Multisensory Interactions between Auditory and Haptic Object Recognition

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Object manipulation produces characteristic sounds and causes specific haptic sensations that facilitate the recognition of the manipulated object. To identify the neural correlates of audio-haptic binding of object features, healthy volunteers underwent functional magnetic resonance imaging while they matched a target object to a sample object within and across audition and touch. By introducing a delay between the presentation of sample and target stimuli, it was possible to dissociate haptic-to-auditory and auditory-to-haptic matching. We hypothesized that only semantically coherent auditory and haptic object features activate cortical regions that host unified conceptual object representations. The left fusiform gyrus (FG) and posterior superior temporal sulcus (pSTS) showed increased activation during crossmodal matching of semantically congruent but not incongruent object stimuli. In the FG, this effect was found for haptic-to-auditory and auditory-to-haptic matching, whereas the pSTS only displayed a crossmodal matching effect for congruent auditory targets. Auditory and somatosensory association cortices showed increased activity during crossmodal object matching. Based on our previous results, we defined the left FG as a higher order convergence zone for conceptual object knowledge.

Keywords: auditory perception, functional magnetic resonance imaging, multisensory interactions, object recognition, touch perception

Introduction

The visual appearance, haptic properties (including shape and surface), and characteristic sounds associated with the manipulation of an object provide both redundant and complementary features that facilitate object recognition. Several functional magnetic resonance imaging (fMRI) studies have investigated the neural correlates that are implicated in the integration of object-specific sensory input across vision and touch and across vision and audition (for reviews, see Amedi et al. 2005; Beauchamp 2005). For instance, in the ventral visual pathway, the lateral occipital cortex (LO) processes visual and haptic object-related information when individuals grasp or see man-made tools, animal models, or toys (e.g., Amedi et al. 2001, 2002). On the other hand, the posterior superior temporal sulcus (pSTS) and adjacent middle temporal gyrus have been implicated in audio-visual integration of object-specific input across a wide range of living (e.g., voices/faces, animals) or nonliving objects (e.g., tools, weapons, musical instruments; Beauchamp, Argall, et al. 2004; Beauchamp, Lee, et al. 2004; Sestieri et al. 2006).

While previous neuroimaging work on multisensory object recognition has mainly focused on visuo-haptic and audio-visual integration, little is known about the convergence of object processing across audition and touch. Studies using low-level auditory and tactile stimuli (e.g., vibrotactile stimuli and related sounds) have implicated audio-tactile integration mechanisms in the posterior superior temporal gyrus (pSTG) and adjacent pSTS (Foxe et al. 2002; Schurmann et al. 2006; Beauchamp et al. 2008) as well as the anterior insula (Renier et al. 2009). Consistent with these results, data from a recent high-density electroencephalography (EEG) study using a haptic-to-auditory priming paradigm with meaningful object stimuli suggests integration-related neuronal activity in the left lateral temporal cortex (Schneider et al. 2011). In line with these data, we have recently shown in a fMRI study that the left fusiform gyrus (FG) and pSTS preferentially process manipulable object stimuli relative to nonobject control stimuli within both the auditory and the tactile modalities (Kassuba et al. 2011). These data showed that both regions were also activated during actual multisensory audio-haptic object processing (i.e., when the objects were presented simultaneously in both modalities). However, the contribution from either modality within these regions to higher order aspects of object recognition remains unclear.

Here, we employed event-related fMRI to study audio-haptic interactions in higher order object recognition. We opted for a delayed-match-to-sample paradigm in which participants had to match object-related sensory input within and across audition and touch. We manipulated the semantic congruency of sample and target stimuli to identify brain regions that are selectively responsive to semantically congruent crossmodal input. We reasoned that these cortical regions host unified object representations and, thus, subserve the binding of auditory and haptic input that refers to the same object concept (cf. Schneider et al. 2011). We chose familiar real manipulable objects and typical sounds created by a manipulation on them as sample and target stimuli because audio-haptic interactions are especially relevant in the context of object manipulation. Based on our previous results, we defined the left FG and pSTS as candidate regions to host unified audio-haptic object representations. We expected that crossmodal matching effects in these association regions would critically depend on the semantic congruency between sample and target objects.

Materials and Methods

Participants

Twenty-one healthy young volunteers (10 females) participated in the fMRI study. The age of the participants ranged from 21 to 33 years.
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Handedness was tested with the short form of the Edinburgh Handedness Inventory (Oldfield 1971); all participants were consistently right handed with a Laterality Index \( > 0.87 \) (scaling adapted from Annett 1970). All participants had normal or corrected-to-normal vision, normal hearing ability, and normal tactile acuity, and none had a history of psychiatric or neurological disorders (self-report). Each individual gave written informed consent. The study was approved by the local ethics committee (Arztetkammer Hamburg) and in accordance with the Declaration of Helsinki.

**Stimuli**

Stimuli consisted of 24 different objects. These objects were man-made objects, including tools, toys, and musical instruments. All objects were fully graspable with one hand. Each object was associated with one characteristic sound derived from a manipulative action on it (e.g., a rubber duck was connected to its typical squeak). All sounds were cut to clips of 2 s of duration and equated for root-mean-square power using MATLAB (The MathWorks, Natick, MA).

**Experimental Design and Procedure**

We adopted an event-related fMRI paradigm. Examples of the different trials are depicted in Figure 1. Each trial consisted of a sample object stimulus (S1) and a target object stimulus (S2) presented successively. Upon presentation of S2, participants had to indicate whether the object pairs were either semantically congruent (50\%) or incongruent (50\%), that is, representing conceptually the same object or different objects. The object stimuli could be presented either haptically (actively palpating the object) or acoustically (hearing a typical sound of an object), and S1 and S2 were either of the same modality (unimodal) or of the different modalities (crossmodal). This resulted in 2 levels of sensory matching (unimodal and crossmodal), 2 levels of semantic congruency (congruent and incongruent), and 2 levels of sensory matching (unimodal and crossmodal).

S1 and S2 were both presented for 2 s and preceded by a low-pitched (550 Hz) or high-pitched (800 Hz) tone or a double tone, respectively, to announce the sensory modality of the forthcoming stimulus. A single tone lasted for 300 ms and indicated S1, a double tone consisted of 2 short tones of 120 ms each with an intertone interval of 60 ms and indicated S2. After the instruction tone, there was a short pause of 100 ms before the stimulus was presented. The interstimulus and intertrial intervals (i.e., from the offset of a stimulus to the onset of the next indicating tone) varied between 2 and 6 s, jittered in steps of 1 s. Trials were presented in pseudorandomized order so that the same objects would not occur in successive trials and that the S1-S2 sensory modality combination would be repeated maximally once in successive trials. Every object appeared once as S1 and once as S2 in each condition, the combination of S1 and S2 objects in incongruent trials was randomized.

It is important to note that the participants did not know whether the S2 would be presented as an auditory or a haptic stimulus until 400 ms before onset of S2. At this point in time, an instruction tone informed the participants about the modality of S2. This means that all trials with an auditory S1 and all trials with a haptic S1 were identical, respectively, until shortly before the onset of S2. Thus, during the delay period, participants had to prepare for potential processing of both an auditory and a haptic object stimulus. Therefore, it is very likely that S1 was indeed encoded up to the semantic level.

One day before the fMRI experiment, participants were trained to recognize the 24 object stimuli by hearing the related sound and by haptic exploration (without ever seeing the objects). The training was repeated until the object stimuli were identified with an accuracy of 100\% (0–2 repetitions per participant). Then, they were familiarized with the experimental task and trained to haptically explore the objects with an appropriate speed. A short recall of the training immediately before scanning showed that the objects were still identified with an accuracy of 100\% within each modality. Throughout the whole experiment, participants were not able to hear the sound that their haptic exploration produced. In addition, participants were blindfolded during the training as well as during the fMRI experiment.

The event-related fMRI experiment consisted of 192 trials (24 trials per condition), separated into 2 runs of scanning of about 20 min (96 pseudorandomized trials per run). Haptic stimuli were placed on a board which was fixed onto the participants’ waist by a vacuum cushion. Participants were asked to rest their right hand on the board and were instructed to wait for a presented tone. In case of a low-pitched tone, they were asked to make a slight movement to the left, palpate the haptic stimulus, and then to reposition their hand. Participants were trained to use no more than 2 s for hand movements and exploration. The participants’ right upper arm was completely fixed in a slightly elevated position and supported by cushions to minimize arm movements during haptic exploration. In case of a high-pitched tone, they were asked to wait for the auditory presentation of the following object. Upon the presentation of S2, participants were instructed to indicate by button press as fast and as accurately as possible whether both objects were the same or different. Responses were given by the middle and the index finger of the left hand, using an MR-compatible button device. The finger-response assignment was counterbalanced across participants, and the responses were recorded with Presentation software (Neurobehavioral Systems, Albany, CA).

Auditory stimuli were presented using Presentation running on a Windows XP professional SP3 PC. During MR image acquisition, they were presented at approximately 75 dB using an MR-compatible headphone system with gradient noise suppression (MR confon GmbH, Magdeburg, Germany). Sound volume was adjusted individually before the beginning of the first experimental run. Participants reported no problems with hearing the sound stimuli during MR image acquisition.

**Magnetic Resonance Imaging**

MRI measurements were carried out on a 3-T MRI scanner with a 12-channel head coil (TRIO, Siemens, Erlangen, Germany). To measure task-related changes in the blood oxygen level-dependent (BOLD) signal, we acquired 38 transversal slices (216 mm field of view [FOV], 72 \( \times \) 72 matrix, 3 mm thickness, no spacing) covering the whole brain using a fast gradient echo T2*-weighted echo planar imaging sequence (repetition time [TR] 2480 ms, echo time [TE] 30 ms, 80° flip angle). Two functional runs with 505 \( \pm \) 1 brain volumes each were acquired with each volume consisting of 38 slices. The exact number of volumes per run depended on the individual randomization of the jittered time...
interaction between our experimental factors. Thus, according to the 2 Subject factor (modeling the participants' constants) and the in-
75% of the length of the trials) was noninformative in relation to the S2-
and the delay period until the S2 instruction tone (i.e., on average about 
modeling the whole trials as compound events as the processing of S1 
by a high-pass filter with a cutoff period of 128 s. We refrained from 
first derivative. Low-frequency drifts in the BOLD signal were removed 
for the onset of S2 within each trial condition. Only correct trials 
(fxements explained by the onset of S1) and the delay period between S1 
S2. For brain activity related to auditory and haptic sample encoding 
auditory and haptic object-specific processing in our previous study 
because we had not included null-events in our design. A description of 
auditory and haptic processing of the object stimuli used in this study 
in relation to fixation baseline can be found in Kassuha et al. (2011). 
In this recent study, we have shown that the left pSTS and FG are 
both consistently stronger activated during the processing of auditory 
as well as haptic object stimuli relative to nonobject control stimuli 
(Kassuha et al. 2011). Therefore, we report voxelwise familywise error 
rate (FWE) corrected *p*-values as obtained from small volume correction 
in these regions of interest (P < 0.05). Correction was based on spheres 
centered over peak coordinates obtained from the conjunction of 
auditory and haptic object-specific processing in our previous study 
(Kassuha et al. 2011). Based on the size of the regions identified in the 
previous study, the radius of the sphere was set to 8 mm: The FG was 
corrected using a sphere centered at x = −39, y = −45, z = −18 and the 
pSTS was corrected using a sphere centered at x = −51, y = −57, z = 12 
for all other voxels in the brain, voxelwise whole-brain FWE 
correction was applied (P < 0.05). Anatomical labeling was done 
by using the probabilistic stereotactic cytoarchitectonic atlas imple-
mented in the Anatomy Toolbox version 1.7 (Eickhoff et al. 2005) and 
adjusted according to anatomical landmarks of the average structural 
T1-weighted image of all participants. The percent signal change 
plotted for visualization of the results was extracted using the rfxplot 
toolbox (Gläscher 2009).

Results

Task Performance
All participants solved the task with high accuracy. Mean frequency of correct responses over all conditions was 96.97 ± 
2.63% with no significant differences between conditions (P > 0.05). Response latencies are displayed in Figure 2. Participants 
overall responded faster to auditory than haptic S2 (main effect of 
S2-Modality: F_{1,20} = 238.89, P < 0.001). Both, differences in 
RTs related to sensory matching as well as related to semantic 
congruency, were modulated by the modality in which S2 
was processed (S2-Modality × Sensory-Matching interaction: 
F_{1,20} = 14.21, P < 0.01; S2-Modality × Congruency interaction: 
F_{1,20} = 15.53, P < 0.001).

We examined these interactions further by exploring the 
effects separately for each S2 modality. For auditory S2, faster 
responses were noted for unimodal as compared to crossmodal 
matching (main effect of Sensory-Matching for auditory S2: F_{1,20} 
= 60.96, P < 0.001), and this effect was larger for congruent than 
incongruent stimulus pairs (Sensory-Matching × Congruency 
interaction for auditory S2: F_{1,20} = 11.66, P < 0.01). In trials with
haptic S2, participants responded consistently faster to congruent than incongruent object pairs (main effect of Congruency for haptic S2: \( F_{1,20} = 39.58, P < 0.001 \)). In addition, even though not significant when corrected for post hoc testing, participants tended to react faster to unimodal than crossmodal matching, but only when S1 and S2 were semantically congruent (\( t_{20} = 2.39, P = 0.027 \), uncorrected; see Fig. 2). Accordingly, there was a significant Sensory-Matching by Congruency interaction for haptic S2 (\( F_{1,40} = 4.65, P < 0.05 \)).

**Functional MRI**

Crossmodal Matching by Semantic Congruency Interactions

As outlined in the introduction, we defined the left FG and pSTS as candidate regions where semantically congruent object features are unified across audition and touch. According to our hypothesis, we expected that the left FG and pSTS would show an increase in activation when semantically congruent auditory and haptic information had to be matched relative to unimodal matching. Moreover, we expected this crossmodal matching effect to occur only when both objects were semantically congruent, as only congruent information can be unified to a coherent representation. Analysis of task-related activation in response to S2 revealed an involvement of the left FG and the left pSTS in crossmodal matching of semantically congruent object features.

In both cortical regions, we found the expected interaction between crossmodal matching and semantic congruency (crossmodal > unimodal × congruent > incongruent), but only in the left FG, this interaction was independent of the direction of crossmodal matching (see Table 1 and Figs 3 and 4).

In the left FG, regional activity increased during crossmodal matching of auditory and haptic object features, but only if auditory and haptic stimuli referred to the same object. Thus, the left FG showed a stronger BOLD response in semantically congruent but not in semantically incongruent conditions if an auditory S2 had to be matched with a haptic S1 (HA trials) or vice versa (AH trials) as compared to when S2 and S1 were presented within the same modality (AA and HH trials, respectively; peak \( x, y, z \) in millimeters: \(-39, -13, -23\); \( t_{140} = 3.93, P < 0.01, \) corrected). Most importantly, a direct comparison of the crossmodal matching effect within congruent and incongruent trials (i.e., Sensory-Matching × Congruency interaction) revealed significantly greater BOLD changes in the left FG in congruent trials. This was indicated by a conjunction analysis which considered the Sensory-Matching by Congruency interaction term for auditory and haptic S2 (see Table 1). A possible concern may be that the interaction pattern was driven by the fact that in the unimodal congruent condition, the 2 objects were identical, whereas they were different (although conceptually matching) in the crossmodal congruent trials. Thus, a crossmodal matching effect that only exists in congruent but not incongruent trials may not be surprising. However, the FG interaction patterns were not only defined by a relative increase during crossmodal as compared to unimodal congruent matching but in addition by a relative increase during crossmodal congruent as compared to incongruent matching (see Fig. 3). Planned \( t \)-contrasts confirmed the observed interaction patterns (auditory S2: HA > AH: \( t_{140} = 1.89 \), haptic S2: AH > HA: \( t_{140} = 1.42 \); both \( P < 0.05 \), uncorrected).

In contrast to the left FG, the left pSTS was only involved in matching an auditory S2 to a haptic S1 (HA trials) but not vice versa (AH trials; see Fig. 4). The left pSTS and adjacent pSTG showed a stronger activation when an auditory S2 had to be matched to a congruent haptic S1 as compared to unimodal auditory–auditory matching (\(-54, -52, 13\); \( t_{140} = 4.08, \) \( P < 0.01 \), corrected). Critically, the increased activation during crossmodal as opposed to unimodal matching of auditory S2 was significantly greater in semantically congruent relative to unimodal matching (see Table 1 and Figs 3 and 4).

![Figure 2](http://cercor.oxfordjournals.org/) Mean RTs for auditory and haptic S2 for all 4 matching conditions (unimodal/crossmodal × congruent/incongruent) with error bars indicating the standard error of the mean. RTs were recorded from S2 onset onward. Responses to auditory S2 were significantly faster when they were matched with an auditory S1 (unimodal) compared to a haptic S1 (crossmodal). On the other hand, participants responded faster to haptic S2 if they referred to the same object concept as the S1 (congruent) than when they did not (incongruent). In general, it took participants longer to match congruent stimuli across than within modalities, and this difference was less (auditory S2) or not at all (haptic S2) pronounced for incongruent stimuli (sensory matching by semantic congruency interaction). Sample–target (S1–S2) conditions: A = auditory, H = haptic. *\( P < 0.05 \), **\( P < 0.01 \), ***\( P < 0.001 \), Bonferroni corrected.

![Table 1](http://cercor.oxfordjournals.org/) Crossmodal matching by semantic congruency interaction (crossmodal > unimodal × congruent > incongruent)

<table>
<thead>
<tr>
<th>Region</th>
<th>( x )</th>
<th>( y )</th>
<th>( z )</th>
<th>( t_{\text{peak}} )</th>
<th>( P_{\text{corrected}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auditory and haptic S2 conditions: ([HA &gt; AA] &gt; [HA &gt; AA]) ( \cap ) ([AHc &gt; HHc] &gt; [AH &gt; HH])</td>
<td>LM FG</td>
<td>-39</td>
<td>-37</td>
<td>-20</td>
<td>3.08 0.026</td>
</tr>
<tr>
<td>Only auditory S2 conditions: ([HA &gt; AA] &gt; [HA &gt; AA])</td>
<td>L pSTS</td>
<td>-48</td>
<td>-52</td>
<td>7</td>
<td>3.16 0.021</td>
</tr>
<tr>
<td>Only haptic S2 conditions: ([AHc &gt; HHc] &gt; [AH &gt; HH])</td>
<td>L anterior cingulum</td>
<td>-6</td>
<td>11</td>
<td>52</td>
<td>5.44 0.004</td>
</tr>
<tr>
<td>Only haptic S2 conditions: ([AHc &gt; HHc] &gt; [AH &gt; HH])</td>
<td>L LO</td>
<td>-48</td>
<td>-64</td>
<td>-5</td>
<td>5.00 0.022</td>
</tr>
</tbody>
</table>

Note: Coordinates are denoted by \( x, y, z \) in millimeters (Montreal Neurological Institute [MNI] space) and indicate the peak voxel. Strength of activation is expressed in \( t \) and \( P \) values corrected for the whole brain (df = 140). All contrasts are masked inclusively with the contrast of the respective condition of interest versus baseline \( P < 0.05 \), uncorrected. There were no significant results for the reversed interaction contrasts (crossmodal > unimodal × incongruent > congruent) that surpassed a correction for multiple comparisons. L = left, R = right. Sample–target (S1–S2) conditions: A = auditory, H = haptic, c = congruent, i = incongruent.

*Corrected for the left FG.

*Corrected for the left pSTS.
congruent matching as compared to incongruent matching of auditory S2. This interaction effect did not occur for the analog comparisons when haptic S2 were matched (blue bars). Moreover, activation was relatively decreased during unimodal congruent matching and relatively increased during crossmodal congruent matching as compared to incongruent matching. L = left. Sample-target (S1-S2) conditions: A = auditory, H = haptic, c = congruent, i = incongruent. *P < 0.05, corrected.

In fact, the left LO showed rather a decrease in the BOLD response in all conditions in which an auditory S2 was presented haptically (left LO: \( t_{110} = 8.96 \), right LO: \( t_{110} = 7.00 \); both \( P < 0.001 \), corrected). In contrast, the activation in the LO did not differ between unimodal and crossmodal matching if S1 and S2 were semantically incongruent. Accordingly, the crossmodal matching effect (crossmodal > unimodal) in the left LO was significantly greater for semantically congruent than incongruent stimulus pairs (Sensory-Matching × Congruency within haptic S2; see Table 1 and Fig. 5). However, in contrast to the interaction patterns found in the FG and pSTS, the activation in the LO did not differ between crossmodal matching of congruent versus incongruent input. The left LO was rather strongly activated whenever a haptic S2 was processed and, thereby, less activated when the same haptic input was matched (see Fig. 5). For auditory S2, on the other hand, there was no interaction effect at all (\( P > 0.5 \), uncorrected). In fact, the left LO showed rather a decrease in the BOLD response in all conditions in which an auditory S2 was processed (see Fig. 5).
The reversed interaction contrast (crossmodal > unimodal × incongruent > congruent) did not yield any significant results in any of the S2-modality conditions. Taken together, the left FG showed a crossmodal matching effect that was specific to semantically congruent object features and independent of the direction of matching. In contrast, the left pSTS displayed the interaction only for auditory and the left LO only for haptic S2. Most important, this selectivity to congruent crossmodal matching cannot be explained by task difficulty because the RT data indicated that in general, crossmodal incongruent matching was at least as difficult or more difficult than crossmodal congruent matching.

Main Effects of Crossmodal Matching

In addition to the effects modulated by semantic congruency, we also found crossmodal matching effects that were independent of semantic congruency (for a summary, see Table 2). The left inferior frontal gyrus (IFG) and a region in the left posterior FG (pFG; posterior to the region reported above) were both stronger activated during crossmodal than during unimodal matching for both auditory and haptic S2 and independent of semantic congruency. In addition, modality-specific main effects for crossmodal matching were found in secondary sensory cortices and association cortices such as in bilateral pSTG for auditory S2 and bilateral precentral and postcentral gyri as well as the anterior insula for haptic S2. The reversed contrast (unimodal > crossmodal) did not yield any significant results.

Main Effects of Semantic Congruency

Effects of semantic congruency or incongruency that were not modulated by crossmodal matching were only found for haptic S2 (see Table 3). A region in left pFG and the IFG displayed an incongruency effect (i.e., stronger activation for incongruent vs. congruent matching) for haptic S2, independent of S1-modality. A reversed effect (congruent > incongruent) was found in the left angular gyrus, but only when the threshold was lowered to P < 0.001, uncorrected.

The activations found in the left IFG and pFG for the incongruency contrast partially overlapped with the activations found for the main effect of crossmodal matching contrast (see Supplementary Figs S5 and S6). Thus, the 2 regions showed increased BOLD responses in S2-matching conditions in which participants needed longer to react (crossmodal matching for auditory S2, incongruent matching for haptic S2), suggesting that the activation of these regions might have been driven by task difficulty. An additional analysis supported this assumption by showing that the individual trialwise BOLD responses in these regions (but not in the more anterior mid-FG region showing the crossmodal matching by semantic congruency interaction effect) were positively correlated with individual trialwise response latencies (see Supplementary Methods, Results, and Figs S5 and S6).

Discussion

Here, we performed fMRI during a delayed-match-to-sample task to investigate audio-haptic interactions during higher order object recognition. In secondary auditory and somatosensory cortices and association cortices, the BOLD response increased during crossmodal as opposed to unimodal object matching independently of semantic congruency. In contrast, the left FG and pSTS were specifically activated during crossmodal matching of a target (S2) with a congruent sample (S1). The direction of the observed activity patterns suggests complementary functions of these regions: Only the left FG showed a stronger differential activation during both haptic-to-auditory and auditory-to-haptic matching of semantic congruent object features. By contrast, the left pSTS was selectively engaged in matching auditory targets. The present results extend previous findings that the left FG and pSTS are object selective within the auditory and tactile modalities (Kassuba et al. 2011). Together, the results show that auditory and haptic object processing interact at various stages of object recognition and putatively converge into a higher order conceptual object representation hosted by the FG.
coherence in content is a relevant factor for the binding of meaningful information (Laurienti et al. 2004), we speculate that the specific neuronal enhancement for congruent cross-modal matching might imply multisensory binding mechanisms related to a coherence in meaning. The current results extend previous data showing that the FG processes object stimuli across vision and touch (e.g., James et al. 2002; Stevenson et al. 2009), vision and audition (e.g., Adams and Janata 2002; Naumer et al. 2009; Stevenson and James 2009), or all 3 modalities (Binkofski et al. 2004; Kassuba et al. 2011) and suggest that the FG is a higher order convergence zone for multisensory object information.

Recognizing an object by a related sound is arbitrary and depends on prior associations of a given object and the respective sound (Griffiths and Warren 2004). Accordingly, in order to match object information across audition and touch, associative object knowledge needs to be accessed. Together with other studies of semantic processing (Wagner et al. 1998; Price 2000; Wheately et al. 2005; Gold et al. 2006), the current data indicate a contribution of the FG when detailed semantic interpretations are involved such as linking the typical sound, shape, and consistency associated with a given object. In line with this view, learning audio-visual associations between artificial objects results in an integration-related recruitment of the left FG (Naumer et al. 2009). Therefore, the left FG might provide conceptual object representations (Martin 2007) in the form of associative object knowledge that is accessible via different senses.

Given the current data, we can only speculate about how the FG convergence zone might bind information from different senses at the level of single neurons. Whereas the gold standard for multisensory integration advancing the orientation of behavior is neuronal convergence (i.e., synaptic convergence of modality-specific processing onto individual neurons; Meredith 2002), much less is known about the physiological principles underlying higher order processes such as perceptual binding (Stein and Stanford 2008). Theories on semantic memory suggest that multisensory associative object knowledge is implemented as supramodal nodes linking features provided by different senses (Damasio 1989; Mesulam 1998; Meyer and Damasio 2009). Thereby, the neurons within associative areas code a record to reconstruct an approximation of the original perceptual representations. At the level of single neurons, such records might be formed by strengthened synaptic interconnections due to Hebbian learning (Cruikshank and Weinberger 1996) between neurons preferentially connected to specific unisensory areas or lower order convergence areas (Mesulam 1998; Meyer and Damasio 2009). Given the role of the FG in processing meaning (Martin and Chao 2001; Martin 2007), it is tempting to speculate that the FG directly links semantic attributes related to manipulable objects (e.g., graspable shape, nonbiological motion pattern, related action) extracted by lower order convergence zones rather than perceptual features per se.

An alternative account is that the activation of the left FG might be related to visual imagery (Ishai et al. 2000; O’Craven and Kanwisher 2000), used by the participants as strategy to accomplish task performance. Although, visual imagery might have occurred to some degree, 2 reasons render it unlikely that it explains the differential activation pattern in the FG. First, if auditory and haptic object input were translated into visual images, one might expect the visual images to be particularly maintained during the delay period. However, there was no activation in the FG nor in any other visual region during the delay period (see Supplementary Table S2). The fact that in

### Table 2

<table>
<thead>
<tr>
<th>Region</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t_{peak}</th>
<th>P_{uncorrected}</th>
<th>P_{corrected}</th>
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<tbody>
<tr>
<td>Crossmodal &gt; unimodal, auditory and haptic S2 conditions: (AHc &gt; AHi) ∩ (AHi &gt; HHc)</td>
<td>L pFG</td>
<td>-42</td>
<td>-49</td>
<td>-11</td>
<td>3.43</td>
<td>0.013*</td>
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<td></td>
<td>L IFG (pars triangularis)</td>
<td>-45</td>
<td>14</td>
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<td>-28</td>
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<tr>
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<td>-19</td>
<td>4</td>
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<td>0.027</td>
</tr>
<tr>
<td></td>
<td>R superior temporal gyrus</td>
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<td>-22</td>
<td>4</td>
<td>6.03</td>
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<td>R superior temporal gyrus</td>
<td>-51</td>
<td>-10</td>
<td>1</td>
<td>5.92</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>R superior temporal gyrus</td>
<td>-57</td>
<td>-4</td>
<td>-8</td>
<td>4.99</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td>L cerebellum</td>
<td>-33</td>
<td>-46</td>
<td>-29</td>
<td>4.89</td>
<td>0.033</td>
</tr>
</tbody>
</table>

Note: Coordinates are denoted by x, y, z in millimeters (Montreal Neurological Institute [MNI] space) and indicate the peak voxel. Strength of activation is expressed in t and P values corrected for the whole brain (df = 140). All contrasts are masked inclusively with the contrast of the respective condition of interest versus baseline (P < 0.05, uncorrected). There were no significant results for the respective reversed contrasts (unimodal > crossmodal) that surpassed the above used thresholds. L = left, R = right, pFG = postcentral gyrus, AHi = auditory, haptic, c = congruent, i = incongruent.

*Corrected for the left FG.

### Table 3

<table>
<thead>
<tr>
<th>Region</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t_{peak}</th>
<th>P_{uncorrected}</th>
<th>P_{corrected}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congruent &gt; incongruent, haptic S2 conditions: (AHc &gt; AHi) ∩ (HHi &gt; HHc)</td>
<td>L angular gyrus</td>
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<td>-58</td>
<td>37</td>
<td>4.07</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>L IFG (pars opercularis)</td>
<td>-42</td>
<td>-52</td>
<td>-14</td>
<td>3.51</td>
<td>0.019*</td>
</tr>
<tr>
<td></td>
<td>L IFG (pars triangularis)</td>
<td>-42</td>
<td>8</td>
<td>25</td>
<td>4.90</td>
<td>0.032</td>
</tr>
<tr>
<td></td>
<td>R superior medial gyrus</td>
<td>-6</td>
<td>17</td>
<td>46</td>
<td>4.52</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>R IFG (pars triangularis)</td>
<td>51</td>
<td>20</td>
<td>28</td>
<td>4.13</td>
<td>&lt;0.001</td>
</tr>
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<td>R IFG (pars triangularis)</td>
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<td>16</td>
<td>3.45</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>L middle frontal gyrus</td>
<td>42</td>
<td>29</td>
<td>25</td>
<td>3.55</td>
<td>&lt;0.001</td>
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<tr>
<td></td>
<td>L middle OccG/psPS</td>
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<td>-70</td>
<td>37</td>
<td>3.88</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Note: Coordinates are denoted by x, y, z in millimeters (Montreal Neurological Institute [MNI] space) and indicate the peak voxel. Strength of activation is expressed in t and P values corrected for the whole brain (df = 140). All contrasts are masked inclusively with the contrast of the respective condition of interest versus baseline (P < 0.05, uncorrected). Additional clusters at P < 0.001, uncorrected, are listed if they exceed a cluster size ≥15 voxel (P values in parentheses). There were no significant results for the respective contrasts in auditory S2 conditions that surpassed the above used thresholds. L = left, R = right, pFG = postcentral gyrus, AHi = auditory, H = haptic, c = congruent, i = incongruent.

*Corrected for the left FG.
contrast the left IFG showed a sustained activation during the delay period might suggest that participants encoded the samples semantically (e.g., as object names) rather than visually. Second, if the differential activation pattern in the left FG at time point of matching was driven by differences in visual imagery, we would expect to find a similar differential activation pattern in the left LO which had shown a similar object-specific activation pattern as the left FG in a previous study using the visual counterparts of the currently used object stimuli (Kassuba et al. 2011; see also Lacey et al. 2010). The left LO, however, showed no clear selectivity for congruent over incongruent crossmodal matching and was rather deactivated during auditory target conditions.

**Audio-haptic Interactions in the pSTS**

The left pSTS only showed a greater crossmodal matching effect for congruent as opposed to incongruent matching when auditory but not when haptic targets had to be processed. This observation extends our previous findings that the left pSTS is increasingly activated during processing of action-related object sounds as well as during active palpation of the respective manipulable objects as compared to non-object control stimuli (Kassuba et al. 2011). The current results tie in with previous work in nonhuman primates (Kayser et al. 2005) and humans (Foxe et al. 2002; Beauchamp et al. 2008) showing that multisensory interactions between low-level auditory and tactile input in the pSTG sometimes extend into the pSTS, as well as with human EEG data source reconstruction suggesting that haptic-to-auditory object priming effects are located in the left lateral temporal cortex (Schneider et al. 2011). The pSTG and adjacent pSTS have been particularly implicated in higher order auditory functions such as storing auditory knowledge related to objects (Wheeler et al. 2000; James and Gauthier 2003) and making semantic judgments on those auditory object features (Kellenbach et al. 2001; Goldberg et al. 2006). We, therefore, argue that conceptual object representations in left pSTS are primarily auditory in nature but can be activated in a top-down fashion by crossmodal feedback loops.

Previous work has implicated the pSTS in audio-visual integration of meaningful information (for reviews, see Amedi et al. 2005; Doehrmann and Naumer 2007). For instance, Tanabe et al. (2005) found a similar asymmetry of the activation in left pSTS in a delayed-match-to-sample learning task between the matching of visual and auditory stimuli. More posterior parts in left pSTS showed a decrease in activation when learning audio-visual associations when auditory targets were matched to visual samples but not vice versa (see Fig. 5A/B and Table 2 in Tanabe et al. 2005). Similarly, a region in left pSTG and adjacent pSTS/middle temporal gyrus was selectively modulated by auditory but not by visual or action-related conceptual features, suggesting that this region codes higher order acoustic object information contributing to a concept (Kiefer et al. 2008). Together with the current results, these previous data support the assumption that the left pSTS is primarily tuned to auditory object information. It has to be kept in mind that matching of an auditory target with a haptic sample is constantly taking place in daily life since it is usually the manipulation of the object that produces a characteristic sound. Thus, the interactions of auditory and haptic object features in left pSTS might be strongly influenced by the pragmatic use of these manipulable objects in daily life.

**Audio-haptic Interactions in the LO**

The left LO might be the counterpart of the pSTS hosting haptic object representations that are crossmodally modulated by sounds because the left LO displayed the opposite interaction, namely a selective increase in activity during crossmodal matching of a haptic target to a congruent auditory sample. In contrast to the left pSTS, the LO was only activated when processing haptic but not auditory targets, which ties in with previous studies (e.g., Amedi et al. 2001, 2002; Kassuba et al. 2011). This may be explained by the particular involvement of the LO in processing the shape of objects, a feature which is normally derived from visual and haptic but not auditory input (Amedi et al. 2002, 2007). In a recent fMRI-adaptation (fMRI-A) study, Doehrmann et al. (2010) found an enhanced response to auditory object repetitions in contrast to auditory object changes in the left LO, thus, demonstrating that the object-related left LO is able to discriminate auditory object stimuli at least along the dimension “same” versus “different.” Together with the crossmodal matching by semantic congruency interaction for haptic targets found in the current results, we speculate that the LO is able to extract conceptual object information from auditory input to a certain degree without being an auditory region per se (cf. Amedi et al. 2007; Kim and Zatorre 2011).

**Audio-haptic Interactions at Different Hierarchical Stages of Object Processing**

Regardless of semantic congruency, a wide range of regions along the proposed auditory (Ahveninen et al. 2006) and haptic (Reed et al. 2005) “what” pathways displayed an increased activation during crossmodal as compared to unimodal matching of auditory and haptic targets. For instance, crossmodal matching of auditory targets involved the bilateral pSTG, whereas crossmodal matching of haptic targets activated the bilateral anterior insula as well as the bilateral precentral and postcentral gyri. The pSTG, a part of the auditory association cortex (Kaas and Hackett 2000) and the possible homologous area in the caudal auditory belt in monkeys, has previously been shown to integrate low-level auditory and somatosensory information in humans (Foxe et al. 2002; Schurmann et al. 2006) and monkeys (Kayser et al. 2005), respectively. Similarly, auditory and somatosensory “what” information have been shown to converge in the bilateral anterior insula (Renier et al. 2009). Thus, the effects for crossmodal matching independent of semantic congruency found in these regions in the current data might very likely reflect early audio-haptic interactions that are not related to object recognition per se. However, the crossmodal interactions in auditory-related cortices were clearly biased toward auditory object processing, whereas the crossmodal interactions in somatosensory-related cortices were biased toward haptic object processing.

Modality-independent crossmodal matching effects that were not affected by semantic congruency were found in the left IFG and a left pFG region, just posterior to the region where crossmodal matching was modulated by semantic congruency. The same regions were also stronger activated during matching of incongruent as opposed to congruent haptic targets. Similar incongruency effects have been found in the left IFG for audio-visual interactions (Hein et al. 2007; Noppeney et al. 2008). These findings imply that in the context of the current task, the left IFG might have played the role of a
semantic working memory or executive system (Poldrack et al. 1999), possibly involved in semantic conflict monitoring and modulated by task difficulty. In line with this interpretation is also that the left IFG showed a sustained activation during the delay period between sample and target and that its activity was in general positively correlated with individual response latencies (see Supplementary Results, Table S2, and Fig. S5). The left pFG region might similarly be involved in higher order processes and probably converge auditory and haptic information, which is then unified into coherent object representations in the more anterior FG region (Martin 2007). Apart from the main effects for crossmodal matching, a main effect of semantic congruency that was independent of crossmodal matching was only found as a trend for haptic targets in the left angular gyrus. This trend might reflect the involvement of the angular gyrus in semantic retrieval (Noppeney and Price 2003, 2004; Noppeney et al. 2008).

Together, the results suggest multisensory interactions between audition and touch at various stages of object recognition: Object processing in secondary sensory cortices and association cortices is crossmodally modulated independent of semantic congruency, whereas crossmodal interactions in the left FG and the left pSTS are highly dependent on semantic congruency between object pairs, indicating that they unify crossmodal information into a coherent object representation. The left pSTS might host auditory-dominated object concepts that can be crossmodally modulated. Conversely, the left FG might represent a higher order convergence zone for object concepts independent of the modality the object is perceived in.

**Methodological Considerations**

In addition to more basic binding cues like temporal and spatial coherence (Stein and Stanford 2008), the correspondence in content (semantic congruency) is an important factor for the binding of object features provided by different senses into a unified conceptual representation (Laurienti et al. 2004). However, unlike most previous studies comparing the effects of semantic congruency during simultaneous processing of input from different senses (for review, see Dochermann and Naumer 2008), we introduced a temporal delay between the processing of the 2 object stimuli. Consequently, the study focused on the effects of audio-haptic interactions on higher order object recognition rather than basic multisensory integration processes (Stein and Stanford 2008). Our choice was based on the following considerations: simultaneous processing of 2 auditory stimuli potentially reduces their perceptibility and the concurrent haptic exploration of 2 objects would have been hardly possible with one hand. By employing a delayed-match-to-sample paradigm, we kept the perceptual requirements constant across the unimodal and crossmodal conditions. Moreover, the use of a semantic task guaranteed a conceptual processing of both target and sample.

The sequential stimulus presentation employed in the present study resembles fMRI-A or priming paradigms. fMRI-A studies revealed a crossmodal adaptation effect for semantic congruent information in multisensory integration regions such as in the pSTS across audition and vision (van Atteveldt et al. 2010) and in the LO across vision and touch (Tal and Amedi 2009). In the current study, we rather found the opposite effect, namely increased activation during crossmodal congruent matching instead of an adaptation in object-specific audio-haptic regions of interest. Furthermore, we did not find any general habituation effects of the BOLD response during target matching due to the repeated presentation of the same object stimuli throughout the experiment (see Supplementary Results and Fig. S5). There are 2 major differences between the present study and typical fMRI-A paradigms that might account for this difference. First, the stimulus onset asynchronies between sample and target were rather long and might have favored a semantic encoding of the sample. Second, instead of using a task orthogonal to the effect of interest (crossmodal matching) such as a detection task (Dochermann et al. 2010; van Atteveldt et al. 2010) or passive recognition (Tal and Amedi 2009), our task explicitly required a semantic decision on whether the 2 objects matched onto the same concept.

The exact neuronal mechanisms underlying BOLD adaptation are not readily understood and probably vary as a function of the time scale of adaptation, task and stimuli, and brain regions (Griff-Spector et al. 2006). Thus, the task demands in the current study might have overruled general effects of stimulus habituation. In line with this account, the BOLD response during sample encoding but not during target matching in the left pFG and pSTS habituated as a function of how often an object had already been presented throughout the experiment (see Supplementary Figs S4 vs. S3). Similarly in the visuo-haptic domain, for instance, the left anterior intraparietal sulcus showed crossmodal adaptation effects in an fMRI-A paradigm (Tal and Amedi 2009), whereas it showed enhanced responses to crossmodal matching in a delayed-match-to-sample task (Grefkes et al. 2002). Yet, our findings are consistent with the results from crossmodal fMRI-A studies (e.g., Tal and Amedi 2009; van Atteveldt et al. 2010), namely by showing a selectivity of occipitotemporal regions for crossmodal associations of meaningful object information.

Finally, the comparison of conditions with versus without haptic exploration differs in terms of manual activity, which is present during the former but not the latter (Naumer et al. 2010). Therefore, we refrained from directly comparing haptic versus auditory target conditions but rather limited the comparison to modality-specific differential effects across modalities. This enabled us to eliminate motor specific effects related to haptic target processing by the applied contrasts. While we cannot completely rule out that motor activity during sample encoding had affected target processing (HA > AA and HH > AH, respectively), this is unlikely for 2 reasons. First, the jittered delay of 2–6 s between sample and target objects allowed us to model BOLD responses to the target objects independently from BOLD responses to the sample objects. Second, if a differential pattern of hand movements due to haptic sample encoding had affected target processing, we would expect to consistently find increased activations in motor cortex for conditions with versus without haptic samples, which was not the case (see Supplementary Results).

**Conclusions**

In summary, the present fMRI study used a delayed-match-to-sample paradigm to study audio-haptic interactions in higher order object recognition. We propose that audition and touch interact at various stages of object processing with crossmodal modulations at early association cortices and intermediate modality-sensitive object recognition regions (pSTS and putatively LO) and converge into higher order conceptual object representations in the left FG. Future studies might shed more
light on the role of the FG in multisensory object recognition and particularly in the integration of auditory and haptic object information.

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

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Notes
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References