Compositional Neural Representations in the Hippocampal Formation and Prefrontal Cortex Underlie Visual Construction and Planning

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Abstract:

The hippocampal formation is critical for spatial and relational inference in navigation problems. The neural code underlying such inference is factorized in the entorhinal cortex (EC) and conjunctive in the hippocampus (HC). A factorized code implies a separate encoding of sensory and relational knowledge, which can be flexibly conjoined to an object representation that reflects both sensory and relational properties. We hypothesize that the same neural mechanisms are employed in complex decision-making and compositional planning, which requires the flexible generalization of knowledge to novel instances. We tested this hypothesis in a task where subjects had to construct novel visual objects based on a set of basic visual building blocks and relations. We found behavioral evidence that subjects form a hierarchical representation of this task that allows them to flexibly apply compositional knowledge to novel stimuli. Using fMRI adaption, we found evidence that the construction of novel objects depends on compositional neural representations in HC-EC and medial prefrontal cortex (mPFC). Further, we found that these structures also encode purely relational information, indicative of a factorized representation. These results suggest that compositional neural representations in the hippocampal formation and prefrontal cortex enable the generalization of abstract knowledge to novel stimuli during visual construction.

Keywords: hippocampal formation; cognitive map; compositional planning; combinatorial generalization

Introduction

The hippocampal formation encodes a ‘cognitive map’ that allows animals to navigate successfully (Behrens et al., 2018; Tolman, 1948). A cognitive map provides an efficient neural representation of knowledge about the structure of the world that enables flexible and generalizable behavior. In the context of spatial navigation, the instantiation of a cognitive map has been associated with place cells in the HC (O’Keefe & Nadel, 1978) and grid cells in the EC (Hafting, Fyhn, Molden, Moser, & Moser, 2005). It has been suggested that place cells encode individual states within a task, such as a particular location in a maze, whereas grid cells encode relational information about those states, such as likely transitions between locations (Momennejad et al., 2017; Stachenfeld, Botvinick, & Gershman, 2016). Recently, the same neural architecture has been implied in non-spatial navigation based on a cognitive map of task structure (Aronov, Nevers, & Tank, 2017; Constantinescu, O’Reilly, & Behrens, 2016; Garvert, Dolan, & Behrens, 2017).

A key principle of functional organization within the hippocampal-entorhinal system is a factorized and conjunctive neural code (Behrens et al., 2018; Manns & Eichenbaum, 2006), which can be organized hierarchically (Stachenfeld et al., 2016). This implies that
EC encodes a separate or factorized representation of sensory and relational components of a stimulus, whereas HC encodes its conjunction and thus forms a high-dimensional representation of a stimulus within a specific structure.

We hypothesized that the same neural mechanisms are at play during complex decision-making and flexible planning behavior. A central aspect of flexible behavior is the generalization of abstract knowledge to novel instances. For example, if we would offer you some tea jelly for dessert, there is a fair chance that you would be able to express your preference for this food item even though you have probably never tasted it before. Importantly, previous work has shown that the construction of such novel goods relies on neural representations in the HC and mPFC (Barron, Dolan, & Behrens, 2013).

In the present study, we investigated the neural representations that underlie such compositional reasoning and the generalization of abstract knowledge to novel instances, called combinatorial generalization (Battaglia et al., 2018). Our key hypothesis was that the generalization of knowledge critically depends on compositional neural representations, where basic building blocks and relational knowledge can be flexibly combined to form novel conjunctive representations (Behrens et al., 2018; Battaglia et al., 2018).

### Results

To investigate the neural representations underlying compositional planning, we developed a task in which subjects learned to construct visual objects using a toolkit of building blocks and relations. As illustrated in Figure 1A, subjects were trained to combine different building blocks by putting them on top or beside each other, without worrying about the physical stability of the resulting object. This task allowed us to probe whether a compositional neural representation in terms of a cognitive map of task structure would emerge after training, based on which a given visual object can be decomposed into its constituent building blocks and relations.

To test whether compositional neural representations can be organized hierarchically, we added an additional layer to the task. During early training, subjects were repeatedly tasked to build specific visual objects that were combinations of two basic building blocks. Later on, they were tasked to build larger visual objects, which were decomposed into two of the smaller visual objects from early training. Thus, although never instructed explicitly, subjects were exposed to a set of ‘compositional building blocks’, which allowed an efficient decomposition of larger visual objects (see bottom panel of Figure 1A). Analysis of participants’ behavior revealed that when they had to construct large visual objects, they relied on ‘compositional building blocks’ more often than predicted by chance (Figure 1B). These findings suggest that the participants indeed formed a hierarchical representation of this task.

In the scanner, subjects passively viewed novel visual objects. These objects were either a combination of two basic or two compositional building blocks connected via ‘on-top-ness’ or ‘besideness’, or one basic or compositional building block alone (see Figure 1C). Participants were tasked to think about the construction of these objects. To ensure that subjects engaged in this task, ten percent of stimuli were followed by a ‘catch trial’, in which subjects were asked about the construction of the previous object.

First, we asked whether this visual construction task depended on neural representations in the hippocampal formation and prefrontal cortex. To test this, we
analyzed fMRI adaption for individual building blocks followed by compounds that contained these particular building block (or vice versa). The key comparison was between conditions where compound AB was preceded by building block A (or vice versa) compared to building block C. Higher adaption for AB when preceded by A compared to when preceded by C would reflect a neural representation for building blocks within a compound, and thus a compositional representation necessary for constructing a novel visual object.

This analysis revealed effects in the HC underlying this visual construction task. Specifically, we found strong basic building block–compound adaption in the medial temporal lobe (Figure 2A). In a region of interest analysis, we found a bilateral effect for basic building block-compound adaption in HC extending into EC; right: \( p=0.001, t_{peak}=5.95 \ [16 \ -38 \ -6] \); left: \( p=0.004, t_{peak}=5.26 \ [20 \ -8 \ -12] \). More exploratory whole-brain analysis revealed an effect for compositional building block-compound adaption in mPFC; \( p<0.001, t_{peak}=5.95 \ [16 \ -38 \ -6] \), cluster size = 8797.

We also found a small-volume corrected effect for compositional building block–compound adaption in the bilateral hippocampus: right: \( p=0.045, t_{peak}=4.27 \ [20 \ -30 \ -6] \) and left: \( p=0.005, t_{peak}=5.25 \ [20 \ -36 \ -20] \). More exploratory whole brain analysis revealed an effect for compositional building block-compound adaption in the bilateral hippocampus: right: \( p=0.006, t_{peak}=5.13 \ [-24 \ 2 \ -16] \) and left: \( p=0.031, t_{peak}=4.39 \ [-12 \ -14 \ -24] \). We also found a whole brain effect for this analysis in a large cluster containing both left and right HC, \( p<0.001, t_{peak}=5.95 \ [16 \ -38 \ -6] \), cluster size = 8797.

To test this, we probed for fMRI adaption that reflects a purely relational representation, namely contrasting same relation transitions (i.e. an object constructed with ontopness/besideness preceded by an object with the same relation) with different relation transitions. In this analysis, we found a small-volume corrected effect in left anterior EC for basic compounds, \( p=0.052, t_{peak}=4.17 \ [-16 \ 4 \ -30] \). We did not detect an effect in right HC–EC or any other whole brain effects for basic compound relation adaption, or any effects for compositional compound relation adaption.

Further, we tested the same question in a second analysis probing for adaption ‘within’ a compositional compound. As displayed in Figure 1A, compositional compounds are constructed with two compositional building blocks, which themselves consist of two basic building blocks. Importantly, these two compositional building blocks can be built with the same or different relations. Consequently, we expect stronger threshold of \( p<0.001 \), family-wise error corrected at the peak level.
(parametric) adaption for a higher proportion of ‘same’ relation solutions within a compositional compound. We tested this hypothesis and found a small-volume corrected effect in the HC that reflected ‘within object’ relational adaption; right: p=0.028, t_{peak}=4.56 [24 -36 -4] and left: p=0.039, t_{peak}=4.37 [-20 -36 -4]. On a whole-brain level, we also detected parametric effects in parietal cortex, p<0.001, t_{peak}=5.90 [-12 -36 44], cluster size = 4247; callosal body, p=0.007, t_{peak}=5.65 [4 -14 24], cluster size = 348; inferior frontal gyrus, p=0.008, t_{peak}=5.07 [-36 34 6], cluster size = 333; and mPFC p=0.012, t_{peak}=4.50 [0 46 2], cluster size = 308.

Figure 3: A) Adaption effect in anterior EC for basic ‘same relation’ compound transitions (light blue). Overlaid is a previously reported effect in EC for relational navigation in conceptual space (Constantinescu et al., 2016, pink). Effects thresholded at T=2.3. B) Parametric adaption effect in HC reflecting the proportion of ‘same relation’ solutions in compositional compounds (green), overlaid on Figure 2A. Effects thresholded at T=3.5. All effects masked for HC and EC.

Conclusion

We developed a paradigm in which subjects had to use abstract compositional knowledge to construct novel visual objects. We found evidence for such compositional neural representations in the hippocampal formation and mPFC. The latter effect was specifically pronounced for compositional building blocks within larger compounds, suggestive of a hierarchical organization in line with behavioral measures. Further, we found evidence suggesting that objects were encoded purely in terms of their relational properties in the hippocampal formation and mPFC. Taken together, our results suggest that compositional representations in the HC-EC system and mPFC underlie the flexible construction of novel stimuli, which is a central aspect of flexible decision-making and compositional planning.

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References


