Reduction of Theta Rhythm Dissociates Grid Cell Spatial Periodicity from Directional Tuning
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cy of the hippocampo-entorhinal circuitry (25) when pacing by septal inputs was reduced.

Hippocampal place cells are also spatially modulated cells without periodicity in their firing pattern. Their spatial firing patterns are partially affected in the radial maze after septal inactivation and in the open field during reduced cholinergic neurotransmission (26, 27). For comparisons with MEC cells, we therefore recorded from hippocampal place cells in the same experimental design while reducing theta oscillations to the same extent as in MEC recordings (fig. S2) (20). We observed a substantial decrease in the firing rate of hippocampal place cells, and the spike trains of individual hippocampal cells showed reduced theta modulation (Fig. 3A and fig. S2). Place fields remained well-defined and at corresponding locations (Fig. 3, B and C, figs. S9 and S10, and table S1).

By silencing the septal area, we diminished theta oscillations in the entorhino-hippocampal circuitry and showed that the periodic firing of grid cells does not persist. Subcortical inputs to hippocampus and parahippocampal cortices are thus essential not only for theta oscillations but also for sustaining the spatial periodicity of grid cells. These findings are consistent with the theory that grid cells emerge from the interference between multiple precisely tuned theta oscillations within individual cells (12–14). Alternatively, the silencing of septal inputs to the MEC might result in the desynchronization of grid cells, so that the local network of cortical cells can no longer generate oscillatory interference (12, 25). Our data also identified a subpopulation of grid cells that do not regain their spatial regularity when theta oscillations recover (fig. S6). This suggests that a fraction of grid cells might not be directly participating in the generation of grid patterns, but rather becomes associated with other grid cells by plasticity-dependent mechanisms.

A parallel study (29) has independently discovered that grid cell firing does not persist during reduced theta oscillations and that other cell types are less affected. Together, our results show that the neuronal network mechanisms that sustain the periodic spatial firing of grid cells are different from those required for other firing correlates in the entorhino-hippocampal circuitry, including head-direction cells and place cells. The effect on grid cells is likely not mediated through effects of septal silencing on the firing of hippocampal place cells, because it has been shown that grid cell firing initially remains intact after the hippocampus has been silenced (30). Subcortical inputs are therefore necessary for the neural computations in the MEC that generate grid-like, periodic, spatial firing patterns, whereas the firing locations of place cells largely persist after inputs from grid cells and from subcortical areas to the hippocampus have substantially changed.

References and Notes
19. Materials and methods are available as supporting information on Science Online.
20. Hippocampal and MEC cells were recorded in separate experiments (except for the data in fig. S10). In the MEC, grid cells (n = 261) and other cell types (n = 27) simultaneously with grid cells, and n = 37 without grid cells) were recorded.
21. The firing fields of a subset of MEC cells had the stripe-like appearance of boundary vector cells, but only 6 of 19 fired maximally at the border. A field size criterion (>625 cm²) rather than the border score was thus used to include these cells (see also fig. S8).
31. The authors thank T. Solstad and E. Mankin for Matlab scripts and M. Hasselmo, L. Squire, and M. Scanziani for comments and discussions. Supported by the Walter F. Heiligenberg Professorship, NSF/NIH/Bundesministerium für Bildung und Forschung grant 1010463, Ellison Medical Foundation grant AG-NS-0724-10, the Alfred P. Sloan Foundation, and Alzheimer’s Association grant NIGR-09-133414.

Supporting Online Material
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Reduction of Theta Rhythm Dissociates Grid Cell Spatial Periodicity from Directional Tuning

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Grid cells recorded in the medial entorhinal cortex of freely moving rats exhibit firing at regular spatial locations and temporal modulation with theta rhythm oscillations (4 to 11 hertz). We analyzed grid cell spatial coding during reduction of network theta rhythm oscillations caused by medial septum (MS) inactivation with muscimol. During MS inactivation, grid cells lost their spatial periodicity, whereas head-direction cells maintained their selectivity. Conjunctive grid–by–head-direction cells lost grid cell spatial periodicity but retained head-direction specificity. All cells showed reduced rhythmicity in autocorrelations and cross-correlations. This supports the hypothesis that spatial coding by grid cells requires theta oscillations, and dissociates the mechanisms underlying the generation of entorhinal grid cell periodicity and head-direction selectivity.

The role of oscillations in neural coding is controversial. Theta frequency oscillations (4 to 11 Hz) play an important role in memory behavior (1–4) and code spatial location by the precession of spike timing relative to theta oscillations (theta phase precession) in the hippocampus (5, 6) and medial entorhinal cortex (MEC) (7). However, disagreement remains about whether theta oscillations are critical to spatial coding by neurons. Grid cells (8, 9) in the MEC provide a powerful example for testing the theoretical role of oscillations in neural coding. Some computational models of grid cells use network theta rhythm oscillations to generate grid cell spatial periodicity (10, 11). These models simulate the phase of spike timing in grid cells (7)
and have successfully predicted that the spatial scale of grid cell firing correlates with measures of intrinsic rhythmicity (12, 13). Recent models also show the potential role of theta oscillations for updating position in attractor dynamic models of grid cells (14). In rats, we tested the role of theta rhythm oscillations in the spatial coding of grid cells by testing the spatial periodicity of grid cells during pharmacological disruption of theta rhythm oscillations.

Lesions or inactivation of the medial septum (MS) in rats cause a disruption of theta oscillations in the entorhinal-hippocampal system (2, 4, 15, 16) and cause spatial memory impairments (1–4, 17, 18). We performed microinfusions of muscimol to pharmacologically inactivate the MS bilaterally (1, 17). Simultaneously, we used tetrodes in the MEC to monitor the spiking activity of grid cells, head-direction cells, and conjunctive grid–by–head-direction cells.

After infusions of muscimol into the MS (Fig. 1A, left), recordings in the MEC (Fig. 1A, right) demonstrated a clear decrease in the power of theta oscillations in the MEC local field potential (Fig. 1B) and a strong reduction in the spatial periodicity of grid cells (Figs. 1, C and D, and 2A and fig. S3). Recovery of theta rhythm and spatial periodicity occurred in recordings 3 to 6 hours and 24 hours after the infusion. The “gridness” (hexagonal regularity of firing) score measured before the MS inactivation showed a significant decrease after the inactivation (n = 29 cells; baseline: mean ± standard error: 0.64 ± 0.05 gridness score; inactivation: −0.27 ± 0.06 gridness score, P < 0.001) (Fig. 3A) that recovered at 3 to 6 hours and 24 hours (3 to 6 hours: n = 26, 0.35 ± 0.10 gridness score; 24 hours: n = 21, 0.46 ± 0.09 gridness score). Because the firing rates of grid cells were also reduced (n = 29; baseline: 1.88 ± 0.20 Hz; inactivation: 1.17 ± 0.21 Hz, P < 0.001), we subsampled the spiking of the baseline recordings before infusion to match the same overall firing rate after infusion to confirm that the reduction of gridness still appeared when compared to subsampled data (n = 29; subsampled: 0.58 ± 0.05 gridness score; inactivation: −0.27 ± 0.06 gridness score, P < 0.001) (column 2 in Fig. 1, C and D, and fig. S3).

There were no differences in running speeds between the baseline, MS inactivation, 3- to 6-hour, or 24-hour recovery periods (Fig. 4C and fig. S8). Control infusions of phosphate-buffered saline (PBS) into the MS did not alter theta oscillations or the grid cell spiking patterns in the MEC (Fig. 1E and fig. S7).

Many neurons in the MEC are conjunctive grid–by–head-direction cells (19, 20), head-direction cells (19, 21), or cells showing spatial selectivity without grid cell periodicity, such as border cells (22, 23). We observed a dissociation in the effect of MS inactivation on the spatial periodicity and head-direction specificity of entorhinal neurons. After muscimol infusion, conjunctive grid–by–head-direction cells fired with reduced spatial periodicity (Fig. 2A)}
and fig. S3) but retained their directional preference, \( \phi \) (\( n = 8 \), 1.72 ± 1.15 ° (ΔΦ) degrees, not significant (n.s.)) (Figs. 2A and 3B and fig. S3). After infusion, head-direction cells retained their directionality (\( n = 11 \); baseline: 37.89 ± 8.9° Watson \( U^2 \) statistic; inactivation: 39.55 ± 14.54 Watson \( U^2 \) statistic, n.s.; 24.64 ± 9.17 ° (ΔΦ) degrees, n.s.) (Figs. 2B and 3B and fig. S4) and as a population retained their firing rates (\( n = 11 \); baseline: 2.7 ± 0.72 Hz; inactivation: 1.9 ± 0.65 Hz, n.s.) (Fig. 3A). Cells with positive gridness scores showed a reduction of spatial information (grid cells: \( n = 29 \); baseline: 1.08 ± 0.07 bits per spike; inactivation: 0.56 ± 0.05 bits per spike, \( P < 0.001 \)) (Fig. 3A). Cells with negative gridness scores but high spatial information also showed a reduction of spatial information (spatially modulated non-grid cells: \( n = 9 \); baseline: 1.3 ± 0.15 bits per spike; inactivation: 0.70 ± 0.10 bits per spike, \( P < 0.01 \)) (Fig. 3A and fig. S5).

The slow onset and long half-life that are characteristic of muscimol effects (1, 17) permitted us to quantify the relationship between theta power and grid cell spiking. Using 10-min sliding window calculations, we found a clear theta power and grid cell spiking. Using 10-min

\[ \text{Power} = \frac{\sum (\text{spike} \times \text{theta})}{\sum \text{spike}} \]

characteristic of muscimol effects (\( P < 0.01 \)) (Fig. 3A and fig. S5).

- **Fig. 2.** Head-direction selectivity is spared. (A) Examples of four conjunctive grid-by-head-direction cells. The selectivity for head direction observed before infusion (Baseline) remains during MS inactivation despite the loss of grid cell spatial periodicity in the spatial maps. (B) Four head-direction–selective cells without spatial periodicity before infusion (Baseline) maintain head-direction preference during MS inactivation. Overall firing rates are not reduced in either cell type.
Fig. 3. Statistics for grid cells, head-direction cells, and spatially modulated cells. (A) (Top) Three gridness measures show a sharp reduction in gridness 15 to 75 min after muscimol infusion (MS Inactivation), with recovery at 3 to 6 hours (3-6hr) and 24 hours (24hr) after infusion. (Middle) Spatial map correlations and spatial information for grid cells, PBS infusions, and spatially modulated cells for each period. (Bottom) Average firing rate during each period for all cell types. (B) Head-direction cell statistics. Mean resultant length and change in head-direction preference angle from baseline (in degrees) for all head-direction cells and conjunctive cells during each period are shown. (C) Average gridness score at different levels of theta power. There is a consistent relationship between gridness and theta power. (D) Time course of the effect of MS inactivation on theta power and gridness score computed in a sliding 10-min window at 1-min intervals for each period during the experiment, showing similar reduction and recovery.
Simultaneously recorded neurons during MS inactivation. The combined data suggest that grid cell spatial periodicity is not essential for place cell responses in familiar environments, which is consistent with studies suggesting that the development of place cell responses does not depend on the development of grid cell responses (27, 28).

These data support the hypothesized role of theta rhythm oscillations in the generation of grid cell spatial periodicity (10–12) or at least a role of MS input. The loss of grid cell spatial periodicity could contribute to the spatial memory impairments caused by lesions (3, 4, 18, 29) or inactivation (1, 2, 17) of the MS. These data support a role of neuronal oscillations in the coding of spatial information.

Fig. 4. Temporal coordination of local field potential and spike timing disrupted during MS inactivation. (A) Top) Raw electroencephalogram shows reduction of theta rhythm during MS inactivation. (Bottom) Power spectrum of MEC local field potential for 13 sessions during periods before (Baseline), 15 to 75 minutes after (MS inactivation), 3 to 6 hours after, and 24 hours after infusion. (B) Top) For each period, the left graph shows temporal cross-correlations and autocorrelations during each period. There is a reduction of theta rhythm after infusion in cross-correlations suggests loss of timing relationships between cells. There is a decrease in frequency during MS inactivation. (Middle) Temporal autocorrelations of spiking show reduction in theta power and frequency during MS inactivation. (Bottom) Percentage of cells showing theta rhythmicity in cross-correlation for each period did not change during infusion. (C) Top) Cross-correlation of theta power (the ratio between power in the theta and delta bands) decreases during MS inactivation. (Right) Average running speed for each period did not change during infusion. (D) Top) Cross-correlation of theta power for each cell pair or cell during each period shows a reduction in frequency during MS inactivation and an increase 3 to 6 hours after infusion. (Bottom) Percentage of cells showing theta rhythmicity in cross-correlations and autocorrelations during each period. There is a reduction of theta rhythmic cells and cell pairs during MS inactivation.

Referenices and Notes
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References
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