Functional organization of the hippocampal longitudinal axis

Bryan A. Strange^{1,2}, Menno P. Witter³, Ed S. Lein⁴ and Edvard I. Moser³

Abstract | The precise functional role of the hippocampus remains a topic of much debate. The dominant view is that the dorsal (or posterior) hippocampus is implicated in memory and spatial navigation and the ventral (or anterior) hippocampus mediates anxiety-related behaviours. However, this 'dichotomy view' may need revision. Gene expression studies demonstrate multiple functional domains along the hippocampal long axis, which often exhibit sharply demarcated borders. By contrast, anatomical studies and electrophysiological recordings in rodents suggest that the long axis is organized along a gradient. Together, these observations suggest a model in which functional long-axis gradients are superimposed on discrete functional domains. This model provides a potential framework to explain and test the multiple functions ascribed to the hippocampus.

Hippocampus

In animal studies, the term describes dentate gyrus (DG) and CA subfields. In human functional MRI studies, the term typically includes the DG, CA subfields and subiculum (except in high-resolution functional MRI)

¹Laboratoru for Clinical Neuroscience. Centre for Biomedical Technology Technical University of Madrid, Campus de Montegancedo, 28223 Pozuelo de Alarcón, Spain. ²Department of Neuroimagina. Alzheimer's Disease Research Centre Reina Sofia-CIFN Foundation, Calle Valderrebollo 5 28071 Madrid Spain ³Kavli Institute for Systems Neuroscience and Centre for the Biology of Memory, MTFS, Olav Kyrres gate 9, Norwegian University of Science and Technology, NO-7489 Trondheim, Norwau, ⁴Allen Institute for Brain Science, Seattle, Washington 98103, USA. Correspondence to B.A.S. e-mail: bryan.strange@ctb. upm.es doi:10.1038/nrn3785

The hippocampus is a medial temporal lobe structure that is critically involved in episodic memory and spatial navigation¹⁻⁷. Its long, curved form is present across all mammalian orders and runs along a dorsal (septal)-toventral (temporal) axis in rodents, corresponding to a posterior-to-anterior axis in humans (FIG. 1a,b). The same basic intrinsic circuitry is maintained throughout the long axis and across species (FIG. 1c). Despite this conserved intrinsic circuitry, the dorsal and ventral portions have different connectivities with cortical and subcortical areas, and this has long posed a question as to whether the hippocampus is functionally uniform along this axis. In this article, we review cross-species data that show how the seemingly disparate functions ascribed to the hippocampus can be accommodated by a model in which different functional properties exist along the longitudinal axis.

The severe memory impairment suffered by patient H.M. after a bilateral hippocampal resection¹ led to intensive study⁸ of patients and animal models with hippocampal damage, with an ensuing characterization of hippocampal function in terms of declarative memory², encompassing both episodic and semantic memory. At the same time, however, evidence emerged for a hippocampal role in spatial memory, based on the discovery of hippocampal place cells^{9,10} and the demonstration that hippocampal lesions impair spatial memory⁴. Both the declarative memory hypothesis¹¹ and the spatial mapping hypothesis¹² of hippocampal function proposed a unitary model in which the entire hippocampus

is dedicated to a single, general type of memory. In light of subsequent evidence for a hippocampal role in emotional memory¹³, an alternative model that could account for different types of memory is that each type of memory depends on separate intrahippocampal circuits; this raises the question of whether these circuits are segregated or superimposed¹⁴.

In one anatomical framework, functionally distinct hippocampal circuits are segregated along the dorsoventral hippocampal axis. Indeed, early rodent electrophysiological studies indicated dissociable response properties in the dorsal versus ventral hippocampus^{15,16}, and early lesion studies suggested that behaviour was differentially affected by dorsal and ventral hippocampal lesions¹⁷⁻²⁰. These early studies did not, however, distinguish between the location and the size of the lesion. Subsequent work^{21,22}, which did make this distinction, showed that lesions restricted to the dorsal hippocampus, but not similarly sized ventral lesions, impaired spatial learning. It was proposed that the more ventral parts of the hippocampus mediate emotional responses²³, on the basis of more dense ventral than dorsal connectivity with the amygdala^{24,25} and hypothalamic endocrine and autonomic nuclei²⁶, and the selective ventral hippocampal role in the endocrine stress response²⁷. The ensuing view, which has dominated the field ever since, has been that dorsal parts of the hippocampus mediate cognitive functions — particularly spatial memory — whereas ventral portions of the hippocampus are involved in emotional responses^{28,29}.

REVIEWS

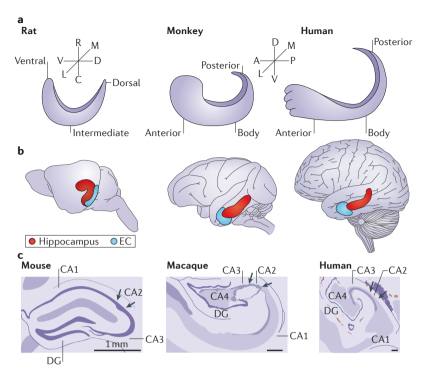


Figure 1 | Cross-species comparison of hippocampal anatomy. a | Schematic illustrations of the orientation of the hippocampal long axis in rats, macaque monkeys and humans. The longitudinal axis is described as ventrodorsal in rodents and as anteroposterior in primates (also referred to as rostrocaudal in non-human primates). There is currently no precise anatomical definition for a dorsal (or posterior) portion of the hippocampus relative to a ventral (or anterior) one, although in general, topologically, the former is positioned close to the retrosplenial cortex and the latter close to the amygdaloid complex. Note that a 90-degree rotation is required for the rat hippocampus to have the same orientation as that of primates. In primates, the anterior extreme is curved rostromedially to form the uncus. $\boldsymbol{b}\,|\,\text{The full long}$ axis of the hippocampus (red) can be seen in brains of rats, macaque monkeys and humans, with the entorhinal cortex (EC) shown in blue. c | Drawings of Nissl cross-sections of mouse, rhesus and human hippocampi. A, anterior; C, caudal; D, dorsal; DG, dentate gyrus; L, lateral; M, medial; P, posterior; R, rostral; V, ventral. Panel a is adapted with permission from REF. 171, Copyright © 1993 Wiley-Liss, Inc., A Wiley Company. Panel c is from REF. 54, Nature Publishing Group.

Episodic memory

Long-term memory for events or episodes that is accessible to conscious recollection.

Semantic memory

Long-term memory for facts that is accessible to conscious recollection.

Place cells

Pyramidal cells that fire in specific locations with spatially restricted firing patterns that are maintained on memory retention trials. This 'dorsal-ventral dichotomy view' was, in part, based on observations that emphasized the segregation of inputs to the hippocampus. However, differences in connectivity with cortical and subcortical structures along the dorsoventral axis of the hippocampus are gradual rather than absolute³⁰, which suggests that functional differences along the long axis may also exhibit a gradient-like organization³¹. Furthermore, recent gene expression data indicate that there are multiple, discretized dorsal-ventral subdivisions along the hippocampal long axis³². Thus, given this potentially more complex hippocampal long-axis functional organization¹⁴, the currently accepted dorsal-ventral dichotomy model requires revision.

In this Review, we first describe anatomical findings in rodents that suggest that there are multiple long-axis functional gradients. We then review evidence from rodent gene expression data indicating that discrete genetic domains are superimposed on this graded long-axis organization. We then discuss — using data from

studies in animals and humans — how these anatomical and genetic patterns may result in patterns of long-axis functional specialization, particularly in terms of spatial processing, emotional responses, action and episodic memory. The evidence for multiple levels of longitudinal functional organization should change our view of the hippocampus and is crucial for understanding the role of the hippocampus in cognition.

Hippocampal long-axis anatomy in rodents

Gradients in hippocampal-cortical connectivity. In terms of cortical input in rodents, a dorsolateral-toventromedial gradient of origin in the entorhinal cortex (EC) corresponds to a dorsoventral axis of termination in the hippocampus^{33–35} (FIG. 2a). This topography is smooth, without abrupt transitions in EC-hippocampus projections. The cortical input to the EC is itself topographically arranged (FIG. 2a), and this mapping is maintained in EC-hippocampus inputs. Using the rat cingulate cortex as an example³⁶, information arising from the infralimbic and prelimbic cortices will, via input to the ventromedial parts of the EC, primarily reach ventral parts of the hippocampus. By contrast, projections from the prelimbic cortex targeting intermediate parts of the EC influence the hippocampus at intermediate dorsoventral levels. The remaining parts of the cingulate cortex — anterior cingulate and retrosplenial cortices - primarily target dorsal and lateral parts of the EC, which subsequently project to dorsal parts of the hippocampus³⁶. The hippocampus thus receives a transition of projections from the cingulate cortex along its long axis: cingulate areas involved in emotional regulation (infralimbic and prelimbic cortices) project to more ventral regions, and cingulate areas involved in spatial processing (the retrosplenial cortex) project to more dorsal regions. Importantly, this transition of projections is continuous rather than discretized. Furthermore, reciprocating projections from the CA1 and subiculum to the EC show a topographical organization similar to that of the EC-hippocampus inputs³⁷.

Gradients in hippocampal-subcortical connectivity.

Hippocampal connectivity with multiple subcortical structures also shows dorsoventral topographical gradients. Taking the topography of the major hippocampal output to the lateral septum (LS)26 as an example, the dorsal half of the hippocampus projects to a very small dorsal part of the LS, whereas progressively more ventral parts of the hippocampus innervate progressively larger parts of the LS more ventrally (FIG. 2b). Adjacent hippocampal areas along the longitudinal axis innervate distinct but overlapping regions of the LS³⁸. Thus, although individual LS neurons receive inputs from a dorsoventral 'patch' of hippocampal pyramidal cells³⁸, the projection on the whole has a topographically graded organization. Crucially, this topographically graded organization is preserved in LS projections to the hypothalamus. This implies that different hippocampal regions along the longitudinal axis topographically map onto different hypothalamic regions involved in behavioural, endocrine and

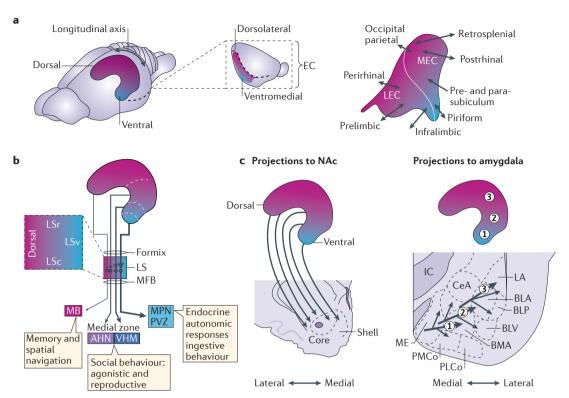


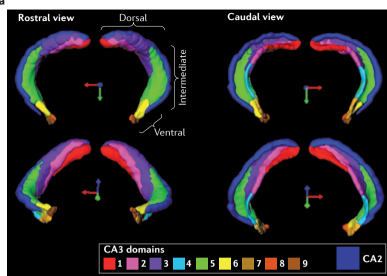
Figure 2 | Extrinsic connectivity gradients. a | The left panel shows a representation of the topographical arrangement of entorhinal-hippocampal reciprocal connections in rodents. A dorsolateral band of the entorhinal cortex (EC) (magenta) is preferentially connected to the dorsal hippocampus. Increasingly more ventral and medial bands of the EC (purple to blue) are connected to increasingly more ventral levels of the hippocampus. The right panel shows an enlarged EC, indicating the topology of its major cortical connectivity. The white line indicates the border between the lateral EC (LEC) and medial EC (MEC). b | The hippocampal output to the lateral septum (LS) and hypothalamus. The LS can be divided into rostral (LSr), caudal (LSc) and ventral (LSv) parts. The most ventral tip of the CA1-subiculum (blue) projects to LSv, which projects to the medial preoptic nucleus (MPN) and hypothalamic periventricular zone (PVZ). More dorsal parts of the CA1-subiculum field project to the LSr, which in turn projects to hypothalamic medial zone nuclei, including the anterior hypothalamic nucleus (AHN) and the ventromedial hypothalamic nucleus (VMH). The dorsal subiculum sends a small projection to the dorsal LS, which is relayed to the mammillary body (MB). The thickness of the arrows indicates the projection density. c | Topographical gradient of projections from the hippocampus to the medial (shell)-to-lateral (core) portions of the nucleus accumbens (NAc) and the medial-to-lateral portions of the amygdala. Note the absence of projections from the dorsal hippocampus and the relative lack of innervation of the central nucleus of the amygdala (CeA). BLA, basolateral amygdala; BLP, posterior basolateral nucleus of the amygdala; BLV, ventral basolateral nucleus of the amygdala; BMA, basomedial nucleus of the amygdala; IC, internal capsule; LA, lateral amygdala; ME, medial nucleus of the amygdala; MFB, medial forebrain bundle; PLCo, posterolateral cortical nucleus of the amygdala; PMCo, posteromedial cortical nucleus of the amygdala. The right panel of part a is adapted with permission from REF. 183, Hindawi. The bottom right panel of part c is adapted with permission from REF. 40, Copyright © 2006 Wiley-Liss, Inc.

autonomic responses associated with specific goaloriented behaviours²⁶ (FIG. 2b). Hippocampal connectivity with the nucleus accumbens (NAc)³⁹ and amygdala⁴⁰ also follows a topographical pattern, with progressively more ventral hippocampal portions projecting to progressively more medial parts of both of these subcortical structures (FIG. 2c).

Interestingly, these topographical gradients seem to arise during embryonic neurogenesis⁴¹. Although neurogenesis occurs simultaneously along the hippocampal dorsoventral axis, the dorsal hippocampus projects to those zones in target structures in which cells were generated earlier, whereas progressively more ventral parts project to zones in which cells were generated later. For example, the dorsal hippocampus projects to a zone in

the LS that contains earlier-formed, medially placed LS cells, whereas the ventral hippocampus — which is geometrically further away from the LS — projects to LS zones containing later-formed, laterally placed cells⁴¹.

The density of neuromodulatory projections to the hippocampus also changes along the long axis (<u>Supplementary information S1 (box)</u>). Whether these changes are gradual, step-like or abrupt has not been studied in detail, but a clear pattern of stronger projections of monoamine systems to more ventral parts of the hippocampus is apparent. Thus, in general, the dorsoventral organization of extrinsic connectivity is one of gradual transitions of topographically organized projections, which does not show a dichotomous segregation into discrete dorsal versus ventral portions.



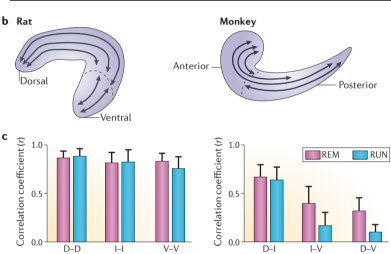


Figure 3 | Discrete transitions in the molecular, anatomical and functional organization of the hippocampal long axis. a | Discrete gene expression domains in CA3 are defined by reciprocal, non-overlapping boundaries. Colour-coded three-dimensional models of nine gene expression-based subdivisions of CA3 are shown in rostral and caudal views at two different orientations (three-dimensional orientation bars: lateral is red; ventral is green; and rostral is blue). Suggested boundaries for collapsing the nine domains into three domains (ventral, intermediate and dorsal) are indicated in the top left three-dimensional model. Note, however, that there are substantially different patterns within each of the dorsal, intermediate and ventral domains, and that these are sharp boundaries in some cases. CA2 is indicated in dark blue. **b** | Extensive versus limited intrinsic connections in the rat hippocampus and monkey hippocampus. In rats, the longitudinal ipsilateral extent of associational fibres from the dentate hilus is shown. In monkeys, projections from CA3 to CA1 and CA3 at the level of the uncus are restricted to the anterior portions of the hippocampus. The boundary between the posterior (dorsal) two-thirds versus anterior (ventral) one-third of the hippocampus is indicated schematically by dashed lines. Note that this line is interrupted in the right panel to indicate that this boundary is less discrete in monkeys than in rodents 58,59. c | Coherence decreases along the longitudinal axis. Theta-power correlations between dorsal (D), intermediate (I) and ventral (V) sites in the CA1 pyramidal layer during running (RUN) and rapid eye movement (REM) sleep. Powerpower correlations are high within the same portions (left) and significantly decrease between ventral versus intermediate and dorsal sites (right). Panel a is based on data from REF. 32. Panel **b** is adapted with permission from REF. 59, Copyright © 2009 Wiley-Liss, Inc. Panel c was published in Neuron, 75, Patel, J., Fujisawa, S., Berényi, A., Royer, S. & Buzsáki, G., Traveling theta waves along the entire septotemporal axis of the hippocampus, 410–417, Copyright Elsevier (2012)⁶¹.

Gene expression along the long axis

The development of an unbiased transcriptional map of the mouse hippocampus, using genome-scale in situ hybridization⁴², has provided detailed molecular evidence for a discretized dorsal-ventral pattern of gene expression^{29,32,43}. Importantly, genetic domains are not defined by the expression of any single gene but, rather, by the combined overlap of many gene expression domains³². Thus, the overlap of the expression of many genes with common expression boundaries gives rise to genetic domains with clearly demarcated borders³². Boundaries between domains can be reciprocal, in that individual genes delineate a given boundary from each side³² (FIG. 3a). Multiple segregated molecular subdomains, each containing a unique complement of expressed genes, have been demonstrated along the long axis. One study showed that there are nine domains within area CA3 (REF. 32), and two other studies showed that the dentate gyrus (DG)²⁹ and area CA1 (REF. 43) are segregated into three major molecular domains: dorsal, intermediate and ventral (with the ventral CA1 domain comprising four subdomains). Importantly, the molecular differentiation along the longitudinal axis is not simply dorsal versus ventral: that is, there is no evidence for a boundary that divides the long axis into two portions. If the nine expression domains in the CA3 can be simplified into dorsal, intermediate and ventral parts, similarly to the CA1 and DG domains29, this could suggest a tripartite model of the long axis. Such a tripartite model has been recently corroborated in a developmental gene expression study in rats44. Nevertheless, the exact number of domains along the long axis, and whether these are hierarchically organized, is currently unknown¹⁴.

The interesting challenge ahead will be to assess whether these patterns of molecular expression translate into specific functional properties along the hippocampal long axis. The expression profiles of genes encoding adhesion molecules and ion channels32,43 may determine intrinsic electrophysiological properties of discrete hippocampal neuronal populations, such as the differences in neuronal excitability45 and synaptic plasticity46,47 that have been detected along the long axis. For example, hyperpolarization-activated cyclic nucleotidegated channel 1 (HCN1) and HCN2, which mediate hyperpolarization-activated currents (I_L) currents, are differentially expressed along the dorsoventral axis48 and are important for a spatial function that is dorsoventrally graded⁴⁹⁻⁵¹. In general, neurotransmitter receptor expression varies across the long axis for the majority of transmitter systems (Supplementary information S2 (table)). Studies combining genetic and anatomical techniques in the rodent brain have begun to reveal that neuronal circuits, both within the hippocampus^{52,53} and between the hippocampus and LS43, share common gene expression patterns, which indicates overlap between anatomical and genetic levels of organization along the long axis. Importantly, however, in contrast to the anatomical homologies between the rodent hippocampus and primate hippocampus described in BOX 1, the recently developed transcriptional atlas of the adult human brain⁵⁴ indicates that there are differences in

Callosal mammals

Mammals with a corpus callosum. In acallosal mammals, such as the opossum, the dorsal portion of the hippocampus extends into the frontal lobe.

gene regulation between the mouse hippocampus and human hippocampus. The molecular organization along the hippocampal long axis in primates, and whether this is similar to that in mice^{32,43}, remains to be examined.

Reconciling molecular and anatomical data

How can the molecular data indicating sharp expression boundaries along the long axis that are common to many genes be reconciled with the anatomical data showing extrinsic connectivity gradients along the long axis? Two points are important in answering this question. First, at

the level of individual genes, there are various long-axis expression patterns, including gradual changes, steplike changes and sharp transitions³². Second, although extrinsic hippocampal connectivity appears to follow a smooth, graded topographical organization, sharp demarcations of intrinsic connectivity along the long axis have also been observed. For example, the two major longitudinal association fibre systems in the hippocampal formation — the longitudinal axon collaterals of CA3 pyramidal cells and the longitudinally oriented axons of DG mossy cells — show extensive axon

Box 1 | Is the rodent ventral-dorsal axis homologous to the primate anterior-posterior axis?

There are obvious macroscopic differences between the rodent hippocampus and the primate hippocampus. Therefore, we consider whether the rodent ventral-dorsal axis is homologous to an anterior-posterior axis in non-human primates and humans (FIG. 1).

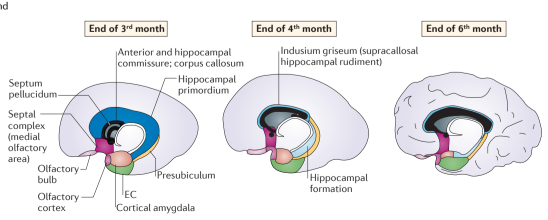
One obvious difference lies in the orientation of the hippocampal long axis in rodents versus humans. This difference probably relates to the fact that in non-primate callosal mammals the major portion of the dorsal hippocampus is tucked under the caudal section of the corpus callosum, whereas this subcallosal flexure diminishes from prosimian to simian species and is practically absent in humans, presumably because of forward growth of the temporal lobe 170. That is, the ventral hippocampus appears to have been 'pulled' downwards and forwards in primates to occupy a position in the anterior medial temporal lobe, thereby changing the long-axis orientation.

A second macroscopic difference is that the rodent hippocampus cross-sectional area is relatively uniform along the long axis, whereas the anterior hippocampus has expanded relative to the posterior hippocampus in primates, particularly in humans¹⁷¹. One speculative phylogenetic account for this involves the entorhinal cortex (EC), which in all mammals has a close topological relationship with the ventral or anterior hippocampus (FIG. 1b). With forward growth of the temporal lobe, the EC moved from its occipital lobe position in lower-order mammals to a rostral location in the primate anterior medial temporal lobe, where it has expanded considerably compared with other components of the uncus¹⁷⁰. Thus, the expansion of the EC and its more anterior position in the temporal lobe in primates may have accompanied the expansion of the anterior hippocampus, such that a greater portion of hippocampal tissue became located anteriorly. This observation poses several currently unanswered questions, such as what is the functional gain or loss of the increased size of the anterior hippocampus, and is this at the expense of the functions of the posterior hippocampus in humans? What would an increased number of anterior cells be useful for? Can the posterior functions be carried out with the small number of cells that, for example, a rodent dorsal hippocampus has?

The rodent hippocampus and primate hippocampus also differ in terms of embryonic development¹⁷². Species that have an evolutionary relationship typically share the early stages of embryonic development but differ in later stages. Indeed, during early embryonic development, the human hippocampus resembles that of the rat, running dorsal to ventral, with the dorsal portion lying above the diencephalon¹⁷³.

At approximately the 14-week stage and coincident with the development of the corpus callosum, the dorsal (supracallosal) hippocampus in humans begins massive involution and remains only as a rudimentary thin band above the corpus callosum (the indusium griseum)^{173,174}. By contrast, the ventral embryological portion develops to form the length of the human $hippocampus ^{173,174}. \ The \ figure \ illustrates \ the \ embryological \ development$ of the human hippocampus. Note massive involution of the dorsal (supracallosal) hippocampal primordium. Involution of the supracallosal part of the hippocampus also occurs in rodents, although the indusium griseum is far less conspicuous than in humans. This leaves open a possibility that the extent of involution of the dorsal embryological hippocampal portion differs between species, and one may therefore wonder whether a homologue of the rat dorsal hippocampus is present in the human brain or whether the human posterior hippocampus instead corresponds, phylogenetically, to rodent intermediate hippocampal portions.

Notwithstanding these differences, a cross-species comparison of anatomical connectivity provides evidence that the primate hippocampal long axis may be homologous to that of the rat. Indeed, output connectivity of the primate hippocampus with subcortical areas - including the nucleus accumbens¹⁷⁵ — follows a graded topography that is similar to that in rodents¹⁷⁶ (but note longitudinally restricted versus distributed hippocampus-amygdala projections in primates and rodents, respectively 108,177). Input connectivity from the EC to the dentate gyrus also follows a graded mapping that is analogous to that in rats^{34,35}, with an anteromedial–posterolateral EC axis corresponding to an anterior-posterior dentate gyrus termination¹⁷⁸⁻¹⁸⁰. For example, the pattern of connectivity between the cingulate cortex and hippocampus in primates is similar to that in rats, in the sense that the anterior hippocampus is more strongly connected with anterior regions and the medial frontal cortex, whereas the posterior hippocampus is more strongly connected with the posterior cingulate (including the retrosplenial cortex)^{181,182}. Figure is adapted with permission from REF. 173, Copyright © 1951 The Wistar Institute of Anatomy and Biology.



divergence within the dorsal two-thirds and within the ventral one-third of the rat hippocampus, but few fibres cross between these subdivisions^{30,55–58} (FIG. 3b). That is, the division between these areas in terms of intrinsic connectivity is relatively abrupt. Similarly, in monkeys, there are extensive versus limited interconnections in the posterior two-thirds versus the anterior one-third of the hippocampus, respectively⁵⁹ (FIG. 3b), although the boundary, in terms of intrinsic connectivity, between these hippocampal portions is less clearly demarcated than that in rodents.

In humans, discrete changes in molecular or anatomical organization along the hippocampal long axis have yet to be examined. However, one study showed abrupt transitions in electrophysiological properties along this axis in humans⁶⁰. Specifically, measurements at adjacent contacts (on multicontact depth electrodes) showed an abrupt decrease in coherence at approximately the transition between the anterior one-third and posterior two-thirds of the hippocampus⁶⁰. Similarly, in rats, theta-wave coherence is relatively high between dorsal and intermediate sites but substantially lower between dorsal and ventral sites⁶¹ (FIG. 3c). It will be important to determine whether this decrease in coherence coincides with the locus on the long axis at which intrinsic connectivity shows the partition described above58,59.

Together, the data suggest that there are different types of longitudinal organization — both gradual gradients and discrete, sharply demarcated domains — that seem to be superimposed at both the anatomical and mRNA levels (FIG. 4). Next, we review how these various patterns of long-axis organization may be expressed functionally.

Functional organization of the long axis

Spatial processing in rodents. The representation of location by hippocampal place cells is non-topographic³. A local cluster of place cells in the rodent dorsal hippocampus can cover most of a spatial environment⁶². Initial evidence suggested that relatively small segments of the dorsal hippocampus (a quarter or less of total hippocampal volume) are sufficient to encode spatial memory²². However, if the original spatial encoding occurs in the context of a normal hippocampus, retrieval requires the entire dorsal two-thirds of the hippocampus (that is, including parts of the ventral hippocampus), suggesting a more distributed — or graded — mode of action in a normal hippocampus during spatial learning⁶³. Thus, these lesion studies suggested the possibility that normal rats engage an extensive hippocampal network — located in the dorsal 70% of the hippocampus — during encoding and retrieval of spatial memory, whereas more limited networks within this dorsal region can be used for encoding in rats with partial hippocampal lesions⁶³.

Does the ventral hippocampus have a role in spatial processing? Initial data indicated that the proportion of ventral hippocampal cells that express place fields was markedly lower than that of dorsal hippocampal cells expressing place fields and that ventral place cells have lower spatial selectivity⁶⁴. More recent data demonstrate that the relative size of place fields in area CA3 increases almost linearly with position from the dorsal hippocampal pole (where place fields are ~1 metre) to the ventral pole (where place-field size approaches 10 metres)³¹ (FIG. 5a). This finding not only highlights a role for the ventral hippocampus in the processing of large-scale spatial information, it also implies that there is a functional gradient along the hippocampal longitudinal axis

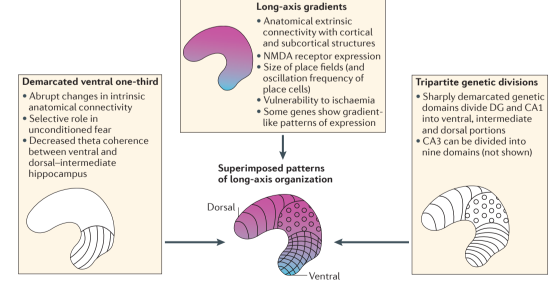


Figure 4 | **Schematic of superimposed patterns of long-axis organization.** Behavioural lesion, electrophysiological recording and intrinsic-connectivity studies have suggested a functional distinction between the ventral one-third of the hippocampus versus the dorsal two-thirds. Other studies have revealed gradual changes along the hippocampus in terms of extrinsic connectivity, receptor expression and place field size, whereas recent gene expression studies indicate that there are three sharply demarcated portions of the hippocampus. Superimposing these three organizational patterns results in a new model of functional organization along the hippocampal long axis.

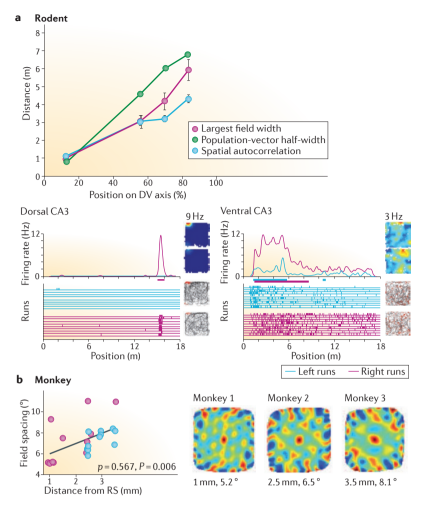


Figure 5 | Gradients for space in the medial temporal lobe in rodents and monkeys. a | The graph in the top panel shows the monotonic relationship between spatial scale and position along the dorsoventral (DV) hippocampal axis. Spatial scale is expressed as the half-width of the correlated band of the population vector, the average width of the largest place field of individual cells and the estimated field width. The lower panels show place fields of example pyramidal cells in the dorsal and ventral CA3 of rats during running on an 18 m track. Smoothed spike-density function indicates that the firing rate is a function of position. The horizontal bar indicates the estimated place field. Below the graphs are raster plots showing the density of spikes on individual laps. Each vertical tic indicates one spike, and each horizontal line shows one lap. To the right of each panel are rate maps and trajectories (top pairs and bottom pairs, respectively), with individual spikes from repeated trials in two-dimensional enclosures (1 m by 1 m). Rate maps are colour-coded, with red as maximum and blue as 0 Hz and the peak rate indicated at the top. Trajectories are shown as black traces and the positions of individual spikes are shown as red dots on top of the trajectory. **b** | The graph on the left shows that, in monkeys, grid-cell spacing increases with distance from the rhinal sulcus (RS). Blue and magenta circles identify the grid cells from each of two monkeys. Note that these grid cells are from head-fixed monkeys and that firing is defined by view position rather than by the position in the room. Shown on the right are autocorrelations for representative grid cells recorded at different locations medial to the RS in three monkeys. The distance from the RS (mm) and field spacing (degrees) are indicated below. Part a is from Kjelstrup, K. B. et al. Finite scale of spatial representation in the hippocampus. Science 321, 140-143 (2008). Reprinted with permission from AAAS³¹. Part **b** is from REF. 95, Nature Publishing Group.

(as opposed to a dorsal-ventral dichotomy). That is, the ventral hippocampus may subserve similar spatial processing functions as the dorsal hippocampus but at a larger spatial scale. Such a representation of space at multiple scales has computational advantages in the

sense that a gradient for space accommodates both spatial resolution and spatial contiguity. A further recent observation regarding place cells is that when rodents locomote in a constant location, the firing fields of place cells are defined by time^{65,66} or a moment within a sequence⁶⁷ instead of by location, posing the interesting question of whether such 'time fields' expand from dorsal to ventral regions⁶⁸ in a similar manner to place fields.

Place cells participate in multiple, independent spatial representations^{69,70}, whereas the more recently discovered entorhinal grid cells71 encode a universal metric of the spatial map. Grid-cell firing locations define a periodic triangular or hexagonal array that represents the animal's entire environment⁷¹, and they are anchored to external cues and maintained when cues are removed and with ongoing changes in the animal's speed and direction⁷¹. The spatial selectivity of place cells may be linked to inputs from grid cells⁷²⁻⁷⁶. Crucially, the increase in the size of place fields along the hippocampal dorsoventral axis^{31,64,77} is mirrored by an increase in the spacing between grid-cell firing locations from the dorsomedial to the ventrolateral medial EC71,78,79. In contrast to the gradual dorsoventral increase in placefield size, the observed spatial gradient in the medial EC grid size shows discrete, step-like increases80. However, although the scale of place cells increases gradually from dorsal to ventral on average, this does not rule out the existence of discrete transitions such as those observed in the EC80. If it is found that the increase in place-field scale is not continuous, it will be important to determine whether this scale changes abruptly with transitions between genetic domains. Assuming for now that the place-field scale is indeed continuous, inputs from different medial EC functional modules could, in theory, be combined76,81 to give rise to the observed longitudinal spatial gradient75. Specifically, EC modules of increasing spatial scale show considerable anatomical overlap in the dorsoventral axis of the EC80, suggesting that there may be overlap of module inputs to the hippocampus, even if inputs come from the same dorsoventral EC level. Future studies will determine whether grid-cell modules in the medial EC distribute evenly across the hippocampus, whether they connect to modules in the hippocampus or whether there is complete convergence. The fact that rescaling of grid fields in response to environment compression is observed in modules with large, but not small, gridscales⁸⁰ raises the question of whether similar dissociations exist between large and small place fields in the ventral and dorsal hippocampus, respectively. Recordings from the dorsal hippocampus indicate that compression can take place at relatively small scales82, which is evidence for independence between grid and place cells. However, a systematic comparison of rescaling in dorsal and ventral place fields has not been conducted. With respect to the organization of the hippocampal long axis more generally, the gradient in place-field size illustrates that, despite numerous molecular and anatomical domains having distinct boundaries, a combination of hippocampal afferent signals may engender gradually changing functional properties (FIG. 4).

Spatial processing in primates. Does the dependence of spatial processing on dorsal portions of the hippocampus in rodents extrapolate to posterior portions of the hippocampus in primates? The majority of primate data pertaining to functional long-axis organization come from human structural and functional MRI (fMRI) studies. It should be kept in mind that technical factors may differentially influence fMRI and voxel-based volumetric measures for anterior versus posterior portions of the human hippocampus. The susceptibility artefact and signal drop-out of fMRI may affect the anterior medial temporal lobe more than the posterior medial temporal lobe83 (although protocols exist to correct this84). In addition, the cross-sectional area of the posterior hippocampus is approximately 50% less than that of the larger anterior hippocampal head, such that the activated cluster size and degree of post-acquisition spatial smoothing may differentially influence statistical effects along the long axis.

Despite these potential limitations, MRI studies in humans have demonstrated a relationship between activation^{7,85,86} and structural change⁸⁷ in the posterior hippocampus with navigation, a finding that is broadly in keeping with the dependence of spatial function on dorsal portions of the hippocampus in rodents. However, neuroimaging results are typically reported as anatomically focal effects that exceed a particular statistical threshold, and simply demonstrating that an effect is located at a specific long-axis locus does not exclude the possibility that there may be an effect just below that statistical threshold at another locus. Although most studies report responses in either the anterior or posterior hippocampus, some studies demonstrate functional double dissociations between long-axis loci, and these are particularly informative^{6,88}. Thus, in further support of a posterior hippocampal specialization for space processing, one reported double dissociation is that accurate way-finding activated the posterior, but not anterior, hippocampus⁸⁵, whereas activity in the anterior, but not posterior, hippocampus correlated with the formation of a survey representation of a new virtual-reality environment85 (see also REFS 89,90). A recent fMRI study91 reported a long-axis dissociation in terms of spatial size and complexity. Participants navigated through three virtual mazes (small with 6 corridors, large with 6 corridors and large with 14 corridors) and then, during scanning, were presented images of landmarks from these mazes and asked to retrieve to which maze they pertained. Anterior hippocampal activation scaled with the number of corridors (complexity), whereas posterior responses were larger for larger mazes. The interpretation of these data is limited, however, by the fact that for all mazes, participants navigated almost exclusively along border paths, and there was no measure of how much spatial retrieval was evoked by correct landmark retrieval (see REF. 92).

Electrophysiological evidence for posterior hippocampal involvement in spatial processing in primates is limited. One non-human primate study using a spatial delayed matching-to-sample task demonstrated greater activity during the delay period in the posterior

hippocampus than in the anterior hippocampus⁹³. Although intracranial recordings in humans have provided evidence for place-cell-like responses during navigation⁹⁴, the relative distribution of these cells along the long axis, and how their responses vary as a function of environment size, has yet to be determined. However, recent electrophysiology data from non-human primates reveal grid-like cell properties in the posterior EC. The study showed that spatial scale varied as a function of distance from the rhinal sulcus⁹⁵ (which is equivalent to the dorsomedial-to-ventrolateral axis in the rodent medial EC), suggesting that spatial scale may also vary along the primate hippocampal long axis (FIG. 5b).

How does the spatial scale representation observed in rodents increase or change across species? Developments in human fMRI, such as hippocampal MRI unfolding% and high-resolution fMRI techniques^{97,98}, combined with within-scanner virtual-reality applications^{85,86}, may provide the technical advances that are required to confirm whether there is a linear representation of spatial scale along the human hippocampal long axis. It will be particularly interesting to assess whether humans show the same spatial precision as that expressed in the rat dorsal hippocampus (<1 metre at the dorsal pole³¹) and, conversely, whether spatial scale extends beyond the 10 metres expressed in the rat ventral hippocampus³¹, given the much larger home range of humans versus rats. By contrast, no species may need grids larger than a few metres because grid maps are likely to be local and fragmented in all realistic environments99. An important point to note, however, is that humans are obviously not locomoting during fMRI scanning, and this might influence the scale of the place fields during scanning, similar to what has been observed in rodents locomoting by train instead of walking themselves¹⁰⁰. Studies in monkeys have also been limited for practical reasons: so far, these have involved head fixation95. As a result, the grid-like cells in monkeys95 observed in these studies differ from rodent grid cells⁷¹ in that they follow eye position rather than the animal's movement in space. Thus, future studies could carry out recordings in freely moving monkeys to test whether spatial scale is comparable to that observed in rodents.

Emotion. Anatomical evidence demonstrates that the reciprocal connectivity between the amygdala on the one hand and the CA1 and the subiculum on the other hand is largely confined to the ventral two-thirds $^{40,101-103}$. This is particularly striking in a study that reported that only the dorsal-most portion of the hippocampus does not innervate the amygdala⁴⁰. This connectivity is topographically organized along the longitudinal hippocampal axis so that the ventral-to-dorsal axis of origin of the projection in the CA1 and the subiculum is associated with a medial-to-lateral axis of termination in the amygdala^{40,103} (FIG. 2c). In view of the specific roles for individual amygdala nuclei, this pattern of connectivity could explain the emotion-related functions of hippocampal regions along the dorsoventral axis. For example, the basolateral amygdala, which has a crucial role in fear learning 104, receives inputs from a considerable portion of the dorsoventral

Susceptibility artefact

The different magnetic susceptibility of air and tissue cause inhomogeneities in the magnetic resonance scanner's static magnetic field at the air—tissue boundaries. These inhomogeneities result in geometrical distortion and reduced sensitivity of functional images, particularly in the orbitofrontal cortex and anterior medial temporal lobe.

Hippocampal MRI unfolding

The application of cortical unfolding techniques to high-resolution magnetic resonance images of the hippocampus. Structural images are segmented and the grey matter surface is extracted and stretched until it is a two-dimensional, flat surface.

long axis^{40,103} (FIG. 2c). This may explain the inconsistent findings in rodent fear-conditioning studies in which either the dorsal or ventral hippocampus is lesioned or inactivated^{29,105-107} (some studies find effects of dorsal not ventral lesions, and vice versa). That is, the effects on conditioned fear may be a function of the locus of the hippocampal lesion with respect to the ventrodorsal hippocampus-to-mediolateral amygdala connection topography. It should be noted that the origin of the homologous hippocampus-amygdala topographical projections in primates is more restricted in the sense that amygdala-projecting neurons are focally restricted to the most anterior (uncal) CA1 and prosubiculum¹⁰⁸. This may explain why fMRI activations associated with emotional memory in humans are primarily in anterior regions^{109,110} (but see REF. 111).

In contrast to evidence for the involvement of both the dorsal hippocampus and ventral hippocampus in conditioned fear, there is growing evidence that the ventral hippocampus, but not the dorsal hippocampus, plays a part in mediating unconditioned fear behaviour^{28,112-114}. An initial study¹¹² demonstrated that ventral, but not dorsal, hippocampal lesions reduce defensive fear responses during exposure to the elevated plus maze (an unconditioned threatening environment). Another early study114 demonstrated that the effects of ventral hippocampal lesions on unconditioned or ethologically based tests of fear extend to non-spatial tasks, such as latency to eat novel foodstuffs, and the degree of social interaction in a familiar environment²⁸. In the initial study112, the fact that selective amygdala lesions did not reduce defensive responses suggests that the ventral hippocampus may influence unconditioned fear expression independently of the amygdala, namely through direct ventral hippocampal projections to downstream neuroendocrine and behavioural control systems in the hypothalamus²⁶ (FIG. 2b). With respect to longitudinal organization, the critical observation is that lesion data for unconditioned emotional responses show an anatomically marked ventral-dorsal hippocampal distinction: lesions of the dorsal two-thirds of the hippocampus did not affect fear expression, whereas small lesions in the ventral one-third did¹¹². The hippocampal role in unconditioned emotional responses may thus be segregated to a ventral functional portion.

Given the model of longitudinal organization we propose, in which demarcated domains are superimposed on functional gradients (FIG. 4), it is particularly interesting to consider the role of this ventral portion in unconditioned emotional responses in the context of the hippocampal gradient for space processing³¹. In non-spatial tasks, such as tone-shock fear conditioning, place-cell responses to non-spatial stimuli, such as the auditory tone that predicts the shock¹¹⁵, are only observed when the animal is in that cell's place field. Thus, having larger place fields in the ventral hippocampal portion, which is strongly linked to defensive behaviour-related circuitry of the hypothalamus^{26,102}, may be evolutionarily advantageous. That is, it is obviously advantageous to detect approaching danger from as far away as possible, and distant danger may require fewer computational steps within these larger fields of the ventral hippocampus. However, it is not yet known whether this 'emotional' portion of the hippocampus has a dorsal border that is defined by molecular transitions^{29,32,43} or abrupt changes in longitudinal association fibre anatomy⁵⁸, or is anatomically circumscribed to a particular level of topographical ventral hippocampus–LS–hypothalamus connections²⁶.

Action and motivation. Although no gross, permanent motor deficits arise after bilateral hippocampal lesions, an association between hippocampal activity and motor acts has long been described^{3,116}. In non-human primates, movement-related responses have been reported in anterior, but not middle or posterior, portions of the hippocampus⁹³. Although human intracranial recordings117 and fMRI118 studies have demonstrated various motor-evoked hippocampal responses, differences in these responses along the human long axis have yet to be examined. In rodents, ventral, but not dorsal, hippocampal stimulation increases locomotion 119,120 by engaging the NAc and mesolimbic dopamine system¹²¹⁻¹²³, whereas inhibiting the ventral hippocampus decreases locomotion¹⁰⁵. This relationship with the NAc is also relevant to the observation that reward- or goaldirected functions localize to ventral parts of the rodent hippocampus^{124,125} and to the anterior human hippocampus¹²⁶, given that the ventral striatum — in particular the NAc — is considered to be the 'limbic-motor interface' at which motivation- and emotion-related processing gains access to the motor system^{127,128}.

The rodent studies discussed above¹¹⁹⁻¹²³ examined dorsal versus ventral functional dissociations, but the anatomical connectivity between the hippocampus and NAc in fact shows a graded topography³⁹. In view of this topography, it was suggested that the intermediate hippocampus, lying between the dorsal and ventral poles, is the site where accurate place encoding (which is strongest in the dorsal hippocampus) 'meets' connections (which are strongest in the ventral hippocampus) with behavioural control areas, including the prefrontal cortex and NAc129 (FIG. 2). Selective lesions along the long axis have demonstrated that the intermediate hippocampus is critical for rapid place learning and the subsequent use of this encoded information to guide navigational performance¹²⁹. However, it should be noted (in view of the anatomical orientation of the intrinsic hippocampal circuitry) that after selective lesioning, the remaining dorsal, intermediate and ventral portions of the hippocampus will differ in their composition of subfields. Thus, intermediate tissue blocks are more likely to comprise complete trisynaptic circuits than blocks from the poles, and this could bias the interpretation of such studies in terms of the functional relevance of the intermediate hippocampus.

Episodic memory. An early suggestion¹³⁰, based on human positron emission tomography (PET) data, proposed a dissociation between anterior and posterior portions of the hippocampus for episodic-memory encoding and retrieval, respectively (but

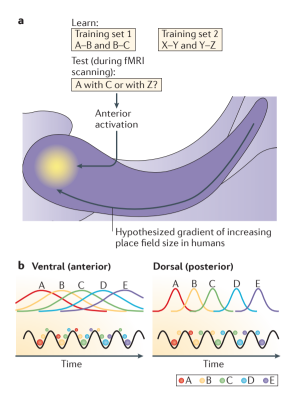


Figure 6 | Forming non-sequential, higher-order connections in the human hippocampus. A schematic to illustrate a putative mechanism by which the human anterior hippocampus is able to form non-sequential connections that enable flexible cognitive processes such as transitive inference. a | Functional MRI (fMRI) studies demonstrate increased anterior hippocampal responses when subjects infer the correct transitive inference (for example, A-C is correct if previous pairings were A-B and B-C) relative to simple recognition of previously learned pairs of non-overlapping visual stimuli. **b** | Interleaved neuronal sequences in the ventral (anterior) and dorsal (posterior) hippocampus. The coloured Gaussian curves represent place fields of five cell assemblies in the ventral (anterior) hippocampus and five cell assemblies in the dorsal (posterior) hippocampus. Together, the place fields could pertain to locations A-E or (speculatively) a sequence of items A-E. Note the longer 'tails' of the fields in the ventral (anterior) hippocampus. Shown below the place fields are circles representing spiking activity from each cell assembly that represents an item in the sequence A-E. The spiking activity gradually precesses from the end to the beginning of the theta cycle (the black oscillating line), and the size of the circles indicates the firing rates of the hypothesized assemblies. Each item is defined by the most active cell assembly that fires at the trough of the theta cycle (for example, C is defined by the assembly depicted by the green place field) and is embedded in the temporal context of previous and subsequent items. Portions of the sequence A-E are replicated repeatedly within individual theta cycles. Note that longer sequences are accommodated ventrally (anteriorly). The formation of assembly sequences within theta cycles could reflect a strengthening of connections not only between adjacent items (for example, C-D) but also between nonadjacent items (for example, A-E and B-D), thereby enabling transitive inference to be made. Part a is based on data from REFS 150, 151. Part b as discussed in REFS 5,157.

see REF. 131). Furthermore, anterior hippocampal responses to novel (versus familiar) stimuli have been frequently reported^{6,132-134} (but see REF. 135) and some studies showed a double dissociation between anterior responses to novelty and posterior responses to previously encountered stimuli^{6,132}. Given that novelty and familiarity detection may be components of memory encoding and retrieval processes, respectively 6,132,136, these data could be taken as support for a dissociation between encoding and retrieval within the hippocampus¹³⁰. However, a caveat to these proposed dissociations is that single-unit data and neuronal-network models indicate that it is extremely unlikely that different hippocampal cells — that is, anterior versus posterior cells — are involved in encoding versus retrieval of a particular memory. This is because a memory is recalled by reactivating the very same neuronal network that was formed during the encoding of the event^{92,137–139}, so that encoding and retrieval occur in parallel, possibly on alternating theta cycles 139-141.

One recent suggestion¹⁴² — based on an extrapolation of the ventral–dorsal increasing resolution gradient in the rodent representation of topographical space — is that, in humans, episodic memories follow a similar gradient in terms of level of detail: that is, the degree of context specificity and/or richness in detail with which that memory can be retrieved. Indeed, it has been observed that retrieval of detailed spatial¹⁴³ or autobiographical¹⁴⁴ memory engages the posterior hippocampus, whereas the anterior hippocampus may be more involved in coarse, 'gist-like' memory¹⁴². A demonstration that this organization follows a gradient-like pattern akin to place representation has yet to be provided, and a clear challenge will be how to define a metric by which to quantify the richness or detail of episodic memories.

Forming non-sequential, higher-order connections. A highly consistent observation in neuroimaging studies of human memory is that tasks requiring semantic processing engage the anterior hippocampus^{131,145,146}. There is evidence of double dissociation between semantic processing in the anterior hippocampus and non-semantic processing in the posterior hippocampus^{145,147} (the term 'relational' memory 148 has been used for the former, but we use semantic memory here, given that all memory could be viewed as relational). One example of semantic processing that requires flexible expression of memory is transitive inference¹⁴⁹. Human studies showing that transitive inference activates the anterior hippocampus^{150,151} (FIG. 6a) are underpinned by an earlier study showing that hippocampal lesions impair transitive inference in rodents¹⁴⁹. This led to the suggestion that the hippocampus is critical for the linking of episodic memories into semantic networks in order to extract the common features — spatial and non-spatial — among related memories and to mediate flexible memory expression and inferential reasoning¹⁵². Although initial lesion data linking the hippocampus to transitive inference involved the entire dorsoventral axis149, a recent electrophysiological study reported that neurons in the ventral CA3 possess the response characteristics that are required to enable flexible encoding of

Transitive inference
If A is paired with B, and B
paired with C, the transitive
inference is A with C.

Theta rhythm

A prominent 4-10 Hz oscillation in the hippocampal local field potential. It is studied mostly in rodents but is also present in humans.

Phase precession

The phenomenon that when a rat first enters the field of firing of a place cell, spiking occurs at late phases but shifts to earlier theta phases as the rat moves through the place field.

Adult neurogenesis

The production of new neurons within the brain of an adult animal. Adult neurogenesis is primarily confined to the subventricular zone and the subgranular zone of the dentate gyrus.

Ischaemia

A restriction in blood supply. which leads to lack of oxygen delivery

memories that span different contexts¹⁵³. Whereas neuron ensembles in the dorsal CA3 rapidly associated the identity of specific objects with locations, successively more ventral neurons were reported to increasingly generalize over object-sampling events involving specific objects and locations within a spatial context, while still distinguishing between different spatial contexts¹⁵³.

What response properties of ventral hippocampal neurons might facilitate the formation of higher-order memory representations? One possible mechanism emerges from the relationship between place-cell oscillating frequency and place-field size¹⁵⁴. Every place cell oscillates faster than the population theta rhythm, which brings about a frequency interference pattern known as phase precession ¹⁵⁵. Phase precession enables a compressed representation of temporal structure to be expressed within single theta cycles (the compression dynamic¹⁵⁶). Given the size of place fields, several place cells are active together in each theta cycle, such that the compression dynamic potentially enables not only adjacent but also more distant neuronal assemblies to be linked, as long as they consistently co-occur in the same theta cycles. The oscillation frequency of place cells decreases along the dorsoventral axis, whereas the size of place fields increases^{31,77,124}. Thus, larger place-field size ventrally theoretically provides more opportunities for neurons with distant place fields (that is, in the ventral hippocampus) to fire together in the same theta cycle than in the dorsal hippocampus^{5,157}. As such, the ventral hippocampal portion may be specially suited for the formation of non-sequential or higher-order links between memory representations that could provide the flexibility needed for efficient navigation and detour planning^{5,157}. Although this suggestion is derived from

studies on spatial processing, it could be extrapolated to semantic function: if locations are assumed to be analogous to items, and we assume that dorsal-ventral differences in place-cell properties extrapolate to the human anteroposterior axis, a larger field size anteriorly provides a potential explanation for the anterior locus of semantic processing responses in the human hippocampus (FIG. 6b). Semantic memory involves considerably more than just the linking of remote locations or time points, but this mechanism for creating higher-order memory representations potentially underpins aspects of semantic memory formation.

Clinical implications

We propose a model of hippocampal functional organization that superimposes long-axis gradients and discrete functional domains (FIG. 4). Can we use this model of longitudinal organization to make specific predictions about the clinical manifestations of hippocampal damage along the long axis in humans? Hippocampal structural abnormalities are observed in a wide range of diseases¹⁵⁸. With developments in human hippocampal volumetric techniques159 and the application of functional imaging to patient populations, evidence is emerging for anterior-posterior differences in the relative severity of hippocampal structural and functional changes in various psychiatric and neurological conditions¹⁶⁰ (although the caveats in interpreting long-axis differences described earlier also apply here). For a number of these conditions, preclinical animal models have considerable predictive value regarding the relative severity of anterior versus posterior pathology observed in patients (TABLE 1). In addition, the locus of pathology on the long axis is associated with specific cognitive

Condition	Abnormality along hippocampal long axis	
	Animal	Human
Medial temporal lobe epilepsy	Greater spontaneous epileptiform bursting in the ventral hippocampus than in the dorsal hippocampus 15,184	 Chronic intracranial recordings in patients indicate that seizure initiation is more frequent in the anterior hippocampus than in the posterior hippocampus ¹⁸⁵ Neuronal loss is greater in the anterior hippocampus than in the posterior hippocampus ¹⁸⁶⁻¹⁸⁸ (expressed as an anterior–posterior gradient ¹⁸⁶)
Depression	Behavioural effects of chronic antidepressant treatment are critically dependent on adult neurogenesis in the hippocampus ¹⁸⁹ , and this has been suggested to occur specifically in the ventral hippocampus ¹⁹⁰	Post-mortem studies on patients with major depressive disorder show that antidepressants increase neurogenesis in the anterior dentate gyrus ¹⁹¹
Schizophrenia	 Lesioning of the ventral hippocampus is used to model several features of schizophrenia¹⁹² Schizophrenia-related biomarkers are present in the ventral hippocampus at birth⁴⁴ 	Increasingly thought that the primary pathology is in the anterior hippocampus ¹⁶⁰ , but there is also considerable evidence for abnormalities in the posterior hippocampus (for example, see REFS 193–194)
Ischaemia	 Ventral-to-dorsal increase in hippocampal vulnerability to ischaemia¹⁹⁵ May be related to an increasing gradient for NMDA receptor expression from ventral to dorsal in area CA1 (REF. 196), as NMDA receptor activation has been proposed to have a role in hypoxic excitotoxicity¹⁹⁷ 	Posterior hippocampus volume is decreased in patients who have had cardiac arrest with successful subsequent resuscitation ¹⁹⁹ (but note previous reports of cardiac arrest-induced ischaemia affecting the entire hippocampal long axis ²⁰⁰)

Table 1 | Preclinical animal studies provide insights into the locus of hippocampal damage in different patient populations

• Cerebral blood flow is greater in the ventral hippocampus than in the dorsal hippocampus during reperfusion following ischaemia, which may contribute to dorsal hippocampus damage¹⁹⁸

impairments (for example, schizophrenia is associated with anterior hippocampal pathology and with impaired transitive inference^{161,162}) as well as with clinical manifestations of particular diseases. For example, in view of the greater connectivity between the ventral (anterior) hippocampus and endocrine hypothalamic nuclei²⁶, impaired hormonal regulation by the hypothalamus (such as hyponataraemic polydypsia reported in patients with schizophrenia who have decreased anterior hippocampal volume¹⁶³⁻¹⁶⁵) may be a common finding in patients with anterior hippocampal damage — this is something that has been relatively under-investigated in medial temporal lobe epilepsy^{166–168}. Furthermore, given the role of the ventral hippocampus¹¹² — and the ventral DG in particular 169 — in models of innate anxiety, this region could prove to be an important future target for a range of neurotic disorders. Last, assuming that genetic subdomains are found in the human hippocampus, one important future challenge for clinical research will be to determine whether these subdomains can be characterized non-invasively with current MRI techniques and whether the genetic composition of these subdomains can be related to specific pathologies.

Conclusions and future directions

Two patterns of functional organization appear to be superimposed on the hippocampal long axis: gradual and discrete transitions. At present, this framework can accommodate some of the multiple, and disparate, functions that have been ascribed to the hippocampus. However, for future studies to disambiguate the relative contributions of different genetic domains and different levels along functional gradients to a given behaviour, a novel approach with high anatomical precision is required. The huge advance in understanding hippocampal molecular anatomy enables this information to be used to allow highly specific targeted genetic manipulation of a particular region of the hippocampus (for example, the ventral one-third of a specific CA subfield). A range of transgenic tools can be applied to stimulate or block activity in that region with tight temporal control relative to an experimental paradigm. Thus, this experimental approach provides an avenue towards functional manipulation that could determine whether a specific domain of the hippocampus is necessary or sufficient to subserve a particular behaviour and the mechanism through which this is achieved.

- Scoville, W. B. & Milner, B. Loss of recent memory after bilateral hippocampal lesions. *J. Neurol. Neurosurg. Psychiatry* 20, 11–21 (1957).
- Squire, L. R. Memory and the hippocampus: a synthesis from findings with rats, monkeys, and humans. *Psychol. Rev.* 99, 195–231 (1992).
- O'Keefe, J. & Nadel, L. The Hippocampus as a Cognitive Map (Clarendon, 1978).
- Morris, R. G., Garrud, P., Rawlins, J. N. & O'Keefe, J. Place navigation impaired in rats with hippocampal lesions. *Nature* 297, 681–683 (1982).
- Buzsáki, G. & Moser, E. I. Memory, navigation and theta rhythm in the hippocampal-entorhinal system. *Nature Neurosci.* 16, 130–138 (2013).
- Strange, B. A., Fletcher, P. C., Henson, R. N. A., Friston, K. J. & Dolan, R. J. Segregating the functions of human hippocampus. *Proc. Natl Acad. Sci. USA* 96, 4034–4039 (1999).
 - The first experimental demonstration, using fMRI, of a double dissociation in anterior versus posterior human hippocampal responses.
- Maguire, E. A. et al. Knowing where and getting there: a human navigation network. Science 280, 921–924 (1998)
- Corkin, S. Acquisition of motor skill after bilateral medial temporal-lobe excision. *Neuropsychologia* 6, 255–265 (1968)
- O'Keefe, J. & Dostrovsky, J. The hippocampus as a spatial map. Preliminary evidence from unit activity in the freely-moving rat. *Brain Res.* 34, 171–175 (1971).
- O'Keefe, J. & Speakman, A. Single unit activity in the rat hippocampus during a spatial memory task. Exp. Brain Res. 68, 1–27 (1987).
- Cave, C. B. & Squire, L. R. Equivalent impairment of spatial and nonspatial memory following damage to the human hippocampus. *Hippocampus* 1, 329–340 (1991).
- 12. Nadel, L. The hippocampus and space revisited. *Hippocampus* 1, 221–229 (1991).
- Kim, J. J. & Fanselow, M. S. Modality-specific retrograde amnesia of fear. Science 256, 675–677 (1992)
- Moser, M. B. & Moser, E. I. Functional differentiation in the hippocampus. *Hippocampus* 8, 608–619 (1998)
- Elul, R. Regional differences in the hippocampus of the cat. I. Specific discharge patterns of the dorsal and ventral hippocampus and their role in generalized seizures. Electroencephalogr. Clin. Neurophysiol. 16, 470–488 (1964).
- Racine, R., Rose, P. A. & Burnham, W. M.
 Afterdischarge thresholds and kindling rates in dorsal

- and ventral hippocampus and dentate gyrus. *Can. J. Neurol. Sci.* **4**, 273–278 (1977).
- Hughes, K. R. Dorsal and ventral hippocampus lesions and maze learning: influence of preoperative environment. *Can. J. Psychol.* 19, 325–332 (1965).
- Nadel, L. Dorsal and ventral hippocampal lesions and behavior. *Physiol. Behav.* 3, 891–900 (1968).
- Stevens, R. & Cowey, A. Effects of dorsal and ventral hippocampal lesions on spontaneous alternation, learned alternation and probability learning in rats. Brain Res. 52. 203–224 (1973).
- Sinnamon, H., Freniere, S. & Kootz, J. Rat hippocampus and memory for places of changing significance. J. Comp. Physiol. Psychol. 92, 142–155 (1978).
- 21. Moser, E., Moser, M. B. & Andersen, P. Spatial learning impairment parallels the magnitude of dorsal hippocampal lesions, but is hardly present following ventral lesions. *J. Neurosci.* 13, 3916–3925 (1993). This study, along with that reported in reference 22, made an important advance in localizing rodent spatial function predominantly to the dorsal hippocampal. By controlling the size and locus of hippocampal lesions, this work demonstrated that restricted dorsal hippocampal lesions, but not similarly sized ventral lesions, impaired spatial learning.
- impaired spatial learning.

 22. Moser, M. B., Moser, E. I., Forrest, E., Andersen, P. & Morris, R. C. Spatial learning with a minislab in the dorsal hippocampus. *Proc. Natl Acad. Sci. USA* **92**, 9697–9701 (1995).
- Gray, J. A. & McNaughton, N. The Neuropsychology of Anxiety: An Enquiry into the Functions of the Septo-Hippocampal System (Oxford Univ. Press, 1982).
- van Groen, T. & Wyss, J. M. Extrinsic projections from area CA1 of the rat hippocampus: olfactory, cortical, subcortical, and bilateral hippocampal formation projections. J. Comp. Neurol. 302, 515

 –528 (1990).
- Canteras, N. & Swanson, L. W. Projections of the ventral subiculum to the amygdala, septum, and hypothalamus: a PHA-L anterograde tracing study in the rat. J. Comp. Neurol. 324, 180–194 (1992).
- Risold, P. Y. & Swanson, L. W. Structural evidence for functional domains in the rat hippocampus. Science 272, 1484–1486 (1996).
 A beautiful demonstration of topographical hippocampal connectivity with subcortical areas,
 - hippocampal connectivity with subcortical areas, this is one of the earliest studies to imply that the hippocampal long axis could comprise multiple domains as opposed to a ventral—dorsal dichotomy.
- Henke, P. G. Hippocampal pathway to the amygdala and stress ulcer development. *Brain Res. Bull.* 25, 691–695 (1990).

- Bannerman, D. M. et al. Regional dissociations within the hippocampus—memory and anxiety. Neurosci. Biobehav. Rev. 28, 273–283 (2004).
- Fanselow, M. S. & Dong, H.-W. Are the dorsal and ventral hippocampus functionally distinct structures? *Neuron* 65, 7–19 (2010).
- 30. Amaral, D. G. & Witter, M. P. The three-dimensional organization of the hippocampal formation: a review of anatomical data. *Neuroscience* 31, 571–591 (1989). A landmark synthesis of anatomical data, in which the organization of inputs along the hippocampal long axis was emphasized, as well as the longitudinal distribution of intrinsic hippocampal pathways.
- Kjelstrup, K. B. et al. Finite scale of spatial representation in the hippocampus. Science 321, 140–143 (2008).
 - The first evidence for a gradient of function in the rodent hippocampal long axis. A gradual increase in place field size from the dorsal to ventral hippocampus also indicates that the ventral hippocampus may subserve similar spatial processing functions as the dorsal hippocampus, but at a larger spatial scale.
- 52. Thompson, C. L. et al. Genomic anatomy of the hippocampus. Neuron 60, 1010–1021 (2008). A major advance in our understanding of functional divisions of the hippocampal long axis. The techniques developed in reference 42 were applied to show that CA3 can be divided into nine discrete domains on the basis of gene expression.
- Dolorfo, C. L. & Amaral, D. G. Entorhinal cortex of the rat: topographic organization of the cells of origin of the perforant path projection to the dentate gyrus. J. Comp. Neurol. 398, 25–48 (1998).
- Witter, M. P., Wouterlood, F. G., Nabér, P. A. & Van Haeften, T. Anatomical organization of the parahippocampal-hippocampal network. *Ann. NY Acad. Sci.* 911, 1–24 (2000).
- van Strien, N. M., Cappaert, N. L. & Witter, M. P. The anatomy of memory: an interactive overview of the parahippocampal-hippocampal network. *Nature Rev. Neurosci.* 10, 272–282 (2009).
- Jones, B. F. & Witter, M. P. Cingulate cortex projections to the parabippocampal region and hippocampal formation in the rat. *Hippocampus* 17, 957–976 (2007).
- Witter, M. P. Organization of the entorhinal
 hippocampal system: a review of current anatomical
 data. *Hippocampus* 3, 33–44 (1993).
- Risold, P. & Swanson, L. Connections of the rat lateral septal complex. *Brain Res. Rev.* 24, 115–195 (1997).

- Groenewegen, H. J., Vermeulen-Van der Zee, E., te Kortschot, A. & Witter, M. P. Organization of the projections from the subiculum to the ventral striatum in the rat. A study using anterograde transport of *Phaseolus vulgaris* leucoagglutinin. *Neuroscience* 23, 103–120 (1987).
- Kishi, T., Tsumori, T., Yokota, S. & Yasui, Y. Topographical projection from the hippocampal formation to the amygdala: a combined anterograde and retrograde tracing study in the rat. *J. Comp. Neurol.* 496, 349–368 (2006).
- Bayer, S. A. & Altman, J. Directions in neurogenetic gradients and patterns of anatomical connections in the telencephalon. *Prog. Neurobiol.* 29, 57–106 (1987).
- Lein, E. S. et al. Genome-wide atlas of gene expression in the adult mouse brain. Nature 445, 168–176 (2007)
- Dong, H. W., Swanson, L. W., Chen, L., Fanselow, M. S. & Toga, A. W. Genomic-anatomic evidence for distinct functional domains in hippocampal field CA1. *Proc. Natl Acad. Sci. USA* 106, 11794–11799 (2009).
- O'Reilly, K. C. et al. Identification of dorsal—ventral hippocampal differentiation in neonatal rats. Brain Struct. Funct. http://dx.doi.org/10.1007/s00429-014-0851-8 (2014).
- Dougherty, K. A., Islam, T. & Johnston, D. Intrinsic excitability of CA1 pyramidal neurones from the rat dorsal and ventral hippocampus. *J. Physiol.* 590, 5707–5722 (2012).
- Maggio, N. & Segal, M. Striking variations in corticosteroid modulation of long-term potentiation along the septotemporal axis of the hippocampus. J. Neurosci. 27, 5757–5765 (2007).
- Papatheodoropoulos, C. & Kostopoulos, G. Decreased ability of rat temporal hippocampal CA1 region to produce long-term potentiation. *Neurosci. Lett.* 279, 177–180 (2000).
- Dougherty, K. et al. Differential expression of HCN subunits alters voltage-dependent gating of h-channels in CA1 pyramidal neurons from dorsal and ventral hippocampus. J. Neurophysiol. 109, 1940–1953 (2013).
- Hussaini, S. A., Kempadoo, K. A., Thuault, S. J., Siegelbaum, S. A. & Kandel, E. R. Increased size and stability of CA1 and CA3 place fields in HCN1 knockout mice. *Neuron* 72, 643–653 (2011).
- knockout mice. *Neuron* **72**, 643–653 (2011).
 50. Giocomo, L. M. *et al.* Grid cells use HCN1 channels for spatial scaling. *Cell* **147**, 1159–1170 (2011).
- Garden, D. L., Dodson, P. D., O'Donnell, C., White, M. D. & Nolan, M. F. Tuning of synaptic integration in the medial entorhinal cortex to the organization of grid cell firing fields. *Neuron* 60, 875–889 (2008).
- Deguchi, Y., Donato, F., Galimberti, I., Cabuy, E. & Caroni, P. Temporally matched subpopulations of selectively interconnected principal neurons in the hippocampus. *Nature Neurosci.* 14, 495–504 (2011).
- 53. Moser, E. I. The multi-laned hippocampus. *Nature Neurosci.* **14**, 407–408 (2011).
- Hawrylycz, M. J. et al. An anatomically comprehensive atlas of the adult human brain transcriptome. *Nature* 489, 391–399 (2012).
- Li, X. G., Somogyi, P., Ylinen, A. & Buzsaki, G. The hippocampal CA3 network: an in vivo intracellular labeling study. J. Comp. Neurol. 339, 181–208 (1994).
- İshizuka, N., Weber, J. & Amaral, D. G. Organization of intrahippocampal projections originating from CA3 pyramidal cells in the rat. J. Comp. Neurol. 295, 580–623 (1990).
- Swanson, L. W., Wyss, J. M. & Cowan, W. M. An autoradiographic study of the organization of intrahippocampal association pathways in the rat. *J. Comp. Neurol.* 181, 681–715 (1978).
 Fricke, R. & Cowan, W. M. An autoradiographic study
- Fricke, R. & Cowan, W. M. An autoradiographic study of the commissural and ipsilateral hippocampodentate projections in the adult rat. *J. Comp. Neurol.* 181, 253–269 (1978).
- Kondo, H., Lavenex, P. & Amaral, D. G. Intrinsic connections of the macaque monkey hippocampal formation: II. CA3 connections. *J. Comp. Neurol.* 515, 349–377 (2009).
- Staresina, B. P., Fell, J., Do Lam, A. T., Axmacher, N. & Henson, R. N. Memory signals are temporally dissociated in and across human hippocampus and perirhinal cortex. *Nature Neurosci.* 15, 1167–1173 (2012).
- Patel, J., Fujisawa, S., Berényi, A., Royer, S. & Buzsáki, G. Traveling theta waves along the entire septotemporal axis of the hippocampus. *Neuron* 75, 410–417 (2012).

- Wilson, M. A. & McNaughton, B. L. Dynamics of the hippocampal ensemble code for space. Science 261, 1055–1058 (1993).
- Moser, M. B. & Moser, E. I. Distributed encoding and retrieval of spatial memory in the hippocampus. *J. Neurosci.* 18, 7535–7542 (1998).
- Jung, M. W., Wiener, S. I. & McNaughton, B. L. Comparison of spatial firing characteristics of units in dorsal and ventral hippocampus of the rat. *J. Neurosci.* 14, 7347–7356 (1994).
- MacDonald, C. J., Lepage, K. Q., Edén, U. T. & Eichenbaum, H. Hippocampal "time cells" bridge the gap in memory for discontiguous events. *Neuron* 71, 737–749 (2011).
- Kraus, B. J. et al. Hippocampal "time cells": time versus path integration. Neuron 78, 1090–1101 (2013)
- Pastalkova, E., Itskov, V., Amarasingham, A. & Buzsáki, G. Internally generated cell assembly sequences in the rat hippocampus. *Science* 321 1322–1327 (2008).
- Pilly, P. K. & Grossberg, S. How do spatial learning and memory occur in the brain? Coordinated learning of entorhinal grid cells and hippocampal place cells. *J. Cogn. Neurosci.* 24, 1031–1054 (2012).
 Muller, R. U. & Kubie, J. L. The effects of changes in
- Muller, R. U. & Kubie, J. L. The effects of changes in the environment on the spatial firing of hippocampal complex-spike cells. *J. Neurosci.* 7, 1951–1968 (1987).
- Colgin, L. L., Moser, E. I. & Moser, M. B. Understanding memory through hippocampal remapping. *Trends Neurosci.* 31, 469–477 (2008).
- Hafting, T., Fyhn, M., Molden, S., Moser, M. B. & Moser, E. I. Microstructure of a spatial map in the entorhinal cortex. *Nature* 436, 801–806 (2005).
- O'Keefe, J. & Burgess, N. Dual phase and rate coding in hippocampal place cells: theoretical significance and relationship to entorhinal grid cells. *Hippocampus* 15, 853–866 (2005).
- McNaughton, B. L., Battaglia, F. P., Jensen, O., Moser, E. I. & Moser, M. B. Path integration and the neural basis of the 'cognitive map'. *Nature Rev. Neurosci.* 7, 663–678 (2006).
- 74. Fuhs, M. C. & Touretzky, D. S. A spin glass model of path integration in rat medial entorhinal cortex.
- Neurosci. 26, 4266–4276 (2006).
 Solstad, T., Moser, E. I. & Einevoll, G. T. From grid cells to place cells: a mathematical model. *Hippocampus* 16, 1026–1031 (2006).
- Zhang, S.-J. et al. Optogenetic dissection of entorhinal-hippocampal functional connectivity. Science 340, 1232627 (2013)
- Science **340**, 1232627 (2013).

 77. Maurer, A. P., Vanrhoads, S. R., Sutherland, G. R., Lipa, P. & McNaughton, B. L. Self-motion and the origin of differential spatial scaling along the septotemporal axis of the hippocampus. *Hippocampus* **15**, 841–852 (2005).
- Fyhn, M., Molden, S., Witter, M. P., Moser, E. I. & Moser, M. B. Spatial representation in the entorhinal cortex. *Science* 305, 1258–1264 (2004).
- Brun, V. H. et al. Progressive increase in grid scale from dorsal to ventral medial entorhinal cortex. Hippocampus 18, 1200–1212 (2008).
- 80. Stensola, H. et al. The entorhinal grid map is discretized. Nature 492, 72–78 (2012). The discovery of discrete modules of grid-cell clusters in the rat EC that maintain grid scale and respond independently to the environment is important for understanding the increase in place field size along the hippocampal dorsoventral axis, as it raises the possibility that the latter is also discretized.
- Fyhn, M., Hafting, T., Treves, A., Moser, M. B. & Moser, E. I. Hippocampal remapping and grid realignment in entorhinal cortex. *Nature* 446, 190–194 (2007).
- O'Keefe, J. & Burgess, N. Geometric determinants of the place fields of hippocampal neurons. *Nature* 381, 425–428 (1996).
- Greicius, M. D. et al. Regional analysis of hippocampal activation during memory encoding and retrieval: fMRI study. Hippocampus 13, 164–174 (2003).
- Preston, A. R., Thomason, M. E., Ochsner, K. N., Cooper, J. C. & Glover, G. H. Comparison of spiral-in/ out and spiral-out BOLD fMRI at 1.5 and 3 T. Neuroimage 21, 291–301 (2004).
- Hartley, T., Maguire, E. A., Spiers, H. J. & Burgess, N. The well-worn route and the path less traveled: distinct neural bases of route following and wayfinding in humans. *Neuron* 37, 877–888 (2003).

- Suthana, N. A., Ekstrom, A. D., Moshirvaziri, S., Knowlton, B. & Bookheimer, S. Y. Human hippocampal CA1 involvement during allocentric encoding of spatial information. J. Neurosci. 29, 10512–10519 (2009).
- Maguire, E. A. et al. Navigation-related structural change in the hippocampi of taxi drivers. Proc. Natl Acad. Sci. USA 97, 4398

 –4403 (2000).
- Acad. Sci. USA 97, 4398–4403 (2000).
 88. Nadel, L., Hoscheidt, S. & Ryan, L. R. Spatial cognition and the hippocampus: the anterior–posterior axis.
 J. Cogn. Neurosci. 25, 22–28 (2013).
- Wolbers, T. & Buchel, C. Dissociable retrosplenial and hippocampal contributions to successful formation of survey representations. J. Neurosci. 25, 3333–3340 (2005)
- Doeller, C. F., King, J. A. & Burgess, N. Parallel striatal and hippocampal systems for landmarks and boundaries in spatial memory. *Proc. Natl Acad. Sci. USA* 105, 5915–5920 (2008).
- Baumann, O. & Mattingley, J. B. Dissociable representations of environmental size and complexity in the human hippocampus. *J. Neurosci.* 33, 10526–10533 (2013).
- Miller, J. F. et al. Neural activity in human hippocampal formation reveals the spatial context of retrieved memories. Science 342, 1111–1114 (2013).
 Colombo, M., Fernandez, T., Nakamura, K. &
- Colombo, M., Fernandez, T., Nakamura, K. & Gross, C. G. Functional differentiation along the anterior-posterior axis of the hippocampus in monkeys. J. Neurophysiol. 80, 1002–1005 (1998).
- Ekstrom, A. D. et al. Cellular networks underlying human spatial navigation. *Nature* 425, 184–188 (2003).
- Killian, N. J., Jutras, M. J. & Buffalo, E. A. A map of visual space in the primate entorhinal cortex. *Nature* 491, 761–764 (2012).
- Zeineh, M. M., Engel, S. A., Thompson, P. M. & Bookheimer, S. Y. Dynamics of the hippocampus during encoding and retrieval of face-name pairs. *Science* 299, 577–580 (2003).
 - This important methodological advance in human MRI of the hippocampus was the forerunner for now a large number of high-resolution hippocampal fMRI studies that localize activity to different hippocampal subregions.
- Ekstrom, A. D. et al. Advances in high-resolution imaging and computational unfolding of the human hippocampus. *Neuroimage* 47, 42–49 (2009).
- hippocampus. *Neuroimage* **47**, 42–49 (2009).

 8. Carr, V. A., Rissman, J. & Wagner, A. D. Imaging the human medial temporal lobe with high-resolution fMRI. *Neuron* **65**, 298–308 (2010).
- Derdikman, D. et al. Fragmentation of grid cell maps in a multicompartment environment. Nature Neurosci. 12, 1325–1332 (2009).
- Terrazas, A. et al. Self-motion and the hippocampal spatial metric. J. Neurosci. 25, 8085–8096 (2005).
- 101. Pikkarainen, M., Rönkkö, S., Savander, V., Insausti, R. & Pitkänen, A. Projections from the lateral, basal, and accessory basal nuclei of the amygdala to the hippocampal formation in rat. J. Comp. Neurol. 403, 229–260 (1999).
- 102. Petrovich, G. D., Canteras, N. S. & Swanson, L. W. Combinatorial amygdalar inputs to hippocampal domains and hypothalamic behavior systems. *Brain Res. Rev.* 38, 247–289 (2001).
- 103. Pitkänen, A., Pikkarainen, M., Nurminen, N. & Ylinen, A. Reciprocal connections between the amygdala and the hippocampal formation, perirhinal cortex, and postrhinal cortex in rat: a review. *Ann. NY Acad. Sci.* 911, 369–391 (2000).
 104. Fanselow, M. S. & LeDoux, J. E. Why we think
- 104. Fanselow, M. S. & LeDoux, J. E. Why we think plasticity underlying Pavlovian fear conditioning occurs in the basolateral amygdala. *Neuron* 23, 229–232 (1999).
- 105. Bast, T., Zhang, W. N. & Feldon, J. The ventral hippocampus and fear conditioning in rats. Different anterograde amnesias of fear after tetrodotoxin inactivation and infusion of the CABA_A agonist muscimol. Exp. Brain Res. 139, 39–52 (2001).
- 106. Maren, S., Aharonov, G. & Fanselow, M. S. Neurotoxic lesions of the dorsal hippocampus and Pavlovian fear conditioning in rats. *Behav. Brain Res.* 88, 261–274 (1997)
- 107. Carvalho, M. C., Masson, S., Brandao, M. L. & de Souza Silva, M. A. Anxiolytic-like effects of substance P administration into the dorsal, but not ventral, hippocampus and its influence on serotonin. *Peptides* 29, 1191–1200 (2008).
- 108. Fudge, J. L., Decampo, D. & Becoats, K. Revisiting the hippocampal–amygdala pathway in primates: association with immature-appearing neurons. *Neuroscience* 212, 104–119 (2012).

REVIEWS

- Dolcos, F., LaBar, K. S. & Cabeza, R. Interaction between the amygdala and the medial temporal lobe memory system predicts better memory for emotional events. *Neuron* 42, 855–863 (2004).
- events. Neuron 42, 855–863 (2004).

 110. Murty, V. P., Ritchey, M., Adcock, R. A. & LaBar, K. S. fMRI studies of successful emotional memory encoding: a quantitative meta-analysis.

 Neuropsychologia 48, 3459–3469 (2010).
- 111. Kensinger, E. A. & Schacter, D. L. Amygdala activity is associated with the successful encoding of item, but not source, information for positive and negative stimuli. J. Neurosci. 26, 2564–2570 (2006).
- 112. Kjelstrup, K. G. et al. Reduced fear expression after lesions of the ventral hippocampus. Proc. Natl Acad. Sci. USA 99, 10825–10830 (2002).
- Pentkowski, N. S., Blanchard, D. C., Lever, C., Litvin, Y. & Blanchard, R. J. Effects of lesions to the dorsal and ventral hippocampus on defensive behaviors in rats. *Eur. J. Neurosci.* 23, 2185–2196 (2006).
 Bannerman, D. *et al.* Double dissociation of function
- Within the hippocampus: spatial memory and hyponeophagia. *Behav. Neurosci.* 116, 884–901 (2002).
- 115. Moita, M. A., Rosis, S., Zhou, Y., LeDoux, J. E. & Blair, H. T. Hippocampal place cells acquire locationspecific responses to the conditioned stimulus during auditory fear conditioning. *Neuron* 37, 485–497 (2003).
- 116. Vanderwolf, C. H. Hippocampal electrical activity and voluntary movement in the rat. *Electroencephalogr. Clin. Neurophysiol.* 26, 407–418 (1969).
 117. Halgren, E. Firing of human hippocampal units in
- Halgren, E. Firing of human hippocampal units in relation to voluntary movements. *Hippocampus* 1, 153–161 (1991).
- Thoenissen, D., Zilles, K. & Toni, I. Differential involvement of parietal and precentral regions in movement preparation and motor intention. J. Neurosci. 22, 9024–9034 (2002).
- Zhang, W. N., Bast, T. & Feldon, J. Effects of hippocampal N-methyl-D-aspartate infusion on locomotor activity and prepulse inhibition: differences between the dorsal and ventral hippocampus. Behav. Neurosci. 116, 72–84 (2002).
- Peleg-Raibstein, D. & Feldon, J. Effects of dorsal and ventral hippocampal NMDA stimulation on nucleus accumbens core and shell dopamine release. Neuropharmacology 51, 947–957 (2006).
- Neardyndamacology 31, 341–337 (2000).
 Bardgett, M. E. & Henry, J. D. Locomotor activity and accumbens Fos expression driven by ventral hippocampal stimulation require D1 and D2 receptors. Neuroscience 94, 59–70 (1999).
- 122. Wu, M. & Brudzynski, S. M. Mesolimbic dopamine terminals and locomotor activity induced from the subiculum. *Neuroreport* 6, 1601–1604 (1995).
- 123. Legault, M., Rompre, P. P. & Wise, R. A. Chemical stimulation of the ventral hippocampus elevates nucleus accumbens dopamine by activating dopaminergic neurons of the ventral tegmental area. J. Neurosci. 20, 1635–1642 (2000).
- 124. Royer, S., Sirota, A., Patel, J. & Buzsáki, G. Distinct representations and theta dynamics in dorsal and ventral hippocampus. J. Neurosci. 30, 1777–1787 (2010).
- 125. Ruediger, S., Spirig, D., Donato, F. & Caroni, P. Goaloriented searching mediated by ventral hippocampus early in trial-and-error learning. *Nature Neurosci.* 15, 1563–1571 (2012).
- 126. Viard, A., Doeller, C. F., Hartley, T., Bird, C. M. & Burgess, N. Anterior hippocampus and goal-directed spatial decision making. J. Neurosci. 31, 4613–4621 (2011).
- 127. Mogenson, G. J., Jones, D. L. & Yim, C. Y. From motivation to action: functional interface between the limbic system and the motor system. *Prog. Neurobiol.* 14, 69–97 (1980).
- 128. Pennartz, C., Ito, R., Verschure, P., Battaglia, F. & Robbins, T. The hippocampal–striatal axis in learning, prediction and goal-directed behavior. *Trends Neurosci.* 34, 548–559 (2011).
- 129. Bast, T., Wilson, I. A., Witter, M. P. & Morris, R. G. From rapid place learning to behavioral performance: a key role for the intermediate hippocampus. *PLoS Biol.* 7, e1000089 (2009).
- Lepage, M., Habib, R. & Tulving, E. Hippocampal PET activations of memory encoding and retrieval: the HIPER model. *Hippocampus* 8, 313–322 (1998).
- Schacter, D. L. & Wagner, A. D. Medial temporal lobe activations in fMRI and PET studies of episodic encoding and retrieval. *Hippocampus* 9, 7–24 (1999).

- Daselaar, S. M., Fleck, M. S. & Cabeza, R. Triple dissociation in the medial temporal lobes: recollection, familiarity, and novelty. *J. Neurophysiol.* 96, 1902–1911 (2006).
- 133. Kohler, S., Crane, J. & Milner, B. Differential contributions of the parahippocampal place area and the anterior hippocampus to human memory for scenes. *Hippocampus* 12, 718–723 (2002). 134. Duzel, E. et al. Human hippocampal and parahippocampal activity during visual associative recognition memory for spatial and nonspatial stimulus configurations. *J. Neurosci.* 23, 9439–9444 (2003).
- 135. Kirchhoff, B. A., Wagner, A. D., Maril, A. & Stern, C. E. Prefrontal-temporal circuitry for episodic encoding and subsequent memory. *J. Neurosci.* 20, 6173–6180 (2000).
- Ranganath, C. & Rainer, G. Neural mechanisms for detecting and remembering novel events. *Nature Rev. Neurosci.* 4, 193–202 (2003).
- 137. Marr, D. Simple memory: a theory for archicortex. *Philos. Trans. R. Soc. Lond. B* **262**, 23–81 (1971).
- McNaughton, B. L. & Morris, R. G. Hippocampal synaptic enhancement and information storage within a distributed memory system. *Trends Neurosci.* 10, 408–415 (1987).
- Paulsen, O. & Móser, E. A model of hippocampal memory encoding and retrieval: GABAergic control of synaptic plasticity. *Trends Neurosci.* 21, 273–278 (1998).
- 140. Hasselmo, M. E., Bodelón, C. & Wyble, B. P. A proposed function for hippocampal theta rhythm: separate phases of encoding and retrieval enhance reversal of prior learning. *Neural Comput.* 14, 793–817 (2002).
- 141. Jezek, K., Henriksen, E. J., Treves, A., Moser, E. I. & Moser, M. B. Theta-paced flickering between place-cell maps in the hippocampus. *Nature* 478, 246–249 (2011)
- Poppenk, J., Evensmoen, H. R., Moscovitch, M. & Nadel, L. Long-axis specialization of the human hippocampus. *Trends Cogn. Sci.* 17, 230–240 (2013).
- 143. Hirshhorn, M., Grady, C., Shayna Rosenbaum, R., Winocur, G. & Moscovitch, M. Brain regions involved in the retrieval of spatial and episodic details associated with a familiar environment: an fMRI study. *Neuropsychologia* 50, 3094–3106 (2012).
- 144. Addis, D. R., Moscovitch, M., Crawley, A. P. & McAndrews, M. P. Recollective qualities modulate hippocampal activation during autobiographical memory retrieval. *Hippocampus* 14, 752–762 (2004).
- 145. Chua, E. F., Schacter, D. L., Rand-Giovannetti, E. & Sperling, R. A. Evidence for a specific role of the anterior hippocampal region in successful associative encoding. *Hippocampus* 17, 1071–1080 (2007).
- 146. Davachi, L. Item, context and relational episodic encoding in humans. *Curr. Opin. Neurobiol.* **16**, 693–700 (2006)
- 693–700 (2006).

 147. Giovanello, K. S., Schnyer, D. & Verfaellie, M. Distinct hippocampal regions make unique contributions to relational memory. *Hippocampus* 19, 111–117 (2009).
- 148. Cohen, N. J. & Eichenbaum, H. Memory, Amnesia,
- and the Hippocampal System (MIT Press, 1993).
 149. Bunsey, M. & Eichenbaum, H. Conservation of hippocampal memory function in rats and humans Nature 379, 255–257 (1996).
 - This important rodent lesion study extended the hippocampal role in spatial memory to a non-spatial task transitive inference that can also be studied in humans.
- Heckers, S., Zalesak, M., Weiss, A. P., Ditman, T. & Titone, D. Hippocampal activation during transitive inference in humans. *Hippocampus* 14, 153–162 (2004).
- 151. Preston, A. R., Shrager, Y., Dudukovic, N. M. & Gabrieli, J. D. E. Hippocampal contribution to the novel use of relational information in declarative memory. *Hippocampus* 14, 148–152 (2004).
- 152. Eichenbaum, H. Hippocampus: cognitive processes and neural representations that underlie declarative memory. *Neuron* 44, 109–120 (2004).
- 153. Komorówski, R. W. et al. Ventral hippocampal neurons are shaped by experience to represent behaviorally relevant contexts. J. Neurosci. 33, 8079–8087 (2013).
- 154. Geisler, C. et al. Temporal delays among place cells determine the frequency of population theta oscillations in the hippocampus. Proc. Natl Acad. Sci. USA 107, 7957–7962 (2010).

- 155. O'Keefe, J. & Recce, M. L. Phase relationship between hippocampal place units and the EEG theta rhythm. *Hippocampus* 3, 317–330 (1993).
- 156. Skaggs, W. E., McNaughton, B. L., Wilson, M. A. & Barnes, C. A. Theta phase precession in hippocampal neuronal populations and the compression of temporal sequences. *Hippocampus* 6, 149–172 (1996).
- 157. Buzsáki, G. Neural syntax: cell assemblies, synapsembles, and readers. *Neuron* 68, 362–385 (2010)
- 158. Geuze, E., Vermetten, E. & Bremner, J. D. MR-based in vivo hippocampal volumetrics: 2. findings in neuropsychiatric disorders. Mol. Psychiatry 10, 160–184 (2005).
- 159. Geuze, E., Vermetten, E. & Bremner, J. D. MR-based in vivo hippocampal volumetrics: 1. review of methodologies currently employed. Mol. Psychiatry 10, 147–159 (2005).
- 160. Small, S. A., Schobel, S. A., Buxton, R. B., Witter, M. P. & Barnes, C. A. A pathophysiological framework of hippocampal dysfunction in ageing and disease. Nature Rev. Neurosci. 12, 585–601 (2011).
- 161. Titone, D., Ditman, T., Holzman, P. S., Eichenbaum, H. & Levy, D. L. Transitive inference in schizophrenia: impairments in relational memory organization. Schizophr Res 68, 235–247 (2004)
- Schizophr. Res. **68**, 235–247 (2004). 162. Ongur, D. et al. The neural basis of relational memory deficits in schizophrenia. Arch. Gen. Psychiatry **63**, 356–365 (2006).
 - Although this study does not report a double dissociation between anterior and posterior hippocampal responses, it links deficits in transitive inference to abnormal anterior hippocampal function in schizophrenia.
- 163. Goldman, M., Marlow-O'Connor, M., Torres, I. & Carter, C. S. Diminished plasma oxytocin in schizophrenic patients with neuroendocrine dysfunction and emotional deficits. Schizophr. Res. 98, 247–255 (2008).
- 164. Goldman, M. B. *et al.* Reduced anterior hippocampal formation volume in hyponatremic schizophrenic patients. *Hippocampus* **17**, 554–562 (2007)
- patients. *Hippocampus* **17**, 554–562 (2007).
 165. Luchins, D. J., Nettles, K. W. & Goldman, M. B.
 Anterior medial temporal lobe volumes in polydipsic schizophrenic patients with and without hypoosmolemia: a pilot study. *Biol. Psychiatry* **42**, 767–770 (1997).
- 166. Gallagher, B. B. Endocrine abnormalities in human temporal lobe epilepsy. *Yale J. Biol. Med.* 60, 93–97 (1987).
- 167. Herzog, A. G., Seibel, M. M., Schomer, D. L., Vaitukaitis, J. L. & Geschwind, N. Reproductive endocrine disorders in women with partial seizures of temporal lobe origin. *Arch. Neurol.* 43, 341–346 (1986).
- 168. Quigg, M., Kiely, J. M., Shneker, B., Veldhuis, J. D. & Bertram, E. H. Interictal and postictal alterations of pulsatile secretions of luteinizing hormone in temporal lobe epilepsy in men. *Ann. Neurol.* 51, 559–566 (2002).
- 169. Kheirbek, M. A. et al. Differential control of learning and anxiety along the dorsoventral axis of the dentate gyrus. Neuron 77, 955–968 (2013).
- 170. Gloor, P. *The Temporal Lobe and Limbic System* (Oxford Univ. Press, 1997).
- Insausti, R. Comparative anatomy of the entorhinal cortex and hippocampus in mammals. *Hippocampus* 3, 19–26 (1993).
- 172. Frotscher, M. & Seress, L. in *The Hippocampus Book* (eds Andersen, P., Morris, R., Amaral, D., Bliss, T. & O'Keefe, J.) 115–132 (Oxford Univ. Press, 2007).
- 173. Macchi, G. The ontogenic development of the olfactory telencephalon in man. *J. Comp. Neurol.* 95, 245–305 (1951)
- 174. Kier, E. L., Fulbright, R. K. & Bronen, R. A. Limbic lobe embryology and anatomy: dissection and MR of the medial surface of the fetal cerebral hemisphere. AJNR Am. J. Neuroradiol. 16, 1847–1853 (1995).
- 175. Friedman, D. P., Aggleton, J. P. & Saunders, R. C. Comparison of hippocampal, amygdala, and perirhinal projections to the nucleus accumbens: combined anterograde and retrograde tracing study in the Macaque brain. J. Comp. Neurol. 450, 345–365 (2002).
- 176. Saunders, R. C. & Aggleton, J. P. Origin and topography of fibers contributing to the fornix in macaque monkeys. *Hippocampus* 17, 396–411 (2007).
- Aggleton, J. P. A description of the amygdalohippocampal interconnections in the macaque monkey. Exp. Brain Res. 64, 515–526 (1986).

- 178. Witter, M. P., Van Hoesen, G. W. & Amaral, D. G. Topographical organization of the entorhinal projection to the dentate gyrus of the monkey.
 1 Navges 19, 216, 2238 (1989)
- J. Neurosci. 9, 216–228 (1989). 179. Mohedano-Moriano, A. et al. Topographical and laminar distribution of cortical input to the monkey entorhinal cortex. J. Anat. 211, 250–260 (2007).
- Mohedano-Moriano, A. et al. Convergence of unimodal and polymodal sensory input to the entorhinal cortex in the fascicularis monkey. Neuroscience 151, 255–271 (2008).
- Aggleton, J. P. Multiple anatomical systems embedded within the primate medial temporal lobe: implications for hippocampal function. *Neurosci. Biobehav. Rev.* 36, 1579–1596 (2012)
- 36, 1579–1596 (2012).
 182. Aggleton, J. P., Wright, N. F., Vann, S. D. & Saunders, R. C. Medial temporal lobe projections to the retrosplenial cortex of the macaque monkey. Hippocampus 22, 1883–1900 (2012).
- 183. Canto, C. B., Wouterlood, F. G. & Witter, M. P. What does the anatomical organization of the entorhinal cortex tell us? *Neural Plast.* 2008, 381243 (2008).
- 184. Bragdon, A. C., Taylor, D. M. & Wilson, W. A. Potassium-induced epileptiform activity in area CA3 varies markedly along the septotemporal axis of the rat hippocampus. *Brain Res.* 378, 169–173 (1986)
- 185. Engel J. Jr et al. Presurgical evaluation for partial epilepsy: relative contributions of chronic depthelectrode recordings versus FDG-PET and scalpsphenoidal ictal EEG. Neurology 40, 1670–1677 (1990).
- 186. Mouritzen Dam, A. Epilepsy and neuron loss in the hippocampus. *Epilepsia* 21, 617–629 (1980).

- 187. Babb, T. L. et al. Temporal lobe volumetric cell densities in temporal lobe epilepsy. *Epilepsia* 25, 729–740 (1984).
- O'Connor, W. M. et al. Hippocampal cell distributions in temporal lobe epilepsy: a comparison between patients with and without an early risk factor. *Epilepsia* 37, 440–449 (1996).
- 189. Santarelli, L. *et al.* Requirement of hippocampal neurogenesis for the behavioral effects of antidepressants. *Science* **301**, 805–809 (2003)
- antidepressants. *Science* **301**, 805–809 (2003). 190. Sahay, A. & Hen, R. Adult hippocampal neurogenesis in depression. *Nature Neurosci.* **10**, 1110–1115 (2007)
- Boldrini, M. et al. Antidepressants increase neural progenitor cells in the human hippocampus. Neuropsychopharmacology 34, 2376–2389 (2009)
- 192. Lipska, B. K., Jaskiw, G. E. & Weinberger, D. R. Postpubertal emergence of hyperresponsiveness to stress and to amphetamine after neonatal excitotoxic hippocampal damage: a potential animal model of schizophrenia. *Neuropsychopharmacology* 9, 67–75 (1993).
- Benes, F. M., Sorensen, I. & Bird, E. D. Reduced neuronal size in posterior hippocampus of schizophrenic patients. *Schizophr. Bull.* 17, 597–608 (1991).
 Maller, J. J. et al. Hippocampal volumetrics in
- 194. Maller, J. J. et al. Hippocampal volumetrics in treatment-resistant depression and schizophrenia: the devil's in De-Tail. Hippocampus 22, 9–16 (2012).
- 195. Ashton, D., Van Reempts, J., Haseldonckx, M. & Willems, R. Dorsal-ventral gradient in vulnerability of CA1 hippocampus to ischemia: a combined histological and electrophysiological study. *Brain Res.* 487, 368–372 (1989).

- 196. Martens, U., Capito, B. & Wree, A. Septotemporal distribution of [¹H]MK-801, [²H]AMPA and [³H]kainate binding sites in the rat hippocampus. *Anat. Embryol.* 198, 195–204 (1998).
- Rothman, S. M. & Olney, J. W. Excitotoxity and the NMDA receptor. *Trends Neurosci.* 10, 299–302 (1987).
- 198. Smith, M. L., Auer, R. N. & Siesjo, B. K. The density and distribution of ischemic brain injury in the rat following 2–10 min of forebrain ischemia. *Acta Neuropathol.* 64, 319–332 (1984).
 199. Horstmann, A. et al. Resuscitating the heart but losing
- Horstmann, A. et al. Resuscitating the heart but losing the brain: brain atrophy in the aftermath of cardiac arrest. Neurology 74, 306–312 (2010).
- Zola-Morgan, S., Squire, L. R. & Amaraí, D. G. Human amnesia and the medial temporal region: enduring memory impairment following a bilateral lesion limited to field CA1 of the hippocampus. *J. Neurosci.* 6, 2950–2967 (1986).

Acknowledgements

The authors thank L. Giocomo for helpful discussion. B.A.S. is supported by the Spanish Ministry of Science and Innovation (SAF2011-27766) and Marie Curie Career Integration Grant (FP7-PEOPLE-2011-CIG 304248), E.IM. and M.P.W. are supported by the Kavli Foundation and Centre of Excellence grant from the Norwegian Research Council, and E.S.L. by the Allen Institute for Brain Science.

Competing interests statement

The authors declare no competing interests.

SUPPLEMENTARY INFORMATION

See online article: <u>S1</u> (box) | <u>S2</u> (table)

ALL LINKS ARE ACTIVE IN THE ONLINE PDF