Dynamic coding of choice targets in the perirhinal cortex

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Abstract

Cortical neurons flexibly respond to multiple task-events characterized by distinct computational aspects. There are two possible explanatory scales, dynamic population code realizing computational flexibility and single-neuron code associating relevant information across task events. Since these interpretations were independently developed, it is unclear how neurons contribute to population dynamics with their individual ability to integrate relevant information. Here we show systematic changes in single-neuron representations of choice targets shape population dynamics. We analyzed neural activities in the perirhinal cortex while rats performed a two-alternative target-choice task. A time-resolved pattern analysis indicated dynamic population code of choice targets with two time-stable states during the cue and reward periods. The population patterns during these periods were substantially reversed and therefore allowed us to decode task-periods as well as choice targets. Our results clarify the contribution of single-neuron representations to population dynamics, suggesting its potential advantages such as reconciling different computational demands.
Introduction

Individual neurons across cortical areas show temporally flexible responses to multiple task-events characterized by different computational aspects, such as cue, action and reward (e.g., Sakurai et al., 2017; Sakurai, 1990a,b). Recent studies indicate that such complex single-neuron responses are only interpretable in terms of their contribution to population dynamics which flexibly perform different computations, particularly in association and motor cortices (Elsayed et al., 2016; Lara et al., 2018; Mante et al., 2013; Raposo et al., 2014). On the other hand, in some cortical areas, it has been shown that individual neurons respond to multiple task-events while reflecting their associations (Perirhinal cortex: Eradath et al., 2015, Motor cortex: Guo et al., 2017, Insular cortex: Gardner and Fontanini, 2014, Anterior cingulate cortex: Kennerley and Wallis, 2009, and Orbitofrontal cortex: Furuyashiki et al., 2008; Hirokawa et al., 2017; Hosokawa et al., 2005), suggesting that some cortical neurons afford to integrate relevant information across different task-events. Therefore, it is still an open question how such neurons contribute to population dynamics which would be suitable for differentiating task events.

In the present study, we explored neural activities in the perirhinal cortex (PRC) which has been implicated in association memory of objects (Ahn and Lee, 2017, 2015; Naya et al., 2003a, 2003b, 1996; Sakai and Miyashita, 1991). This region receives sensory inputs from almost all modalities, behavioral context information from the prefrontal cortex and reward-related signals from the amygdala (Burwell, 2001; Burwell and Amaral, 1998; Furtak et al., 2007; Tomás Pereira et al., 2016), creating integrated object representations (Ohyama et al., 2012; Qu et al., 2016; Taylor et al., 2006). Some studies show that some PRC neurons respond to the same visual cue across time contexts but in different degrees of strength (Naya et al., 2017; Naya and Suzuki, 2011). Therefore, it is possible that the PRC is able to represent the same target even across explicitly different task-events such as cue, action and reward, while differentiating those task-events.

To determine this possibility and elucidate underlying mechanisms, we analyzed single-unit activities recorded from the PRC while rats performed a two-alternative target-choice task. We used randomly-interleaved visual and olfactory cues to investigate choice-target representations. We found that
dynamic population code for choice targets evolved through the task. Systematic shift of the choice-target code represented task-periods and was meditated by neurons characterized by activation across the task-periods. Our results clarify the contribution of single-neuron responses to population dynamics, suggesting potential advantages of flexible but structured single-neuron responses as encoding strategy.

Results

We trained four rats to perform a two-alternative target-choice task where they chose a target port (left/right) associated with a presented cue to obtain reward (Figure 1a–b). The task performance was similar level regardless of the cue modality (mean correct rate in visual trials = 94.9±5.8 %; olfactory trials = 93.3±3.8 %). We recorded spiking activities from the PRC (n = 182 neurons) during the task performance.

The PRC neurons frequently encoded choice targets during the task performance. We used ROC analysis to generate a target-preference index which quantifies preferential responses of a neuron to a choice target. We focused on two task-periods where prominent target-encoding was observed (Figure 1c–d), the cue period (−400 to 0 ms before withdrawal) and the reward period (200 to 600 ms after choice), and summarized neural responses during these task-periods (Figure 1e). The target preference within each task-period was highly consistent across the cue modalities, indicating target processing beyond sensory modalities (r = 0.59, P < 0.001 for the cue period; r = 0.86, P < 0.001 for the reward period in Figure 1e). Similar amount of neurons activated in either or both of these task-periods (Figure 1f). Interestingly, we found that the majority of the neurons activated across both task-periods (64%; 42 of 66 neurons) dynamically changed their target-preference, while a smaller fraction (36%; 24 of 66 neurons) sustained target preference across the task periods (Figure 1f).

To investigate temporal dynamics of the target code, we employed a time-resolved pattern analysis (Stokes et al., 2013). Using this method, we decoded choice targets from the population response in a modality-independent manner (n = 182 neurons). The population code evolved through the task with two time-stable states (Figure 2a). Target code
was sustained during the presentation of the cue and was followed by a transient response during movement toward a target port. Soon after the rats chose a target port, the target code settled again into a time-stable state (Figure 2a). We computed mean performance of the classifier during the cue and reward periods and compared them with the baseline (−400 to 0 ms before cue onset). As shown in Figure 2b, we successfully decoded choice targets in both task-periods ($P < 0.01$), indicating that the PRC tracks target information across task-periods. Furthermore, in Figure 2c, we found decreased target-code when we decoded choice targets using correct visual trials but erroneous olfactory trials ($n = 89$ neurons). To quantify this effect, we computed the mean classification performance in correct and erroneous trials. As shown in Figure 2d, the mean performance in both task-periods decreased to chance level in erroneous trials despite the fact that animals chose the same target-port as correct trials (cue period: $P \approx 0.10$; reward period: $P \approx 0.24$). These results indicate that the target code was relevant to successful task performance.

Importantly, population patterns during these two task-periods were substantially reversed (white arrows in Figure 2a), suggesting that the PRC distinguished different task-periods by systematic shift of the target code. We computed mean classification performance across the two task-periods and found significant shift of the target code ($P < 0.01$; Figure 2e). To clarify whether the shift depends on single-neuron responses or population-level structure, we shuffled the correspondence between the cue-period and reward-period responses among the neurons. This procedure resulted in collapsed shift patterns (Figure 2e), indicating that changes at each single-neuron level were necessary for the shift. Additionally, the shift decreased to chance level when the 20 most contributed neurons were eliminated from the population (Figure 2f). On the contrary, if the population was pruned randomly, the shift was preserved even after the majority of neurons were removed (Figure 2f). Overall, our data indicated that the PRC dynamically modified target code between task-periods and that it depended on the flexible but structured single-neuron responses.

The shift of target code is possibly signature of the encoding strategy which allows the PRC to integrate the different information, choice targets and task periods. If it is the essential nature of the PRC rather than noise, we should be able to decode richer information from neurons contributing to
the shift than the others. We therefore repeated the pattern analysis for each of the following types (classified in Figure 1f): neurons with shifting response, sustained response, and specific response (for either task-period). As expected, the neurons with shifting response showed clear target-code in both task-periods, as well as a dramatic change of it between the task periods (Figure 3a). In Figure 3b, we computed mean classification performance for each type and found that only neurons with shifting response showed significant performance in both task-periods ($P < 0.01$). The other types showed significant performances in the reward period ($P < 0.01$) but not in the cue period (sustained response: $P \approx 0.21$; specific response $P \approx 0.44$). Only neurons with shifting response showed significant discrimination of the task periods ($P < 0.01$; Figure 3c). These results underscore capacity of the PRC to integrate information across task-periods, by temporally flexible neural responses.

Discussion

In this study, we showed that the PRC dynamically encoded choice targets during task performance. We demonstrated that the target code was highly consistent across cue modality and reduced in erroneous trials. These results suggest that neural responses during the cue period might reflect retrieved target as shown by electrophysiological studies, with visual tasks, both in rodents (Ahn and Lee, 2017) and non-human primates (Naya et al., 2001; Naya et al., 1996). Our results thus provide substantial evidence for integration across sensory modalities which might support object memory.

We also found that the PRC reversed its target code between cue and reward periods. This allowed us to decode task periods as well as choice targets from the neural population. The decodability was achieved by flexible but structured single-neuron responses. Recent studies highlighted neurons with non-linear mixed selectivity which integrate different task-parameters at a given time (Pagan et al., 2013; Raposo et al., 2014) or across similar task-periods (e.g., Cue1 and Cue2) (Rigotti et al., 2013), suggesting that such complex selectivity supports functional flexibility of a brain region. Unexpectedly, our results indicate simply structured single-neuron responses across task-events and underscore their potential advantage as an
encoding strategy which reconciles different demands, keeping target 
information and differentiating task-events. This suggests that functional 
flexibility could be achieved by explicitly structured single-neuron responses. 
Moreover, this structured code has additional benefits. First, it can be easily 
read-out by downstream neurons in a consistent manner, enabling stable 
recognition. Second, it would support both generalization and discrimination, 
conflicting computations (Barak et al., 2013). Third, it is suitable for linking 
relevant information together through different time-points (Eradath et al., 
2015), an essential computation for associative learning.

Materials and Methods

Subjects
Four male Long-Evans rats weighting 315–360 g at the beginning of training 
were individually housed and maintained on a laboratory light/dark cycle 
(lights on 8:00 A.M. to 9:00 P.M.). Rats were placed on water restriction with 
ad libitum access to food. The animals were maintained at 80% of their 
baseline weight throughout the experiments. All experiments were 
implemented in accordance with the guidelines for the care and use of 
laboratory animals provided by the Animal Research Committee of the 
Doshisha University.

Behavioral apparatus
The behavioral apparatus (Figure 1a) has been previously described 
(Hirokawa et al., 2017; Osako et al., 2018). Briefly, an operant chamber 
(O’Hara, Tokyo, Japan) which had three ports in the front wall for nose-poke 
response was enclosed in a soundproof box (Brain Science Idea, Osaka, 
Japan). Each port was equipped with an infrared sensor to detect animals’ 
response. Visual cues were presented using white LEDs (4000 mcd; RS 
components, Yokohama, Japan) placed on the left and right walls of the 
operant chamber. Odors were presented via the central port through a 
stainless tube. Odors were mixed with pure air to produce 1:10 dilution at a 
flow rate of 600 ml/min using a custom-build olfactometer (AALBORG, 
Orangeburg, NY). Water rewards were delivered from gravity-fed reservoirs 
regulated by solenoid valves (The Lee Company, Westbrook, CT) through
stainless tubes placed inside of the left and right target-ports. We controlled
stimulus and reward deliveries and measured behavioral responses using
Bpod and Pulsepal (Sanworks LLC, Stony Brook, NY: Sanders and Kepecs,
2014).

Two-alternative target-choice task
Each trial started when the rats poked their snout into the central port
(Figure 1b). After a variable delay (200–600ms), a cue pseudo-randomly
selected from four stimuli (left/right LED for visual modality,
S(+)/R(−)-2-octanol for olfactory modality) was presented for 1 s. If the rats
successfully maintained their nose in the central port, go sound was
delivered, and animals were allowed to withdraw from the central port and
to choose either the left or right target-port based on the task rule (Figure
1a). When the rats left the center port without waiting for the go sound, the
trial was canceled and followed by a 5 s punish intertrial-interval. Only
correct choices were immediately rewarded by a drop of water (0.013 ml),
from the target port. Rats performed typically 1018±34 trials in a daily
recording session.

Training
We trained the rats step-by-step to perform the task described above. The
training period typically lasted 4 to 7 weeks. First, rats were trained to poke
into the central port and then collect the water reward (0.02 ml) from the left
or right port. We gradually extended the duration of the poke by delaying the
go sound up to 1 s after the poke onset. Next, the rats were trained to
discriminate visual cues based on the same contingencies as recording
sessions. A variable delay (200–600 ms) was inserted before the cue onset.
After the rats became able to successfully discriminate visual cues (> 80%),
they were also trained to discriminate odors based on the same contingencies
as recording sessions (> 80%). Finally, we interleaved visual and olfactory
trials within a session and trained the animals to perform the task to a
training performance criterion (> 80%).

Mixture of odors
Cue odors, S(+)/R(−)-2-octanol, were mixed together in a subset of sessions in
order to increase olfactory discrimination difficulty and thereby obtain a
sufficient number of erroneous trials. For instance we used a 60/40 ratio in a
given session, delivering odor mixture of 60% S(+)-2-octanol and 40%
R(−)-2-octanol. We kept odor discrimination accuracy constant (> 80%)
throughout recording sessions by adjusting the degree of odor mixing before
each session.

Surgery
Rats were anesthetized with 2.5% isoflurane before surgery, and it was
maintained throughout surgical procedures. We monitored body temperature,
movements and hind leg reflex and adjusted the depth of anesthesia as
needed. To keep the eyes moistened throughout the surgery, we used eye
ointment. Subcutaneous scalp injection of a lidocaine 1% solution provided
local anesthesia before the incision. The left temporalis muscle was retracted
to expose the skull during the surgery. A craniotomy was made over the
anterior part of the left PRC (AP −3.5 to −3.24 mm, ML 6.6 to 6.8 mm
relative to bregma, 3.5 to 4.0 mm below the brain surface) and a
custom-designed electrode was vertically implanted using a stereotactic
manipulator. A stainless screw was placed over the cerebellum and served as
ground during the recordings. We used the mean response of all electrodes as
reference. During a week of postsurgical recovery, we gradually lowered the
tetrodes to detect unit activities in the PRC. Electrode placement was
estimated based on depth and histologically confirmed at the end of the
experiments.

Electrophysiological recordings
A custom-designed electrode composed of eight tetrodes (tungsten wire, 12.5
µm, California Fine Wire, Grover Beach, CA) was used for extracellular
recordings. Tetrodes were individually covered by a polyimide tube (A-M
Systems, Sequim, WA), were placed with a 100 µm separation and typically
had an impedance of 250–600 kΩ at 1 kHz. Signals were recorded with
OpenEphys (Cambridge, MA) at a sampling rate of 30 kHz and
bandpass-filtered between 0.6 and 6 kHz. Tetrodes were lowered
approximately 80 µm after each recording session, and thereby independent
populations of neurons were recorded across sessions.

Spike sorting and screening criteria of units
All analyses were performed in Matlab (Mathworks, Natick, MA). To detect single-neuron responses, spikes were manually clustered with MClust (A.D. Redish) for Matlab. Only neurons met the following criteria in further analyses: (1) units with reliable refractory period (violations were less than 1% of all spikes); and (2) units with sufficient mean firing rate in the 1 s after the cue onset (> 0.5 Hz).

Selective response to choice targets

In order to evaluate selective responses to choice targets, we calculated a target-preference index (Hirokawa et al., 2011). We first grouped correct trials into four types based on cue modality (visual or olfactory) and target choice (left or right). For each modality, we independently calculated the target preference, using ROC analysis (Green and Swets, 1966). The target-preference index was obtained from the area under ROC curve (AUC) and defined as $2 \times (\text{AUC} - 0.5)$ ranging from $-1$ to $1$. In our analysis, a positive value indicated a neuron selectively fired to left target-choice, and a negative value indicated the opposite. A value of zero showed the absence of selective responses. To determine statistical significance (one-tailed, $P < 0.05$), we used bootstrapping (100 interactions).

Decoding analysis statistics

We evaluated statistical significance of decoding analysis with bootstrapping procedure (Ojala and Garriga, 2010; Parthasarathy et al., 2017). Two bootstrapped distributions whose 95th ranges for mean did not overlap were considered as significantly different. We estimated the $P$ value for this bootstrapping procedure by computing the ratio $(1+X)/(N+1)$ between the number $X$ of overlapping data points between the two distributions and the number $N$ of interactions (Ojala and Garriga, 2010; Parthasarathy et al., 2017). Since we used 100 bootstraps throughout all decoding analyses, two distributions with no overlap resulted in $P < 0.01$, and two distributions with $x \%$ overlap resulted in $P \approx x/10$.

Decoding analysis

We employed cross-temporal pattern analysis (Stokes et al., 2013; Wasmuht et al., 2018) to investigate the temporal dynamics of the target code by neural population in the PRC. Neural responses were pooled across
recording sessions to maximize the number of neural responses included in the decoding analysis. Here we refer to the pooled pseudo-population of the PRC neurons (n = 182) as full population. The instantaneous firing rate of each neuron was estimated by spike counts in a 150 ms sliding window (10 ms increment). We converted the instantaneous firing rates for each neuron into the target-preference index independently calculated for visual and olfactory trials. In this manner, we generated two independent population vectors for the full population (cell×time vector each for the cue modalities).

We obtained a pattern similarity index by calculating the Fisher-transformed Pearson correlation ($r'$) between these two population vectors. This index provided the pattern similarity for both equivalent and different time points (e.g., Figure 2a). A positive correlation was interpreted as an evidence for reliable target code irrespective of the cue modality.

To estimate mean performance values for the pattern classification analysis, we pseudo-randomly resampled neurons (the same number of neurons as neural population analyzed) and computed the target preference in visual and olfactory trials. Neural responses were aligned to withdrawal onset (from the central port) and target-choice onset (left or right). This enabled us to calculate pattern similarity indices within and across the two task-periods that were main focus of our study, the cue and the reward periods. We repeated this process 100 times to obtain a distribution of 100 different measurements of the pattern classification. To investigate neural responses during the task periods, we averaged the performance within the cue period and reward period (e.g. Figure 2b). To obtain a baseline performance, we averaged the classification performance during the baseline period, −400 to 0 ms before the cue onset. We also quantified the temporal change of neural responses between these task-periods by averaging the following two pattern classifications: pattern classification for cue-period responses in visual trials and reward-period responses in olfactory trials, and cue-period responses in olfactory trials and reward-period responses in visual trials (e.g. Figure 2e). We calculated the 95th percentile range for each of distribution. If the 95th range of 100 different pattern classification measurements within a task period did not overlap with the 95th range of baseline distribution, we considered the choice targets to be decodable from the neural population response during the task event. We considered zero to be chance level instead of the baseline distribution when we checked
Decoding analysis of erroneous trials
We included 89 neurons recorded in sessions with sufficient number of erroneous choices (at least 11 trials for both choice targets) in all analyses for erroneous trials. Due to the similarity of the target preference between different cue modalities in correct trials, we calculated the pattern similarity index using correct visual trials and erroneous olfactory trials to evaluate influence of erroneous behavioral performance on target code (Figure 2c). We computed mean performance of the pattern classification in correct and erroneous trials (Figure 2d).

Contribution of single-neuron responses to population level shift
We quantified changes of the target code between the task-periods with n−1 neural population by excluding one neuron from the full population and repeating this procedure 182 times. This enabled us to directly quantify how strongly each neuron contributed to the shift and to sort the neurons in decreasing order of contribution level. We gradually eliminated neurons that highly contributed to the shift from the full population. In this manner, we computed the classification performance across the task-periods as a function of the number of neurons excluded from the neural population (Figure 2f). This was compared with the 95th percentile range of a control distribution obtained by randomly eliminating neurons from the population (100 repetitions).

Role of different types of neurons
We randomly selected 24 neurons each for three different types of the target code (defined by target preference in the cue period and the reward period) and performed the pattern classification analysis for each type of neuron: (i) neurons with shifting response, (ii) neurons with sustained response, and (iii) neurons with specific response (Figure 3b–c). We combined 12 neurons which showed significant target preference in the cue period and 12 neurons which showed significant target preference in the reward period into subpopulations of neurons with specific response.
Author contributions
T.O., Y.S., and J.H. designed the experiments. T.O. performed the experiments and analyzed data. Y.O. analyzed data. H.M, Y.S. and J.H. supervised the project. All authors contributed to writing the manuscript.

Conflict of Interest Statement
The authors declare no competing interests.

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Figure 1. The PRC neurons encoded choice targets. (a) Schematic drawing of behavioral apparatus and cue-target associative relationships. (b) Schematic of the task timeline. Bars represent two 400 ms task-periods. (c) Peri-stimulus time histograms showing the activity of a representative neuron throughout the task. Trial types are classified according to cue.
modality and target choice as follows: blue, left target-choice; red, right target-choice; solid line, visual trials; dashed line, olfactory trials. Mean firing rates for each of the four trial types were computed in 10 ms time windows (smoothed with a Gaussian, $\sigma = 30$ ms). Neural responses were independently aligned to cue, withdrawal and target-choice onset and then reconstructed because of variable time between them. Only correct trials were included. (d) Temporal patterns of choice-target selective responses of the PRC neurons ($n = 182$) in visual (left) and olfactory trials (right). Mean firing rates were smoothed with a Gaussian ($\sigma = 150$ ms) and then converted into a target-preference index. Positive values (blue) indicate selective firings to leftward choices, and negative values (red) indicate selective firings to rightward choices. For each modality, neurons were independently sorted according to the time of their peak response. Only segments with significant target preference were shown ($P < 0.05$; 1,000 bootstraps). (e) Scatter plots showing target preference of the neural population in the cue (left) and reward (right) periods. Each point corresponds to values of a single neuron. Colors indicate significance: light gray, no selectivity; deep gray, significant in either cue modality; purple significant in both cue-modalities. (f) Distribution of neurons with different types of choice-target selective response across the cue and reward periods.
Figure 2. Time-resolved pattern analysis for choice-target code. (a) Heat map showing performance of the pattern classification analysis. White lines correspond to the onset time of cue, withdrawal, and target choice. We trained classifiers using neural responses in correct visual trials to discriminate choice targets and tested them with neural responses in correct olfactory trials (n = 182 neurons). (b) Mean classification performance during the baseline, cue and reward periods. (c) A classifier tested with erroneous olfactory trials (n = 89 neurons). (d) Mean classification performance for erroneous trials. Dashed lines indicate the 97.5th percentile value of a baseline distribution. (e) Mean classification performance across the cue and reward periods for original data and data with shuffled correspondence of those periods (n = 182 neurons). (f) Classification performances with decreasing number of neurons, removed from most to least contributed to the shift of target code. Original data is indicated as a black line. Gray shaded
area indicates 95th percentile range of data with shuffled order of neurons (100 shuffles). In box plots: center orange line, median; box limits, 25th and 75th quartiles; notch limits, \((1.57 \times \text{interquartile range})/\sqrt{n}\); whiskers, 95th percentile range (two-sided) of the distribution. Asterisks indicate statistical significance \((P < 0.01)\): the 95th percentile range of the performance distribution did not overlap with the 95th percentile range of a baseline distribution (for classification performances within a task period) or zero (for classification performances across two task-periods).

**Figure 3. Roles of different types of neurons in target code.** (a) Heat map showing performance of a classifier trained and tested using neural responses with shifting target preference between the cue- and reward period \((n = 42 \text{ neurons})\). (b) Mean classification performance with three different types of neurons. 24 neurons were included in each different type. Dashed lines indicate the 97.5th percentile value of the baseline distribution. (c) Mean classification performances across the two task-periods. In box plots: center orange line, median; box limits, 25th and 75th quartiles; notch limits, \((1.57 \times \text{interquartile range})/\sqrt{n}\); whiskers, 95th percentile range (two-sided) of the distribution. Asterisks indicate statistical significance \((P < 0.01)\): the 95th percentile range of a performance distribution did not overlap with the 95th percentile range of a baseline distribution (for classification performances within a task period) or zero (for classification performances across two task-periods).