Dynamic firing patterns of ventral hippocampal neurons govern anxiety-dependent behaviour and predict the extent of exploration of an anxiogenic location

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Abstract

The ventral hippocampus (vH) plays a key role for anxiety-related behaviour and vH neurons increase their firing when animals explore anxiogenic environments. However, if and how such neuronal activity induces or restricts the exploration of an anxiogenic location remains unexplained. Here, we developed a novel behavioural paradigm to motivate rats to explore an anxiogenic location. Rats ran along an elevated linear maze with protective sidewalls, which were temporarily removed in parts of the track to introduce a novel anxiogenic location. We recorded neuronal action potentials during task performance and found that vH neurons exhibited remapping of activity, overrepresenting novel anxiogenic locations. Direction-dependent firing was homogenised by the anxiogenic experience. We further showed that the activity of vH neurons predicted the extent of exploration of the anxiogenic location. Our data suggest that anxiety-related firing does not solely depend on the exploration of anxiogenic environments, but also on intentions to explore them.
Introduction

In the *Epistulae Morales ad Lucilium*, Seneca wrote: “There are more things, Lucilius, likely to frighten us than there are to crush us; we suffer more often in imagination than in reality”. This sentence, from one of the key figures of the school of stoicism describes inner fear within our imagination in the absence of a direct fear-provoking stimulus. Nowadays, even though anxiety and fear correspond to the same theoretical construct in some literature, anxiety differentiates from the latter based on the potential nature of the threat in the absence of an imminent harmful stimulus. Anxiety disorders are becoming more commonly reported: 12-month prevalence estimates on mental disorders show that at least 14% of people in the European Union suffer from anxiety disorders, and around 31% of people in the United States have experienced some type of anxiety disorder in their lifetime. Different brain areas play a role in the underlying circuitry of anxiety. Stimulations in the brainstem, more precisely in the periaqueductal grey matter (PAG) or the locus coeruleus are specifically involved in the symptomatology of anxiety. Some studies have also shown how the amygdala plays a role in humans suffering from anxiety disorders or in animal models with generalised fear responses associated with anxiety. Also, by using the elevated plus maze (EPM) as an anxiety task, Tye and colleagues induced anxiolytic effects by targeting projections from the basolateral amygdala (BA) to the central nucleus of the amygdala. Additional structures are involved in anxiety and include the bed nucleus of the stria terminalis, whose subdivisions are differentially involved in anxiety responses, and the medial prefrontal cortex (mPFC), which has also been directly linked to anxiety processing in both humans and rodents.

Last but not least, the ventral hippocampus (vH) plays a critical role in anxiety behaviour. Lesions in the vH induce anxiolysis during the exploration of elevated open arenas (e.g. EPM), or generally in tasks associated with approach-avoidance conflicts. Neurons recorded in the vH showed increased firing in locations with elevated anxiety. Likewise, projections neurons from- and to the vH exhibit anxiety-related activity: amygdala projections to the vH are specifically shaping anxiety-related behaviour during the exploration of an EPM. Similarly, the reciprocal connection (from vH to BA) is involved in the expression of context-dependent fear memories. Furthermore, information related to anxiety in the vH is directly routed to the mPFC, and synchronised activity in this monosynaptically connected long-range circuit is essential for the expression of anxiety behaviour. Motivated by the critical role of the dorsal hippocampus region (dH) in the encoding of spatial information, several studies focusing on the vH have described its involvement in spatial coding. Yet, the neuronal dynamics in the vH associated with anxiety during spatial navigation remain poorly understood.
To study the neuronal dynamics associated with anxiety behaviour in the vH, we simultaneously recorded the activity of individual neurons during the exploration of the EPM, as well as during the exploration of a novel behavioural paradigm, the elevated linear maze (ELM). During the same recording session, we modified the ELM from a non-anxiogenic to an anxiogenic configuration, while recording the activity of the same individual vH neurons. This enabled the investigation of the neuronal dynamics within the vH underlying different anxiety states. We specifically examined the remapping of neuronal activity at the single neuron- and population-level as animals transitioned from one ELM configuration to another. Collectively, the results of this study show that the neuronal activity in the vH does not simply reflect anxiogenic locations but that it is dynamically modulated by the experience and expectation of anxiety during spatial navigation.

**Results**

**The firing activity of ventral hippocampal neurons is dynamically modulated during EPM exploration.**

Rats \((n = 6)\) freely explored an EPM consisting of two opposite arms with protective side-walls (closed arms) and two opposite arms without sidewalls (open arms) (Fig. 1a). The rats exhibited strong anxiety-related behaviour by spending most of the time in the closed arms, avoiding the more anxiogenic open arms and the centre (Fig. 1a, bottom. Closed vs open, \(p = 9.5615e-10\); closed vs centre, \(p = 9.5657e-10\). One-way ANOVA, Tukey-Kramer for multiple comparisons). While rats explored the EPM, we recorded neuronal activity in the vH with tetrodes (Fig. 1b) and isolated individual spikes from different single neurons \((n = 98)\). We identified vH neurons with previously described activity patterns\(^{25}\), exhibiting preferential firing in the open arms, closed arms, or centre of the EPM (Fig 1c). To understand the effect of the open areas on the neuronal activity, we defined six possible trajectories taken by animals during EPM exploration (from one closed arm to any other arm). After linearizing the trajectories (see methods), we organised the activity of the recorded neurons based on the spatial location of their peak (i.e. maximal) firing activity (Fig. 1d). We observed that the peak firing activities of individual neurons spanned the entire maze even though the exploration of open arms was minimal. Nonetheless, the peak firing activity of vH neurons was elevated in the open areas of the EPM and the centre of the maze, even when animals shuttled from one closed arm to the other one (Fig. 1e).

To further investigate the evolution and dynamics of the neuronal activity during the exploration of open areas, we analysed changes in activity of individual vH neurons while animals were navigating from a closed to an open arm (Fig. 1f). We then calculated the correlation between the furthest location reached by animals in the open arms and the location of the peak firing activity of each neuron (Fig. 1f). Around 20% of the recorded neurons (19 out of 98 neurons) showed a significant
correlation indicating a dynamic modulation of the neuronal activity during the exploration of an anxiogenic location (Fig. 1g).

However, these results suffer from important limitations: First, the extremely low and sporadic exploration of the open arms does not provide robust data sampling to test hypotheses regarding the neuronal computations associated with the exploration of an anxiogenic location. Second, the non-anxiogenic and the anxiogenic areas of the EPM are constantly present. This means that the characterization of the anxiety states experienced by animals is not trivial as these may feel continuously anxious not solely in open spaces but also when considering to visit an open arm, while being in the closed arms of the EPM. For these reasons, we developed a novel behavioural paradigm to better control the transitions between anxiety states and the extent of exploration of anxiogenic areas.

Removal of protective sidewalls along an ELM induces anxiety behaviour

We developed an ELM, which consisted of an elevated linear track with removable protective sidewalls. This allowed to have sidewalls either all along the entire ELM, called closed-closed configuration (CC), or to have the sidewalls removed from half of the track, called closed-open (CO) configuration (Fig. 2a). The rapid removal of the sidewalls enabled the alternation between a non-anxiogenic and an anxiogenic configuration for the same maze and in a single behavioural session with continuously exploring rats. To control for novelty effects, we modified the visual appearance and floor texture of sidewalls in one half of the track. This configuration was termed closed-texture (CT) configuration (Fig. 2b). Rats (n = 6) were motivated to fully explore the ELM, by shuttling from one end of the track to the other one over numerous trials, to receive food rewards (Fig. 2c). To assess whether the removal of sidewalls induces behavioural readouts of anxiety similar to those of the EPM, we calculated the percentage of time spent on each of the arms for each configuration (Fig. 2d). No differences were observed in the time spent by rats on the different arms during non-anxiogenic explorations (CC, CT). On the contrary, a significant difference was found in the configuration with sidewalls removed (CO exploration, p = 1.45e-05, Wilcoxon signed-rank). The time spent in the centre area (11.5 cm around the middle of the track) was also longer during CO exploration compared to CC or CT explorations (Fig. 2e, p = 2e-05 and p = 0.034 against CC and CT respectively, one-way ANOVA, Turkey-Kramer for multiple comparisons), suggesting of a hesitation to enter this open area of the maze.

Overall, the removal of sidewalls along the ELM induced anxiety behaviour that evolved during single behavioural sessions according to the anxiety content of each maze configuration.
**Overrepresentation and remapping of vH activity during anxiety**

We recorded a total of 133 neurons with tetrodes in the vH (Fig. 1b), while the rats were exposed to the different configurations of the ELM. When the activity of individual neurons was sorted according to the spatial location of their peak firing activity, we observed that it spanned over the entire extent of the ELM for the three different configurations (Fig. 3a). However, during the CO exploration, the entire distribution of peak firing activity was skewed toward the open area. Furthermore, the activity of vH neurons appeared to be spatially broader during CO exploration, suggesting that the anxiogenic area in the CO configuration selectively affect the spatial features of vH neurons. To validate these two observations, first, we assessed the proportion of vH neurons with peak firing activity located on the different halves of the ELM during CC exploration. We found no differences in the proportion of peak firing activity located in each half of the CC configuration, even for the closed half that was going to be opened in the CO configuration (Fig. 3b, left). Subsequently, when one half was opened, a remapping of the neuronal activity was induced in the anxiogenic location. We observed a higher proportion of peak firing activities located on the open arm (Fig. 3b left, \( p = 0.001 \), chi-square test). In a subsequent analysis, we detected that a significant proportion of vH neurons changed the location of their peak firing activities from the closed area to the novel open area after sidewall removal (Fig. 3b right, \( p = 0.015 \), chi-square test). Importantly, no differences were found for the location and location changes of peak firing activity during the exploration of the CT configuration using a novel texture and visual cues in the closed arm (Fig. 3b right), suggesting that dynamic remapping of vH activity is contingent to the experience of anxiety rather than novelty.

Next, we quantified the remapping of peak firing activity for vH neurons that changed their firing peak location from the closed arm (during CC exploration) towards the open arm (in the CO exploration). Similarly, we analysed the remapping of peak firing locations from the arm to be opened (during CC exploration) towards the closed arm (during CO exploration) (Fig. 3d left). We found a difference (\( p = 0.0013 \), Wilcoxon signed-rank test) in the location of peak firing activity during the transition from the CC to CO configuration, but not when the configuration was changing from CC to CT (Fig. 3d right). This supported the view of a remapping of neuronal activity in vH with neurons dynamically shifting their peak firing preference towards the anxiogenic location.

To validate our second observation of a spatial broadening of the neuronal activity in the CO configuration, we used a coverage index (also known in the literature as sparsity index, see methods) and compared the change of this index during changes in ELM configuration. We observed a selective spatial broadening of neuronal activity from the CC to the CO configuration (Fig. 3e top, \( p = 8.237e^{-06} \), Wilcoxon signed-rank), which was absent during the transition between the CC and the CT configuration.
Collectively, these data indicate a dynamical recruitment of vH neurons when swapping between a non-anxiogenic (CC) to an anxiogenic configuration (CO), generating not solely a remapping of activity but also an overrepresentation of the anxiogenic area. As these effects were not detected during a novel, but non-anxiogenic experience, we concluded that they could be attributed to the processing of anxiety during open arm exploration rather than novelty perception per se.

The direction-dependent activity of vH neurons becomes homogenised following the introduction of an anxiogenic location

Hippocampal place cells have been reported to exhibit direction-specific spatial modulation of activity as animals run along a linear maze\textsuperscript{35-37}. This raises the question as to whether the activity of vH neurons, recruited by areas with increased anxiety content, are also modulated by the direction of the journey along a linear track. When monitoring the activity of individual vH neurons recorded during the exploration of the CC configuration (Fig. 4a), we expectedly observed a profound direction-dependent difference. However, this direction-dependent neuronal firing of the same vH neurons became homogenised, meaning that it was very similar in both directions, once the sidewalls were removed and the rats were exploring the CO configuration. To quantify this phenomenon, we calculated the place field similarity (PFS, see methods) of a neuron between its activity while the animal moved in one direction vs. the other direction. Values closer to one imply that both neuronal activities are very similar in both directions. As expected, the PFS index of animals exploring the CC configuration was different from the PFS index of animals exploring the CO configuration (Fig. 4b, \( p = 7.622\times10^{-04} \), two-sample Kolmogorov-Smirnov test). Based on the values of the PFS indices (median PFS\textsubscript{CC} = -0.0182, median PFS\textsubscript{CO} = 0.3351), we inferred that the significant difference between the PFS distributions is due to a similar neuronal activity in both directions during the exploration of the CO configuration.

To confirm that the spatial activity of vH neurons is similar during the exploration of CO configuration and that this activity is higher in the open arm, we averaged the neuronal activity in the two types of arms (i.e. the arm always closed and the arm to be open, see methods). For both configurations (CC and CO), we calculated the sum and the difference of the averaged activities for both directions. When we summed the averaged activity for one direction with the averaged activity for the opposite, we found that the highest neuronal activity was located in the open arm of the CO configuration, but not in the CC configuration (Fig. 4c top, \( p = 2.16e-06 \), Wilcoxon signed-rank test). When we calculated the absolute difference between the averaged activity of both directions, the result was significantly lower for the both arms of the CO configuration when tested against the CC configuration, indicating that the direction-dependent neuronal activity of individual vH neurons is more similar in the CO than in the CC configuration (\( p = 5.12e-12, p = 1.29e-04 \), first arm and second arm respectively, Wilcoxon signed-rank test). To corroborate that such a phenomenon was not due to
the novelty of the arm, we calculated the PFS indexes of the neuronal activity during the exploration of a novel arm (CT), instead of an anxiogenic location (i.e. CO configuration). The PFS indexes during the CT configuration were significantly lower than during the exploration of the CO configuration and more similar to the PFS indexes during the exploration of the CC configuration (called CC2 when it was immediately before CT) (Fig. 4d, CO vs CC, \(p = 7.622\times10^{-4}\); vs CT, \(p = 0.0033\) and vs CC2, \(p = 0.0166\); two-sample Kolmogorov-Smirnov test).

In conclusion, even though there was direction-dependent neuronal activity in the vH during the exploration of a non-anxiogenic linear maze, this spatial dependency was reduced when animals encountered an anxiogenic location. In addition, the activity of individual vH neurons tended to cluster on the open area of the ELM, independent of the animal’s direction of travel.

**The activity of vH neurons predicts the extent of exploration of an anxiogenic location**

We have shown that neurons of the vH change their activity patterns during the experience of anxiety. However, the question remains if and how this anxiety-related firing predicts the extent of exploration of an anxiogenic location. In accordance with the EPM findings (Fig. 1f), we observed also a trial-by-trial dynamic modulation of the neuronal activity during the exploration of the open area of the ELM (Fig. 5a). We found individual vH neurons whose location of peak firing activity was modulated by the extent of exploration of the open area on a trial by trial basis. The location of the peak firing activity was related to how far animals would venture during each trial. 59 out of 133 neurons (44% of the neurons) exhibited a correlation between peak firing activity and extent of exploration (Fig. 5b, Spearman rank correlation).

Next, we asked whether neuronal firing in the closed area might be predictive of the extent of an upcoming exploration in the anxiogenic area. To do so, we divided spatial explorations into two groups depending on how far animals explored the open arm during a particular trial: proximal and distal exploration trials were defined using an individual threshold for each session \(\text{Threshold} = \frac{\text{Furthest spatial bin} - \text{Nearest spatial bin}}{2}\) (Fig. 5c). We then tested if the neuronal activity before entering the open arm was informative of how far animals would venture into the open arm (proximal or distal explorations). By using the firing activity of each vH neuron prior to the entry to the open arm in the CO configuration, we represented each trial by a population vector built with the activity of all the co-recorded neurons during single recording days by \(T_n = \langle FR_{1n}, FR_{2n}, ..., FR_{mn} \rangle\) where FR is the firing rate, \(n\) is the total number of trials, and \(m\) is the total number of neurons. If some anxiety-related information is computed during the exploration of the closed part of the ELM, we then anticipated that this neuronal activity would predict the extent of explorations (proximal or distal explorations) in the open anxiogenic arm. To better visualise the population activity in the close arm related to the future extent of exploration of the open arm, we calculated the principal components (PCA, see methods) of
the population vectors and plotted the two first principal components (Fig. 5c). We observed a clear separation between the two types of trials (proximal exploration, red; distal exploration, blue) based on the neuronal activity recorded. To further strengthen this observation, we trained a support vector machine (SVM, Fig. 5d) using the neuronal activity in the closed arms for each recording day and were able to predict the extent of exploration for single trials (see methods). We found that during individual recording sessions (Fig. 5d) the performance of the SVM was above chance levels in 8 out of 11 sessions.

These analyses indicate that the neuronal representation of the anxiogenic location was not only dynamically modulated by the direct experience of anxiety, but already existed in the closed arm, possibly reflecting the intention to venture into the open arm. This implies that neuronal activity within the vH can predict upcoming anxiogenic situations, even when animals are still located in a safer environment without a direct exposure to an anxiety-inducing location.
Discussion

To investigate the neuronal dynamics governing anxiety behaviour in the vH, we recorded the activity of individual neurons while rats explored anxiogenic locations. In addition to the classical EPM, we used a novel ELM, which allowed us to rapidly change the anxiety content of the maze to expose rats to non-anxiogenic or anxiogenic configurations. We found that the neuronal activity of the vH exhibited a uniform spatial representation in the non-anxiogenic configuration of the ELM. However, vH neurons displayed direction-dependent spatial firing when shuttling from one end of the ELM to another. When the anxiogenic location was introduced by removing the protective sidewalls from half of the track, the peak firing activity remapped towards the newly introduced anxiogenic location. Importantly, direction-dependent firing was homogenised, and spatially-modulated firing patterns of individual vH neurons were similar in both trajectories. In addition, the activity of vH neurons was modulated by the distance of exploration in the anxiogenic environment. We observed that for a population of vH neurons, the location of their peak firing activity was strongly related to how far animals would explore the anxiogenic location during each individual trial. Of important note, the population activity predicted the extent of the upcoming exploration in the open arm even before rats entered into the open anxiogenic location.

Much of the research on anxiety in freely-moving rodents has been relying on the EPM. Using the EPM, it has been shown that: amygdala projections to the vH control the expression of anxiety; there is an anxiety-associated neuronal activity in the vH routed to the mPFC; and that the vH-prefrontal pathway is critical for anxiety behaviour. The first part of this manuscript focuses on the activity of neurons recorded in the vH during the exploration of an EPM. We divided the exploration of the EPM into different trajectories. We observed a localised increase in the density of peak firing activity when rats crossed the centre in all the different C-C or C-O trajectories (Fig. 1e). This might be explained by the fact that not only the open arms of an EPM are anxiogenic, but also the centre zone. Strikingly, we found evidence of a group of neurons with modulated firing during the exploration of the open arms. Unfortunately, it proved difficult to analyse these effects quantitatively, because rats explored the open arms only minimally and sporadically, consistent with the anxiogenic nature of the EPM paradigm.

To overcome this problem, we introduced a novel ELM on which rats were motivated to shuttle from one extremity to the other to receive rewards. The ELM had three different configurations: non-anxiogenic (CC configuration); anxiogenic (CO configuration); and a configuration with new texture and visual cues (CT). Configurations could be quickly switched within a session. At the behavioural level, we observed anxiety-related behaviour during ELM exploration comparable to the ones during EPM exploration (Fig. 2 d, e). However, the main advantage of the ELM was the possibility to record the
neuronal activity of the same vH neurons while rapidly modifying ELM configurations and motivating rats to spend more time in the anxiogenic location. After animals transited from a non-anxiogenic to an anxiogenic configuration, we observed a significant increment in peak firing activities predominately located in the open area. We attributed this recruitment or remapping of the neuronal activity to the novel anxiogenic location. The neuronal mechanisms during spatial remapping remain largely elusive. Global remapping is a phenomenon observed in the dH when animals move from one environment to a different one. One could argue that the simple removal of the walls is changing the environment of the ELM and therefore animals could perceive the open arm as a completely novel environment, inducing global remapping. Nevertheless, remapping of neuronal activity in a new environment is expected to be random and independent of a previously explored environment, even when emotional contexts are introduced. However, the remapping of neuronal activity observed in ventral hippocampal neurons is not arbitrary as it is most prominent in the anxiogenic location. Nonetheless, the opening of walls per se, independent of the anxiogenic experience, could cause changes in vH neuronal activity. Previous work has shown that novel environments elicit the activation of vH neurons to a similar extent as an aversive stimulus. To show that the remapping of neuronal activity relies on anxiety, we changed the ELM from a CC to CT configuration during the same recording session. In the CT configuration, the previously open location changed to a novel one with a different floor texture and visual patterns in the inner part of the walls, but with protective sidewalls kept all along the track (Fig. 2b). As anticipated, these changes equally induced a remapping of neuronal activity in the vH. Yet, in this case, the remapping was random and the novel area did not show an increased number of peak firing activity (Fig. 2c). Overrepresentation of behaviourally relevant locations is not novel in the hippocampus research field. It has been previously demonstrated that hippocampal place cells fire preferentially at reward locations during goal-directed tasks. Furthermore, as the vH is strongly associated with anxiety, the anxiogenic location represents a highly salient environment for vH-dependent computations. This is supported by our finding that the anxiogenic area is overrepresented by neurons of the vH, further strengthening its role in the emotional processing of information.

Another relevant observation relates to the directional firing of vH neurons. In the non-anxiogenic configuration of the ELM, the spatial activity of single vH neurons varied depending on the direction of exploration. Similar observations have been made in both the dH and vH for linear mazes, suggesting of a common principle underlying spatial information along the dorso-ventral axis of the hippocampus. With respect to the vH, Royer et al. hypothesised that the differential activity between inbound and outbound trajectories might be caused by the reward delivered at the end of the arm implying some reward-associated value coding. In contrast to Royer et al., we placed rewards at both extremities of the ELM. Although this does not invalidate the view of Royer et al., in our study,
the direction-dependent firing was homogenised in the anxiogenic configuration of the ELM, with vH neurons exhibiting similar firing independently of the direction of exploration in the open arm (Fig. 4). We attribute the homogenisation of the direction-dependent firing in single vH neurons to the relevance of the anxiogenic location for vH-dependent computations. This is further supported by the results of the control experiments, in which we introduced a new, but not anxiogenic, environment to the ELM (Fig. 4d) leading to fewer changes in vH neuronal activity.

We further showed that the vH neuronal activity is related and even predictive of the extent of exploration of the open anxiogenic area. This view of vH function in anxiety is also supported by human studies in which an EPM was recreated in virtual reality while registering respiration, heart rate and skin conductance levels. In that study, higher anxiety levels associated with greater exploration of an open area during virtual reality navigation. This implies that anxiety levels are modulated upon the experience of an anxiogenic location, suggesting a conserved neuronal mechanism and circuit to monitor the extent of exploration during anxiety. Of particular note, in our experiments with rats, we identified a group of neurons in both the EPM and the ELM exhibiting peak firing activity at the furthest locations on the open area, indicating that different anxiety levels were reflected in the activity of vH neurons (Fig. 1f, g and Fig. 5a, b).

Furthermore, the modulation of the neuronal activity in the anxiogenic location was not exclusively observed while the rats explored the open area. The CO configuration of the ELM contained protective walls in half of the maze, while the other half was entirely open. The neuronal population activity was a good predictor of the extent of the exploration of the open location, even when rats explored the closed arm before entering to the open location. Indeed, the neuronal activity in the closed arm was sufficient to infer whether rats would perform proximal or distal explorations of the open arm (Fig. 5c, d, e). Anxiety-related modulation of vH neuronal activity in both the closed and the open locations implies that not only a direct experience of anxiety enhances neuronal activity in the vH, but also its anticipation without a direct confrontation to an anxiety-inducing situation.

Overall, we provided evidence that the neuronal dynamics within the vH are subjected to the experience of anxiety. When an anxiogenic situation was encountered, vH neurons, first, over-represented this location (Fig. 3); Second, their activity was tuned to the anxiogenic environment, impairing previous direction-dependent neuronal activity manifesting in the absence of anxiety (Fig. 4); Third, the neuronal activity of vH neurons reflected and predicted the exploration of an anxiogenic location (Fig. 5). Collectively, these results expand our view of vH function by highlighting dynamic and predictive computations during anxiety.
Methods

Experimental animals

In total, nine long Evans rats from Charles River Laboratories (male, 250–600 g), were kept in 12 h light cycles during behavioral experiments (performed during the light cycle). All experimental procedures were performed under an approved licence of the Austrian Ministry of Science and the Medical University of Vienna.

Surgery and microdrive implantation

Using isoflurane, animals were anaesthetised (induction 4%, maintenance 1–2%; oxygen flow 2 l/min) and fixed to a stereotaxic frame. The body temperature was controlled using a heating pad. Iodine solution was applied to disinfect the surgery site and eye cream was used to protect the eyes. Local anaesthetic (xylocain® 2%) was used before the incision. Saline solution was injected subcutaneously every 2 h, to avoid dehydration. Seven stainless steel screws were anchored into the skull to improve the stability of the construct, and two of the screws were placed above the cerebellum as references for the electrophysiological recordings. Next, based on the rat brain atlas, a craniotomy was performed above the ventral hippocampal area (from bregma: -4.8 mm anterior, 4.5 mm lateral, right hemisphere). After removal of the dura mater, an array of 12 independently movable, gold plated (100–500 kΩ) wire tetrodes (13 µm insulated tungsten wires, California Fine Wire, Grover Beach, CA) mounted in a custom-made microdrive (Miba Machine Shop, IST Austria) were implanted (Dorso-ventral: -6.5 mm). Paraffin wax was then applied around the tetrode array, and the lower part of the microdrive was cemented (Refobacin® Bone Cement) to the scalp. At the end, the surgery site was sutured, and systemic analgesia (metacam® 2 mg/ml, 0.5 ml/kg) was given. Animals were allowed at least 7 days of recovery time.

Mazes description and behavior

The elevated plus-maze (EPM) consisted of two closed (with protective side walls) and two open (without sidewalls) arms. The dimensions of the arms were 9 x 50 cm, the walls in the closed arms were 40 cm high, and the EPM was elevated 70 cm above the floor. Rats were placed on the EPM facing the open arm distal to the experimenter. Sessions lasted 5 - 8 min and were done at 200 lux of room light intensity.

The elevated linear-maze (ELM) consisted of a linear track of 120 cm length and 8 cm in width. The maze was elevated by 105 cm above the ground. A reward was given at both endpoints (two 20 mg sugar pellets). Three possible configurations were presented during the EPM exploration: A Closed-
Closed configuration (CC), consisting of 4 black panels acting as side walls which covered the entire length of the track and prevented the animal from experiencing the height; a Closed-Open configuration (CO) which consisted on 2 black panels acting as walls covering one half of the maze and leaving the other half completely open, resulting in an anxiogenic area; and a third configuration was called Closed-Texture (CT) which consisted on 4 black panels acting as walls covering the entire length of the maze. The difference with the CC configuration was that in half of the maze (the half which was open in the CO configuration) coloured geometrical figures were added to the sidewalls as new visual patterns and the texture of the floor was changed. The ELM sessions were composed by the presentation of the CO and CT configurations, each preceded by a CC configuration. Depending on the motivation of animals to explore, CC and CT configurations lasted 5 – 15 minutes while CO configurations lasted between 5 – 20 minutes. When a CC configuration was preceding a CT configuration, we refer to the configuration as CC2.

Tracking of the rats’ movement was monitored by triangulating the signal from three LEDs (red, blue, green) placed on the implanted headstage and recorded at 25 frames per second by an overhead video camera (Sony).

In vivo electrophysiology

Either an Axona headstage (HS-132A, 2 x 32 channels, Axona Ltd) or Intan headstage (2 x RHD32-channel headstage) were used to pre-amplify the extracellular electric signals from the tetrodes. For the Axona headstages, output signals were amplified 1000 x via a 64-channel amplifier and then digitised continuously with a sampling rate of 24 kHz at 16-bit resolution, using a 64-channel analogue-to-digital converter computer card (Axona Ltd). For the Intan headstages, signals were acquired with the RHD32 channel headstage and directly sent to the Intan 512ch/1024ch recording controller. Single-unit offline detection was performed by thresholding the digitally filtered signal (0.8 – 5 kHz) over 5 standard deviations from the root mean square in 0.2 ms sliding windows. For each single-unit, 32 data points (1.33 ms) were sampled. A principal component analysis was implemented to extract the first three components of each spike waveform for each tetrode channel51.

Spike waveforms from individual neurons were detected using the KlustaKwik automatic clustering software52. Using the Klusters software53, single units were isolated manually by verifying the waveform shape, waveform amplitude across tetrode’s channels, temporal autocorrelation (to assess the refractory period of a single-unit) and cross-correlation (to assess a common refractory period across single-units). The stability of single-units was confirmed by examining spike features over time.

Histology
To confirm the position of the recording sites, rats were deeply anaesthetised with urethane and lesions were made at the tip of the tetrodes using a 30 µA unipolar current for 5 - 10 s (Stimulus Isolator, World Precision Instruments). Then, rats were perfused with saline followed by 20 min. fixation with 4% paraformaldehyde, 15% (v/v) saturated picric acid and 0.05% glutaraldehyde in 0.1 M phosphate buffer. Serial coronal sections were cut at 70 or 100 µm with a vibratome (Leica). Sections containing a lesion were Nissl-stained. One rat, for which histological data could not be confirmed, was included based on: insertion coordinates, oscillatory LFP profile and similarity of neuronal activity.

**Firing rate maps and trajectory linearisation**

To compute firing rate maps found in figure 1, bins of 10x10 cm were created. For each bin, the total number of spikes was divided by the rat’s occupancy (in seconds): the firing rate maps were smoothed by convolving them into two dimensions with a Gaussian low-pass filter. For the EPM, by using the geometry of the maze, the centre and the arms were defined. Trajectories were then found by demarcating the consecutive tracked positions going from the furthest point reached on the arm, to the furthest point reached on the next visited arm. To linearise this position, each two-dimensional point \((x, y)\) was projected to the directional vector describing the arm to which that point belongs. Each projection was made by using the following equation:

\[
\text{Projection} = \frac{P_{x,y} \cdot D_{arm}}{||D_{arm}||}
\]

Were \(P_{x,y}\) is the position to be projected, \(D_{arm}\) is the directional vector of the corresponding arm and \(||D||\) denotes the norm of the vector \(D\). Each trajectory was composed of three parts: Starting arm, centre and ending arm. For the starting and ending arm, the activity was calculated over the space by dividing the total number of spikes on each linear bin (5 cm) over the occupancy (in seconds) on that particular bin. However, due to the different possible trajectories that the animal can follow in the centre, the activity there was divided into five fixed time bins. Then, the linear firing rate maps (activity in the starting, centre and ending arm) were smoothed by convolving them with a 1-D Gaussian function. Linear firing rate maps on the ELM were calculated by dividing the space into bins (2.5 cm each) and for each bin, the corresponding spikes of each neuron were summed and divided by the occupancy (in seconds).

**Coverage index**

The coverage term used is the same as the sparsity term used by Skaggs in 1996\(^4\). The formula is:

\[
\text{Coverage} = \frac{(\sum p_i \mu_i)^2}{\sum p_i \mu_i^2}
\]
where the spatial bins of the environment are denoted by \(i\), \(p_i\) is the probability of occupying that bin and \(\mu_i\) is the mean firing rate of the neuron in that particular bin.

**Place field similarity and direction-dependent analyses**

The place field similarity was calculated by using the Spearman correlation between the z-scored linear firing rate map of a neuron while the animal is moving in one direction and the z-scored linear firing rate map of the same neuron while the animal is moving in the other direction.

For the analyses found in the figure 4c, the procedure was the following: for each neuron, the z-scored linear firing rate map was calculated for both directions (from left to right and from right to left). Then, for each bin, the sum and the difference between the z-scored activities was calculated. All the values, for all the neurons, were separated corresponding to the bins belonging to the non-changing arm (i.e. the arm that is never opened or changed) and those values corresponding to the bins belonging to the changing arm (i.e. the arm that will be open). For the entire dataset (with all the neurons) 8 distributions are obtained: 1. Sum of the activity for the non-changing arm during CC configuration; 2. Sum of the activity for non-changing arm during CO configuration; 3. Sum of the activity for the changing arm during CC configuration; 4. Sum of the activity for the changing arm during CO configuration; 5. Difference of the activity for the non-changing arm during CC configuration; 6. Difference of the activity for non-changing arm during CO configuration; 7. Difference of the activity for the changing arm during CC configuration; 8. Difference of the activity for the changing arm during CO configuration.

**Neuronal state-space**

By using the firing rate of each neuron prior to the entry to the open arm in the CO configuration, each trial was represented by a population vector built with the activity of all the co-recorded neurons during that particular day \(T_n = \langle FR_{1n}, FR_{2n}, ..., FR_{mn}\rangle\) where FR is the firing rate, \(n\) is the total number of trials, and \(m\) is the total number of neurons.

**Classifier**

The exploration of the open arm on the ELM was divided into either proximal exploration or distal exploration using a threshold per session \(\text{Threshold} = \frac{\text{Furthest spatial bin} - \text{Nearest spatial bin}}{2}\). A support vector machine classifier with a linear kernel was used to determine if, based on the neuronal activity in the other trials, the identity of a given trial (proximal or distal exploration) was possible. To do so, in a “one-leave out cross-validation” fashion, the identity of each trial was predicted by training the classifier with the activity of all the other trials.
All calculations were made in MATLAB (Mathworks, version R2015b and R2019b) and statistical analyses were performed with MATLAB and Microsoft Excel. All the statistical tests used in this manuscript were non-parametric unless stated otherwise.

Lead contact

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Materials availability

This study did not generate new unique reagents.

Data and code availability

Any original code or additional information required to reanalyse the data reported in this paper is available from the lead contact upon request.

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Contributions

H. M-V., S.C. and T.K. contributed to experiments, data analysis and the preparation of the manuscript.

Competing interests

The authors declare no competing interests.
References


Figure 1. The activity of ventral hippocampal neurons is dynamically modulated during elevated-plus-maze exploration. 

a). Top, picture of the EPM. Bottom, the percentage of the time spent in different areas by the rats (n = 6) during the exploration of the EPM. The time spent in closed arms is significantly higher than in the open arms and the centre (p = 9.5615e-10, p = 9.5657e-10, respectively. One-way ANOVA, Tukey-Kramer for multiple comparisons) 

b). Location of the recording in the ventral hippocampus are indicated by red dots in three consecutive coronal sections (n = 47, number of rats = 8, one additional rat, for which histological location could not be confirmed, was included based on: insertion coordinates, oscillatory LFP profile and similarity of neuronal activity) 

c). Firing rate of 6 individual neurons during the exploration of the EPM. Full black lines denote a closed arm while the dotted lines indicate an open arm. Note three different anxiety-related activity patterns: increased...
firing in the open arms (left) or in the closed arms (centre) or in the centre (right). **d)** Z-transformed firing rates (colour-coded) of ventral hippocampal neurons during the exploration of the EPM, separated by trajectories and sorted by the spatial location of their peak firing activity. Dotted lines indicated the centre area. **e)** Top, density plot of the peak firing activity location for all neurons recorded during the journeys from a closed arm to the other during EPM exploration. Note the increased number of peak activity at the centre (i.e. the only open area for these trajectories). Bottom, same as on the top, but for the trajectories between a closed arm and an open arm. The dotted lines denote the beginning and end of the centre area. **f)** Z-transformed firing rates (colour-coded) of individual neurons during the exploration of the open arms. The grey line shows the peak firing activity. The insert shows the correlation between the furthest spatial bin reached (bin) and the location of the peak firing activity of the neuron during that particular trial (peak). *p*-values and *r*-values are noted in the figure (Spearman’s correlation). **g)** Histogram of correlation values (significant = black line, not significant = dotted line) for single neurons between the furthest spatial bin reached by the animal in a given trial vs. the location of the peak firing activity of the neuron during that particular trial on the EPM.
Figure 2. Removal of protective sidewalls along an elevated-linear-maze induces anxiety behaviour. 

a). ELM configurations with sidewalls along the entire track (CC, both arms closed, left), and with 
sidewalls removed for half of the track (CO, called one arm closed – one arm open, right). Note the 
presence of a non-changing arm, while the other arm changes from a closed to an open configuration. 
R, indicates locations of food reward. b). ELM configuration with both arms closed, but the floor 
texture and visual cues on inner-walls are changed in one of the arms (CT, closed and texture arm). c). 
Linearised trajectories of a rat running in the three different ELM configurations during a single 
behavioural session. The grey line denotes the centre of the linear maze and the division between the 
two arms. d). Time spent in both the non-changing and the changing arm in each configuration. 
Significant differences in the time spent appeared solely in the CO configuration (p = 1.45e-05, 
Wilcoxon signed-rank). e). More time spent in the central zone (defined as 11.5 cm around centre of 
the track) in the CO configuration in comparison to the CC and CT configurations (p = 2e-05 and p = 
0.034 respectively, one-way ANOVA, Tukey-Kramer for multiple comparisons).
Figure 3. Overrepresentation and remapping of ventral hippocampal activity during anxiety. 

a). Z-transformed firing rates (colour coded) of ventral hippocampal neurons during the exploration of the ELM and sorted by the spatial location of their peak firing activity for the three configurations: CC (left); CO (centre); CT (right). The order of neurons is sorted for each configuration independently. Dotted lines indicated the centre of the maze. Note the increased number of neurons with peak firing activity in the open arm. b). Left, comparisons of the percentage of neurons with peak firing activity located in the different arms of the CC and CO configurations of ELM. Note the significant differences of the proportion of neurons with peak firing activity in the open arm ($p = 0.001$, chi-squared test). Right, upon removal of the sidewalls, a larger proportion of neurons change the location of their peak firing activity from a previously closed to a currently opened arm ($p = 0.015$, chi-square test). c). Same analysis as in B for the CT configuration. CC2 is a configuration with sidewalls along the entire track (fully closed) explored right before the presentation of the CT configuration. d). Changes of the spatial location of the peak firing activity for individual neurons between different configurations. Each arrow
denotes the remapping of the location of the peak firing activity between the CC (base of the arrow) and the CO configurations (arrowhead). The red line indicates the centre of the linear maze. Note that the peak activity of the neurons shifted towards the open arm when changing from the CC to CO configuration (left, \( p = 0.0013, n = 133, \) Wilcoxon signed-rank). This shift was not present after change from the CC2 to the CT configuration (right, \( p = 0.3213, n = 75, \) Wilcoxon signed-rank). Comparison between the coverage index of the activity of single neurons (see methods) recorded during ELM exploration. Top, the cumulative probability of the distribution generated by subtracting the coverage indexes of the CC configuration from the coverage indexes of the CO configuration. The plot indicates that the firing rate’s coverage index of individual neurons significantly increased when the rat explored the CO configuration in comparison with the CC configuration (\( p = 8.237\times 10^{-06}, \) Wilcoxon signed-rank). Bottom, same as left but without differences between the CCT and CT configurations.
Figure 4. The direction-dependent activity of vCA1 neurons is homogenised in an anxiogenic location.

a). Neuronal activity of individual vH neurons while animals explored the ELM in both the CC and the CO configuration. Blue lines denote when animals headed towards the arm that will be open (in the case of the CC configuration) or is open (in the case of the CO configuration). Red lines denote when animals returned from this arm. Correlation values (Spearman correlation) indicate the similarity between the neuronal activities of both trajectories. b). Histograms of the firing rate maps similarity index (PFS, see methods) for the activities of single neurons calculated in the two possible directions on the ELM (left to right and right to left). PFS index is higher during the CO configuration (cyan) compared to the CC (black) configuration ($p = 7.622 \times 10^{-4}$, two-sample Kolmogorov-Smirnov test). c). Top, plots depicting the overall sum of the z-scored firing rate between the two possible trajectories (left to right or right to left) for all the neurons in the first arm (not changing arm, left of the dotted line) and the second arm (changing arm, right of the dotted line). Significant differences are encountered in the second arm between the CC configuration (black) and the CO configuration (Cyan) ($p = 2.16e-06$, Wilcoxon signed-rank test). Bottom, plots depicting the overall difference of the z-scored firing rate between the two possible trajectories (left to right or right to left) for all the neurons in the first arm (not changing arm, left of the dotted line) and the second arm (changing arm, right of the dotted line). Significant differences are encountered in both arms between the CC configuration (black) and the CO configuration (Cyan) ($p = 5.12e-12$, $p = 1.29e-04$, first arm and second arm respectively, Wilcoxon signed-rank test). d). Cumulative distribution of the PFS indexes for the CC, CO, CT and CC2 configurations. Note that during the CO configuration the PFS index of neurons is significantly higher compared to the other configurations (vs CC, $p = 7.622e-04$; vs CT, $p = 0.0033$ and vs CCT, $p = 0.0166$; two-sample Kolmogorov-Smirnov test).
Figure 5. The activity of ventral hippocampal neurons predicts the extent of exploration of an anxiogenic location.

a). z-transformed firing rates (colour-coded) of individual vH neurons during the exploration of the open arm on the ELM. The grey line indicates the peak firing activity. The inserts show the correlation between the furthest spatial bin reached (bin) and the location of the peak firing activity of the neuron during that particular trial (peak). p-values and r-values are noted in the figure (Spearman’s correlation).

b). Histogram of correlation values (significant = black line, no significant = dotted line) for single neurons between the furthest spatial bin reached by the animal in a given trial vs. the location of the peak firing activity of the neuron during that trial in the open arm of the ELM.

c). For two behavioural sessions, the furthest spatial bin visited on each trial (left) and the single-unit activity of all co-recorded neurons during the run on the closed arm projected onto their two highest principal components (right) are shown. The dotted line indicates the spatial bin set as a criterion to define two different types of trajectories: proximal (red dots) and distal (blue dots) exploration.

d). Predictions for three individual sessions. The blue line shows the furthest spatial bin reached during specific trials. The dotted line indicates the middle of the exploration of the open arm and divides the trials into proximal (left) and distal (right) explorations (used for the SVM-classifier). Dots at the right of each plot show the trial by trial accuracy of the SVM-classifier (see methods). Full dots show trials with distal explorations while circles show trials with proximal explorations. Red colour implies inaccurate...
prediction of the SVM classifier. e). Normalised distributions of the SVM-classifier performance (observed data – shuffled data) for each of the 11 sessions used. The green line marks the SVM-classifier performance on the observed data. Green dots show sessions for which the SVM-classifier successfully predicted proximal or distal exploration (higher than the 95% percentile of the shuffled data) based on neuronal firing activity during the exploration of the closed arms, while grey dots show sessions with inaccurate prediction. Box plots show median, 25th and 75th percentile.