Short report

Preservation of musical memory in Alzheimer’s disease

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SUMMARY An 82 year old musician with Alzheimer’s disease (AD) showed a preserved ability to play previously learned piano compositions from memory while being unable to identify the composer or titles of each work. He also showed a preserved ability to learn the new skill of mirror reading while being unable to recall or recognise new information. Both anterograde and retrograde procedural memory may be relatively spared in AD.

Recent investigations of memory functioning in both normal and amnesic subjects distinguish between procedural memory or information based on skills learned implicitly and without awareness, and declarative memory or information based on specific facts acquired explicitly and with deliberate intention. Studies have shown that procedural and declarative memories are not uniformly impaired in subjects with neurological disease. Amnesic subjects with Korsakoff’s syndrome demonstrated normal learning of the skills required to read reversed mirror print, despite profound amnesia for the words they had read and for the details of the test. Previous studies documented preservation of anterograde procedural memory shown by the ability to acquire new skills. We report the preservation of retrograde procedural memory in a patient with Alzheimer’s disease who despite severe impairment in declarative memory showed intact ability to play previously learned piano compositions from memory.

At the age of 77, this subject volunteered to participate in a longitudinal study of risk factors for dementia and was tested annually for the next seven years. He started to complain of memory problems two years before entry into the study.

Case report

The subject was a musicologist with over 12 years of formal musical training who had worked for over 40 years as a music editor. Throughout his adult life he had practised the piano two hours a day. Concerned about instructing his children in music, he made his family listen to classical music every evening during dinner, and quizzed his children on the composer of each composition.

His past medical history recorded a head injury with concussion one year before entry into the study. In the fourth year of the study he was involved in a motor vehicle accident with transient alteration in consciousness. CT performed two days after the accident was normal. A family history showed that his father became demented at the age of 70.

Table 1 summarises his neuropsychological test results in years 1 to 7. When the study began, his score on recall 1 of the Full Object Memory Evaluation (FOME) showed marked impairment in recent verbal memory. (Normal subjects, ages 75–85 score 7–8 items correct on this test). Impairment in verbal memory was corroborated by his score of 24/144 possible items on a 12-item, 12-trial selective reminding test. He scored a 5 on the Blessed test because he was unable to recall the memory phrase. He scored within the superior to very superior range on all subtests of the WAIS. He easily solved all 12 items in set A of the Ravens coloured progressive matrices, a test of spatial reasoning ability. His ability to copy line drawings was intact.

Verbal IQ remained stable until year 6; performance IQ declined a year earlier. An impairment in confrontation naming was noted in year 6. At that time, cued recall on a 16 item list learned under controlled conditions was quite impaired.

Table 1 summarises his neuropsychological test results in years 1 to 7. WAIS-PIQ = performance IQ from the Weschler Adult Intelligence Scale (WAIS). WAIS-VIQ = verbal IQ from the WAIS. FOME = score on recall 1 on the Full Object Memory Evaluation. Raven = score on set A of the Raven coloured progressive matrices. Sum recall = recall on a 12-item, 12-trial selective reminding test.

<table>
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<th>Year</th>
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<th>WAIS-PIQ</th>
<th>WAIS-VIQ</th>
<th>FOME</th>
<th>Raven</th>
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NA = Not available.
In years 5 and 6 he showed intact skill learning as measured by the speed of reading mirror reversed words. Nonrepeated words presented in session 2 were read faster than nonrepeated words presented in session 1 (t = 3.29, p < 0.01, year 5; t = 3.33, p < 0.01, year 6). In contrast, he failed to show priming effects in reading words which were repeated across sessions. Words which were repeated in session 2 were not read faster than nonrepeated words in session 2 (t = 1.26, year 5; t = 0.68, year 6). Recognition for both repeated and nonrepeated words was poor; the usual advantage for repeated words was not observed.

In year 5 his ability to recognize and play well known works of classical music was tested. The first two minutes of commercial recordings of six well known works were played on a cassette player. The patient was unable to recall the composer or title of any of the six works including the first movement of Beethoven's fifth symphony. Given a forced choice test with four alternatives per musical selection, he made one correct selection in six attempts. Next he was brought to the piano, and the examiner (HC) would play the first few bars of fifteen different pieces of popular classical music. In thirteen cases, the patient was able to continue playing the music as it had originally been written. He had, however, no idea of the composer of any of this music, nor could he select the correct composer on a forced choice test. Furthermore, if told the name of the composer, he could not identify the title of the music nor select it from a list of four possible titles.

In year 5, head CT, EEG, and routine blood studies for dementia evaluation were normal. He was diagnosed as having an amnesic disorder because he had no evidence of cognitive dysfunction other than memory impairment. Two years later, when intellectual functioning declined and impairment in confrontation naming became apparent, his diagnosis was changed to AD. In year 8, he could continue to play the passages on the piano but only for a few bars.

Discussion

Although there have been several previous demonstrations that neurological disease can dissociate declarative and procedural memory, our present case is the first to demonstrate that the dissociation occurs for both anterograde and retrograde memories. Impaired anterograde declarative memory was shown by his very poor scores on tests of free recall, cued recall, and recognition for new information. Impaired retrograde declarative memory was shown by an inability to recall or recognize previously learned classical compositions. In contrast, there was relative preservation of anterograde and retrograde procedural memory. Though unable to recall or recognize previously learned musical compositions, he could continue playing these compositions after cueing. However, cueing was not helpful in aiding retrieval of declarative memories. Relative preservation of anterograde procedural memory was demonstrated by his ability to acquire the skill of mirror reading.

This dissociation between procedural and declarative memory has been shown in other Alzheimer patients who could learn new procedures involving tapping, rotary pursuit, and pattern learning while showing no declarative memory for the new information. Parkinson's disease produces the opposite pattern with subjects performing poorly on procedural memory tests, but well on declarative memory tests. Korsakoff and temporal lobectomy amnesic patients show a pattern of deficits similar to our subject. Huntington patients show a mixed pattern of deficits.

Based on the pattern of dissociation caused by these different neurological diseases, the anatomical substrate of procedural memory is presumed to involve striato-prefrontal circuits, whereas the substrate of declarative memory is presumed to involve circuits connecting the hippocampus, subiculum, and association cortex. Much of the pathology in AD is found in these latter regions which is probably responsible for the impairment in declarative memory in AD patients. The skill of playing previously learned piano compositions from memory represents a special kind of procedural memory and is stored at least in part in the neostriatum, an area of the brain spared until the latest stages of AD.

References