Successful choice behavior is associated with distinct and coherent network states in anterior cingulate cortex

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Successful decision making requires an ability to monitor contexts, actions, and outcomes. The anterior cingulate cortex (ACC) is thought to be critical for these functions, monitoring and guiding decisions especially in challenging situations involving conflict and errors. A number of different single-unit correlates have been observed in the ACC that reflect the diverse cognitive components involved. Yet how ACC neurons function as an integrated network is poorly understood. Here we show, using advanced population analysis of multiple single-unit recordings from the rat ACC during performance of an ecologically valid decision-making task, that ensembles of neurons move through distinct coherent and dissociable states as the cognitive requirements of the task change. This organization into distinct network patterns with respect to both firing-rate changes and correlations among units broke down during trials with numerous behavioral errors, especially at choice points of the task. These results point to an underlying functional organization into cell assemblies in the ACC that may monitor choices, outcomes, and task contexts, thus tracking the animal’s progression through “task space.”

There is now accumulating evidence that the transient organization of neurons into dynamic ensembles and the sequential transitions among them may form the basis for cortical information processing (2, 3, 5, 7, 13–15).

Although such processes have been investigated in some depth for perceptual and spatial domains, much less is known regarding the network dynamics that govern higher-order cognitive processes, such as action and outcome monitoring. The anterior cingulate cortex (ACC) has been the subject of increased interest as a region that plays a role in action monitoring, a supervisory cognitive function that is especially important for optimal decision making in challenging and novel situations (16–19). A number of diverse single-unit correlates accompanying these processes have been observed in the ACC (20–27), yet it is unknown how single neurons organize into functional networks that could serve these functions.

In the present study, we investigated functional ensemble dynamics within the ACC during the performance of the delayed win-shift radial-arm maze, a task with distinct cognitive phases and high ecological validity (28). The locations of rewards changed between and within trials, such that the animal had to continually monitor its own actions and track the changing environmental contingencies as it progressed through the task. Using sophisticated statistical methods, we found that the recorded population appeared to track each aspect of the task by entering distinct and separable network states that broke down on trials with numerous errors.

Results

Population Activity Organizes into Specific Patterns During Distinct Task Epochs. Multiple single-unit recordings from the rat ACC (dorsal medial agranular prefrontal cortex; see supporting information [SI] Fig. S1A) were obtained from 10 rats over a total of 27 trials of the delayed-win-shift radial-arm-maze task. The task consisted of the following behaviorally dissociable epochs illustrated in Fig. 1A: the periods surrounding correct arm choices during the training phase (TrC) and test phase (TsC), respectively; periods surrounding incorrect choices during the training phase (TsI; there were very few incorrect choices during the training phase); the period surrounding the point when the animal reached a food cup (TrR, during the training phase, and TsR, during the test phase); the entire delay phase (D); and all of the remaining periods intervening between arm choices and reward epochs during the training (Tr) and test phases (Ts).

Consistent with previous studies (24, 29–33), we observed a number of single-unit correlates that were associated with arm entries, reward processing, specific movements, and behavioral errors (Fig. 1E). However, the focus of the present study was the functional patterns of network activity associated with the distinct task epochs rather than single-unit correlates.

To analyze population activity, spike trains of all units were first convolved with Gaussian functions and binned at 200 ms, yielding smoothed instantaneous firing rates (iFR) for each unit as a function of time. For each time bin, the iFR of the N simultaneously recorded units were combined into population vectors embedded within an N-dimensional state space, here termed multiple-unit activity (MUA) space. For the purpose of visualization, we used multidimensional scaling to obtain 3D projections of these N-dimensional MUA spaces as shown in Fig. 1 B and C. In these graphs, each dot represents the entire state of the recorded network within one 200-ms bin of the task, and all population vectors (dots) belonging to different 200-ms bins

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of the same task epoch are shown in the same color. As Fig. 1B illustrates (see also Movies S1 and S2 and Fig. S2), different task epochs segregate in MUA space, i.e., MUA vectors belonging to the same task epoch tend to cluster within similar regions of MUA space, whereas MUA vectors belonging to different task epochs populate different regions, implying that the population as a whole differentiates between individual task epochs. Averaging across all points in MUA space belonging to the same task epoch yields an N-dimensional “prototype vector” representing the mean iFRs of all units within this epoch, illustrating the differential patterns of activation as a function of task epoch (Fig. 1D).

This organization into distinct patterns appeared to be functionally relevant to the successful completion of a trial. Fig. 1B illustrates population activity for an animal that performed perfectly on a trial. However, when the population activity for the same animal was reexamined during a trial in which six errors were committed, the segregation in MUA space broke down (Fig. 1C; see also Movies S1 and S2 and Fig. S2). This collapse of segregation in MUA space was also reflected in the lack of differentiation between task-epoch iFR prototypes (Fig. 1D). On this trial, only one unit showed clear differential activity, and it appeared to be most tightly linked to the signaling of behavioral errors (Fig. 1E). Therefore, successful task performance was characterized by the neural population attaining distinct task-epoch-specific patterns that changed dynamically with the cognitive requirements of the task. A failure to exhibit such distinct population patterns was correlated with errors in choosing the correct arms.

To quantify these phenomena statistically, activity patterns were analyzed across all animals used in this study by computing a separation error for each trial and every pair of task epochs. This was done by fitting a hyperplane (linear discriminant function) that optimally separated the clouds of points associated with each pair of task epochs using discriminant analysis and by determining for all of these pairs the relative number of population vectors (i.e., dots in MUA space) that were assigned to the wrong task epoch according to this linear classifier. Because the separability of task-epoch points is also affected by other factors such as the dimensionality of the MUA space and the total number of points/epoch, conservative surrogate data, obeying the same temporal continuity constraints as the original data, were constructed and used to test the significance of separation between any pair of task epochs (see Methods for details). Fig. 2A shows the separation errors for each of the original pairwise comparisons and for the respective surrogates constructed for each pair of task epochs averaged across all trials and animals. Each of the pairwise comparisons reached significance except for the comparison between test-phase incorrect (TsI) and correct (TsC) choices (paired t tests using corrected α*-levels according to the Holm–Bonferroni method, (34), α = 0.05; see SI Text and Fig. S3 for additional separation measures and

Fig. 1. Population analysis of ACC neuronal firing properties with respect to task epochs on a trial when an animal performed with no errors and a trial when the same animal made six errors. (A) Illustration of the delayed win-shift radial arm maze with the separate task epochs chosen. (B, C) MUA space representation. Each color-coded dot represents the state of the network (i.e., the iFR population vector of all recorded neurons) in a 200-ms time bin at some time point during the task. The blue and green dots are from 1-s periods centered around the points of arm entries (threshold crossings) for correct arm choices during the training phase (TrR) and test phase (TsR), respectively. The yellow dots are from 1-s time periods centered around the points of arm entries for incorrect choices during the test phase (TsI). The brown and red dots are from 1-s periods starting 200 ms before the points where the animal’s nose reached a food cup during the training phase (TrR) and test phase (TsR), respectively. The smaller light gray, dark gray, and black dots represent delay (Dl), training (Tr), and test phase (Ts) epochs, respectively, excluding those periods defined by the choice and reward events. The axes of this 3D projection correspond to the simultaneous activation and mutual suppression of several units. (D) iFR prototypes showing for each task epoch the average deviation of the iFR of each neuron from the grand average across all task epochs from the same trials as in B and C with no errors (D, Left) and six errors (D, Right). (E) White lines represent the Gaussian smoothed iFRs of 5 units from the 0-error trial (Upper) and 1 unit from the six-error trial (Lower) as the trial progressed from the training phase (dark gray) through the delay (light gray) to the test phase (black). Units were selected based on the prototypes in D. Blue and green vertical lines represent the points of arm entries, whereas brown and red lines indicate points of reward during the training and test phase, respectively.
analysis). Moreover, in >40% of all trials (averaged across all pairwise comparisons), a significant separation could be achieved on a single trial basis (with significant contributions coming from all animals and trials; Fig. S3A). Hence, the statistical analysis confirmed what visual inspection of MUA spaces in Fig. 1 suggested: in many more single trials than expected by chance (5% according to a $P < 0.05$ criterion), and overall on average, all pairs of task epochs significantly differed in terms of population behavior, except for one comparison involving incorrect choices (which is related to the general breakdown of MUA space separation especially at choice points in high-error trials as shown below).

**Individual Neurons Are Generally not Selective for Single-Task Epochs.**

As described above, ensemble activity significantly differentiated between almost all pairs of task epochs. To address the question to what extent this differentiation may be rooted in highly task-epoch selective responses of single ACC neurons, a separability index for each unit and task epoch was computed that compared the average activity of a unit during a given epoch with the average activity across all other epochs (see Methods for details), and the significance of this index was again tested using surrogates. A significant selectivity index of a unit for a particular epoch implies that its average activity during that epoch significantly deviated (positively or negatively) from the average activity across all remaining epochs. For each of the eight task epochs, between 13% and 27% of all units were found to be task-epoch-selective according to this definition (Fig. 2B). However, cells may significantly modulate their average iFR compared with the grand mean during more than one task epoch (e.g., units 5 and 9 in Fig. 1E). Indeed, $\approx$80% of task-epoch-selective units exhibited a significant modulation in at least one other epoch (Fig. 2C, gray bars). Conversely, this implies that <4% of all units recorded were highly selective for just one epoch. On the other hand, if any two of the epochs were compared with each other, 72% of the epoch-selective units were active in only one of the two instances and could therefore account for their separation (Fig. 2C, white bars). Thus, although each unit was generally modulated during more than one epoch, all pairs of task epochs differed in their constitution of selective units.

**Different Task Epochs are also Associated with Unique Coalitions Among Neurons.**

Our results thus far show that different cognitive phases of the task are associated with specific patterns of iFR activity across all recorded units. To examine whether these iFR patterns are also accompanied by epoch-specific correlations among units, the absolute (zero time lag), iFR Pearson correlation coefficients averaged across all pairs of units as a function of task epoch were computed (using 200-ms bins as before; Fig. 2D, dark bars). For each task epoch, except for the incorrect choices, iFR comodulations were significantly above chance ($t$ tests, all $P < 0.005$) based on comparisons with data sets where we shuffled the iFR bins (Fig. 2D, light gray bars). To assess whether these correlated activity changes were indeed task-epoch-specific or simply a general feature of the neuronal pairs recorded, we extracted the pairs with the 20% highest correlations among units, the absolute (zero time lag), iFR Pearson correlation coefficients averaged across all pairs of units as a function of task epoch with regard to the most highly correlated pairs. A given task epoch shared only $\approx$20–30% on average of its most highly correlated pairs with any other task epoch (Fig. 2E; Fig. S4), suggesting that coalitions among neurons formed and dispersed with each task epoch. As a further confirmation of this observation, we constructed surrogates by recombining iFR bins from different task epochs while maintaining the temporal relation among units (i.e., no shuffling of iFR bins). In all cases except for the TsR epochs ($t$ test, $P < 0.1$), these across-task-epoch iFR correlations were significantly lower than the within-task-epoch correlations ($t$ tests, all $P < 0.005$), even after correction by the shuffle predictor (Fig. S5). Hence, different task epochs are not only differentiated by unique patterns of changes in firing rate but also through task epoch specific coalitions among units.

**Ensemble Organization Is Diminished on Trials with Numerous Behavioral Errors.**

To address the functional importance of the distinct network patterns, changes in population activity as a function of behavioral errors were examined. For the entire dataset, trials were divided according to a median split based on the number of incorrect choices. The resulting groups agreed with previously defined criteria of asymptotic performance on this task (0–1 vs. 2 + incorrect choices; ref. 35). For the combined delay and test phases, separability was significantly worse for trials with many incorrect choices versus trials with 0–1 incorrect choices (Fig. 3A; $t$ test, $P < 0.01$). Furthermore, Fig. 3A shows that the breakdown in MUA space affected mostly comparisons involving test-phase choice and reward epochs (TsC and TsR), and less
Fig. 3. Breakdown of separability in MUA space and decrease of correlations are associated with a rise in behavioral errors. (A) Separability of task epochs in MUA space as given by the normalized (to the surrogates) classification errors for the low (0–1; blue bars) and high (≥2; red bars) behavioral error trials. The graph shows overall separability averaged across all pairs of delay and test phase comparisons and separately for all epoch-pair comparisons from the delay + test phases. Asterisks denote a significant (P < 0.05) difference in separability for high and low behavioral error trials. (B) Percentage of individual datasets for which a significant separation of task epochs was reached for the low (blue bars) and high (red bars) behavioral error groups, averaged across all delay + test phase comparisons, and separately for all delay + test phase comparisons. (C) Absolute iFR correlations averaged across all cell pairs in the low (blue bars) vs. high (red bars) behavioral error groups as a function of task epoch, corrected by subtracting the average iFR correlations obtained for the respective iFR-shuffled surrogates.

Fig. 4. Schema of a functional architecture that could explain the present results. Shown are three cell assemblies corresponding to correct choice (green), reward epochs (red), and basic test epochs (black). Neurons enclosed by the same colored outline are embedded within the same assembly, with neurons shared among assemblies being located within the intersections of the colored outlines. The firing rate of a neuron is indicated by its gray level. As the task progresses, first choice assemblies are activated (Left), followed by reward assemblies (Center), followed by a general test phase assembly (Right). Neurons 1 and 2 give an example of a pair that would be correlated during activation of the choice-assembly, as they form part of the same activated assembly but may be uncorrelated during the reward and test epochs. Neuron 3 is an example for a cell that is suppressed during correct choices due to mutual inhibition among assemblies (not illustrated for clarity), activated during the reward epoch, and fires at its average baseline activity in the test epoch. Neuron 2, in contrast, would be activated in both the choice and reward epochs. Recording from such a neuron would therefore allow one to differentiate between choice or reward epochs and the test phase, but not between choice and reward epochs. Hence, depending on a neuron’s participation in various assemblies, numerous patterns of differential activity may be observed for single units, whereas at the same time, the cell assembly gives rise to a unique pattern at the network level.

Discussion

The present study explored population coding within the ACC while animals foraged for food in an ecologically valid radial arm maze task involving distinct cognitive phases. Many single-unit correlates (e.g., of choices or rewards) were observed. In most cases, however, single units did not limit their firing rate changes to one specific event type but were active across multiple cognitively defined task epochs. On the other hand, each task epoch was characterized by a unique pattern of firing-rate changes across units and correlations among units. The behavioral importance of this functional organization was supported by the fact that these population patterns tended to break down on trials with numerous errors.

Cell Assemblies and Dynamic Population Patterns. The cell assembly framework first introduced by Hebb (1) could provide a mechanistic basis for the observed ensemble patterns. A cell assembly is defined by a functional group of neurons entertaining relatively strong recurrent excitatory connections among each other, while potentially inhibiting pyramidal cells of other assemblies. ACC networks may consist of various partially overlapping cell assemblies encoding various cognitive events within the decision making process as illustrated in Fig. 4. As a result of such functional arrangements, at the single-neuron level, there would be numerous combinations of enhanced and depressed activity changes across different cognitive events, whereas each assembly as a whole would be associated with a unique pattern of firing rate changes and a unique pattern of within-assembly correlations (due to the recurrent excitatory connections within but not across assemblies), as observed here (Figs. 1 and 2).

Many other approaches, both similar to and different from ours, have been used to identify ensemble organization in multiple single-unit data. These include Hidden Markov Models, which also revealed distinct patterns and transitions among them in vivo (14, 36), and principal component or cluster analysis to unravel functional groupings (37), hierarchical organization of task-coding assemblies (7), or simply as a means for visualizing distinct population states and their connecting trajectories (15). At the level of precise spiking times, task-phase-specific alliances among neurons were demonstrated in primate prefrontal areas using spike train cross-correlograms (13) and repeating patterns of spikes aligned among multiple neurons (6, 38) were also taken as evidence for ensemble organization. Hence, a number of...
studies employing different analysis methodologies provide converging evidence for the organization of neurons into functional ensembles at different temporal scales.

**ACC and Functional Assemblies in Behavioral Monitoring.** Transient lesions of the rat ACC produced significant impairments on the same task used here (35), causing numerous revisits to previously baited arms indicative of an inability to update responding as reward contingencies changed. Consistent with the putative function of the ACC in monitoring and updating response policies (39), single-unit activity in this area is most often related to specific movements, movement sequences, or response choices, as well as to rewards, specific action-reward pairs, or to the detection of response errors (21–27, 29–33). One important aspect of these single-unit correlates is their highly dynamical nature, with neuronal firing rates changing as a function of task context and reward magnitude even for the very same movements (24, 32, 40), or vice versa, with the same neurons coding for very different movements in different tasks (29, 40). This high degree of flexibility and context dependence supports the view of the ACC as a device for monitoring or attending to actions (16, 17).

Within this framework, the functional organization into assemblies and the transitions among them may help to bind different cognitive attributes of a response strategy together, and to connect actions to subsequent outcomes (39). Performing a complex or novel task requires an animal to integrate various sensory, motor, reward, and memory aspects, and the transient formation of dynamic assemblies in the ACC as observed here may reflect this process. In this sense, the assemblies monitor the animal’s progression in a task-dependent frame of reference or “task space.” The sequential transitions among ACC assemblies and the transecho activity of some of the units recorded may in addition help to connect the temporally separated components of action-outcome chains. Sequential transitions among active populations as well as persistent activity of single frontal neurons have been proposed previously as a means to link events through different action-outcome chains. Sequential transitions among active populations as well as persistent activity of single frontal neurons have been proposed previously as a means to link events through different action-outcome sequences (39).

**Methods**

**Data Acquisition.** Male Long-Evans rats (225–250 g, Harlan) were deeply anesthetized with 100 mg/kg ketamine and 10 mg/kg xylazine and placed into a stereotaxic device where multielectrode arrays were inserted through a cranial hole centered at A/P 2.2; M/L 0.8; D/V 2.5 relative to bregma and offset 1 mm. The arrays were inserted at the stereotaxic coordinates and analyzed and separately for all task epochs. To test significance nonparametrically, 99 surrogates were constructed for each comparison across-task-epoch correlations, equal time slices were first drawn from all task epochs by determining an optimally separating (N–1)-dimensional hyperplane via discriminant analysis (e.g., ref. 44). The relative number of misclassified points was taken as an index of separability and was compared to surrogates constructed in the following way: Pairs of task epochs were first combined, and from these unions of points k contingent segments were randomly drawn, where k is the number of contingent segments for the original task epoch. To determine significance for each dataset individually, a nonparametric test was applied: significance at the 5% level was assigned by establishing whether the original classification error was among the 5% lowest within the set of 1 original and 99 surrogate classification errors.

To examine percentages of epoch-selective cells, for each cell n and task epoch p, a selectivity index was computed as

\[
s_n(p) = \frac{|\langle \text{IFR}_n(t) \rangle_p - \langle \text{IFR}_n(t) \rangle_q|}{\langle \text{IFR}_n(t) \rangle_p + \langle \text{IFR}_n(t) \rangle_q},
\]

where \( \langle \cdot \rangle \) denotes the average across segments of epoch p, or across all of the combined other epochs q, respectively. Significance of each index \( s_n(p) \) at the 5% level was again established by a nonparametric comparison with 99 surrogates, where in this case k contingent segments, with k being the number of segments making up epoch p, were drawn at random from the combination of all epochs in the task.

For the correlation analysis, absolute standard zero-lag Pearson correlation coefficients \( r_p(n,m) \) were computed among scalar times series \( \text{IFR}_n(t) \) and \( \text{IFR}_m(t) \) for all pairs of simultaneously recorded neurons \( n,m \in \{1..N\}, n \neq m \), and separately for all task epochs. To test significance nonparametrically, 99 surrogates were constructed for each \( r_p(n,m) \) by randomly shuffling the bins within series \( \text{IFR}_n(t) \) and \( \text{IFR}_m(t) \). For comparison of within-task-epoch to across-task-epoch correlations, equal time slices were first drawn from all task epochs to ensure that the surrogates drew from every task epoch with equal likelihood. Surrogates were then constructed by drawing for each comparison at random 20 bins from the union of time-equaled task epochs, original correlations were also recomputed for the time-equaled task epochs, and both original and across-epoch surrogate correlations were corrected by the mean from 99 shuffles.

For the comparison of the two behavioral error groups, all separation errors for each dataset were first normalized to the average separation error within the respective set of surrogates. However, the two behavioral error groups based on the median split did not differ with regard to either the average number of recorded neurons (ME = 13.8 for the low and ME = 14.2 for the high error group, two-sided t test: \( P > 0.86 \)) or the standard deviation of this number (SD = 5.2 for the low and SD = 5.8 for the high error group), i.e., the dimensionality of the MUA spaces for those two groups was the same.

Further details of methods and methodological considerations can be found in SI Text.

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