Research

Adult but not aged C57BL/6 male mice are capable of using geometry for orientation

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Geometry, e.g., the shape of the environment, can be used by numerous animal species to orientate, but data concerning the mouse are lacking. We addressed the question of whether mice are capable of using geometry for navigating. To test whether aging could affect searching strategies, we compared adult (3- to 5-mo old) and aged (20- to 21-mo old) C57BL/6 male mice. We established a water maze task in which spatial information is provided by one landmark proximal to the target (featural information) and by the rectangular shape of the maze (geometric information). By means of probe trials in which we manipulated the presence of these two kinds of information, we show that adult mice can use both geometry and landmark to orientate. By contrast, aged mice do not use geometry and rely exclusively on the landmark to locate the platform. This study provides the first evidence that mice are capable of using geometric information for orientation and that this ability declines in aged animals.

Place learning is one of the most common approaches to study mechanisms of memory in animals. Place learning assumes that an invisible target can be found by using allothetic navigation, namely by relying on the diverse information (visual, auditory, and olfactory cues) emerging from the milieu. Thus, specific features in the environment (featural information) can be used as landmarks to orientate. In addition to featural information, the shape of the environment (that is, geometry) can be important information used during allothetic navigation. The studies of Cheng (1986) and Margules and Gallistel (1988) were the first to demonstrate that rats can process geometry and even use it preferentially as compared with landmarks to orientate themselves in an enclosed environment. These studies have been followed by others showing that not only rats (Sakamoto and Okaichi 1996; Ramos 2000; Pearce et al. 2001; McGregor et al. 2004) but also fish (Sovrano et al. 2002, 2003), birds (Vallortigara et al. 1990; Kelly et al. 1998; Vargas et al. 2004), and monkeys (Gouteux et al. 2001) are capable of relying on geometry to orientate. Various species have been studied for their ability to navigate by means of geometric information, but there are no data concerning the mouse, although it has become an important species in cognitive neuroscience due to the generation of transgenic lines. As pointed out by Frick et al. (2000b), “mice are not little rats,” and differences have been found between the two species, notably in terms of the searching strategies used to solve the water maze task (Frick et al. 2000b; Whishaw et al. 2001). Therefore, information acquired from the rat cannot be applied to the mouse without previous experimental evidence. Similar to aged humans (Perlmutter et al. 1981; Sharps and Gollin 1987; Uttl and Graf 1993), aged mice exhibit memory deficits in spatial learning tasks (Lamberty and Gower 1993; Bach et al. 1999; Frick et al. 2000a). However, other studies showed that place learning is unaffected or only partially altered in aged mice and rats (Rapp et al. 1987; Nicolle et al. 2003), for instance when spatial information provided to animals is simplified (Lamberty and Gower 1991). These observations suggest that aging affects some but not all cognitive abilities required in a water maze task. This study aims to test whether mice are capable of using geometry to orientate and whether aging affects this ability. We designed a water maze task in which spatial information is controlled and modifiable so that, depending on the task conditions, mice are differentially exposed to featural and/or geometric information.

Results

Acquisition

Throughout the learning phase, aged mice showed slower swimming velocity than adult mice (Fig. 1A) as indicated by the significant effect of the factor age \((F_{1,32} = 94.09; P < 0.001)\). Since a slower swimming velocity can lead to higher escape latencies independently of cognitive abilities, distance covered was used to evaluate learning success. There was no effect of age nor of the interaction between age and session on distance covered during acquisition, whereas there was an effect of the factor session \((F_{3,260} = 14.75; P < 0.001)\) indicating that both aged and adult mice improved their performance by decreasing over consecutive training days the path length to reach the platform (Fig. 1B). Post hoc analyses indicated that distance covered on day 3 did not differ from distances covered during the last days of training, showing that the performance of aged and adult mice quickly reached the asymptotic level. As performances were stable during the last four learning days in both groups, we assumed that aged and adult mice had learned the task on day 6 of training. All mice showed neither floating, thigmotaxis, nor jumps out of the water throughout the experiment.

First probe trial

During the first probe trial, performed under the same conditions used during training (rectangular maze with one landmark on the short wall nearby the platform, see scheme in Fig. 2A), both aged and adult mice searched in the proximity of the platform position. To test spatial preference for the platform location, we compared the number of entries and time spent in the zone corresponding with the platform position (“T” in Fig. 2A) with three control circular zones of the size of the platform located at the three other corners of the maze (zones 1–3 in Fig. 2A). There was no effect of the factor age on number of entries and time spent in zones, whereas there was an effect of the factor zone on the number of entries \((F_{1,156} = 69.77; P < 0.001)\) and time spent in zones \((F_{1,156} = 49.00; P < 0.001)\); both groups entered more
mice in their performance in this first probe trial (Fig. 2D,E,F) for any of the parameters analyzed besides mean velocity (adult: $22.93 \pm 0.52$ cm/sec; aged: $17.67 \pm 0.58$ cm/sec) and distance covered (adult: $1310 \pm 31.64$ cm; aged: $988.3 \pm 34.58$ cm), which showed higher values in the selected adult mice as compared with the selected aged mice ($P < 0.001$ for both parameters). Interestingly, adult mice spent more time in a corridor linking the target zone and the diagonally opposite one ("T" and "Z" in scheme in Fig. 2A) as compared with aged mice (adult: $43.22 \pm 2.94$%; aged: $33.93 \pm 2.38$%; $P < 0.05$). In contrast, the two groups spent the same amount of time in the corridor linking the two incorrect zones ("1" and "3" in scheme in Fig. 2A) (adult: $30.00 \pm 1.07$%; aged: $25.16 \pm 2.16$%). The qualitative observation of the swimming paths of the selected adult and selected aged mice is in agreement with this result; in fact, in addition to a clear preference for the target zone, adult mice swam along the line connecting the target zone and its diagonally opposite one (Fig. 3A). In contrast, the swimming path of the aged group was focused in the area proximal to the target and less on the diagonal between the two geometrically correct zones (Fig. 3B). Both these quantitative and qualitative observations suggested that adult mice, but not the aged ones, orientated themselves using not only the landmark but also the shape of the maze during the first probe trial.

**Geometry probe trial**

The “geometry probe trial” was performed to test the ability of mice to use the shape of the maze to locate the platform. In this probe trial, two pairs of corners can be distinguished from each other (Fig. 2A) and spent more time (Fig. 2B) in the target zone as compared with the other three zones. There was no effect of the interaction between age and zone on time spent in zones, whereas there was an effect on entries into zones ($F_{1,156} = 3.20$; $P < 0.05$); adult mice entered more often into the target zone than aged mice. Since no difference was observed in time spent in the target zone between the two groups, it is likely that the higher number of entries into the target zone shown by adult mice as compared with aged mice was due to differences in swimming velocity and distance covered. Indeed, as observed during learning and in all following probe trials, adult mice swam faster (adult: $22.65 \pm 0.41$ cm/sec; aged: $18.39 \pm 0.36$ cm/sec; $P < 0.001$) and covered longer distances (adult: $1294 \pm 24.38$ cm; aged: $1043 \pm 21.90$ cm; $P < 0.001$) than aged mice. They thus increased their chance to cross the target platform as compared with aged mice. The evidence that aged and adult mice equally preferred the target zone was confirmed by the analysis of the two preference indices, measures that are independent of distance covered and swimming velocity. There was no difference between aged and adult mice for these indices calculated for entries and time. Both indices were higher than chance level in the two groups, indicating that adult and aged mice preferentially searched the platform at its correct location (Fig. 2C). This first probe trial was also used to select, for the further trials, those mice that displayed high spatial selectivity for the platform by reaching the criterion that we had determined (i.e., mice that spent at least twice as much time in the target platform as compared with the second preferred platform). Less than 50% of the aged mice reached our criterion (15 mice out of 32) versus nearly 70% of the adult animals (15 mice among 22) ($\chi^2$ test: $P = 0.122$; Fisher exact probability test: $P = 0.166$). No difference was detected between the selected aged and adult mice in their performance in this first probe trial (Fig. 2D,E,F) for any of the parameters analyzed besides mean velocity (adult: $22.93 \pm 0.52$ cm/sec; aged: $17.67 \pm 0.58$ cm/sec) and distance covered (adult: $1310 \pm 31.64$ cm; aged: $988.3 \pm 34.58$ cm), which showed higher values in the selected adult mice as compared with the selected aged mice ($P < 0.001$ for both parameters). Interestingly, adult mice spent more time in a corridor linking the target zone and the diagonally opposite one ("T" and "Z" in scheme in Fig. 2A) as compared with aged mice (adult: $43.22 \pm 2.94$%; aged: $33.93 \pm 2.38$%; $P < 0.05$). In contrast, the two groups spent the same amount of time in the corridor linking the two incorrect zones ("1" and "3" in scheme in Fig. 2A) (adult: $30.00 \pm 1.07$%; aged: $25.16 \pm 2.16$%). The qualitative observation of the swimming paths of the selected adult and selected aged mice is in agreement with this result; in fact, in addition to a clear preference for the target zone, adult mice swam along the line connecting the target zone and its diagonally opposite one (Fig. 3A). In contrast, the swimming path of the aged group was focused in the area proximal to the target and less on the diagonal between the two geometrically correct zones (Fig. 3B). Both these quantitative and qualitative observations suggested that adult mice, but not the aged ones, orientated themselves using not only the landmark but also the shape of the maze during the first probe trial.
Geometry and navigation in adult and aged mice

other using the geometric properties of the maze, since the corners connected by one diagonal of the rectangular maze are identical to each other in terms of geometric properties. Therefore, the corner in which the platform was located during training and the diagonally opposite corner are both considered correct during the geometry probe trial ("T" in Fig. 4A), whereas the other two corners are considered incorrect ("O" in Fig. 4A). Adult mice, but not the aged ones, showed a preference for the two geometrically correct target zones versus the two incorrect opposite zones (the absolute values for the two target corners and the two opposite corners, respectively, were added up and used for statistical analysis and graphical representation of entries into zones, time in zones, and preference indices; Fig. 4). There was an effect of the interaction between age and zone on number of entries into zones ($F_{1,28} = 8.05; P < 0.01$) and a tendency on time spent in zones ($F_{1,28} = 2.72; P = 0.11$): While aged mice did not show any difference between entries made and time spent in the target zones as compared with the opposite zones, adult mice entered more often into and spent more time in the target zones than in the opposite ones. In addition, adult mice entered into the target zones more often than aged mice; conversely, they spent less time and tended to enter less often ($P = 0.06$) into the opposite zones as compared with aged mice (Fig. 4A,B). Since aged mice swam more slowly (adult: $22.22 \pm 0.52$ cm/sec; aged: $16.73 \pm 0.66$ cm/sec; $P < 0.001$) and covered less distance (adult: $1265 \pm 31.63$ cm; aged: $940.80 \pm 39.33$ cm; $P < 0.001$) as compared with the adult ones, the results described above could be misleading. Indeed, due to their slower velocity, aged mice had less chance to enter into and to spend time in the correct areas as compared with the adult mice. However, the comparisons between preference indices, which are independent of swimming velocity and distance covered, confirmed the differences observed between groups, as preference indices for entries and time in target zones were higher for adult mice than for aged mice. Moreover, the indices for entries and time in target zones of the adult group, but not of the aged group, were higher than chance level (Fig. 4C). Thigmotaxis and floating were not observed in the adult group, but not of the aged group, were higher than chance ($P < 0.001$) as compared with the adult mice. However, the comparisons between preference indices, which are independent of swimming velocity and distance covered, confirmed the differences observed between groups, as preference indices for entries and time in target zones were higher for adult mice than for aged mice. Moreover, the indices for entries and time in target zones of the adult group, but not of the aged group, were higher than chance level (Fig. 4C). Thigmotaxis and floating were not observed in the two groups. The better ability of adult mice to use geometric information as compared with aged mice was also shown by the spatial histograms representing the swimming paths (Fig. 3C,D). Interestingly, adult mice seemed to swim mostly along the diagonal connecting the two correct corners.

**Landmark probe trial**

Apart from the fact that the aged group swam more slowly (adult: $20.64 \pm 0.55$ cm/sec; aged: $15.90 \pm 0.50$ cm/sec; $P < 0.001$) and covered less distance than adult mice (adult: $1142 \pm 34.21$ cm; aged: $882.3 \pm 29.81$ cm; $P < 0.001$), no difference between groups was found in the "landmark probe trial," and both groups explored preferentially the correct target zone (Figs. 3E,F, 5). There was no effect of age on entries into zones, but a tendency for an effect on time spent in zones ($F_{1,28} = 4.05; P = 0.054$): Aged mice spent in general more time in the four zones than adult mice. A significant effect of the factor zone was found on entries ($F_{3,84} = 18.57; P < 0.001$): Both aged and adult mice entered more often into (Fig. 5A) and spent more time in (Fig. 5B) the target zones more often than aged mice; conversely, they spent less time and tended to enter less often ($P = 0.06$) into the opposite zones as compared with aged mice (Fig. 4A,B). Since aged mice swam more slowly (adult: $22.22 \pm 0.52$ cm/sec; aged: $16.73 \pm 0.66$ cm/sec; $P < 0.001$) and covered less distance (adult: $1265 \pm 31.63$ cm; aged: $940.80 \pm 39.33$ cm; $P < 0.001$) as compared with the adult ones, the results described above could be misleading. Indeed, due to their slower velocity, aged mice had less chance to enter into and to spend time in the correct areas as compared with the adult mice. However, the comparisons between preference indices, which are independent of swimming velocity and distance covered, confirmed the differences observed between groups, as preference indices for entries and time in target zones were higher for adult mice than for aged mice. Moreover, the indices for entries and time in target zones of the adult group, but not of the aged group, were higher than chance level (Fig. 4C). Thigmotaxis and floating were not observed in the two groups. The better ability of adult mice to use geometric information as compared with aged mice was also shown by the spatial histograms representing the swimming paths (Fig. 3C,D). Interestingly, adult mice seemed to swim mostly along the diagonal connecting the two correct corners.

**Figure 3.** Spatial histograms of the mouse locations during the probe trials. Schemes of the maze and rainbow scales are shown on the right side for each probe trial. (A,C,E,G) Adult group; (B,D,F,H) aged group. (A,B) "First probe trial." The target is depicted with a thin dotted circle on the histograms and "T" in the corresponding scheme. (C,D) "Geometry probe trial." The two target zones are depicted with black line circles on the histograms and "T" in the corresponding scheme. (E,F) "Landmark probe trial." The target location is depicted with a thick dotted circle and "T" in the corresponding scheme. (G,H) "Competition probe trial." Geometric zones (black line circles, histogram; "G" in the corresponding scheme); landmark zone (thick dotted circle, histogram; "L" in the corresponding scheme); incorrect zone (thin white circle, histogram; "I" in the corresponding scheme).
zone as compared with the other three zones. No effect of the interaction between age and platform was detected for any parameter. The analysis of preference indices did not reveal any difference between aged and adult mice, and these indices were higher than chance level in both groups (Fig. 5C).

**Fourth probe trial**

The results from the fourth probe trial, which aimed to check whether mice were still motivated to find the platform and focused their search at the correct location after the “landmark” and “geometry” probe trials, replicated those obtained in the first probe trial (data not shown). Both groups preferentially searched the platform at its correct location as indicated by number of entries into and time spent in zones and by the corresponding preference indices. There was no difference between aged and adult mice, with the exception that adult mice swam faster and covered longer distances than aged mice.

**Competition probe trial**

During the “competition probe trial,” the landmark was located on the right side of one of the long walls, instead of on one of the short walls as during acquisition (see scheme in Fig. 6A). Thus, if featural information, namely the landmark, was used to locate the platform, then mice would have searched in the “landmark” zone (L). If geometric information, i.e., the rectangular shape of the maze, was used, then mice would have searched in the “geometry” zones (G). (The mean of the two “geometry” zones was used for statistical analysis and graphical representation of entries into zone, time in zone, and preference indices.) The remaining zone (I) was incorrect regarding information provided either by geometry or landmark. As during the previous probe trials, neither thigmotaxis nor floating was observed, indicating that aged and adult mice actively searched the platform. Despite a slower velocity (adult: 21.73 ± 0.65 cm/sec; aged: 16.85 ± 0.53 cm/sec; P < 0.001) and a shorter distance covered (adult: 1219 ± 41.05 cm; aged: 950.10 ± 31.85 cm; P < 0.001) in the aged group as compared with the adult group, there was no effect of age on either entries into zones or time spent in zones, but there was an effect of the interaction between age and zone on the preference index for time (F2,56 = 4.63; P < 0.05) and on the entry index (F2,56 = 7.92; P < 0.001). Aged mice entered more often into and spent more time in the landmark zone than in the geometry or incorrect zones, while there was no difference between the geometry and incorrect zones, indicating that aged mice preferentially used the landmark to navigate (Fig. 6A,B). In contrast, adult mice searched the platform by using both geometry and landmark: They entered into and spent time in the geometry zones as much as in the landmark ones and entered into the incorrect zone less frequently than into the geometry zone and, although not significantly, into the landmark zone (P < 0.07). Adult mice also tended to spend less time in the incorrect zone as compared with the geometry zones (P < 0.09). In addition, adult mice entered less frequently into and spent less time in the landmark zone as compared with aged mice (Fig. 6A,B). To further test age-dependent differences in the preference for the geometry and landmark zones, we compared preference indices for entries and time calculated for landmark, geometry, and incorrect zones in aged and adult mice (Fig. 6C,D). There was no effect of age but a tendency for an effect of the interaction between age and zone on the preference index for entries (F2,56 = 1.94; P = 0.15) and a significant effect on the index for time (F2,56 = 3.97; P < 0.05). Post hoc analysis revealed that in aged mice, the preference indices for entries and time for the landmark zone were higher than those for the geometry and incorrect zones. Moreover, the preference indices for entries and time for the landmark zone were higher than chance level, whereas those for the geometry zones were equivalent to chance level, confirming that aged mice preferentially searched in the landmark zone but not in the geometry zones (Fig. 6C,D).
Geometry and navigation in adult and aged mice

During the "competition probe trial," adult mice (solid bars in A, B and left side of C, D) flexibly relied either on landmark or on geometry, while aged mice (open bars in A, B and right side of C, D) followed only the landmark indication. (A, inset) A scheme of the maze. (Thick bar) Landmark position, (G) zones designed by geometry, (L) zone indicated by landmark, (I) zone that was incorrect regarding landmark and geometric information. Values represent means plus S.E.M. +P < 0.05, ++P < 0.01, as compared with the incorrect zone within each group; $$$P < 0.01, $$$$P < 0.001, as compared with the geometry zones within the aged group; *P < 0.05, **P < 0.01, ***P < 0.001, as compared with the adult group within the same zone (Newman-Keuls post hoc after a significant effect of the interaction between group and zone); #P < 0.05, ##P < 0.01, ###P < 0.001, as compared with chance level (25%) (Wilcoxon signed-rank test).

Control probe trial

We performed a final probe trial to test whether uncontrolled cues could have influenced the spatial selectivity of the mice. In the absence of landmark and of geometric information, mice did not show any preference for any of the four corners, indicating that no uncontrolled cue had influenced the searching strategies throughout the experiment. There was an effect of the factor age on number of entries (F₁,25 = 4.32; P < 0.05) (Fig. 7A) but not on time spent in zones (Fig. 7B); Adult mice crossed the four zones more often than the aged, probably as a consequence of their higher swimming velocity (adult: 19.81 ± 0.65 cm/sec; aged: 14.99 ± 0.44 cm/sec; P < 0.001) and longer distance covered (adult: 1084 ± 40.61 cm; aged: 826.50 ± 25.22 cm). However, neither effect of zone nor effect of the interaction between group and zone was found for any parameter.

Discussion

This study was designed to assess whether mice are capable of using geometry, in this case the shape of an enclosed environment, to orientate themselves and, if so, whether they can interchangeably use geometric and featural information, namely a prominent landmark. In addition, we wanted to test whether the use of geometric and featural information changes with aging. We showed that adult mice flexibly use geometric and featural information to navigate and do not display preferred strategy, while aged mice do not use geometry and rely on featural information.

Mice process geometry and use it for navigating

Our data demonstrate that adult mice can use the shape of the environment to find a hidden goal, indicating that they detect

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<th>Table 1. Individual performance of adult and aged mice in the competition probe trial</th>
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<td><strong>Entries into zone (n)</strong></td>
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*Geometry zone values correspond to the mean of number of entries into and time spent in the two geometrically correct zones.

**Landmark zone.

***Incorrect zone.

According to chance, time spent in one zone would be 0.5%.
the geometric properties of the environment and use them to navigate. How this information is used is, however, unclear. Three possibilities are conceivable to interpret these observations. First, mice process the global shape of the environment (i.e., “this environment is a rectangle”). Second, they could rely only on the local properties of the environment (i.e., “the platform is situated in the corner that presents a long wall on the right hand”). Or, finally, they could use a basic procedural strategy by learning to turn left when they are facing a long wall. We exclude this last possibility: Adult mice swam along the line connecting the two geometrically correct corners, suggesting that they are capable of extrapolating the diagonal of the rectangular maze by processing either the global shape of the environment or its local properties.

It is noteworthy that geometric information was acquired by mice even if it was not crucial to solve the task. In fact, during learning, the landmark provided a more precise information than geometry since it indicated without any ambiguity the platform position among the four possible corners; in contrast, geometric information led to two possibilities, namely the target corner and its diagonally opposite one. These results add further to those obtained in other species and provide evidence for the first time that mice perceive and are capable of using geometry for orientation. In addition, we show that in mice, as in other species such as fish (Sovrano et al. 2003), chicks (Vallortigara et al. 1990), pigeons (Vargas et al. 2004), and rats (Pearce et al. 2001; Wall et al. 2004), the presence of a landmark does not prevent animals from acquiring geometric information. From an ecological point of view, it is likely that geometry is more important than landmarks: As argued by Sovrano et al. (2002), in wildlife the large-scale shape of the environment provides stable and thus reliable information to orientate, as opposed to featural information such as vegetation that undergoes seasonal changes. This could explain that mice learn to use geometry to orientate also when it is superfluous, as in our study in which the platform was reliably indicated by a prominent landmark.

Interestingly, in a conflictual situation in which geometry and landmark provide contradictory indications, mice rely flexibly on one or the other information. As shown by the analysis of the individual performance during the “competition probe trial,” some mice rely preferentially on the landmark while others use geometry, and some switch between the two. These results are in contrast with those demonstrating that a preference emerges in conflictual situations as observed in pigeons (Vargas et al. 2004; but see also Kelly et al. 1998), chicks (Vallortigara et al. 1990; Tommasi et al. 2003), and monkeys (Gouteux et al. 2001). It has been proposed that the size of the apparatus could be important for choosing between geometric and landmark information, as observed in young children who preferentially use landmarks in large environments and geometry in small ones (Learnmonth et al. 2002). Sovrano et al. (2005) and Vallortigara et al. (2005) have also suggested that this phenomenon could occur in fish and chicks. In our case, it could be argued that our maze has intermediate dimensions, preventing a clear preference for geometry or landmark information in adult mice.

Aging impairs the flexible use of geometry and landmark for navigating

No difference could be detected between adult and aged mice during learning and the first probe trial performed in the presence of congruent landmark and geometric information. Interestingly, the difference between the two groups became apparent when we manipulated spatial information (landmark and/or geometry). In fact, aged mice were impaired when only geometry was accessible but not when the landmark was present. This inability to use geometry for navigation was confirmed in the “competition probe trial,” in which aged mice followed only the information indicated by the landmark. It has been shown previously that cognitive decline differs among individuals (Bach et al. 1999; Schulz et al. 2002; Nicholson et al. 2004; but see also Frick et al. 1995). In our experiment we observed few aged mice performing as well as adult ones in the “geometry probe trial.” Nevertheless, the majority of the aged group was not able to use geometry to navigate, an inability that is unlikely due to lower motivation, swimming impairments, or sight problems, since these mice performed well when the landmark was available. It is possible that the impairment of aged mice is due to their altered stress response, as it is well known that stress can affect the learning rate (for reviews, see Bowman 2005; Shors 2006). Indeed, aged animals display a dysregulation of the hypothalamo-pituitary-adrenal axis (Dalm et al. 2005); notably, they show longer responses to stressors as compared with adult animals (Bizon et al. 2001). However, the stressfulness of an experimental paradigm also depends on the control that animals have on the stressor. For instance, Kavushansky et al. (2006) showed that a forced swim test, but not a standard Morris water maze task, increases the plasma level of corticosterone in rats as compared with home-cage control rats. Thus, if being placed in a water tank can be stressful for mice, the possibility of finding a platform strongly reduces the stressfulness of the task. Indeed, we did not observe any stress-related behavior in the two groups during all experiments, such as floating or thigmotaxis. On the contrary, aged and adult mice actively searched for the platform and stopped on it quietly until they were removed from the apparatus.

We cannot exclude the possibility that all aged mice would have learned to use geometry with extensive training, meaning that age-dependent cognitive decline prevents only a fast acquisition of geometric information. Moreover, it is possible that in the presence of both landmark and geometry, aged mice simply rely directly on the easiest information for them (i.e., the landmark). Therefore, it is conceivable that geometry is overshadowed or blocked by featural information specifically in aged but not in adult mice. Indeed, it has been reported that place learning is blocked in aged rats previously trained with a cued platform (Rapp et al. 1987).

Electrophysiological and lesion studies demonstrated that the hippocampus plays a key role in geometry-based navigation (Mulder and Kubie 1987; Sakamoto and Okaichi 1996; Lever et al. 2002; Tommasi et al. 2003; McGregor et al. 2004; Vargas et al. 2004). More specifically, the hippocampus could be important for processing local metric properties such as angles and dis-
tances to create a cognitive representation of the environment’s shape. Indeed, it has been demonstrated that hippocampal place cells are sensitive to the length of a wall and to the distance from a wall (O’Keefe and Burgess 1996) and that the hippocampus is essential for processing metric properties of the environment (Sakamoto and Okaichi 1997). However, the hippocampus is not the only structure implicated in this cognitive process, since it has been shown that hippocampal lesions do not prevent cells of other structures (i.e., the postsubiculum and the anterior thalamic nucleus) to be sensitive to changes in the environmental shape (Golob and Taube 1997). We tend to consider that the impaired ability of aged mice to navigate by means of geometry is related to age-dependent hippocampal alterations (for review, see Rosenzweig and Barnes 2003). Notably, aged mice could be incapable of processing metric properties of the environment such as the length of the walls, which is crucial information for defining the shape of the rectangular maze. It is also conceivable that, although being capable to detect and process geometric information, aged mice are not able to use this information to navigate. In any case, it is possible that an impaired function of the hippocampus is compensated for by a stronger activation of other brain regions that are also involved in navigation, with the result that the particular form of navigation controlled by these structures will be predominantly used. For instance, aged mice tend to use preferentially cue-based strategies (Nicolle et al. 2003), which are mainly controlled by the striatum (Packard and McGaugh 1996; Oliveira et al. 1997; Colombo et al. 2003). It is interesting in this context that rats displaying a poorer performance in the spatial version of the Morris water-maze task show higher expression of NMDA receptors in the striatum (Nicolle et al. 1996). We therefore hypothesize that aged mice are not capable of using geometry to navigate due to an impaired hippocampal function, and that this inability is compensated for by activation of other regions, such as the striatum, that allows aged mice to navigate by means of cue-based strategies.

Conclusions

This study shows for the first time that mice are capable of detecting and using geometry during navigation, an ability that degrades with aging. Whereas adult mice can face environmental changes by relying either on geometry or landmarks to orientate, aged mice depend on featural information. More specific experiments are, however, needed to gain insights into the mechanisms underlying navigation based on geometry. Place cells recordings, lesion studies, and pharmacological studies in behaving mice could be a means to achieve this goal. Finally, the water maze task established in this study can be a sensitive tool not only to investigate spatial learning and memory processes in wild-type animals, but also to characterize cognitive abilities of mutant mice.

Materials and Methods

Animals and husbandry

C57BL/6j male mice were transferred from a pathogen-free breeding facility into a vivarium with an inverted 12 h light/12 h dark cycle (lights off at 7:00 a.m) and maintained in groups of two to five siblings under standard housing conditions (23 ± 1°C, 40%-50% humidity, food and water ad libitum). After 2 wk of acclimatization, mice underwent the experiments, which were performed between 9:00 and 15:00 in a room adjacent to the vivarium illuminated by red light. At the beginning of the experiment, 32 mice composed the aged group (20- to 21-mo old) and 22 mice composed the adult group (3- to 5-mo old).

Apparatus

A rectangular maze (126 × 66 cm and 50 cm high) made of white polyvinylchloride easily convertible into a square (66 × 66 cm and 50 cm high) was used. The maze could be rotated on its central axis. A white platform (diameter 7.4 cm; 1 cm below the water surface) was placed in a fixed corner with its center located 18 cm from the two walls. The apparatus was filled up to 9 cm from its bottom with water (20°C ± 1°C) made opaque by the addition of non-toxic white paint. To provide supplemental markers of the shape of the apparatus, black metal stripes (10 cm high and 6 cm wide) were located at the four corners of the maze, 20 cm above the water surface. A black and white landmark (21 cm high and 50 cm wide) was placed in the middle of the short wall proximal to the platform, 8 cm above the water surface. This maze was circled by dark curtains hanging from the ceiling to the floor. The ceiling was covered by a white curtain with a hole in its center to allow the view from a video camera placed behind the curtain. Light was provided by four white light neon tubes placed on the floor around the swimming pool, which provided a homogeneous illumination of the maze (50 lux on the water surface). Electronic equipment for video recording and automatic tracking of the swimming paths of the mice was placed in a room adjacent to the experimental one and illuminated by dim red light. Mice were kept in this room between trials while waiting to be tested.

Protocol

Before the experiment started, mice were familiarized for 3 d to swim and climb onto a platform (four trials per day; 60 sec maximal duration; 10-min inter-trial interval). To facilitate learning, familiarization took place in a different room and apparatus (42.5 × 26.5 cm and 15.5 cm high, circled by a black curtain) than that used for learning in the water maze. The position of the platform (diameter of 10 cm, 1 cm below the water surface) was unpredictable, since its location was randomized and we did not provide any information that could guide the animals. Thus, mice had to swim to random escape from the water. From experimental day 1 to day 6, mice were trained in the rectangular maze described previously (see scheme in Fig. 1) with four trials per day (inter-trial interval of 10 min). Four starting positions (two were situated in the middle of the long walls and two at the most distant corners to the platform) were pseudo-randomized and the apparatus was rotated by 90° between trials, clockwise or counter-clockwise depending on the day, to avoid that mice could use extra-maze cues (e.g., sounds or odors) to orientate. Animals were released into the water facing the wall and were allowed 60 sec to find the platform (“P” in scheme in Fig. 1) and climb onto it. After staying on the platform for 30 sec, mice were returned to their home cage and kept warm under red light until the next trial. From days 7–11, mice underwent one probe trial (60 sec in duration) per day. During each probe trial, the platform was removed and mice were released from the center of the maze. To avoid extinction, mice underwent three learning trials starting 10 min after the probe trial under the same conditions as those used during learning on days 1–6. Depending on the day, spatial information provided during the probe trial was manipulated. The first probe trial (day 7) took place with the same spatial information available during learning (geometry and landmark information were the same as during learning, see scheme in Fig. 2) and aimed to evaluate spatial selectivity for the platform position. Only mice showing spatial selectivity for the platform position (determined by a criterion described below under “Behavioral Analysis”) underwent the probe trials performed from days 8–11. On days 8 and 9, the so-called “landmark” and “geometry” probe trials were carried out. During the “geometry probe trial,” the landmark was removed, while the maze had the same rectangular shape as during learning (see scheme in Fig. 4); under these conditions the geometrically correct platform position was in the two corners indicated with “T.” Indeed, the corners identical to each other in terms of geometric properties are those connected by one diagonal of the rectangular maze. Therefore, the corner in which the platform was located during training and the diagonally opposite corner are both considered correct during this probe trial. Conversely, during the “landmark probe trial,” geometric informa-
tation was absent since the rectangular maze was converted into a square while the landmark was still present on one of the four walls (see scheme in Fig. 5); based on the landmark, the correct position of the platform was in corner “L”. To avoid possible bias due to the temporal order of the two probe trials, half of the mice from each age group were tested in the “landmark probe trial” on day 8 and in the “geometry probe trial” on day 9, whereas the other half was tested in the “geometry probe trial” on day 8 and in the “landmark probe trial” on day 9. To control whether mice still had a spatial preference for the target after the “landmark” and “geometry” probe trials, a fourth probe trial was performed on day 10, which took place under the same conditions used for the probe trial on day 7. On day 11, a “competition probe trial” was performed in the rectangular maze, where both landmark and geometric information were available but provided conflicting information since the landmark was placed on one of the long walls of the maze (see scheme in Fig. 6). In fact, based only on the landmark, the platform should be searched at corner “L”, while the geometrically correct corners would be those indicated as “O”. Finally, we performed a “control probe trial” on day 12, where both landmark and geometric information were removed (see scheme in Fig. 7) and mice were expected to equally search in all corners.

Behavioral analysis

All trials were video recorded, and animal paths were digitalized and analyzed with the software Ethovision (Noldus). For the learning trials, velocity, distance covered, escape latency, and thigmotaxis (time spent in an 8-cm width zone that borders the wall) were calculated. For all the probe trials, we calculated number of entries into and time spent in four circular zones of size of the platform located at the previous platform location and at its virtual mirror positions in the three other corners. Distance covered, velocity, and thigmotaxis were also calculated for all probe trials. In addition, we calculated two preference indices for the target platform that are independent of swimming velocity and distance covered. One index (preference index for entries) corresponded to the total number of entries into the target zone relative to the total number of entries into all zones; the second index (preference index for time) was calculated as the previous one, but using time spent in zone as the parameter. For the first probe trial, we evaluated the spatial preference of mice and selected for the further trials only those mice reaching a criterion of high spatial selectivity for the target zone; namely, mice that during the probe trial spent at least twice as much time in the target platform as compared with the second preferential platform. We calculated also for this first probe trial the time spent in a corridor linking the target zone and its diagonally opposite one. This corridor was 20 cm in width and did not include the two platform zones. For a qualitative analysis of the swimming paths of the two groups during the probe trials, we generated spatial histograms using the software IgorPro (WaveMetrics).

Statistical analysis

Learning trials were analyzed with a two-way ANOVA for repeated measures (having “age” as the between-groups factor and “session” as the within-groups factor), followed by Newman-Keuls post hoc analysis when appropriate. Number of entries and time spent in zones calculated during probe trials were also analyzed with a two-way ANOVA for repeated measures (having “age” as the between-groups factor and “zone” as the within-groups factor), followed by Newman-Keuls post hoc analysis when appropriate. These data are presented as means plus S.E.M. T test comparison between groups was performed using Student’s t-test. Correlation analysis using Pearson correlation coefficient was performed to test the relationship between variables. For all analyses, statistically significant differences were considered for p < 0.05. In vitro autoradiography of iotopic glutamate receptors in hippocampus and striatum of aged Long-Evans rats: Relationship to spatial learning. Neuroscience 74: 741–756.

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