
Organic psychosis due to hypoparathyroidism in an older adult: a case report

We present a case of psychosis secondary to iatrogenic hypoparathyroidism and hypocalcemia in a 73-year-old woman. At age 18, she had undergone a total thyroidectomy and developed severe psychotic symptoms due to unintended removal of the parathyroid glands, but had recovered fully after a hospital stay. For the last 15 years, she had been receiving periodic psychiatry follow-up for mild depressive and psychotic symptoms, consisting of auditory hallucinations (children’s voices calling her “Mommy”). For years, she had been stable on a combination of fluoxetine 20 mg/day and olanzapine 10 mg/day. She had worked as a nurse with no functional impairment until retirement at age 65.

The patient presented to the emergency department reporting anguish, auditory hallucinations of male voices that insulted her (which, according to the patient, came from electronic devices), and auditory misperceptions, with intolerance to electronic noises. The situation had worsened in the previous 6 months, and the patient now exhibited marked social withdrawal. She also presented a secondary depressed mood.

She was admitted to the psychiatric department. Initial treatment was olanzapine 10 mg/day, with no therapeutic response. The patient crawled into the observation bay, stating the noise of the air conditioner was unbearable, with no interference with function or behavior. The patient was found to have a mild ataxic gait worsened by extrapyramidal symptoms, with no clear improvement despite anticholinergic therapy. No other neurological signs of hypocalcemia were found. Blood tests showed hypocalcemia, hyperphosphatemia, and mild hypoparathyroidism (Table 1). A severe vitamin D deficiency was also noticed (Table 1). We noticed that the patient had been refusing to take the calcium and calcitriol supplementation prescribed by her endocrinologist, leading to chronic hypocalcemia. Computed tomography scans showed bilateral calcifications in the globi pallidi and cerebellar dentate nuclei (Figure 1). Neuropsychological assessment showed mild cognitive impairment in some areas (attention/concentration, verbal initiative, and motor function).

Additional calcium supplementation was prescribed (calcium carbonate 4,000 mg daily) and calcitriol was reintroduced (0.25 μg twice daily), with subsequent improvement in laboratory values (Table 1). The patient’s atactic gait became barely perceptible and her auditory hallucinations, although persistent, became quieter and more bearable, with no interference with function or behavior. Accordingly, her secondary depressed mood improved to euthymia. Interestingly, motor function and attention, which were impaired at the first neuropsychological assessment, also improved after calcium and calcitriol supplementation. Symptomatic treatment of psychosis with risperidone 2 mg/day was reduced to 1 mg/day and later switched to paliperidone 3 mg/day due to intolerable extrapyramidal symptoms. No worsening was noticed after the dose reduction and switch in antipsychotic agent.

Few cases of hypoparathyroidism and hypocalcemia with neuropsychiatric manifestations have been reported.1,2 Basal ganglia calcification involving the globus pallidus and cerebellum is typical, perhaps due to hyper-

Table 1 Blood test results at admission and after 6 months of calcium and vitamin D supplementation

<table>
<thead>
<tr>
<th>Value</th>
<th>Admission</th>
<th>Follow-up</th>
<th>Normal range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ionized calcium (mmol/L)</td>
<td>0.84</td>
<td>1.24</td>
<td>1.13-1.32</td>
</tr>
<tr>
<td>Phosphorus (mmol/L)</td>
<td>5.40</td>
<td>4.3</td>
<td>2.70-4.5</td>
</tr>
<tr>
<td>Magnesium (mmol/L)</td>
<td>0.68</td>
<td>0.80</td>
<td>0.78-1.02</td>
</tr>
<tr>
<td>Parathyroid hormone (pg/mL)</td>
<td>8.40</td>
<td>9.40</td>
<td>10-55</td>
</tr>
<tr>
<td>Vitamin D (ng/mL)</td>
<td>13</td>
<td>27</td>
<td>&gt; 30</td>
</tr>
</tbody>
</table>

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phosphatemia, resulting in psychosis, mood disorders, or both, as well as neurological manifestations. The basal ganglia play important roles in mood, cognition, motivation, and motor control; their function includes motor learning, attention allocation and filtering, and working memory. Psychotic symptoms in such cases often include auditory hallucinations, delusions of influence, paranoid states, and complex perceptual distortions. Hypocalcemia also causes cognitive impairment. Neurological manifestations tend to improve with calcium correction, but psychiatric symptoms do not improve substantially.

We believe this case report illustrates the need for an exhaustive workup of possible organic etiologies when psychiatric patients present with atypical symptoms. All symptoms presented by the patient could be explained by her hypoparathyroidism, hypocalcemia, and consequent typical brain calcifications, justifying the diagnosis of organic psychosis. In addition, vitamin D deficiency may worsen psychotic symptoms.

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Disclosure

The authors report no conflicts of interest.

References