Lateralized task shift effects in Broca's and Wernicke's regions and in visual word form area are selective for conceptual content and reflect trial history

Mikkel Wallentin a,b,⁎, Jákup Ludvík Dahl Michaelsen b, Ian Rynne b, Rasmus Høll Nielsen b

a Center of Functionally Integrative Neuroscience, Aarhus University Hospital, Building 10-G-5, Nørrebrogade, 8000 Aarhus C, Denmark
b Center for Semiotics, Aarhus University, Denmark

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A B S T R A C T

We investigated whether lateralized BOLD-fMRI activations in Broca's region, Wernicke's region and visual word form area (VWFA) reflect task shift costs and to which extent these effects are specific to language related task shifts. We employed a linguistic one-back memory paradigm where participants (n = 58) on each trial responded to whether a given word was the same as the previous word. In concordance with previous findings we found that conceptual shifts (CS), i.e. new words, elicited a strongly left-lateralized response in all three regions compared to repeat words. Words were sometimes presented through the visual modality (read) and sometimes through the auditory modality (spoken). This enabled the study of perceptual modality shifts (PS) relative to trials that stayed in the same modality as the previous trials. Again, we found a strongly left-lateralized effect in all regions. This was independent of whether the word was a CS or not, suggesting that linguistic translation across modalities taxes the same system as CS. Response shifts (RS), on the other hand, when shifting from one response (e.g. reporting a repeat word) to another (e.g. reporting a new word) did not yield an observable left lateralized response in any of the regions, suggesting that the lateralized task shift cost effects in these regions are not shared by all types of task shifts. Lateralization for individual tasks was found to be correlated across brain regions, but not across tasks, suggesting that lateralization may not be a unitary phenomenon, but vary across participants according to task demands. Both response time and lateralization were found to reflect the demands not only of the current trial but also of the previous trial, illustrating the context dependency of even simple cognitive tasks.

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Introduction

Broca's region

Broca's region (Amunts and Zilles, 2012; Broca, 1861), defined as Brodmann areas 44 and 45 of the left inferior frontal gyrus (LIFG) is known to play a key role in speech production as well as in the processing of language input (e.g. see Kristensen and Wallentin, in press for a review). The exact function (or functions) of Broca's region, however, remains disputed (Grodzinsky and Santi, 2008; Novick et al., 2010). Recently, it has been suggested that activation in this area reflects general linguistic processing costs (Kristensen and Wallentin, in press). In support of this are findings that syntactic manipulations yield increased Broca's region activation (Ben-Shachar et al., 2004; Christensen and Wallentin, 2011; Kristensen et al., 2013; Tettamanti et al., 2009) and findings showing that Broca's region responds to unpredicted word order, rather than syntactic manipulations per se, as defined by working memory demands (Fiebach et al., 2005), cloze probability (Obleser and Kotz, 2010) or context (Kristensen et al., in press).

Broca's region activation can also be observed in the absence of word order manipulations. Single word studies have demonstrated that Broca's region activity is correlated with working memory demand (Cohen et al., 1997), and displays an inverse correlation with word frequency (Fiebach et al., 2002), word generation practice (Raichle et al., 1994) and priming (Thompson-Schill et al., 1999). Pseudowords, on the other hand, generate greater activation than real words (Fiebach et al., 2002), again suggesting that unexpected words or words that are difficult to understand yield greater activation. Along the same lines, Broca's region has been found to increase activation to degraded visual word features (visual word form) as well as to decreased plausibility based on world knowledge (van de Meerendonk et al., 2013). More generally, it has been found that Broca's region activation is positively correlated with response time during linguistic processing (Christensen and Wallentin, 2011; Kristensen and Wallentin, in press; Orfanidou et al., 2006; Wallentin et al., 2006).

⁎ Corresponding author at: Center of Functionally Integrative Neuroscience, Aarhus University Hospital, Building 10-G-5, Nørrebrogade, 8000 Aarhus C, Denmark.
E-mail address: mikkel@cfins.dk (M. Wallentin).

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The question remains, however, at what level of abstraction this effect occurs (Fedorenko et al., 2013). Is this a general linguistic phenomenon as the diverse findings seem to suggest, and if so, is it limited to language or can it be observed with tasks that are non-linguistic as well? Recent proposals suggest that Broca's region serves a more domain general purpose during cognitive control (Fedorenko et al., 2013; Novick et al., 2010).

**Wernicke's region**

Wernicke's region, usually defined as the posterior part of the superior temporal cortex behind the primary auditory cortex (Geschwind, 1970; Perry and Lupyan, 2014; Price, 2000), is traditionally thought to be involved in the recognition of patterns of spoken language (Geschwind, 1970; Wernicke, 1874). However, discussions similar to those for Broca's region have been led on the exact function, location and even existence of Wernicke's region (e.g. Binder et al., 1996; Bogen and Bogen, 1976; Dronkers et al., 2004; Turken and Dronkers, 2011; Wise et al., 2001). Recent studies have suggested that Wernicke's region plays a role in the mapping between visual and linguistic representations of categories (Perry and Lupyan, 2014) and indeed perhaps the most compelling argument in favor of the posterior temporal cortex being involved in linguistic processing is the location right between auditory and visual cortices (Geschwind, 1970), making it intuitively appealing to believe that it serves a function as an associative region for input that link the two perceptual modalities.

**Visual word form area**

In the posterior part of the inferior temporal gyrus lies a region that has repeatedly been found to be associated with reading words, so much that it has been named the visual word form area — VWFA (Dehaene and Cohen, 2011; Dehaene et al., 2002). VWFA has been found to be functionally specific to words relative to objects (Szewd et al., 2011) and relative to mirror images of words (Dehaene et al., 2010a). It has also been found to be sensitive to bigram frequency, i.e. the statistics of letter pairs in a particular language (Vinckier et al., 2007). However, controversies also exist about this alleged specificity (Price and Devlin, 2003) with some also finding context dependency of the VWFA's response pattern (Starrfelt and Gerlach, 2007). Additionally, VWFA has been found to be involved in the processing of both spoken words and pseudo-words (Orfanidou et al., 2006). Although more ventrally placed than Wernicke's region, the VWFA may thus also be a region relevant for the interface between visual and auditory aspects of word forms.

**Lateralization of Broca's and Wernicke's regions and VWFA**

The question about the specificity of Broca's region, Wernicke's region and VWFA activity corresponds to a similar discussion about lateralization in cognitive tasks and language in particular. Language function has been found to be strongly left lateralized when studied through deficits elicited either by brain injury (Dronkers et al., 2004; Pedersen et al., 1995; Pedersen et al., 2004) or by experimentally induced temporary deficits (Rasmussen and Milner, 1977; Woermann et al., 2003). Wernicke's region has long been known to be structurally asymmetric (Gannon et al., 1998; Geschwind and Levitsky, 1968). Activation in the VWFA has, similar to Broca's region, been found to be left-lateralized (Szewd et al., 2011). But the functional effects observed for all regions using functional imaging have often been much less lateralized than the patient studies would predict (e.g. Binder et al., 2000; Brown et al., 2006; Cardillo et al., 2012; Ferstl et al., 2008; Petersen et al., 2013; Vigneau et al., 2006, 2011). One reason for this discrepancy between patient and neuroimaging findings might be that lateralization indices from fMRI studies often are not based on close contrasts between linguistic inputs that differ in terms of processing cost. Indeed, when contrasting linguistic tasks to baseline, a bilateral pattern is often observed (Ferstl et al., 2008). Another reason might be that fMRI based lateralization indices are most often derived from a count of activated voxels in each hemisphere (Bethmann et al., 2007; Binder et al., 2000; Cai et al., 2010; Clements et al., 2006; Deppe et al., 2000; Lehéricy et al., 2000; Pujol et al., 1999; Springer et al., 1999; Szaflarski et al., 2002; Szaflarski et al., 2012; Woermann et al., 2003), a method that is very sensitive to the chosen significance threshold (Jones et al., 2011; Seghier, 2008; Suarez et al., 2009). But the divergent findings also point towards the possibility that lateralization may not be uniform across tasks and brain regions. One example is music. Lateralization for music has been found to depend on musical competence, both when studied using fMRI and MEG (Vuust et al., 2005, 2006). Similarly, it has been found that activation and lateralization in the VWFA change as a function of increased literacy (Dehaene et al., 2010b). But literacy is independent of musical competences and the lateralization changes relevant for the two task types therefore must be hypothesized to happen independently. This can only be the case if there is more than one type of lateralization. This calls for an investigation of whether lateralization is linked to individual brain regions (region specific) or whether lateralization is rather linked to the type of task at hand (task specific) or both.

**Hypotheses**

In the current experiment we investigated to which extent Broca's region (BA 44 & 45), Wernicke's region (posterior part of the superior temporal gyrus) and VWFA activity are specific to different types of processing costs elicited by more or less language related task shifts. To this end, we employed a single word one-back memory paradigm where participants responded to whether a given word was the same as the previous word or not (i.e. a conceptual shift), regardless of the perceptual features alone. But since trials in the current one-back paradigm shifted between visual and auditory formats, this was not always possible. A perceptual shift (PS: new modulation vs same modulation) from one modality to another would therefore increase the load on the linguistic system, because it would force the participant to re-encode the words at a higher level than the mere perceptual surface, i.e. by translating the perceptual features into the other perceptual modality or by remembering conceptual content. We hypothesized that this process would also result in an increased lateralized Wernicke's region and/or VWFA activity region (Dehaene and Cohen, 2011; Dehaene et al., 2002). The randomized string of stimuli made it possible for a number of new words or repeat words to appear in succession. These trains of repeated trial types would thus allow the participants to respond with the same finger on a number of trials, but whenever a trial went from being a new word to being a repeat word or vice versa this required the participants to perform a response shift (RS: new response vs same response). We hypothesized that these shifts, similar to the two other types of shifts, would elicit prolonged response times, but that this type of added processing cost would not be linguistic in nature and therefore would not show up as strongly left-lateralized responses in Broca's region, Wernicke's region or VWFA. For quantification of lateralization we employed an analysis based on averaging beta estimates across the entire regions of interest. Lastly, we wanted to investigate the possible relationship between lateralization, task and region. Is lateralization task- and/or region-specific? If lateralization is task specific then we would expect lateralization indices across the different regions to be correlated within a given task, whereas if it is region...
specific then we would expect lateralization within a region to be correlated across tasks. Lastly, if lateralization is unitary, we should expect it to be correlated across both tasks and regions.

**Material and methods**

**Participants**

Permission for the study was obtained from the local ethics committee (Region Midtjylland, Denmark) and written informed consent was obtained from each participant. Sixty participants with no known neurological history took part in the experiment (28/32 males/females; median age: 22 years, range: 19–38 years). All participants reported being right-handed and having Danish as their primary language. Each received a 150 DKK payment for participation. One participant was excluded due to a misunderstanding of the instructions and one due to a technical problem during fMRI data acquisition (n = 58).

**Stimuli and procedure**

Stimuli consisted of 250 concrete nouns, either displayed on a screen or played through headphones. The words were a subset of a larger collection of words (Sudre et al., 2012) that could be unambiguously translated into Danish. Word meaning spanned a wide area of the semantic space of concrete nouns, but the content is of lesser importance for the present purpose, since all contrasts are based on differences between the same words displayed in different contexts within and across participants. The experiment was a one-back word comparison task with 400 trials consisting of 250 new word trials and 150 repeat word trials. All new words were considered as concept shifts since the word was different from the previous. 200 trials consisted of written words and 200 trials heard. Perceptual shift (PS) trials occurred whenever a trial was presented in a different modality compared to the previous trial, regardless of whether it was a CS trial or not. Participants responded as fast as possible (using right index/middle finger) to whether the current word was the same (index finger response) as the previous or not (middle finger response), regardless of perceptual modality (see Fig. 1). Response shifts (RS) occurred whenever the required response was different from the previous trial, regardless of whether the shift went from new to repeat or from repeat to new and regardless of modality. Order of stimuli and perceptual modality of presentation were randomized within and across participants. All words were recorded in a neutral male voice (44,100 Hz sample rate, 16 bit, mono) using the microphone from a laptop computer and Audacity software (http://audacity.sourceforge.net/). The auditory stimuli had a mean duration of 638 ms (std: 172 ms; range: 255–1068 ms). Written words were displayed for 700 ms, and stimulus onset asynchrony was jittered using a skewed distribution (median 2975 ms; range: 2500–5000 ms).

Stimuli were presented and responses obtained using Cogent 2000 (http://www.vislab.ucl.ac.uk/cogent_2000.php), executed in MATLAB. Sounds were presented through a pair of electrodynamically driven headphones (MR confon, Magdeburg, Germany). These also helped to attenuate scanner noise. During post scan debriefing, all participants reported being able to hear the words through the scanner noise. Visual stimuli were projected onto a screen at the head end of the scanner bed and viewed through a mirror mounted on the head coil.

The whole experiment lasted approximately 20 min. Before entering the scanner, participants were given a trial run of the task with 10 words not used in the experiment in order to get accustomed to the study design and response procedure.

**Fig. 1.** Experimental setup. The experiment was a one-back memory task for words. Words were either displayed on a screen or played through headphones. Participants responded by buttonpress to whether the current word was the same as the previous word, regardless of perceptual modality. No response was given to the first trial (e.g. the word “hand” played through headphones). On the 2nd trial an index finger response was given if the word was a novel word (e.g. “cat”), regardless of which modality it was displayed in (e.g. on the screen). If the word was new, it was categorized as a concept shift (CS). If the perceptual modality was new, it was a perceptual shift (PS). From the third trial onwards the response might or might not be the same as on the previous trial (e.g. if the previous trial was a CS and the current was a nonCS). In this case the participant performed a response shift (RS). 250/400 trials were novel words (CS), 200/400 words were visual and auditory trials. Order was randomized, yielding a variable number of perceptual shifts (PS) and response shifts (RS) (50% on average).
**fMRI acquisition**

A 3 T Siemens Magnetom MRI Trio system (Erlangen, Germany) with a 12 channel matrix head coil was used to acquire both T2*-weighted gradient echo, echo-planar images (EPI) with Blood oxygenation level-dependent (BOLD) contrast and T1-weighted structural images. 610 EPI volumes were acquired per participant, not including the first 5 volumes that were discarded to allow for effects of T1 equilibrium. Whole brain coverage was achieved using 40 axial slices (interleaved acquisition) of 3 mm thickness with an in-plane resolution of 3 × 3 mm in a 64 × 64 voxel matrix (FOV 192 mm). Images were obtained with a TR of 1990 ms, a 27 ms TE and a 90° flip angle.

**Data analysis**

Pre-processing of fMRI data

All fMRI image pre-processing and data analyses were performed using Statistical Parametric Mapping software (SPM8; Wellcome Department of Imaging Neuroscience, www.fil.ion.ucl.ac.uk/spm), implemented in MATLAB. Functional images were motion corrected and registered to the first EPI image. The mean of the motion-corrected images was then co-registered to the individual’s structural MRI using a 9-parameter affine transformation. The anatomical image was spatially normalized to the standard MNI space using the unified segmentation–normalization algorithm within SPM (Ashburner and Friston, 2005) with the resulting deformation field subsequently applied to the fMRI data. Finally, data were spatially smoothed with an isotropic 8 mm full width at half maximum (FWHM) Gaussian kernel to account for differences between participants.

Statistical analyses of fMRI data

Statistical analyses of fMRI data were performed using a two-level general linear model approach (Penny and Holmes, 2007; Worsley and Friston, 1995). In order to control false positives, we thresholded results at p < 0.05, family-wise error corrected for multiple comparisons. Main effects were additionally thresholded to only display beta-estimates > 1 or < -1 with a cluster size of > 10 voxels (at the 2nd contrast level) in order to only report robust effects. In subsequent conjunction analyses we were also concerned with false negatives, so we therefore relaxed the statistical threshold to p < 0.001, uncorrected for multiple comparisons, and looked for voxels that would be significant at this level over more than one contrast. Lastly, we performed three regions of interest (ROI) analyses, by averaging beta-estimates across 1) Broca’s region and its right hemisphere homologue, 2) Wernicke’s region and its right hemisphere homologue and 3) VWFA and its right hemisphere homologue and conducting within-participant analyses of variance on the resulting data for each region. The Broca’s region ROI was defined using the Wake Forest University Pickatlas (Tzourio-Mazoyer et al., 2002). Wernicke’s region and its right hemisphere homologue were defined according to the tradition stemming from Geschwind (1970) as the posterior portions of the superior temporal gyri (but see Bogen and Bogen, 1976; Dronkers et al., 2004; Ojemann, 1991 for conflicting views). The superior temporal gyri were again found using Wake Forest University Pickatlas (Tzourio-Mazoyer et al., 2002) and the posterior portions were defined as Y ≤ -20 in MNI coordinates. The VWFA was defined as a sphere with a 10 mm radius around the MNI coordinates: [-43, -54, -12] and for the right hemisphere an additional sphere was created around the coordinates: [43, -54, -12]. This location has been reported to be the average peak coordinate across 25 word reading studies, with a standard deviation of 5 mm (McCandliss et al., 2003). See Fig. 6 for an illustration of the three regions.

**Results**

**Behavioral results**

Accuracy

Behavioral measures were Bonferroni-corrected for multiple comparisons. A 2 × 2 × 2 factorial repeated-measures ANOVA revealed a main effect of conceptual shift (CS – new vs repeat words, F(1,57) = 163.4, p(bonf) < 0.001), a main effect of perceptual modality shift (PS – new modality vs same modality, F(1,57) = 65.1, p(bonf) < 0.001), a main effect of response shift (RS – new response vs same response, F(1,57) = 20.8, p(bonf) < 0.001), several interaction effects: CS × PS (F(1,57) = 84.6, p(bonf) < 0.001), CS × RS (F(1,57) = 84.618, p(bonf) < 0.001) and PS × RS (F(1,57) = 20.3, p(bonf) < 0.001). No three-way interaction was observed for accuracy (F(1,57) = 1.5, p > 0.2). See Fig. 2.

Response time

Only correct responses were analyzed. A main effect of CS (F(1,57) = 83.02, p(bonf) < 0.001, 93.6 ms difference) was observed, a main effect of PS (F(1,57) = 167.47, p(bonf) < 0.001, 52.2 ms) and a main effect of RS (F(1,57) = 29.8, p(bonf) < 0.001, 22.9 ms), two two-way interaction effects: CS × RS (F(1,57) = 212.03, p(bonf) < 0.001, 62.8 ms) and PS × RS (F(1,57) = 77.5, p(bonf) < 0.001, 25.0 ms). We also observed a three-way interaction for response time (F(1,57) = 17.9, p(bonf) < 0.001, 11.9 ms). No CS × PS interaction was seen (F(1,57) = 0.91, p = 0.343, 3.1 ms). See Fig. 2.

**fMRI results**

Whole brain data

Main effects

Conceptual shift (CS). CS was accompanied by increased activation in a network of largely left-lateralized frontal and temporal brain regions including orbitofrontal, inferior frontal (BA 45), premotor, inferior temporal and middle temporal areas (see Fig. 3-top and Table 1). Negative

![Fig. 2. Behavioral effects. Response time effects of CS (93 ms), PS (52 ms) and RS (23 ms) as well as interactions were observed.](image-url)
effects were observed primarily in the parietal lobes, bilaterally, right lateralized in frontal regions including the right inferior frontal region (BA 45) and in the right inferior temporal gyrus (see Table 1).

Perceptual shift (PS). PS was accompanied by primarily left lateralized temporal regions, including inferior and middle temporal regions. But effects were also seen in the left inferior frontal part (see Fig. 3-middle and Table 2). No regions were found to display negative effects of PS.

Response shift (RS). RS was accompanied with activations primarily in motor and parietal regions. A number of midline effects were observed as well (see Fig. 3-bottom and Table 3). No regions were found to display negative effects of RS.

Interactions. A number of two-way interactions were observed at the whole brain level (Fig. 4, Table 4), including both highly significant positive and negative interactions between CS and RS. The negative effect was observed in motor and premotor regions as well as in anterior insula and in the posterior parietal lobe, including precuneus. The negative CS × RS interaction observed reflects regions of the brain that respond more if the current trial is a CS without RS and if the current trial is a nonCS but with RS (i.e., $-CS/RS + CS/\text{nonRS} + \text{nonCS}/RS - \text{nonCS}/\text{nonRS}$). In both cases this involves the presence of a CS in the previous trial. A similar negative interaction between PS and RS was observed. This reflects trials where the previous trial was a PS. Again, effects were observed in the precuneus among other places (Table 4). No 3-way interaction was observed.

Conjunctions. To investigate if Broca’s region, Wernicke’s region and VWFA are involved in shifts per se or if the processing is specific to conceptual processing we looked for overlapping responses for the three main effects. Here, in contrast to the main effects analysis, we were also concerned about false negatives. We therefore relaxed the statistical threshold to $p < 0.001$, uncorrected for multiple comparisons, and looked for voxels that would be significant at this level over more than one contrast (see Fig. 5).

CS&PS. Similar to the corrected threshold, both Broca’s region and VWFA activations were found for both CS and PS at this lower threshold, whereas they were not found for RS. A number of other regions, including Wernicke’s region were observed to be shared among CS and PS.

CS&RS. Overlaps between CS and RS were observed primarily in the premotor regions, bilaterally and in the supplementary motor region extending into the cingulate. Additional overlap was seen in the retrosplenial region and cerebellum.
Table 1
Effects of conceptual shift (CS).

<table>
<thead>
<tr>
<th>Putative anatomical region</th>
<th>Peak MNI</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Positive effect of conceptual shift [CS–nonCS]</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sup. orb. frontal R — BA 11</td>
<td>–8, 38, –24</td>
<td>5.57</td>
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<tr>
<td>Inf./mid. orb. frontal L — BA 47/11</td>
<td>–28, 34, –18</td>
<td>7.64</td>
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<tr>
<td>Inf. frontal tri. L — BA 45</td>
<td>–46, 18, 22</td>
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<tr>
<td>Supp. motor area L — BA 6</td>
<td>–6, 10, 54</td>
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<tr>
<td>Precentral L — BA 6</td>
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<tr>
<td>Precentral R — BA 6</td>
<td>60, 0, 46</td>
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<tr>
<td>Precentral R — BA 6</td>
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<td>6.37</td>
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<tr>
<td>Paracentral lobule L — BA 4/6</td>
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<td>Mid. temporal L — BA 22</td>
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<tr>
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<tr>
<td>Retrosplenial/calcarine L — BA 30</td>
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<td>Fusiform L — BA 20</td>
<td>–40, –36, –20</td>
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<tr>
<td>Inf. temporal L — BA 20</td>
<td>–50, –54, –18</td>
<td>5.64</td>
</tr>
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**Negative effect of conceptual shift [nonCS–CS]**

<table>
<thead>
<tr>
<th>Putative anatomical region</th>
<th>Peak MNI</th>
<th>Z-score</th>
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</thead>
<tbody>
<tr>
<td>Mid. frontal L — BA 11</td>
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<td>5.37</td>
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<tr>
<td>Supp. frontal R — BA 10</td>
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<td>Supp. frontal R — BA 8</td>
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<tr>
<td>Supp. medial frontal R — BA 9</td>
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<td>5.46</td>
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<td>Supp. medial frontal R — BA 8</td>
<td>8, 30, 54</td>
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<td>Inf. frontal tri. R — BA 45</td>
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<tr>
<td>Insula R</td>
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<td>5.51</td>
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<tr>
<td>Outside brain/inf. parietal L — BA 40</td>
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<td>Precuneus R — BA 7</td>
<td>10, –76, 50</td>
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</tbody>
</table>

P-FWE < 0.05, beta > 1 or < –1, cluster > 10 voxels, peak dist > 12 mm.

**PS&RS**

A large posterior overlap was seen for PS and RS extending over much of the parietal, occipital and posterior temporal lobes as well as cerebellum.

**CS&PS&RS**

Task general shift effects were observed in the premotor cortex, in the retrosplenial/calcarine region and in the cerebellum, all in the left hemisphere.

Table 2
Effects of perceptual shift (PS).

<table>
<thead>
<tr>
<th>Putative anatomical region</th>
<th>Peak MNI</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Positive effect of perceptual shift [PS–nonPS]</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Supp. med. frontal L — BA 8</td>
<td>–2, 54, 44</td>
<td>4.99</td>
</tr>
<tr>
<td>Precentral/inf. front. L — BA 6/45</td>
<td>–52, 12, 30</td>
<td>5.02</td>
</tr>
<tr>
<td>Postcentral L — BA 4/5</td>
<td>–40, –36, 60</td>
<td>4.92</td>
</tr>
<tr>
<td>Fusiform/inf. temporal L — BA 20</td>
<td>–42, –44, –28</td>
<td>5.78</td>
</tr>
<tr>
<td>Inf. temporal L — BA 37</td>
<td>–52, –58, –18</td>
<td>6.33</td>
</tr>
<tr>
<td>Supp. temporal sulcus L — BA 22/39</td>
<td>–44, –54, 20</td>
<td>5.72</td>
</tr>
<tr>
<td>Retrosplenial/calcarine R — BA 30</td>
<td>–45, 60, –10</td>
<td>5.06</td>
</tr>
<tr>
<td>Lingual L — BA 18</td>
<td>–20, 60, 0</td>
<td>4.97</td>
</tr>
<tr>
<td>Lingual R — BA 18</td>
<td>6, –74, –10</td>
<td>5.25</td>
</tr>
<tr>
<td>Precuneus — BA 7</td>
<td>–2, –60, 68</td>
<td>4.95</td>
</tr>
<tr>
<td>Cuneus — BA 19</td>
<td>2, –32, 40</td>
<td>5.18</td>
</tr>
<tr>
<td>Cerebellum R</td>
<td>28, –76, –22</td>
<td>5.07</td>
</tr>
</tbody>
</table>

P-FWE < 0.05, beta > 1, cluster > 10 voxels, peak dist > 12 mm.

ROI: Broca's region and right hemisphere homologue

To investigate lateralization effects and further limit potential false negatives, we investigated the effects of words against the non-modified baseline averaged across Broca's region (left inferior frontal gyrus (LIFG), i.e. BA 44 & 45) and its right hemisphere homologue (RIFG; see Fig. 6). When comparing average beta-estimates for all conditions in LIFG and RIFG with a paired t-test this revealed a weak non-significant lateralization trend (t(57) = 1.67, P = 0.1). 35/58 participants displayed a left-lateralized effect.

Next, we investigated the lateralization effects for CS, PS and RS. We conducted a $2 \times 2 \times 2 \times 2$ ANOVA (CS × PS × RS × hemisphere) on these data and, after correcting for multiple comparisons (using Bonferroni correction), we found no significant main effects, but a strong two-way interaction between CS and cerebral hemisphere ($F(1,57) = 130.1, p(bonf) < 0.001$), a strong interaction between PS and hemisphere ($F(1,57) = 42.5, p(bonf) < 0.001$), and an interaction between CS and RS ($F(1,57) = 12.854, p(bonf) < 0.02$). As can be seen from Fig. 6, the regional effects found in Broca's region for CS and RS are countered by negative effects in the right hemisphere homologue of Broca's region, explaining why we find no main effect of Cs or PS in the ANOVA and also explaining the strong interaction between hemisphere and the two shift effects. Crucially, no such interaction effect is observed for RS and hemisphere ($F(1,57) = 0.3, p > 0.5$), thus limiting the scope of the lateralized shift processing conducted in Broca's area. No other significant effects or interactions were observed in the ROI ANOVA. A lateralization index was computed within the ROI for each contrast separately by a simple left minus right hemisphere subtraction. The lateralization effects for CS and PS were very robust. For CS, 55/58 participants displayed lateralization indices above zero (i.e. left-lateralized); for PS, 46/58 participants were found to be left-lateralized, while for RS, 32/58 participants showed a left-lateralized effect. No effects of gender (Wallentin, 2009) were observed for any of the three lateralization indices in Broca's region ($F(1,57) < 1.3, p(uncorr.) > 0.25$ for all three tests).

ROI: Wernicke's region and right hemisphere homologue

We conducted another $2 \times 2 \times 2 \times 2$ ANOVA (CS × PS × RS × hemisphere) on the average beta estimates for the Wernicke's region ROI and its right hemisphere homologue. We found a significant main lateralization effect ($F(1,57) = 37.96, p(bonf) < 0.001$). No significant main effects of CS, PS or RS when correcting for multiple comparisons ($p(bonf) > 0.05$ for all). We found significant interaction between CS and hemisphere ($F(1,57) = 25.27, p(bonf) < 0.001$) and a significant interaction between PS and hemisphere ($F(1,57) = 18.42, p(bonf) < 0.001$). No interaction between RS and hemisphere was observed ($F(1,57) = 1.67, p(bonf) > 0.05$). We also observed a significant three-way interaction between CS, RS and hemisphere ($F(1,57) = 18.09, p(bonf) < 0.001$), showing that lateralization is greater for CS in the absence of RS and for NonCS if an RS is present (Fig. 7). No other interactions were observed ($p(bonf) > 0.05$ for all tests). For CS, 47/58 participants displayed a left-lateralized effect, for PS, 41/58 participants were found to be left-lateralized, while for RS, 35/58 participants showed a left-lateralized effect. No effects of gender (Wallentin, 2009).
were observed for any of the three lateralization indices in Wernicke’s region ($F(1,57) < 2$, $p(\text{uncorr.}) > 0.1$ for all three tests).

ROI: visual word form area and right hemisphere homologue

We again conducted a $2 \times 2 \times 2 \times 2$ ANOVA ($CS \times PS \times RS \times FL$) involving a new concept but not a response shift or if a trial is a repeat concept but a response shift (see Table 4). In effect this means that these regions display increased response if the previous trial was a conceptual shift. This may reflect history effects or be a down-stream (venous) response. The map is thresholded voxelwise at $p < 0.05$, FWE corrected.

**Table 4** Interaction effects.

<table>
<thead>
<tr>
<th>Putative anatomical region</th>
<th>Peak MNI</th>
<th>Z-score</th>
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</thead>
<tbody>
<tr>
<td><strong>2-way interactions</strong></td>
<td></td>
<td></td>
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<tr>
<td>Positive interaction $CS \times PS$</td>
<td></td>
<td></td>
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<tr>
<td>N.S.</td>
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<td></td>
</tr>
</tbody>
</table>

**Negative interaction $CS \times PS$**

$\left[ -CS / PS + CS / nonPS + nonCS / PS - nonCS / nonPS \right]$

Initial occipital L — BA 19

40, −68, −10

4.85

Mid. occipital R — BA 39

42, −76, 26

5.25

**Positive interaction $CS \times RS$**

$\left[ CS / RS - CS / nonRS - nonCS / RS + nonCS / nonRS \right]$

Frontal sup med L — BA 8

−10, 50, 50

4.78

Frontal sup med R — BA 8

12, 52, 48

5.31

Frontal inf orb L — BA 47

−32, 32, −18

5.19

Rectus/midline

0, 36, −28

5.14

**Negative interaction $CS \times RS$**

$\left[ -CS / RS + CS / nonRS + nonCS / RS - nonCS / nonRS \right]$

Frontal mid L — BA 46

−36, 46, 12

5.59

Frontal mid L — BA 46

−40, 28, 30

5.63

Frontal mid R — BA 46

40, 36, 22

5.31

Frontal sup L — BA 6

−14, −8, 76

5.28

Cingulum mid R — BA 24

6, 30, 32

5.00

Insula L — BA 13

−36, 14, −4

6.09

Insula L — BA 13

−28, 28, 10

5.25

Insula R — BA 13

32, 24, 4

5.38

Insula R — BA 13

40, 20, 0

4.9

Rolandic oper L — BA 13

−50, −2, 4

5.46

Postcentral R BA 2/3/4

−36, −18, 44

6.43

Postcentral R BA 2/3/4

54, −26, 50

6.08

Temporal inf L

−62, −36, −16

5.06

Temporal mid L — BA 21/22

−50, −52, 16

5.41

Parietal inf L — BA 40

−42, −42, 20

6.07

Parietal inf R — BA 40

40, −56, 48

5.63

Precuneus/midline — BA 7

0, −68, 46

6.21

Precuneus R — BA 7

16, −72, 40

5.54

Brain stem

−2, −16, −20

5.26

**Positive interaction $PS \times RS$**

N.S.

**Negative interaction $PS \times RS$**

$\left[ -PS / RS + PS / nonPS + nonPS / RS - nonPS / nonPS \right]$

Parietal inf L — BA 40

−30, −56, 46

5.02

Putamen R

28, 20, −2

4.50

P-FWE < 0.05, peak dist > 12 mm.

**Fig. 4.** Interaction between conceptual shift ($CS$) and response shift ($RS$). The red/yellow effects depict negative interactions, i.e. regions that have increased BOLD response if a trial involves a new concept but not a response shift or if a trial is a repeat concept but a response shift (see Table 4). In effect this means that these regions display increased response if the previous trial was a conceptual shift. This may reflect history effects or be a down-stream (venous) response. The map is thresholded voxelwise at $p < 0.05$, FWE corrected.

We found that conceptual shifts (CS) and perceptual shifts (PS) but not response shifts (RS) in relation to a linguistic one-back task resulted in a strongly left-lateralized BOLD response in Broca’s region (LIFG; BAs 44 & 45). Word stimuli as such (relative to baseline) did not yield a significant lateralized effect in Broca’s region (Fig. 6, top). Increased processing costs were observed in terms of prolonged response times for all types of task shifts, but only CS (peak coordinates: −46, 18, 22) and PS (peak coordinates: 52, 14, 30) yielded significant left-lateralized Broca’s region responses (see Tables 1 and 2 and Figs. 3, 5 and 6). This demonstrates that shift cost effects in this region are not task-general, but limited to certain processes, most likely those that are conceptual in nature. Both PS and nonPS conditions contain CS trials to an equal amount due to randomization of the stimuli within and
Fig. 5. Conjunctions. Red: conceptual shift (CS) & perceptual shift (PS). Green: CS & response shift (RS). Cyan: PS & RS. Yellow: CS & PS & RS. Each effect was thresholded at $p < 0.001$, uncorrected, to balance false positives and false negatives. Unspecified shift costs are primarily seen in premotor and cerebellar regions, suggesting a link to motor output. More widespread overlap is seen for conceptual shift and perceptual shift in frontal and temporal regions, including Broca’s area, Wernicke’s region and VWFA. No effect for RS is seen in either region, indicating that it is not task shifts as such, but more language related shifts that evoke responses here.

Fig. 6. Lateralization effects in regions of interest: yellow panel: Broca’s region (Brodmann areas 44 and 45); red panel: Wernicke’s region (posterior part of superior temporal gyrus) and green panel: visual word form area (10 mm radius spheres around MNI: $-43, -54, -12$) and the right hemisphere homologues of these regions. Average effects across each region in each hemisphere and the within subject difference between the two effects (lateralization index). Top row: all words against the unmodeled baseline. 2nd row: conceptual shift versus no shift. 3rd row: perceptual modality shift versus no shift. 4th row: response shift versus no shift.
across participants. The PS effect (PS–nonPS) therefore cannot be said to reflect CS. However, PS trials are more conceptual than nonPS trials in the sense that the one-back task within perceptual modality can be solved based solely on perceptual surface features whereas a word comparison across modalities requires a semantic representation or at least a translation/recoding of linguistic features (letters to phonemes or vice versa). One might argue that a beneficial task solving strategy would be to encode all trials at the conceptual level. In that case, however, one would not expect to see any difference between PS and nonPS. This is not supported by data. Both the fMRI data and the response time pattern clearly show that PS trials are more costly to process than nonPS trials, suggesting a difference in strategy for solving the two trial types with PS perhaps relying more on conceptual content than nonPS.

A response shift, on the other hand (going from a “repeat” response to a “new” response or the other way), does not on average have any added conceptual content compared to a non-shift (going from “repeat” to “repeat” or from “new” to “new”). Task general shift effects were observed more dorsally in premotor cortex (Fig. 4), consistent with a prolonged response time.

The lateralization for the CS contrast was not only visible in the left hemisphere. The CS–nonCS contrast yielded a negative effect in the right hemisphere homologue of Broca’s region both in the whole brain data and in the ROI analysis, suggesting that this effect does not originate in one hemisphere but reflects a genuine lateralization of processes (Fig. 6, Table 1). For the PS contrast this effect could also be observed in Broca’s region, however only in the ROI analysis (see Fig. 6).

The regions observed for CS are overlapping with those observed in studies of other types of linguistic manipulations, such as semantic integration errors (Christensen and Wallentin, 2011), semantic competition (January et al., 2009; Schnur et al., 2009; Snyder et al., 2007), syntactic complexity (Christensen and Wallentin, 2011; Christensen et al., 2013; Kristensen and Wallentin, 2009), syntactic working memory (Fiebach et al., 2005) and linguistic working memory in general (Kristensen and Wallentin, in press; Wallentin et al., 2006). Wallentin et al. (2006) found that Broca’s region activity correlated with response time in a working memory task where a visual scene was probed by linguistic cues. This finding was recently replicated (Kristensen and Wallentin, in press) in a reanalysis of a study of working memory for previously read sentences probed with linguistic cues (Wallentin et al., 2008). Again, Broca’s region activity was found to be correlated with response time, suggesting that increased linguistic working memory demands yield increased Broca’s region activation. In their study of semantic and syntactic manipulations, Christensen and Wallentin (2011) found that Broca’s region activity was also correlated with response time in a sentence acceptability judgment task, and crucially, in this experiment responses were only collected on every other task, making the neural effect independent of whether a response was given or not. This demonstrates that this is not a motor effect. Broca’s region activation has also been observed for linguistic comprehension where no overt responses were produced, such as for increased emotional intensity during listening to a narrative (Wallentin et al., 2011).

Together, the above-mentioned findings and those from the present experiment are compatible with theories suggesting that Broca’s region

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**Table 5**

<table>
<thead>
<tr>
<th>Lateralization</th>
<th>CS-Br-We</th>
<th>CS-VW-Br</th>
<th>PS-Br-We</th>
<th>PS-VW-Br</th>
<th>RS-Br-We</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS Broca</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CS Wernicke</td>
<td><strong>0.38</strong></td>
<td><em>0.26</em></td>
<td>0.11</td>
<td>0.10</td>
<td>0.12</td>
</tr>
<tr>
<td>CS VWFA</td>
<td><strong>0.32</strong></td>
<td><em>0.13</em></td>
<td>0.08</td>
<td>0.07</td>
<td>0.11</td>
</tr>
<tr>
<td>PS Broca</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PS Wernicke</td>
<td>0.12</td>
<td>0.13</td>
<td>0.08</td>
<td>0.07</td>
<td>0.11</td>
</tr>
<tr>
<td>PS VWFA</td>
<td>0.10</td>
<td>0.06</td>
<td>0.08</td>
<td>0.14</td>
<td>0.08</td>
</tr>
<tr>
<td>RS Broca</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RS Wernicke</td>
<td>0.11</td>
<td>0.13</td>
<td>0.08</td>
<td>0.11</td>
<td>0.08</td>
</tr>
<tr>
<td>RS VWFA</td>
<td></td>
<td></td>
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<td></td>
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</table>

Pearson correlation between lateralization indices for conceptual shift (CS), perceptual shift (PS) and response shift (RS) in Broca’s region, Wernicke’s region and visual word form area (VWFA).

Significant correlations are written in bold.

* p < 0.05 uncorrected.
** p < 0.005 uncorrected.
is involved in processing linguistic processing costs (Kristensen and Wallentin, in press). They may also to a certain extent be compatible with theories positing a role for Broca’s region in cognitive control (Novick et al., 2010). However, a key limitation demonstrated by the present results, is that the region is not involved in processing all kinds of cognitive control. Some conceptual content may be needed in order for Broca’s region to become activated.

The present study includes a relatively large number of participants (n = 58) and a large number of trials per participant (400). This ensures that our results are robust and replicable. Whole brain effects are corrected at the family wise error level for multiple comparisons and for main effects (CS, PS and RS) effects are further thresholded at a beta-estimate level larger than one (at 2nd level). This ensures that the large number of participants do not lead to significant, but negligible effect sizes. Subsequently, we conducted two more liberal analyses in an attempt to rule out false negatives. These show that the lateralization effect for CS in Broca’s region is present in 95% of our participants, a distribution similar to that seen in aphasia studies (Pedersen et al., 1995) and in auditory word form area (Dehaene and Cohen, 2011; McCandliss et al., 2003) and in sodium amytal (Wada) tests of cerebral speech dominance (Rasmussen and Milner, 1977; Woermann et al., 2003). The degree to which lateralization indices found in the present experiment can be predictive of aphasia symptoms remains to be studied.

We have focused our Broca’s region analyses on Brodmann areas 44 and 45. In our ROI analysis, we do not distinguish between these two regions. This is based on our prior findings that the effect observed with linguistic fMRI contrasts are usually located close to the border of BAs 44 and 45, extending into both regions (e.g. see Kristensen and Wallentin, in press for a review). Apart from the BA 44/45 effect, however, we also see a strong effect in BA 47 for CS. BA 47 has recently been shown to be structurally dissimilar from BAs 44 and 45 (Amunts and Zilles, 2012) and has, based on this, been suggested also to be functionally separate. Meta-analyses have pointed towards a role for BA 47 in semantic processing (Binder et al., 2009), which would be consistent with our findings. In the processing of polyrhythms (Vuust and Roepstorff, 2008), an increase in BA 47 has also consistently been observed, both for listening to and for producing polyrhythms compared to simple rhythms (Vuust et al., 2006, 2011). Interestingly, polyrhythm is a strong example of a cognitive conflict (Novick et al., 2010) where musicians play one rhythm (e.g. a 3/4 beat) on top of another rhythmic meter (e.g. a 4/4 beat). This imposes a strong sense of tension both in the listener and in the performer. This would suggest that this region is involved in the processing of cognitive conflicts, both at the conceptual (in the CS contrast) and at the non-conceptual level (as with music). This interpretation, however, would lead to the prediction that PS and RS should also evoke activation in BA 47, which we do not see in this study. At present, we are therefore not able to make a coherent interpretation of the pattern of activations observed in BA 47. Studies that separate cognitive conflict and semantic load are needed to shed light on these apparent inconsistencies. But we note that the results in the present study are very strong with a z-score of 7.6 in BA 47. We are therefore confident that the BA 47 effect is not a false positive and further that the observed lack of results for PS and RS probably do not reflect false negatives either, as we allowed a search for effects using a lowered statistical threshold in our conjunction analysis.

Wernicke’s region and visual word form area

Apart from the described results in Broca’s region, the main effect of perceptual shift (PS) yielded a predominantly posterior network of activations (Fig. 3, Table 2). These were located in the ventral and superior part of the temporal lobe. The ventral region was overlapping with the visual word form area (Dehaene and Cohen, 2011; McCandliss et al., 2003), while the superior regions were in the vicinity of Wernicke’s region, argued by Wernicke (1874) to be the seat of auditory sound images (“Klangbilder”), i.e. the auditory equivalent of the visual word forms. Since PS involves a translation from visual word forms to auditory word forms or vice versa, it makes sense that both regions are involved in this process. Our findings are thus consistent with the idea that these regions are involved in modality specific processing of words and that this effect is enhanced when the stimuli need to be recoded into a new modality. With the lowered threshold and in the ROI analyses we also saw an effect of CS in both regions.

Similar to the effect in Broca’s region, activation in Wernicke’s region and VWFA were found to be highly left lateralized, although the lateralization effects did not reach as high as CS did in Broca’s region. In Wernicke’s region 81% were found to be left lateralized in the CS task and 71% in the PS task. In VWFA 78% were found to be left lateralized in the CS task and 74% in the PS task. Another difference was that while the lateralization in Broca’s region was composed of both an increase of activity in the left hemisphere and a decrease in activity in the right hemisphere (Fig. 6) for CS and PS, this was only the case for PS in Wernicke’s region and for neither CS nor PS in VWFA. Both left and right hemispheres were found to yield positive responses to CS on average and the lateralization thus came about as a relative difference in positive responses (see Fig. 6).

Lateralization across regions and tasks

When comparing lateralization across tasks and regions of interest, it was found that lateralization indices for Broca’s region, Wernicke’s region and VWFA were correlated within tasks whereas lateralization for the different tasks was not correlated within regions (see Table 5). Thus, although both task contrasts produce highly lateralized effects, these may not be directly comparable. One previous study found that the hemispheric lateralization of the VWFA was correlated with the lateralization of the LIFG (Cai et al., 2010). In our study we thus replicate this result, but crucially, only within task.

Apart from the fact that the BOLD signal is a composite measure that depends on a number of factors other than brain activity (Gusnard and Raichle, 2001; Logothetis, 2008), the lack of correlation across tasks suggests that lateralization may be a complex phenomenon where an individual may show more or less lateralization for different conceptual tasks, depending on e.g. language proficiency and prior experience. Differences in lateralization for the PS contrast may thus reflect reading abilities and reading strategies whereas lateralization for CS may reflect other more or less orthogonal skills, such as vocabulary size. Other language tasks may display different lateralization patterns and this leads to the hypothesis that different brain regions within the language network may be differently linked for different tasks and that these links form functional specialization rather than individual brain regions by themselves. Further studies are needed in order to test these hypotheses.

Lateralization has been found to correlate with handedness, both when studied using fMRI (Szaflarski et al., 2002) and functional transcranial Doppler sonography (Knecht et al., 2000a, 2000b). In this study we only included right-handed participants. We therefore cannot say whether the observed lateralization effects would show different relationships to handedness, although it seems likely.

Two-way and three-way interactions

Both in the behavioral and in the neuroimaging data a strong interaction between CS and RS and between PS and RS was found. The CS × RS interaction reflects increased processing costs if the current trial is a CS without RS or if the current trial is a repeat (nonCS) but with RS. In both cases this involves a CS being present in the previous trial. This effect is clearly visible in the response time data (Fig. 2, top) where columns 2 and 4 represent CS in the absence of RS and columns 5 and 7 represent RS in the absence of CS. These columns are all taller than their direct counterparts (columns 1, 3, 6 and 8). This means that there is a lingering response time cost of almost 63 ms if the previous trial was a CS. The PS × RS interaction reflects a similar lingering processing cost (25 ms) if the previous trial was a perceptual shift. Such
short-term dependencies in response times are well-known (Kelly et al., 2001; Laming, 1979; Spivey, 2007; Van Orden et al., 2003), but are not, to our knowledge studied in an fMRI context. In the ROI analysis we see a CS × RS interaction in Broca’s region, but when looking at the more conservative whole brain results we see that the peak activations are outside Broca’s region, suggesting that the ROI result may be a spillover effect from neighboring regions (insula, middle frontal gyrus — see Table 4, Fig. 4). The whole brain effects also reveal that the anterior insula effect has a bilateral distribution (although more pronounced towards the left, see Fig. 4; Table 4). Crucially, the CS × RS interaction was not found to be lateralized in the ROI analysis, i.e. no three-way interaction (CS × RS × hemisphere) was observed (F(1,57) = 0.002, p(bonf) = 1). We therefore conclude that the effect from the previous CS trial in Broca’s region is not similar to the effect observed for the current CS trial.

In Wernicke’s region, however, this exact three-way interaction (CS × RS × hemisphere) was observed in the ROI analysis (Fig. 7). In the posterior temporal area we thus in addition to the increased lateralization for CS and PS in isolation find an effect of the preceding trial. If the current trial is a CS then we find greater lateralization if the trial does not include a response shift (nonRS), indicating that the previous trial was also a CS (columns 2 and 4 vs. 1 and 3 in Fig. 7). If, however, the current trial is a nonCS, then we find greater lateralization if the trial is also an RS, i.e. again indicating that the previous trial was a CS (columns 5 and 7 vs. 6 and 8 in Fig. 7). If the brain thus has a BOLD equivalent of the history effects observed in the behavioral data, our findings suggest that it originates from Wernicke’s region rather than Broca’s region. One problem with this type of analysis in fMRI is the poor temporal resolution of the BOLD response. An interpretation of the regional interaction effects found in the whole brain analysis could be that at least some of them reflect regions with a delayed BOLD response from the previous trial in regions with a different response function from the standard hemodynamic response function incorporated in this analysis. Anterior insula has been found to have a slower HRF, similar to that of a down-stream venous signal (Gonzalez-Castillo et al., 2012; Kruggel and von Cramon, 1999a, 1999b). In theory, the insula signal thus might reflect previous activity in Broca’s region (or elsewhere), reaching the veins in insula with a 2–5 s delay (the intertrial interval in the current experiment) compared to a cortical signal. Similarly, we see interaction effects in posterior brain regions, such as the precuneus (see Tables 3–4). This region also holds large draining veins that may transport blood from posterior perceptual regions. Again, we may be dealing with a down-stream effect of the previous trials (Kruggel and von Cramon, 1999a, 1999b). Further studies are needed to elucidate these claims.

Repetition suppression and prediction error

Task shift effects reflect the inverse of what is known as repetition suppression, i.e. the fact that the neural responses observed for various stimuli are often found to be attenuated when repeated (Dehaene et al., 2001; Orfanidou et al., 2006; Raichle et al., 1998; Summerfeld et al., 2011; Summerfeld et al., 2008; Todorovic et al., 2011). Repetition suppression has been interpreted as an instance of predictive neural coding and predictive coding has been suggested as an overarching framework for how the brain works (Clark, 2013; Friston, 2010). In this sense task shift effects may be hypothesized to reflect a prediction error in relation to a kind of status quo bias. Such a bias has been also observed in relation to certain cognitive tasks (e.g. Kahneman et al., 1991; Nicolle et al., 2011). Our results may thus be interpreted to reflect a very simplistic kind of prediction error, working on a trial by trial basis. Predictive coding within cognition and language, however, of course have to work on multiple timescales from phoneme prediction via words to prediction of word order and pragmatic content (Levy, 2008). Across the present experiment, for example, the CS words were more frequent than the nonCS words (250/150) and in the frame of the whole experiment, the CS trials should thus yield less prediction error than nonCS words. This effect would work in the opposite direction of the trial-by-trial shift effect. Similarly, a CS or PS following another CS or PS should be less costly. In fact we see the opposite. There is a lingering response time penalty for a CS following a CS (62 ms) and for a PS following a PS (25 ms). This history effect is also present in the neuroimaging data in the widespread CS × RS and PS × RS interactions (see above). In the ROI analysis of Wernicke’s region we additionally find the history effect in lateralization where the lateralization level is found to reflect the nature of both the current and the previous trials (Fig. 7). These effects are difficult to reconcile with a priming or prediction error approach. It is therefore not clear how far an interpretation of the activity in the investigated regions as reflecting a more general linguistic prediction error can be taken.

Adding to this complexity is the observation of a widespread set of deactivations, including parietal and right hemisphere regions, as a function of conceptual shift (i.e. regions that are more active during non-shifts than during shifts — see Table 1). These effects are also hard to reconcile within a framework where prediction error is the main driver of neural responses. Considering the memory components of the current experiment, the main difference between CS and nonCS conditions is in the comparison between the previous word (encoded in short-term memory) and the incoming word. In the CS condition there is a mismatch between these two items plus the current word has to be encoded for the next trial. In nonCS trials this is not necessary as it was already encoded during the previous trial. It is difficult to see that there are cognitive operations in the nonCS condition that are not present in the CS, except for recognition. But given that the whole priming literature suggests that recognition is primarily linked to decreases in activation, it seems difficult to argue why it should be responsible for the widespread activations that we see as a negative effect in the CS—nonCS contrast. It becomes even more complicated, since the parietal regions found to be deactivated as a function of CS to a large extent are the same as those that are found in the CS × RS interaction (Fig. 4, Table 4). This means that these regions are not only less activated if the current trial is a CS but they are also more activated if the previous trial was a CS. This type of strange switch behavior is intriguing but very hard to interpret. Further studies are needed to investigate if this effect is real or somehow linked to issues related to the sluggish nature of the BOLD response.

Conclusion

We conclude that Broca’s region (BAs 44 & 45) activation and lateralization for task shift effects is very strong, and selective for task shift effects with a conceptual content. Similar effects are observed in Wernicke’s region and the visual word form area, and lateralization is correlated across regions, however only within tasks. Lastly, response time and lateralization in Wernicke’s region not only reflect trial type, but also trial history.

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