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No Language-Specific Activation during Linguistic Processing of Observed Actions

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Background. It has been suggested that cortical neural systems for language evolved from motor cortical systems, in particular from those fronto-parietal systems responding also to action observation. While previous studies have shown shared cortical systems for action – or action observation - and language, they did not address the question of whether linguistic processing of visual stimuli occurs only within a subset of fronto-parietal areas responding to action observation. If this is true, the hypothesis that language evolved from fronto-parietal systems matching action execution and action observation would be strongly reinforced. Methodology/Principal Findings. We used functional magnetic resonance imaging (fMRI) while subjects watched video stimuli of hand-object-interactions and control photo stimuli of the objects and performed linguistic (conceptual and phonological), and perceptual tasks. Since stimuli were identical for linguistic and perceptual tasks, differential activations had to be related to task demands. The results revealed that the linguistic tasks activated left inferior frontal areas that were subsets of a large bilateral fronto-parietal network activated during action perception. Not a single cortical area demonstrated exclusive – or even simply higher - activation for the linguistic tasks compared to the action perception task. Conclusions. These results show that linguistic tasks do not only share common neural representations but essentially activate a subset of the action observation network if identical stimuli are used. Our findings strongly support the evolutionary hypothesis that fronto-parietal systems matching action execution and observation were co-opted for language, a process known as exaptation.

INTRODUCTION

Since the initial observations on macaque mirror neurons - cells that fire while the monkey performs goal-directed actions and also while the animal observes somebody else’s actions [1] - it has been speculated that these cells may have played a role in the evolution of language [2–4]. The theoretical arguments were substantially two: first, mirror neurons were originally discovered in a macaque brain area (area F5) that seems the homolog of human Broca’s area, a major language center; second, mirror neurons seem to facilitate the parity between the sender and the receiver of a message, a parity that establishes what counts in communication [5,6]. Recently, several labs have investigated shared neural systems between language and motor behavior in general, and language and premotor areas responding to action observation (thus, having mirroring properties) in particular. Taken together, the previous studies have demonstrated shared neural mechanisms - in the form of both activation maps [7–9] and modulation of neural excitability [10–13] - between the domain of language and of motor behavior in general, and action observation in particular. The extent to which neural systems for linguistic processing of visual stimuli is independent from the fronto-parietal mirror neuron system, however, has not been experimentally investigated so far (a completely different issue is obviously related to speech perception and superior temporal cortex: this issue is not investigated here). The experimental conditions of previous studies differed widely (e.g., motor tasks or action observation tasks on one side, and reading words or sentences on the other side), thus making the interpretation of differential activations between language and action (or action observation) quite difficult.

The present study adopted a design in which the experimental stimuli are identical, while task instructions differ, tapping either on linguistic functions or on action perception. By using such design, we believe we are in a position to test the extent to which linguistic processing of visual stimuli concerning actions and objects and human fronto-parietal areas responding to action

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observation overlap or differ. This question seems to us relevant to the hypothesis that mirror neurons played a key role in language evolution. Although evolutionary hypotheses cannot fully be demonstrated in the laboratory, we propose that the mirror neuron hypothesis of language evolution makes a relatively simple, and eminently tractable, prediction. If mirror neurons were initially selected for action observation (and presumably its understanding) and subsequently co-opted for language, - a process also known as exaptation [14] - one would expect that while processing identical visual stimuli, a linguistic task should activate a subset of or even all the areas activated by an action perception task, while no additional areas should be activated by the linguistic task. If additional areas are activated by the linguistic tasks, these additional areas should presumably be exclusively linguistic in nature and may not have evolved from mirror neurons.

METHODS

14 healthy right handed subjects, all of them native english speakers, were investigated (age 25.1±2.6 years, 5 men). The study was approved by the UCLA Institutional Review Board and all subjects gave written informed consent for participation in this study.

Task and stimuli

30 manipulable objects of everyday life were presented either as a picture or in a video showing also a hand manipulating the object in a typical way (e.g. photo of a bell – video of a hand ringing a bell). These two sets of stimuli (video, picture) were crossed with three tasks (see Figure 1) that the subjects were asked to perform:

Perceptual task: when videos were presented, subjects were asked to respond whether all five fingers of the hand manipulating the object touched it (Vd-Perc); when pictures were presented, subjects were asked to respond whether any part of the object was of black color (Pict-Perc).

Conceptual task: while watching videos (Vd-conc) and pictures (Pict-Conc), subjects were asked to respond whether the object presented was a tool (as typically used for craftsmanship).

Phonological task: while watching videos (Vd-Phon) and pictures (Pict-Phon), subjects were asked to respond whether the object’s name started with “/s/” or not.

An overview on the experimental design is given in Fig. 1. Subjects were asked to respond by using the index and middle finger of the right hand to press keys of a MRI-compatible response device. For all tasks, in 30% of the stimuli the correct answer was yes.

Prior to the brain imaging experiment a larger set of stimuli was shown to a separate group of 10 healthy native English speaking subjects. Participants in this pilot behavioral study were asked to evaluate whether the objects presented were tools typically used for craftsmanship and which word would best describe the object. All 10 subjects of this pilot study agreed on the object’s name and whether the object was a tool or not, for the 30 stimuli chosen for the brain imaging imaging experiment.

fMRI study

The fMRI study comprised 6 blocks for each task which were presented in counterbalanced order in a pseudorandom design. The duration of each block was 24 s. Intermixed baseline blocks lasting 20 s involved fixation of a central crosshair (rest condition). Prior to the task blocks, a short sentence presented for 1s indicated the type of task which was used in the following block, as picture and video stimuli were identical across tasks.

Functional magnetic resonance imaging (fMRI) based on the blood oxygenation level-dependent (BOLD) contrast was performed using a 3T Siemens Allegra scanner housed in the

Figure 1. Schematic overview on the experimental design. Identical photo and video stimuli were employed across perceptual tasks (action observation/object perception), conceptual and phonological tasks. Thus differential functional imaging activations were not attributable to stimulus differences.

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Subsequently the normalized data were smoothed using a Gaussian threshold for all contrasts were set to p-0.05, uncorrected. Only clusters with at least 20 adjacent voxels are reported.

Data Analysis
The fMRI Data were analysed using Statistical Parametric Mapping software (SPM2, http://www.fil.ion.ucl.ac.uk/spm/, London, UK). The dummy scans were discarded. The remaining scans were realigned and spatially normalized to standard stereotaxic space using the EPI-template of the Montreal Neurological Institute (MNI). The voxel size was 3×3×3 mm. Subsequently the normalized data were smoothed using a Gaussian kernel = 8×8×8 mm in order to improve the signal-to-noise ratio.

For the following parameter estimation, an appropriate design matrix was specified using a box-car function convolved with the hemodynamic response function. Data were high-pass filtered (cutoff period 128 s) to remove low-frequency signal drifts. The voxel-by-voxel parameter estimation for the smoothed data was carried out according to the general linear model. First, single contrasts for each task and comparison across tasks, as outlined in detail below, were computed on the single subject level. These contrast images were the basis of group fMRI activations using the functional relationship between action observation system and the objects. The fMRI data were analyzed as follows to investigate the functional relationship between action observation system and language areas:

Areas responding to action observation were revealed by the contrast Vd-Perc - rest. To account for the decision making process which was involved in the Vd-Perc condition, we also contrasted the activations during Vd-Perc with the perceptual object (Pict-Perc) condition (Vd-Perc – Pict-Perc). The conceptual and phonological networks involved in the linguistic tasks were shown in the conditions (Pict-Conc – rest) and (Pict-Phon – rest). The comparison of linguistic networks with the action observation network, was carried out in two steps: the extent of the phonological and the conceptual network relative to the action observation network was tested by a masking procedure, where the activations revealed by the contrasts (Pict-Phon – rest) and (Pict-Conc – rest) were masked exclusively by the activations revealed by the contrast Vd-Perc. The hypothesis that linguistic processing of observed actions has evolved from the fronto-parietal action observation system predicts that there would be no activated cluster for this contrast, i.e. no part of the conceptual or phonological network tested here extended beyond the action observation system. Further contrasts involving conditions with identical stimuli assessed the degree of activation between the action observation system and linguistic networks: (Vd-Perc – (Vd-Conc+Vd-Phon)) and (Vd-Conc+Vd-Phon) – Vd-Perc. These contrasts were designed for a quantitative comparison of activity in areas involved both during linguistic processing and action perception tasks.

RESULTS
Task performance
The average rate of correct answers across all tasks was 84.26±1.22%. Correctness rates for the six different tasks were as following: Pict-Perc: 84.7±2.9% (S.E.), Pict-Phon: 85.6±2.1%, Pict-Conc: 82.1±3.7%, Vd-Perc: 84.7±2.2%, Vd-Phon: 85.9±1.9%, Vd-Conc: 82.4±2.7% (Fig. 2). This rate of correct answers may be due to two factors. First, in the brain imaging experiment we used a forced-choice task, thus some of the errors were probably due to somewhat speeded responses. Second, some stimuli were also perceptually challenging; for example, only a small part of the object was black or it was not so easy to determine if all fingers or just four fingers held the object.

Pairwise comparisons of correct responses using Student’s t-test (corrected for multiple comparisons) revealed no significant differences between the conditions.

Average reaction time across conditions was 1284.9±31.4 ms. Reaction times for individual conditions relative to stimulus onset ranged between 1096.2±42.4 ms and 1546.2±74.2 ms [Pict-Perc: 1103.0±68.6 ms, Pict-Phon: 1193.6±44.3 ms, Pict-Conc: 1096.2±42.4 ms, Vd-Perc: 1546.2±74.2 ms, Vd-Phon 1395.5±52.7 ms, Vd-Conc: 1375.0±55.4 ms]. Pairwise comparisons of reaction times revealed significantly higher reaction times for the action observation task compared to all other tasks, for the two other tasks involving video stimuli (Vd-Phon and Vd-Conc) compared to the three tasks involving photo stimuli and for Pict-Phon compared to Pict-Conc. These differences make sense. Indeed, the perceptual video task could be ‘solved’ only at the time point in the video when the hand touched the object. In contrast, the information necessary to perform the remaining tasks was present since the very first frame of the video.

Functional imaging results
As expected, the contrast Vd-Perc vs rest revealed signal increases bilaterally in extrastriate visual regions, inferior and superior parietal regions and extensive bihemispheric frontal activations comprising premotor, inferior frontal and prefrontal (dorsolateral and ventrolateral prefrontal cortex) areas (Fig. 3a, table 1). This network is in good accordance with the bilateral network described by previous fMRI studies investigating action observation [7,15–18]. The contrast Vd-Perc - Pict-Perc, which should subtract activity due to object perception and decision making, demonstrated a very similar network of bilateral parietal, premotor and prefrontal areas (Fig 3a).

The contrast Pict-Phon vs rest revealed mainly left frontal regions including anterior inferior frontal and adjacent middle frontal gyrus, ventral premotor cortex and dorsolateral prefrontal cortex, and a left hemispheric activation of the supramarginal gyrus (Fig. 3b, table 2). The conceptual task (Pict-Conc vs rest) likewise activated a predominantly left hemispheric network including a cluster within left posterior middle frontal gyrus extending into dorsal inferior frontal gyrus and ventral premotor cortex and a parietal cluster mainly covering the angular gyrus (Fig. 3b, table 3). These areas are in good accordance with core regions described in a recent metaanalysis of fMRI studies for phonological and conceptual processes [19]. There were no significant temporal activations at the chosen threshold for both linguistic tasks, a result which is probably related to the fact that the experimental design required photos of object stimuli instead of written text or speech as basis for the linguistic tasks.

The second step of the data analysis aimed at systematic comparison of cortical networks involved in action observation and in conceptual/phonological processing in light of the...
hypothesis of a common evolutionary process of language networks and the mirror neuron system. Exclusive masking of the activations for the phonological network (Pict-Phon - rest) with the activation map of the action perception task (Vd-Perc - rest) did not reveal any remaining activation clusters. The corresponding masking analysis for the conceptual network (Pict-Conc – rest) revealed the same result. These results demonstrate that there were no cortical regions exclusively activated either during the conceptual task or the phonological task which were not part of the network activated by action perception.

Further statistical comparisons were related to the degree of activation during phonological or conceptual processing compared to action perception. For this comparison, the three conditions involving identical video stimuli were employed (Vd-Perc, Vd-Phon and Vd-Conc). The comparative analyses of the fMRI activations during these tasks ((Vd-Conc+Vd-Phon) vs. Vd-Perc, Vd-Conc vs Vd-Perc, Vd-Phon vs Vd-Perc) revealed no significantly higher fMRI activation for the conceptual or the phonological task than for the action perception task.

The reverse contrast (Vd-Perc vs (Vd-Conc+Vd-Phon)) showed a widespread bilateral parietal and mainly right hemispheric fronto-parietal network of regions exhibiting higher fMRI activation during action perception than linguistic analysis, given identical video stimuli (Fig. 4, table 4). There were only small clusters in the left frontal lobe showing higher activation for action perception than for conceptual or phonological processing. Taken together with the previous analyses this result indicates a similar level of activation of the left inferior frontal cortex during action perception and linguistic tasks.

DISCUSSION

The experiment reported here adopted a design that required subjects to process identical visual stimuli while performing different tasks: an action perception task, and two ‘linguistic’ tasks, a phonological task and a conceptual task. With this experimental design, it is possible to test whether the overt linguistic processing of observed object-oriented action recruits cortical areas not engaged by action perception, and/or activates fronto-parietal action perception areas to a higher degree. Both results would support the hypothesis of some independence of linguistic processing of visual stimuli with respect to fronto-parietal areas concerned with action perception. However, we did not find...
any area specifically activated during the two linguistic tasks, and we did not find any area with higher activity during the linguistic tasks. We argue that these results are more readily compatible with the hypothesis that language—as far as linguistic processing of visual stimuli is concerned, at the very least—evolved by co-opting the hypothesis that language—as far as linguistic processing of tasks. We argue that these results are more readily compatible with any area specifically activated during the two linguistic tasks, and dissimilar with possible automatic linguistic processing, but dissimilar with regard to action perception itself. Thus, the deflationary explanation that invokes automatic linguistic processing in left inferior frontal cortex in all tasks does not easily account for all experimental results presented here. Furthermore, a virtual lesion study using repetitive TMS has shown that a transient disruption of neural activity in the left (and right) inferior frontal cortex results in imitation deficits, but not in more general visuo-motor deficits [22]. This result can hardly be reconciled with a purely linguistic property of left inferior frontal cortex.

It could also be argued that the increased signal in fronto-parietal areas is only due to the increased attentional demands of the action perception task, given the increased RT for this task. It should be noted, however, that the increase in reaction time for this task was very small in relation to the overall duration of each task block during scanning. Therefore, we consider it unlikely that the difference in reaction time across tasks was of substantial influence on the fMRI activations. Moreover, the “attention”

Table 1. Peak voxel coordinates in MNI space and z-values for the fMRI contrasts revealing the action observation network.

| Video Perception | region | BA | x   | y   | z   | Z-score |
|------------------|--------|----|-----|-----|-----|--------|
|                  | right dorsal premotor | 6   | 30  | 8   | 62  | 4.49   |
|                  | right ventral premotor | 6/9 | 44  | 8   | 32  | 4.5   |
|                  | right inferior frontal gyrus, pars opercularis | 44  | 46  | 12  | 22  | 4.49   |
|                  | right inferior frontal gyrus, pars orbitalis | 47  | 34  | 26  | –4  | 3.29   |
|                  | right dorsolateral prefrontal cortex | 46  | 52  | 40  | 15  | 4.95   |
|                  | right ventrolateral prefrontal cortex | 10  | 44  | 58  | 10  | 3.22   |
|                  | left dorsal premotor | 6   | 28  | –4  | 56  | 4.32   |
|                  | left inferior frontal junction | 9   | 38  | 18  | 30  | 4.45   |
|                  | left inferior frontal gyrus, pars opercularis | 44  | 46  | 6   | 20  | 4.68   |
|                  | left inferior frontal gyrus, pars triangularis | 45  | 60  | 20  | 16  | 3.52   |
|                  | left inferior frontal gyrus, pars orbitalis | 47  | 47  | 20  | 16  | 3.55   |
|                  | left pre-SMA | 6   | 4   | 16  | 52  | 3.59   |
|                  | right IPS | 7   | 26  | –58 | 50  | 5.21   |
|                  | right inferior parietal | 39  | 32  | –68 | 40  | 5.06   |
|                  | left precuneus | 7   | 30  | –46 | 40  | 5.38   |
|                  | left anterior IPS | 7   | 26  | –60 | 44  | 4.66   |
|                  | right fusiform gyrus | 20  | 40  | –44 | 20  | 4.85   |
|                  | right occipital lobe | 18  | 10  | –98 | 16  | 5.58   |
|                  | right occipito-temporal junction | 39  | 44  | –72 | 10  | 5.21   |
|                  | left occipital lobe | 19  | –50 | –58 | 14  | 5.02   |
|                  |               | 17  | 10  | –90 | 0   | 4.94   |
|                  | left cerebellum, lobule V, VI and crus I | –12 | –74 | –50 | 52.7 |
|                  | right cerebellum, lobule V, VI and crus I | 12  | –78 | –50  | 46.3 |
|                  | right thalamus | 26  | –30 | 0   | 4.07 |
|                  | left thalamus | –16 | –24 | 16  | 4.01 |

Table 2. Peak voxel coordinates in MNI space and z-values for the fMRI contrasts revealing the phonological network, as tested in the present study.

| Photo Phonological | region | BA | x   | y   | z   | Z-score |
|--------------------|--------|----|-----|-----|-----|--------|
|                    | right ventrolateral prefrontal cortex | 11  | 42  | 54  | –14 | 3.81   |
|                    | right pre-SMA | 6   | 4   | 16  | 52  | 3.23   |
|                    | left ventral premotor cortex | 6/9 | –46 | 6   | 34  | 4.36   |
|                    | left inferior frontal gyrus/middle frontal gyrus | 45/46 | –48 | 46  | 8   | 3.94   |
|                    | left middle frontal gyrus | 9   | –48 | 28  | 38  | 3.96   |
|                    | left pre-SMA | 6   | 6   | 38  | 44  | 3.75   |
|                    | left inferior frontal gyrus, pars orbitalis | 47  | 32  | 22  | –2  | 3.46   |
|                    | left supramarginal gyrus | 40  | –50 | –36 | 50  | 4.06   |
|                    | left fusiform gyrus | 37  | –52 | –64  | 16  | 4.85   |
|                    | right fusiform gyrus | 20  | 34  | –44 | –22 | 4.58   |
|                    | right occipital lobe | 18  | 14  | –98 | 12  | 4.42   |
|                    | left occipital lobe | 18  | –32 | –88 | 18  | 4     |

Table 3. Peak voxel coordinates in MNI space and z-values for the fMRI contrasts revealing the conceptual network, as tested in the present study.

| Photo conceptual | region | BA | x   | y   | z   | Z-score |
|------------------|--------|----|-----|-----|-----|--------|
|                   | right ventrolateral prefrontal cortex | 11  | 42  | 54  | –14 | 3.81   |
|                   | left sensorimotor cortex | 1/4 | –50 | 28  | 50  | 3.08   |
|                   | left inferior frontal junction | 9   | –50 | 14  | 34  | 4.11   |
|                   | left ventral premotor cortex | 6   | –42 | 0   | 38  | 3.38   |
|                   | left superior frontal gyrus | 8   | –4  | 42  | 58  | 3.8   |
|                   | left anterior IPS | 7   | –28 | –50 | 42  | 3.66   |
|                   | left angular gyrus | 39  | –28 | –68 | 36  | 3.48   |
|                   | left fusiform | 37  | –50 | –64  | 12  | 4.88   |
|                   | left occipital | 18  | –32 | –86 | 12  | 4.62   |
|                   | right occipital | 18  | 16  | –102| 16  | 4.64   |
argument cannot account for the lack of increased signal in the left inferior frontal cortex during action perception, compared to the linguistic tasks. The selectivity of the effect argues against a non specific attention effect.

Our design also allowed us to compare activity during action perception and during perception of static pictures that comprised all the visual elements of the action stimuli. Thus, this comparison reveals brain activity that is quite specific to action observation, rather than to the complex visual elements that invariably go together with observed actions. This comparison in our experiment shows robust bilateral activation in fronto-parietal areas, revealing that this large network is indeed specifically concerned with action perception. Thus, the result of our specific contrast support the ‘mirror neuron’ interpretation of the vast number of previously published papers showing similar fronto-parietal activations in a variety of experimental conditions [7,18,23–26].

Fronto-parietal areas concerned with action perception are bilateral, whereas our linguistic tasks recruited exclusively left hemisphere areas. This shift from bilateral activity for action perception to a predominantly left lateralized language system may have been favored by a lateralization, in humans, of ‘mirroring’ responses to action sounds, as shown by single pulse TMS [27] and fMRI [28].

To conclude, when visual stimuli concerning object-oriented actions are processed perceptually, they activate a large bilateral fronto-parietal network. When the same stimuli are processed linguistically, they activate only a subset of this network and no additional areas. This pattern of activity supports the evolutionary hypothesis that neural mechanisms for language in humans co-opted phylogenetically older fronto-parietal neurons concerned with action perception, such as mirror neurons in macaques.

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Author Contributions

Conceived and designed the experiments: MI IM. Performed the experiments: IM. Analyzed the data: IM. Wrote the paper: MI IM.
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