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Okatan, Murat

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Boston University
Hippocampal cell assemblies: time encoding neurons or goal representations?

Murat Okatan*

Boston University, Boston, MA, USA
*Correspondence: okatan@bu.edu

Recent evidence obtained in rats suggests that hippocampal neurons become sequentially activated for brief time intervals that are time-locked to task-relevant events while the animals perform reinforcement learning tasks. Similar sequentially activated cells have also been found in the medial prefrontal cortex. It has been suggested that these activity patterns may correspond to goal representations. Recently, a different function has been proposed for these cell assemblies. According to this proposal, these neurons may be viewed as "time encoding" neurons that provide the neural substrate for temporal order memory. This theory was motivated by the observation that hippocampal neurons become sequentially activated during the trace period of an object-odor paired associate task. Here it is pointed out that these neurons also find a functional interpretation within a theory that views such activity patterns as signatures of goal representations. New experiments are proposed to test this theory.

Evidence suggests that hippocampal neurons become activated in a temporal sequence during the 10s-long trace period separating the object stimulus and the odor in a version of Kesner et al.'s (2005) object-odor paired associate task (MacDonald and Eichenbaum, 2009). In this task, rats are presented with one of two objects (A or B) for a brief time period. Then, they spend a trace period that lasts ten seconds. After the trace period the rats are presented with a sand cup that contains one of two odors (1 or 2). The sand cup contains a food reward for object-odor pairs A1 and B2, but not for A2 or B1 (Kesner et al., 2005). Two theories exist concerning the functional role of the hippocampal neurons that are activated in a temporal sequence during the trace period of this task. One theory suggests that these neurons may be viewed as "time encoding" neurons that provide the neural substrate for temporal order memory (Rolls, 2010). Another theory views these cells as an integral component of a goal representation network. Here the latter theory is explained in detail and new experiments are proposed to test it.

According to this theory, sequentially activated hippocampal neurons are the hippocampal branch of a goal representation network that involves the coordinated activity of prefrontal, limbic, basal ganglia and midbrain circuits. The theory proposes that, during reinforcement learning, hippocampal neurons receive inputs from prefrontal goal representations, and that they project information about the reward-predictive value of these goals back to the prefrontal goal representations to mediate a reward-based action selection mechanism (Okatan, 2007a,b, 2009). The theory suggests that, as a result of this prefrontal-hippocampal communication, different cell assemblies are activated in the hippocampus during trips to different goals (Okatan, 2009). Using a statistical neural model, the theory argues that, in a task where a cue presentation signals one of N goals, the presentation of the cue triggers N distinct and simultaneous sets of cue-selective cells in the hippocampus, each set being driven by a distinct prefrontal goal representation and competing to become a set of winner cells (Okatan, 2009). The winner cells probabilistically influence the subsequent choice behavior of the animal, and the activity of the hippocampal winner cells acquires the reward-predictive value of the associated prefrontal goal representation through training by the mesolimbic dopaminergic input, which serves as a reward-prediction error signal (Okatan, 2007a,b, 2009).

For Kesner et al.'s (2005) task, this theory proposes the following: (i) the presentation of one of the objects A or B on a given trial launches object-specific sequentially activated hippocampal cell assemblies, (ii) with learning, the cell assembly that reliably gets rewarded becomes strengthened such that it reliably wins competitive interactions and dominates other cell assemblies, (iii) this strengthening brings with it odor selectivity, which manifests itself first near the time of reward delivery and, with learning, propagates backward in time toward the cue presentation. Thus, the theory proposes that the cells of such winner hippocampal cell assemblies have object and odor selectivity, and influence the animal's upcoming response. The results of MacDonald and Eichenbaum (2009) provide support for some of these proposals. In addition, the theory suggests that the activity of cue and response selective cells is modulated by the magnitude of the reward expected or received by the animal (Okatan, 2007a,b, 2009). Preliminary evidence supporting this has been found in a conditional spatial goal selection task on a T-maze, where the magnitude of the reward delivered to the rats was changed within each session (Okatan et al., 2009a,b).

In this theory sequentially activated cell assemblies are viewed as signatures of how goals are maintained in the working memory (Fujisawa et al., 2008; Pastalkova et al., 2008; Okatan, 2009). This hypothesis may be tested using a modified version of Kesner et al.'s (2005) task. The structure of the new task would be identical to the original task, except that on some randomly selected trials, x seconds after the onset of the trace period, a light would be turned on, where 0 < x < 10 s, would be random. Here, the light signal acts as a "game changer" that indicates that the trial will be aborted at the end of the usual 10-s trace period (i.e. the response period will be skipped), and that a new trial will begin by randomly selecting A or B as the object to be presented. The theory suggests that as soon as the light is turned on, the cell assemblies that are active in the hippocampus and the prefrontal cortex will disappear until the animal's working memory is loaded with information about the next goal, such as the information provided at the next object presentation. Thus, the theory proposes that the cells whose "episode field" (Pastalkova et al., 2008)
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These theories may help design new experiments to further characterize the functional role played by sequentially activated cell assemblies within and outside the hippocampus.

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