Anchoring the neural compass: coding of local spatial reference frames in human medial parietal lobe

Steven A Marchette, Lindsay K Vass, Jack Ryan & Russell A Epstein

The neural systems that code for location and facing direction during spatial navigation have been investigated extensively; however, the mechanisms by which these quantities are referenced to external features of the world are not well understood. To address this issue, we examined behavioral priming and functional magnetic resonance imaging activity patterns while human subjects recalled spatial views from a recently learned virtual environment. Behavioral results indicated that imagined location and facing direction were represented during this task, and multivoxel pattern analyses indicated that the retrosplenial complex (RSC) was the anatomical locus of these spatial codes. Critically, in both cases, location and direction were defined on the basis of fixed elements of the local environment and generalized across geometrically similar local environments. These results suggest that RSC anchors internal spatial representations to local topographical features, thus allowing us to stay oriented while we navigate and retrieve from memory the experience of being in a particular place.

To be oriented in the world, an organism must know where it is and which direction it is facing. In rodents, location information is encoded by place and grid cells in the hippocampal formation1,2, and directional information is coded by head direction (HD) cells in Papez circuit structures3; recent work has described similar spatial codes in these regions in humans4–8. These cellular populations are coordinated with each other, such that if the directional signal coded by HD cells rotates, the firing fields of place and grid cells rotate by a corresponding amount (an example of this is reported in ref. 9). This suggests that HD cells support an internal compass that represents an animal’s heading (i.e., facing direction) and updates this quantity as the animal moves.

However, for a neural compass to be useful, it must do more than represent direction in arbitrary coordinates—the heading must be defined relative to fixed features of the environment, just as the heading revealed by a magnetic compass is defined relative to the north-south axis of the earth. This presents a challenge: in the absence of magnetoeception or a sidereal or solar compass, there is no single perceptible feature that consistently indicates direction across all terrestrial environments. A possible solution is to use one’s perceived orientation relative to local (currently visible) landmarks to anchor one’s sense of direction, which can then be maintained during navigation through idiographic cues10–12. To do this, however, one must have a representation of one’s heading relative to local environmental features that is at least potentially separable from the internal sense of direction supported by the HD cells. Although there is considerable evidence that both rodents and humans use allocentric information to orient themselves, a neural locus for this separate representation of locally referenced facing direction has not yet been identified.

On the basis of previous neuroimaging and neuropsychological results4,13–16, as well as theoretical models17,18, we hypothesized that a medial parietal region referred to here as the RSC might support this locally referenced representation of heading, thus providing a dial for the neural compass. Moreover, we hypothesized that recovery of this heading representation might be an essential element of spatial memory retrieval. To test these ideas, we collected behavioral and functional magnetic resonance imaging (fMRI) data while subjects re-instantiated views of a newly learned virtual environment during performance of a judgment of relative direction (JRD) task. This task requires subjects to imagine themselves in a specific location facing a specific direction, which means that they must mentally reorient themselves (i.e., re-establish their sense of direction) in each trial. Critically, the virtual environment was divided into separate ‘museums’ that had distinguishing visual features but identical internal geometries, which were set at different angles relative to each other within a larger ‘park’ (Fig. 1a). This design allowed us to distinguish between three kinds of spatial representations: (i) spatial representations that used a single global reference frame that applied across the entire environment, (ii) spatial representations that used locally anchored reference frames that were unique to each museum and (iii) spatial representations that used locally anchored reference frames that generalized across different museums with similar geometry.

Previous behavioral work with the JRD task has found evidence for coding of locations and directions in spatial reference frames aligned to local features of the environment such as room geometry or the arrangement of objects within a room (and also in frames aligned to egocentric axes when external features are absent)19,20. Our primary concern was to test the hypothesis that RSC (or other brain regions) might mediate these locally referenced spatial frames. To this end, we used behavioral priming (experiment 1) and similarities and differences between multivoxel patterns elicited in different retrieval trials (experiment 2) to identify location and direction codes and then tested whether these spatial representations were aligned to the local...
features. Our results indicate that RSC codes for facing direction and location of imagined viewpoints and that these are coded in a reference frame that is anchored to environmental features but generalizes across local environments with similar geometry.

RESULTS

Behavioral evidence for direction coding

We first used cognitive behavioral testing to establish that directional codes are used during our version of the JRD task (location codes are considered below). In experiment 1, participants learned a virtual environment consisting of four museums, which were visually distinguishable but had similar internal geometry (Fig. 1b). After reaching a criterion during a training phase in which subjects were required to navigate from a starting point outside the museums to individual objects within (Online Methods), they were tested on their knowledge of the objects’ locations. In each trial, the subjects were verbally cued to imagine themselves facing a reference object (Fig. 1c), and they used the keyboard to indicate whether a target object would be to their left or right. Because each object was placed in such a way that it could be viewed only from a specific direction, specifying a reference object implicitly specified both a facing direction and a location (Fig. 1b,c). Participants responded accurately in most trials (mean 89.8% correct, s.d. 6%). Only correct trials were entered into analyses of reaction time.

We hypothesized that directional representations used for mental orientation in each trial would be revealed through behavioral priming; specifically, reaction times would be speeded in trials that had the same implied direction as the immediately preceding trial. For this first set of analyses, we defined direction locally. That is, “north” for each museum was defined as the direction facing the wall opposite the doorway (Fig. 2), and repetition of this orientation (or other orientations) across museums was considered to be a repetition of the same direction. To ensure that priming effects were attributable to repetition of direction rather than repetition of location, we excluded from analysis cases in which the facing object in the immediately preceding trial was drawn from the same corner of the same museum or the geometrically equivalent corner of a different museum (Fig. 2a). This restriction imposes a criterion of abstractness on direction coding: representations of imagined direction must generalize across different views and different locations.

A repeated-measures analysis of variance (ANOVA) with direction (same or different as the preceding trial) and museum (same or different as the preceding trial) as factors revealed a main effect of direction (F(1,21) = 11.63, P = 0.003), with faster reaction times when the imagined direction was repeated over successive trials (3.31 ± 0.20 s (mean ± s.e.m.) for same compared to 3.56 ± 0.18 s for different) (Fig. 3a). We also observed a main effect of repeating museum (F(1,21) = 11.80, P = 0.002), with faster reaction times when reference objects in successive trials were located within the same museum (3.27 ± 0.19 s for same compared to 3.59 ± 0.19 s for different). There was no significant interaction between direction and museum (F(1,21) = 0.73, P = 0.401). The absence of an interaction is important because it suggests that direction was defined consistently across all four museums. That is, “local north” in one museum was equivalent to local north in another, even though these were different directions in the global reference frame. In contrast, if direction had been coded in a local manner that was unique to each museum or had used a global system that applied across all museums, then we would not have seen this equivalence between locally defined directions.

These data indicate that participants represented their facing direction during the JRD task and updated this representation when the direction changed from trial to trial, thus leading to a reaction time cost. Moreover, this representation of facing direction generalized across different museums with similar local geometry. Further analyses indicated that reaction times were faster for imagined views facing the wall opposite the doorway (Supplementary Fig. 1), suggesting that this direction served as a ‘conceptual north’ within each environment and providing additional evidence for spatial codes that vary with direction.
Figure 2 Summary of the analysis scheme. (a) Contrasts used to test for coding of facing direction. Top, all views for two of the four museums; views that face the same direction as defined by the local museum frame are the same color. Middle, comparisons between view 1 and other views that face the same or different local direction within and across museums. To partially control for location, view 1 is never compared to views in the same corner (i.e., views 1, 8, 9 and 16 are excluded). Bottom, test for direction coding that controls completely for location: in this case, the same-direction comparison view is located in the same corner as the different-direction comparison view. (b) Contrasts used to test for coding of location. Top, all views for two of the four museums; views located in the same corner of the environment (defined by the local museum frame) are the same color. Middle, comparisons between view 1 and other views located in the same or different corners within and across museums. To partially control for direction, view 1 is never compared to views facing the same local direction (i.e., views 1, 2, 9 and 10 are excluded). Bottom, test for location coding that controls completely for direction: in this case, the same-location comparison view faces the same direction as the different-location comparison view.

Behavioral evidence for location coding
When reorienting themselves in each trial of the experiment, participants established an imagined location in front of the reference object in addition to establishing an imagined direction. We hypothesized that, as with facing direction, this location code would reveal itself through priming. For this analysis, objects in the same corner of a museum were considered to be in the same location, as were objects in geometrically equivalent corners in different museums (where geometrical equivalence was defined in terms of the local reference frame—thus, for example, the corner opposite the door on the left in one museum was considered equivalent to the corner opposite the door on the left in the other museums). To ensure that priming effects reflected repetition of location rather than repetition of direction, we only analyzed cases for which the implied direction differed across successive trials (Fig. 2b).

A repeated-measures ANOVA with location (same or different) and museum (same or different) as factors found a significant main effect of location \( (F(1,21) = 5.695, P = 0.026) \), indicating that—as predicted—reaction times were faster when the imagined location was repeated over successive trials \( (3.56 \pm 0.20 \text{ s for same compared to } 3.31 \pm 0.20 \text{ s for different}) \) (Fig. 3b). There was a marginal effect of repeating museum \( (F(1,21) = 4.227, P = 0.052) \). Critically, there was no interaction of location repetition and museum repetition \( (F(1,21) = 0.012, P = 0.915) \), indicating that priming occurs between geometrically equivalent locations, not only when these locations are physically the same but also when they are different locations in different museums. These results showed that location was represented in this task as well as direction and—like direction—was coded using a locally anchored reference frame that generalizes across geometrically equivalent environments.

Comparing local and global reference frames
Although our results strongly implicate coding with respect to local reference frames, we also tested whether there might also be residual coding of location and direction within the global reference frame. Because museums were oriented orthogonally or oppositely within the park, it was possible to dissociate local and global spatial quantities by examining response priming across trials from different museums.

To test for global coding of direction, we divided trials in which museum and direction both differed from the preceding trial into two groups: (i) trials for which local direction differed from that in the immediately preceding trial but global direction was the same (for example, in Fig. 2a, views 1 and 15 face different local directions but both face global north) and (ii) trials for which the local and global direction both differed. There was no difference in reaction time between these two trial types \( (t(21) = 1.283, P = 0.214) \); hence, there was no evidence of global direction priming (Fig. 3a). Indeed, the trend was in the opposite direction: reaction time was slower when global direction was repeated \( (3.74 \pm 0.20 \text{ s}) \) than when it changed \( (3.64 \pm 0.18 \text{ s}) \). To test for global coding of location, we performed an analogous analysis. That is, we divided different-museum and different-location trials into (i) trials for which the location was in a different corner of the museum as defined by the local frame but in the same corner as defined by the global frame and (ii) trials for which there was no spatial equivalence in either the local or global frame. There was no significant difference in reaction times between these two kinds of trials \( (t(21) = 0.976, P = 0.340) \), and hence there was no evidence for location priming in the global frame (Fig. 3b).

We then considered the strongest case for priming in the global reference frame by examining trials for which both location and direction, defined in this frame, were repeated across museums. These were compared to trials for which both location and direction defined locally were repeated across museum. For example, we compared reaction times when view 1 was followed by view 15 (repetition of global location and direction) to reaction times when view 1 was followed by view 9 (repetition of local location and direction). Reaction times were significantly faster for repetitions of the locally analogous view than for repetitions of the globally analogous view \( (t(21) = 4.242, P = 0.00036) \), once again suggesting that location and direction were coded in the local rather than global reference frame.

Directional coding in fMRI
We next turned to fMRI to understand the neural basis of the spatial codes revealed in the first experiment. Participants in experiment 2 underwent a modified version of the training procedure (Online Methods) and then performed the JRD task in the scanner while fMRI data were obtained. To keep the scan sessions to a manageable length of time, locations being tested were drawn from only two of the four museums shown at training; in addition, the timing and sequence of trials were adjusted to maximize fMRI signal detection. Most other aspects of the testing procedure were identical between the two experiments (Online Methods).
Participants achieved a high level of accuracy in the JRD task in experiment 2 (mean 94.4% correct, s.d. 6%). Reaction times were significantly faster in experiment 2 than in experiment 1, probably because fMRI participants received an additional training session on the day of the scan (2.60 s compared to 3.39 s, unequal sample sizes t-test: t(44) = 4.599, P = 0.00008). Directional and location priming effects were not significant in experiment 2 (direction: F(1,23) = 1.501, P = 0.233; location: F(1,23) = 1.190, P = 0.287), but other behavioral effects were consistent across the two experiments (Supplementary Fig. 1). Although this lack of priming might suggest that qualitatively different representations were used to solve the task in the two experiments, a more plausible explanation is that priming effects were weakened in experiment 2 by the use of a longer intertrial interval (Online Methods). As fMRI data did not depend on obtaining differences in reaction times, we proceeded to look for neural signals corresponding to coding of direction.

We focused our initial analyses on RSC, as previous work has suggested that this region might be critically involved in the coding of spatial quantities such as facing directions. This region was functionally defined in each subject on the basis of greater activity to scenes than to objects in independent localizer scans. To investigate whether RSC represented participants’ imagined direction, we calculated similarities between multivoxel activity patterns elicited by all 16 possible views (8 reference objects × 2 museums) across two independent data sets corresponding to the first and second half of the experiment. We then compared the pattern similarity for views facing the same locally defined direction to the pattern similarity for views facing different directions. As in the behavioral experiment, we only analyzed view pairs for which the implied location was different (Fig. 2a). Thus, same-view pairs were excluded, as were view pairs that faced different directions from the same location (i.e., views in the same corner).

The results indicated that RSC coded imagined facing direction (Fig. 4a). A two-by-two repeated-measures ANOVA with factors for direction (same or different) and museum (same or different) found a significant main effect of direction (F(1,23) = 20.009, P = 0.00017). There was no effect of museum identity (F(1,23) = 2.450, P = 0.131) and no interaction between direction and museum (F(1,23) = 0.005, P = 0.942). Thus, just as in experiment 1, direction was coded in a manner that generalized across equivalent local directions in different museums. These data suggest that RSC encodes imagined facing direction during memory retrieval and does so in a way that is referenced to the local environment and is consistent across geometrically similar environments.

The analysis described above potentially confounds direction coding with location coding, because implied locations were, on average, closer together for same-direction views than for different-direction views. To ensure that the results described above truly reflected coding of direction, we performed an additional test of direction coding for which location was controlled completely. Specifically, we compared each view to two other views, both of which had the same implied location, one with the same implied direction as the original view and the other with a different implied direction (Fig. 2a). We submitted these controlled comparison pattern similarities to a two-by-two repeated-measures ANOVA with direction and museum as factors. Even when controlling for location in this manner, we observed greater pattern similarity for same-direction views than for different-direction views (F(1,23) = 8.279, P = 0.008), with no main effect of museum (F(1,23) = 0.850, P = 0.366) and no interaction of direction and museum (F(1,23) = 1.043, P = 0.318).

Location coding in RSC

In the analyses described above, we examined direction coding while controlling for location to ensure that the findings could not be explained by location coding. However, the results of these analyses do not preclude the possibility that both location and direction might be encoded. To test this possibility, we compared pattern similarities for views obtained in the same location (i.e., the same corner) to pattern similarities for views obtained in different locations, with geometrically equivalent corners in different museums (as defined by the local reference frame) counting as the same location. As in experiment 1, we restricted this analysis to pairs of views that differed in direction (Fig. 2b). A two-by-two repeated-measure ANOVA with location and museum as factors found a main effect of location (F(1,23) = 7.162, P = 0.013), indicating that patterns were more similar for views in the same (or geometrically equivalent) corner than for views in different corners (Fig. 4b). There was no main effect of museum (F(1,23) = 2.719, P = 0.113) and no interaction of location with museum (F(1,23) = 0.284, P = 0.600). Thus, location is encoded in RSC; moreover, like direction, it is encoded using a local reference frame that is anchored to each museum and generalizes across museums.

Although the comparisons described above were restricted to views facing different directions, a potential confound is that views in the
same corner face more similar directions than views in different corners. For example, views in different corners can differ by 180 degrees, whereas views in the same corner can differ by, at most, 90 degrees. To control for this, we performed an additional analysis in which we only considered pattern similarities between views that differed by 90 degrees (Fig. 2b). A two-by-two ANOVA revealed a main effect of location (F(1,23) = 4.581, P = 0.043), no main effect of museum (F(1,23) = 1.834, P = 0.189) and no interaction of location by museum (F(1,23) = 0.768, P = 0.390). Thus, just as direction coding can be observed in RSC when location is controlled completely, so can location coding be observed when direction is controlled completely.

Comparing local and global reference frames in RSC

Our analyses suggest that RSC coded both direction and location in the local reference frame. Following the same logic as that in the behavioral analyses of experiment 1, we then tested for residual coding of global direction and global location. First, we calculated the average similarity between views in different museums that faced the same global direction but different local directions and compared this to the average similarity between views in different museums that differed in both global and local direction. This analysis revealed no difference in pattern similarity relating to global direction (t(23) = 0.724, P = 0.476) (Fig. 4a). Second, we calculated the average similarity between views in different museums that were in the same location as defined by the global reference frame, and we compared this to the average similarity between views in different museums whose locations were noncorresponding in both reference frames. We found no difference between these conditions (t(23) = 0.965, P = 0.344) (Fig. 4b), a result that is once again consistent with an absence of global coding.

We then compared the pattern similarity for views that share both location and direction across museums, as defined by either the local or global reference frame. Pattern similarity was greater for locally analogous views than for globally analogous views (t(23) = 3.495, P = 0.002). In sum, our results indicate that RSC coded spatial quantities that were referenced to the internal geometry of the museums rather than to the global geometry of the park.

Visualization and reconstruction of spatial similarities

To better understand the nature of the spatial representations revealed above and qualitatively assess the full pattern of spatial coding in RSC, we visualized similarities between views by overlaying them on a map of the environment (Fig. 5). To create this map, we first created a pattern similarity map for each of the 16 views that indicated the similarity between that view (starting view) and the 15 other views (comparison views). We then combined these maps by aligning the starting views to one another using rotations and reflections that maintained the relative spatial relationships between the starting and comparison views. We then averaged the aligned maps to produce a composite similarity map.

Inspection of the similarity map confirmed and extended the previous findings. The starting view was maximally similar to itself (scaled to 1.0), indicating consistency of the response pattern across the first and second half of the experiment. Within a museum, the most similar nonidentical views were the view facing the same direction in the adjacent location and the view facing the orthogonal direction in the same location. Across museums, the most similar views were the locally analogous view and the view facing the same local direction; there was also some similarity for the view facing the orthogonal direction in the equivalent location. Comparison views in adjacent corners to the starter view were more similar than views in the opposite corners, indicating the possibility that RSC may represent location in a continuous gradient corresponding to distance or may represent discrete boundaries shared among views.

To evaluate these last two possibilities, we analyzed similarities between views in terms of the number of shared boundaries (same corner: two shared boundaries; adjacent corner: 1 shared boundary; opposite corner: 0 shared boundaries), excluding view pairs that faced the same direction. Because the number of shared boundaries is highly correlated with Euclidean distance, these two factors are virtually indistinguishable in our current design. A three-by-two repeated-measures ANOVA with factors for shared boundaries and museum (same or different) found a significant main effect of shared boundaries (F(2,46) = 6.675, P = 0.003), a marginal main effect of museum (F(1,23) = 3.200, P = 0.087) and no interaction between the two (F(2,46) = 0.316, P = 0.730). The main effect of shared boundaries was explained by a significant linear contrast (F(1,23) = 8.615, P = 0.007), reflecting the fact that pattern similarity increased continuously as a function of the number of shared boundaries and/or distance between the views. This sensitivity to boundaries raised the question of whether RSC represents the structure of the environment near the imagined view rather than the heading and location of the imagined view relative to that structure. Whereas location coding would be predicted under both accounts, direction coding would be predicted only under the latter. To examine this idea, we compared...
the main effects of direction and location from the earlier analyses. A paired \( t \) test confirmed reliably stronger coding of facing direction than of location (\( t(23) = 2.652, P = 0.014 \)), suggesting that RSC is more concerned with situating imagined views relative to local geometric structure than in representing that structure itself.

These results provide another demonstration that activity patterns in RSC contain rich information about the implied headings and locations of imagined views that could be used to link views to long-term spatial knowledge and, thus, to localize and orient oneself within a mental map of local space. To further test this idea, we examined whether pattern similarities in RSC were sufficient to reconstruct the relative positions of views within a museum. We calculated each subject’s correlation matrix for the eight different views that occurred within the same museum and then averaged them to produce a grand correlation matrix. This grand correlation matrix was submitted to multidimensional scaling to produce a two-dimensional map of the views, which was then aligned with a Procrustes analysis to a schematic of our museums (Fig. 6a). Patterns of similarity in RSC created a reconstruction of the views within a museum that was significantly less distorted than predicted by chance (\( D = 16\%, P = 0.00008 \)). To assess the regional specificity of this finding, we repeated this procedure with the other functionally defined regions of interest (ROIs) (parahippocampal place area, occipital place area and early visual cortex); none was able to perform reliable spatial reconstruction (all \( D > 58\%, \text{all } P > 0.1 \)).

We then performed the same analysis on the across-museum view similarities (Fig. 6b). Again, the reconstruction was significantly less distorted than predicted by chance (\( D = 47\%, P = 0.015 \)), although it tended to be less accurate than the reconstruction based on within-museum view similarities (difference in \( D = 31\%, P = 0.09 \)). Successful spatial reconstruction across museums suggests that RSC’s representation of views in one museum can accurately predict the relations among views in another museum. This generalization is consistent with the idea that RSC represents views relative to the frame of the local environment and can generalize these relationships to different environments with similar geometry.

Whole-brain analyses

Having discovered RSC representations that coded for direction and location within the local space in our ROI analysis, we then used a whole-brain searchlight analysis\(^{21} \) to discover other regions that might code these quantities (Fig. 7). Consistent with our ROI analysis, local direction could be decoded in the bilateral RSC at the juncture of the calcarine sulcus and parietal occipital sulcus (Montreal Neurological Institute (MNI) coordinates: right 14 −68 11; left −5 −65 20). This locus is just posterior to retinocortical cortex proper (Brodmann area 29/30), which has been previously implicated in the coding of spatially fixed objects\(^{59} \). Beyond this region, we also observed significant local direction decoding in the left superior parietal lobule (SPL) (MNI coordinates: −19 −67 54; \( P = 0.033 \) corrected for multiple comparisons), an area that has been previously implicated in the coding of directional information\(^{60,61} \). No other brain regions exhibited local direction coding at either corrected significance levels or the more liberal uncorrected threshold of \( P < 0.001 \). An analogous searchlight analysis for local location coding found no brain regions at corrected significance levels, although coding in the bilateral RSC, left SPL and other regions of the posterior parietal cortex were observed at uncorrected significance levels. We did not observe evidence for coding of global direction in any region of the brain, even at relatively liberal thresholds (uncorrected \( P < 0.005 \)). Detailed analyses of the responses...
in the left SPL are presented in Supplementary Figure 2, and results from other ROIs are presented in Supplementary Figures 3 and 4.

**DISCUSSION**

Our results indicate that RSC, a region in the human medial parietal lobe, represents imagined facing direction and imagined location during spatial memory recall; moreover, it does so using a reference frame that is anchored to local environmental features and generalizes across local environments with similar geometric structures. These findings suggest that RSC is centrally involved in a critical component of spatial navigation: establishing one's position and orientation relative to fixed elements of the external world.

Location and direction codes revealed themselves in two ways. First, they were revealed through behavioral priming: reaction times in the JRD task were speeded when imagined facing direction or imagined location was repeated across successive experimental trials (experiment 1). Second, they were revealed through multivoxel pattern analysis: distributed fMRI activity patterns in RSC during the JRD task were more similar for imagined views corresponding to the same facing direction or location than for imagined views corresponding to different facing directions or locations (experiment 2). The fact that direction and location were defined relative to local environmental features was established by examining priming and decoding across views in different local environments (i.e., museums) that were geometrically similar but oriented orthogonally to each other. Views facing the same local direction (as defined by local geometry) in different museums were treated as equivalent across museums if they were in the equivalent position within the room but not if they were in the equivalent position as defined by the global reference frame. Thus, both behavioral priming and fMRI signals revealed spatial codes that were anchored to the local environmental features and generalized across geometrically similar local environments rather than codes that were idiosyncratic to each local environment or codes that were anchored to a global reference frame. To our knowledge, this is the first evidence for a spatial representation that uses the same alignment principles across different environments, such that local north (i.e., facing the back wall) in one local environment is equivalent to local north in another. A geometry-dependent but environment-independent representation of this type might be critical for translating viewpoint-dependent scene information (‘I am in a rectangular room, facing the short wall’) into allocentric spatial codes (‘I am facing to the north’) and, conversely, for recreating viewpoint-dependent scenes from allocentric memory traces.

The finding that RSC represents imagined direction and imagined location in a local spatial reference frame during spatial recall has been anticipated by previous neuroimaging and neuropsychological results. This region has been implicated previously in spatial navigation and spatial memory. It responds strongly during passive viewing of environmental scenes in fMRI studies, especially when the scenes depict familiar places. Moreover, its response is increased when subjects retrieve spatial information and move through an environment for which they have obtained survey knowledge. fMRI adaptation is observed in RSC when successively presented scenes face the same direction, and we previously reported that multivoxel patterns elicited in RSC when viewing familiar scenes contain information about location and direction that is independent of the specific view. Damage to RSC produces a profound form of topographical disorientation in which patients can accurately recognize landmarks but cannot use these landmarks to orient themselves in space. The current results extend this previous work by showing that RSC codes directions and locations while subjects perform a behavioral task that elicits direction-dependent and location-dependent effects. In addition, we demonstrate that these spatial codes are anchored to environmental features in a consistent manner that applies across different local environments—a crucial element of a spatial referencing system that was not demonstrated by previous studies. The reliability of these neural codes was evidenced by the fact that we were able to recreate accurate maps of the spatial relationships between views based on activity patterns in RSC. To our knowledge, this is the first case in which the spatial structure of an environment could be reconstructed purely from the neural correlates of human imagination.

Our findings also dovetail with previous behavioral research on spatial memory recall. When subjects perform a JRD task, they typically exhibit orientation-dependent (i.e., direction-dependent) performance. In cases where subjects have experienced only one view of an environment, accuracy is greater and reaction times are faster for imagined views that face the same direction as the learned view, irrespective of the imagined location. In cases where subjects have learned multiple views of an environment or have extensive experience over time, performance is typically best for a single imagined facing direction that acts as a conceptual north, and it is better for facing directions that are aligned or orthogonal to this primary reference direction than for directions that are misaligned. The factors that influence the alignment of the reference axes are a subject of much investigation, but previous work has indicated that spatial geometry can be a dominant influence under many circumstances. Indeed, geometry has long been known to guide reorientation, often to the detriment of other cues, and geometry is a strong controller of HD cells when animals cannot rely on a pre-existing sense of direction. Consistent with these previous results, we observed...
orientation-dependent priming and orientation-dependent activity patterns when orientation was defined relative to local geometry. Moreover, we also observed location-dependent priming and location-dependent activity patterns when location was defined relative to the same spatial reference frame. In sum, our results build on the classic findings from behavioral literature by showing that priming, like overall performance, is orientation dependent and by demonstrating that RSC encodes a spatial reference frame that may underlie this orientation-dependent behavior.

An open question is how the spatial codes in RSC are implemented neurally. HD cells have been identified previously in the retrosplenial cortex of rodents33,34. Thus, one possibility is that the HD code at encoding is re-instantiated at retrieval. However, there are reasons to doubt that the firing of HD cells, in and of themselves, can explain our current results. To our knowledge, there is little evidence that HD cells code direction in a manner that is consistent across different environments with similar geometries, as we observed here in RSC. Indeed, the fact that HD cells maintain stable preferred directions when traveling across distinct environments11,35,36 suggests that these cells do not directly represent local geometric relationships. Rather, they maintain a consistent sense of direction through path integration37, only referring back to the environment periodically to recalibrate when error has accumulated10. This means that a different mechanism that codes heading relative to local features is necessary to provide the calibration. Moreover, we found that spatial codes in RSC do not represent only facing direction but also include information about location, which is not encoded by HD cells. This finding is consistent with previous work indicating that cells in the retrosplenial and medial parietal region encode a variety of spatial quantities, not just heading direction35,38.

One possibility is that the responses we observed in RSC reflect the firing of neurons that encode the egocentric bearings to specific landmarks. In principle, any landmarks could be represented by such reference vector cells, but to explain our result, at least some of the landmarks must correspond reliably to Euclidean directions and have a clear analogy across the different museums. This suggests that the primary landmarks in the current environment are the walls, although it is possible that other features such as the doorway might be used, whose bearings would only approximately correspond to Euclidean direction. To account for location sensitivity, we must further posit that reference vector cells preferentially encode directions to nearby rather than distant landmarks during the JRD task, which would lead to location decoding because the identities of the nearest walls (as well as the door) would differ as a function of location. Under this account, both location and direction should be decodable in RSC, with the relative balance of these two kinds of information determined by the overlap in the identities of the landmarks used to define direction at each location3. Such a landmark referencing system could be used to anchor the HD system and thus provide a sense of direction. Alternatively, the results might be explained by the firing of orientation-modulated boundary vector cells39–41, which would fire when facing a given direction at a certain distance from specific walls. It is also possible that RSC represents heading and location using a mixed ensemble code that is not easily interpretable at the single-unit level42.

Whatever the underlying neural code, it is clear that spatial representations in RSC were tied to local environmental features in the current experiment. That said, we must caveat the use of the word ‘local’. Reference objects and target objects were always located in the same museum. If subjects had been asked to imagine themselves within a museum and report the bearing to a target outside, they might have accessed a global rather than local spatial frame. Previous behavioral work with non-nested environments has indicated that reference frames can be observed across many different scales, from tabletops to rooms43 to campuses44. Whether RSC represents spatial reference frames at all of these scales is an important question for future investigation—for tasks that involve memory recall, the local scene might potentially expand to include landmarks that can be encompassed by the imagination, not just landmarks that are actually visible from a particular point. An additional point is that subjects learned all the museums simultaneously, which might have prompted them to encode a single spatial frame or spatial schema44 that was applicable to all of them rather than separate spatial frames that would apply to each uniquely. We observed some anecdotal evidence in support of this idea during the pre-scan encoding phase: when subjects were asked to find an object, they sometimes traveled to the right location in the wrong museum. This suggests that they knew where the object was within the spatial frame but did not remember the museum to which the frame should be applied. One possibility for this is that the absence of proprioceptive and vestibular cues during virtual navigation may have made it difficult for RSC to use path integration to distinguish between museums. However, similar results were observed in a recent rodent neurophysiology study, which found that hippocampal place cells fired in analogous positions in geometrically identical environments that were learned in a single training session, even though these proprioceptive and vestibular cues were present45.

Nevertheless, a different training regime in which subjects learned one museum thoroughly before learning another might have resulted in less generalizability of responses between local environments; furthermore, it is possible that the local environments might dissociate after further experience46. We should also note that the present results leave it unclear whether the critical environmental factor for defining the reference frame was the shape of the museum as defined by the walls, the shaping of egocentric experience by the single door, or the principal axis defined by both factors.

Beyond its role in providing a system of reference for defining heading and location, our results also illuminate the contribution of RSC to memory retrieval more broadly. Much recent interest has focused on the striking overlap among brain regions involved in the retrieval of autobiographical memories, the imagination of novel events, and navigation1,4,47,48. These regions, including RSC, are believed to form a network that mediates both prospective thought and episodic memory49. Our results identify a specific role for RSC in the construction and reconstruction of episodes: by situating our imagined position and heading relative to stable spatial reference elements, RSC allows us to mentally place ourselves within an imaginary world built on the foundations of our remembered experience. This conclusion is consistent with previous work indicating that RSC is strongly activated when imagining changes in one’s viewpoint50 or when retrieving spatial relationships after first-person navigation27. Moreover, the fact that RSC codes a spatial schema that can apply across different local environments suggests the intriguing possibility that it might also be involved in coding memory schemas that are not explicitly spatial.

In summary, our experiments demonstrate a neural locus for the local spatial reference frames used for mental reorientation during spatial recall. RSC represents the locations and directions of imagined views with respect to these local reference frames, and this spatial code is rich enough to allow recreation of accurate maps of the local space. Like a compass rose, RSC anchors our sense of direction to the world, thus allowing us to stay oriented while we navigate and to reconstruct from memory the experience of being in a particular place.
METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

ACKNOWLEDGMENTS

This research was supported by the US National Institutes of Health (EY022350) and the National Science Foundation (SBE-0541957 and SBE-1041707). We thank A. Stigliani for assistance with data collection and three anonymous referees for useful comments on an earlier version of the manuscript.

AUTHOR CONTRIBUTIONS

S.A.M., L.K.V. and R.A.E. designed the experiments. S.A.M. and J.R. collected the data. S.A.M. analyzed data with input from L.K.V. and R.A.E. S.A.M. and R.A.E. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

Reprints and permissions information is available online at http://www.nature.com/reprints/index.html.

1. O’Keefe, J. & Dostrovsky, J. The hippocampus as a spatial map. Preliminary evidence from unit activity in the freely-moving rat. Brain Res. 34, 171–175 (1971).
2. Hafting, T., Fyhn, M., Molden, S., Moser, M.-B. & Moser, E.I. Microstructura spatial of the entorhinal cortex. Nature 436, 801–806 (2005).
3. Taube, J.S., Muller, R.U. & Ranck, J.B. Head-direction cells recorded from the postsubiculum in freely moving rats. I. Description and quantitative analysis. J. Neurosci. 10, 420–435 (1990).
4. Vass, L.K. & Epstein, R.A. Abstract representations of location and facing direction in the human brain. J. Neurosci. 33, 6133–6142 (2013).
5. Hassabis, D. et al. Decoding neuronal ensembles in the human hippocampus. Curr. Biol. 19, 546–554 (2009).
6. Doeller, C.F., Barry, C. & Burgess, N. Evidence for grid cells in a human memory system. Nature 463, 579–582 (2009).
7. Ekstrom, A.D. Direct recordings of grid-like neuronal activity in human spatial navigation. Nature 436, 1188–1190 (2013).
8. Kinneri, J.J., Kudrimoti, H.S. & McNaughton, B.L. Place cells,head direction cells, and the learning of landmark stability. J. Neurosci. 15, 1648–1659 (1995).
9. Valero, S. & Taube, J.S. Path integration: how the head direction signal maintains and corrects spatial orientation. Nat. Neurosci. 15, 1445–1453 (2012).
10. Yoder, R.M. et al. Both visual and idiothetic cues contribute to head direction cell stability during navigation along complex routes. J. Neurophysiol. 105, 2989–3001 (2011).
11. Yoder, R.M., Clark, B.J. & Taube, J.S. Origins of landmark encoding in the brain. Trends Neurosci. 34, 561–571 (2011).
12. Epstein, R.A. Parahippocampal and retrosplenial contributions to human spatial navigation. Trends Cogn. Sci. 12, 388–396 (2008).
13. Vann, S.D., Aggleton, J.P. & Maguire, E.A. What does the retrosplenial cortex do? J. Exp. Psychol. Learn. Mem. Cogn. 28, 162–170 (2002).
14. Epstein, R.A., Higgins, J.S., Jablonski, K. & Feiler, A.M. Visual scene processing in familiar and unfamiliar environments. J. Neurophysiol. 97, 3670–3683 (2007).
15. Epstein, R.A., Parker, W.E. & Feiler, A.M. Where am I now? Distinct roles for parahippocampal and retrosplenial cortices in place recognition. J. Neurosci. 27, 6141–6149 (2007).
16. Baumann, O. & Mattingley, J.B. Medial parietal cortex encodes perceived heading direction in humans. Proc. Natl. Acad. Sci. USA 101, 14046–14051 (2004).
17. Kriegeskorte, N., Goebel, R. & Bandettini, P. Information-based functional brain mapping. Proc. Natl. Acad. Sci. USA 103, 3863–3868 (2006).

22. Maguire, E.A. The retrosplenial contribution to human navigation: a review of lesion and neuroimaging findings. Scand. J. Psychol. 42, 225–238 (2001).
23. Burgess, N. Spatial cognition and the brain. Ann. NY Acad. Sci. 1124, 77–97 (2008).
24. Epstein, R.A., Higgins, J.S., Jablonski, K. & Feiler, A.M. Visual scene processing in familiar and unfamiliar environments. J. Neurophysiol. 97, 3670–3683 (2007).
25. Epstein, R.A., Parker, W.E. & Feiler, A.M. Where am I now? Distinct roles for parahippocampal and retrosplenial cortices in place recognition. J. Neurosci. 27, 6141–6149 (2007).
26. Rosenbaum, R.S., Ziegler, M., Winocur, G., Grady, C.L. & Moscovich, M. “I have often walked down this street before”: fMRI studies on the hippocampus and other structures during mental navigation of an old environment. Hippocampus 14, 826–835 (2004).
27. Zhang, H., Copara, M. & Ekstrom, A.D. Differential recruitment of brain networks following route and cartographic map learning of spatial environments. PLoS ONE 7, e44886 (2012).
28. Wolters, T. & Böchel, C. Dissociable retrosplenial and hippocampal contributions to successful formation of survey representations. J. Neurosci. 25, 3333–3340 (2005).
29. Burgess, N., Spiers, H.J. & Paleologou, E. Orientational manoeuvres in the dark: dissociating allocentric and egocentric influences on spatial memory. Cognition 94, 49–80 (2004).
30. Cheng, K. A purely geometric module in the rat’s spatial representation. Cognition 23, 149–178 (1986).
31. Galič, C.R. The Organization of Learning (The MIT Press, Cambridge, MA, 1990).
32. Knight, R., Hayman, R., Lin Grizberg, L. & Jeffery, K. Geometric cues influence head direction cells only weakly in nondirosrected rats. J. Neurosci. 31, 15681–15692 (2011).
33. Cho, J. & Sharp, P.E. Head direction, place, and movement correlates for cells in the rat retrosplenial cortex. Behav. Neurosci. 115, 3–25 (2001).
34. Chen, L.L., Lin, L.H., Green, E.J., Barnes, C.A. & McNaughton, B.L. Head-direction cells in the rat posterior cortex. Exp. Brain Res. 101, 8–23 (1994).
35. Taube, J.S. & Burton, H.L. Head direction cell activity monitored in a novel environment and during a cue conflict situation. J. Neurophysiol. 74, 1953–1971 (1995).
36. Dudchenko, P.A. & Zinyuk, L.E. The formation of cognitive maps of adjacent environments: evidence from the head direction cell system. Behav. Neurosci. 119, 1511–1523 (2005).
37. McNaughton, B.L., Battaglia, F.P., Jensen, O., Moser, E.I. & Moser, M.-B. Path integration and the neural basis of the ‘cognitive map’. Nat. Rev. Neurosci. 7, 663–678 (2006).
38. Sato, N., Sakata, H., Tanaka, Y.L. & Taira, M. Navigation-associated medial parietal neurons in monkeys. Proc. Natl. Acad. Sci. USA 103, 17001–17006 (2006).
39. Lever, C., Burton, S., Jeewaiaje, A., O’Keefe, J. & Burgess, N. Boundary vector cells in the subiculum of the hippocampal formation. J. Neurosci. 29, 9771–9777 (2009).
40. Solstad, T., Boccara, C.N., Kropff, E., Moser, M.-B. & Moser, E.I. Representation of geometric borders in the entorhinal cortex. Science 322, 1865–1868 (2008).
41. Sharp, P.E. Subicular place cells generate the same “map” for different environments: comparison with hippocampal cells. Behav. Brain Res. 174, 206–214 (2006).
42. Hung, C.P., Kreiman, G., Poggio, T. & DiCarlo, J.J. Fast readout of object identity from macaque inferior temporal cortex. Science 310, 863–866 (2005).
43. Mehta, S.A., Yerasansetti, A., Burns, T.J. & Shelton, A.L. Spatial memory in the real world: long-term representations of everyday environments. Mem. Cognit. 39, 1401–1408 (2011).
44. Tse, D. et al. Schemas and memory consolidation. Science 316, 76–82 (2007).
45. Spiers, H.J., Hayman, R.M.A., Jovalekic, A., Marozi, E. & Jeffery, K.J. Place field repetition and purely local remapping in a multicompartment environment. Cereb. Cortex doi:10.1093/cercor/bht198 (13 August 2013).
46. Lever, C., Wills, T., Cacucci, F., Burgess, N. & O’Keefe, J. Long-term plasticity in hippocampal place-cell representation of environmental geometry. Nature 416, 90–94 (2002).
47. Ranganath, C. & Ritchey, M. Two cortical systems for memory-guided behaviour. Nat. Rev. Neurosci. 13, 713–726 (2012).
48. Hassabis, D., Kuman, D. & Maguire, E.A. Using imagination to understand the neural basis of episodic memory. J. Neurosci. 27, 14365–14374 (2007).
49. Schacter, D.L. et al. The future of memory: remembering, imagining, and the brain. Neuron 76, 677–694 (2012).
50. Lambyre, S., Doeller, C., Berthoz, A. & Burgess, N. Imaging being somewhere else: neural basis of changing perspective in space. Cereb. Cortex 22, 166–174 (2012).
ONLINE METHODS

Participants. Forty-six healthy subjects (23 female; mean age, 21.2 ± 1.7 years (s.d.)) with normal or corrected-to-normal vision were recruited from the University of Pennsylvania community. Twenty-two participated in the behav- ioral study (experiment 1), and 24 were scanned with fMRI (experiment 2). Sample size was determined based on a pilot behavioral experiment. Subjects provided written informed consent in compliance with procedures approved by the University of Pennsylvania Institutional Review Board. Five participants were replaced in experiment 1 for scoring less than 70% correct overall. Six additional participants from experiment 2 were excluded before scanning for scoring less than 70% on the behavioral practice trials after the first day of training (as described below). Three additional subjects were scanned in experiment 2 but were excluded before analysis, the first for technical difficulties in fMRI acquisition, the second because she requested to terminate the scanning session early and the third for sleeping.

MRI acquisition. Scanning was performed at the Hospital of the University of Pennsylvania using a 3 T Siemens Trio scanner equipped with a 32-channel head coil. High-resolution T1-weighted images for anatomical localization were acquired using a three-dimensional magnetization-prepared rapid acquisition gradient echo pulse sequence (repetition time (TR), 1,620 ms; echo time (TE), 3.09 ms; inversion time, 950 ms; voxel size, 1 mm × 1 mm × 1 mm; matrix size, 192 × 256 × 160). T2*-weighted images sensitive to blood oxygenation level-dependent contrasts were acquired using a gradient echo echoplanar pulse sequence (TR, 3,000 ms; TE, 30 ms; flip angle 90°, voxel size, 3 mm × 3 mm × 3 mm; field of view, 192; matrix size, 64 × 64 × 44).

Virtual environment. A virtual environment consisting of a park containing four large rectangular buildings, or museums (Fig. 1a), was constructed with the Source SDK Hammer Editor (http://www.valvesoftware.com, Valve Software, Bellevue, WA). Distal orientational cues surrounded the park, such as a mountain range and the sun at a high azimuth to the north, apartment buildings and refineries to the east and west and two high rise apartment buildings to the south. The museums had identical interior geometry, with an aspect ratio of 0.84 and an axis of elongation running parallel to the direction of entry, but they were distinguishable interiorly and exteriorly by textures and architectural features, such as facades, columns and plinths (Fig. 1b). Large windows were set in the back wall of three of the four museums. Museums were placed in the environment so that each one was entered from a unique direction. This had the effect of dissociating direction within a museum as defined by the axis of entry and internal geometry (for example, facing the back wall, or local north) from direction in the exterior park (for example, facing the mountains, or global north).

Each museum contained eight distinct, nameable objects that were arranged in alcoves along the walls (Fig. 1b). Because of the shape of the alcoves, objects could only be viewed head on from a specific direction. In experiment 2, we randomized the assignment of objects to alcoves in the environment across participants to ensure that reliable decoding of direction and location could not be attributed to decoding of object identity.

Training session. All subjects underwent an initial training session in which they learned the layout of the virtual environment and the location of objects within it. This session was divided into a free exploration period (15 min) and a guided learning period (approximately 45 min). Subjects in experiment 2 performed an additional training session 2 d later, immediately before the fMRI scan, consisting of the guided learning period only.

In the free exploration period, subjects were placed at the entrance to the park (‘start’; Fig. 1a). After receiving instructions on how to navigate using the arrow keys, they were allowed to explore the environment at will. The only guidance provided by the experimenter was that they should make sure that they visited each of the buildings during this phase, a criterion met by every subject. The environment was rendered and displayed on a laptop running the commercial game software Portal (http://www.valvesoftware.com, Valve Software, Bellevue, WA).

The guided learning phase of the training session was divided into eight self-paced blocks. At the beginning of each, the name of an object was presented on the screen, and the subject was asked to navigate from the entrance to the park to the object as directly as possible. Once the subject had navigated to the correct location of the object, the name of another target object was presented on the screen, and the subject was asked to navigate directly to it. After locating eight objects (two from each museum), the subject was teleported back to the entrance of the park, and the next block began. All of the objects remained visible throughout the task to afford subjects additional opportunities to learn their locations while simultaneously challenging them to apply this knowledge. Targets were drawn randomly, with the constraint that no target object was repeated until all objects had been found once. Each object was searched for twice for a total of 64 trials.

Testing session. Immediately after the initial training session, subjects in experi- ment 1 completed 384 trials of a JRD task. In each trial, subjects used the keyboard to report whether a target object from the training environment would be located to their left or right if they were standing in front of and facing a reference object. The names of the reference object and target object were presented visually and simultaneously in gray letters on two different lines at the center of a black screen (for example, “Facing the Bicycle,” “Lamp”; Fig. 1c). Subjects were instructed to imagine themselves looking directly at the reference object while making their judgment; this required them to imagine themselves in a specific location while facing a specific direction (Fig. 2). They were told to report left and right broadly, including anything that would be on that side of their body and not just directly to the left (for example, in Fig. 2a, when facing view 1, view 7 would be to the left). Trials ended as soon as the participant responded, and the next trial began after a 750-ms intertrial interval. The target object was always within the same museum as the reference object. No feedback was given. One 60-s break was given halfway through the experiment to mitigate mental fatigue.

The 384 trials in experiment 1 were ordered randomly but with the constraint that successive trials faced the same direction 50% of the time and that direction changed at least once every six trials. The probability of repeating museum or corner across successive trials was not manipulated (25% for each based on chance). These transition probabilities were adopted to ensure that we would have a sufficient number of trials to detect priming of direction within a given museum. Because reaction times often have an asymmetric distribution, reaction times that were 2.5 s.d. above the mean were removed (5% of responses in experiment 1 and 1% of responses in experiment 2), and median reaction times were calculated for each condition.

Subjects in experiment 2 only completed 20 trials of the JRD task after the initial training session, after which they received feedback on their performance. They then returned 2 d later for a second training session (described above), after which they completed 544 trials of the JRD task in the fMRI scanner. The experi- mental paradigm was the same as that in experiment 1, except that the timing was modulated to fit the requirements of fMRI acquisition: rather than disappear- ing after the subject's response, as in experiment 1, word cues remained on the screen for 5,000 ms followed by a 1,000-ms gap before the start of the next trial. This meant that there was a substantially longer interval between a participant's response and the subsequent trial, which might have reduced behavioral priming effects. In addition, a grid of jittered gray lines was superimposed behind the word cues to limit differences in visual extent. Testing was performed in 8 scan runs, each of which was 8 min 24 s and consisted of 68 JRD trials and 5 null tri- als, during which the subject viewed a fixation cross and made no response for 12 s. To keep the scan sessions to a manageable length of time, only 16 of the 32 reference objects shown at training were used, drawn from 2 of the 4 museums, which were chosen to be adjacent and, thus, oriented orthogonally to one another (Fig. 2). The four possible museum pairs meeting this requirement (1-2; 2-3; 3-4; and 4-1) were randomly counterbalanced across subjects.

JRD trials were presented within a continuous carryover sequence51, which ordered the 16 imagined views (corresponding to the 16 reference objects) in a serially balanced design for which each view preceded and followed every other view, including itself, exactly once. Two unique carryover sequences were gener- ated for each subject, with an entire sequence shown over four scan runs. Each view was tested 17 times within each carryover sequence.

Functional localizer. In addition to the main experiment, subjects also completed two functional localizer scans, each lasting 5 min 32 s, which consisted of 16-s blocks of scenes, objects and scrambled objects. Images were presented for 600 ms with a 400-ms interstimulus interval as subjects performed a one-back task on image repetition.
Data preprocessing. Functional images were corrected for differences in slice timing by resampling slices in time to match the first slice of each volume. Images were then realigned to the first volume of the scan run, and subsequent analyses were performed within the subjects’ own space. Motion correction was performed using MCFLIRT. Data for the functional localizer scan were smoothed with a 6-mm full-width at half-maximum Gaussian filter; data for multivoxel pattern analyses were not smoothed.

Functional ROIs. Data from the functional localizer scan were used to identify RSC, an area in the retrosplenial and parietal-occipital sulcus region that has been implicated previously in spatial memory and navigation. RSC was defined for each subject individually and in their own space using a contrast of greater activity to scenes than to objects in the subjects’ functional localizer data and a group-based anatomical constraint of scene-selective activation derived from a large number (42) of localizer subjects in our lab. To select a constant number of voxels in a threshold-free manner, an individual’s RSC within a hemisphere was defined as the top 100 voxels that responded more strongly to scenes than to objects and fell within the group-parcel mask warped to the subject’s own space with a linear transformation. This method ensures that RSC can be defined in both hemispheres for every subject. Scene-selective ROIs corresponding to the objects and fell within the group-parcel mask warped to the subject’s own space with a linear transformation. This method ensures that RSC can be defined in both hemispheres for every subject. Scene-selective ROIs corresponding to the parahippocampal place area and occipital place area were defined in the same manner, and the early visual cortex was defined based on a contrast of scrambled-objects > objects.

Multivoxel pattern analysis. To test whether an ROI contained spatial information about the direction and location of views within the local environment, we calculated the correlation between multivoxel patterns associated with each of the 16 views (corresponding to the 16 reference objects) across different continuous carryover sequences. This procedure allowed us to test whether the information carried by patterns of responses in the first half of the experiment (i.e., the first carryover sequence) generalized to patterns in the second half. First we estimated the multivoxel activity pattern reflecting the response to each view within a carryover sequence. To do this, we applied a general linear model (GLM) with 16 regressors, one for each view, to the time course of functional activity within each voxel. Separate GLMs were performed on each of the four runs corresponding to a continuous carryover sequence, and the resulting parameter estimates were averaged to provide an estimate of the average response to a view in that half of the experiment. Multivoxel patterns were represented by a vector concatenating the responses across the voxels of an ROI. GLMs were implemented in FSL and included highpass filters, which removed low temporal frequencies, and nuisance regressors for motion parameters and outlier volumes discovered using the Artifact Detection Toolbox (http://www.nitrc.org/projects/artifact_detect/). The first three volumes of each run were discarded to ensure data quality.

Next we measured the similarity between activity patterns corresponding to the 16 views by calculating the Pearson correlations between patterns in the two halves of the data corresponding to the two carryover sequences. Individual pattern were normalized before this computation by subtracting out the grand mean pattern (i.e., the cocktail mean) for each continuous carryover sequence. This calculation created a $16 \times 16$ correlation matrix for each subject. To test for coding of a particular quantity, such as location or direction, we computed the average correlation for views that shared that quantity (for example, faced the same local direction) and compared this to the average correlation for views that did not share that quantity (for example, faced different directions), taking into account various restrictions in the other spatial quantities, as described in the main text (Fig. 2). These values were computed for each subject and then submitted to statistical analysis of cross-subject reliability. For four subjects, the 16th view was not represented within the continuous carryover sequence because of technical errors. For these subjects, the analysis was performed as normal but was adjusted to exclude this view.

Visualization and reconstruction of spatial similarities. To visualize pattern similarity qualitatively in RSC, we created a pattern similarity map for each of the 16 views by calculating the similarity between that view (starting view) and the other 15 other views (comparison views). Each map was subjected to rotations and reflections that aligned the starter views to one another while maintaining the spatial relations between the starter and comparison views. These maps were then averaged together to create a composite similarity map. Pattern similarities were converted to a range of 0 to 1 and colored according to this value.

To create reconstructions of the museums, multivoxel patterns were compared across carryover sequences to calculate a correlation matrix for the eight unique view positions in a museum for each subject. These correlation matrices were then averaged to produce a grand correlation matrix that was submitted to multidimensional scaling to produce a two-dimensional map of the views and then aligned with a Procrustes analysis to a schematic of our museums. Distortion of the reconstructed map was defined as the sum of squared errors between the reconstructed and true points after optimal linear alignment, and significance was calculated by randomly shuffling the correlation matrix and applying multidimensional scaling to the result over 100,000 iterations to produce a chance distribution for distortion. A similar reconstruction was performed based on similarities between corresponding views in different museums.

Searchlight analysis. To test for coding of local direction and other spatial quantities outside of our predefined ROIs, we implemented a whole-brain searchlight analysis. This analysis stepped through every voxel of the brain and centered a small spherical ROI (radius of 5 mm) around it. The comparisons among views for discriminating different sources of spatial information (Fig. 2) were then performed within the spherical neighborhood. The central voxel of the sphere was assigned the pattern discrimination score (for example, the difference in pattern similarity between views that face the same or different local directions) calculated over its neighborhood. For each subject, this procedure generated a map associated with the pattern discriminability of a particular spatial quantity. These maps were then aligned to the MNI template with a linear transformation and submitted to a second-level random-effects analysis to test the reliability of discrimination for that quantity across subjects. To find the true type I error rate, we performed Monte Carlo simulations that permute the sign of the data (i.e., local or global coding). Nonparametric Monte Carlo simulations were used to determine significance for the reconstructions of spatial similarities, as well as for the whole-brain searchlight analyses.

Statistics. Data distributions were assumed to be normal, but this was not formally tested. Repeated-measures ANOVAs were used compare reaction times (experiment 1) and pattern similarities (experiment 2) as a function of imagined direction and location. Where appropriate, paired-sample t-tests were used to directly compare different possible coding schemes that could explain the data (i.e., local or global coding). Nonparametric Monte Carlo simulations were used to determine significance for the reconstructions of spatial similarities, as well as for the whole-brain searchlight analyses.

A Supplementary Methods Checklist is available with the supplementary materials.
Corrigendum: Anchoring the neural compass: coding of local spatial reference frames in human medial parietal lobe

Steven A Marchette, Lindsay K Vass, Jack Ryan & Russell A Epstein
Nat. Neurosci. 17, 1598–1606 (2014); published online 5 October 2014; corrected after print 12 January 2015

In the version of this article initially published, three citations present in an earlier draft were omitted. To correct this, the following sentence and phrase have been inserted in the first paragraph of the Results section “Whole-brain analyses,” before the third sentence and at the end of the fourth, respectively: “This locus is just posterior to retrosplenial cortex proper (Brodmann area 29/30), which has been previously implicated in the coding of spatially fixed objects59. “... an area that has been previously implicated in the coding of directional information60,61. “In the sentence before these inserts, the corresponding description "and just posterior to the retrosplenial cortex proper” has been deleted. The references are as follows:

59. Auger, S.D. & Maguire, E.A. Assessing the mechanism of response in the retrosplenial cortex of good and poor navigators. Cortex 49, 2904–2913 (2013).
60. Schindler, A. & Bartels, A. Parietal cortex codes for egocentric space beyond the field of view. Curr. Biol. 23, 177–182 (2013).
61. Spiers, H.J. & Maguire, E.A. A navigational guidance system in the human brain. Hippocampus 17, 618–626 (2007).

The error has been corrected in the HTML and PDF versions of the article.