) analysis was applied to these contrast images in a new GLM. Statistical threshold was set at $p < 0.05$, FDR-corrected (Worsley, 2002).

In addition to using GLM analysis, we quantified the temporal similarity of brain activity across subjects during mismatching and matching cue conditions using inter-subject correlation (ISC) analysis. ISC focuses on temporal similarity rather than activation amplitude of the signals in single voxels. As was shown in previous studies (Hasson, Nir, Levy, Fuhrmann, & Malach, 2004; Jäskeläinen et al., 2008; Stephens, Silbert, & Hasson, 2010; Wilson, Molnar-Szakacs, & Iacoboni, 2008), ISC can reveal consistent patterns of brain activation within a group of subjects during naturalistic stimulation, for which explicit stimulus model cannot necessarily be applied. Pearson’s correlation coefficient was employed to derive between-subjects voxel-wise similarity measures at the group level using the toolbox developed by Kauppi, Jäskeläinen, Sams, and Tohka (2010). Data were preprocessed similarly as for GLM analyses. Motion parameters were regressed out from the data, and time series were shifted by three TRs to account for hemodynamic lag. ISC was calculated separately for each of the 20 narratives for the two counterbalancing, including all time points when the story was told (35–41 TRs), resulting in a total of 40 ISC group maps. ISC values across conditions were linearly regressed with the average behavioural comprehensibility ratings and thresholded with $p < 0.01$, cluster-corrected with cluster size of 27 voxels.

2.6. Functional connectivity analysis

Functional connectivity was estimated using psychophysiological interactions (PPPIs; Friston et al., 1997) implemented in SPM8 software (www.filion.ucl.ac.uk/spm/software/spm8). Broca’s area was selected as a seed ROI for the PPI analysis, given its profound role in speech comprehension (Bookheimer, 2002) and its spm/software/spm8/). Broca’s area was selected as a seed ROI for the PPI analyses, linearly regressed with the average behavioural comprehensibility ratings and preprocessed similarly as for GLM analyses. Motion parameters were regressed out from the data, and time series were shifted by three TRs to account for hemodynamic lag. ISC can reveal consistent patterns of brain activation within a group of subjects during naturalistic stimulation, for which explicit stimulus model cannot necessarily be applied. Pearson’s correlation coefficient was employed to derive between-subjects voxel-wise similarity measures at the group level using the toolbox developed by Kauppi, Jäskeläinen, Sams, and Tohka (2010). Data were preprocessed similarly as for GLM analyses. Motion parameters were regressed out from the data, and time series were shifted by three TRs to account for hemodynamic lag. ISC was calculated separately for each of the 20 narratives for the two counterbalancing, including all time points when the story was told (35–41 TRs), resulting in a total of 40 ISC group maps. ISC values across conditions were linearly regressed with the average behavioural comprehensibility ratings and thresholded with $p < 0.01$, cluster-corrected with cluster size of 27 voxels.

2.7. Eye movement recordings

As pupil dilation is associated with processing and memory load (Beatty, 1982), we recorded participants’ pupil size during the narrative listening phase of the experiment. Data were recorded successfully from 17 participants with a SMI 60Hz Eye Tracking long-range eye tracking system (SensoMotoric Instruments GmbH, Germany). To get an absolute scale for pupil diameter, we used a reference “fake pupil”, a piece of paper with a black circle of known diameter painted on it. This fake pupil was placed in front of participants’ eye at the beginning of the experiment, and a conversion factor was acquired. This conversion factor was subsequently used to convert the pupil diameter from pixels to millimetres. Only horizontal diameter was used for the analysis, as it is not confounded by subject’s position or eyelid movement. Pupil size measures were cleared from blink artefacts and horizontal pupil diameters were averaged for each trial for every subject. The resulting pupil size measurements were compared between matching and mismatching context conditions.

3. Results

3.1. Behavioural results

Comprehension of narratives preceded by matching context picture was significantly better than comprehension of narratives preceded by mismatching picture. $t(18)=9.47$, $p < 0.0001$, $d = 2.74$, means: 2.89 (72%) and 1.66 (42%) for matching and mismatching context conditions, respectively (Fig. 2). More idea units were also recalled for the matching context condition, $t(18)=10.21$, $p < 0.0001$, $d = 2.94$, means: 1.78 (39%) and 0.19 (4%) for matching and mismatching context conditions, respectively. Pupil diameter was not significantly influenced by context, $F(16)=0.94$, $p = 0.89$ (means: 4.70 and 4.72; std: 0.75 and 0.78 for matching and mismatching context conditions, respectively).

3.2. Functional MRI

GLM analysis revealed significantly stronger ($p < 0.05$, FDR corrected) task-evoked BOLD responses when listening to narratives preceded by contextually mismatching compared to matching pictures in the cluster that covered both left BA 44 and BA 45, as well as a portion of the left dorsal and ventral premotor cortex (see Fig. 3A and B, in blue, and Table 1). No significant differences were observed in this analysis or in the opposite contrast (matching versus mismatching).

PPI analysis revealed that in matching versus mismatching contrast connectivity increased between Broca’s area and bilateral AG/SMG (extending to right PCC), anterior and posterior cingulate cortices, and an area in the left anterior superior frontal gyrus (Fig. 3A and B, in red, and Table 1). There were no brain regions showing increased connectivity with Broca’s area in mismatching versus matching contrast.

Next we studied the functional connectivity of the anterior and posterior parts of Broca’s area separately in PPI analysis. Although
the connectivity of both subregions with right AG/SMG increased while listening to matching versus mismatching narratives, the analysis revealed a clear dissociation in the regional connectivity profiles (Fig. 4 and Table 1). Left BA 44 showed additional increased connectivity with left AG, left inferior/middle temporal gyrus and left postcentral gyrus, whereas left BA 45 showed increased connectivity with right PCC, right anterior IFG, lateral occipital cortex and ACC.

Regressing ISC against comprehension ratings revealed stronger ISC for lower comprehension ratings in anterior part of Broca’s area, left lobule VIIa of the cerebellum and left middle temporal gyrus (Fig. 5). This cluster in Broca’s area overlaps with the corresponding cluster identified in the GLM analysis described above, i.e. 69% of the voxels showing significant ISC showed statistically significant activity in the GLM analysis. For higher comprehension ratings ISC was stronger in right cerebellum crus II and V1/V2 (Table 2).

4. Discussion

In the present study we demonstrate that a fronto-parietal functional network supports context-dependent narrative comprehension while participants listen to spoken narratives preceded by matching versus mismatching contextual cues. Matching contextual cues improved the comprehension and subsequent recall
of the narratives (c.f. Bransford & Johnson, 1972; Dooling & Lachmann, 1971), and increased functional connectivity within a fronto-parietal functional network comprising Broca’s area, bilateral AG/SMG, as well as anterior and posterior cingulate and left superior frontal cortices. Subjects’ pupil sizes did not differ between matching and mismatching conditions, suggesting that cognitive load in the two conditions was comparable (Beatty, 1982; Kahneman & Beatty, 1966; Stanners, Coulter, Sweet, & Murphy, 1979) and that the observed differences across conditions likely cannot be explained by differences in allocation of attention or level of vigilance. Increased narrative comprehension and recall were paralleled by enhanced functional connectivity between Broca’s area and bilateral inferior parietal cortex, anterior and posterior cingulate cortices, and left anterior superior frontal gyrus. Importantly, in spite of its enhanced connectivity in the matching condition, hemodynamic activity in Broca’s area was stronger in conflicting condition.

4.1. A fronto-parietal network supporting context-dependent narrative comprehension

Functional connectivity analysis revealed that Broca’s area interacts with the linguistic processing nodes in bilateral AG and extra-linguistic ACC and PCC areas during context-dependent speech comprehension. Previously IPC regions have been associated with both local and global semantic functions, e.g. accessing word meanings and higher level semantic integration (Binder et al., 2009; Harpaz, Levkovitz, & Lavidor, 2009; Humphries et al., 2007; Price, 2000) and left AG has been shown to exhibit enhanced activity when subjects listen to titled compared to untitled narratives, possibly reflecting its involvement in processing contextual information (Martin-Loeches et al., 2008). Furthermore functional connectivity between the components of the language network increases as a function of speech intelligibility (Abrams et al., 2013). Our findings are in line with these

Table 1
MNI coordinates of peak activations obtained in the GLM and PPI analyses. The data are thresholded at p < 0.05 (FDR corrected at cluster level).

<table>
<thead>
<tr>
<th>Region</th>
<th>Hemisphere</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>Z</th>
<th>Voxels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Broca’s area (BA 44/45)</td>
<td>L</td>
<td>−43</td>
<td>18</td>
<td>18</td>
<td>5.24</td>
<td>1251</td>
</tr>
<tr>
<td>Inferior parietal cortex, posterior cingulate cortex</td>
<td>R</td>
<td>24</td>
<td>−46</td>
<td>32</td>
<td>4.66</td>
<td>4864</td>
</tr>
<tr>
<td>Lateral occipital cortex</td>
<td>R</td>
<td>44</td>
<td>−68</td>
<td>−22</td>
<td>4.31</td>
<td>280</td>
</tr>
<tr>
<td>Anterior cingulate/Paracingulate cortex</td>
<td>L</td>
<td>−12</td>
<td>22</td>
<td>28</td>
<td>4.03</td>
<td>313</td>
</tr>
<tr>
<td>Angular gyrus</td>
<td>L</td>
<td>−36</td>
<td>−68</td>
<td>46</td>
<td>3.92</td>
<td>732</td>
</tr>
<tr>
<td>Anterior superior frontal gyrus</td>
<td>L</td>
<td>−18</td>
<td>44</td>
<td>32</td>
<td>3.91</td>
<td>455</td>
</tr>
<tr>
<td>Middle frontal gyrus</td>
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<td>28</td>
<td>20</td>
<td>34</td>
<td>3.66</td>
<td>383</td>
</tr>
<tr>
<td>Postcentral gyrus</td>
<td>L</td>
<td>−42</td>
<td>−26</td>
<td>36</td>
<td>4.53</td>
<td>498</td>
</tr>
<tr>
<td>Angular gyrus</td>
<td>L</td>
<td>−56</td>
<td>−54</td>
<td>42</td>
<td>4.17</td>
<td>538</td>
</tr>
<tr>
<td>Inferior/Middle temporal gyrus</td>
<td>L</td>
<td>−50</td>
<td>−18</td>
<td>−26</td>
<td>4.15</td>
<td>203</td>
</tr>
<tr>
<td>Supramarginal gyrus/Angular gyrus</td>
<td>R</td>
<td>48</td>
<td>−42</td>
<td>34</td>
<td>3.91</td>
<td>396</td>
</tr>
<tr>
<td>Inferior parietal cortex</td>
<td>R</td>
<td>38</td>
<td>−50</td>
<td>28</td>
<td>6.52</td>
<td>2091</td>
</tr>
<tr>
<td>Anterior cingulate/Paracingulate cortex</td>
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<td>4</td>
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<td>768</td>
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<td>Inferior frontal gyrus</td>
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<td>30</td>
<td>4.82</td>
<td>364</td>
</tr>
</tbody>
</table>

Fig. 4. (A) and (B) Brain regions showing significant functional connectivity with posterior (hot colours) and anterior (cold colours) Broca’s area (seeds indicated by orange and blue circles respectively) during matching versus mismatching trials. Overlap is indicated in purple. Connectivity from both seeds increases only in the right inferior parietal cortex (AG/SMG, indicated by purple). Data are thresholded at Z > 2.3, and FDR corrected (p < 0.05) at the cluster level.
observations, showing that functional connectivity between frontal and parietal nodes of the language network is enhanced when contextual cues are available for supporting language comprehension. We addressed here specifically the global aspect of contextual processing; therefore we can only speculate about processes that took place at the local semantic level. We propose that when a contextual cue is presented, a hierarchical representation of the context is activated. Depending on its relevance, it either facilitates or impairs processing of the following narrative. The observed functional connectivity between Broca’s area and AG is likely related to both local and global stages of semantic processing. At the local level, the meaning of incoming word is accessed and the specific meaning is further selected based on the contextual information. At the global level, the linguistic information is integrated into a coherent narrative facilitated by the matching context.

Given that previous studies have shown that posterior cingulate cortex contributes to episodic memory retrieval (for review see Cabeza and Nyberg (2000)), the enhanced coupling between PCC and Broca’s area during matching context conditions might reflect access to contextual knowledge in long-term memory (Binder et al., 2009). Posterior midline regions, including precuneus, medial frontal cortex and paracingulate cortex, were also shown to reliably respond only to linguistic information, which was coherent over long time scales (~30 s; Lerner, Honey, Silbert, & Hasson, 2011). Moreover, PCC was implicated in a coherence building process (Ferstl, Neumann, Bogler, & von Cramon, 2008), which may also take place during contextual understanding, when more information becomes available as the narrative develops. The relatively low recall scores may have been caused by the behavioural task taking place after the scanning session at which point details of the stories may have been forgotten; ACC, in turn, supports various executive control functions (Abutalebi et al., 2012; Paus, 2001) such as error monitoring (Botvinick et al., 2004; Carter, Braver, Barch, Botvinick, Noll & Cohen, 1998). Thus, enhanced coupling between Broca’s area and ACC could support tracking of the consistency between the incoming narrative and the active schema provided by the context-matching picture.

We observed an increase of connectivity between left BA 45 and ACC in the matching condition. This finding might seem counterintuitive, because one would expect to see greater error monitoring activity in mismatching condition. To explain this finding, we tentatively suggest that in the matching condition consistency of the on-going narrative needs to be constantly monitored to build a coherent representation of its semantics, and this is reflected in stronger functional connectivity between left BA 45 and ACC. In mismatching case, once the narrative has been judged as inconsistent, there is no longer any need for consistency monitoring. Because of the short duration of this conflict, it probably was not reflected in the connectivity over the whole story time course. In sum, the observed functional connectivity pattern suggests that the prefrontal regions (Ye, Doñamayor, & Münte, 2012), Connectivity of the anterior Broca’s area with PCC and dACC suggests that this brain region is specifically associated with semantic processing (Hagoort, 2005) and monitors the consistency of the narrative via its connections with the dACC. However, activations in Broca’s area (mainly BA 45, extending into 44 and 47) and left inferior parietal areas (including AG and SMG) have also been reported while processing multiple syntactic

4.2. Functional connectivity of the anterior and posterior portions of Broca’s area

Functional connectivity analysis of left posterior (BA 44) and anterior (BA 45) subregions of Broca’s area revealed that these areas have differential functional connectivity profiles (see Fig. 4). While both subregions showed enhanced connectivity with right AG/SMG during matching condition – possibly reflecting interactions between sensory representation of the words and their meanings – the posterior part was also functionally coupled with left AG. This connection could reflect the initial linguistic processing, which requires both phonological and semantic components. It has been proposed that left IPC is involved in retrieval of the details of the stories may have been forgotten; ACC, in turn, supports various executive control functions (Abutalebi et al., 2012; Paus, 2001) such as error monitoring (Botvinick et al., 2004; Carter, Braver, Barch, Botvinick, Noll & Cohen, 1998). Thus, enhanced coupling between Broca’s area and ACC could support tracking of the consistency between the incoming narrative and the active schema provided by the context-matching picture.

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**Table 2**

<table>
<thead>
<tr>
<th>Region</th>
<th>Hemisphere</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>R</th>
<th>Voxels</th>
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<tr>
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<td>mismatching versus matching</td>
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<tr>
<td>V1/V2</td>
<td>LR</td>
<td>12</td>
<td>86</td>
<td>18</td>
<td>0.55</td>
<td>736</td>
</tr>
<tr>
<td>Middle temporal gyrus/SMG</td>
<td>L</td>
<td>48</td>
<td>32</td>
<td>8</td>
<td>0.58</td>
<td>80</td>
</tr>
<tr>
<td>Middle temporal gyrus</td>
<td>LR</td>
<td>48</td>
<td>32</td>
<td>8</td>
<td>0.58</td>
<td>80</td>
</tr>
<tr>
<td>Broca’s area</td>
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<td>40</td>
<td>22</td>
<td>6</td>
<td>0.58</td>
<td>80</td>
</tr>
<tr>
<td>Crus II</td>
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<td>-32</td>
<td>-8</td>
<td>-0.59</td>
<td>109</td>
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<tr>
<td>Middle temporal gyrus</td>
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<td>78</td>
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<tr>
<td>Superior temporal gyrus</td>
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<td>66</td>
<td>52</td>
<td>6</td>
<td>-0.58</td>
<td>109</td>
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<td>V1/V2</td>
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Fig. 5. Brain regions where ISC was modulated by contextual cues. Hot colours denote regions where ISC was significantly higher for contextually mismatching trials. The only area where ISC was higher in contextually matching trials was Crus II of cerebellum (in blue). Data are thresholded at cluster significance threshold of $p < 0.01$. 

**Table 2**

MNI coordinates of peak activations obtained in ISC analysis. The data are thresholded at $p < 0.01$, cluster-corrected.
representations in ambiguous phrases (Tyler et al., 2011). Results of Humphries et al. (2007) imply that left AG is also sensitive to a combinatorial semantic structure, arising from possibility to align individual words to form a more complex meaning. These authors suggested that AG integrates semantic and syntactic information to produce the overall sentence meaning. Since activation in AG was found for cases not involving syntactic processing, they imply that AG is involved primarily in semantic integration.

We also observed increased connectivity of BA 45 with right anterior IFG and AG. These regions are activated during figurative speech processing (Bookheimer, 2002), e.g. bilateral IFG and middle temporal gyrus during processing idioms (Zempleni et al., 2007a, 2007b). It is thus possible that contextual processing involves similar global linguistic functions as comprehending less transparent figurative language.

4.3. Dissociation of response amplitudes and functional connectivity of Broca’s area

The effects of contextual information on the response amplitudes and functional connectivity were dissociated in Broca’s area: while the matching context markedly increased connectivity of Broca’s area with the bilateral inferior parietal cortex, cingulate gyrus and left anterior superior frontal gyrus, we failed to see enhanced hemodynamic activity in these areas in the corresponding GLM analysis. Instead, when semantic coherence of the narrative was broken due to mismatching rather than matching context, we failed to see connectivity between the areas, suggested by the correlated inter-area activity, only takes place when proper contextual information is available.

ISC was negatively rather than positively associated with comprehension ratings in Broca’s region (BA 44 and BA 45). This finding seems contradictory, as one would expect the matching context to result in higher degree of inter-subject similarity of stimulus processing. However, we suggest that the mismatching context led participants into a shared state of increased semantic selection and integration demands, which is reflected in higher synchrony in Broca’s area. The only regions where we observed higher ISC for higher comprehension ratings were V1/V2, possibly reflecting similarity of visually processed information, since only fixation cross was presented on the screen, and right cerebellum’s crus II that might reflect supporting functions of cerebellum during linguistic processing (Stoodley & Schmahmann, 2009).

Cerebellar activity was shown to scale with complexity of language task (Xiang et al., 2003), and greater activation during semantic processing, compared to phonological processing (McDermott, Petersen, Watson, & Ojemann, 2003), as well as involvement in semantic decision process (Noppeney & Price, 2002).

The cluster observed in our GLM analyses encompassed both BA 44 and BA 45. In the literature, BA 44 is typically reported to participate in phonological processing, and we can only speculate about its involvement in context-dependent processing. It may be possible that while listening to contextually matching narrative, the context provides a more accurate predictive model of the upcoming phonetics of the speech, thus facilitating phonological processing. Another subregion of Broca’s area that is likely involved in semantic processing is BA 47, activation of which we would have observed with a slightly more lenient threshold.

4.4. Context-dependent language processing in the brain

Our findings significantly extend the results of earlier studies addressing contextual effects and coherence of text and context (Maguire et al., 1999; Martin-Loeches et al., 2008; St George et al., 1999). St George et al. (1999) contrasted reading titled and untitled paragraphs word by word and found right middle temporal regions being particularly important for integrative processes during discourse processing. In PET study of Maguire et al. (1999) the subjects were presented with auditory passages that were either readily comprehensible or unusual stories that required prior knowledge to comprehend. Prior knowledge was manipulated by presenting relevant, irrelevant or no visual cue before the story. The authors found involvement of anterior and ventral parts of the medial parietal and posterior cingulate cortices when unusual story was given with relevant prior knowledge.

Finally, a more recent study used fMRI to address effects of story coherence on brain activation, employing textual long narratives with or without a title that made paragraphs globally coherent (Martin-Loeches et al., 2008). This study found left AG activation while processing titled paragraphs and linked it to global coherence processing. While our results are in line with this interpretation, some other studies did not find the same regions involved in contextual understanding process. Methodological differences are a possible source of discrepancy, e.g., selection of a priori regions of interest (St George et al., 1999) or use of PET (Maguire et al., 1999) that has lower temporal resolution than fMRI, or the fact that two of these earlier studies presented discourses as text and context as a title (Martin-Loeches et al., 2008; St George et al., 1999), in contrast to our study that involved spoken narratives and pictorial context. Indeed, several studies have compared neural mechanisms of written and spoken language comprehension and have shown activation of different cortical areas depending on modality of stimulation (Buchweitz, Mason, Tomitch, & Just, 2009; Constable et al., 2004; Michael, Keller, Carpenter, & Just, 2001).

Listening to versus reading sentences results in activation of more anterior sites of Broca’s area, possibly reflecting more intensive semantic processing (Hagoort, 2005) during listening comprehension: it was specifically suggested that spoken language provides listener with prosodic cues, while written language offers punctuation as a means of information parsing (Michael et al., 2001).

Finally, it has been suggested that language comprehension can be treated as language-guided mental simulation of the described situation (Martin-Loeches et al., 2008; Zwaan & Rapp, 2006). From this point of view, different situational models could be created as a function of narrative content, and consequently, result in different patterns of brain activity. Even the same content can lead to different situational models being constructed due to level of global coherence achieved by reader and listener and lead to variable brain activation patterns (Martin-Loeches et al., 2008). Given the profound role of Broca’s area in particularly spoken language processing, our study, manipulating the contextual information of spoken instead of textual narratives, is possibly more sensitive for detecting contextual effects in Broca’s area.

Narative comprehension requires active prediction, selection, and integration of task-relevant information (Desimone & Duncan, 1995; Gibson, 1979). If the context is unknown, several alternative explanations of the current discourse segment can be prepared in parallel when listening to the unfolding story (Cisek & Kalaska, 2010; Wlotko & Federmeier, 2012). Given that Broca’s area is engaged by tasks where response is selected among several
linguistic regions such as ACC or PCC that contribute to specific aspects of semantic processing such as error monitoring or episodic memory retrieval respectively.

5. Conclusions

We conclude that Broca’s area and its connections with the inferior parietal and cingulate cortices play a critical role in context-dependent narrative comprehension. Listening to narrative disambiguates with appropriate context facilitates access to and selection of relevant semantic information and further integration of words and sentences into a coherent narrative. We propose that at the neural level, such contextual understanding is supported by enhanced connectivity of posterior and anterior portions of Broca’s area with fronto-parietal network of brain regions. Speculatively, we suggest that within this network the semantic processing involves AG, which participates in access to semantic representation of incoming information. This semantic representation is further selected and unified into larger units in Broca’s area on the basis of available context. In the next stage these larger units are matched and updated to a new representation in AG. The whole process might be monitored by dACC for consistency of narrative in relation to context.

Acknowledgements

We thank Marita Kattelus for her help with the data acquisition. The calculations presented above were performed using computer resources within the Aalto University School of Science “Science-IT” project.

Appendix A. Sample narrative translated to English.

If the environment were polluted, the whole business would be useless. If our spot is very popular, the results may not be as good as expected. Of course, it is nice to have companions but others may also compete for the same limited shares. Accidents are also possible, so one should be prepared to provide first aid for the common injuries. Of course, one can use a phone to call for help, but as we are far out from the hospital the cell phone might not work or it may run out of battery. Operating the equipment takes a good deal of practice and skill in the labile environment. Performing the actions in the right sequence is critical; otherwise one simply ends up tangling everything. Even when everything is done right, a great deal of patience and attention are required to achieve the best results, even though on many occasions one must return to home without having any success at all.

Appropriate contextual cue = Doing laundry. Inappropriate contextual cue = Fishing.

References


Author contributions

DS: Designed the experiments, acquired and analysed fMRI and behavioural data, wrote the manuscript
EG: Analysed fMRI data.
JL: Analysed fMRI data.
JS: Designed the experiment, wrote the manuscript.

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