Odor quality coding and categorization in human posterior piriform cortex

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Efficient recognition of odorous objects universally shapes animal behavior and is crucial for survival. To distinguish kin from nonkin, mate from nonmate and food from nonfood, organisms must be able to create meaningful perceptual representations of odor qualities and categories. It is currently unknown where and in what form the brain encodes information about odor quality. By combining functional magnetic resonance imaging (fMRI) with multivariate (pattern-based) techniques, we found that spatially distributed ensemble activity in human posterior piriform cortex (PPC) coincides with perceptual ratings of odor quality, such that odorants with more (or less) similar fMRI patterns were perceived as more (or less) alike. We did not observe these effects in anterior piriform cortex, amygdala or orbitofrontal cortex, indicating that ensemble coding of odor categorical perception is regionally specific for PPC. These findings substantiate theoretical models emphasizing the importance of distributed piriform templates for the perceptual reconstruction of odor object quality.

An important property of the brain is its ability to create coherent, meaningful perceptual constructs from the complexity of the outside world. These internalized representations of the external environment provide a neural basis for object recognition, identification and categorization, enabling organisms to focus cognitive resources, optimize behavioral responses and generalize past experiences to novel events^{1,2}.

Research on object processing has traditionally focused on visual object processing^{2,3}, which tends to overshadow the critical ecological role of 'odor objects', which we define as the perceptual quality or character of a smell emitted from an odorous substance. Efficient recognition of odor objects universally shapes animal behavior and is crucial for survival. Indeed, the ubiquity of olfactory-guided adaptive behaviors across vertebrate and invertebrate species, including maternal bonding⁴, kinship recognition⁵, mate selection⁶ and territorial defense⁷, makes it clear that olfactory systems are important for recording and classifying odor objects.

Where and in what form does the brain encode the perceptual quality of an odor object? Several elegant studies have demonstrated that odor-evoked spatial activity in the rodent olfactory bulb correlates with behavioral measures of odor similarity^{8–11}, as shown by habituation or reinforcement learning procedures, leading to the suggestion that neural representations of odor quality are reflected in ensemble bulbar activity. However, the obligatory use of indirect perceptual assays in animal models hampers efforts to establish an explicit link between neural response patterns and odor quality perception. Such limitations accentuate the unique advantages of studying human

subjects, whose ability to verbalize their percepts¹² and use perceptual rating scales offers an ideal research alternative for clarifying the neuroscientific underpinnings of odor quality perception.

One plausible location for the encoding and classification of odor quality information is primary olfactory (piriform) cortex, which receives direct input from the olfactory bulb and has extensive interconnections with intrinsic and extrinsic fiber systems, including amygdala, hypothalamus, entorhinal cortex and orbitofrontal cortex (OFC)^{13,14}. The privileged access that the piriform cortex has to sensory, affective, physiological and motivational features of an olfactory stimulus perfectly endows it with the capacity for weaving together odor representations with direct relevance for perception and behavior. This hypothesis receives support from both anatomical 15,16 and computational^{14,17,18} models, which predict that odor percepts are embodied in spatially distributed patterns of piriform activity, providing a robust neural substrate for odor coding, memory and recall. However, direct neurobiological evidence for this idea is currently not available and the functional architecture of odor object qualities in the olfactory brain remains unknown.

We combined high-resolution fMRI with olfactory multivariate analysis techniques¹⁹ (**Fig. 1**) to investigate spatial ensemble coding of odor qualities and categories in human PPC. Multivariate fMRI methods^{20–22} differ from conventional (univariate) fMRI analyses, in which data are averaged over space (voxels), time (scans) and subjects, obscuring potentially important information that may be contained at the level of individual voxels, scans and subjects. These pattern-based approaches provide a robust new method for

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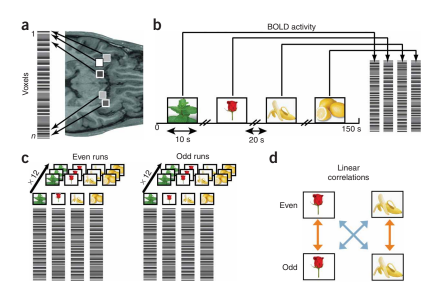


Figure 1 Schematic diagram of the correlation analysis. (a) The condition-specific spatial patterns of voxel activity in the PPC were transformed into linear vectors of voxel activity (pattern vectors). Voxels are represented by shaded squares on an axial slice of an anatomical MRI scan. The level of grayscale intensity represents the BOLD signal intensity. (b) Pattern vectors were composed of the peak BOLD activity across the stimulus presentation, shown here in the context of Experiment 1, for one run (150 s). (c) The entire dataset of pattern vectors was split into halves, with one half containing data from the 12 even runs and the other half containing data from the 12 odd runs and then averaged across runs, producing one mean pattern vector per odorant in each half of the data. (d) Averaged pattern vectors were used to calculate within-odorant (orange arrow) and across-odorant (blue arrow) pair-wise correlation coefficients.

characterizing how (rather than just where) perceptual information is represented in the human brain²³.

Our study consists of two independent experiments centered on the hypothesis that odor qualities and categories are encoded as distributed spatial ensembles in human PPC. In Experiment 1, we combined multivariate and cortical flattening techniques to test whether qualitatively distinct odorants are associated with unique multi-voxel fMRI patterns in PPC in the absence of mean activation differences, odorevoked fMRI representations in PPC are distributed and overlapping, without evidence for topographical clustering; and olfactory ensemble codes of odor quality are regionally specific for PPC. In Experiment 2, we extended these hypotheses to a wider set of odorants, which, in combination with multidimensional scaling techniques, enabled us to examine whether fMRI ensemble patterns in PPC constitute an olfactory code that aligns with categorical perception of odor quality.

RESULTS

Experiment 1: odor-specific ensemble codes in PPC and OFC

We set out to test the hypothesis that neural codes of odor quality take the form of ensemble fMRI activity in PPC. We decided to target PPC as a primary region of interest because recent animal and human studies have suggested that neural representations of odor quality are stored in this brain region^{24,25}. During fMRI scanning, six subjects sniffed four easily distinguishable odorants (R-(-)-carvone (mint), phenethyl alcohol (rose), amyl acetate (banana) and citronellol (lemon)) across a total of 24 imaging runs. On each odorant presentation, subjects were visually cued to make a sniff and to tacitly identify the quality of the odor.

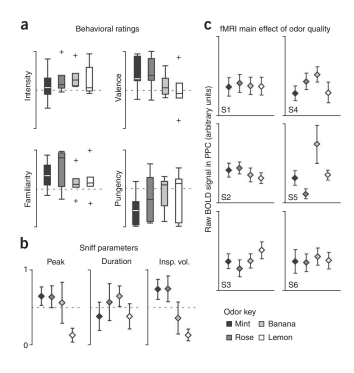
The behavioral ratings of the four odorants, acquired before scanning, did not significantly differ in odor intensity ($F_{3,20} = 0.39$,

Figure 2 Behavioral data and univariate imaging analysis for Experiment 1.

(a) Group-averaged behavioral ratings of odor intensity, pleasantness, pungency and familiarity are depicted as box plots indicating median (central line) and upper and lower quartiles (top and bottom of box, respectively) for each odorant. Whiskers denote the extent of data between the tenth and 90th percentiles. Outliers are indicated by crosses. Ratings did not differ across any of these measures. (b) Mean normalized values (± between-subjects s.e.m.) for sniff peak amplitude, duration and inspiratory volume (insp. vol.) did not differ between the four odorants. (c) Plots of mean fMRI signal in PPC for each subject and odorant (± within-subject s.e.m.) revealed no difference except for subject 5 (S5).

P=0.76), pleasantness ($F_{3,20}=1.49$, P=0.25), pungency ($F_{3,20}=0.73$, P=0.55) or familiarity ($F_{3,20}=1.00$, P=0.41) (**Fig. 2a**). Moreover, pair-wise ratings of odor quality similarity (total of six pair-wise comparisons) indicated that the odorants were equally discriminable ($F_{5,35}=1.73$, P=0.15). There were also no systematic differences in sniff peak amplitude ($F_{3,20}=2.78$, P=0.11), duration ($F_{3,20}=0.53$, P=0.67) or inspiratory volume ($F_{3,20}=2.04$, P=0.14) between odorants (**Fig. 2b**) that might have otherwise confounded the imaging analysis.

We first considered whether wholesale differences in PPC activity between odorant conditions were detectable using a conventional (univariate) fMRI analysis. After all, if odor quality–specific information in PPC merely amounts to global changes in mean fMRI activity, then multivariate methods would not be necessary to confirm our original hypothesis. We averaged the fMRI signal in PPC across the set of voxels for each subject and found no significant difference between



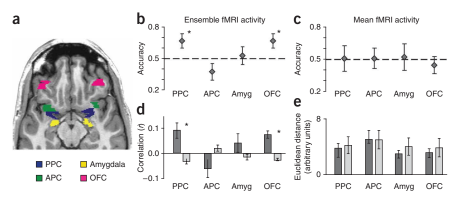


Figure 3 Pattern discrimination of odor quality in human PPC and OFC. (a) Axial slice of a T1-weighted structural scan showing anatomically defined regions of interest. Subsets of voxels from these brain regions (see Online Methods) were used in the pattern analyses. (**b**–**e**) Odor identification accuracy (mean \pm between-subjects s.e.m., n=6, **b**) calculated using fMRI patterns of ensemble activity exceeded chance (dashed line) across subjects in PPC and OFC and the within-odor correlations (dark gray bars) were greater than the across-odor correlations (light gray bars) in both regions (**d**). In contrast, identification accuracy on the basis of mean fMRI activity levels did not significantly differ from chance in any of the measured regions (**c**), nor were there differences between within-odor and across-odor Euclidean distances (**e**). * P < 0.05.

odorants for five out of six subjects (P > 0.05, one-way ANOVAs, four levels (odorants), one ANOVA per subject; **Fig. 2c**). The mean fMRI signal significantly differed between odorants only in subject 5 ($F_{3,92} = 3.71$, P = 0.014), although follow-up pair-wise t tests indicated that not all odorants could be significantly discriminated from the others (for example, mint versus rose, $T_{23} = 1.86$, P = 0.076; mint versus lemon, $T_{23} = 0.28$, P = 0.76; banana versus lemon, $T_{23} = 1.67$, P = 0.11). These results indicate that if a neural code for odor quality exists in PPC, it might be contained in a more fine-grained pattern of voxel activity that cannot be seen using standard imaging techniques.

To test this hypothesis directly, we extracted odorant-specific voxelwise patterns of fMRI ensemble activity in PPC, as determined by an independent odor localizer task. The 24-run set of spatial patterns, organized as linear vectors of voxel activity, was divided into 12 even and 12 odd runs²⁰ and used to calculate linear correlations (Fig. 1) for all within-odor (for example, rose/even versus rose/odd) and all across-odor (for example, rose/even versus banana/odd) pairs (Fig. 3). Odor identification accuracy, calculated as the proportion of withinodor correlations greater than across-odor correlations, significantly exceeded 50% chance in PPC across subjects ($T_5 = 2.42$, P = 0.030; Fig. 3a,b). In addition, the average within-odor correlation was significantly greater than the average across-odor correlation across subjects in PPC ($T_5 = 2.99$, P = 0.030; Fig. 3d), an effect that was equally observed across the four odors ($F_{3,20} = 2.10$, P = 0.13, one-way ANOVA, four levels (difference between within- and across-odor correlations for each odorant)). These results indicate that odorspecific information is contained in multi-voxel PPC activity patterns and that these effects were not driven by any one particular stimulus.

These findings indicate that fMRI representations of odor quality are embedded in PPC ensemble activity. However, these analyses alone cannot reveal the specific topographical organization of brain activity underlying these putative odor codes (for example, local versus distributed, discrete versus overlapping). To visualize the complex PPC anatomy in a single plane, we used cortical flattening (unfolding) techniques²⁶ to generate odor 'flat maps' in PPC. These maps were comprised of the flattened two-dimensional patterns of blood oxygen level–dependent (BOLD) signal in all odor-active piriform voxels, averaged across trials for each of the four odorants. Data from two

representative subjects (**Fig. 4**) indicate that the topographical arrangement of left PPC activity was spatially distributed and unique for each odorant in the absence of any obvious local clustering and without topographical consistency between subjects. The implication is that olfactory percepts are represented in overlapping, but distinct, response profiles across subsets of voxels in PPC.

Is odor quality information encoded in PPC selectively or are other cortical areas involved? To answer this question, we extended the analysis to three additional olfactory brain regions. In anterior piriform cortex (APC) and amygdala, odor identification accuracy did not differ from chance (APC: $T_5 = 1.57$, P = 0.089; amygdala: $T_5 = 0.34$, P = 0.37), but accuracy in OFC rivaled that in PPC and significantly exceeded chance ($T_5 = 2.48$, P = 0.028) (Fig. 3a,b). A one-way ANOVA testing for an effect of region on the identification accuracy was significant ($F_{3.20} = 3.45$, P = 0.036), implying

that the observed results are specific for PPC and OFC. Similarly, the within- versus across-odor correlation difference was not significant in APC ($T_5 = 1.70$, P = 0.15) or amygdala ($T_5 = 1.12$, P = 0.31), but was highly significant in OFC ($T_5 = 5.40$, P = 0.0029) (**Fig. 3d**). Once again, a one-way ANOVA testing for region effects was significant ($F_{3,20} = 4.91$, P = 0.010), indicating that the correlation findings are specific to PPC and OFC. These additional results indicate that PPC and OFC both contain distributed odor-specific ensemble representations, consistent with prior animal and human data highlighting the role of OFC in olfactory coding^{27–30}. That these patterns were only observed in PPC and OFC underscores the regional specificity of olfactory ensemble coding and validates the efficacy of multivariate techniques to delineate odor information processing in the human brain.

The above results indicate that PPC ensemble patterns can be used to discriminate odorant identity, but it remains possible that the mean fMRI activity in PPC could also contain discriminating information. Therefore, to directly compare classification performance on the basis of fMRI multivariate and univariate datasets, we performed a complementary analysis of the mean fMRI amplitudes, following the same classification methods used to analyze the fMRI patterns. Specifically, the global fMRI activity level was computed across all voxels in PPC for each odorant and each run. The data were then split into even and odd runs and pair-wise Euclidean distances (as a proxy for linear correlations) were calculated between the means of withinodorant and across-odorant conditions. Identification accuracy (proportion of within-odor distances shorter than across-odor distances) did not differ from chance across the group in PPC, APC, amygdala or OFC (P = 0.47, 0.48, 0.42 and 0.33, respectively; **Fig. 3c**). Furthermore, the within-odor distance was not significantly smaller than the across-odor distance in any of these regions (P = 0.73, 0.99, 0.47 and 0.67; Fig. 3e). These additional data suggest that mean fMRI amplitude does not contain sufficient information to distinguish between the odorants.

Finally, we applied split-half classification methods to the trialspecific respiratory parameters (split into even and odd runs) to rule out potential contributions of sniffing to the observed PPC patterns. Odor identification accuracy was not significant across the group when classification was performed on sniff peak, duration, volume or a linear



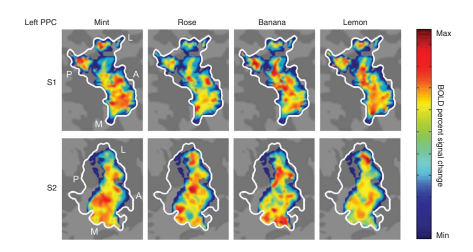


Figure 4 Odorant-specific spatial maps in PPC. The three-dimensional representations of odorantevoked activity in PPC from two subjects were projected onto two-dimensional (flat) maps, allowing visualization of voxel-wise odor patterns in a single plane. Maps depict the odorant-evoked BOLD percent signal change in all odor-active voxels (liberally thresholded at P < 0.5), averaged across trials for each of the four odorants. The pseudo-color scale spans the full range of BOLD percent signal change in each map, from minimum (deep blue) to maximum (bright red). Each odorant elicited a distributed pattern of fMRI activity in left PPC (white outline) that overlapped with, but was distinct from, the other odorants. Unique distributed, overlapping profiles were also observed in right PPC (data not shown). A, anterior; L, lateral; M; medial; P, posterior.

combination of the three parameters (P > 0.18; Supplementary Fig. 1 online), demonstrating that any differences in sniff performance cannot account for the differential activity found in the PPC.

Experiment 2: categorical odor quality coding in PPC

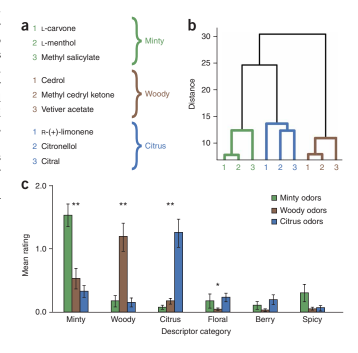
The results from Experiment 1 are consistent with the idea that information about odor quality can be extracted from multi-voxel patterns of fMRI activity in human PPC (and OFC). However, to the extent that only one stimulus exemplar per odor category was used, the possibility remains that the observed fMRI effects may largely reflect odorant-specific differences rather than more generalized differences in odor qualities and categories. Therefore, we conducted a second independent experiment using a more diverse set of odorants. This study, in conjunction with additional psychophysical measures and multidimensional scaling techniques, enabled us to definitively characterize ensemble coding of odor quality categories and to quantify how well these fMRI codes coincide with perceived odor quality on an odorant × odorant and subject × subject basis.

In this experiment, four additional participants smelled three exemplars for each of three odor quality categories: minty (R-(-)carvone, L-menthol and methyl salicylate), woody (cedrol, methyl cedryl ketone and vetiver acetate) and citrus (citral, citronellol and R-(+)-limonene) (Fig. 5a). Pair-wise similarity ratings of odor quality verified that subjects successfully grouped this nine-odorant set into three odor categories, as assessed using hierarchical cluster analysis (Fig. 5b). In addition, another ten independent participants, using a standardized questionnaire³¹ to rate the applicability of 146 odor descriptors to each stimulus (see Online Methods), robustly classified the odorants into the relevant perceptual categories chosen a priori (minty category: $\chi^2 = 16.63$, P = 0.0002; woody: $\chi^2 = 16.62$, P = 0.0003; citrus: $\chi^2 = 16.92$, P = 0.0002; Friedman test; **Fig. 5c**). A weaker category effect was also observed for the floral descriptors $(\chi^2 = 6.74, P = 0.034)$, although the 'floral' ratings were significantly

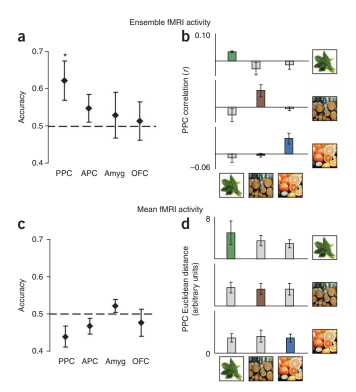
Figure 5 Odor stimuli and psychophysical ratings for Experiment 2. (a) The nine odorants included three stimuli for each of three quality categories (minty, woody and citrus). (b) A dendrogram plot obtained from cluster analysis of the mean pair-wise similarity ratings of odor quality revealed that the nine odorants sorted into three quality categories. Shorter distances indicate greater similarity. (c) Ratings of the applicability of 146 odor descriptors to the odor stimuli (descriptors pre-sorted into six different quality categories) indicated that subjects classified the odorants into the appropriate categories (mean \pm between-subjects s.e.m.; n = 10). Nonparametric Friedman tests for related samples were conducted separately on each category (* P < 0.05, ** P < 0.005).

lower than the 'minty' ratings for minty odorants (Z = 3.67, P = 0.0002, Wilcoxon test), the 'woody' ratings for woody odorants (Z = 3.87, P = 0.0001) and the 'citrus' ratings for citrus odorants (Z = 2.88, P = 0.004). In fact, the floral ratings were not different from the woody (Z = 0.51, P = 0.61) or citrus (Z = 0.34, P = 0.73) ratings for the minty odorants, nor were they different from the minty (Z = 0.57, P = 0.57) or woody (Z = 0.99, P = 0.32) ratings for the citrus odorants. Together, these data show that the stimulus set closely conforms to the perceptual categories of odor quality proposed here. Finally, there were no substantial category differences in behavioral ratings, sniffing or mean PPC activity (Supplementary Fig. 2 online).

During fMRI scanning, the nine odorants were presented to the subjects 24 times each, spread over 3 d. Subjects tacitly identified the quality of the odor on each trial, and we organized patterns of odorevoked brain activity, extracted from the same four regions as in Experiment 1, into vectors of voxel activity. The entire set of pattern vectors was divided into even and odd runs and linear correlations were then calculated between every possible odorant pair (total of 72 unique correlations). Finally, classification accuracy was computed by testing the proportion of within-category correlations (for example, mint-1/







even versus mint-2/odd) that were greater than across-category correlations (for example, mint-1/even versus woody-1/odd). Classification was significantly above chance across the group in PPC ($T_3 = 2.34$, P = 0.050), but not in other regions (APC: $T_3 = 1.31$, P = 0.14; amygdala: $T_3 = 0.48$, P = 0.33; OFC: $T_3 = 0.27$, P = 0.40) (**Fig. 6a**). In addition, the average within-category correlation was greater than the across-category correlation only in PPC ($T_3 = 3.46, P = 0.041$; **Fig. 6b**), but not in APC ($T_3 = 0.78$, P = 0.49), amygdala ($T_3 = 0.47$, P = 0.67) or OFC ($T_3 = 0.47$, P = 0.67). A one-way ANOVA testing for an effect of quality category on the within versus across correlation differences in PPC was not significant ($F_{2.9} = 0.30$, P = 0.75), implying that the above results were not driven by one particular category.

Following the same methods outlined in Experiment 1, we performed a split-halves classification analysis to determine whether mean fMRI activity contains reliable information about olfactory perceptual categories. Identification accuracy, calculated as the proportion of within-category distances that were shorter than across-category distances, was not significantly different from chance in any of the four regions of interest (P = 0.44, 0.39, 0.16 and 0.22 for PPC, APC, amygdala and OFC, respectively; Fig. 6c). Furthermore, there were no significant differences in PPC between within-category and acrosscategory distances ($T_3 = -1.23$, P = 0.31; Fig. 6d). These data provide persuasive support for the idea that local (ensemble) fMRI activity in PPC, but not global (mean) fMRI activity, contains discriminating information about odor quality categories. At the same time, odor classification accuracy was not greater than chance when trial-specific respiratory parameters were entered into split-halves analyses (Supplementary Fig. 1), indicating that sniff peak amplitude, duration and volume (and their combination) are unable to account for the categorical effects in PPC.

In a subsequent analysis, we used multidimensional scaling (MDS) techniques³² to test the hypothesis that multi-voxel patterns of PPC activity coincide with perceptual ratings of odor quality on an odorant \times odorant basis. There were two related predictions. First, odorants sharing a high (or low) degree of spatial overlap in PPC should be

Figure 6 fMRI pattern discrimination of odor categorical perception in PPC. (a,b) Classification performance calculated using fMRI patterns of ensemble activity. Odor identification accuracy (mean \pm between-subjects s.e.m; n = 4) was significantly greater than chance in PPC only (a). * P < 0.05. The within-category correlation was greater than the across-category correlation in PPC for all three odor quality categories, an effect that was separately observed for each category (b). (c,d) Classification performance calculated using fMRI mean activity. Identification accuracy (c) did not significantly differ from chance in any of the four regions, and there was no group difference between within-category and across-category Euclidean distances in PPC (d) or in APC, amygdala or OFC (data not shown).

perceived as smelling more (or less) similar in quality. Second, odorants showing greater spatial overlap in PPC should be perceived as belonging to the same odor category. To implement this procedure, we assembled two 'distance' matrices out of the nine-odorant dataset: a 9×9 imaging matrix composed of the multi-voxel fMRI correlations (averaged across subjects) in PPC for every odorant pair and a 9×9 perceptual matrix composed of perceived differences in odor quality between every odorant pair. We then used classical MDS to project the distance matrices onto a common three-dimensional space, followed by a standard linear transformation algorithm (Procrustes alignment) to determine how well the imaging matrix aligned with the perceptual matrix.

The MDS analysis demonstrated robust spatial correspondence between the projected PPC imaging map and the projected perceptual map (Fig. 7a); odorants belonging to the same category clustered together in each map, and clustering of odor quality categories (as well as of individual odorants) was closely aligned between the imaging and perceptual maps. The statistical significance of this latter effect was tested by calculating the 'goodnessof-fit' (that is, the sum of squared errors arising from alignment of the perceptual and imaging datasets, where lower values indicate stronger fits). This parameter was plotted against a distribution of goodness-of-fits, generated by randomly shuffling the identities of each of the nine odors (10,000 iterations). The observed goodnessof-fit in PPC (0.53) was situated outside the lower bound of the 95% confidence interval of the randomly generated distribution (Fig. 7b), demonstrating a significant alignment between imaging and perception in this region. In contrast, when imaging matrices of APC, amygdala and OFC were each aligned to the perceptual matrix, the goodness-offit (using the same methods) was not statistically significant (P > 0.05; Fig. 7b). These findings reinforce the idea that ensemble coding of odor categorical perception is regionally specific for PPC.

It is important to note that the three stimulus exemplars in each of the three odor categories are chemically and structurally diverse. Nonetheless, the possibility that unforeseen molecular similarities could explain some of the observed correlation effects still remains. To examine this issue, we obtained a set of 32 optimized molecular descriptors for each odorant that have been shown to account well for variations in neural activity observed in the olfactory epithelium and olfactory bulb of different animal species³³. Pair-wise Euclidean distances between each odorant were calculated from the set of molecular descriptors and used to assemble a 9 × 9 molecular-distance matrix, similar to the ones assembled for the imaging and perceptual data. We then implemented the permutation analysis as described above, but this time using the molecular-distance matrix as the alignment target in place of the perceptual matrix. The observed value of goodness-of-fit between imaging and molecular features fell inside of the 95% confidence interval of the randomly generated distribution in PPC, APC, amygdala and OFC (Supplementary Fig. 3 online), indicating that relationships between odorants in molecular space cannot easily



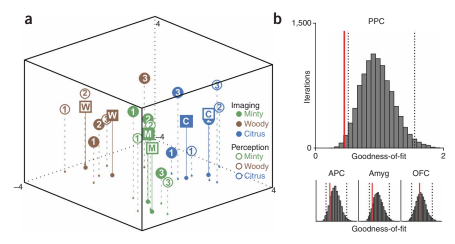


Figure 7 Alignment of fMRI spatial patterns and perceived odor quality. (a) The group-averaged imaging and perceptual datasets were each projected onto a common three-dimensional space using MDS, indicating that the imaging maps of PPC linear correlations (filled circles) closely overlapped with the perceptual maps of odor quality similarity (empty circles), both for individual odorants and for odor quality categories. Squares labeled 'M' (minty), 'W' (woody) and 'C' (citrus) represent centroids of each category for the imaging (solid squares) and perceptual (empty squares) data. (b) The observed goodness-of-fit in PPC (red line) fell outside the lower bound of the 95% confidence interval (dashed lines) of a randomly permuted distribution of goodness-of-fits, demonstrating a significant alignment between PPC activity and perception in this region. Alignment between imaging and perception was not significant in APC, amygdala or OFC (P > 0.05).

account for relationships between odorants in imaging space, at least with respect to these particular olfactory brain regions.

Finally, we combined MDS with multiple linear regression analysis to evaluate whether there is a consistent predictive relationship between PPC ensemble activity and odor quality perception across subjects. This question was tested by considering the three-dimensional MDS imaging projections of the PPC correlations as perceptual estimates of odor quality, closely following previously described methods for predicting odor perception from olfactory bulb (or olfactory mucosal) activity in rats^{11,34}. Notably, we used a 'leave-one-subject-out' approach to maintain data independence, such that the average PPC imaging correlation matrix of three subjects was used to estimate the perceptual matrix of a fourth (test) subject, with fourfold crossvalidation, preventing the test set from influencing its own prediction. Multiple linear regression analysis showed that there was a highly significant predictive relationship between estimated odor quality and actual odor quality ($R = 0.44, F_{1,24} = 18.84, P = 0.001$) in the absence of significant across-subject variability (R = 0.16, $F_{3,24} = 0.61$, P =0.615), suggesting that the capacity of PPC to encode information about odor quality generalizes across subjects.

DISCUSSION

Here, we integrated high-resolution olfactory fMRI, cortical flattening techniques, sensory psychophysical assays and multivariate analyses to provide measurements of odor-evoked piriform spatial activity patterns and odor quality perception in the same set of subjects. Until now, pattern-based techniques have not been successfully implemented in the context of human olfactory imaging (but see ref. 19), in spite of their widespread use to delineate odorant spatial mapping in animal imaging studies of the olfactory bulb^{35–37}. The methods outlined here have enabled us to consider a set of research questions that were not previously testable and are centered about the idea that ensemble fMRI activity patterns may represent a viable signature of sensory perception that can be used to infer olfactory perceptual experience. That odor classification could be reliably attained from multi-voxel fMRI patterns,

but not from mean fMRI responses (Figs. 3 and 6), indicates that multivariate measures of fMRI ensemble activity are particularly suited to extract discriminating information about odor qualities and categories from the human olfactory brain.

These results show that the spatial arrangement of fMRI activity in PPC is distributed and overlapping for each odorant, without any obvious local clustering (Fig. 4), consistent with the known anatomical organization of this region^{15,16}. Odorants belonging to the same perceptual category showed similar pattern topography (Fig. 7), suggesting that perceptual information about odor quality may be reflected in distributed ensemble activity in PPC. Overall, our findings suggest that olfactory codes of odor object categories are arranged in much the same way that visual object categories (for example, houses, cows and chairs) are organized in inferotemporal cortex²⁰, highlighting the critical sensoryassociative nature of PPC. It is reasonable to speculate that the degree of correlation between odor-evoked PPC spatial patterns and preexisting odor 'templates'38 would provide a

convenient metric by which the brain could infer similarities among odorants and classify odor objects into discrete, meaningful categories.

As a point of clarification, references here to 'spatially distributed' codes and patterns are not meant to imply that the brain is abstracting and codifying information about the spatial location of odors in the external world. Such definitions are highly relevant to neural coding of visual and auditory stimuli, but are probably less pertinent to the olfactory modality (for further discussion of this topic see ref. 39). Rather, as used in the present context, a spatially distributed olfactory code refers to a spatial ensemble representation of odor-evoked activity dispersed across a set of fMRI voxels. Ultimately, what matters is the formation of a unique spatial (or spatiotemporal) pattern of cortical activity that reliably encodes the perceptual identity of an odorant, consistent with 'content-addressable memory' models that are predicated on unique distributed neuron ensembles in piriform cortex^{14,18,40,41}.

Notably, odor-specific fMRI ensemble patterns were also reliably observed in OFC, but this finding was restricted to the first experiment (Fig. 3b,c). Considering that Experiment 1 included only one stimulus exemplar per odor category, the multivariate data across both experiments suggest that PPC and OFC each contain fMRI ensemble representations of individual odorants (odorant identity), but PPC alone contains ensemble representations of odor perceptual category (odor quality). That we observed effects across diverse odor classes (minty, woody and citrus) substantiates the idea that PPC is an important substrate where categorical perceptions of odor objects are maintained (see Supplementary Discussion online). On the other hand, it remains unclear whether OFC may encode some discriminating perceptual feature other than quality, but the idea that the neural fidelity of individual odorants is preserved in OFC is consistent with single-unit recordings in monkeys³⁰, showing that the specificity of odor tuning also happens to be highest in this region.

Data from Experiment 2 demonstrated a statistically significant relationship (P < 0.05) between PPC spatial patterns and subjective similarity ratings of odor qualities and categories. Indeed, to the extent that PPC ensemble maps correlate with subjective perceptions of odor

quality, these results suggest that it might be possible to read out olfactory percepts from spatially distributed fMRI activity in PPC. The pattern regression analysis further extended these findings by showing that perceptual estimates of odor quality can be inferred from group-based three-dimensional MDS projections of PPC imaging correlations. That the perception of odor object categories can be estimated from group-averaged PPC correlations would fulfill an important criterion⁴² for establishing an olfactory code with direct relevance for odor quality perception. The ability of multivariate techniques to draw out these predictive relationships highlights a previously unknown and robust method for characterizing the direct links between olfactory codes and odor percepts, and perhaps for assessing (or predicting) individual human variability in how an odorant's quality is perceived.

Several influential models of olfactory perception 14,17,18,38,40 posit that a spatially (or spatiotemporally) distributed pattern of brain activity satisfies the requirements of an odor detection system faced with the challenge of extracting perceptual constancy from environmental inconstancy. Ecological variations in odor background, wind direction and sniff sampling phase, as well as physical alterations in the odor source itself, can all degrade the stimulus fidelity of the original input. The prevailing idea that pattern-completion mechanisms and content-addressable memory systems^{14,18} can resolve corrupted odor inputs hinges on the presence of ensemble activity patterns in olfactory cortical structures, but these theories are principally derived from anatomical data and network simulations, in the absence of confirmatory functional data. Our findings are among the first to verify the presence of pattern-based perceptual representations of odor qualities and categories in human PPC, which would be indispensable for the reconstruction of odor objects, ensuring perceptual invariance in the wake of fragmentary inputs.

METHODS

Methods and any associated references are available in the online version of the paper at http://www.nature.com/natureneuroscience/.

Note: Supplementary information is available on the Nature Neuroscience website.

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AUTHOR CONTRIBUTIONS

J.A.G. conceived the experiment, with contributions and methodological suggestions from J.-D. Haynes. J.D. Howard and J.P. collected the imaging and behavioral data. J.D. Howard, J.P. and J.A.G. analyzed the data. M.G., J.-D. Haynes and J.D. Howard implemented the flat map analysis. J.A.G., J.D. Howard and J.P. wrote the manuscript.

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ONLINE METHODS

Subjects. We obtained informed consent from 20 subjects (age range, 22–35 years) to participate in this study, which was approved by the Northwestern University Institutional Review Board. Six subjects (four women) participated in Experiment 1, four subjects (three women) participated in Experiment 2 and ten subjects (eight women) completed odor questionnaires for Experiment 2.

Behavioral ratings. Prior to the first scanning day, subjects rated odor intensity, valence, pungency and familiarity using visual analog scales^{43,44}. Ratings were analyzed in Matlab (Mathworks) using one-way ANOVAs. Subjects also rated odor quality similarity between all possible odorant pairs on a visual analog scale (anchored by 'not alike at all' and 'identical'). In Experiment 2, the 36 pair-wise similarity ratings, averaged across subjects, were transformed into a hierarchical cluster tree using a single-linkage algorithm in Matlab and plotted as a dendrogram, with linkage distances (reflecting degree of quality similarity between odorants) indicated on the y axis.

Ten independent participants completed a 146-item Odor Quality Evaluation questionnaire³¹ for each odorant in Experiment 2. Ratings of the applicability of odor descriptors on the questionnaire ranged from 0 ('absent') to 5 ('extremely'). Before acquiring ratings, and following prior techniques^{24,45}, we identified the descriptors best fitting the three perceptual categories (minty, woody and citrus) and three control categories (floral, berry and spicy) (**Supplementary Table 1** online). For each descriptor category, odorant-specific ratings were averaged across descriptors belonging to that category and were then averaged across the three odorants pre-assigned to each perceptual group. For example, all of the minty descriptor ratings were averaged together for each odorant and were then averaged across the three putative minty odorants, the three putative woody odorants and the three putative citrus odorants to form mean ratings for each odorant perceptual category. The applicability of each descriptor rating to each perceptual category was tested (Friedman tests for related samples and Wilcoxon sign-rank tests for paired samples).

The citronellol odorant used in both Experiments 1 and 2 was a racemic mixture of the (R)-(+)- β -citronellol isomer, which is reportedly 'citrus like' in quality, and the (S)-(-)- β -citronellol isomer, which is reportedly 'geranium like'. We acquired additional behavioral ratings from these same ten independent participants to confirm that the odor quality of this racemic mixture most closely resembled the citrus character (**Supplementary Fig. 4** and **Supplementary Data** online).

Respiratory monitoring and analysis. Subjects were affixed with breathing belts to monitor respirations during scanning⁴³. Sniff peak amplitude, duration and inspiratory volume were computed for each trial, averaged across conditions and runs, normalized within subjects by subtracting the mean parameter value (across conditions) from each condition-specific value, and then entered into individual one-way ANOVAs for statistical analysis.

fMRI data acquisition. Gradient-echo T2-weighted echoplanar images were acquired with a Siemens Trio 3T scanner using parallel imaging and an eight-channel head-coil (repetition time, 2 s; echo time, 20 ms; matrix size, 128×120 voxels; field-of-view, 220×206 mm; in-plane resolution, 1.72×1.72 mm; slice thickness, 2 mm; gap, 1 mm; acquisition angle, 30° rostral to the intercommissural line). A 1-mm T1-weighted MRI scan was obtained to define anatomical regions of interest (ROIs).

Multivariate fMRI procedure. In both experiments, there were eight fMRI runs on each of 3 consecutive d (total of 24 runs). In Experiment 1, each odorant was presented once per run for 10 s, with a 20-s rest between odorants (Fig. 1). In Experiment 2, each odorant was presented once per run for 6 s with a 12-s rest. Odorants were presented using an MRI-compatible olfactometer (airflow, 2.5 l min⁻¹)²⁴. On odorant presentation, subjects were visually cued to sniff and to identify the odor quality covertly. Each run was separated by approximately 180 s to minimize olfactory fatigue. Odorant presentation was counterbalanced across runs such that each possible stimulus order was used once in Experiment 1, and each odorant appeared only once per run and a different stimulus order was used for each run in Experiment 2.

Odor localizer scan. We used an independent 'odor localizer' fMRI scan (using the same imaging parameters as the main experiment) as an unbiased way to

identify odor-active voxels⁴⁴. In Experiment 1, subjects undertook an odor detection task on each scanning day consisting of 12 odor and 12 no odor (air only) trials using four odorants (butanol, anisole, heptanol and α -ionone). In Experiment 2, two odor localizer runs were performed on a fourth day of scanning. Odor and no odor trials were presented for 2 s (stimulus-onset asynchrony, 12 s), followed by a button press to indicate whether an odor was present or absent. Images were spatially realigned, smoothed (6-mm full-width half-maximum) and then analyzed using the general linear model (GLM) in SPM2 (http://www.fil.ion.ucl.ac.uk/spm/). After model estimation, we contrasted odor versus no odor conditions, collapsed across runs, to generate subject-specific statistical parametric maps of odor-active cortex.

Voxel selection procedure. Voxels included in the multivariate analyses were selected from anatomically defined ROIs, manually drawn on each subject's structural MRI scan using MRIcroN software (http://www.mricro.com/) (Supplementary Fig. 5 online). Anatomical definition of PPC, APC and amygdala was guided by a human brain atlas⁴⁶. Delineation of OFC was based on an olfactory fMRI meta-analysis²⁷. Subsequently, the voxels in each subject's ROI were functionally ranked according to their T values (obtained from the independent odor localizer scans), specifically without reference to the pattern-based fMRI data or to odor classification performance. For each ROI, we considered the N-most odor-active voxels for which anatomical data were maximally available from all subjects, such that N was set on the basis of the subject with the smallest anatomical ROI. Thus, if the PPC of subjects 1, 2 and 3 contained 200, 150 and 300 voxels, respectively, then N=150 voxels. In this manner, the size of each ROI was different across regions, but the same across subjects for a given ROI. The respective voxel numbers used for analysis of PPC, APC, amygdala and OFC were 186, 138, 312 and 195 (Experiment 1), and 228, 155, 339 and 239 (Experiment 2).

Multivariate fMRI data pre-processing. After discarding the first six 'dummy' volumes of each run, we spatially realigned all functional images to the first volume of the first run using SPM2. Subsequent steps, including extraction of fMRI signal intensity from each voxel in an ROI for each run, temporal detrending with a second-order polynomial⁴⁷ and assembly into linear vectors of odor-specific voxel activity, were carried out outside of SPM2, using customized scripts in Matlab. To preserve the native spatial fidelity of the fMRI signal, we did not perform image normalization and smoothing. Control analyses indicated that spatial realignment across the three scanning days was robust (Supplementary Fig. 6 and Supplementary Data online).

Multivariate fMRI analysis. We extracted 96 pattern vectors (Fig. 1) for Experiment 1 (one vector for each of the four odorants across 24 runs) and 216 pattern vectors for Experiment 2 (nine odorants, 24 runs). In both experiments, pattern vectors were split into even and odd halves and the mean activity across all conditions was subtracted from each vector in that half²⁰. These mean-corrected vectors were averaged across runs in each half, producing one pattern vector per condition per half, which were then used to calculate pair-wise correlation coefficients within and between odorants (Experiment 1) and odor quality categories (Experiment 2).

Within-odor and across-odor correlations were compared directly and were also used to estimate identification accuracy 20 . In Experiment 1, identification accuracy was calculated such that, for odor A, if $(A_{\rm even}$ versus $A_{\rm odd})$ was greater than $(A_{\rm even}$ versus $B_{\rm odd})$, it was counted as a correct identification. There were six such comparisons for each odor (given four unique odor types), resulting in 24 comparisons. In Experiment 2, identification accuracy was calculated such that if a within-category correlation (for example, minty_even versus minty_odd) was greater than an across-category correlation (for example, minty_even versus woody_odd), it was counted as a correct identification. Thus, for each of the three odor quality categories, there were three within-category correlations and 18 across-category correlations, resulting in 54 comparisons per quality category. The pair-wise correlations (total of 72 correlations, excluding on-diagonal correlations to prevent estimation bias of the within-category correlations) were calculated before mean within- and across-category correlation coefficients were computed. Chance identification accuracy was 50%.

Piriform flat maps. In the first step, we specified a GLM using SPM2 software that included odor onset times from Experiment 1 as regressors of interest

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(collapsed across runs). Percent signal change was then estimated for each condition. Flattened cortical maps were made using mrVista software (http://white.stanford.edu/software/). In this process, each subject's T1-weighted anatomical image was segmented to isolate gray matter in PPC, followed by spatial flattening into a two-dimensional map. The functional volumes containing voxel-wise percent signal change information from the GLM were then coregistered to the T1 volume and flattened to two dimensions using the same transformation parameters.

MDS. MDS is commonly employed to measure similarities among complex, high-dimensional datasets containing non-independent elements³². We used it to facilitate comparisons and statistical analysis between fMRI and perceptual data acquired in Experiment 2. First, a 9 × 9 imaging dissimilarity (distance) matrix for each ROI was created for each subject by subtracting the pair-wise linear correlation coefficients from a value of 1 and scaling from 0 to 10. Second, a 9 × 9 perceptual dissimilarity (distance) matrix was created using each subject's similarity ratings of perceived odor quality for every odorant pair. These distance matrices were each averaged across subjects and entered into classical MDS analyses (Matlab). This generated three-dimensional maps (one for the imaging data, one for the perceptual data), with each odorant being represented by a unique coordinate in xyz space. Finally, the three-dimensional map of imaging coordinates was aligned to the perceptual map using Procrustes analysis, which provides a quantitative measure of similarity (goodness-of-fit) between two sets of coordinates (ranging between 0 and 1, where lower values indicate better alignment). Selection of a three-dimensional MDS space was based on a 'scree plot' of the perceptual dataset indicating that a three-dimensional projection best captured the variance in the nine-dimensional matrix.

To estimate the significance of goodness-of-fit, we implemented a random permutation (Monte Carlo) procedure, whereby the actual (measured) fit was compared with a distribution of fits generated by randomizing the odor condition assignments before MDS analysis and Procrustes alignment (10,000 iterations). This procedure scrambles the assignment of odorant condition (and category) in advance of MDS, preserving the values of all pair-wise distances and minimizing overfitting of the data. Randomization statistics have been applied to studies comparing multivariate response profiles between different brain regions or between neural and perceptual datasets $^{48-50}$. An observed fit between imaging and perception was considered to be significant at P < 0.05, indicating that the observed fit was smaller (better fit) than 95% of randomly generated fits.

For presentation, the goodness-of-fits, d, were transformed to normal values using Fisher z transformation:

$$z = \frac{1}{2} \ln \frac{(1+d)}{(1-d)}$$

Odor classification: mean fMRI activity. To test whether odor-specific information is reliably contained in the mean fMRI signal amplitude, we performed a split-halves classification analysis that closely paralleled the methods employed to analyze fMRI ensemble activity. First, the raw BOLD signal elicited by each odorant presentation was averaged across all voxels from each ROI for each trial, resulting in 24 values per condition (corresponding to the 24 data acquisition runs). Mean signal across conditions in each run was subtracted from the condition-specific values in that run for normalization. Normalized signals were split into halves, one containing values from even runs and one from odd runs. Data in each condition and half were averaged across runs and pair-wise Euclidean distances were calculated within and between conditions.

Odor classification: sniff parameters. To test whether trial-specific sniff parameters could be used to successfully classify the odors, a split-halves analysis was performed on sniff peak, duration and inspiratory volume. In addition, the three sniff parameters were combined into one three-component vector (analogous to vectors of voxel activity used in the pattern-based fMRI analyses) to test whether the combination of these parameters might contain odor-specific information. Data were normalized by subtracting the mean parameter value across conditions in each run from the condition-specific values. Data were then split into halves according to even and odd runs and averaged across runs in each half. Pair-wise Euclidean distances were calculated within and across conditions.

Statistical analyses. Results are shown as mean \pm s.e.m. for participants or conditions. Unless otherwise indicated, statistical significance was determined using one-tailed t tests (when comparing identification accuracy to chance), two-tailed t tests (when comparing two conditions) or ANOVAs (when comparing more than two conditions). Results were considered to be significant at P < 0.05.

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