Understanding Effector Selectivity in Human Posterior Parietal Cortex by Combining Information Patterns and Activation Measures

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The posterior parietal cortex (PPC) has traditionally been viewed as containing separate regions for the planning of eye and limb movements, but recent neurophysiological and neuroimaging observations show that the degree of effector specificity is limited. This has led to the hypothesis that effector specificity in PPC is part of a more efficient than strictly modular organization, characterized by both distinct and common activations for different effectors. It is unclear, however, what differentiates the distinctions and commonalities in effector representations. Here, we used fMRI in humans to study the cortical representations involved in the planning of eye, hand, and foot movements. We used a novel combination of fMRI measures to assess the effector-related representational content of the PPC: a multivariate information measure, reflecting whether representations were distinct or common across effectors and a univariate activation measure, indicating which representations were actively involved in movement preparation. Active distinct representations were evident in areas previously reported to be effector specific: eye specificity in the posterior intraparietal sulcus (IPS), hand tuning in anterior IPS, and a foot bias in the anterior precuneus. Crucially, PPC regions responding to a particular effector also contained an active representation common across the other two effectors. We infer that rostral PPC areas do not code single effectors, but rather dichotomies of effectors. Such combinations of representations could be well suited for active effector selection, efficiently coding both a selected effector and its alternatives.

**Key words:** effector selectivity; fMRI; foot; MVPA; parietal cortex; pointing

**Introduction**

The ability to generate an appropriate response in a complex environment is of utmost importance for the survival of any organism. A critical aspect is the selection of the effectors involved, such as eyes, hand, and foot. While the posterior parietal cortex (PPC) has been implicated in this process, it is debated how this structure maps multiple effectors onto the 2D cortical surface.

PPC organization could separate motor responses in relation to the effector that is to be moved. This would lead to separate neural modules for the control of different effector systems (Cui and Andersen, 2007). To date, this has been the prevailing concept of PPC organization (Colby and Goldberg, 1999; Matelli and Luppino, 2001; Fillimon et al., 2009). For instance, the lateral intraparietal area (LIP) is thought to encode eye movements (Gnadt and Andersen, 1988), the parietal reach region (PRR) during reaching movements (Snyder et al., 1997), and the anterior intraparietal sulcus (aIPS) grasping movements (Murata et al., 2000). Recently, it has been argued that effector specificity in PPC is part of an efficient coding scheme, representing effectors in terms of commonalities in the behavioral repertoire (Graziano, 2006; Levy et al., 2007; Jastorff et al., 2010). For example, because eye and hand are often moved together, the efficient coding principle proposes a shared neural substrate for both effectors. Accordingly, both LIP and PRR respond during preparation of hand as well as eye movements (Snyder et al., 1997; Calton et al., 2002). Likewise, human fMRI studies have revealed large overlap in PPC during the planning of eye, hand (Beurze et al., 2007; Levy et al., 2007), and foot movements (Heed et al., 2011). However, it remains unclear which characteristic of those effector representations drives distinct and overlapping representations in PPC.

Here, we characterize identities and commonalities in the neural representations of eye, hand, and foot movements in PPC by using a novel combination of multivariate and univariate fMRI measures. Multivariate pattern analysis (MVPA) tests spatial patterns of fMRI activity to infer informational content (Fig. 1A; Haxby et al., 2001; Haynes et al., 2005; Gallivan et al., 2011, 2013). The underlying logic is that distinct patterns encode effector identities, whereas a shared pattern across effectors codes...
The region might code for eye movements, it might integrate hand and foot movements, or it might distinguish activity pattern of a region differs between eye movements and both hand and foot movements, but not between hand and foot movements. These possibilities can be disentangled by considering the degree of activation of that region for each effector, that is, whether the region is activated for the eye only, for the limbs, or for all three, respectively. Activation adds direction to the pattern information results. Crucially, we combine information and activation measures into one score, showing which distinct and common representations (Fig. 1D) are present as well as activated across PPC during movement planning.

Materials and Methods

Participants

Twenty-three healthy, right-handed participants with normal or corrected-to-normal vision participated in this study. Data of six participants were not included in further analyses due to poor task performance (see below). The remaining 17 participants (9 female) were aged 19–33 years (mean 23.5). Participants gave their written consent in accordance with the local ethics committee (CMO Committee on Research Involving Human Subjects, region Arnhem-Nijmegen, The Netherlands).

Experimental setup

Participants lay supine in the scanner, with their head inside a phased-array receiver head coil. The head and neck were stabilized within the coil using foam blocks and wedges. The limbs used in the task (right arm, right leg) were additionally cushioned for stabilization. Upper arms and legs were cushioned and strapped to minimize head movements during task execution. Visual stimuli were controlled using Presentation software (Version 14.7; Neurobehavioral Systems), projected onto a screen and viewed by the participant using a mirror, giving the perception that the stimuli were approximately above the participants’ head. Responses to stimuli were made using the eyes, right hand, or right foot.

Eye tracking. To track eye movements, the position of the left eye was recorded using a long-range infrared, video-based eye tracker (SMI) at a frequency of 50 Hz. Saccades were identified by detecting a 2% change in eye position, relative between baseline and the maximum amplitude of the trial. Results from the automatic analysis were verified visually.

Hand and foot movement recording. Hand and foot pointing was performed with the right limbs. Hand pointing involved rotating the wrist, and pointing at the target with the index finger. Foot pointing involved rotating the ankle to point at the target with the big toe. To track hand and foot movements, infrared light-emitting diodes (LEDs) were taped to the right index finger and right big toe. The infrared light, while invisible to the participants, was visible to a camera, mounted to the ceiling at a distance of 1–2 m of the scanner. One of the two LED lights was shortly interrupted in parallel with presentation of the stimulus and movement cue, respectively, for alignment of movement data with the experiment’s events. The LED locations were extracted from the video footage frame by frame using MATLAB (MathWorks) as done...
Movements were identified by detecting a 5% change from baseline of hand or foot position, relative to the maximum amplitude of the current trial, and were verified visually.

**Experimental paradigm**

Participants performed a delayed movement task using three possible effectors (eyes, right hand, and right foot) to six potential targets (4, 6, and 8 degrees to the left and right of the fixation point; Fig. 2). Participants started all trials pointing to a central fixation point (white dot, size 1°) with all three effectors. The hand and foot pointed to the fixation dot in the horizontal (left–right) direction only, as the degrees of freedom of wrist and ankle limit vertical movement. Trials started with a change of the fixation dot’s color to indicate the effector, together with the presentation of a movement target (400 ms). The fixation dot could change to red, blue, or orange to instruct the effector for the upcoming trial. After a delay period of 1.6–5.6 s, the fixation dot turned purple to indicate movement execution (400 ms). During the delay period, participants had to plan the instructed movement to the remembered target, and execute this movement at the presentation of the purple dot. After 1.6 s, the next trial started.

Trials were grouped in runs of 43 trials. Trial order was arranged such that each of the six trial types [resulting from three effectors (eye, foot, and hand) and two target sides (left and right)] followed each other equally often, while appearing random to the participant (Brooks, 2012). Each run started and ended with 20 and 8 s of fixation, respectively. These intervals served as baseline in the GLM analysis. The length of the breaks between runs was determined by the participant. The total duration of the experiment, consisting of 18 runs, 774 trials (43 of each type), excluding the breaks, was 86 min, split into two sessions.

We used eye and limb movement data to verify that participants held fixation during the delay period with all three effectors and only moved the cued effector during the execution period. Six participants made a large number of errors (performance < 75%) and were excluded from further analysis. The excluded participants mostly failed to keep eye fixation during the execution of hand or foot movement. For the remaining 17 participants, the average number of trials that entered analysis was 87% (SD: 6.7).

**fMRI analysis**

A recent report analyzed the data obtained in the present paradigm using a repetition suppression approach (Heed et al., 2013) and contains a detailed analysis of the behavioral data. Here, we developed and applied a new analysis approach that integrates univariate and multivariate analysis of these data to examine the cortical representations involved in the planning of eye, hand, and foot movements.

**Pattern information versus activation measures.** In the current study, we combined pattern information and activation measures into a pattern-activation framework. Pattern information analysis distinguishes regional representations (Kriegeskorte and Bandettini, 2007; Raizada and Kriegeskorte, 2010), focusing on differences in the activation pattern across voxels in that region. Activation analysis, on the other hand, is based on statistic parametric mapping, and typically tests average activation relative to a baseline in a given region. Activation is believed to represent the differential metabolic response of a region to different sensorimotor or cognitive challenges (Friston et al., 2011; Fig. 1A). The metabolic demand, in turn, is thought to indicate the involvement of a region in current processing. Note that the two measures are independent, similar to how a mean and SD are independent, or how a correlation measure is mean independent: a pattern can be highly informative, but, on average, might not be activated above baseline. Similarly, an activated region can contain the same pattern of activation across conditions. As such, they have often been explicitly contrasted (Peelen et al., 2006; Peelen and Downing, 2007; Jimura and Poldrack, 2012; Coustanche, 2013). In the present framework, the two methods are combined, rather than contrasted: patterns (i.e., representations) can be activated (i.e., involved) to different degrees.

The rationale of the framework is as follows. Consider patterns of activation in a region for eye, hand, and foot (Fig. 1D). Patterns for two of the three effectors can be equal (high correlation or low classification) or different (low correlation or high classification). Equal patterns indicate a common representation across effectors. Different patterns indicate a distinct representation per effector. With three effectors, a combination of distinct and common is also possible, e.g., eye versus hand and foot. For each pattern, its mean activation, as tested by univariate analysis, represents the pattern’s level of involvement. In our framework, patterns for each effector are weighted based on mean activation per effector: the more activated, the higher the combined value. This combination of pattern and activation again results in a separation of common and distinct representations (Fig. 1D). Representations, which are not activated, are effectively masked out because the low activation weight leads to negligible combination scores.

We first performed separate pattern information and activation analyses, followed by the combination of the two measures into a common pattern-activation measure. Following the conventions from MVPA (Kriegeskorte et al., 2006), we refer here to pattern information simply as “information,” although, in a mathematical sense, mean activation can also be considered information, albeit on a different spatial scale.
GLM. Both information and activation analyses were based on the output of a participant-specific GLM. Six regressors of interest captured variance during the planning period, specific to effector (three: eye, hand, foot) and target side (two: left, right). Side was included because PPC is highly sensitive to target direction (for review, see Silver and Kastner, 2009). Thirteen additional regressors were used to constrain the variance explained by the planning regressors: six location-dependent spike regressors for the information analysis, correlated pattern in t values between effectors, and for the activation analysis we averaged the values (Fig. 1A). We chose t values over β-values from the GLM because they are the more informative measure (Misaki et al., 2010).

Searchlight analysis. Both information and activation analysis were performed within a searchlight sphere (Kriegeskorte et al., 2006) with a radius of three voxels. Searchlight sphere size varied at the brain’s border and was on average 83 voxels. Searchlight analysis was performed in native space.

Pattern information analysis. Pattern information analysis was based on a correlation analysis (Haxby et al., 2001; Kriegeskorte et al., 2008) of the three effector pairs: eye–hand, eye–foot, hand–foot. The spatial distribution of effector-specific t values across a searchlight sphere was taken as local effector-specific pattern, which were then compared per effector pair using correlations. Low correlations between patterns reflect dissimilarity or information (akin to classification performance), meaning that patterns form distinct effector-specific representations. In addition, we take into account the alternative possibility: high correlations between patterns, indicative of similarity or lack of information, mean that patterns (for hand–man in Fig. 1D, distinct representation). However, combination with the effector-specific activation scores requires an effector-specific information score of one effector versus the other two (e.g., eye vs hand and foot, blue in Fig. 1D). First, we converted RGB values to HSB (hue, saturation, brightness) color space. Second, we rotated hue by 1/6π (effectively rotating color by 60° along the color circle, that is, half the distance between the colors of two respective effectors: green becomes cyan, blue turns magenta, and red transforms to yellow. Finally, we converted the rotated HSB color value back to RGB. After rotation, each of the three values (R, G, and B) indicated the amount of information on one effector versus the other two effectors in a [−1 1] range. We split the range in two [0 1] intervals to differentiate heuristically between common representations (similarity, left side of dissimilarity distribution, flipped sign) and distinct representations (dissimilarity, right side of dissimilarity distribution).

Activation analysis. Within a sphere, the activation analysis resembled an ROI analysis (Poldrack, 2007): we averaged the pattern of t values within a searchlight sphere for the effector-specific regressor of interest. Contrasts were first computed against baseline to detect common activations across effectors. We regard this as a measure of absolute activation, even though baseline measures are inherently relative in fMRI (Stark and Squire, 2001). We also calculated the mean t contrasts for one effector versus the other two effectors as a measure of relative activation. Both types of activation values were normalized to a [0–1] range by considering only positive values and dividing by the maximum value across effectors. For the combined pattern-activation analysis, we converted the two activation measures into one activation measure, consisting of three numbers (one for each effector) in a [0–1] range. The score was defined as the maximum of the absolute and relative activations per effector, indicating the region was activated, regardless whether it was absolute or relative.

Combination of pattern information and activation measures. We subsequently combined the pattern information and activation measures into a combined pattern-activation measure by elementwise multiplication of the activation scores with either the dissimilarity or similarity scores. Multiplying the activation scores with the full [−1 1] information range and splitting the range afterward gives the same result. The active common and distinct scores are the left and right side of the pattern-activation distribution, which allows a straightforward inspection of common and distinct representations. Subsequently we calculated square root values of the multiplied values to correct for multiplication damping. The multiplication of pattern information and activation scores attenuates nonactive patterns and increases the strength of active patterns. To determine areas encoding active distinct representations, we multiplied activation and dissimilarity scores. Thus, the distinction score will be high only when both activation and dissimilarity scores are high for a given effector. To determine areas encoding active common representations, we multiplied the activation scores with the similarity scores. As a result, the value for the common representation will be high only if information on one effector exceeds that of the other two (e.g., eye vs hand and foot, blue in Fig. 1D). The reference distribution of the permutation test was created by repeating the searchlight procedure using the same number of spheres and sphere sizes as in the
Table 1. The MNI coordinates of the regions used in the ROI analysis and their normalized peak values for activation, information, and combination for eye (E), hand (H), and foot (F)

<table>
<thead>
<tr>
<th>ROI</th>
<th>Reference</th>
<th>Peak</th>
<th>Activation</th>
<th>Information</th>
<th>Combination</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>x</td>
<td>y</td>
<td>z</td>
<td>x</td>
<td>y</td>
</tr>
<tr>
<td>pIPS</td>
<td>24</td>
<td>81</td>
<td>29</td>
<td>21</td>
<td>87</td>
</tr>
<tr>
<td>mIPS</td>
<td>19</td>
<td>69</td>
<td>57</td>
<td>21</td>
<td>63</td>
</tr>
<tr>
<td>aIPS</td>
<td>32</td>
<td>38</td>
<td>57</td>
<td>39</td>
<td>42</td>
</tr>
<tr>
<td>aSPL</td>
<td>27</td>
<td>42</td>
<td>70</td>
<td>24</td>
<td>54</td>
</tr>
<tr>
<td>aPCu</td>
<td>9</td>
<td>54</td>
<td>60</td>
<td>15</td>
<td>45</td>
</tr>
</tbody>
</table>

Coordinates were originally reported in Talairach space. Bold numbers indicate significant effects, p < 0.05, permutation test.

**Original Procedure**

original procedure, but with the voxel data permuted randomly across the entire brain. As in the original analysis, information, activation, and pattern-activation scores were calculated for each of three effectors within each searchlight sphere. The information scores were calculated by subtracting the mean information score. P values were calculated as the percentage of permuted spheres with a score higher (one-sided test, for activation) or different (two-sided test, for information and pattern activation) than that of the original, nonpermuted sphere, and determined separately for each effector. This procedure tests whether the continuous group of voxels in a search sphere gives a significantly different test score than a random set of voxels taken from the same brain.

**Group Analysis**

For analysis across participants, searchlight sphere results were first converted to MNI space after which information, activation, and pattern-activation scores were averaged across participants. Participant-level p values were accumulated by applying Fisher’s method (Fisher, 1925), which combines multiple p values into one aggregate p value based on a χ² distribution. Spheres were regarded significant for a given test if at least one of the three effectors was significant across participants.

**ROIs**

To facilitate reporting, and to allow direct linking of our results to previous, region-specific findings, we defined five ROI search spheres in PPC, corresponding to areas often implicated in motor planning (Levy et al., 2007; Filimon et al., 2009; Heed et al., 2011): posterior intraparietal sulcus (pIPS), medial IPS (mIPS), aIPS, anterior superior parietal lobe (aSPL), and anterior precuneus (aPCu).

Area pIPS, also known as V7 or IPS1 (Swisher et al., 2007), has previously been implicated in saccade control (Levy et al., 2007). Area mIPS, also referred to as IPS2 (Silver and Kastner, 2009), has been found active both for saccade and reach control (Levy et al., 2007; Heed et al., 2011). Area aIPS is involved in hand movements (Culham et al., 2003; Heed et al., 2011; Konen et al., 2013), whereas aPCu (within area 5L) seems to be involved in both hand and foot control (Filimon et al., 2009; Heed et al., 2011). Last, aSPL (area 5L7A) has recently been found to contain a shared representation for hand and foot (Heed et al., 2013).

The location of the ROIs was determined based on the peak of the activation for any of the effectors or combination thereof closest to the reference coordinates, restricted to fall within a posterior parietal and precuneus surface mask, as defined by FreeSurfer cortical parcellation. Thus, the ROIs are based on functional results and, hence, are used for descriptive purposes only (Kriegeskorte et al., 2009). We report the properties of the search sphere centered on the activation peaks of the respective areas. The complete overview of ROIs is shown in Table 1.

**Results**

The goal of this study was to characterize commonalities and distinctions in parietal effector representations involved in movement planning. Participants performed a delayed movement task involving goal-directed movements of the eye, hand, and foot. A novel searchlight-based pattern-activation approach (see Materials and Methods; Fig. 1) was developed to disentangle representations committed to the three effectors. Pattern information measures (dissimilarity and similarity) determined whether regions contained distinct or common representations for the different effectors. Activation measures tested the strength of the activation of these effector-specific patterns. The combined pattern-activation measure was used to delineate active distinct representations (significant activation as well as information) and active common representations (significant activation, but lack of information), revealing the active representational content of a region. We start with a description of the results of the left hemisphere, contralateral to the used limb.

**Pattern Information**

Pattern information analysis was applied to determine which representations are present in PPC, coding effector distinctions and/or commonalities (Figs. 3A, 4A).

Information was present within a parietofrontal network, including different types of effector-specific representations in PPC (Fig. 3A, straight arrow, top and bottom). Caudorostral along the IPS, we found a large and continuous, distinct representation related to eye movement planning (Fig. 3A, blue, left). This was combined with strong common representation of the hand and foot to different relative degrees (Fig. 3A, yellow/orange/light green, right). More specifically, pIPS, mIPS, and aSPL represented primarily an eye versus limb distinction (Fig. 4A, blue vs yellow colors).

Furthermore, we found a lateromedial hand-eye-foot information gradient in rostral PPC. The gradient for common representation ranged from foot and eye (Fig. 3A, cyan, right), to foot (green), to hand and foot (yellow) to hand (red), and to hand and foot (magenta). Note that the regional boundaries differ between the two information measures.

Regarding the predetermined ROIs (Fig. 4A), aIPS represented hand movements versus foot and eye movements (red vs cyan); aSPL represented the eye versus the limbs (blue vs yellow), and aPCu represented foot versus hand and eye movements (green vs magenta). The distinct representation of aIPS and aPCu continued well into lateral and medial somatosensory and motor cortex, respectively.

In summary, information analysis revealed a caudorostral distinct eye representation and a rostral, lateromedial distinct hand-eye-foot representational gradient, combined with mostly common representations for the complementary effectors. Thus, most PPC areas represented a dichotomy between a distinct representation for a single effector and a representation common to the other two effectors.
Whereas the information analysis revealed distinct and common representations across PPC, it does not show which of the representations of a given area was most activated by the task, and whether, perhaps, some representations were on average not activated at all. The activation analysis examined such regional involvement (Figs 3B, 4B).

Activation

Whereas the information analysis revealed distinct and common representations across PPC, it does not show which of the representations of a given area was most activated by the task, and whether, perhaps, some representations were on average not activated at all. The activation analysis examined such regional involvement (Figs 3B, 4B).

Activation of the different effectors against the fixation baselines was evident widely across a parietofrontal network, showing strong involvement of the PPC (Fig. 3B). Within PPC, there was a caudorostral eye-to-limb gradient: caudally, eye activation was dominant, whereas more rostral regions were more activated by all three effectors, both in a shared fashion (Fig. 3B, gray-white colors, left, straight arrow), and separately (Fig. 3B, red and green). At the level of the preselected ROIs (Fig. 4B), pIPS showed activation for eye movements only, mIPS demonstrated also activation for the other two effectors, and aSPL showed comparable activations for the three effectors.

We also observed a rostral, lateromedial hand–foot gradient: a hand bias (Fig. 3B, red, curved arrow) was evident lateral to aSPL, a foot bias (green) more medially, whereas all three effectors were represented in the in-between region (yellow or gray-white). For the ROIs (Fig. 4B) along the postcentral sulcus, aIPS was predominantly activated for the hand, aSPL for the hand and foot, and aPCu for the foot. In aIPS the dominant hand activation was accompanied by activation for the foot and eye, and in aPCu the dominant foot activation was accompanied by activation for the hand. Thus, all rostral PPC ROIs were activated for multiple effectors. The relative hand and foot dominance in aIPS and aPCu continued in lateral and medial somatosensory and motor cortex.
In sum, the activation analysis revealed a caudorostral eye–limb gradient and a lateromedial hand–foot gradient, combined with significant secondary activations in rostral PPC.

**Combined activation and information**

The information-based analysis revealed mostly dichotomies of distinct and common effector representations across PPC, but did not allow conclusions about which representations were actually activated by the task. Activation analysis, on the other hand, showed general activation with relative effector-specific peaks, but did not allow conclusions about whether the general activations held effector-specific representations. Combining the two measures in the pattern-activation framework allows examination of both of these aspects simultaneously, that is, to observe which unique representations are present as well as which of them are activated by the task (Figs. 3C, 4C).

The combined pattern-activation analysis revealed active distinct representations across the frontoparietal network, including the PPC (Fig. 3C, left). Caudal PPC was dominated by eye-related representations, whereas rostral PPC contained active distinct representations for the hand and the foot, separated by an active representation distinct for planning eye movements. Note that all combined activation (Fig. 3B, gray, left) has been filtered out, revealing clear patches of representations. ROI analysis revealed that pIPS, mIPS, and aSPL all actively distinguished movements of the eyes, whereas aIPS encoded movements of the hand and aPCu represented planned movements of the foot (Fig. 4C).

Of the common representations revealed by the information analysis (Fig. 3A, right), especially the rostral PPC representations were activated (Fig. 3B), resulting in rostral patches of active common representations (Fig. 3C, right). Specifically, the combined analysis uncovered three distinct representations that were hidden in the overlapping activations: an activated limb representation (Fig. 3C, right, yellow/orange), foot and eye representation (green/cyan), and hand representation (red). The selected ROIs are covered by these patches (Fig. 4C): in addition to the active distinct representations, mIPS contained an active common representation for the limbs (circle diagram, yellow), aIPS coded an active common representation for the foot and eyes (green and cyan), and aPCu contained a large activated hand representation (red). Furthermore, in aSPL, we found an integrated hand–foot representation at the cross section of the aIPS and aPCu regions.

Both active distinct and active common representations were found within the same regions in PPC (compare Figs. 3C, left and right, Fig. 4C), but with different amounts of activation. For most regions, the distinct representation was also the most activated representation, that is, most regions were dominated by one effector. aSPL formed an exception, in that both the eye and limb representation were equally activated (dominance indicated by brightness of colors in Fig. 3C, size of dot in 4C).

In summary (Fig. 5), the pattern of representations for effector planning in PPC emerging from the combined information-activation framework is mostly that of active dichotomies: mIPS, eye versus the limbs (most activated first); aSPL, limbs versus eye (equal dominance); aIPS hand versus eye and foot; aPCu, foot versus hand. Area pIPS appears as an exception, with only an active representation of the eye.

**Right hemisphere**

Analysis was focused on the contralateral (left) hemisphere, which is thought to be most involved in coding right hand and foot movement (Beurze et al., 2009; Gallivan et al., 2011; Heed et al., 2011). However, the hemisphere ipsilateral to the moving limb can also contain information about planned movements (Gallivan et al., 2013). As Figure 6 shows, eye-specific information was present ipsilaterally in caudal PPC, and hand- and foot-specific information was present in the motor cortex. In rostral PPC regions, effector-specific informational content was reduced compared with the contralateral hemisphere. Furthermore, activation of the right hemisphere was attenuated compared with the left hemisphere, particularly in motor and somatosensory cortices. The combined analysis revealed that the eye-specific posterior representations were active and distinct, whereas no clear,
active distinct limb-specific representations emerged in rostral PPC. Similarly, we did not observe any strong active common representations.

Thus, the only representation active in ipsilateral PPC was for the one effector for which the brain is not lateralized: the eyes.

Control measures

Relation activation and information

Results of the pattern information and activation analysis show a remarkable consistency: in a given region, the most distinct representation was, at the same time, the most strongly activated. However, this consistency may in fact be artifactual: more strongly activated voxels may dominate the correlation calculation and thus increase the likelihood that higher activated representations turn out distinct. If this is the case, positive correlations between information and activation should also be present in the random data generated in the permutation test. If not, it should be unique to the actual results, more specifically to the effectors represented in each hemisphere: all effectors in the left hemisphere, only the eyes in the right hemisphere. We tested this potential confound by comparing the relationship between mean relative activation per effector and information per effector in the actual data, separately for each hemisphere, with the same measures calculated for random data.

In the left hemisphere, correlations between activation and information were positive for all three effectors (mean correlation: 0.68 (hand), 0.39 (foot), 0.47 (eye), all \( p < 0.05 \)). In the right hemisphere, the correlations dropped for the hand (\( 0.02, p > 0.05 \)), but not for the eyes and foot (0.38 and 0.45, respectively, both \( p < 0.05 \)). In contrast, for the random data, activation and information correlated negatively (\( r = -0.19, -0.18, -0.09, \) all \( p < 0.05 \)). Thus, we conclude that the higher relative activations predict higher information measures, but that the relationship is not due to a confound in our analyses.

Unsmoothed group GLM

All our analyses are based on tests using search spheres of 83 voxels on average, which is comparable in size to a standard smoothing kernel. It is possible that this approach has hindered precise spatial localization, with spheres capturing information and activation from neighboring areas. To control for such effects, we performed a standard group GLM on spatially unsmoothed data (Fig. 7). We predicted that there should be effector-specific voxels within PPC areas, validating the spatial sensitivity of our analyses.

The prediction was confirmed, as activation patterns were consistent with those determined by the searchlight procedure: aIPS was most active for hand planning; mIPS was most active for eye planning; and aPCu was most active for foot planning. Unsurprisingly, activations were more scattered, and patches were not as consistent as those detected by the pattern-activation analysis, which underlines the high sensitivity of the pattern-activation analysis for regional representations.

Discussion

This study characterized distinctions and commonalities across effector representations in PPC. The motor planning responses of most regions were dominated by one particular effector: eye movements in pIPS, hand movements in aIPS, and foot movements in aPCu. The new observation is that several PPC regions also actively coded for commonalities between the other effectors, distinct from the dominant effector. This finding provides empirical evidence for the notion that effector selection is based on an efficient neural code that distinguishes an effector from other potential effectors (Medendorp et al., 2005; Cui and Andersen, 2011).

Pattern-activation gradient

The pattern-activation measure revealed a caudorostral gradient in PPC. This finding extends previous reports, showing that activation of a particular region by the planning of several effectors (Levy et al., 2007; Heed et al., 2011) does not necessarily imply identical representational content. Furthermore, we extend previous MVPA studies (Gallivan et al., 2011a, b; Gallivan et al., 2013) by specifying, per region, the dominant representation and testing for differences between limbs, instead of only eye–hand or hand–hand distinctions.

Previous reports have shown that, along the caudorostral axis, pIPS is informative for the eye–hand distinction (Gallivan et al., 2011). Here we show that the eye representation is dominant, that there is no information on the distinction between the hand and foot (limbs), and that limb movements do not lead to activation. We hence conclude that pIPS is involved in saccade planning (Levy et al., 2007). Following the same logic, the pattern-activation findings suggest that areas mIPS and aSPL specifically distinguish between eye and limbs. Eye, hand, and foot planning all activate these regions, but the informational content is restricted to a distinction between eyes and limbs, but not between the limbs. Only the distinct representations for eye movements are active in both hemispheres, in line with the bilateral organization of saccade preparation (Sereno et al., 2001).

In rostral PPC, aIPS coded for the hand and aPCu coded for the foot. The common representations underlying this dominant hand–foot gradient largely complemented the dominant representations: the foot area also represents the hand and the hand area also represents the foot and eye. This corepresentation of the different limbs may explain why others have found hand-tuning in the aPCu region (Fernandez-Ruiz et al., 2007; Filimon et al., 2009), which we find to be dominated by foot planning (Heed et al., 2011). Importantly, the combined pattern-activation measure allowed us to detect two separate representations in aPCu (one foot, one hand), of which the foot representation is dominant in activation. We hence conclude that aPCu is predominantly responsible for foot movement planning, while aIPS is responsible for hand movement planning.
In addition to motor planning, rostral PPC is important for somatosensation (Sereno and Huang, 2014). The rostral effector gradient may hence stem from somatosensory activations. However, the current analysis specifically focused on the planning period, during which there is no somatosensory feedback. Rather, the activation could be related to the prediction of the somatosensory consequences of the movement from the start of the trial onward (MacKay and Crandall, 1987), consistent with the effector-specific involvement of SI and SII in the current task (SII not shown; Eickhoff et al., 2007) and the predictive coding framework (Friston and Kiebel, 2009). Furthermore, knowing the position of the hand and foot, in addition to their target positions, is essential for the planning of movements with the respective effectors (Buneo et al., 2002; Beurze et al., 2009). Such an integrative role would fit with the position of the regions next to the somatosensory homunculus (Seeleke et al., 2012), in line with continuous topographic coding (Graziano and Aflalo, 2007a), similar to the location of the parietal saccade areas near regions containing visual maps (Aflalo and Graziano, 2010).

Nature of coding
The continuous topographic organization of different combinations of active distinct and active common effector representations suggests a more complex organization than modular effector specificity. The common representations could code movements with similar computational constraints (cf. Heed et al., 2011), and the combination of distinct and common representations might allow combining effectors toward a common functional goal (Graziano and Aflalo, 2007b); e.g., hand and foot in aPCu for walking or climbing (Abdollahi, 2013). Such classes of movement based on coding commonalities and identities may represent a more efficient organization of motor preparation than effector-specific coding per se (Levy et al., 2007; Jastorff et al., 2010).

The fact that the rostral planning-related PPC representations are consistently dichotomous suggests that these representations could additionally be used in the selection of effectors or classes of movements. For example, in single-effector movements, aPCu could select foot movements over hand movements, while in actual climbing or walking it could represent both effectors (Graziano, 2006). This would imply that rostral PPC regions can make effector “choices,” rather than representing one or the other effector (Cisek and Kalaska, 2010; Cui and Andersen, 2011). The activated distinct representation would then be related to the selected effector, the less-activated common representation to the alternative effector(s). Which alternative effectors are considered could depend on the class of movement represented in a region (e.g., walking or climbing in aPCu: foot and hand) and/or on the combination of effectors tested in the task. Whether the basis of selection is single effectors or classes of movement could be studied by testing movements involving combinations of effectors or movements belonging to different movement classes.

Alternatively, the selection process could reflect motor inhibition of the alternative effectors (Brown et al., 2006). In the present study, participants were explicitly instructed to move only one effector, while, in nonexperimantal circumstances, some effectors (e.g., eye and hand) are often moved together. Moreover, the task context itself, with equal probability for each effector, might have induced coactivation of effectors based on contingency (Chang et al., 2008; Gallivan et al., 2013). Testing movement planning with different effector probabilities should allow the separation of these accounts.

Finally, the active common representations could be caused by a general effect, such as spatial attention. We cannot rule this out, given that we did not test non-motor trials, but the specificity of our analysis to the delay period, the specific parietofrontal representations in the ipsilateral hemisphere, and the rich gradient of secondary representations all speak against a general origin of the reported common representations.

Pattern-activation framework
The pattern-activation framework makes explicit the notion that information and activation are complementary measures (Peelen and Downing, 2007; Jimura and Poldrack, 2012). Patterns are seen as representations contained in an area (MVPA assumption), whereas the mean activity shows how involved the region is in representing the condition (univariate analysis assumption). In essence, the framework assumes that patterns can be active to differing degrees, allowing quantification of the representation’s involvement in the current cognitive process: we interpret weakly activated representations as weakly involved in the task. For example, limb representations in the ipsilateral hemisphere were activated for the current task to some extent, but our framework makes the testable prediction that comparable representations should be evident, though at higher activation, during movement planning of the left limbs.

There are several alternatives with which the pattern-activation framework may be implemented. An alternative for the information measure (that is, correlation) is classification (Cox and Savoy, 2003). The advantage of a correlative measure as implemented here is the possibility to test whether two patterns are less distinct than expected by chance. This allowed us to identify common representations; in contrast, in a classification approach, nondistinctiveness would be a null result. Alternatively, we could have used cross-classification: if patterns for two classes are indistinguishable (low interclass classification score), they could either be noise (no cross-classification possible) or truly the same pattern (cross-classification possible). The current results have large predictive power for such classification results. Another alternative for correlation is component analysis (Beckmann et al., 2005). This approach has the attractive feature of allowing detection of partly shared representations, such as a common hand–foot representation and a distinct hand representation within the same region. Any alternative to the activation measure should be based on a clean baseline and ideally be sensitive to both absolute and relative activation effects.

Activation is essentially information at a different spatial scale. If one took a larger ROI, some activation results would translate into information results, as they would become a subpart of the region, and vice-versa, a smaller ROI would lead to a shift from information to activation results, when local patterns no longer activate parts but the whole region. The combined measure we propose allows taking this into account by combining the two measures.

Conclusion
We conclude that PPC represents effector dichotomies of distinct and common representations. The distinct representations are organized along a posterior-to-anterior, visuosomatic gradient, which splits into a second, lateral-to-medial hand–foot gradient in rostral PPC. This appears to reflect a topographically continuous organization, from visual perception and saccades, along lateral hand and medial foot planning pathways, to the somatosensory hand and foot regions. The combination of distinct and common representations in rostral PPC makes this region ideally
suited to function as a selector, either of effectors or of classes of movements.

References


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