One-trial perceptual learning in the absence of conscious remembering and independent of the medial temporal lobe

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A degraded, black-and-white image of an object, which appears meaningless on first presentation, is easily identified after a single exposure to the original, intact image. This striking example of perceptual learning reflects a rapid (one-trial) change in performance, but the kind of learning that is involved is not known. We asked whether this learning depends on conscious (hippocampus-dependent) memory for the images that have been presented or on an unconscious (hippocampus-independent) change in the perception of images, independently of the ability to remember them. We tested five memory-impaired patients with hippocampal lesions or larger medial temporal lobe (MTL) lesions. In comparison to volunteers, the patients were fully intact at perceptual learning, and their improvement persisted without decrement from 1 d to more than 5 mo. Yet, the patients were impaired at remembering the test format and, even after 1 d, were impaired at remembering the images themselves. To compare perceptual learning and remembering directly, at 7 d after seeing degraded images and their solutions, patients and volunteers took either a naming test or a recognition memory test with these images. The patients improved as much as the volunteers at identifying the degraded images but were severely impaired at remembering them. Notably, the patient with the most severe memory impairment and the largest MTL lesions performed worse than the other patients on the memory tests but was the best at perceptual learning. The findings show that one-trial, long-lasting perceptual learning relies on hippocampus-independent (nondeclarative) memory, independent of any requirement to consciously remember.

Significance

Studies of memory have established a distinction between ordinary recollection of the past (declarative memory), which depends on medial temporal lobe (MTL) structures and other (nondeclarative) forms of memory that are expressed through performance and depend on other brain systems. One phenomenon that has eluded classification is one-trial perceptual learning, whereby a degraded image of an object, which is difficult to identify, becomes recognizable after a single exposure to the original image. This effect was fully intact in memory-impaired patients with hippocampal or larger MTL lesions and persisted undiminished for more than 5 mo, despite impaired memory for the test format and the images themselves. Perceptual learning is MTL-independent (nondeclarative) and occurs without a requirement to consciously remember.

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improved when degraded images had been followed at study by matching, intact images but not when they had been followed by nonmatching, intact images. Finally, the patients were not only as successful at naming degraded images as controls at each of the three delays, they also were as confident as controls in their responses to the degraded images (patients: 3.0 ± 0.4, 3.1 ± 0.4, 3.0 ± 0.4 at 1 d, 7 d, and 5.4 mo, respectively; controls: 2.9 ± 0.3, 3.1 ± 0.3, and 3.0 ± 0.2).

It is of interest that the ability to identify the degraded images that had been followed at study by nonmatching, intact images improved a little across test sessions (Fig. 3), presumably because the same degraded images were presented a total of four times: at study and at each of three delays [analysis of linear trend across the four tests; controls, F(1, 10) = 7.9, P = 0.02; patients, F(1, 4) = 14.4, P = 0.02]. This finding reflects a gradual, albeit modest, improvement in the ability to perceive degraded images simply as a result of repeated exposure to them. Note that this nonspecific effect was distinct from the robust, long-lasting facilitation of naming that was specific to, and dependent on, single exposures at study to matching, intact images that revealed what was depicted in the degraded images.

Fig. 4 shows directly the amount of facilitation of naming in each group at each delay. At each delay, the percent correct identification score for the 20 degraded images that had been followed at study by nonmatching, intact images was subtracted from the percent correct identification score for the 20 degraded images that had been followed at study by matching, intact images. The patients exhibited overall as strong a facilitation as controls, and the facilitation in each group persisted for at least 5.4 mo. These findings were documented by an ANOVA (Group and Delay), which yielded no effect of Group [F(1, 14) = 0.0, P = 0.90], no effect of Delay [F(2, 28) = 2.5, P = 0.10], and no interaction of Group × Delay [F(2, 28) = 2.6, P = 0.09]. The finding that the Delay effect and the Group × Delay interaction nevertheless approached significance likely reflects the fact that the facilitation exhibited by controls weakened across time [documented by an analysis of linear trend, F(1, 10) = 8.9, P = 0.01]. In contrast, the facilitation exhibited by the patients was sustained without decrement for as long as 5.4 mo [F(1, 4) = 0.0, P = 0.91]. The controls, but not the patients, may have drawn in part on declarative, conscious memory in order to name degraded images at the shorter delays. That is, shortly after study, controls not only perceived degraded images successfully but also explicitly remembered some of the solutions. This advantage became less available to controls as the delay increased but was not available to patients at any delay.

Remembering.

Test 1. Despite the fact that patients improved at identifying degraded images for as long as 5.4 mo after a single exposure to the corresponding, intact images, the patients had difficulty remembering intact images after only 1 d (Fig. 5, Left). Controls scored 98.0 ± 1.1% correct 1 d after studying 20 intact images, but patients scored only 79.5 ± 4.2% correct [t(14) = 5.86, P < 0.001; d* = 3.7 ± 0.1 and 2.1 ± 0.2, respectively]. The controls were also more confident of their responses than patients [4.8 ± 0.1 vs. 4.2 ± 0.2; t(14) = 2.82, P = 0.01].

Test 2. The patients had difficulty remembering facts about the naming test, which they had last encountered 6 d earlier (Fig. 5, Right). Whereas the controls scored 81.8 ± 3.9% correct on multiple-choice questions about the naming test format (chance = 33.3%), the patients scored only 55.0 ± 9.4% correct [t(14) = 3.18, P = 0.007].
impaired at remembering the format of the naming test (Fig. 5, Right). Notably, patient G.P., who has the most severe memory impairment and the largest MTL lesions, performed worse than the other four patients on the memory tests but was the best of the patients at perceptual learning (Fig. 4).

In a direct comparison of perceptual learning and remembering, at 7 d after learning patients and controls with identical testing histories took either a naming test with degraded images or a recognition memory test with degraded images. The patients improved as much as controls on the naming test but were severely impaired on the memory test (Fig. 6). Thus, patients could identify the degraded images, benefiting as much as controls from having previously seen the “solutions,” but the patients could not recognize the degraded images as familiar.

The successful performance of patients with MTL lesions suggests that the one-trial and long-lasting learning demonstrated here relies on nondeclarative (hippocampus-independent) memory. As a result of their experience, patients improve at perceiving degraded images but without remembering them. Note that at early intervals after learning controls may draw on declarative memory, thereby further improving their performance. In an earlier study (3), performance was much higher at 15 min after learning than after 1 d and continued to decline a little from 1 d to 5.4 mo (Fig. 4). In contrast, the patients exhibited consistent performance across the same time intervals, falling a little short of the controls at 1 d (albeit, not significantly), and even exceeding the controls by a little after 5.4 mo. Presumably, at shorter retention intervals,

That is, prior to the 7-d memory test, participants first tried to name 40 degraded images, 20 of which were followed by the intact, matching image, and they then took a naming test at 1 d. Not surprisingly, scores on these tests largely recapitulated the scores in Fig. 3. At study, controls correctly named 34.5 ± 4.2% of the 20 degraded images that were followed by intact, matching images and correctly named 28.6 ± 4.1% of the other 20 degraded images. For patients, the corresponding scores were 29.0 ± 7.0% and 25.0 ± 3.9% correct (compare to Fig. 3). At 1 d, the controls correctly named 59.1 ± 5.0% of the 20 images that had been matched at study and 31.8 ± 5.0% of the other 20 degraded images (patients, 47.0 ± 10.1% and 24.0 ± 6.6% correct) (compare to Fig. 3). The amount of facilitation at 1 d was robust and similar in the two groups (controls, 27.3 ± 5.2%; patients, 23.0 ± 4.9%) (compare to Fig. 4). Finally, the patients and controls expressed similar confidence in their responses throughout this stage of testing (patients at study, 2.9 ± 0.4; at 1 d, 3.0 ± 0.4; controls at study, 2.6 ± 0.3; at 1 d, 3.0 ± 0.2).

The finding of interest was obtained at 7 d after study, instead of taking another naming test, participants now took a yes/no recognition memory test for the 40 degraded images and correctly named 47.0 ± 10.1% of the other 20 degraded images (patients, 47.0 ± 10.1% and 24.0 ± 6.6% correct) (compare to Fig. 3). At 1 d, the controls correctly named 59.1 ± 5.0% of the 20 images that had been matched at study and 31.8 ± 5.0% of the other 20 degraded images (patients, 47.0 ± 10.1% and 24.0 ± 6.6% correct) (compare to Fig. 3). The amount of facilitation at 1 d was robust and similar in the two groups (controls, 27.3 ± 5.2%; patients, 23.0 ± 4.9%) (compare to Fig. 4). Finally, the patients and controls expressed similar confidence in their responses throughout this stage of testing (patients at study, 2.9 ± 0.4; at 1 d, 3.0 ± 0.4; controls at study, 2.6 ± 0.3; at 1 d, 3.0 ± 0.2).

The ability to identify degraded images substantially improved after single, brief exposures to the original, intact images. The improvement was evident at 1 d and persisted for more than 5 mo (Fig. 3). Memory-impaired patients with MTL lesions exhibited this effect at full strength and without decrement across the same time period (Fig. 4). Improved perceptual performance was unrelated to the ability to remember the images that had been presented. Thus, despite a robust and long-lasting improvement at identifying previously studied degraded images, the patients were severely impaired at remembering intact images even after 1 d (Fig. 5, Left), and in another session were...
controls can remember some of the images and benefit by using declarative memory in the naming task.

Informal demonstrations using a single pair of images, as in Figs. 1 and 2, often begin with failure to identify the hidden object, followed some time later by confident and successful identification, as if the ability to identify degraded images might typically move from 0% correct initially to 100% correct on a later test. However, performance on tests involving multiple images does not behave this way. First, participants inevitably identify some degraded images spontaneously without benefit of seeing the solutions (see scores at study in Fig. 3) (3). Second, while presentation of the original, intact image directly after the degraded image often results in insight as to what the degraded image represents, sometimes participants do not see the connection between a degraded image and its original, so that there is no basis for identifying the degraded image when it is presented later. In our study, naming performance improved to 45.7% correct, averaged across both groups and three retention intervals (Fig. 3). The study by Ludmer et al. (3) reported a similar performance score across the interval 1 d to 21 d after learning.

An early hint of good performance by memory-impaired patients on perceptual learning came from a task in which drawings of common objects were first presented briefly in fragmented form and then in progressively more recognizable form until the object was identified (24, 25). When tested a second time, both patients and controls identified the objects at an earlier point in the series. Even the noted patient H.M. (26, 27) improved his performance when tested after 1 h, though he did not remember having taken the test before. However, the controls performed far better than the patients. As noted (25), the short retention interval and the small number of objects likely allowed a substantial contribution of declarative memory to task performance. That is, the controls were likely advantaged because they could remember some of the solutions or have available in memory many of the correct names, and thereby be aided in their guessing. Accordingly, at the time of this work it seemed possible that conscious remembering might be an important part of perceptual learning, and it was unclear if this should count as an example of learning that lies outside the province of the MTL, as had been demonstrated a few years earlier in the case of motor skill learning (28).

The present study demonstrates that when the possible contribution of declarative memory is limited by using a large number of images and long retention intervals, robust one-trial perceptual learning relies fully on nondeclarative memory. Participants are not asked to remember anything and are asked only to report what they see. Perceptual learning occurs without conscious control (8) and independent of any requirement to consciously remember. Brain activity elicited by successfully identified degraded images is sharpened in regions of the neocortex, including in the ventral visual stream (29). A similar idea involving sharpening has been suggested to underlie perceptual priming (30, 31). This pattern of activation is distinct from the activity associated with the same degraded images when they are not identified (5, 32, 33), and by 800 ms after image onset is similar to the activity associated with the corresponding, intact images (34). These cortical changes underlying one-trial perceptual learning occur independently of the MTL.

Methods

Participants. Five memory-impaired patients participated, who have also been studied previously (33) (mean age = 66.0 ± 8.2 y; mean education = 13.1 ± 0.8 y). Four have bilateral lesions thought to be limited to the hippocampus (CA fields, dentate gyrus, and subiculum complex), and one (G.P.) has larger MTL lesions (Table 1). For the five patients, the summed score for delayed recall (30 min) of two short prose passages (Weschler Memory Scale-Revised, WMS-R) averaged 1.2 segments (25 segments per passage). The average score for delayed reconstruction (10 to 15 min) of a complex diagram (36) was 5.8 (maximum score = 36). Paired-associate learning of 10 unrelated noun–noun pairs summed across each of three successive trials was 3.0 pairs (30 pairs total). Eleven healthy controls (four females) also participated (mean age = 72.8 ± 2.5 y; mean education = 14.4 ± 0.7 y). They scored 28.5 for the prose passages, 19.6 for the diagram, and 24.6 for paired-associate learning.

Patients D.A. and G.W. became amnestic in 2011 and 2001, respectively, following a drug overdose and associated respiratory failure. K.E. became amnesic in 2004 after an episode of ischemia associated with kidney failure and toxic shock syndrome. L.J. (the only female) became amnesic during a 6-mo period in 1988 with no known precipitating event. Her memory impairment has been stable since that time. G.P. has severe memory impairment resulting from viral encephalitis in 1987.

Estimates of MTL damage were based on quantitative analysis of magnetic resonance images from the patients and from 19 age-matched, healthy males for K.E., G.W., and G.P., 11 age-matched, healthy females for patient L.J. (37), and 8 younger healthy males for D.A. Patients D.A., K.E., L.J., and G.W. have an average bilateral reduction in hippocampal volume of 35%.
Fig. 6. Remembering in contrast to naming. The percent correct naming scores are reproduced from the 7-d test in Fig. 4, which shows the amount of facilitation in naming (i.e., how much the naming of degraded images benefited from earlier presentation of their intact, matching images). G.P.’s score was 23.3%, the best of all the patients. For remembering (d’), the procedure at study and after 1 d was the same as for the naming test (Fig. 3). However, at 7 d after study, instead of taking another naming test, participants took a yes/no recognition memory test for the 40 old degraded images and 40 new degraded images. G.P. obtained a d’ score of 0.3 (55.0% correct), the poorest of all the patients. CON = 11 controls; MTL = 5 patients with MTL lesions. Error bars show SEM.

Materials. Images in grayscale and degraded images were constructed as previously described (8). Briefly, images were generated from photographs of single, real-world animate and inanimate objects selected from the Caltech database, the Pascal VOC database, and online search engines. Using MATLAB, images were first constructed in grayscale by resizing the original image to 9 × 9 cm and 500 × 500 pixels, and then applying a box filter (initially set at 10 × 10 pixels) for low-pass spatial filtering. Black-and-white degraded images were generated by thresholding the grayscale image to binarize it into black or white pixels. The threshold was set at the median intensity of each image. Images were then selected that were judged difficult to identify but that could be identified correctly when they were compared to the matching grayscale image. Next, groups of 40 to 60 degraded images were screened in pilot testing to construct sets of 40 where the mean probability of identification was 20 to 30% and where identification of each image improved after the matching, intact image was presented; 320 different images were used in the experimental conditions described below.

Procedure.

Naming. The task began with nine practice trials in which a degraded image was presented on a computer screen, followed after 1 to 2 s of blank screen by its matching, intact image. Participants tried to name each image. The experimenter then presented the degraded image again, directing attention to the relationship between the two images. On a final (10th) practice trial, the degraded image was followed by a nonmatching, intact image, and the experimenter explained that sometimes the degraded image and the intact image would not match in this and all other tasks. The experimenter pressed a key to advance to the next item.

Participants were next told that they would see new images on the screen for 6 s each and should name each image, guessing if necessary. They were also asked to provide a confidence rating after each response (1 to 5 scale; 1 = pure guess, 5 = very confident). Participants then saw 40 degraded images: 20 were followed by the matching, intact image and 20 others, intermixed with the first 20, were followed by a nonmatching, intact image (6 s per image with a 1- to 2-s blank screen between images). Which 20 images were paired with their matching image and which 20 were paired with a nonmatching image was balanced across participants. The following day, again after 7 d, and again after 4.2 to 7.5 mo (mean = 5.4 mo), participants took four practice trials and then saw the same 40 degraded images for 6 s each with instructions to name them and provide a confidence rating. For each testing session (at study and at delays of 1 d, 7 d, and 5.4 mo), three different orders of the 40 degraded images were available, and these were assigned pseudorandomly across participants. No feedback (correct, incorrect) was given for any of the tests. The full task (study + three delays) was given a total of three times using different sets of material across a period of 1.5 y.

Remembering.

Test 1. Participants saw 20 novel grayscale images on a computer screen, each for 6 s followed by a blank screen for 1 to 2 s. They were instructed to name the images and to remember them for a later test. One day later, they took a naming test described above. Thus, participants began by trying to name 40 new images: 20 were followed by a nonmatching, intact image (6 s per image with a 1- to 2-s blank screen between images). After each response, participants provided a confidence rating from 1 to 5 (1 = pure guess, 5 = very confident).

Test 2. Participants were presented with eight three-alternative, multiple-choice questions about the format of the naming test: for example, what color was the computer screen after the item disappeared (off-white, dark gray, or black)? At the time of this test, participants had previously encountered the naming test either 9 or 10 times, most recently 6 d earlier.

Remembering in contrast to naming. To contrast remembering and naming directly, a remembering test was constructed using new materials. Up to 7 d after study, the experimental design and procedure were identical to the naming test described above. Thus, participants began by trying to name 40 degraded images and provide confidence ratings (1 to 5) for their responses. Twenty images were followed by the matching, intact image and 20 others, intermixed with the first 20, were followed by a nonmatching, intact image. Which 20 images were paired with their matching images and which were

Table 1. Characteristics of memory-impaired patients

| Patient | Age (y) | Education (y) | WAIS-III IQ | Attention | Verbal | Visual | General | Delay |
|---------|---------|---------------|-------------|-----------|--------|--------|---------|-------|
| D.A.    | 37      | 12            | 95          | 104       | 90     | 91     | 90      | 56    |
| K.E.    | 78      | 13.5          | 108         | 114       | 64     | 84     | 72      | 55    |
| L.J.    | 82      | 12            | 101         | 105       | 83     | 60     | 69      | <50   |
| G.W.    | 60      | 12            | 108         | 105       | 67     | 86     | 70      | <50   |
| G.P.    | 73      | 16            | 98          | 102       | 79     | 62     | 66      | 50    |

WAIS-III is the Wechsler Adult Intelligence Scale-III and WMS-R is the Wechsler Memory Scale-Revised. The WMS-R does not provide numerical scores for individuals who score <50. The IQ score for D.A. is from the WAIS-IV.
paired with a nonmatching image was balanced across subjects. The following day, again just as in the test of naming, participants saw the same 40 degraded images with instructions to name them and provide confidence ratings. Then, at 7 d after study, instead of taking another naming test, participants took a yes/no recognition memory test for the 40 old images intermixed with 40 new images. Which 40 images served as the 40 old images and which served as the 40 new images was balanced across participants. Memory performance at 7 d after study was compared to naming performance at 7 d after study (Fig. 4). Part of this taking this remember test this for all other testing was completed.

1. K. M. Dallenbach, A puzzle-picture with a new principle of concealment. Am. J. Psychol. 64, 431–433 (1951).
2. P. B. Porter, Another puzzle-picture. Am. J. Psychol. 67, 550–551 (1954).
3. R. Ludmer, Y. Dudai, N. Rubin, Uncovering camouflage: Amygdala activation predicts long-term memory of induced perceptual insight. Neuron 69, 1002–1014 (2011).
4. R. J. Dolan et al., How the brain learns to see objects and faces in an impoverished context. Nature 389, 596–599 (1997).
5. C. González-García, M. W. Flounders, R. Chang, A. T. Baria, B. J. He, Content-specific activity in frontoparietal and default-mode networks during prior-guided visual perception. eLife 7, e36068 (2018).
6. T. D. Albright, On the perception of probable things: Neural substrates of associative memory, imagery, and perception. Neuron 74, 227–245 (2012).
7. S. Gorlin et al., Imaging prior information in the brain. Proc. Natl. Acad. Sci. U.S.A. 109, 7935–7940 (2012).
8. R. Chang, A. T. Baria, M. W. Flounders, B. J. He, Unconsciously elicited perceptual prior. Neurosci. Consious. 2016, niv008 (2016).
9. P. Graf, L. R. Squire, G. Mandler, The information that amnesic patients do not forget. J. Exp. Psychol. Learn. Mem. Cogn. 10, 164–178 (1984).
10. E. Tulving, D. L. Schacter, Priming and human memory systems. Science 247, 301–306 (1990).
11. D. L. Schacter, L. A. Cooper, M. Tharan, A. B. Rubens, Preserved priming of novel objects in patients with memory disorders. J. Cogn. Neurosci. 3, 117–130 (1991).
12. C. B. Cave, L. R. Squire, Intact and long-lasting repetition priming in amnesia. J. Exp. Psychol. Learn. Mem. Cogn. 18, 509–520 (1992).
13. C. B. Cave, Very long-lasting priming in picture naming. Psychol. Sci. 8, 322–325 (1997).
14. C. L. Wiggs, J. Weisberg, A. Martin, Repetition priming across the adult lifespan—The long and short of it. Neuropsychol. Dev. Cogn. B Aging Neuropsychol. Cogn. 13, 308–325 (2006).
15. L. R. Squire, Memory and the hippocampus: A synthesis from findings with rats, monkeys, and humans. Psychol. Rev. 99, 195–231 (1992).
16. L. R. Squire, S. M. Zola, Structure and function of declarative and nondeclarative memory systems. Proc. Natl. Acad. Sci. U.S.A. 93, 13515–13522 (1996).
17. L. R. Squire, S. Zola-Morgan, The medial temporal lobe memory system. Science 253, 1380–1386 (1991).
18. E. H. Eichenbaum, N. J. Cohen, From Conditioning to Conscious Recollection: Memory Systems of the Brain (Oxford University Press, New York, 2004).
19. L. R. Squire, S. Zola-Morgan, Memory: Brain systems and behavior. Trends Neurosci. 11, 170–175 (1988).
20. D. L. Schacter, E. Tulving, “What are the memory systems of 1994?” in Memory Systems 1994, D. L. Schacter, E. Tulving, Eds. (The MIT Press, Cambridge, MA, 1994), pp. 1–39.
21. L. R. Squire, Memory systems of the brain: A brief history and current perspective. Neuropsychol. Learn. Mem. 82, 171–177 (2004).
22. H. F. Crovitz, M. T. Harvey, S. McClanahan, Hidden memory: A rapid method for the study of amnesia using perceptual learning. Cortex 17, 273–278 (1981).
23. V. S. Ramachandran, “2-D or not 2-D.—That is the question” in The Artful Eye, R. L. Gregory, J. Harris, Eds. (Oxford University Press, Oxford, 1994) pp. 249–267.
24. E. K. Warrington, L. Weiskrantz, New method of testing long-term retention with special reference to amnesic patients. Nature 217, 972–974 (1968).
25. B. Milner, S. Corkin, H. L. Teuber, Further analysis of the hippocampal amnesic syndrome: 14 year follow-up study of H.M. Neuropsychologia 6, 215–234 (1968).
26. W. B. Scoville, B. Milner, Loss of recent memory after bilateral hippocampal lesions. J. Neurol. Neurosurg. Psychiatry 20, 11–21 (1957).
27. L. R. Squire, The legacy of patient H.M. for neuroscience. Neuron 61, 6–9 (2009).
28. B. Milner, “Les troubles de la memoire accompagnant des lesions hippocampiques bilaterales” in Physiologie de l’Hippocampe, B. Passouant, Ed. (Centre National de la Recherche Scientifique, Paris, 1962), pp. 257–272.
29. C. González-García, B. J. He, A gradient of sharpening effects by perceptual prior across the human cortical hierarchy. J. Neurosci. 41, 167–178 (2021).
30. K. Grill-Spector, R. Henson, A. Martin, Repetition and the brain: Neural models of stimulus-specific effects. Trends Cogn. Sci. 10, 14–23 (2006).
31. C. L. Wiggs, A. Martin, Properties and mechanisms of perceptual priming. Curr. Opin. Neurobiol. 8, 227–233 (1998).
32. A. M. van Loon et al., NMDA receptor antagonist ketamine distorts object recognition by reducing feedback to early visual cortex. Cereb. Cortex 26, 1986–1996 (2016).
33. P. J. Hsieh, E. Vul, N. Kanwisher, Recognition alters the spatial pattern of fMRI activation in early retinotopic cortex. J. Neurophysiol. 103, 1501–1507 (2010).
34. M. W. Flounders, C. González-García, R. Hardstone, B. J. He, Neural dynamics of visual ambiguity resolution by perceptual prior. eLife 8, e41861 (2019).
35. C. N. Smith, L. R. Squire, Awareness of what is learned as a characteristic of hippocampus-dependent memory. Proc. Natl. Acad. Sci. U.S.A. 115, 11947–11952 (2018).
36. R. A. Osterricht, Le test de copie d’une figure complexe. Arch. Psychiat. 30, 206–356 (1944).
37. J. J. Gold, L. R. Squire, Quantifying medial temporal lobe damage in memory-impaired patients. Hippocampus 15, 79–85 (2005).
38. N. L. Rempel-Clover, S. M. Zola, L. R. Squire, D. G. Amaral, Three cases of enduring memory impairment after bilateral damage limited to the hippocampal formation. J. Neurosci. 16, 5233–5255 (1996).
39. E. Franko, A. M. Insauti, E. Artacho-Pérala, R. Insauti, C. Chavoix, Identification of the human medial temporal lobe regions on magnetic resonance images. Hum. Brain Mapp. 35, 248–256 (2014).
40. R. Insauti et al., MR volumetric analysis of the human entorhinal, perirhinal, and tempopolar cortices. AJNR Am. J. Neuroradiol. 19, 659–671 (1998).
41. Z. J. Urgolites, C. N. Smith, L. R. Squire, Eye movements support the link between conscious memory and medial temporal lobe function. Proc. Natl. Acad. Sci. U.S.A. 115, 7599–7604 (2018).
42. L. Fei-Fei, R. Fergus, P. Perona. Learning generative visual models from few training examples: an incremental Bayesian approach tested on 101 object categories. IEEE CVPR 2004, Workshop on Generative-Model.