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Many animals, on air, water, or land, navigate in three-dimensional (3D) environments, yet it remains unclear how brain circuits encode the animal’s 3D position. We recorded single neurons in freely flying bats, using a wireless neural-telemetry system, and studied how hippocampal place cells encode 3D volumetric space during flight. Individual place cells were active in confined 3D volumes, and in >90% of the neurons, all three axes were encoded with similar resolution. The 3D place fields from different neurons spanned different locations and collectively represented uniformly the available space in the room. Theta rhythmicity was absent in the firing patterns of 3D place cells. These results suggest that the bat hippocampus represents 3D volumetric space by a uniform and nearly isotropic rate code.

Navigation is crucial for survival, and the need to navigate cuts across the animal kingdom. Several navigational strategies are used by animals, among them maplike navigation (1). This strategy relies on a set of brain structures, at the hub of which is the hippocampus (1, 2). This brain area contains “place cells,” neurons that activate when the animal enters a restricted region of the environment, the place field (1, 2). Since the discovery of place cells in rodents, these neurons have been reported across mammalian species (1–9), and their functional properties have been extensively researched (1, 2). However, the spatial and temporal properties of place cells have never been studied in animals moving freely through 3D volumetric space, without any constraints to particular planes of motion. Indeed, in all studies to date, animals have always been navigating on one- or two-dimensional (2D) planes (1, 10)—either horizontal (3, 7, 11–15), tilted (3, 4, 16–18), or vertical (9, 13, 16)—and thus it remains unresolved how place cells encode the animal’s position throughout the entire volume of a 3D volumetric space. We developed wireless recording methodology for freely flying animals (19) and recorded single-neuron activity from hippocampal area CA1 of Egyptian fruit bats flying through 3D space.

Bats were tested in one of two setups: either a large cuboid-shaped flight room, where bats performed a naturalistic foraging task (Fig. 1A and fig. S1; room size 580 × 460 × 270 cm; n = 3 bats) or a 3D cubic flight arena of smaller dimensions, where bats searched randomly for food (Fig. S2; room size 290 × 280 × 270 cm; n = 2 bats) (19). In both setups, bats exhibited complex naturalistic flight trajectories, during which they moved through all the three dimensions of the room (Fig. 1B and fig. S3), traversed large distances (Fig. 1C, left), and flew at high speeds (Fig. 1C, right; and fig. S4). These flight maneuvers resulted in dense and rather uniform coverage of the environment’s 3D volume (figs. S5 and S6).

Single-unit activity was recorded from freely flying bats, using a tetrode-based microdrive and a custom lightweight four-channel neural telemetry system designed for flying bats (Fig. 1, D to F, and figs. S7 to S10). The telemetry system allowed the transmission of action potentials from the four channels of one tetrode, with high fidelity (Fig. 1, D, E, and G, and figs. S7 to S9), throughout all the locations in the flight room (figs. S7 and S8) and with very little interference from movement-related noise (fig. S11) (20).

The ability to monitor 3D spatial position and record the activity of individual neurons in freely flying bats allowed studying the spatial coding of 3D volumetric space by hippocampal neurons. We recorded a total of 139 well-isolated neurons from five bats in the dorsal CA1 region of the hippocampus (19). About half of the cells (73 out of 139 (73/139) or 53%) were active during flight, and 75% of the active cells (55/73) were classified as place cells (fig. S12), becoming active when the bat flew through a restricted volume of the available environment (19). Figure 2, A to E, shows the spatial spiking activity of a single hippocampal neuron recorded during flight. This neuron fired nearly exclusively in a confined region of the environment, and this region was restricted in all three dimensions (Fig. 2, A and B, and fig. S13). The firing field of this neuron remained highly stable across the recording session, both in its spatial location and its firing rate (Fig. 2, C and D, and fig. S14A); furthermore, the cell was reliably activated on most of the individual flight passes through the place field (Fig. 2E). Likewise, the majority (75%) of neurons that were active during flight...
exhibited strong spatial tuning (Fig. 2, F to K, and fig. S15), and their firing fields were highly stable throughout the recording session (fig. S14, B and C)—similar to previous reports for rat CA1 place cells (1, 22–24).

In both setups, the place fields of different neurons varied in size and location and spanned much of the available 3D space (Fig. 3, A to D). Most place cells discharged within a single 3D firing field (62%; Fig. 3, E and F). The fraction of cells with two or more fields was higher in the large cuboid flight room (43% or 15/35, Fig. 3E) than in the smaller cubic enclosure (30% or 6/20 cells; Fig. 3F; bootstrap test, $P < 0.001$) (19), which is in agreement with previous reports in rats, where the number of 2D firing fields increased with the size of the environment (21).

Next we asked whether 3D space is represented isotropically in the hippocampus of flying bats; namely, do place fields have the same size in all dimensions? Across the population, the sizes of individual place fields along each of the three cardinal axes of the room were quite similar (Fig. 3, G and H). Utilizing the 3D volumetric coverage of the room by the bats (fig. S1C and fig. S15), we then asked whether 3D place fields are spherical (isotropic), or perhaps anisotropy might be found along some noncardinal axis. We fitted a 3D ellipsoid to the place-field shape, where the ellipsoid was not constrained to be parallel to any of the room walls, and an “elongation index” was defined as the ratio of the longest to the shortest axes of the ellipsoid (fig. S16). To test for the significance of the computed elongation index, we implemented a shuffling procedure in which we randomly redistributed the spikes of the place field along the bat’s flight trajectory, constraining them to be within a perfect sphere of the same volume as the ellipsoid, and tested whether the real elongation index was outside the shuffling-based confidence interval (19); that is, whether the actual elongation was significantly different from that expected from an underlying perfect sphere, given the animal’s behavior (Fig. 3I). Only 9% of the fields (Fig. 3I, red dots; 7/82 fields) showed significant elongation, exhibiting compression along the $z$ axis of the room (e.g., neuron 1 in fig. S17C), but sometimes also diagonal field shapes (e.g., neuron 2 in fig. S17C). However, 91% of the place fields in both arenas (75/82 fields) were not significantly different from an underlying sphere (Fig. 3I, green dots; examples in fig. S17C, neurons 3

Fig. 1. Behavioral and neural recordings from freely flying bats. (A) Egyptian fruit bats were flying freely in a large cuboid-shaped room, searching for food hidden in the arms of an artificial tree in the center of the room. cam1 and -2, cameras; mic1 and -2, ultrasonic microphones; ant1 and -2, receiving antennas for neural telemetry signal. The room, tree, and bat are drawn to scale; cameras, antennas, and microphones are enlarged here for display purposes. (B) Examples of 3D flight trajectories of bats. Circles, video frames (25 Hz); the arrowhead indicates the start of flight. Flight durations, from left to right, were 8.0, 7.7, and 6.8 s, respectively. (C) Distribution of total flight distances (left), mean velocity (center), and peak flight velocity (right) across all recording sessions in the cuboid setup. (D and E) Simultaneous wired (tethered) and wireless neural recordings. (D) Top: Raw neural trace recorded from bat hippocampal area CA1 from a single channel of the tetrode, before transmission (top), after transmission (bottom), and overlay (center). Bottom: Zoom-in onto a single action potential. (E) 2D projections of spike energies of CA1 neurons, before (top) and after (bottom) wireless transmission. Red and blue dots, clusters corresponding to two neurons; red dots in (E) are spikes from the red cluster. (F) Telemetry system on a flying bat, drawn to scale [Illustration: S. Kaufman]. (G) Neural traces (right) from the four channels of a tetrode, recorded from bat CA1 during one flight segment (left, gray; the magenta portion corresponds to the duration of the neural traces shown on the right).
and 4). Further, comparing the field sizes specifically along the $x, y, z$ cardinal axes of the room showed that, for these 91% of the fields, there was no significant elongation in any of the cardinal directions (fig. S17A; Sign test, $P > 0.15$ for all comparisons). Thus, for the large majority of neurons, 3D place fields were indistinguishable from isotropic spheres.

We next examined whether the locations and sizes of place fields are distributed uniformly across space, as indicated by the population plots in Fig. 3, A to D. Indeed, the size of place fields did not show significant correlation with the distance from the flight-enclosure center, in either of the setups, indicating similar resolution at room center and near the walls (Fig. 3J; correlation coefficient: $r = -0.20$, $P = 0.13$ for the rectangular cuboid flight-room; $r = -0.02$, $P = 0.91$ for the cubic enclosure). Place fields were slightly bigger in the large cuboid flight room than in the smaller cubic flight enclosure (102.3 ± 5.5 cm compared to 94.3 ± 6.9 cm, respectively). Field locations were distributed quite uniformly, both along the radial

![Fig. 2. Examples of 3D place cells recorded from the hippocampus of flying bats. (A to E) An example cell. (A) 3D representation of the neuron’s spatial firing. Top left: Spikes (red dots) overlaid on bat’s position (gray lines); shown also are the spike waveforms on the four channels of the tetrode (mean ± SD). Top right: 3D color-coded rate map, with peak firing rate indicated. Bottom: Convex hull encompassing the neuron’s place field (red polygon) and the volume covered by the bat during flight (gray polygon). (B) 2D projections of the raw data (top) and color-coded rate maps (bottom). (C and D) Stability of the neuron’s spatial firing for the first versus second half of the recording session; shown are 3D rate maps (C) and 2D projections (D). Peak firing rate is indicated for each map. (E) Reliability of firing across 61 flight passes through the place field. Bottom: raster plot showing spikes during individual passes (time 0, point closest to field center). Top: Spike density function, unsmoothed; bin size, 40 ms. (F to K) Six additional place cells from the hippocampus of different bats. Same notation as in (A). When a neuron had more than one place field, different fields were marked with different colors (K). The neuron in (J) was recorded in the cubic enclosure; the other neurons are from the rectangular-cuboid room.](https://www.sciencemag.org)
dimension (Fig. 3J) and along each of the three major axes (Fig. 3, K and L), indicating that the entire available environment was represented rather uniformly by the population of place cells.

Most hippocampal neurons in rodents exhibit strong theta modulation of their firing rate, which can be readily observed in spike-train temporal autocorrelograms (22–25). However, it has been difficult to find firing-rate oscillations in the hippocampus of several other mammals, including wheeling monkeys (26) and crawling bats [(3, 5); fig. S18]. But could theta rhythmicity occur during flight? We hypothesized this might be the case, because flight entails several conditions favorable for the generation and detection of theta rhythmicity, such as high-velocity movements (fig. S4), high firing rates (Fig. 2 and fig. S19), and strong oscillatory behaviors by the animal (fig. S20). Surprisingly, however, we found the opposite. The theta rhythmicity of hippocampal place cells was assessed by computing...
a standard “theta index” (Fig. 4, A to C, and fig. S21) (19), which has been used in previous studies in rats and bats to assess theta rhythmicity (5, 23, 24). A statistical shuffling procedure was applied for each individual neuron to test for significance (or lack thereof) of theta modulation in bat hippocampal neurons (19). Only 3.6% of the 3D place cells in both experimental setups (2/55 neurons) exhibited significant theta-modulated firing (19), but this rhythmicity was very weak (fig. S21) and, at the population level, was not different than would be expected by chance (binomial test, \( P = 0.77 \)) (19). The degree of theta modulation of bat hippocampal neurons was stably low when comparing crawling versus the two flight setups (Fig. 4D), despite dramatic increases in movement velocity (speeding up almost two orders of magnitude: Fig. 4, D and E, and figs. S4 and S19) and a much higher firing rate of neurons during rapid flight (Fig. 4E and fig. S19). Theta rhythmicity was not correlated with the bat’s echolocation rate (fig. S21H). Thus, we found a near-absence of spike-train theta rhythmicity in the bat hippocampus, both during crawling and during free flight. These results are strikingly different from the robust theta-rhythmic spiking of place cells in rodents (1) but are consistent with our previous reports in crawling bats (3, 5). The near absence of theta modulation in the firing of hippocampal place cells in bats (this study) and monkeys (26) is incongruent with dual-oscillatory models of spatial coding (1, 2).

What factors may shape the nearly isotropic 3D coding that we found in >90% of bat hippocampal place cells during flight? Bats are mammals that naturally move and navigate in 3D volumetric environments, and additionally their echolocation system provides them with volumetric 3D sensory information, including depth (10, 27). Thus, isotropic 3D spatial coding might have evolved in bats because of the pressures to encode and retrieve 3D sensory information. This interpretation also implies possibly different encoding of 3D space in land-dwelling mammals that typically do not move volumetrically in 3D during ontogeny but instead rely mainly on horizontal self-motion cues. Indeed, a recent study (16) reported anisotropic coding in rat hippocampal neurons, with strong elongation of place fields in the vertical dimension [see isotropic coding reported in a different study in rats (9) and discussion in (10, 28)]. However, movement through 3D volumetric spaces is characteristic for many nonflying mammals, and we hypothesize that, as in bats, this could have created an evolutionary pressure to represent 3D volumetric position in the brain circuits of many species. The extent to which this prediction applies to humans is an intriguing question, because on one hand we evolved from monkeys that jumped across 3D volumes, but on the other hand we typically locomote along 2D surfaces. This will have to be studied in humans and monkeys in real and virtual 3D settings.

Finally, would these findings extend also to real-life navigation in the wild (29)? The average size of 3D place fields found here (~1 m) was consistent with the predictions of a computation model of 3D place cells, for room-sized environments (30); the same model also predicts place fields on the order of kilometers

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**Fig. 4. Near-absence of theta rhythmicity in the firing of bat hippocampal place cells during flight.** (A and B) Two example cells. From left to right: 3D color-coded rate maps, with peak firing rate indicated; spike-train temporal autocorrelograms; power spectrum of autocorrelogram; and shuffled distribution of theta indexes (red lines, 95% confidence intervals; blue line, actual theta index of the neuron) (19). There was a near-absence of power in the theta band (5 to 11 Hz, gray rectangle in the third column; power normalized to 1). See fig. S21 for additional examples. (C) Theta indexes for all place cells from the rectangular-cuboid flight room (left) and cubic flight enclosure (right). Neurons are sorted by theta-index values. Shaded area, 95% confidence intervals. Only a small fraction of place cells were significantly theta-modulated (red dots: 3.6%, or 2/55 of neurons in both setups together), whereas nearly all the place cells were not modulated (green dots: 96.4%, or 53/55 of neurons). (D and E) Theta indexes (D) and peak firing rates (E) as a function of the bat’s velocity, under three different conditions (crawling, flying inside the cubic enclosure, or flying in the cuboid flight room).
in bats flying in their natural habitat. Further, navigation requires more than the hippocampal spatial signal: It also entails decision-making, goal-directed behaviors, sensory-motor integration, and other cognitive processes (mediated by brain structures such as the striatum, cerebellum, and prefrontal and parietal cortices). Thus, to elucidate the neural basis of real-life navigation in bats (27, 29), it would be essential to record neuronal activity from the hippocampal formation (and additional brain structures) in bats navigating over distances of kilometers.

References and Notes

Developmental Decline in Neuronal Regeneration by the Progressive Change of Two Intrinsic Timers

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Like mammalian neurons, Caenorhabditis elegans neurons lose axon regeneration ability as they age, but it is not known why. Here, we report that let-7 contributes to a developmental decline in anterior ventral microtubule (AVM) axon regeneration. In older AVM axons, let-7 inhibits regeneration by down-regulating LIN-41, an important AVM axon regeneration–promoting factor. Whereas let-7 inhibits lin-41 expression in older neurons through the lin-41 3′ untranslated region, lin-41 inhibits let-7 expression in younger neurons through Argonaute ALG-1. This reciprocal inhibition ensures that axon regeneration is inhibited only in older neurons. These findings show that a let-7–lin-41 regulatory circuit, which was previously shown to control timing of events in mitotic stem cell lineages, is utilized in postmitotic neurons to control postdifferentiation events.

We use Caenorhabditis elegans to study developmental decline in neuronal regeneration (Fig. 1A) (1). As in vertebrates, advancing development leads to decreased axon regenerative capacity in C. elegans (Fig. 1C) (2–4). The timing mechanism that controls developmental decline in neuronal regeneration is poorly understood (2–5). Because heterochronic genes are implicated in regulating developmental timing and aging in C. elegans (6), we hypothesized that they might regulate developmental decline in neuronal regeneration. The heterochronic pathway involves a number of microRNA–regulated posttranscriptional genetic circuits (7, 8), including an important interaction between the let-7 microRNA and its direct target, lin-41, which encodes a tripartite motif (TRIM) protein (9, 10). We show here that let-7 and lin-41 function in postmitotic neurons to time their differentiation and postdifferentiation events. Our study reveals that the intrinsic timing mechanism that controls developmental decline in neuronal regeneration depends on the progressive increase of let-7 and the progressive decrease of lin-41 in neurons, with let-7–lin-41 reciprocal inhibition having a role in this process.

MicroRNA expression is either spatially restricted or temporally regulated in neuronal development (11–14). To explore the role of microRNAs in neuronal regeneration, we examined anterior ventral microtubule (AVM) axon regeneration in mutants defective in microRNA biogenesis, dcr-1 and alg-1 (15). Although dcr-1 hypomorphic mutant animals displayed normal AVM axon regeneration (fig. S1A), regenerating AVM neurons of adult alg-1 mutants extended axons 2.5 times the length of those in adult wild-type animals (Fig. 1, B to D). In addition, regenerating axons of alg-1 adults often displayed compact growth cones (fig. S1C). These regeneration phenotypes in adult alg-1 mutants are reminiscent of those in wild-type animals at an earlier developmental stage (Fig. 1, B to D, and fig. S1), which suggests that alg-1 mutations may retard a normal developmental decline in axon regeneration. Consistent with this conclusion, we observe that although AVM axon regeneration in alg-1 mutants is similar to that of wild-type animals at the larval L2