A patient with Korsakoff syndrome of psychiatric and alcoholic etiology presenting as DSM-5 mild neurocognitive disorder

Georgios Nikolakaros, Timo Kurki, Arttu Myllymäki, Tuula Ilonen

Introduction

Thiamine deficiency can cause Wernicke’s encephalopathy (WE) and chronic cognitive impairment, Korsakoff syndrome (KS). Alcohol abuse may result in thiamine deficiency, but conditions unrelated to alcohol may cause malnutrition and WE. Major depression with associated malnutrition may cause WE in non-alcoholic patients or in the presence of alcohol abuse. KS has usually been associated with alcoholic WE, and only recently it has been fully appreciated.
that also non-alcoholic WE can cause KS.\textsuperscript{4,6,8,22} To diagnose KS, the DSM-IV\textsuperscript{23} and DSM-5\textsuperscript{24} require functional impairment. Recently, the need for a comprehensive definition of KS has been noted.\textsuperscript{5}

The core neuropsychological feature of KS is memory impairment.\textsuperscript{2,3,5,25} Executive functions are impaired in alcoholic KS,\textsuperscript{26–32} but they may be normal in non-alcoholic KS.\textsuperscript{4,6,33} It has been recently reported that in alcoholic KS the executive functions of shifting and updating are affected, whereas inhibition may be spared.\textsuperscript{32}

Conventional brain MRI of patients with neuropsychologically documented non-alcoholic KS has been normal in most cases, but frontal lobe and vermis atrophy have been reported.\textsuperscript{4} Mammillary body (MB),\textsuperscript{34–38} central,\textsuperscript{38–41} vermis,\textsuperscript{40,41} and thalamic atrophy\textsuperscript{38,42} have been shown in patients with cognitive symptoms following non-alcoholic WE. Alcoholic KS is associated with brain atrophy in cortical areas (particularly the frontal lobes), the MBs, the amygdala, the thalamus, the hippocampus, the corpus callosum, and the cerebellum.\textsuperscript{43,44}

Diffusion Tensor Imaging (DTI) in KS has shown abnormalities in frontotemporal tracts (uncinate, cingulum), the fornix, the corpus callosum, the inferior longitudinal fasciculus, and the corona radiata.\textsuperscript{4,6,45,46}

We describe Wernicke-Korsakoff syndrome (WKS) in a patient with major depression and alcohol use disorder. We describe neuropsychological, MRI, and DTI findings. We discuss functional impairment and the diagnostic use of the DSM-5.

**Material and methods**

**Clinical description**

This male patient has had panic disorder and generalized anxiety disorder since the age of 31, and major depression since the age of 51. The patient had been using large amounts of alcohol since his teen years. He had been on long-term disability leave from his job as shop manager. There was no history of traumatic brain injury or substance abuse other than alcohol.

At the age of 54, the patient’s depression worsened for several months. He had poor appetite, and on several occasions, he did not eat anything for two to three consecutive days. His weight dropped by 11 kg. During this period of malnutrition, heavy alcohol use continued. The patient was examined at the neurology outpatient clinic because of pain and weakness in the lower limbs. There was ataxia (wide-based walking, abnormal heel-shin test) and tremor, muscle weakness, and muscle wasting of the lower limbs. The patient walked with the help of a rollator. Electromyoneurography and muscle biopsy of the lower limbs were normal.\textsuperscript{4} Brain MRI showed some small diffuse vascular lesions. Cerebrospinal fluid examination was normal except for a slightly elevated protein concentration. Concentrations of gamma-glutamyl transpeptidase and carbohydrate-deficient transferrin were elevated (76 U/L and 48\%, respectively), magnesium was low-normal (0.73 mmol/L, normal range 0.7–1.1), whereas the rest of the blood tests (full blood count, sodium, potassium, calcium, thyroid tests, creatine kinase, alkaline phosphatase, 25-hydroxy vitamin D, ferritin, B12-vitamin, folate) were normal. The diagnosis was leg weakness, and the patient was advised to stop using alcohol.

Five months later the patient’s condition worsened, and he was treated daily with thiamine intramuscularly for several days as part of alcohol detoxification treatment. At the same time, he considerably reduced his alcohol use. During the next months, the lower limb symptoms partially resolved, and the patient was able to walk independently. However, one year after thiamine treatment, there was still muscle weakness. The patient was able to walk a maximum of four kilometers and could not bicycle uphill or run. He complained of severe memory problems. He had difficulties remembering past events, to which photographs helped. He could not resume watching a recorded TV-program after a 30-min break. There was a mild confabulation tendency presenting as wrong recollection of conversations with friends and relatives. The patient was living alone in his own home and was independent in basic everyday activities. The patient was evaluated for possible KS.

**Neuropsychological assessment**

The Vocabulary, Digit Symbol, Information, and Letter-Number Sequencing subtests of the Wechsler Adult Intelligence Scale-III (WAIS-III) were used to assess general cognitive level, psychomotor speed, general knowledge, and verbal working memory, respectively. General attention and divided attention were tested with the Trail Making A and B tests, respectively. We used the Wechsler Memory Scale-III (WMS-III) to test memory. In testing auditory/verbal memory, we used the Logical Memory subtest and both word list subtests: the Word List test that contains 12 unrelated words and the Verbal Pair Associates (VPA) test that contains 8 word pairs. The words of each pair are semantically unrelated, but it is possible for the test subject to form associations that
facilitate remembering. Testing of executive functions comprised set shifting (Wisconsin Card Scoring Test/WCST: number of categories achieved and number of perseverative errors) and task switching (the score difference between Trail Making Tests B and A). Results at least 1.5 standard deviations (SDs) below the mean of normative values were considered abnormal for all tests except the WCST, for which we used the −1 SD threshold suggested in the test manual. Intrusions were defined as false words and statements in the WMS-III.

**DTI and conventional brain MRI**

MRI and DTI were performed as previously described. Volumetric analysis was performed with the FDA-approved cNeuro cMRI software (Combinostics, Finland). Brain area volumes were measured in milliliters, and age- and sex-adjusted volume z-scores were calculated based on the comparison with controls. Z-scores ≤ 1 were considered indicative of atrophy. The neocortical areas (frontal, temporal, parietal, and occipital lobes separately) were evaluated with the Pasquier’s global cortical atrophy (GCA) scale. In addition, the MBs were manually outlined in 1.0 mm thick coronal reconstructions of 3D T1 images, and their volume in milliliters was compared to 20 age- and sex-matched controls.

<table>
<thead>
<tr>
<th>Cognitive domain</th>
<th>Test</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Working memory</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Logical Memory I</td>
<td>−1.67</td>
</tr>
<tr>
<td>Verbal</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Verbal Paired Associates</td>
<td>−0.67</td>
</tr>
<tr>
<td>Visual</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Word Lists</td>
<td>−2.67</td>
</tr>
<tr>
<td>Episodic memory</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Visual Reproduction I</td>
<td>−0.67</td>
</tr>
<tr>
<td>Immediate recall</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (logical)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (word list)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (word list)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visual</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Delayed recall</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (logical)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (word list)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal (word list)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visual</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Recognition</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verbal</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Logical Memory II + Verbal</td>
<td>−3</td>
</tr>
<tr>
<td>Visual</td>
<td>WMS-III&lt;sup&gt;b&lt;/sup&gt;, Visual Recognition</td>
<td>−2</td>
</tr>
</tbody>
</table>

Table 1 Results of memory tests for a patient with Korsakoff syndrome

Notes: Values of −1.5 or less (Z-score) are in bold. The patient’s results were compared to age-adjusted normative values. <sup>a</sup>Wechsler Adult Intelligence Scale–III. <sup>b</sup>Wechsler Memory Scale–III.

Consent to participate

The study complied with the Declaration of Helsinki. The patient gave written informed consent for the use of his medical information in this publication. The study was approved by the Satakunta Central Hospital. The Finnish legislation does not require any further permissions.

Results

Abnormal memory results were found in tests of verbal working memory, immediate and delayed verbal recall, and in verbal and visual recognition (Table 1). The WMS-III Word List subtest was abnormal, but the VPA test was normal. There were two non-semantic intrusions in the
VPA test, and in the Logical Memory test, there were five semantic and one non-semantic intrusion. In the other neuropsychological tests (Table 2), abnormal results were found in the Information subset of the WAIS-III and in the set-shifting test of executive functions (WCST).

In brain volumetric analysis, atrophy changes were found in the hypothalamus, the MBs, the left hippocampus, the nucleus accumbens, the caudate, the pallidum, the thalamus, and the vermis (Table 3). In addition, there was general reduction of white-matter volume. CGA values were well within normal limits. Abnormal FA-values were found in several white matter tracts, including frontotemporal tracts (uncinate fasciculi, cingulum) (Table 4). Figure 1 shows atrophy changes. Figure 2 shows a DTI reconstruction of damaged white-matter tracts.

**Discussion**

We describe a patient with WKS caused by depression-related malnourishment and alcohol. At the time of WE, the patient had lost weight, and he was ataxic. Thus, the WE diagnostic threshold of two out of four diagnostic criteria (nutritional deficiency, ataxia, oculomotor abnormalities, altered mental state or mild memory impairment) proposed by Caine et al\textsuperscript{52} was reached.

**Neuropsychological findings**

The neuropsychological examination showed cognitive impairment, memory being preferentially affected. Logical memory and learning of unrelated words were most severely affected, whereas learning of words that can be semantically interconnected (VPA test) was normal. Patients with KS may have partially preserved anterograde semantic memory and be able to recruit associative mechanisms.\textsuperscript{53} Preferential impairment in the VPA test has been demonstrated in individuals with agenesis of the corpus callosum with normal interhemispheric connectivity being a prerequisite for a strong self-generated semantic network that supports encoding.\textsuperscript{47} Interestingly, the corpus callosum FA-values of our patient were normal.

We have recently shown that in patients with non-alcoholic KS the logical memory test may be abnormal, and the word list test may be normal and vice versa.\textsuperscript{6} Taken together with findings from the present patient, these results corroborate our previous suggestion on the need for a comprehensive neuropsychological evaluation of patients with suspected KS.\textsuperscript{6} Among word list tests, those containing unrelated words might have a better sensitivity. Had this patient been tested only with the VPA test or another word test that uses semantically related words, it could have been erroneously concluded that there is no memory impairment.

Similarly to previous studies, the patient made intrusions both in the logical memory test\textsuperscript{4,6,54} and the word list test.\textsuperscript{4,6} This further supports the usefulness of intrusions in examining patients with suspected KS.

In the WCST, both orbital prefrontal and dorsolateral prefrontal functions were affected. This is in accordance with previous findings in KS.\textsuperscript{27,32}

In alcoholic KS, executive dysfunction is common,\textsuperscript{26–32} but in non-alcoholic KS, executive functions appear to be largely preserved.\textsuperscript{4,6,33} There are several possible explanations for this discrepancy. Executive dysfunction in alcoholic KS could be due to a toxic effect of alcohol to the frontal lobes,\textsuperscript{26} and coexistence with thiamine-deficiency

**Table 2 Results of neuropsychological tests other than memory for a patient with Korsakoff syndrome**

<table>
<thead>
<tr>
<th>Cognitive domain</th>
<th>Test</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>General cognitive level</td>
<td>Vocabulary subset (WAIS-III\textsuperscript{a})</td>
<td>–0.67</td>
</tr>
<tr>
<td>General knowledge</td>
<td>Information subset (WAIS-III\textsuperscript{a})</td>
<td>–2.33</td>
</tr>
<tr>
<td>General attention</td>
<td>Trail Making Test A</td>
<td>0.82</td>
</tr>
<tr>
<td>Divided attention</td>
<td>Trail Making Test B</td>
<td>0</td>
</tr>
<tr>
<td>Psychomotor speed</td>
<td>Digit Symbol subset (WAIS-III\textsuperscript{a})</td>
<td>–1</td>
</tr>
<tr>
<td>Executive functions I (set shifting)</td>
<td>WCST\textsuperscript{b}, number of perseverative errors</td>
<td>–3</td>
</tr>
<tr>
<td>Orbital prefrontal functions</td>
<td>WCST\textsuperscript{b}, number of categories completed</td>
<td>–1.41</td>
</tr>
<tr>
<td>Dorsolateral prefrontal functions</td>
<td>Trail B – Trail A score</td>
<td>–0.82</td>
</tr>
</tbody>
</table>

Notes: Z-score values ≤ −1.5 (≤ −1 for the WCST test) are in bold. The patient’s results were compared to age-adjusted normative values. \textsuperscript{a} Wechsler Adult Intelligence Scale–III. \textsuperscript{b} Wisconsin Card Scoring Test.
induced memory impairment may be incidental. Another possible explanation is a selection bias, due to recruitment of KS study subjects from nursing facilities. Residents of nursing facilities are very probable to have executive dysfunction (since they cannot live independent lives). Van Oort and Kessels have shown that KS patients residing in nursing facilities have executive dysfunction,\(^4\) and Bowden has noted that studies of alcoholic KS are biased toward non-alcoholic KS without executive dysfunction.\(^4\) In contrast, our previously reported patients with non-alcoholic KS and normal or minimally impaired executive functions were identified by screening psychiatric outpatients for previously undiagnosed WE.\(^4\) Alternatively, repeated episodes of WE may cause executive dysfunction, and patients with long-term alcoholism are more probable to have repeated WE episodes compared to non-alcoholic WE patients. However, we have recently reported on a patient with two separate WE episodes caused by hyperemesis gravidarum who developed non-alcoholic KS without executive dysfunction.\(^4\)

One of the tests we used to assess executive functions was the WCST. Similarly to previous studies,\(^25,27,56,57\) both the number of categories completed and the number of perseveration errors were abnormal. This dual abnormality may differentiate alcoholic KS patients from patients with other causes of organic amnesia.\(^56,57\) Although frequently associated with frontal lobe atrophy, abnormal

### Table 3

<table>
<thead>
<tr>
<th>Brain area</th>
<th>Volume (ml)</th>
<th>% value</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain stem</td>
<td>18.2</td>
<td>29</td>
<td>-0.55</td>
</tr>
<tr>
<td>Hemispheric gray matter total</td>
<td>598</td>
<td>56</td>
<td>0.15</td>
</tr>
<tr>
<td>Hemispheric gray matter right</td>
<td>300</td>
<td>57</td>
<td>0.18</td>
</tr>
<tr>
<td>Hemispheric gray matter left</td>
<td>298</td>
<td>54</td>
<td>0.10</td>
</tr>
<tr>
<td>Hemispheric white matter</td>
<td>404</td>
<td>7</td>
<td>-1.48</td>
</tr>
<tr>
<td>Cerebrosinal fluid</td>
<td>49.2</td>
<td>94</td>
<td>1.55</td>
</tr>
<tr>
<td>Entorhinal area, right</td>
<td>2.71</td>
<td>85</td>
<td>1.04</td>
</tr>
<tr>
<td>Entorhinal area, left</td>
<td>2.25</td>
<td>51</td>
<td>0.03</td>
</tr>
<tr>
<td>Hippocampus, right</td>
<td>3.67</td>
<td>27</td>
<td>-0.61</td>
</tr>
<tr>
<td>Hippocampus, left</td>
<td>3.22</td>
<td>10</td>
<td>-1.28</td>
</tr>
<tr>
<td>Amygdala, right</td>
<td>1.17</td>
<td>47</td>
<td>-0.08</td>
</tr>
<tr>
<td>Amygdala, left</td>
<td>1.13</td>
<td>32</td>
<td>-0.47</td>
</tr>
<tr>
<td>Nucleus accumbens, right</td>
<td>0.34</td>
<td>4</td>
<td>-1.75</td>
</tr>
<tr>
<td>Nucleus accumbens, left</td>
<td>0.33</td>
<td>1</td>
<td>-2.33</td>
</tr>
<tr>
<td>Caudate nucleus, right</td>
<td>2.79</td>
<td>7</td>
<td>-1.48</td>
</tr>
<tr>
<td>Caudate nucleus, left</td>
<td>2.65</td>
<td>3</td>
<td>-1.88</td>
</tr>
<tr>
<td>Pallidum, right</td>
<td>1.29</td>
<td>9</td>
<td>-1.34</td>
</tr>
<tr>
<td>Pallidum, left</td>
<td>1.29</td>
<td>12</td>
<td>-1.17</td>
</tr>
<tr>
<td>Putamen, right</td>
<td>4.74</td>
<td>66</td>
<td>0.41</td>
</tr>
<tr>
<td>Putamen, left</td>
<td>4.63</td>
<td>48</td>
<td>-0.05</td>
</tr>
<tr>
<td>Thalamus, right</td>
<td>7.13</td>
<td>7</td>
<td>-1.48</td>
</tr>
<tr>
<td>Thalamus, left</td>
<td>7.32</td>
<td>8</td>
<td>-1.41</td>
</tr>
<tr>
<td>Hypothalamus, right</td>
<td>4.17</td>
<td>7</td>
<td>-1.48</td>
</tr>
<tr>
<td>Hypothalamus, left</td>
<td>4.21</td>
<td>4</td>
<td>-1.75</td>
</tr>
<tr>
<td>Cerebellar vermal lobules I-V</td>
<td>3.41</td>
<td>10</td>
<td>-1.28</td>
</tr>
<tr>
<td>Cerebellar vermal lobules VI-VII</td>
<td>1.61</td>
<td>13</td>
<td>-1.13</td>
</tr>
<tr>
<td>Cerebellar vermal lobules VIII-X</td>
<td>2.19</td>
<td>9</td>
<td>-1.34</td>
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<tr>
<td>Cerebellum exterior, right</td>
<td>49</td>
<td>23</td>
<td>-0.74</td>
</tr>
<tr>
<td>Cerebellum exterior, left</td>
<td>49.8</td>
<td>36</td>
<td>-0.36</td>
</tr>
<tr>
<td>Cerebellum white matter, right</td>
<td>11.7</td>
<td>15</td>
<td>-1.04</td>
</tr>
<tr>
<td>Cerebellum white matter, left</td>
<td>11.9</td>
<td>23</td>
<td>-0.74</td>
</tr>
<tr>
<td>Mammillary bodies (right + left)</td>
<td>112</td>
<td>9</td>
<td>-1.32</td>
</tr>
</tbody>
</table>

### Table 4

<table>
<thead>
<tr>
<th>White matter tract</th>
<th>Tractography</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corpus callosum genu</td>
<td>0.50/27</td>
<td>-1.2</td>
</tr>
<tr>
<td>Corpus callosum body</td>
<td>0.50/27</td>
<td>-1.1</td>
</tr>
<tr>
<td>Corpus callosum splenium</td>
<td>0.50/27</td>
<td>-0.8</td>
</tr>
<tr>
<td>Superior cingulum, right</td>
<td>0.15/27</td>
<td>-2.5</td>
</tr>
<tr>
<td>Superior cingulum, left</td>
<td>0.15/27</td>
<td>-2.0</td>
</tr>
<tr>
<td>Inferior cingulum, right</td>
<td>0.15/27</td>
<td>-2.2</td>
</tr>
<tr>
<td>Inferior cingulum, left</td>
<td>0.15/27</td>
<td>-2.3</td>
</tr>
<tr>
<td>Subgenual cingulum, right</td>
<td>0.15/27</td>
<td>-2.8</td>
</tr>
<tr>
<td>Subgenual cingulum, left</td>
<td>0.15/27</td>
<td>-2.5</td>
</tr>
<tr>
<td>Uncinate fasciculus, right</td>
<td>0.15/27</td>
<td>-2.2</td>
</tr>
<tr>
<td>Uncinate fasciculus, left</td>
<td>0.15/27</td>
<td>-2.1</td>
</tr>
<tr>
<td>Superior longitudinal fasciculus, right</td>
<td>0.15/27</td>
<td>-3.0</td>
</tr>
<tr>
<td>Superior longitudinal fasciculus, left</td>
<td>0.15/27</td>
<td>-2.1</td>
</tr>
<tr>
<td>Inferior longitudinal fasciculus, right</td>
<td>0.15/27</td>
<td>-1.3</td>
</tr>
<tr>
<td>Inferior longitudinal fasciculus, left</td>
<td>0.15/27</td>
<td>-1.4</td>
</tr>
<tr>
<td>Arcuate fasciculus, left</td>
<td>0.15/27</td>
<td>-1.9</td>
</tr>
<tr>
<td>Inferior fronto–occipital fasciculus, right</td>
<td>0.15/27</td>
<td>-1.3</td>
</tr>
<tr>
<td>Inferior fronto–occipital fasciculus, left</td>
<td>0.15/27</td>
<td>-1.6</td>
</tr>
<tr>
<td>Anterior corona radiata, right</td>
<td>0.15/27</td>
<td>-2.0</td>
</tr>
<tr>
<td>Anterior corona radiata, left</td>
<td>0.15/27</td>
<td>-1.5</td>
</tr>
<tr>
<td>Frontostriatal projection fibers, right</td>
<td>0.15/27</td>
<td>-1.6</td>
</tr>
<tr>
<td>Frontostriatal projection fibers, left</td>
<td>0.15/27</td>
<td>-1.9</td>
</tr>
<tr>
<td>Mesencephalic projections, right</td>
<td>0.30/27</td>
<td>-1.5</td>
</tr>
<tr>
<td>Mesencephalic projections, left</td>
<td>0.30/27</td>
<td>-0.3</td>
</tr>
<tr>
<td>Fornix, right</td>
<td>0.15/40</td>
<td>-1.8</td>
</tr>
<tr>
<td>Fornix, left</td>
<td>0.15/40</td>
<td>-2.0</td>
</tr>
<tr>
<td>Descending fornix, right</td>
<td>0.15/40</td>
<td>-2.2</td>
</tr>
<tr>
<td>Descending fornix, left</td>
<td>0.15/40</td>
<td>-1.7</td>
</tr>
</tbody>
</table>

Notes: The volume of the mammillary bodies was compared to 20 healthy controls. For the rest of the brain areas, the cNeuro cmri software was used. Values <=1 (for gray or white matter brain areas) and >1 (for CSF areas) are in bold.

Notes: Z-scores represent the difference from the mean of controls with one standard deviation as the unit. Values >=2 are in bold.
results in the WCST may also be due to lesions in the thalamus, the basal ganglia, the cerebellum, and frontostriatal white matter tracts. The thalamus, the basal ganglia, and the cerebellum, but not the frontal lobes, showed signs of atrophy in our patient. The frontostriatal tracts had reduced FA-values but below the –2 SD threshold we used. Similarly to previous studies, the task switching test (Trail B – Trail A score) was normal.

Neuroimaging findings
MRI showed more than 1 SD volume reduction in the thalamus, the MBs, and the vermis, as severe as in typical alcoholic KS. Atrophy was also found in the basal ganglia, a brain area that can be affected in WKS. However, contrary to findings in patients with alcohol use disorder, there was no cortical atrophy. This underscores the importance of thiamine deficiency in the development of KS in the present patient as opposed to a direct effect of alcohol.

DTI showed extensive damage in white matter tracts. Similarly to previous KS studies, damage was shown in the Papez circuit (fornix), frontotemporal tracts (uncinate fasciculus, cingulum), the inferior longitudinal fasciculus, and the anterior corona radiata. There was also damage in the superior longitudinal fasciculus, which has been recently shown to be affected in alcoholic men. Together with results from previous studies, our results support the importance of DTI in exploring brain damage in WKS. Damage in the uncinate fasciculus and the temporal cingulum supports our previous suggestion on the importance of frontotemporal tract involvement in WKS.

KS diagnosis
The clinical history and the results of the neuropsychological and radiological examinations support the diagnosis of KS. However, when considering functional impairment, the patient can be diagnosed with KS according to the DSM-IV but not according to the DSM-5.

The DSM-IV mentions KS as alcohol-induced persisting amnestic disorder due to thiamine deficiency. The DSM-5 changed the classification of neurocognitive disorders (NCDs) by introducing a baseline distinction between major and mild NCDs. A defining characteristic of major NCD is inability to function independently in everyday life. The DSM-5 mentions thiamine deficiency and alcohol as independent causes of major and mild...
NCD. KS is mentioned as alcohol-induced amnestic confabulatory major NCD with prominent amnesia and a tendency to confabulate.

Therefore, the DSM-5 has raised the threshold to diagnose KS. A person with memory impairment that cannot work but is independent in everyday life could have been diagnosed with KS in DSM-IV but cannot be diagnosed with KS with DSM-5. Interestingly, the other NCDs may manifest both as major and mild, and KS appears to be the only exception.

The DSM-5 inappropriately restricts the use of the term KS to cases were the memory impairment was caused by alcohol and the patient confabulates (“alcohol-induced amnestic confabulatory NCD”). However, it has been shown that KS may occur as a result of non-alcoholic WE. Spontaneous confabulations are uncommon in chronic KS.

As illustrated by the present patient, patients with partially preserved functioning after WE may fit diagnostically within the current clinical and research framework of KS. It is possible to use the DSM-5 terms mild NCD due to a medical disorder (thiamine deficiency), mild NCD due to alcohol, or (as in the current patient) use both diagnoses.

According to the DSM-5, neuropsychological test results between –1 and –2 SDs below the normative mean are associated with mild NCD, whereas more deviant test results (at least – 2 SDs below the normative mean) suggest major NCD. Our patient had test results compatible with major NCD on most abnormal memory test results.

Three considerations further support the use of the KS diagnosis for patients with DSM-5 mild NCD. First, the DSM-5 encourages a disorder conceptualization based on empirical validation of biological determinants. Researchers and clinicians are encouraged to use the Alzheimer’s disease diagnosis instead of Mild Cognitive Impairment for patients with cerebrospinal fluid, positron-emission tomography, and neuropsychological findings compatible with Alzheimer’s disease, even with not severe functional impairment. Second, thiamine deficiency can vary in severity and duration, and so can vary the adequacy of thiamine treatment. Presumably, the ensuing brain damage and functional impairment will vary in severity. Mild memory impairment after WE has been mentioned in many previous reports. We have previously reported on a patient with non-alcoholic KS and atypical neuropsychological presentation that also presented with mild NCD. Third, WKS is greatly underdiagnosed, particularly in non-alcoholic patients. Given the devastating effects of undiagnosed and untreated WKS, it is not wise to raise the diagnostic threshold in the absence of supporting empirical evidence.

Major depression may cause cognitive impairment, but comorbid affective disorder does not exacerbate the effects of alcohol use on cognition. Chronic alcoholism may cause cognitive problems, but abstinence is associated with substantial recovery. The present patient presented memory complaints only after the WE. The presence of intrusions in memory testing is suggestive of organic memory impairment. Deficiency of other micronutrients may have contributed to the cognitive problems.

Other clinical aspects

The patient presented with severe muscle weakness that partially improved after thiamine treatment. Partially improved muscle weakness has been previously described in WKS.

The underdiagnosis of KS has been attributed to patients being diagnosed with alcoholic dementia when the cognitive impairment has been in fact caused by thiamine deficiency. Psychiatric patients with non-alcoholic KS are another underdiagnosed group. Our results, together with results from our previous reports, may help identify KS patient who despite clear memory impairment live relatively independent lives.

The patient in the present report developed thiamine deficiency as a result of psychiatric disease (major depression) and alcohol use. Major depression is associated with weight loss, and comorbid alcohol use disorder is common. Thus, depressed patients may be at high risk for WKS. To our knowledge, there are 12 previous reports of major depression causing malnutrition and WE. Two of these patients had concomitant alcohol use disorder. In our patient, treatment with parenteral thiamine was delayed, which possibly resulted in long-term memory impairment and muscle weakness. This underlines the importance of early recognition and treatment of WE but also the usefulness of routine administration of parenteral thiamine in alcohol detoxification.

The continuity theory claims that alcohol affects memory along a continuum with KS patients being gravely impaired. However, as the present report illustrates, it is not possible to know whether a patient with clear but not debilitating memory impairment has mild KS or is an “uncomplicated alcoholic.” This supports the notion
that previous results in support of the continuity theory may instead reflect a continuum of memory impairment caused by thiamine deficiency.\(^5\)

We have previously reported on six psychiatric patients with KS and previously undiagnosed non-alcoholic WE, which were identified through systematic consideration of WKS in psychiatric patients with prominent memory complaints.\(^4,6\) The present patient extends the usefulness of this clinical approach to patients with previously undiagnosed WE of mixed etiology (alcoholic and psychiatric).

**Limitations**

Our study has several limitations. Data come from only one patient. Larger studies are needed to further explore imaging and neuropsychological features of relatively mild WKS. We assessed executive functions with only two tests. A more thorough examination could have provided a more complete picture of impairment and preserved skills. We did not examine the frontocerebellar tracts that may be affected in WKS.\(^45,82\)

**Conclusions**

In summary, we present a patient that developed WKS as a result of depression and alcohol use. The patient’s chronic memory impairment has compromised his working ability, but he does not need assistance in everyday life. The patient can be diagnosed with KS according to the DSM-IV but not according to the DSM-5. However, all the core neuropsychological and radiological findings of KS were present. Strict use of the DSM-5 in evaluating patients suspected of having KS may lead to underdiagnosis. Finally, the memory testing of patients with suspected KS should be comprehensive and include word list tests containing semantically unrelated items.

**Abbreviation list**

DTI, diffusion tensor imaging; FA, fractional anisotropy; KS, Korsakoff syndrome; MB, mammillary body; NCD, neurocognitive disorder; SD, standard deviation; VPA, verbal pair associates; WCST, Wisconsin Card Scoring Test; WE, Wernicke’s encephalopathy; WKS, Wernicke-Korsakoff syndrome; WMS-III, Wechsler memory scale-III.

**Disclosure**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

**References**
