The activity of thalamic nucleus reuniens is critical for memory retrieval, but not essential for the early phase of “off-line” consolidation

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Spatial navigation depends on the hippocampal function, but also requires bidirectional interactions between the hippocampus (HPC) and the prefrontal cortex (PFC). The cross-regional communication is typically regulated by critical nodes of a distributed brain network. The thalamic nucleus reuniens (RE) is reciprocally connected to both HPC and PFC and may coordinate the information flow within the HPC–PFC pathway. Here we examined if RE activity contributes to the spatial memory consolidation. Rats were trained to find reward following a complex trajectory on a crossword-like maze. Immediately after each of the five daily learning sessions the RE was reversibly inactivated by local injection of muscimol. The post-training RE inactivation affected neither the spatial task acquisition nor the memory retention, which was tested after a 20-d “forgetting” period. In contrast, the RE inactivation in well-trained rats prior to the maze exposure impaired the task performance without affecting locomotion or appetitive motivation. Our results support the role of the RE in memory retrieval and/or “online” processing of spatial information, but do not provide evidence for its engagement in “off-line” processing, at least within a time window immediately following learning experience.

[Supplemental material is available for this article.]
2012; Hallock et al. 2013, 2016; Layfield et al. 2015). Nonetheless, the role of RE for spatial learning remains controversial as several studies were unable to demonstrate any effects of RE lesion or inactivation on the rat ability to acquire a spatial task (Dolleman-van der Weel et al. 2009; Loureiro et al. 2012; Ito et al. 2015), but see also (Davoodi et al. 2009). Furthermore, despite the lack of evidence that RE neurons encode spatial representations, the RE output appears to affect the hippocampal spatial coding (Ito et al. 2015). The “head direction” coding by a population of RE neurons may also contribute to spatial navigation (Jankowski et al. 2014). At present, little is known about the role of RE for the HPC–PFC interaction occurring “off-line.” Experimental evidence exists that the RE may be involved in the spatial memory consolidation (Davoodi et al. 2009) and/or its long-term maintenance (Loureiro et al. 2012; Ali et al. 2017). The results from studies on aversive learning also suggest that the RE may be an intermediate processing step within the PFC–HPC pathway that enables fear memory consolidation and retrieval (Davoodi et al. 2011; Xu and Sudhof 2013). Yet, in contrast to the well-established role of the HPC–PFC interactions for both “online” and “off-line” memory-related processing, the exact role and the functional significance of RE activity within this highly interconnected neural circuit in different stages of memory formation remains insufficiently understood.

The present study was thus designed to examine the extent to which the RE is involved in the consolidation of spatial memory. To this end, we trained rats to perform a spatial task and evaluated the behavioral effects of post-learning reversible inactivation of the RE. From a methodological perspective, inactivation of a brain region after encoding phase (or “online” processing) tests its involvement in the consolidation phase, taking place “off-line.” We also tested rat spatial memory after 20-d “forgetting” period; any deviation from the rat performance on the remote memory test would indicate the involvement of post-learning RE activity in memory retention.

Rats were trained on an elevated crosswall-like maze (Fig. 1A), the configuration of which resembled a multiple-unit T-alley maze, originally designed to study rodent spatial cognition (Tolman 1948). In the beginning of each trial a rat was randomly released from one of the two start locations and allowed to reach reward by navigating along maze alleys (Fig. 1A). Two salient distal visual cues were fixed on the black curtains surrounding the maze. The experimental design, namely, availability of distal cues, variable start positions, and goal-oriented navigation along the two rather complex trajectories, assumed that the rat’s performance will be relying on allocentric cues and a path integration, both known to depend on HPC (McNaughton et al. 2006; Buzsáki and Moser 2013). Performing this maze task relying on procedural memory (e.g., acquired motor habit) is unlikely at the early stages of learning, but it may eventually prevail following extensive training (Packard and McGaugh 1996), which however did not take place in the present study. Based on the literature reviewed above, efficient learning and successful performance of such a spatial task most likely require a coordinated interplay between HPC and PFC, whereas the “off-line” HPC–PFC interactions may be essential for the stabilization of encoded information.

Our present findings demonstrate that RE activity is essential for the performance of a spatial target-oriented spatial task, however not required for the early phase of “off-line” processing enabling memory stabilization and long-term storage. In other words, the contribution of the RE is likely limited to the network that is activated “online” or during active phases of information encoding.

Results

Rats were trained on the crossword-like maze (Fig. 1A). The rat performance on the maze was evaluated using the trial latency (time required to reach reward), the trajectory length (total number of maze alleys visited by a rat before reaching reward), and the number of errors. Each deviation from the “correct” (shortest) path was considered an error, regardless of how many maze alleys the rat would cross before returning to the “correct” path (see example on Supplemental Fig. S1A). Traversing along the “correct” path, but in the opposite direction was also classified as an error (Supplemental Fig. S1B). Entering a “wrong” maze alley with all four paws was considered an error.

Immediately after each of the five learning sessions, muscimol (MUS) or saline (SAL) was injected via chronically implanted cannulas targeting the RE (Fig. 1B). On days 6 and 7, MUS or SAL was injected prior to the maze exposure to test the effect of RE inactivation on the performance of a recently acquired spatial task (Fig. 1B). On day 8, we tested the drug-free rats on the maze again, but included two probe trials to verify that rats used distal visual cues for navigation on the maze. Specifically, the first three trials were run under the standard conditions, then all lights were turned off and rats had to perform the trial 4 in the darkness. Next trials (5–7) were standard and on the trial 8 rats were tested in the

![Figure 1](image-url)
darkness and without any vertical barriers that prevented them entering some maze alleys during the initial learning (Fig. 1A); the trials 9 and 10 were again standard. On trial 4 (darkness test), rats made significantly more errors (Supplemental Fig. S2), which was suggestive that rats used allocentric, but not procedural (e.g., sequence of turns) strategy to perform the task. On trial 8 (no barriers test), all rats easily reached reward, but followed completely different trajectories from the “correct” ones (Supplemental Fig. S3); thus rats did not appear to rely on the acquired motor habit (e.g., sequence of turns), but likely used path integration. On day 9, rats were trained in the standard conditions to stabilize the task performance and were tested on the maze again after 20-d “forgetting” period (Fig. 1B).

The histological examination confirmed the location of the cannula tip in the direct proximity to the RE in 17 rats (Fig. 2). The cases with the cannula placement outside the RE served as additional control. The histology result was in agreement with a post hoc analysis of the presence or absence of a motor deficit following intrabrain MUS injection. Typically, the RE inactivation did not affect motor activity. In contrast, MUS injection outside the RE or into the third ventricle often produced a motor deficit of different degrees. To quantify motor activity, for each of 10 trials after the MUS injections we extracted the rat maximal speed using video recording and a custom software (MathWorks). The K-means clustering analysis revealed three main patterns with motor activity stable across trials (Cluster1, Supplemental Fig. S4), gradually decreasing over time (Cluster2), and severely suppressed (Cluster3). In two rats with the cannula tip placed in the RE, the onset of motor deficit was delayed, probably due to MUS diffusion outside the RE. Conservatively, the data from these two rats were excluded from the analysis of the effects of the RE inactivation on memory consolidation. Hence, the rats were distributed between three experimental conditions with MUS injections restricted to the RE (RE-MUS, n = 7), with MUS injections outside of RE (notRE-MUS, n = 7) and SAL injections (SAL, n = 16). During the first training session (before any intrabrain injections), the rats’ behavior on the maze was similar across three experimental groups (one-way analysis of variance [ANOVA]; errors: F(2,27) = 0.39, ns; trial latency: F(2,27) = 1.32, ns; trajectory length: F(2,27) = 0.24, ns).

The RE is not essential for “off-line” memory consolidation or long-term storage

To assess the effect of post-learning inactivation of the RE on the learning rate, we submitted behavioral variables across five training days to the repeated-measures ANOVA and compared across three experimental conditions. There was a significant day effect for the trial latency (F(2,7,71.6) = 154.5, P < 0.001; Greenhouse–Geisser correction was applied whenever the assumption of sphericity was violated), the trajectory length (F(2,4,65.3) = 84.3, P < 0.001) and the number of errors (F(4,108) = 184.5, P < 0.001), yet no significant interaction or group effect. The task performance gradually improved over time as reflected by decreasing number of errors, but the learning rate was equal for all three groups (Fig. 3A). Post hoc multiple comparisons (Bonferroni corrected) between training days showed that rats learned the task during four sessions; the number of errors reached an asymptote level on day 4 and there was no further improvement on day 5 (Fig. 3B). There was also no between-group difference on the last day of training (one-way ANOVA, errors: F(2,27) = 1.25, ns; trial latency: F(2,27) = 1.44, ns; trajectory: F(2,27) = 1.40, ns). Since the rats had to learn two partially overlapping trajectories (Fig. 1A), we also compared the learning rate between Start1- and Start2-trials. No trajectory preference was revealed in any experimental group by any of the behavioral variables analyzed (not shown). Finally, to evaluate the efficiency of spatial memory consolidation, we compared the rat performance during the first trial on days 2–5, which is equivalent to the conventional memory retrieval test. We found a significant day effect (latency: F(3,81) = 34.14, P < 0.001; errors: F(3,81) = 19.79, P < 0.001; trajectory: F(3,81) = 11.04, P < 0.001), but no interaction or group effect. Overall, our results did not provide any evidence that post-learning inactivation of the RE affected the learning rate. Notably, post-learning MUS injections outside the RE, while caused a transient motor deficit, did not affect the next day task performance; the latter result suggests that brain regions surrounding the RE are not likely to contribute to the “off-line” consolidation.

We also tested if post-learning inactivation of the RE affected the long-term stability of acquired memory. After the initial learning phase was completed (days 1–5), rats were additionally tested.
on the maze on days 6–9 and then were allowed 20 d of “forgetting” period, during which rats were kept in their home cages with unlimited access to food and water (Fig. 1B). At the end of the forgetting period rats were food deprived and tested again on the maze. The remote memory retrieval was evaluated by rat performance on the first trial. No intrabrain injections were made during “forgetting” period or prior to the maze exposure on day 30. Substantial forgetting was clearly evident as rats made significantly more errors on day 30 than on day 5, when the asymptote performance was reached (repeated-measures ANOVA, day effect: $F_{(1,26)} = 40.607, P < 0.001$) (Fig. 4). However, memory decay was equal in all experimental groups ($F_{(2,26)} = 0.960, \text{ns}$), there was also no significant interaction ($F_{(2,26)} = 1.886, \text{ns}$) (Fig. 4). Although, on day 30 rats made even more errors as on day 1 ($6.4 \pm 2.5$ vs. $4.9 \pm 2.9$ $P < 0.05$), more detailed analysis of rat behavior revealed some important differences. First, on day 30 rats actively explored the maze and most of them reached reward within 3-min cut-off time (SAL: 73.3%, RE-MUS: 85.7%, notRE-MUS: 42.9%). Importantly, the proportion of completed trials on day 30 was much higher than on the very first learning trial on day 1 (6.9%, all rats combined); the latter indicated that rat behavior on day 30 was clearly target-oriented. Second, the average length of correct path (traversing along the correct trajectory without deviation) was much longer on day 30 ($3.8 \pm 1.5$ maze sections) than on day 1 ($1.3 \pm 0.3$ maze sections) and, in fact, it was comparable to the performance on day 3 ($4.4 \pm 1.9$ maze sections). Thus, despite substantial memory decay on day 30, the rat behavior on the maze was not random and indicated that memory about the task was, at least, partially preserved. Finally, the trial latency on day 30 greatly varied across rats. The latencies distribution was bimodal with peaks around 63 and 162 sec. Using the K-means clustering, we assigned the rats to “slow” and “fast” performers. The “fast” performers showed a somewhat better preserved memory as they made fewer errors ($2.55 \pm 1.57$ vs. $6.39 \pm 2.55$ for “fast” and “slow” performers, respectively; $t_{(26.978)} = 5.024, P < 0.001$) and had a shorter trajectory ($18.0 \pm 5.9$ vs. $45.3 \pm 11.2$ maze sections, $t_{(27)} = 7.425, P < 0.001$). However, the proportion of “fast” performers was similar between three groups (SAL: 40.00%, notRE-MUS: 28.57%, and RE-MUS: 42.86%; Kruskal–Wallis test for independent samples, ns).

**Figure 3.** Reversible inactivation of the RE after each of the five task acquisition sessions did not affect the efficiency of spatial learning. (A) The average number of errors in each trial is shown across five training sessions for rats that received intrabrain injections of SAL (open circles, $n = 16$), MUS outside the RE (gray circles, $n = 7$), or MUS in the RE (black circles, $n = 7$). The accuracy of rat performance improved with equal rate in each experimental group. (B) The average number of errors in each trial is shown across five training sessions for all rats tested ($n = 30$). Rats reached asymptote performance on day 4. Error bars represent ±SEM; (*) $P < 0.01$ (Bonferroni corrected).

**Figure 4.** Reversible inactivation of the RE after each of the four task acquisition sessions had no effect on short- or long-term memory retention. The accuracy of task performance during the first trial on day 5 (memory retrieval test 1 d after the task acquisition) and on day 30 (remote memory retrieval) is shown for three experimental conditions. No drug injection was made during 20-d “forgetting” period or prior memory retrieval tests. Memory expression on days 5 and 30 was equal across three experimental groups. Error bars represent ±SEM.

**The RE is critical for spatial memory retrieval**

To assess the effects of RE inactivation on the retrieval of recently acquired spatial memory, we selected rats with injection centers in the RE and with no MUS-induced motor deficits during the retrieval trial ($n = 16$) (Fig. 2). Given that post-learning inactivation of the RE did not affect task acquisition (Fig. 3), we combined rats which received either SAL or MUS injections into the RE during the initial learning phase (days 1–5). After five learning sessions, on days 6 and 7 rats received either MUS or SAL injection 30 min before the maze exposure (Fig. 1B). The RE inactivation dramatically affected the rat behavior on the maze. On the first trial, 9 out of 16 rats (56.3%) did not find the reward location within a 3-min cut-off time. The first trial latency was significantly longer compared to SAL injection ($149.9 \pm 45.4$ sec vs. $41.9 \pm 38.5$ sec after MUS and SAL injection, respectively; $t_{(15)} = -8.4, P < 0.001$). Notably, the MUS-injected rats were actively exploring the maze (Fig. 5A). The
trajectory length was twice as long after the RE inactivation (33.0 ± 17.2 vs. 16.0 ± 11.1 maze sections for MUS and SAL, respectively; t(16) = −4.6, P < 0.001). Remarkably, the MUS-induced performance deficit was preserved during all 10 trials of the maze session (Supplemental Fig. S5). To quantify this observation, we restricted the analysis to rats with unaffected motor activity (Supplemental Fig. S3, Cluster 1). The number of errors was stable across trials and significantly lower on day 5 after MUS injection than after SAL injection (repeated ANOVA, trials: F(1, 419, 21.28) = 0.864, ns).

Thus, the RE inactivation caused a persistent spatial memory deficit as the task performance did not improve with repeated trials.

**Discussion**

The findings of the present study suggest that nucleus RE has a selective regulatory effect on the network that support the active phase of learning (information retrieval and encoding), but not involved in the mechanisms of systems consolidation occurring after learning or “off-line.” Specifically, we found that the RE inactivation after each of the five learning sessions did not affect the rat’s ability to acquire a spatial task, nor did it result in a faster memory decay over a 20-d “forgetting” period. In contrast, the RE inactivation in well-trained rats prior the maze exposure strongly impaired their task performance. Notably, the unaffected motor activity and appetitive motivation were both suggestive of a memory retrieval off-line.

Reuniens inactivation impairs memory retrieval in both spatial and nonspatial tasks. It is also unlikely that the RE inactivation impaired the rat ability for stimulus–response association as an earlier study showed no effect when the RE-lesioned rats were tested in a visually guided task (Hembrook and Mair 2011).

Our findings may appear to contradict the result of other studies that revealed no effects of the RE inactivation on performance of the Morris water maze (MWM) task (Loureiro et al. 2012; Cholvin et al. 2013). Yet, the discrepancy in the results may actually be explained by the task- and time-specific activation of the memory supporting large-scale network. It is well established that learning of the MWM task depends on the HPC, but does not require the PFC (de Bruin et al. 1994; Sloan et al. 2006). Notably, the recruitment of the PFC is required for the retrieval of remote (25–30 d after learning), but not recent (1–5 d) memory in the MWM (Teixeira et al. 2006; Lopez et al. 2012); the latter is consistent with time-dependent reorganization of the circuit supporting memory storage (Frankland and Bontempi 2005). Interestingly, in a modified version of the MWM task, namely, under partial-cue conditions, the PFC is required also for recent memory retrieval (Jo et al. 2007). Furthermore, when rats are tested on a double-H water maze or on a T-maze, both the HPC and the PFC are engaged in the spatial task performance (Cholvin et al. 2013; Layfield et al. 2015). Therefore, the RE inactivation appears ineffective when the HPC–PFC interaction is not required for behavioral execution. Besides, as noted by Hembrook et al. (2012), cognitive demands of the MWM maze task may be insufficient to reveal less pronounced memory deficits like, for example, in cases of lesion of the ventral midline thalamic nuclei. Collectively, our results suggest that the RE may critically contribute to the retrieval of spatial memory and, possibly, to spatial navigation; yet it does not appear to be involved at least in the early phase of “off-line” processing leading to memory stabilization. Our results also support the view that the RE is important for cognitive functions that depend on the HPC–PFC interaction (Hembrook et al. 2012; Cholvin et al. 2013, 2016; Layfield et al. 2015); memory retrieval, working memory, and spatial navigation belonging to such brain functions (Churchwell et al. 2010; Benchenane et al. 2011; Gordon 2011; Eichenbaum 2017).

The behavioral effects of the RE inactivation reported here support the idea that the RE may coordinate the PFC–HPC interactions. Specifically, gamma-range synchronization between the HPC and PFC is thought to mediate encoding and updating task-related spatial information (Spellman et al. 2015). The PFC–HPC synchrony at slower (4–12 Hz) frequency range presumably facilitates integration and maintenance of information in working memory (Jones and Wilson 2005). The spike-phase coherence and cross-frequency coupling between the HPC and PFC have been attributed to memory encoding (Siapas et al. 2005; Benchenane et al. 2010). Finally, both HPC and PFC function is critical for memory retrieval (Churchwell et al. 2010; Lopez et al. 2012; Cholvin et al. 2016).

The following mechanisms described by now in the literature can account for our results: (1) the excitatory input from the RE may facilitate the HPC–PFC coupling (Dollemen-van der Weel et al. 1997; Di Prisco and Vertes 2006; Hallock et al. 2016; Roy et al. 2017); (2) the neural activity in the CA1 may be adjusted via the HPC output to the RE (Dollemen-van der Weel et al. 1997); (3)
the RE output may affect spatial coding in the HPC and/or spatial information from the HPC to the PFC may be transferred via the RE (Jankowski et al. 2014; Ito et al. 2015). Besides, the RE, as a part of the circuitry mediating top-down control of dopamine neurons in the ventral tegmental area (Zimmerman and Grace 2016), may influence reward-motivated behaviors. Therefore, if the RE, indeed, gates the bidirectional information flow within the HPC–PFC pathway, it is not surprising that the inactivation of RE impaired the spatial task performance, which is dependent on this functional circuit.

It is also possible that the RE contributes to the systems consolidation, but the RE inactivation in our experiments was not sufficient to cause interference of “off-line” processing. Most studies are consistent in reporting that the effects of MUS reach maximum within 30 min after injection and last for at least 2 h (Martin 1991; Edeline et al. 2002; Allen et al. 2008). Depending on the injection volume and concentration, it has been shown that the effects of MUS injection can last up to 6 h (Brandon et al. 2011). In our study, we made an effort to make a rather small injection by using MUS concentration of 0.27 pg/μL and volume 0.19 μL. The acute effects of MUS injection (30–60 min) were rather robust as reflected by impaired task performance on the memory retrieval test or by motor deficit in case of injection outside the RE. The diffusion of fluorophore-conjugated MUS was ~1 mm radius, which was likely smaller than the diffusion of unlabeled MUS. Taking into account MUS diffusion over time (Edeline et al. 2002), it is unlikely that inactivation of the RE was insufficient; moreover, the adjacent brain regions including the rhomboid nucleus were likely affected by MUS. A higher concentration of MUS or a larger volume would compromise even more the spatial selectivity of the affected brain area. In our experiments, the RE activity was substantially suppressed within at least the first hour after learning when the experience-induced neuronal ensembles replay (Wilson and McNaughton 1994; Peyrache et al. 2009) and other learning-induced changes of HPC and PFC population activity occur (Eschenko et al. 2006, 2008). Consistently, several studies reported effects on memory consolidation due to manipulation of neural activity during the first hour after learning experience (Girardeau et al. 2009; Ego-Stengel and Wilson 2010; Maingret et al. 2016; Novitskaya et al. 2016). It is possible that the duration of the RE inactivation was insufficient to interfere with the mechanisms of systems consolidation or the RE activity may be critical within a delayed post-learning time window.

Finally, any behavioral study of memory unavoidably faces methodological drawbacks, which may complicate the interpretation of results. We used the crossword-like maze, which belongs to a family of multi-unit T-mazes (Tolman 1948) or can be considered as one of configurations of the Hebb–Williams maze (Hebb and Williams 1946); both mazes have been designed and intensively used for studying spatial cognition in rodents. Our experimental design fulfilled the requirements for allocentric navigation (Vorhees and Williams 2014). Successful task performance required spatial orientation at the start position, retrieval of acquired memory about the maze configuration may facilitate path integration, maintenance in working memory information about visited locations may help for updating the animal current position, and contextual decision-making would help targeted navigation. This scenario is generally consistent with the role of the RE for performance of working memory-dependent spatial tasks (Hembrook and Mair 2011; Hembrook et al. 2012; Hallock et al. 2013, 2016; Layfield et al. 2015). The aforementioned methodological concerns open a possibility that a compromised spatial behavior after the RE inactivation was due to an integrated deficit in the retrieval of spatial memory, in the working memory, in the ability to navigate or make decisions. To further clarify the nature of the behavioral deficit produced by the RE inactivation, testing animals in various situations differing by predominant cognitive demands is crucial.

In all, our findings further support the hypothesis that the RE is critical for spatial navigation and memory retrieval, yet do not provide evidence for the RE involvement in the mechanisms of systems consolidation, at least during the time window immediately following learning experience.

Materials and Methods

Male Sprague Dawley rats (Envigo, Huntingdon, UK) weighing 300–350 g at the beginning of experiment were kept in pairs with food and water ad libitum on 12 h/12 h light–dark cycle. All the experiments were performed during the dark cycle. When rat appetitive behavior was tested, rats were kept on a food-restricted diet to ensure their appetitive motivation at times of behavioral testing. On these days, in addition to the chocolate milk (0.6 mL) obtained as reward during maze exposure, each rat received 15–20 g of food pellets and unlimited access to water in their home cage. Rat weight was monitored on a daily basis and kept at ~90% of ad libitum body weight. All experimental procedures were approved by the local authorities (Regierungspräsidium Tübingen, Germany, Referat 35, Veterinärwesen) in accordance with the regional animal welfare committee pursuant to §15 of the German Animal Welfare Act (Kommission nach §15 des Tierschutzgesetzes), and were in full compliance with the Directive 2010/63/EU of the European Parliament and of the council on the protection of animals used for scientific purposes.

Surgical procedures

We performed surgeries following standard aseptic procedures. Briefly, before surgery rats were deeply anesthetized with isoflurane (4% induction, ~1.5% maintenance) and placed in a stereotaxic frame (David Kopf Instruments). Subsequently, the skull was exposed, a burr hole was drilled to target the RE using the following stereotaxic coordinates: AP/ML = −1.8/−1.5 mm (Paxinos and Watson 2005). Furthermore, a stainless-steel guide tube (22-gauge, Plastics One Inc.) was inserted 4.9 mm deep relative to the surface of the brain and at a medial-lateral angle of 10° to avoid damaging the sagittal sinus. The guide tube was fixed to the skull using dental mountain.
cement and stainless-steel anchor screws. After the end of the surgery we placed a dummy cannula inside the guide tube to prevent the brain tissue growth. Rats were allowed to have a 1-wk postsurgery recovery before behavioral testing began.

Behavioral apparatus and training procedures
The crossword maze (130 × 130 cm) was custom-built from black polyvinyl chloride. The perpendicular maze alleys (4 × 4) formed nine identical square sections (Fig. 1A). Maze alleys were 10 cm wide and had 2-cm high rims on both sides. There were two start locations and one reward port connected via tubing with a pump (Izmatec) releasing liquid reward (chocolate milk). To reduce the number of alternative routes on the maze we placed nine vertical barriers (30-cm high and 25–40-cm wide) at specific crossing points, thus restricting the access of the animal to some maze sections. The maze was elevated 80 cm above the floor and surrounded by black curtains; two posters served as extramaze visual cues. All behavioral procedures were performed under dim light.

For habituation, the maze was converted to a single linear alley. A rat was released from one end of the alley, had to reach the reward port at the other end and obtain reward by nose poking. This procedure was repeated until the rat behavior became consistently reward oriented. The start and reward locations were different from the ones used for the main task; extramaze visual cues were removed. Rats were also habituated to the intrabrain microinjection procedure by handling. After three habituation sessions, rats were trained on a spatial task for five consecutive days. Two start locations were used in pseudo-random order to minimize the procedural component of learning; the two shortest (correct) trajectories leading to reward partially overlapped (Fig. 1A). A rat was released from the start location and allowed 3 min to reach the reward port. After reward consumption or after the trial cut-off time elapsed, the rat was removed from the maze and left in a waiting box for 3–5 min. During each intertrial interval the maze alleys were wiped to minimize local olfactory cues. Each training session consisted of 10 trials. Immediately after the learning session rats received either intrabrain injection of phosphate-buffered saline (SAL-group) or muscimol (MUS-group) and returned to their home cages. On days 6 and 7, rats received either SAL or MUS injection 30 min before the maze exposure (Fig. 1B). On days 8 and 9, rats were trained on the same task (without any drug injections) to further stabilize the acquired spatial memory and then kept in their home cages for 20 d without any behavioral testing; during this “forgetting” period rats had food and water ad libitum, except 24 h before the remote memory retrieval test (Fig. 1B).

Intrabrain microinjection
The MUS powder (Sigma-Aldrich) was diluted in SAL at a final concentration of 0.27 µg/µL. The injections were performed using Hamilton syringe loaded into a microinfusion pump (UMP3, WPI). For the drug injection procedure, a rat was gently restrained by hand, the dummy cannula was removed and the injection cannula was inserted inside the guide tube. The tip of the injection cannula extended 2 mm beyond the tip of the guide tube. We used polyethylene tubing to connect the injection cannula to the Hamilton syringe. MUS or SAL (0.19 µL) was infused over 60 sec. The injection cannula was left in place for additional 2 min before it was slowly retracted. The dummy cannula was then inserted back to the guide tube.

Data analysis
The rat behavior on the maze was video recorded by an infrared camera. The trial time was measured manually using a stopwatch. The rat trajectories on the maze were reconstructed manually from video and the movement speed was extracted and further analyzed using custom made program in MATLAB (MathWorks). Any deviation from the “correct” (shortest) trajectory leading to reward was considered as an error. Specifically, if a rat entered a maze alley, which did not belong to the “correct” path, with all four limbs, it was scored as an error. Further animal passage along other maze alleys outside of the “correct” path after deviating was not scored as additional errors unless the rat returned to the correct path (see examples on Supplemental Fig. S1A). Moving along the correct trajectory, but in opposite direction (away from reward port) was also considered as an error (see examples on Supplemental Fig. S1B).

The trajectory length, the time to reach reward, and the number of errors were extracted for each trial. To assess the “content” of spatial memory from the rat trajectory, maze alleys were arbitrary divided into 24 equal-length sections separated by maze crossings; each maze section was assigned a unique number and the movement sequences analyzed. Thus, the shortest (correct) trajectory from the Start 1 (S1) consisted of nine maze sections (Fig. 1A). For each trial, we extracted the mean length of the rat trajectory (number of maze sections) overlapping with the correct path before deviation from the correct path.

Behavioral variables during five learning sessions were submitted to the repeated-measures ANOVA; post hoc comparisons were made when appropriate using Bonferroni correction. Data were tested for equality of error variance and Greenhouse-Geisser correction was applied whenever the assumption of sphericity was violated. The one-way ANOVA or Student t-test was used for between-group comparisons. The statistical significance α-value was set at P < 0.05 level. The IBM SPSS Statistics (v.22) software package was used for statistical analysis.

Perfusion and histology
At the end of behavioral experiments, we injected each rat with fluorescein-conjugated MUS (Thermo Fisher Scientific). This helped to localize the site of injection and evaluate the extent of drug diffusion. After ~30 min post-injection, the rat was euthanized with a lethal dose of sodium pentobarbital (100 mg/kg i.p.; Narcoren, Merial GmbH) and perfused transcardially with 0.9% saline followed by 4% paraformaldehyde in 0.1 M phosphate buffer (PB, pH 7.4). The brain was removed and stored in the same fixative. Before sectioning, the whole brain was placed in 30% sucrose solution at 4°C until they sank. Serial 50-µm-thick coronal sections were cut on a horizontal freezing microtome (Thermo Fisher Scientific). Every second section was Nissl-stained; adjacent sections were stored for examination under the fluorescent microscope (AxioVision, Carl Zeiss). Position of the injection cannula tip was assessed visually and digitized.

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