Hippocampal and Neocortical Interactions During Context Discrimination: Electrophysiological Evidence From the Rat

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ABSTRACT: There is substantial evidence that hippocampus plays an important role in the processing of contextual information. Its specific role, however, remains unclear. One possibility is that single hippocampal neurons represent context information so that local circuits can construct representations of the current context, and the context that is expected based on past experience. Population codes derived from input by multiple local circuits may then engage match–mismatch algorithms that compare current and expected context information to determine the extent to which an expected context has changed. The results of such match–mismatch comparisons can be used to discriminate contexts. When context changes are detected, efferent messages may be passed on to connected neocortical areas so that informed “decisions” regarding future behavioral and cognitive strategies can be made. Here, a brief review describes evidence that a primary consequence of hippocampal processing is the discrimination of meaningful contexts. Then, the functional significance of neocortical circuits that likely receive hippocampal output messages are described in terms of their contribution to the control of ongoing behavioral and cognitive strategy, especially during active navigation. It is clear from this systems view that studies of spatial navigation continue to provide researchers with an excellent model of hippocampal–neocortical interactions during learning.

KEY WORDS: context; place cells; spatial processing; temporal processing; behavioral intention

INTRODUCTION

In recent years, an increasingly impressive array of methodologies and perspectives has allowed us to gain new, and sometimes surprising, insight into the dynamic (intrinsic) neural operations and behavioral consequences of hippocampal function. Converging evidence has led to the general conclusion that hippocampus is essential for context processing, especially as it relates to episodic memory function (e.g. O’Keefe and Nadel, 1978; Nadel and Wilner, 1980; Nadel and Payne, 2002; Tulving, 2002). It has become equally clear that fully adaptive context analysis and learning requires not only hippocampus, but the larger neural system within which hippocampus operates. In what follows, it is argued that the specific contribution that hippocampus makes to context processing is to detect changes in context so that connected areas (especially neocortex) can make informed “decisions” in terms of future behavioral and cognitive strategy. First, a brief review is provided concerning the role for hippocampus in context discrimination. Then, the function significance of neocortical circuits that likely receive hippocampal output messages are described in terms of their contribution to the control of ongoing behavior and cognitive strategy.

HIPPOCAMPUS IS ESSENTIAL FOR CONTEXT PROCESSING

There is now strong evidence that hippocampus processes contextual information (e.g. Hirsh, 1974; Myers and Gluck, 1994; Anagnostaras et al., 2001; Maren, 2001; Fanselow and Poulos, 2005; Bouton et al., 2006). Conditioned fear responses to contextual stimuli is eliminated with hippocampal lesions even though responses to discrete conditional stimuli remain intact (Kim and Fanselow, 1992; Phillips and LeDoux, 1992, 1994). Also, hippocampal or entorhinal cortical damage produces an insensitivity to changes in context as evidenced by the fact that lesioned animals do not show the normal decrement in conditioned responding when the context is altered (Penick and Solomon, 1991; Freeman et al., 1996a,b). Manipulations that impact hippocampal synaptic plasticity (e.g. LTP) also affect context learning (e.g. Shors and Matzel, 1997). These results not only support the hypothesis that hippocampus processes context information, but more specifically, they indicate that hippocampus plays a particularly salient role in the discrimination of meaningful contexts (Smith and Mizumori, 2006a).

A context processing account of hippocampus is consistent with hypotheses that hippocampus plays a special role in the flexible use of conjunctive, sequential, relational, and spatial information (e.g. O’Keefe and Nadel, 1978; Foster et al., 1987; Eichenbaum et al., 1999; Wood et al., 2000; Eichenbaum and Cohen, 2001; O’Reilly and Rudy, 2001; Fortin et al., 2002). These flexible, spatial, sequential, and relational operations likely enable hippocampus to make accurate context discriminations. While significant evidence supporting the existence of these operations in hippocampus can be derived from single unit recordings in behaving rats, it remains a significant challenge to understand how context processing theories of hippocampus account for the varied hippocampal place cell.
responses that have been reported. The following description of a context discrimination hypothesis (CDH; Smith and Mizumori, 2006a; Mizumori, 2007) provides a theoretical framework to account for a significant amount of the place cell findings, as well as to provide a link to current ideas about the specific role of the hippocampus in learning and episodic memory (Tulving, 2002). While many features of this hypothesis build on concepts discussed by other investigators (as pointed out below), its consideration here provides an opportunity to elaborate specifically on the functional networks that should exist as part of hippocampal neural organization. Furthermore, the CDH makes clear predictions about how hippocampal efferent messages ultimately come to impact ongoing behavior.

Consistent with a large body of electrophysiological evidence, CDH postulates that single hippocampal neuronal representations of context provide data to population-based network computations that ultimately determine whether expected contextual features of a situation have changed (e.g. Mizumori et al., 1999b, 2000, in press; Smith and Mizumori, 2006a,b; Mizumori, 2007). That is, hippocampal representations of multiple aspects of the current context (Fig. 1; O’Keefe and Nadel, 1978; Nadel and Wilner, 1980; Nadel and Payne, 2002) may contribute to a match–mismatch type of analysis that evaluates the present context according to how similar it is to the context that an animal is expecting based on past experience (e.g. Gray, 1982; Vinogradov, 1995; Mizumori et al., 1999b, 2000; Gray, 2000; Lisman and Otmakhova, 2001; Anderson and Jeffery, 2003; Hasselmo et al., 2002; Jeffery et al., 2004; Hasselmo, 2005; Manns et al., 2007). Detected mismatches can be used to identify novel situations and to distinguish different contexts, functions that are necessary to define significant events or episodes. When a match is computed, the effect of hippocampal output could be to strengthen currently active memory networks located elsewhere in the brain (e.g. neocortex). In this way, hippocampus potentially plays different mnemonic roles depending on whether or not contexts actually change.

The detection of changes in context is fundamentally important for accurate performance in a variety of learning tasks (e.g. navigation-based learning, instrumental conditioning, classical conditioning). This function is critical because of the need to engage cellular mechanisms for new learning at potentially important times (Paulsen and Moser, 1998). Indeed, Smith and Mizumori (2006b) showed that hippocampal neurons develop context-specific responses, but only when rats were required to discriminate contexts. Discriminating neural responses were not observed when rats were allowed to randomly forage for the same amount of time. Most recently, Manns et al. (2007) showed that relative to match trials in an odor cue or object recognition task, CA1 neurons preferentially discharged when animals experienced a nonmatch situation in these same tasks. Moreover, cell firing tended to occur during the “encoding phase” of the ongoing theta rhythm. This interpretation is based on the hypothesis that encoding and retrieval alternates at a frequency that coincides with the theta frequency (see review in Hasselmo, 2005). Thus, detection of a nonmatch situation can change the relationship between cell discharge and the local theta rhythm such that encoding functions

FIGURE 1. Color density plots of spatially localized firing show context-dependent changes in place fields (Panels A and B) and egocentric movement correlates of interneurons (Panels C and D). Panel A shows place field reorganization when the room light conditions change as rats perform the same spatial memory task during both dark and light phases of testing on a radial maze. Such responses are almost always observed when the recording session begins in darkness. Presumably this place field reorganization occurs because the appropriate memory has not yet been recalled. When the lights are turned on, a different (learned) representation appears, and these have been shown to remain stable during subsequent dark periods (e.g. Mizumori et al., 1999b). Panel B demonstrates that the exhibition of a given place field may be conditional depending upon the recent behavioral history of the animal. In this case, place fields were recorded as rats started a plus maze task from either north (Start 1) or south (Start 2) maze arms to obtain food located on the east (right) maze arm. Within session, data were divided according to the start arm location. It was found that the place field was observed only when the rat began a trial from the south location. Importantly, this was the case even though the rat exhibited the same behaviors, and experienced the same external sensory environment as it traversed the place field location. (Panels C and D) Hippocampal interneurons often vary firing rates as a function of the velocity or acceleration of translational movements through extended space (e.g. McNaughton et al., 1983). Panel C (left) illustrates such movement-related firing (open circles) when a rat performed a plus maze task similar to the one described in Panel B. When the reward location changed from the east to west maze arm, it can be seen that the acceleration-correlated firing was sharply attenuated (closed circles) even though the rat continued to engage in the same type of behavior as before the reward location shift (Smith and Mizumori, 2006). Panel D illustrates the effects of changes in cognitive strategy (from spatial to response strategies) on acceleration correlates of a hippocampal interneuron. This cell initially did not show significant acceleration correlated firing. However, after the strategy switch, an acceleration correlate emerged (Eschenko and Mizumori, 2007). Importantly, such changes in movement-correlated firing were observed for many cells even though the sensory and behavioral responses of the animal did not change. Panels C and D, then, illustrate that interneurons represent context-dependent movement information. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
In further support of the CDH, disconnecting hippocampus by fornix lesions impairs context discrimination (Smith et al., 2004), and hippocampal lesions reduce animals' ability to respond to changes in a familiar environment (Good and Honey, 1991; Save et al., 1992a,b). Spatial novelty detection corresponds to selective elevation of the immediate early gene c-fos in hippocampus, and not in surrounding parahippocampal cortical regions (Jenkins et al., 2004). Also, hippocampal neurons show significantly altered firing patterns when rats experience spatial or nonspatial changes in a familiar environment (O'Keefe, 1976; Muller and Kubie, 1987; Wood et al., 1999; Fyhn et al., 2002; Ferbinteanu and Shapiro, 2003; Moita et al., 2004; Yeshenko et al., 2004; Leutgeb JK et al., 2005; Leutgeb S et al., 2005; Puryear et al., 2006; Smith and Mizumori, 2006b; Eschenko and Mizumori, 2007).

The specific patterns of neural change after altering different features of an experimental context tend to be multidimensional and complex (see Special issue on Place Fields and Episodic Memory, vol. 16(9), *Hippocampus*). After considering the diversity of neural responses and test conditions that have been reported, it appears that there are (at least) three qualitatively distinct types of influences on hippocampal neural codes that likely occur automatically regardless of the specific task demands: memory (i.e. past experience), spatial, and temporal (Fig. 2). This proposal is based in part on past suggestions that hippocampal-dependent memory involves associations of spatial, and temporal features (e.g. O'Keefe and Nadel, 1978; Mizumori et al., 2000; Redish et al., 2000; Burgess et al., 2001; Eichenbaum and Cohen, 2001; Morris, 2001; O'Reilly and Rudy, 2001; Buzsáki, 2005). At slight variance with many of these proposals, Mizumori (2007) suggest that the default mode of hippocampal processing is to continually integrate memory-guided perceptions of sensory, movement, and motivational information, or memory (M), within a spatial (S) reference framework as a function of time (T; Fig. 2). An important point here is that hippocampus may receive highly preprocessed M information so that it can be incorporated into specific spatial and temporal frameworks. The integration is proposed to occur according to a hierarchically organized scheme that incorporates M, S, and T information as fundamental inputs. As a result, during unrestrained navigation, place fields appear as neural representations of...
A hierarchical representation of context processing within hippocampus. Each square corresponds to a context matrix that reflects multiple types of information coded by single cells (Level 1; light blue), local ensembles (Level 2; tan), or large regions of hippocampus (Level 3; pink). The width of the lines reflects the relative strengths of the different input. Subscripts correspond to the M, S, or T input for a given cell number. Place cell studies reveal that a number of specific features (e.g., sensory, behavioral, motivation, and knowledge of task rules; see Fig. 2) are represented in hippocampal networks according to past experience or memory (M). The M input to place cells may vary in strength depending on the learned significance of the information that M represents. As a result, some M inputs may disproportionately present current context information to hippocampal networks, while other M inputs may be influenced more by expectations for a given context, such as memory for task rules. Another factor that continually shapes the organization of information in hippocampus is the spatial reference framework (S) provided by entorhinal cortex grid cells (described in text). A third continual influence on hippocampal place fields is one that strives to organize incoming information as a function of time (T). Level 2 neural integration reflects ensemble activity of local networks of cells. Presumably, this level integrates representations by single place cells to define portions of the expected and current context within local ensembles. In this example, Cells 1 and 3 combine to define a part of what will become a component of the definition of the expected features of a context. Cell 3 has stronger influence than Cell 2, perhaps reflecting a stronger history of synaptic activation. Cells 2 and 4 similarly combine to define another portion of the expected context. Cells 5 and 7 combine to define a segment of the current context. The greater influence at Level 2 is depicted by the taller cylinder. Finally, Cells 6 and 8 combine to form another local ensemble that processes current context information. Cell 6 has greater influence on Level 2 integration, and this is shown by the taller cylinder in Local circuit 4. Presumably there are many such local circuits throughout hippocampus, and many more than two individual place cells contribute to each local ensemble. Also, due to the known massive interconnection between cells in a specific hippocampal subregion, it is likely that the activity of one ensemble impacts the activity of connected ensembles (not shown). Level 3 integration compares expected and current context ensemble representations to produce an efferent code that indicates the extent to which the expected context has changed. If the expected and current contexts differ, as in the present case, a signal identifying the nonmatch will be forwarded to cortical systems to notify them of the change. This notification could have the effect of altering cell excitability patterns so that the nature of the change can be evaluated. An example of one such consequence could be an increase in exploratory behaviors. If the ensemble codes for the expected and current context are the same, the expected and current context matrices should overlap. In this case, the output message of hippocampus may signal the continuation of the ongoing behavior, and it may strengthen synaptic connections within the neural network that defines the current activated memory. The result of the pattern of activity shown in this figure exemplifies an output message during a nonmatch condition. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
different types of sensory, behavioral, and intrinsic information that have strong spatial and temporal features, thereby validating the term *spatial context* when referring to the meaning of place field representation (Nadel et al., 1985; Mizumori et al., 1999b, 2000; Jeffery et al., 2004). CDH, then, suggests a more complex definition of context than is typically used in the learning literature—a definition that emphasizes the integration of sensory, motivational, response, and mnemonic input within a spatial framework as a function of time. That is, hippocampal context codes are not just a reflection of the external sensory environment.

**HIERARCHICALLY ORGANIZED CONTEXT PROCESSING WITHIN HIPPOCAMPUS**

Previous hierarchical models of hippocampal processing consider the fundamental elements of the hierarchy to be either stimulus-defining information (e.g. Shapiro et al., 1997) or individual events (e.g. Eichenbaum et al., 1999; Shapiro and Eichenbaum, 1999; Wood et al., 1999). Figure 3 illustrates an alternative hierarchical model that includes as inputs S (spatial) information as a form of organization rather than specific environmental features that individual cells discriminate. Also, M (memory) input from extrahippocampal systems brings preselected task-relevant sensory, behavioral, motivational, and learned units of information into hippocampus. Presumably, such M input is derived from neocortical memory circuits. Finally, it is argued that the T (temporal) organization of information may not necessarily represent only the end goal of hippocampal processing. Rather it may also emerge from intrinsic synaptic plasticity mechanisms that can be guided by extrahippocampal temporal events (e.g. Buzsaki, 2005). These synaptic mechanisms that temporally organize salient information allow hippocampus to ultimately define expected and current contextual features in terms of their duration or sequence. Direct connections to CA1 from entorhinal cortex may provide M inputs to hippocampus that aid in the segregation of sequences into meaningful epochs at multiple levels. It should be pointed out that M, S, and T information may not be mutually exclusive. Rather, there may be moderate to considerable interaction amongst these variables prior to and within hippocampus (Fig. 4). As an example, the reliability or specificity of a particular place field could be based on M inputs. Indeed it has been shown that such measures of place fields vary with experience (e.g. Mizumori and Kalyani, 1997; Puryear et al., 2006). Also, the degree of temporal coherence amongst a group of neurons could depend on their recent activation patterns (e.g. Shapiro and Ferbinteanu, 2006). As a third example, spatial features could conceivably determine the degree of temporally correlated firing one could expect to observe between especially place cell pairs.

Consideration of the details and significance of place field representations within a hierarchically organized scheme suggests three basic levels of integration (Fig. 3; Mizumori, 2007). The basic features of each level are summarized in Table 1. Essentially, Level 1 allows for the initial formulation of the spatial-temporal context features that are typically observed for place fields. Level 2 integrates single neuron context representations into local circuit signals that reflect current context information or expected context information in experience-dependent ways. Local circuit neural codes could be reflected in the firing of one cell relative to the firing of another cell within the same circuit. Moreover, given that Level 2 integrates input from multiple individual neurons, correlated firing between cells may be found to preferentially occur relative to general task events such as the beginning and end of trials, as has been shown by Shapiro and Ferbinteanu (2006). Level 3 integration is considered to progressively carry out the context comparison function amongst an increasingly greater number of local circuit arrays, perhaps according to mechanisms such as pattern completion and pattern separation.

In contrast to M and S information (discussed in more detail in the following section), T information seems to be driven largely (although not exclusively) by circuitry intrinsic to hippocampus. This has been shown primarily from studies of how place cell firing is temporally regulated. This temporal organization may take different forms, perhaps for different purposes. On a single pass through a field, the first spike of successive bursts occurs at progressively earlier phases of the theta cycle (O’Keefe and Recce, 1993). It has been suggested that such temporally dynamic changes in spike-timing (called phase precession) may be a key mechanism by which place fields provide a link between temporally-extended behaviors of an animal and the comparatively rapid synaptic plasticity mechanisms (e.g. LTP) that are thought to subserve learning (e.g. Skaggs et al., 1996). Theoretical considerations have generated detailed models of how phase precession could explain the link between...
behavior and neural plasticity mechanisms (e.g. Buzsáki, 2005; Zugaro et al., 2005).

Another form of temporal-based neuroplasticity within hippocampus involves a change in the timing of spike discharge by one cell relative to spiking of other cells, or relative to prior spike discharges by the same cell. For example, it has been shown that the temporal coherence of place cell discharge is greater in mice with an intact hippocampus compared to mice with deficient NMDA systems (McHugh et al., 1996). Greater synchronization could offer a stronger output signal to efferent structures. Relatively, experience-dependent temporal codes may be found in terms of the temporal relationships between the firing of cells with adjacent place fields. With continued exposure to a new environment, place fields begin to expand asymmetrically such that the peak firing rate occurs with shorter latency upon entrance into the field (Mehta et al., 1997, 2000). The asymmetric backward expansion of place fields is thought to provide a neural mechanism for learning directional sequences. Moreover, it has been suggested that the backward expansion phenomenon may contribute to the transformation of a rate code to a temporal code such as that illustrated in phase precession (Mehta et al., 2002). Perhaps, the backward expansion phenomenon could help to explain other place field phenomenon such as the tendency for place cells to fire in anticipation of entering a specified location within a familiar environment (Muller and Kubie, 1989).

While the dynamic changes in place field shape are intriguing, it remains a challenge to determine whether, and then how, changes in the temporal distribution of cell firing are directly related to spatial learning, and how these alterations are reflected in local circuit codes (Level 2). However, Shapiro and Ferbinteanu (2006) recently reported that the temporal relationship between firing of simultaneously recorded place cells discriminated task phase, suggesting that past experience determines relative spike timing of individual neurons. Experience-dependent changes in spike timing are evident not only when considering cell discharge of one cell relative to another, or to itself, but also timing relative to task-specific events. Lee et al. (2006) described place fields whose center of mass moved in the forward direction (toward a goal) with each successive trial. The direction of place field movement was in the direction opposite to that predicted by spike timing dependent plasticity mechanisms. However, it may be that place fields will move in the forward or backward directions for different purposes, and their relative direction of movement may depend on the task conditions. For example, forward movement of place fields may be more prevalent when choices must be made between multiple goal locations. Backward movement may be more prominent when merely knowing the direction of movement is sufficient to predict future reinforcement. In these cases, a distinguishing feature would not only be the direction of place field movement, but also the temporal resolution of the movement itself. Indeed, forward movement of place fields seemed to occur over a protracted time scale relative to the backward expansion effect. If the above analysis is correct, then a critical issue is to determine the mechanism that regulates place field shifts in the forward and backward directions, and whether these are independent mechanisms. It is also possible that the more protracted forward movement of place fields reflects changes in the spatial code instead of, or in addition to, a temporal shift.

Ferbinteanu and Shapiro (2003) described place fields that were consistently located on the goal arm of a plus maze regardless of the start location. These cells were considered to represent prospective information about the upcoming reward event. An
Entorhinal Cortex-Hippocampal Interactions

The notion of exploration-induced spatial organization of hippocampal representations is consistent with the finding that hippocampal place fields are observed upon first exposure to a new environment (Muller and Kubie, 1987; Wilson and McNaughton, 1993; Markus et al., 1995; O’Keefe and Burgess, 1996a,b; Hetherington and Shapiro, 1997; Frank et al., 2004). Also, compared to passive movement conditions in which rats are made to go through a place field either by being held by the experimenter or by being placed on a moveable robotic device, active and unrestrained movement seems to generate more selective and reliable place fields (Foster et al., 1989; Gavrilov et al., 1998; Song et al., 2005). Furthermore, pyramidal cells fire more robustly when rats run faster across a given location (i.e. running in a running wheel; Czurko et al., 1999). Voluntary locomotion, and the subsequent imposition of a spatial organization to contextual information, appears to have the effect of sharpening the neural image of spatial representations, at least in dorsal hippocampus. A sharper code should relay more specific spatial information to intrinsic and extrinsic computational circuits. The finding that unrestrained movement produced sharper codes than passive movement suggests that learned behavioral responses have more meaning for, and impact on, place fields than random movement of the animal. This may be one way in which learned information helps to define a context code.

The tendency for voluntary navigation to impose a spatial organization of contextual information may also explain why place fields have not been seen as the predominant form of coding in the primate hippocampus. Monkey hippocampal neurons respond primarily when the subject directs its gaze at a particular part of the environment (Rolls, 1999), whereas rodents explore the environment by active locomotion, primes accomplish much of their exploration visually, by directing their gaze about the environment. Using virtual navigation methods, location-selective firing has not only been reported, but it seems selective to contexts in which subjects must search for and identify meaningful locations (Ekstrom et al., 2003). Thus, the apparently discrepant findings between primate, human, and rodent studies may not indicate fundamentally different computations by the hippocampus, but rather they may result from different combinations of information passed on to hippocampus, information that is dictated by the task conditions.

Interestingly, a common feature of the neural organization of most sensory systems is that there is a clear spatial organization (or topography) to neural representations, starting from the sensory receptor to sensory cortex. This form of organization may facilitate adaptive responding since motor output systems (from motor cortical areas through spinal cord) are also organized topographically. Assuming that the use of a spatial reference frame to organize sensory and response information is highly efficient, it seems reasonable to speculate that there was strong evolutionary pressure to process and retain sensory information spatially as sensory association (cortical) regions evolved. Maintaining a spatial reference framework for fundamentally important tasks such as accurate navigation may have facilitated an organism’s ability to rapidly adapt to environmental changes. For this reason, phylogenetically old structures such as hippocampus may be initially predisposed to process information within a spatial framework, regardless of the

**HIPPOCAMPAL AND NEOCORTICAL INTERACTIONS DURING CONTEXT DISCRIMINATION**

**CONTEXT DISCRIMINATION BY HIPPOCAMPUSS IS GUIDED BY, AND ALSO INFORMS, A BROADER CONTEXT PROCESSING SYSTEM**

Information about the stability or changes in context is likely to be useful to insure adaptive behavioral response selection and execution, and to update currently active memory or other intrinsic processing systems. Thus, it should be expected that hippocampal context discrimination functions operate continuously and automatically almost regardless of the task at hand. This is not to say, however, that task demands don’t regulate the efficiency or perhaps even the type of hippocampal context information being compared. In fact, the contrary may be true.

**Entorhinal Cortex-Hippocampal Interactions**

The preponderance of striking place fields regardless of the cognitive strategy required by navigating rats (e.g. Eschenko and Mizumori, 2007) suggests that hippocampus is somehow predisposed to organize contextual information spatially. Place field experiments invariably involve exploration of the environment. Such voluntary movement may direct spatial and movement signals through the medial entorhinal cortex such that networks of spatial grid cells become activated (Haffing et al., 2005; Leutgeb S et al., 2005; Sergolini et al., 2006). Grid cells selectively discharge when rats traverse positions that coincide with the vertices of a regularly tessellating grid. The grid fields appear to represent conjunctions of location, direction, and movement information within a two-dimensional coordinate representation of the environment. Thus, the entorhinal cortex may pass on to hippocampus a spatial framework (McNaughton et al., 2006) within which context-specific M and T information is organized and processed. For this reason, the term spatial context is often preferred when discussing context processing during navigation (Nadel et al., 1985; Mizumori et al., 1999b, 2000; Jeffery et al., 2004).

The time scales for the latter two studies are very different so it remains to be determined if they are derived from the same or different mechanism. One may also consider the backward expansion effect (described above) a form of anticipatory neural code due to M input. It is possible, then, that at least a subpopulation of place cells conveys information about what to expect in the future within the milliseconds to seconds range. Place cells may also code information retrospectively. Place fields have been found at consistent distances from a start location on linear tracks and plus mazes (e.g. Gothard et al., 1996; Ferbinteanu and Shapiro, 2003). In summary, these findings are consistent with the view that changes in spike timing may allow hippocampal cells to encode perhaps simultaneously information about what to expect in a given context (prospective code), as well as current or recent (retrospective) events.

earlier example of prospective coding by place cells was the demonstration that place fields seemed to anticipate the future location of an animal by some 90 ms (Muller and Kubie, 1989). The time scales for the latter two studies are very different so it remains to be determined if they are derived from the same or different mechanism. One may also consider the backward expansion effect (described above) a form of anticipatory neural code due to M input. It is possible, then, that at least a subpopulation of place cells conveys information about what to expect in the future within the milliseconds to seconds range. Place cells may also code information retrospectively. Place fields have been found at consistent distances from a start location on linear tracks and plus mazes (e.g. Gothard et al., 1996; Ferbinteanu and Shapiro, 2003). In summary, these findings are consistent with the view that changes in spike timing may allow hippocampal cells to encode perhaps simultaneously information about what to expect in a given context (prospective code), as well as current or recent (retrospective) events.
task. Indeed, hippocampal place fields are observed with similar abundance in spatial and nonspatial tasks (Yeshenko et al., 2004). The fact that it is possible to break out of the spatial framework if needed may reflect a more recently evolved adaptation.

Temporal organization of spatially organized hippocampal neural codes appears to be reflected in neural signaling by neocortical cells. It was shown that the temporal sequence of neural firing recorded during exploration is essentially “replayed” during slow wave sleep but in a temporally condensed fashion (Wilson and McNaughton, 1994; Hoffman and McNaughton, 2002; Pennartz et al., 2002). Recently, another form of replay of neural activation was reported when rats paused between traversals along an elevated track (Foster and Wilson, 2006). This differed from the replay observed during sleep because the temporal order of neural activation was reversed relative to the order in which locations were visited during locomotion. This reverse replay was interpreted as being useful for a different mnemonic function, one that allows the evaluation of recent behaviors (trajectories in this case) in terms of their reinforcement outcomes. This particular form of temporal organization may be generated within hippocampus, reflecting a retrospective mechanism that is needed to detect changes in the expected situation.

The different forms of temporal coordination can be used to predict behaviors and events in a prospective manner, or to recall recent behaviors or events in a retrospective manner. Thus, T-information can be represented in the multidimensional context matrix (Fig. 2) according to the degree to which the neural code is prospective or retrospective. Cell-to-cell spike timing changes likely emerge from a number of natural synaptic sources that regulate coordinated spike activity within hippocampus. More broadly speaking, rhythmic activity may be regulated by plastic changes within intrahippocampal circuitry and by extrinsic systems responsible for generating rhythmic activity in hippocampus (e.g. brainstem: Buzsáki, 2002, 2005; Vertes et al., 2004; Hasselmo, 2005). The varied sources of temporal regulation may allow for coding such M information as stimulus sequences (Dragoi and Buzsáki, 2006), or for determining stimulus duration.

**Retrospenial (Parietal) Cortex-Hippocampal Interactions**

The parietal cortex is traditionally considered in terms of space-based attention to the external environment as well as attention to oneself (Colby and Goldberg, 1999). Movement intention is another commonly considered function of parietal cortex. Recent findings, however, indicate that at least the retrosplenial sector of parietal cortex may serve more complex cognitive roles. For example, retrosplenial cortex is one of the many posterior parietal structures that become active during correct responses in item recognition tasks (Konishi et al., 2000), suggesting a critical role in memory retrieval (Wagner et al., 2005). In the case of spatial navigation by rats, the retrosplenial cortex has been discussed as a key player in updating visuospatial representations with movement-related (idiothetic) information (Sutherland et al., 1988; Cooper and Mizumori, 2001; Harker and Whishaw, 2004). Anatomical connections of the retrosplenial cortex support such an integrative role between visual and movement-related information since it receives input from, and returns projections to, visual association cortex, medial precentral cortex, motor cortex, anterior cingulate cortex, and the subicular complex (Vogt and Miller, 1983; Wyss and van Groen, 1992; see Fig. 5). Temporary inactivation of the retrosplenial cortex impairs spatial performance on a radial maze when darkness was imposed on a familiar test environment, but not when the same environment was lit (Cooper and Mizumori, 2001). The same inactivation procedure impaired initial learning in a well lit environment. Thus, it was concluded that the retrosplenial cortex contributes to spatial navigation when learned associations between visual and idiothetic information are required to perform the task (as in the case of well trained rats performing without familiar visual cues), or when new associations are being formed between visual and idiothetic information (as in the case of new learning in a lit environment). That retrosplenial information importantly informs hippocampal processing was shown by the finding that retrosplenial cortex inactivation altered hippocampal place fields that were recorded during the same experiment. It appears then that retrosplenial cortex may serve as perhaps one of many portals through which hippocampal information interacts with stored memory representations: Retrosplenial cortex may relay to hippocampus context-specific integrated sensory and movement information. Hippocampus in turn may effectively update the memory representations by providing retrosplenial cortex with feedback about the reliability of the context-based input.

To test the view that retrosplenial cortex provides integrated learned context information to hippocampus in the service of context discrimination, Smith et al. (2003) recorded retrosplenial cortex neurons as rats learned to discriminate behavioral contexts that differed only in terms of the location of food reward. That is, sensory and behavioral aspects were the same for the two contexts. It was found that reward-related retrosplenial cortex neurons developed task-relevant responses when animals learned the context discrimination (see Fig. 5 for examples), but not when rats foraged for randomly located food on the same maze. This result is entirely consistent with Smith et al. (2004) who showed
FIGURE 6. Event histograms illustrating context-specific responding by reward sensitive retrosplenial cortex neurons. Details of the training can be found in the text. Briefly, rats were trained to discriminate contexts according to where (on a plus maze) the animal expected to find reward. Top row: This cell showed elevated firing associated with reward consumption during trials 1–10 trials in one context (Block 1), while showing inhibited responding during trials 11–20 performed in the second context (Block 2). All trials occurred during the same recording session and within the same sensory environment. Bottom row: This cell showed only a slight elevation in firing during reward consumption during Block 1, while showing a marked reduction in firing rate during Block 2. These data demonstrate that, like hippocampus (Smith and Mizumori, 2006b), retrosplenial cortex neurons discriminate learned contexts. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

that disconnection of retrosplenial cortex and hippocampus resulted in a loss of context-specific firing by retrosplenial cortex neurons. This pattern of neural responses, together with similar responses reported for hippocampal neurons (Smith and Mizumori, 2006b), indicates that a context discrimination circuit extends beyond hippocampus. Moreover, hippocampal retrieval of context-specific knowledge and behaviors may be mediated by retrosplenial cortex.

Consistent with previous suggestions (e.g. Sutherland and Hoesing, 1993; McNaughton et al., 1996) that posterior neocortical regions contribute to the determination of the appropriate cognitive and behavioral strategies to be used by the animal, retrosplenial cortex may serve another key function relevant to context discrimination, and that is to facilitate the appropriate behavioral responses when changes in context are detected. From retrosplenial cortex, updated information may be passed on to striatum (Risold and Swanson, 1995), or to premotor areas of cortex (Reep et al., 1990). An important relay in the latter pathway may include the medial precentral nucleus, or PrCm (also referred to as medial agranular cortex, Reep et al., 1987; Reep and Corwin, 1999). Indeed, we have shown that reversible inactivation of the retrosplenial cortex produces significant alteration of PrCm neural correlates (Mizumori et al., 2002; Mizumori et al., under review).

The PrCm may play a pivotal role in the implementation of future cognitive strategies (Mizumori et al., 2002; Mizumori et al., under review) for it projects to the striatum (Reep et al., 1987; Reep and Corwin, 1999) as well as to frontal cortical motor regions (Reep et al., 1990). The PrCm-striatal projection could provide spatial context-dependent movement information to the striatum. The striatum may in turn evaluate the reinforcement consequences of the current behavior that is engaged within a defined context (e.g. Mizumori et al., 1999a; Schultz and Dickinson, 2000). It would be highly adaptive for this striatal analysis to continue through new learning as well as during asymptotic performance. This would allow continual feedback from striatum to cortex regarding the effectiveness of behavioral/cognitive strategies. If the PrCm signal is one that indicates that the context has changed, striatum may become primed for maximal sensitivity to impending input. Such feedback seems critical for an animal’s ability to quickly respond to changes in a familiar environment. For situations in which the strategy remains constant (i.e. during asymptotic performance in a static environment), behaviors can be implemented more quickly via direct PrCm efferent connection to motor cortex. The PrCm, then, may allow organisms to quickly respond to changes in familiar situations as well as to “automate” their responses in well-learned, stable situations.

Connections between PrCm and the prefrontal cortex (PFC) provide what might be considered as the last link in a polysynaptic series of connections between hippocampus and cortical circuitry that is thought to be essential for implementing and orchestrating adaptive behaviors. With this indirect pathway, memory guided interpretations of changes in the expected context (via the indirect path) can be integrated with direct signals from hippocampus (via direct hippocampal-PFC connections; Jay and Witter, 1991) that signal occurrence of a change. Perhaps the latter serves to transform PFC neurons into an “up state” of cell excitability so that it can readily incorporate subsequent information coming in from the indirect pathway. Consistent with this view, PFC neurons exhibit context-dependent anticipatory neural codes for reward information (Pratt and Mizumori, 2001). Also, PFC neural signaling has been found to be conditionally linked to hippocampal output since PFC cell firing can become phase locked to CA1 hippocampal theta rhythms and precession (Hyman et al., 2005; Jones and Wilson, 2005). The entrained PFC neurons tended to be ones whose firing is correlated with specific behavioral acts (Hyman et al., 2005). Furthermore, Hyman et al. showed that testing subjects in different rooms caused some PFC neurons to change the degree to which they were entrained to the hippocampal theta rhythm, suggesting that the degree to which PFC is functionally connected to hippocampus was dependent on the stability of spatial context information.

CONCLUSION

Although it is abundantly clear that the hippocampus engages in context processing, it is less certain how and to what extent this context analysis contributes to the broader cortical learning cir-
cuit. Hippocampus may serve to evaluate the extent to which one’s expectations about contextual features of a situation are met. In navigating animals, such an evaluation requires not only an input from neocortical memory systems to define the expected context, but such information is subject to spatial and temporal organization via temporal (entorhinal) cortical and subcortical input. The outcome of hippocampal analyses is used to update subsequent memory activation networks in cortex and to guide ongoing behavioral acts. Both of these functions may involve the retrosplenial cortex. Recent evidence suggests that retrosplenial cortex is especially related to accurate behavioral choice when current visuomotor input must be linked to past visuomotor associations. Thus, the retrosplenial cortex may serve as one of perhaps many portals through which memory representations can be accessed.

To affect behavioral choice, it is suggested that the direct pathway from hippocampus to PFC serves to change frontal cortex to an excitable state that is ready to quickly respond to the new information arriving via the indirect pathway that includes retrosplenial cortex and the medial precentral nucleus (PrCm). Together, the PrCm and the PFC coordinate a striatal-frontal cortical information system that evaluates the reinforcement consequences of the new context, and then alters the behavioral or cognitive strategy if needed. Many aspects the present proposal are generally in line with different existing theories of hippocampal, entorhinal, retrosplenial or frontal cortical function that are particularly related to accurate behavioral choice when current visuomotor input must be linked to past visuomotor associations. Thus, the retrosplenial cortex may serve as one of perhaps many portals through which memory representations can be accessed.

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REFERENCES


Hippocampus


