Sensory and Motor Systems

Neural Evidence of Hierarchical Cognitive Control during Haptic Processing: An fMRI Study

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Abstract

Interacting with our immediate surroundings requires constant manipulation of objects. Dexterous manipulation depends on comparison between actual and predicted sensory input, with these predictions calculated by means of lower- and higher-order corollary discharge signals. However, there is still scarce knowledge about the hierarchy in the neural architecture supporting haptic monitoring during manipulation. The present study aimed to assess this issue focusing on the cross talk between lower-order sensory and higher-order associative regions. We used functional magnetic resonance imaging in humans during a haptic discrimination task in which participants had to judge whether a touched shape or texture corresponded to an expected stimulus whose name was previously presented. Specialized haptic regions identified with an independent localizer task did not differ between expected and unexpected conditions, suggesting their lack of involvement in tactile monitoring. When presented stimuli did not match previous expectations, the left supramarginal gyrus (SMG), middle temporal, and medial prefrontal cortices were activated regardless of the nature of the haptic mismatch (shape/texture). The left primary somatosensory area (SI) responded differently to unexpected shapes and textures in line with a specialized detection of haptic mismatch. Importantly, connectivity analyses revealed that the left SMG and SI were more functionally coupled during unexpected trials, emphasizing their interaction. The results point for the first time to a hierarchical organization in the neural substrates underlying haptic monitoring during manipulation with the SMG as a higher-order hub comparing actual and predicted somatosensory input, and SI as a lower-order site involved in the detection of more specialized haptic mismatch.

Key words: cognitive control; haptic; manipulation; SI; somatosensory predictions; supramarginal gyrus

Significance Statement

The findings in the present study have important implications for the understanding of the neural architecture that supports haptic monitoring during manipulation. The results point, for the first time, to a hierarchical organization in the neural substrates underlying haptic monitoring during manipulation. In this hierarchy, the supramarginal gyrus (SMG) is positioned as a higher-order region comparing predicted and actual somatosensory input, and primary somatosensory area (SI) as a lower-order site involved in the detection of more specialized haptic mismatches. The increased functional connectivity between the SMG and SI during the processing of unexpected stimuli emphasizes the cross talk between lower-order sensory and higher-order associative regions during manipulation.
Introduction
Interacting with our immediate surroundings requires constant manipulation of objects. Dexterous manipulation (sequences of somatosensory events linked to subgoals) requires that monitoring mechanisms (controllers) adapt motor commands to the relevant physical properties of the target object (Johansson and Westling, 1988). Motor commands are based on both the previously stored sensorimotor memory representations of objects (Johansson and Cole, 1992) and the current state of the motor apparatus (which depends on previous representations of the body in terms of proprioceptive information; Giummarra et al., 2008). Furthermore, information about the surface properties of the object is provided by somatosensory afferents in the hand. In particular, this relies mainly on the timing and firing rate of slow- and fast-adapting afferent populations (FA-I and SA-I) that can discriminate different surface curvatures after as few as five afferents have begun firing (Johansson and Westling, 1987; Johansson and Bizniak, 2004).

To regulate haptic monitoring, controllers compare the actual sensory input with the expected sensory consequences of initiated motor commands by means of the efference copy (Feinberg, 1978; Wolpert and Miall, 1996; Frith, 2014). This copy of efferent motor commands allows calculating how self-generated movement might influence the input sensory signal as well as maintaining performance in the presence of feedback delays. In other words, these predictions are necessary, taking into account that sensorimotor monitoring loops involved in corrective actions have long time delays (~100 ms; Johansson and Flanagan, 2008). Of note, in addition to the efference copy (where a copy of the motor commands issued to an effector is projected to low-level somatosensory neurons; Von Holst and Mittelstaedt, 1950), the information transfer from motor to sensory areas might also occur at multiple levels, leading to the distinction of lower- and higher-order (corollary discharge) CD signals (Crapse and Sommer, 2008). Lower-order CD signals regulate the sensory information that enters the system whereas higher-order CD signals implement adjustments in anticipation of the sensory input, facilitating the contextual interpretation of sensory information. To combine these two types of signals, the interplay between lower-order sensory areas and higher-order associative regions is crucial. Despite previous literature supporting the hierarchical nature of haptic processing (Bodegård et al., 2001; Bohlhalter et al., 2002; Sathian et al., 2011; Kassuba et al., 2013), there is still scarce knowledge regarding the hierarchy of the neural substrates underlying haptic monitoring and how these two types of information are implemented by local and global networks.

The present study assessed this issue by focusing on the cross talk between lower-order somatosensory and higher-order associative regions. To address this, we first identified, using an independent localizer task, lower-order somatosensory regions that process shapes or textures selectively. Afterward, we conducted a haptic discrimination task in which unexpected shapes and textures were presented to trigger monitoring mechanisms involved in the comparison between predicted and actual sensory input. We assessed whether the specialized haptic areas identified with the localizer task distinguished expected from unexpected (60%) stimuli in the category they were suited to process. We also conducted a whole-brain analysis to identify higher-order areas involved in haptic monitoring. We hypothesized that higher-order associative regions supporting haptic monitoring during manipulation would respond similarly regardless of the nature of the haptic mismatch, independently of whether it was a shape or texture mismatch. Some of these areas (the highest in the hierarchy) might even serve to detect mismatches in other sensory modalities. In contrast, we expected to see different patterns of response in specialized somatosensory regions (e.g., somatosensory cortices or areas selectively responding to haptic exploration of shapes and textures) depending on the type of tactile property (shape or texture) violating the expectation.

Materials and Methods
Participants
Twenty-two right-handed participants (13 female, mean age = 23.2 ± 1.4 years) took part in the experiment. For the localization task, the data of 20 subjects were analyzed, since two subjects were excluded from the fMRI analysis. One subject was excluded due to anomalous cortical response to tactile events, which did not elicit activity in the somatosensory cortex. Moreover, the logfile of another participant was not generated. For the haptic discrimination task, three additional subjects were excluded from the fMRI analysis since their responses were not recorded. Hence, the fMRI analysis of the haptic discrimination task included seventeen participants (12 female, mean age = 23.4 ± 1.5 years). The experiment was undertaken with the understanding and written consent of each participant and was approved by the local ethics committee in accordance with the Declaration of Helsinki.

Procedure and general experimental design
Before the single scanning session participants underwent a training phase that lasted ~30 min. During the training phase, they became familiarized with the haptic stimuli (used in both the independent localizer task and the haptic discrimination task) and the experimental procedure. Since inside the scanner all the instructions were vocally presented, the meaning of each auditory cue was explained, and a brief simulation was conducted. Partic-
Participants lay supine in the scanner and were blindfolded during the entire scanning session. First, the T1 structural image was acquired. To avoid circularity (Kriegeskorte et al., 2008), an independent functional localizer task was then conducted to localize areas typically involved in haptic processing shape and texture [from here on referred to as property-selective haptic regions of interest (ROIs); Amedi et al., 2001]. Lastly, the haptic discrimination task was conducted, to assess whether the previously identified haptic-specialized areas distinguished expected from unexpected stimuli and to establish higher-order regions involved in the detection of haptic incongruencies. Note that these higher-order areas are not selective to haptic processing and thus could not be spotted by the independent localizer task.

**Independent functional localizer task**

**Stimuli**

For the localizer task, we used six real 3D objects (pen cap, thread spool, eye drop bottle, clothespin, bottle cork, and mini pencil) and six textures, ~4 × 4 cm (corkoy, cork, sackcloth, sandpaper, sponge, and scourer), presented in a rotating tray (Fig. 1A).

**Design**

Participants were asked to palpate sequences of shapes or textures presented in a rotating tray and covertly recognize them. The task began with a 10-s resting interval that was not analyzed. We used a block design that consisted of a single run with two haptic conditions (shape, Sh; texture, Tx) and a rest condition (Fig. 1B). All blocks lasted 15 s, and Sh and Tx blocks were alternated between each pair of Rest blocks. Each haptic block consisted in palpating (4 s/palpation) three stimuli placed in a quarter of the rotating tray and then spinning the tray to continue with the next quarter (after the resting period; Fig. 1C). The hand always palpated the sequence of objects in the same order since the location of the stimuli was previously known. The task lasted ~5 min.

**Haptic discrimination task**

**Stimuli**

A total of 25 3D haptic stimuli were manufactured, measuring ~4 × 4 × 2 cm and varying in shape and texture. The 25 stimuli comprised a variety of five shapes (flower, circle, heart, square, and star) that were covered with five textures (corkoy, sandpaper, plastic, paper, and expanded polystyrene; Fig. 1A). Thus, each stimulus had two properties: a particular shape and a particular texture (e.g., circle of corkoy, circle of paper, etc.). All the stimuli were chosen so as to be easily identifiable, and the five shapes were previously used in two other studies (Gurtubay-Antolin et al., 2015; Gurtubay-Antolin et al., unpublished observation).

**Design**

Participants were presented with brief sequences of objects or textures to palpate and had to judge whether a touched stimulus corresponded to an expected stimulus whose name had been previously presented orally. Subjects were placed with the right hand facing upward, and the stimuli were placed into the subject’s palm by the experimenter. The experiment was designed as a mixed block/event-related design with two haptic conditions (shape, Sh; texture, Tx) in which subjects were instructed to attend to the shape or the texture of the stimuli (block design), a control motor condition (motor) in which they were asked to move their fingers as they were exploring an imaginary object (and were explicitly instructed not to touch themselves), and a “rest” condition.

The experimental design consisted of four runs and each run comprised 10 haptic blocks (five Sh, five Tx), four motor blocks, and five rest blocks (Fig. 2B). All the runs began with a 10-s resting interval that was not analyzed. The type of haptic block (shape or texture) was randomized and lasted 32 s. Each haptic block consisted of four consecutive somatosensory trials (8 s each if we take into account the ~4-s jittering interval). Each haptic block began with an auditory cue indicating the dimension

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**Figure 1.** Stimuli and procedure used in the localizer task. **A,** Stimuli: rotating tray with three objects (shapes: yellow) or three textures (green) in each quarter. Shapes were real 3D objects: pen cap, thread spool, eye drop bottle, clothespin, bottle cork, and mini pencil. Textures, ~4 × 4 cm, were corduroy, cork, sackcloth, sandpaper, sponge, and scourer. **B,** Block design for the single run. The run comprised 10 haptic blocks (yellow-shape; green-texture) and began with a 10-s interval that was not analyzed. The order of the haptic blocks was alternated. **C,** Procedure for a haptic block. The block began with an auditory cue indicating that the haptic block would begin. After 1 s of silence, the first auditory cue indicated to the participant the first (of the sequence of three) stimulus and to palpate it. After 4 s of exploration, a second (and third) auditory cue indicated to move the hand toward and palpate the second (and third) stimulus. The hand always moved in the same direction when ranging from one stimulus toward the next one since the location of the stimuli was previously known. To conclude, an auditory cue instructed the participants to spin the tray (to prepare it for the next block) and rest.
of the object that had to be attended (50% shape, 50% texture). After a 2-s interval, the first haptic trial started (Fig. 2C). In each haptic trial, the name of an expected stimulus was vocally presented and a second later, an auditory cue indicated that the exploration of the actual item could begin. In half of the trials, the word delivered by the headphones corresponded to the touched object (congruent, C), and in the other 50% of trials, the object did not match the name (incongruent, I). All the possible combinations were presented the same number of times. We therefore had six conditions [four corresponding to all the possible combinations: congruent shape (CSh), incongruent shape (ISh), congruent texture (CTx), and incongruent texture (ITx); and two additional control conditions: motor (M) and rest (R)]. Free exploration was allowed. 2.5 s later, a second auditory cue indicated that the exploration period was over. After a jittering period, the next haptic trial could start. After four haptic trials, the participants were asked about the number of incongruent trials. Using their left hand, participants had to push a button as many times as the incongruent trials that had been presented (ranging from 0 to 4). To conclude, an auditory cue indicated whether a rest or “motor” block followed (a block in which participants were asked to move the fingers as they were exploring an imaginary object).

Behavioral analysis
We considered a correct response block each time the number of incongruencies presented in a haptic block and the number of times the participant had pushed the button matched. The percentage of correct responses served (1) to rule out blocks with erroneous responses from the fMRI analysis, and (2) to obtain an overall estimate of the performance.

Image acquisition
Scanning was performed on a 3-T Siemens Trio System. Functional data were acquired using a gradient echo pulse sequence (32 transverse slices oriented along the anterior-posterior commissural axis with a 30° upward tilt to avoid the eyes, repetition time of 2 s, echo time of 30 ms, 3 × 3 × 3.5 mm voxels, 0.8-mm interslice gap). A high-resolution T1-weighted magnetization-prepared rapid acquisition gradient echo (MPRAGE) image (240 slices sagittal, TR = 2300 ms, TE = 2.98 ms, 1-mm isotropic voxels) was also collected.

Image processing and statistical analysis
fMRI data were analyzed using standard procedures implemented in the Statistical Parametric Mapping software (SPM8, Wellcome Trust Center for Neuroimaging, University College, London, United Kingdom; https://www.fil.ion.ucl.ac.uk/spm/). The preprocessing consisted of several steps: realignment, segmentation, normalization, and smoothing. To correct head-movement artefacts.
based on an affine rigid body transformation, images were spatially realigned with respect to the first volume of the first run (Friston et al., 1996). Each participant’s MPRAGE scan was coregistered to the mean echo-planar imaging (EPI) volume, produced in the previous step during spatial realignment. Each coregistered structural scan was then segmented using New Segment (Ashburner and Friston, 2005). Then the flow fields containing the deformation parameters of this template were used to normalize each participant’s realigned EPIs to MNI space. Finally, normalized EPI images were re-sliced to $2 \times 2 \times 2$ mm and smoothed with an 8 mm FWHM Gaussian kernel (Ripollés et al., 2014, 2016).

### Independent functional localizer task

We aimed to identify specialized haptic regions responding differentially to the exploration of shapes and textures (property-selective haptic ROIs) to analyze their involvement in tactile monitoring. To avoid circularity (Kriegeskorte et al., 2008), the ROIs were created using independent data obtained from the independent localizer task. For the independent functional localizer task, a block design matrix was specified using the canonical hemodynamic response function. Block onsets were modeled at the moment at which participants heard the auditory cue that indicated that they could start palpating the first of the three stimuli placed in a quarter of the rotating tray. First-level statistical analysis was based on a least square estimation using the general linear model. Individual brain responses to the “shape” and “texture” conditions were modeled with a regressor wave form convolved with a canonical hemodynamic response function. Movement parameters (estimated during the realignment phase) were also included in the model as covariates of no interest to correct for motion effects as well as constant vectors. Linear contrast images for the main effect of haptic processing ($Sh + Tx >$ rest) and the main effect of property ($Sh > Tx$ and vice versa) were calculated for each subject, and statistical parametric maps (SPMs) were generated. Group activation was calculated using a random effects model, accounting for intersubject variance. Main effects of haptic processing were only used for sanity checks to confirm that they activated sensorimotor cortical areas. Property-selective haptic ROIs were created entering individual contrast images into a second level one-sample t test to test for (1) main effects of shape ($Sh > Tx$) and (2) main effects of texture ($Tx > Sh$; $P_{uncorrected} = 0.005, k = 50$). To maximize sensitivity within our ROIs, we performed a meta-analysis using NeuroSynth [a platform for large-scale, automated meta-analysis of fMRI data; http://www.neurosynth.org; Yarkoni et al., 2011; for a similar approach, see Ripollés et al., 2016]. We put together a term-based search for “tactile” that resulted in 190 studies (search performed on October 20, 2016). Then, a forward inference mask (which represented the probability that the term tactile was associated with a particular activation) was generated (corrected at $P_{FDR} = 0.01$). We then refined the previously created ROIs by masking them with the results of the NeuroSynth meta-analysis. In other words, each final ROI contained only voxels that were part of the original ROIs and that were also tactile-related according to the meta-analysis.

### Haptic discrimination task

For the haptic discrimination task, an event-related design matrix was specified using the canonical hemodynamic response function. Trial onsets were modeled at the moment at which participants heard the auditory cue that indicated that they could palpate the stimuli. We only analyzed blocks with correct responses (blocks where the number of presented incongruencies matched the times the participant pushed the response button). Blocks with incorrect responses were not analyzed. First-level statistical analysis was based on a least square estimation using the general linear model. Individual brain responses to the different conditions [congruent shape ($CSh$), incongruent shape ($ISh$), congruent texture ($CTx$), incongruent texture ($ITx$), and motor ($M$)] were modeled with a regressor wave form convolved with a canonical hemodynamic response function. Movement parameters (estimated during the realignment phase) were also included in the model as covariates of no interest to correct for motion effects as well as constant vectors.

### ROI analysis

First, a ROI analysis was performed using the results from the independent functional localizer task. To assess whether property-selective haptic ROIs could detect conflicting haptic information in the category they were suited to process, paired t tests were conducted. We compared mean beta values (1) between $CSh$ and $ISh$ conditions in ROIs selective to shape processing and (2) between $CTx$ and $ITx$ in ROIs selective to texture processing. The significance threshold was corrected for multiple comparisons taking into account the number of ROIs.

### Whole-brain analysis

Additionally, to identify higher-order areas involved in haptic monitoring, a whole-brain analysis was conducted. Linear contrast images for (1) the main effect of haptic processing with the related motor component ($CSh + ISh + CTx + ITx > R$), (2) motor processing without somatosensory stimulation ($M > R$), (3) somatosensory stimulation without motor processing ($CSh + ISh + CTx + ITx > M$), (4) the main effect of shape ($CSh + ISh > CTx + ITx$), and (5) the main effect of texture ($CTx + ITx > CSh + ISh$) were calculated to perform some sanity tests. These tests aimed to confirm that the previous contrasts activated (1) sensorimotor cortical areas, (2) motor, but not somatosensory, areas, (3) somatosensory, but not motor, areas, and (4) previously reported areas specifically processing shapes as opposed to textures and (5) vice versa.

To identify regions involved in haptic monitoring, linear contrast images for (1) the main effect of congruency ($CSh + CTx > ISh + ITx$), (2) the main effect of incongruency ($ISH + ITx > CSh + CTx$), and the interaction terms (3) for greater shape incongruency ($ISH > CSh > ITx – CTx$), and (4) for greater texture incongruency ($ITx – CTx > ISh > CSh$), were calculated for each subject. SPMs were generated. Group activation was calculated using a random effects model, accounting for intersubject variance. To
test for the main effect of incongruency (ISH + ITx > CSh + Ctx and the reverse) and the interaction term (ISH – CSh > ITx – Ctx and the reverse), the individual contrast images were entered into a second level repeated measures ANOVA with two within-subjects factors (property and congruency) and two levels each [shape, texture (Sh and Tx) and congruent, incongruent (C and I), respectively]. The results were thresholded at \( p = 0.05 \) FEW-corrected at cluster level, with a cluster-forming (voxel-wise) threshold of \( \rho_{uncorrected} < 0.001 \) (Woo et al., 2014; Flandin and Friston, 2017). In cases where clusters were further FWE-corrected at the voxel-level, the voxel-level and the FWE-corrected \( p \) value are explicitly mentioned.

**Functional connectivity (psychophysiological interaction; PPI)**

PPI analyses (Friston et al., 1997; Gitelman et al., 2003) identify voxels in which activity is more closely related to activity in a seed ROI in a given psychological context. In the present study, PPI analyses were performed to identify brain regions that were more functionally coupled (1) with regions that showed the main effects of incongruency (from now on referred to as iROIs) while processing incongruent versus congruent trials, (2) main effects of congruency (cROIs) while processing congruent versus incongruent trials, and (3) an interaction between incongruency and property (IntROIs) while processing shape incongruences in the opposite direction of texture incongruences (ISh – CSh > Itx – Ctx).

Four mm radius spheres were created around the group peaks obtained for (1) iROIs, (2) cROIs, and (3) IntROIs. For all participants, individual deconvolved time-series were extracted from all voxels within the seed ROIs. New linear models were generated at the individual level, using three regressors. The first regressor was the activity extracted in the seed area. The second regressor represented the condition as a vector that coded (1) the main effect of incongruency (CSh: –1, ISh: 1, CTx: –1, Itx: 1) for iROIs, (2) the main effect of congruency (CSh: 1, ISh: –1, CTx: 1, Itx: –1) for cROIs, and (3) the incongruence by property interaction (CSh: –1, ISh: 1, CTx: 1, Itx: –1) for IntROIs. The third regressor represented the interaction of interest between the first (physiologic) and the second (psychological) regressors. This was calculated as the element by element product of the extracted time-series (the first eigenvariate from each voxel in the sphere) and the second regressor. The result of this product was then reconvolved with the canonical hemodynamic response function to create the final PPI regressor (Gitelman et al., 2003). The design matrix also included movement parameters as a regressor of no interest. A significant PPI indicated a change in the regression coefficients between any reported brain area and the seed area, related to the experimental condition (ISH + ITx > CSh + Ctx for iROIs; CSh + CTx > ISh + Itx for cROIs; and ISh – CSh > Itx – Ctx for IntROIs). The voxels identified in this analysis show a pattern of activity correlated with the seed region. Individual summary statistic images obtained at the first level (fixed effects) analysis were entered in a second-level (random effects) analysis using a one-sample \( t \) test. The results were thresholded at \( p = 0.05 \) FEW-corrected at cluster level, with a cluster-forming (voxel-wise) threshold of \( \rho_{uncorrected} < 0.001 \) (Woo et al., 2014; Flandin and Friston, 2017).

**Results**

**Behavioral results**

Overall percentage of correct response was 91% (SD = 5; considering both types of haptic blocks). The accuracies considering each property separately were 92 ± 5% for shape blocks and 90 ± 9% for texture blocks, with no significant difference between the two (\( t_{(10)} = 0.1; p = 0.3 \)).

**fMRI results**

**Property-selective haptic ROI creation from localizer task**

The results revealed a main effect of shape (Sh vs Tx) in three clusters located in the left anterior intraparietal sulcus [aIPS; –34 –40 46, \( t_{(19)} = 5.49, k = 53044 \), in the left lateral occipital complex [LOC; –48 –66 –6, \( t_{(19)} = 3.65, k = 6044 \), and in the right LOC [52 –56 –6, \( t_{(19)} = 3.10, k = 3324 \)]. Additionally, a cluster located in the left secondary somatosensory area (SII), parietal operculum [–42 –18 24, \( t_{(19)} = 5.01, k = 197 \)] showed a main effect of texture (Tx vs Sh; Fig. 3).

**Haptic discrimination task**

The average number of trials included in the fMRI analysis was 36 ± 1 (mean ± SD, CSh 37 ± 2; ISh 36 ± 3; CTx 35 ± 5; ITx 35 ± 3). The haptic discrimination task activated a network associated with haptic stimulation, comprising bilateral sensorimotor areas, insulae, posterior parietal cortices, LOCs, premotor and supplementary motor areas, prefrontal regions, and thalamus.
**Incongruence in property-selective haptic ROIs**

Results of paired t-tests between the CSh and ISh conditions for the three ROIs showing a main effect of shape (Sh vs Tx) were not significant [left aIPS (p = 0.2), left LOC (p = 0.2), right LOC (p = 0.3), with the significance threshold \( p_{\text{BONFERRONI}} = 0.05/3 = 0.017 \). Results of a paired t-test between the CTx and ITx conditions for the ROI showing a main effect of texture [Tx vs Sh; left SII, parietal operculum (p = 0.06)] were not significant, either.

**Whole-brain analysis: flexible factorial results**

Flexible factorial results revealed important effects of incongruency in a cluster located in the left supramarginal (SMG; and part of angular) gyrus [iROI1; \( -54 -52 24, t_{(48)} = 5.44, \quad p_{\text{FWE cluster}} < 0.001, \quad p_{\text{FWE voxel}} = 0.03, \quad k = 984 \)], in the left middle temporal gyrus [MTG; iROI2; \( -48 -18 -10, t_{(48)} = 5.31, \quad p_{\text{FWE cluster}} < 0.001, \quad p_{\text{FWE voxel}} = 0.04, \quad k = 744 \)], and in the medial prefrontal cortex [mPFC; iROI3; \( -10 54 34, t_{(48)} = 4.64, \quad p_{\text{FWE cluster}} = 0.003, \quad k = 421 \); Fig. 4A]. An important effect of congruency was found in the right primary somatosensory area [SI; cROI; \( 58 -16 32, t_{(48)} = 4.42, \quad p_{\text{FWE cluster}} = 0.04, \quad k = 235 \); Fig. 4B] as well as a significant interaction of incongruency \( \times \) property (Sh + ITx > CSh + Ctx) in the left SI [IntROI; \( -32 -36 64, t_{(48)} = 4.39, \quad p_{\text{FWE cluster}} = 0.03, \quad k = 238 \); Fig. 4C].

**Functional connectivity results**

We ran PPI analyses to further investigate which brain regions showed a significant change in the regression coefficients between that area and the seed area, related to the experimental condition (ISh + ITx > CSh + Ctx for iROIs and ISh – CSh > Itx – CTx for Int ROI), indicating that they were more functionally coupled. The left SMG (iROI1) showed increased incongruency-related connectivity with a cluster that included the left primary motor area (MI), premotor area, SI, SII, and part of the superior parietal lobe [SPL; \( -32 -16 42, t_{(16)} = 6.21, \quad p_{\text{FWE cluster}} < 0.001, \quad k = 2045 \)], as well as with a cluster including regions of the right SI and SPL [\( 50 -42 66, t_{(16)} = 4.93, \quad p_{\text{FWE cluster}} < 0.001, \quad k = 451 \)]. With a more liberal threshold of \( p_{\text{FWE cluster}} = 0.05 \), the left SMG also presented increased connectivity during incongruent trials with a region in the right cerebellum [\( -20 -48 -28, t_{(16)} = 5.04, \quad p_{\text{FWE cluster}} = 0.01, \quad k = 201 \); Fig. 5A]. Increased incongruency-related connectivity was also found between the left MTG (iROI2) and the left MI and SI [\( -46 -22 -62, t_{(16)} = 5.22, \quad p_{\text{FWE cluster}} < 0.001, \quad k = 546 \); Fig. 5B]. The cluster at the left medial frontal cortex (iROI3) did not show any increased incongruency-related connectivity. The cluster in the right primary somatosensory area (cROI) did not show any increased incongruency-related connectivity. Lastly, the left SI (IntROI) presented increased connectivity for higher shape incongruency than texture incongruency with the left SMG area [\( -62 -34 30, t_{(16)} = 5.98, \quad p_{\text{FWE cluster}} = 0.05, \quad k = 152 \); Fig. 5C].

For further information, unthresholded t-maps resulting from the fMRI analysis have been uploaded to NeuroVault (Gorgolewski et al., 2015; https://neurovault.org/collections/4161/).

**Discussion**

The present study aimed to elucidate the hierarchy in the neural substrates underlying haptic monitoring during manipulation, focusing on the cross talk between the lower-order sensory regions and the higher-level associa-
We found that the aIPS, LOC, and SII responded differently to the exploration of shapes and textures, whereas they did not differ between expected and unexpected conditions. This suggests that they are specialized in haptic exploration and processing but are not involved in tactile monitoring. The left SMG, the middle temporal, and the medial prefrontal cortices were activated whenever there was a haptic mismatch, regardless of its nature (shapes and textures alike). In contrast, the activity in the left SI distinguished between unexpected shapes and textures, in line with more specialized haptic mismatch detection. Moreover, the left SMG-Ang area and left SI were more functionally coupled during unexpected trials.

The activity observed in the left SMG-Ang gyrus for unexpected haptic input converges with lesion and functional neuroimaging studies relating this area to comparison processes between predicted and actual sensory consequences of ongoing actions (Sirigu et al., 2004; Desmurget et al., 2009). Prior electrophysiological studies found that the mismatch between internal predictions and reafferent signals elicited a parietally distributed error signal resembling the N400 component (Gurtubay-Antolin et al., 2015; Padrao et al., 2016), whose main neural substrate has been located in the SMG (Lau et al., 2008; Baumgaertner et al., 2002). The functional connectivity analyses revealed that the left SMG-Ang was more functionally coupled during unexpected than expected trials with bilateral sensorimotor regions and the right cerebellum, suggesting a pivotal role of the SMG-Ang area in orchestrating the monitoring of sensory predictions. In this vein, Jenmalm et al. (2006) observed activation in the right SMG area when unexpectedly heavy and light weights were lifted, as well as activity in the left SI and right cerebellum during the lifting of unexpectedly heavy and light weights, respectively. Taken together, these results favor the idea that the controller (SMG-Ang) compares the actual sensory input projected by the early somatosensory cortex with the predictions computed in the cerebellum, made on the basis of proprioceptive information. In fact, the cerebellum is thought to be the keystone computing these predictions since it is widely accepted that it contains internal models of the motor apparatus (Wolpert et al., 1998) and is involved in recruiting internal representations of object properties (Bursztyn et al., 2006).

Furthermore, the preponderant left-lateralized (contralateral) activity of the SMG-Ang when manipulating objects with the right hand contrasts with the study by Jenmalm et al. (2006), who reported activity in the right SMG even when people used the right hand for lifting. Indeed, a general dominance of the right hemisphere for somatosensory functions has been proposed (Naito et al., 2005). However, aspects such as the selection of the hand configuration (Emmorey et al., 2007) might explain the left-lateralized preponderance, which converges with studies reporting greater activation of the left SMG when participants are asked to pantomime object use (Rumiati et al., 2004), or when deaf signers name objects compared to speakers (Emmorey et al., 2007). In this task, the representation of the expected object was available before its perception, so participants might have benefited.

![Figure 5. Functional connectivity results (PPF). A, Connectivity results seeding from ROI that showed main effects of incongruency (iROIs) while processing incongruent versus congruent trials (CSh: –1, ISh: 1, CTx: –1, and ITx: 1). Seeds are 4-mm spheres centered at the peak coordinate of the cluster located in left supramarginal-angular gyrus (iROI1) and in left middle temporal gyrus (iROI2; B). C, Connectivity results seeding from the ROI showing an interaction between incongruency and property (IntROI), while processing shape incongruencies different from texture incongruencies (CSh: –1, ISh: 1, CTx: 1, and ITx: –1). Maps are thresholded at $p_{FWE}$ cluster-level = 0.001 and $p = 0.05$ (with cluster-forming voxel-wise thresholds of $p_{uncorrected} < 0.001$) and a minimum cluster size of 50 voxels. MI, primary motor cortex; MTG, middle temporal gyrus; SI, primary somatosensory cortex; SII, secondary somatosensory cortex; SMG, supramarginal gyrus; R, right hemisphere and L, left hemisphere. MNI coordinates.](image-url)
from preparing a hand configuration specific for the expected stimulus.

Different patterns of response were found for unexpected shapes and textures in regions of the left SI, including areas that were functionally connected to the SMG area during incongruence. This supports the notion that a more specialized mismatch detection occurs in the left SI, in line with the view of SI as a lower-order region in the hierarchy of the neural substrates that underlie tactile monitoring. This finding emphasizes the importance of the contralateral SI not only in early somatosensory processing and short-term maintenance of haptic traces (Kaas et al., 2007; Romo et al., 1999), but also in haptic monitoring. In accordance with our results, SI has been seen to participate in decision-making during the haptic choice in a haptic delay task (Wang et al., 2012). SI cells showed differential neural activity when monkeys had to choose between different haptic objects, and such differential activity diminished significantly in erroneous trials. In addition, our results show increased functional connectivity between SI and a region in the SMG when unexpected shapes but not textures were presented, pointing to a relevant cross talk between these regions. However, the right SI did not show increased functional connectivity related to incongruence, highlighting the notion that the functional connections between the left SI and the SMG region are uniquely enhanced by incongruence processing. On the whole, this interplay seems to be important in the detection of haptic expectancy violations, with SI processing more specialized tactile information than the SMG-Ang area.

Lastly, the results suggest that the MTG and the mPFC are additional higher-order regions that work in parallel with the SMG-Ang. During mismatches, the MTG showed increased functional connectivity with the left sensorimotor cortex. The MTG has been observed to respond to deviant stimuli in a tactile oddball task (Allen et al., 2016) and to unexpected touch sensations in monkeys (Perrett et al., 1990); thus, it seems to be related to violations of predictions selective to the haptic domain. Moreover, the activation found in the mPFC seems to correspond to higher-order cognitive control areas (Rushworth et al., 2004; Nee et al., 2011). This area has been associated with mismatch detection in several sensory modalities, suggesting its multimodal or amodal nature (Gaebler et al., 2015; Blakemore et al., 2000; Malekshahi et al., 2016). This matches well with our results, which show a lack of functional connections between the mPFC and somatosensory areas (while these are present between the somatosensory regions and the SMG and MTG). There is still debate on this question, as some theories state that the mPFC supports conflict monitoring by calling for control processes to resolve discrepancies (Botvinick et al., 2001) while a recent line of research suggests that it responds to unexpectedness (Jessup et al., 2010; Oliveira et al., 2007).

Importantly, the fact that the SMG-Ang, the MTG and the mPFC exhibited negative values in all the haptic conditions might reflect the neuronal inhibition associated with the suppression of items that were not expected (Frankenstein et al., 2003). This inhibition was greater in congruent trials where the actual stimulus matched the expectancy, since the non-expected stimuli had to be inhibited for a longer period. In incongruent trials, the non-expected items were inhibited only until participants realized that the touched stimulus was not the expected one. Subsequently, they might have disinhibited the non-expected items to identify the actual item, even if this was not required in the task. An alternative explanation for these deactivations is that they are associated with the decrease in activity shown by areas of the default mode network (Xu et al., 2016). Recent findings suggest the existence of a gradient in the human cortical organization (which spans from primary sensorimotor cortices to higher-order areas whose activity is not specific to a single sensory modality) that is reflected in cortical microstructure and macroscale connectivity (Huntenburg et al., 2018). According to this view, a continuous pattern of connectivity exists between sensorimotor areas that converge in multimodal integration areas, and higher order regions of the default mode network. Our results fit nicely with this interpretation.

Of note, the localizer task did not show shape-related activations in sites of the IPS that are typically involved (Sathian, 2016), nor texture-selective activity in the early visual cortex (Sathian et al., 2011; Eck et al., 2013). This raises questions about the actual sensitivity of the localizer task, which may have been affected by the limited amount of time for scanning. Despite this potential limitation, the task revealed property-selective haptic ROIs that fit nicely with previous reports of shape-selective activations in the LOC and the IPS (Roland et al., 1998; Sathian et al., 2011), as well as texture-sensitive areas located in the SII, parietal operculum (Roland et al., 1998). Particularly, the LOC and the IPS are involved not only in haptic perception but also in haptically-guided grasping. This is suggested by the fact that the occipital pole (active during haptic exploration of shapes) shows stronger functional connectivity with the LOC and the IPS during haptic than visual exploration (Monaco et al., 2017) and that the aIPS is sensitive to characteristics of the required grasp (Marangon et al., 2016).

Altogether, the results point to a hierarchical organization in the neural substrates underlying haptic monitoring during manipulation, with the SMG as a higher-order region comparing actual and predicted somatosensory input and SI as a lower-order site involved in the detection of more specialized haptic mismatch. We report, for the first time, the functional coupling of these regions during the processing of unexpected tactile stimuli, supporting their pivotal role in haptic monitoring.

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