Pantomimes of object use require accurate representations of movements and a selection of the most task-relevant gestures. Prominent models of praxis, corroborated by functional neuroimaging studies, predict a critical role for left parietal cortices in pantomime and advance that these areas store representations of tool use. In contrast, lesion data points to the involvement of left inferior frontal areas, suggesting that defective selection of movement features is the cause of pantomime errors. We conducted a large-scale voxel-based lesion–symptom mapping analyses with configurational/parietal (CS) and body-part-as-object (BPO) pantomime errors of 150 left and right brain-damaged patients. Our results confirm the left hemisphere dominance in pantomime. Both types of error were associated with damage to left inferior frontal regions in tumor and stroke patients. While CS pantomime errors were associated with left temporoparietal lesions in both stroke and tumor patients, these errors appeared less associated with parietal areas in stroke than in tumor patients and less associated with temporal in tumor than stroke patients. BPO errors were associated with left inferior frontal lesions in both stroke and tumor patients. Collectively, our results reveal a left intrahemispheric dissociation for various aspects of pantomime, but with an unspecified role for inferior frontal regions.

Keywords: frontal, ideomotor apraxia, lesion, pantomime, parietal, voxel-based lesion–symptom mapping

Introduction

Pantomime of object or tool use is the act of pretending to use an object by adopting the same limb configurations and producing the same sequences of movements as if the object were actually held and used. Pantomime of object or tool use is the act of pretending to use an object by adopting the same limb configurations and producing the same sequences of movements as if the object were actually held and used. Within the model of praxis by Rothi et al. (1991, 1997), pantomime to verbal command is distinguished from other types of motor productions based on the fact that it neither requires a visual analysis of the gesture to be produced nor a comparison between the visual input with a lexicon of action (as would be the case for, e.g., imitation of new or familiar gestures). Rather, the analysis of the auditory/verbal command is directly followed by the selection of the spatiotemporal attributes of the gesture to be performed from an action output lexicon and the programming and implementation of the motor action (see also Peigneux and Van der Linden 2000).

Because the production of pantomimes involve semantic, executive, and spatial/configural level of motor processing (to respectively understand the gestures, select the relevant movements representing the action, and represent accurately the relationships between the body parts involved in the movement and of how they interact with the object (Goldenberg 2009)), pantomime constitutes a sensitive task to detect ideomotor apraxia following a brain lesion (Heilman and Rothi 1993). In the current article, we refer to “ideomotor apraxia” using the definition proposed by Rothi et al. (1991, 1997): “an impairment in the timing, sequencing, and spatial organization of gestural movements” (Rothi et al. 1991).

Starting from the seminal hypotheses of Liepmann (1908) in stroke patients, most prominent models of praxia advance that left parietal areas store the motor representations of tool use guiding action and therefore predict that these structures play a central role in pantomime (Moll et al. 2000; Peigneux et al. 2004). Functional neuroimaging studies corroborate these models by consistently observing correlations between left parietal areas activity and pantomiming (Vingerhoets, Acke et al. 2012, Vingerhoets, Vandekerckhove et al. 2011; see Lewis (2006) for a meta-analysis of activation studies).

In contrast, lesion studies report that accurate pantomime depends on the integrity of left inferior frontal areas (Goldenberg et al. 2007) and less consistently of parietal areas (Kertesz and Ferro 1984; Goldenberg and Hagmann 1997; Peigneux et al. 2000). Although not directly for pantomimes, parietal areas have been involved in ideomotor apraxia (Basso et al. 1985; Haaland et al. 2000; Buxbaum et al. 2007) or coordination of arm movements in ideomotor apraxia (Mutha et al. 2010). Therefore, lesion studies conclude that pantomime critically depends on the selection of a limited, task-relevant set of features among the many features involved in the actual tool use to be mimed (Goldenberg et al. 2007; Goldenberg 2009; Bohlhalter et al. 2011). The disparity between the findings of neuroimaging and lesion approaches about the involvement of parietal regions has been hypothesized to follow from the pantomimes being realized under different conditions in each type of study. Because of the constraints induced by the scanner on participant's movements, the pantomimes require additional spatial transformations of movements to unusual reference frames, which in turn increase the involvement of parietal structures (Andersen et al., 1997; Goldenberg et al., 2007). Rumit et al. (2004), however, reported an involvement of left parietal areas for the pantomiming of visually presented objects in patients with deficit in the organization of sequences relative to tool use (ideational apraxia), suggesting that these structures might trigger tool use-related motor programs. Pantomime are...
globally sensitive to left hemisphere lesions (Bickerton et al. 2012), but have been found to be sensitive to lenticular stroke and associated to impaired working memory, suggesting that correct pantomime execution necessitates an efficient lexical route but also a dedicated workplace subserved by subcortical structures (Bartolo et al. 2003). Of note, electroencephalography studies manipulating the production of pantomimes in naturalistic conditions showed evidence for parietal activation in preparing tool-use movements, suggesting that this region is not only involved in spatial transformation, but also in planning tool-related motor actions (Wheaton, Shibasaki et al. 2005; Wheaton, Yakota et al. 2005). During the neuropsychological assessments, the patients produce the movements within a natural body-centered reference frame and with visual feedback (Goldenberg et al. 2007; Goldenberg 2009), which involves only routine support from parietal areas.

However, several other hypotheses could account for the discrepancy between the results of functional and lesion studies. The contribution of parietal structures to pantomime might have been underestimated in previous lesion studies due to the assessment having been conducted in chronic patients, that is, more than 1 month after lesion onset (Goldenberg 2003a, 2003b; Dovern et al. 2011). Specificity of networks may indeed be revealed in the postacute phase only, after the resorption of the ischemic penumbra (Witte et al. 2000) and before the occurrence of major plastic anatomofunctional reorganizations (Adriani et al. 2003; Rey et al. 2007). The type of evaluation (conceptual vs. production components) may be more sensitive to parietal or to frontal lesions (Halsband et al. 2001). Furthermore, previous neuropsychological studies included only a limited number of patients with lesions covering only limited portion of the brain (Goldenberg 2003a, 2003b), patients selected based on a priori hypotheses on the region of interest (Dovern et al. 2011; Hanna-Pladdy et al. 2001) or patients with aphasia (Goldenberg et al. 2007). Finally, previous studies dichotomized behavioral data on apraxia (with vs. without apraxia) instead of considering the scores as continuous data, leading to a loss of power and reduced effect sizes (Cohen 1983). Collectively, these potential caveats could have lead to false-negative results in current lesion data on pantomime, potentially concerning the involvement of parietal areas.

In addition, attempts to find common substrates for different types of pantomime errors and the rarity of some kinds of errors motivated researchers to collapse together various types of error in neuropsychological scoring of pantomime. As lesions to distinct areas may induce distinct types of error (Rumiati and Humphreys 1998; Halsband et al. 2001; Hanna-Pladdy et al. 2001; Rumiati et al. 2001), the use of compound scores might have in turn contributed to obscure putative intrahemispheric dissociations for different types of pantomime errors. For instance, body-part-as-object (BPO) pantomime errors, consisting in representing objects with a part of the body rather than pretending to use an “invisible” object as specified in the test instructions (Goodglass and Kaplan 1965) have been suggested to depend on frontal but not parietal components (Peigneur and Van der Linden 1999; Arzy et al. 2006). The study of BPO errors could thus help to further reveal intrahemispheric dissociations between frontal and parietal contributions to pantomime.

To test these hypotheses, we conducted large-scale retrospective voxel-based lesion–symptom-mapping analyses (VLSM; Bates et al. 2003) on a group of subacute, unselected, hemispheric brain-damaged patients and pantomime scores differentiating the typical spatial/configural (CS) and BPO pantomime error types. We used highly selective inferential statistical analyses of lesion–symptom mapping based on continuous scores rather than descriptive comparisons between lesion patterns of patients’ groups defined by behavioral cutoffs (i.e., with or without apraxia). Because the inclusion of tumor and stroke patients in VLSM analyses might yield different results, we analyzed separately these 2 types of brain-damaged patients (e.g., Karnath and Steinbach 2011).

Methods
Patients
One hundred and fifty right-handed patients with a first right or left unilateral hemispheric lesion (demographic data in table 1) were selected retrospectively from consecutive in-patients admitted to the Neuropsychology and Neurorehabilitation Service of the Centre Hospitalier Universitaire Vaudois or the Hôpitaux Fribourgeois between 2007 and 2011. Patients with bilateral lesions were excluded to facilitate the interpretability of our results in terms of hemispheric specialization of pantomime. On average, the pantomime assessment was conducted 2.3 ± 6.9 weeks (mean ± SD) after the lesion onset or tumor diagnosis or removal and was part of the formal neuropsychological assessment carried out by experienced psychologists specialized in neuropsychology. All patients met the following criteria: 1) first unilateral hemispheric lesion without damage to the brain stem or cerebellum documented by CT-scan and/or MRI; 2) no prior neurological illness; 3) no psychiatric illness; 4) good cooperation and absence of major behavioral or attentional problems; 5) sufficient understanding of the instructions; and 6) assessment of at least 4 pantomimes. Inclusion in the study was neither determined by the lesion characteristics nor by the pattern of behavioral deficit. The study was carried out in agreement with the recommendations of the Ethics Committee of the Faculty of Biology and Medicine, Lausanne. Data were handled according to Swiss-Federal law on data protection.

Neuropsychological Assessment of Pantomime
The production of pantomime on verbal command was assessed by asking the patient to mime the use of an imaginary tool. In the case of hemiparesis, the patient was asked to use only the nonparetic ipsilesional hand. The evaluation of paresis was based on the Medical Research Council Scale for Muscle Strength (e.g., Pizzi et al. 2009). In the present study, if the patient’s strength was below 3, the nondominant (left) hand was used. Pantomimes were tested with items of the

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Demographic data of the 150 patients included in the study</th>
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<tbody>
<tr>
<td>N = 150 Patients</td>
<td></td>
</tr>
<tr>
<td>Age, mean (±SD)</td>
<td>60.5 (±15.3), 16–89 years</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>68</td>
</tr>
<tr>
<td>Male</td>
<td>82</td>
</tr>
<tr>
<td>Damaged hemisphere</td>
<td></td>
</tr>
<tr>
<td>LBD</td>
<td>84 (38 with aphasia: 32 patients with fluent aphasia and 6 with nonfluent aphasia)</td>
</tr>
<tr>
<td>RBD</td>
<td>66</td>
</tr>
<tr>
<td>Etiology</td>
<td></td>
</tr>
<tr>
<td>Stroke (LBD, RBD)</td>
<td>81 (42, 39)</td>
</tr>
<tr>
<td>Tumor (LBD, RBD)</td>
<td>69 (42, 27) (44 before and 25 after surgery)</td>
</tr>
<tr>
<td>Postlesion delay (weeks ± SD)</td>
<td>2.3 (±6.9)</td>
</tr>
</tbody>
</table>

| LBD, left brain-damaged; RBD, right brain-damaged. |
screening batteries by Peigneux and Van der Linden (2000) or Mahieux-Laurent et al. (2009). The former included 4 items: brushing the hair with a comb; brushing the teeth with a toothbrush; planting a nail with a hammer; and sawing a branch of wood (Peigneux and Van der Linden 2000). The latter included 5 items: planting a nail with a hammer, tearing a piece of paper in two, lighting a match, brushing the hair with a comb, and drinking a glass (Mahieux-Laurent et al. 2009). On average 4.3 ± 0.5 (mean ± SD; range 4–5) items were probed for the assessment of pantomimes.

Two different types of errors were documented and analyzed in the present study: First, CS errors refer to inaccurate limb configurations during pantomiming at the level of the sequencing, timing, and/or amplitude of the gestures, and of the relationships between the different body parts engaged in the movements. CS errors result in imprecise or unrecognizable gestures. For example, if the patient mimed brushing his hair with an imaginary comb with one hand but placed the imagined comb far away from his head, the pantomime was considered as incorrect. The second type of errors consisted in the use by the patients of their body parts as the object (BPO errors). BPO were considered as errors only when the patients did not correct it after reinstruction from the examiner. The patient was reinstructed after every BPO error. This reinstruction condition was implemented because BPO are common among healthy controls, but neurologically healthy population correct BPO after being reinstructed (Heilman and Rothi 1993; Raymer et al. 1997; Peigneux and Van der Linden 1999).

For example, for combing the hair, if the patient used his fist as the comb and brushed his hair with it, the examiner reinstructed him to pretend he was holding an imaginary comb in his hand rather than using his forelimb as the comb. If the patient did not correct the error and continued to use his limb as the object, the BPO was considered as pathological and counted as a BPO error (Heilman and Rothi 1993; Raymer et al. 1997). We would note that because the present study was based on a retrospective analysis of data collected during routine neuropsychological assessments, the scoring of pantomime was not as precise and controlled as what could have been obtained with a specifically designed prospective study. For this reason, not all possible error types [including, e.g., semantic content errors (Rumiati and Humphreys 1998); sequence or conceptual errors (Rumiati et al. 2001); or parapraxic errors (Halsband et al. 2001)] were analyzed separately. However, the pantomime errors were scored by trained and experienced specialists in neuropsychology, according to strict published procedures (Peigneux and Van der Linden 2000; Mahieux-Laurent et al. 2009). Because each patient did not have to produce the same number of pantomimes (4 items for Peigneux’s battery and 5 items for Mahieux-Laurent’s battery), the scores used in the VLSM were the standardized numbers of BPO and CS pantomime error types (%). Pantomime scores obtained with Peigneux’s battery or Mahieux-Laurent’s battery did not statistically differ, neither for BPO errors ($t(148) = −0.43; P = 0.66$) nor for CS errors ($t(148) = 0.56; P = 0.57$).

Voxel-Wise Statistical Analysis of Lesion-Symptom Mapping

Brain lesions were manually reported on axial slices of the standard Montreal Neurological Institute’s (MNI) brain template using the MRIcro software (Rorden and Brett 2000), according to previously described methods (Karnath et al. 2004; Spierer et al. 2009). Lesions were reported on the template brain by trained assistants naive to the clinical profiles of the patients (Fiez et al. 2000). These normalized lesions were then submitted to statistical mapping analyses using VLSM algorithms implemented in the MRICroN and NPM softwares (Rorden et al. 2007) to determine brain areas where damage yielded each type of pantomime errors. Because each patient did not have to produce the same number of pantomime, the scores used in the VLSM were the standardized numbers of BPO and CS pantomime error types (%). The t-tests on the continuous CS and BPO scores were performed on a voxel-by-voxel basis to compare performance in patients with versus without a lesion in each voxel, only testing voxels damaged in at least 4 patients. The results of the t-tests were then color-coded and mapped on the MNI template brain using the software package (Rorden and Brett 2000). Only voxels surviving a conservative false discovery rate (FDR) corrected significance threshold of $P < 0.05$ were considered in the results (though a threshold of 0.01 was applied for the analyses of the tumor and stroke patients collapsed together presented in the Fig. 1 because we reached a much larger sample size, see the Results and discussion section).

The overall distribution of lesion among our patient sample is depicted in Supplementary Figure S1a.

Results

We conducted the VLSM analyses on the groups of stroke and tumor patients separately (see Table 1 for demographic information of each subgroup).

In stroke patients, CS pantomime error types were associated with lesions to a network centered on inferior frontal and temporal areas, with sparse evidence for a role of parietal areas (Fig. 1b). In tumor patients, CS error types were associated with lesions to a more posterior network extending from inferior frontal to parietal areas, mostly including parietal white matter (Fig. 1e).

In stroke patients, BPO pantomime errors were associated with lesions to the left middle and inferior frontal gyri, the rolandic and inferior frontal opercula, and the underlying white matter, mainly including the superior longitudinal fasciculus (Fig. 1c). In tumor patients, BPO errors were associated with the same network, but extending higher to the supplementary motor area (Fig. 1f; see Supplementary Fig. S2 for the double dissociation between CS and BPO errors).

We also conducted the same VLSM analyses as above with stroke and tumor patients collapsed together. CS pantomime error types were associated with lesions to the left inferior parietal and angular gyri, postcentral and supramarginal gyri, and portions of the underlying white matter (Supplementary Fig. S1c). BPO pantomime errors were associated with lesions to the left middle and inferior frontal gyri, the rolandic and inferior frontal opercula, and the underlying white matter, mainly including the superior longitudinal fasciculus (Supplementary Fig. S1d) (see Supplementary Fig. S2 for the double dissociation between CS and BPO errors). Comparison between the results of the analyses of the lesions associated with CS and BPO errors revealed that the left inferior frontal regions predicted the occurrence of both types of error (Supplementary Fig. S1cd). We further tested putative effects of lesion size on the occurrence of CS and BPO errors. Lesion size differed between patients with versus without CS errors ($P < 0.05$, uncorrected) but not for patients with versus without BPO errors ($P = 0.29$, uncorrected). However, there was no evidence for correlations between the CS or BPO errors and the size of lesions ($r(26)=0.16; P = 0.40; r(11) = −0.10; P = 0.74$, respectively).

The incidence of the different patterns of error (CS, BPO, or CS+BPO) across patients is displayed in Table 2. The relationship between impaired pantomiming and aphasia is depicted in Figure 2. There were more apraxic patients in the group “left hemispheric with aphasia” than in the group “left hemispheric without aphasia” and in the group “right-hemispheric.” The analysis of the incidence of at least 1 pantomime error in these 3 groups of patients further reveal that very few patients show both CS and BPO errors (Supplementary Fig. S3).
Figure 1. Voxel-based lesion–symptom mapping on the stroke patients and tumor patients separately shows the relationship between performance in pantomime and brain lesions. (a) Overlap lesion plot of the 81 stroke patients. The number of overlapping lesions is coded with colors ranging from dark red (n = 1) to light yellow (n = 18 patients). (b) Only voxels significant at P < .05 FDR corrected are color-coded ranging from red to white. Configural spatial errors were associated with lesions to a network centered on left inferior frontal and temporal areas, with sparse evidence for a role of left parietal areas. (c) Body-part-as-object errors were associated with lesions of the left middle and inferior frontal gyri and the rolandic inferior frontal opercula, and the underlying white matter mainly including the superior longitudinal fasciculus. (d) Overlap lesion plot of the 69 tumor patients. The number of overlapping lesions is coded with colors ranging from dark red (n = 1) to light yellow (n = 11 patients). (e) Only voxels significant at P < .05 FDR corrected are color-coded ranging from red to white. Configural spatial errors were associated with lesions to the left inferior frontal and inferior and superior parietal gyri, angular gyrus, postcentral and supra marginal gyri, largely including the underlying white matter. (f) Body-part-as-object errors were associated to lesions of the left middle and inferior frontal gyri and the underlying white matter mainly including the superior longitudinal fasciculus. Brain slices are displayed from z-coordinates −16 to 48 of the MNI space, with the left hemisphere on the right side.
Incidence and range of each error type

<table>
<thead>
<tr>
<th>Error (≥1)</th>
<th>Total (n = 150), %</th>
<th>Mean % of error (± SD) + range</th>
</tr>
</thead>
<tbody>
<tr>
<td>No error</td>
<td>76</td>
<td>0 ± 0 + (0–15%)</td>
</tr>
<tr>
<td>CS only</td>
<td>15</td>
<td>36.8 ± 18.0% (10–75%)</td>
</tr>
<tr>
<td>BPO only</td>
<td>6</td>
<td>45.4 ± 16.6% (20–75%)</td>
</tr>
<tr>
<td>CS + BPO</td>
<td>3</td>
<td>CS: 43.8 ± 24% (25–75%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BPO: 31.3 ± 12.5% (25–50%)</td>
</tr>
</tbody>
</table>

CS, configural/spatial error; BPO, body-part-as-object error.

Discussion

We conducted VLSM analyses based on a large cohort of 150 unselected patients with unilateral left or right hemispheric brain damage and their continuous scores in BPO and CS errors types in a classical neuropsychological assessment of pantomime. Our results reveal that distinct lesion sites within the left hemisphere predict the occurrence of CS and of BPO pantomime error types. Both types of error were associated with damage to left inferior frontal regions in tumor and stroke patients. CS errors were associated with lesions of left inferior parietal areas, whereas BPO errors were associated with lesions extending from left superior to inferior frontal gyri and a large portion of the underlying white matter in both tumor and stroke patients. Of note, we put forward a differential pattern of deficits according to etiology for CS errors: CS errors were less associated with parietal areas in stroke than in tumor patients, and temporal areas were less associated with CS errors in tumor than stroke patients.

Most of the previous investigations of pantomime in brain-damaged patients included only left brain-damaged patients and/or samples selected based on the presence of aphasia (Schnider et al. 1997; Hanna-Pladdy et al. 2001; Goldenberg 2003a, 2003b; Goldenberg et al. 2007; Dovern et al. 2011). In contrast, our study includes unselected consecutive patients sustaining both left and right unilateral brain damage. Our finding for a left hemispheric dominance in pantomime thus provides robust lesion evidence for the prominent involvement of left but not right hemispheric structures in pantomime and corroborates functional imaging studies documenting a left hemispheric specialization for pantomime (Hermsdorfer et al. 2007; Vingerhoets, Acke et al. 2012, Vingerhoets, Vandekerckhove et al. 2011). A limitation of our results in this regard is that because we included only right-handed patients, the present study cannot disentangle potential interactions between handedness, the hand used for the pantomime and the side of the lesion. However, our results are consistent with previous evidence for a left lateralization in pantomimes in both left- and right-handed individuals (Vingerhoets, Acke et al. 2012) and for a similar left lateralization in studies comparing left- and right-hand pantomimes in right-handed participants (Moll et al. 2000; Choi et al. 2001). Of note, the VLSM analysis revealed the brain regions inducing the “more severe” pantomime impairments (Fig. 1, Supplementary Fig. S1), which does not rule out that other regions play a role in pantomime. Although the left parieto-frontal network revealed in our results is the region which, when damaged, induces the most robust increase in the number of pantomime errors, right-hemispheric brain regions (or left nonfrontal, nonparietal regions) might be involved in praxis as well. Indeed, impaired pantomiming manifest in 8% of the right brain damaged patients (Fig. 2) and the lesion overlap of patients showing a deficit in pantomime reveals associations between right hemispheric lesions and apraxia (see Supplementary Fig. S2b).

A separated VLSM analysis of the lesion sites associated with the commission of distinct types of pantomime errors revealed a left intrahemispheric dissociation between the contribution of parietal and frontal areas to CS and BPO error types, respectively.

In tumor patients, our finding for a role of left parietal areas in CS pantomime errors is in line with findings from functional imaging approaches (Moll et al. 2000; Hermsdorfer et al. 2007; Vingerhoets, Acke et al. 2012, Vingerhoets, Vandekerckhove et al. 2011). These studies interpreted the left parietal activity during pantomime as supporting the storage of knowledge about manipulation of familiar objects and learned gestures, both mechanisms being specific to praxia and necessary for accurate pantomime. Damage to this region is thus conceivably at the origin of the CS error types we observed (Buxbaum et al. 2007; Vingerhoets, Acke et al. 2012, Vingerhoets, Vandekerckhove et al. 2011). However, our analyses revealed a critical role for the parietal white matter in pantomime in tumor patients, suggesting that a disruption of the functional interactions between the subparts of the frontal-parietal network involved in pantomime would yield even more CS errors than focal damage to their constitutive regions (Peigneux et al. 2001; Wheaton, Shibasaki et al. 2005; Wheaton, Yokota et al. 2005). In line with this finding, frontal-parietal and basal ganglia damage induced by corticobasal degeneration have been shown to induce severe apraxic symptoms (Leiguarda et al. 2000).

In contrast to these activation studies and to our finding in tumor patients, lesion data so far mostly report that the integrity of left frontal but less consistently parietal areas are necessary for pantomime (Goldenberg et al. 2007; see also Bohlhalter et al. 2011 for supporting Transcranial Magnetic Stimulation data). Moreover, sparse evidence from single case reports describe patients with left parietal damage but preserved pantomime (Goldenberg and Hagmann 1997; Peigneux et al. 2000). These lesion data have been interpreted in terms of the involvement of inferior frontal regions in the selection of task relevant gestures among all gestures possibly related to a given tool or object use (Goldenberg et al. 2007).

Goldenberg et al. (Goldenberg et al. 2007; Goldenberg 2009)
advanced that the disparities between the findings of functional and lesions studies on the role of parietal regions in pantomime could follow from the gestures being realized under different conditions in the scanners when compared with during neuropsychological assessments. In the scanner, more spatial transformation would be required because the movements have to be performed with constrained limb positions, within unusual portions of space and without visual feedback. These additional demands would have artificially increased the involvement of the parietal structure supporting spatial processing and transformations into coordinates (Sack 2009; though see Rumiati et al. 2004 for evidence of parietal activity even when participant were not instructed to perform the gestures in the scanner). However, our result for a parietal involvement in the absence of any extra demand on spatial transformation calls for additional accounts for the lack of associations between parietal damage and CS pantomime errors observed in previous lesion studies. The following hypotheses could be put forward in this regard. First, previous studies included only stroke patients; our results for a much stronger association between CS errors and parietal areas in the tumor than in the stroke group suggest that the etiology of the lesion might play a role in their functional consequences on apraxia. While some evidence suggest that stroke and tumor results in the same deficits (Haaland and Delaney 1981), other pointed out that these 2 etiologies could yield distinct patterns of deficits, even if lesion size and location is controlled (Anderson et al. 1990). Our results suggest that lesion location associated with pantomime errors might differ depending on whether the VLSM analyses are based on stroke versus tumor lesions. Lesion–symptom mapping based on tumoral lesions has been argued to induce different patterns as when stroke lesions are analyzed. A possible reason for these discrepancies could be that in tumor patients, infiltrations could yield functional loss while being invisible to the MRI or CT scans used to delineate the lesion loci during the lesion reconstruction, in turn confounding the mapping between lesion and symptoms (for discussion, see Anderson et al. 1990; Karnath and Steinbach 2011; Shallice and Skrap 2011). Interestingly, in line with the previous lesion data reviewed above (e.g., Goldenberg et al. 2007), the result for a parietal involvement in CS error almost vanished when analyses were conducted in stroke patients only. However, we would note that specifically designed studies should be conducted to elucidate the differential role of lesion versus tumor patients (Duffau 2011). The results of VLSM analyses are indeed highly dependent on the spatial distribution of the lesion because it not only determines where in the brain the VLSM tests are actually conducted, but also the distribution of the statistical power of the statistical tests conducted at each voxel between the behavioral scores of lesioned versus intact patients (Kimberg et al. 2007; see Method section). This factor possibly account for our differential pattern of results in the 2 groups of patients as evident from the difference in the lesion overlap in Figure 1a,d showing that lesion location are not strictly identical in stroke and tumor patients.

In this regard, the fact our study included a large cohort considerably increased the statistical power of our analyses and the portion of the brain covered by lesions. This factor could also explain why, in contrast to previous lesion studies, we reveal a parietal involvement in pantomime (though mostly in tumor patients). The inclusion of both left and right brain-damaged patients in the VLSM further strengthened the sensitivity of our statistical tests by increasing the number of data-points (i.e., the behavioral scores) in the groups of the intact and lesioned patients compared at each voxel. We also analyzed continuous data instead of dichotomizing the scores into normal versus impaired based on behavioral cutoffs, thereby taking into account the severity of pantomime impairment in the VLSM and maximizing the statistical power of the analyses (Cohen 1983).

Finally, although the following reasoning only applies to stroke patients where the association between parietal areas and CS error was weak, our results might have revealed a parietal involvement because pantomime was assessed during the subacute phase and not during the chronic phase as in most previous studies. In the literature so far, pantomime scores were collected at postlesion delays of about 28 weeks (Goldenberg 2003a, 2003b); 20 weeks (Goldenberg et al. 2007); or 4 weeks (Schnider et al. 1997). In contrast, pantomime scores in our study were collected on average 2 weeks after lesions onset, a period corresponding to the subacute phase. The specificity of parietal networks for pantomime was possibly revealed in the present study because pantomime was assessed after—or at least during—the release of areas surrounding damaged regions from ischemic penumbra (Witte et al. 2000), but before major plastic anatomofunctional reorganization took place (Adriani et al. 2003). Although highly speculative, parsimonious explanations for the fact that the functions preserved by parietal but not frontal areas recovered in chronic patients would be that 1) parietal mechanisms could be hierarchically subordinated to frontal selection processes and more specific to the pantomime task; and/or 2) the largely acquired and mnesic nature of parietal movement representations could be more prone to be recovered and taken over by other areas than frontal executive selection mechanisms. We would note, however, that our interpretation of the results on the influence of the post lesion delay is not made by directly comparing subacute versus chronic patients but in the light of previous literature. Consequently, the present study does not allow drawing definitive conclusions on the influence of postlesion delay functional recovery in pantomime, but rather calls for further investigations specifically designed to disentangle the precise influence of this factor.

The results of the VLSM revealed a role of temporal regions in CS error, mostly in stroke patients. This finding is in line with models positing that the knowledge on tool use required to perform accurate pantomime on verbal command as in the current study depends on the semantic memory, notably instantiated within temporal areas (Kellenbach et al. 2003; Lewis 2006; Frey 2007; Canessa et al. 2008; Goldenberg and Spatt 2009). Our analyses further reveal that left frontal but not parietal lesions correlated with BPO error type. This finding is consistent with previous lesion studies reporting higher rates of BPO in left than right brain damaged patients (Mozaz et al. 1993). We also observed that lesions predicting BPO extended largely to the white matter underlying inferior and middle frontal cortices, including the superior longitudi nal fasciculus (SLF). This finding substantiates the observations by Hanna-Pladdy et al. (2001) of more BPO after left subcortical than cortical damage and evidence that lesions to the SLF induce severe apraxia (Mori et al. 2002; Schmahmann...
and Pandya 2011). Lesions to frontal white matter tracks have been interpreted as inducing apraxia by disconnecting parietal and frontal motor areas (Pramstaller and Marsden 1996).

Several candidate mechanisms have been advanced to explain BPO errors. First, because BPO are considered as errors only when they persist after re-instruction, they could be interpreted as perseveration and be accounted for by mere executive dysfunctions. A neighbor hypothesis by Peigneux and Van der Linden (1999) assumes that BPO could follow from difficulties in inhibiting automatic activations of often used emblematic gestures (e.g., using his hand to represent the handset to signify a phone call). Supporting the hypothesis that BPO are due to a lack of inhibitory control, healthy elderly individuals with weaker inhibitory control also show more BPO errors than young and healthy adults (Peigneux and Van der Linden 1999). However, if BPO errors resulted from perseveration only, they should manifest in both left and right frontal damaged patients and not selectively in left hemispheric patients as in our results (Freedman et al. 1998).

Alternatively, BPO could be committed due to a pathological embodiment of the tool in the patient’s limbs, echoing phenomenon occurring during the rubber hand illusion (Botvinick and Cohen 1998). Kondo et al. (2009) further advanced that the inability to precisely form finger postures to perform the gesture follows from the contamination of the motor command by the information concerning the shape of the objects. Such effect could possibly follow from damage to frontal regions (Arzy et al. 2006; Kondo et al. 2009).

A third candidate mechanism for BPO is advanced by Raymer et al. (1997), who suggests that BPO errors could be linked to difficulties of representing and/or selecting the appropriate object features necessary to produce the correct hand postures used to hold the object. In turn, such deficits would make the patients portraying the object itself instead of imagining it and adapting their gestures accordingly. In this regard, BPO would be in an attempt to circumvent the task difficulty by using limbs as a concrete rather than as an abstract representation of the object (see also Bartolo et al. 2003).

Of note, the overlap between patients’ lesions show that 5 of the 9 patients with only BPO errors are right-brain damaged, suggesting that right hemispheric structures might also play a role in BPO errors (Supplementary Fig. S2b). Because our study includes only unilateral patients, this result might explain the very limited number of patients showing both BPO and CS errors. However, the role of right-hemispheric structures in BPO error does not appear in the VLSM where the severity of the deficits (i.e., the number of BPO) and the patients with both CS and BPO errors are taken into account. Further studies including bilateral patients are required to investigate this question.

Another limitation of the present study is that because it was based on a retrospective approach, only information on CS and BPO error types were available. Previous neuropsychological investigations of pantomime deficits identified several other types of error, which revealed other types of mechanisms involved in pantomime. For instance, investigation of the relationships between pantomime and actual tool use or tool recognition showed that these 2 processes correlated to a certain extent (Bartolo et al. 2003; Rumiati et al. 2004 for discussion), suggesting that pantomime deficits may not solely follow from semantic processing impairments but also from deficits of the output lexicon (Cubelli et al. 2000).

Taken together, our findings reveal that pantomime is subserved by a distributed, left-lateralized, frontoparietal network and that lesions to subparts of this network induce distinct error types. Furthermore, the results point out that the post lesion delay and the etiology of the brain damage might be important to consider in the study of apraxia in brain-damaged patients.

**Supplementary Material**

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/.

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**References**


