Case Report

Catatonia and Mutism: Neurotic, Psychotic, or Organic Disorder?

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Background. Catatonia is caused by a variety of psychiatric and organic conditions. The onset, clinical profile, and response to treatment may vary depending on the underlying cause. Catatonia is more likely to be associated with neurotic and psychotic disorders, but some psychiatric symptoms are key components in the clinical presentation of other medical conditions. Case Report. We report the case of a woman who started showing paroxysmal recurrent episodes since the age of 57 years, characterized by surrounding disconnection, disorientation, and muscle spasm (myoclonus), followed by a postictal state. In the following months, the symptoms evolved to akinetic mutism, catatonia, and rapidly progressive vision and audition loss. She underwent a battery of tests, most of them inconclusive, until a neoplastic meningoencephalitis was diagnosed after more than two years of symptoms. Numerous medical conditions can mimic psychiatric disorders. This uncommon presentation may lead to a late diagnosis and treatment initiation, increasing significantly morbidity and mortality. A differential diagnosis with infectious, autoimmune, and neoplastic etiologies should always be carried out.

1. Introduction

Even though catatonia can be caused by a variety of psychiatric and organic conditions, ranging from neurologic to systemic diseases, it is more likely to be associated with neurotic and psychotic disorder [1]. Catatonia is found in 10% of psychiatric inpatients and is more common in patients with mood disorders, especially mania, than in patients with schizophrenia [2]. The diversity of symptoms often leads to a delay in diagnosis and treatment initiation, increasing morbidity and mortality significantly. Albeit catatonia and akinetic mutism are commonly related to psychiatric diseases, numerous medical conditions can mimic psychiatric disorders [3]. A differential diagnosis with infectious, autoimmune, and paraneoplastic encephalitis should always be carried out [4].

In this report, we describe the case of a patient who consulted due to a two-year history of paroxysmal recurrent episodes, with surrounding disconnection, disorientation, and muscle spasm (myoclonus), followed by a postictal state. Eventually, she ended up in a catatonic state, with rapidly progressive vision and audition loss and almost surrounding disconnection. The patient died seven months later.

The purpose of this report is to describe a case of psychomotor impairment and catatonia due to a neoplastic meningoencephalitis associated with an unknown primary tumor. We also aim to create a differential diagnosis, which should lead us to investigate other causes apart from mental illnesses, and to enumerate the general principles of a comprehensive assessment.

2. Case

The patient, Ms. A., was a 57-year-old married woman and mother to a son. She had no relevant medical history. As surgical background, she had undergone surgery of groin hernia, varicosities in lower limbs, appendectomy, and tonsillectomy. Ms. A. was an ex-smoker (who quit at the age of 39 years). She had a family history of breast cancer in her mother and her sister.

As for her psychiatric history, she suffered from depressive mood after her father’s death, when the patient was 49 years old. She was treated with individual support psychotherapy and venlafaxine 150 mg/day for one year, showing good response and clinical remission.
2.1. First Episodes. In March, 2015, Ms. A. attended the emergency room presenting with a confused state, spatial disorientation, inability to recognize her relatives, and a slurred speech. Due to the patient’s mental condition, the information was provided by her family.

In July, 2015, Ms. A. had a new episode, this time characterized by hyperventilation and upper and lower intermittent shakes. The patient was conscious, and she linked the episode with some stressful familiar situations. In both these episodes, Ms. A. was assessed by a neurologist, who performed blood tests, a computed tomography (CT) scan, an electroencephalogram (EEG), a sleep-deprived EEG, and a brain magnetic resonance imaging (MRI) scan but no abnormalities were detected.

Both neurologist and psychiatrist assessed the patient. Since physical tests were unremarkable, Ms. A. was diagnosed initially with a conversion/dissociative disorder. We referred Ms. A. to neurology and psychiatry departments in outpatient visits. After performing neurological and psychiatric assessments with the consent and collaboration of the patient and her relatives, she was started on an anticonvulsant (levetiracetam 1000 mg/day), an antidepressant (desvenlafaxine 50 mg/day), and a benzodiazepine (alprazolam sustained release tablets 1 mg/day).

2.2. New Episodes Two Years Later. Between July, 2015, and December, 2016, episodes as described above continued, but they were less frequent and self-limiting. From January, 2017, the episodes started to be more severe and disabling. In March 2017, Ms. A. presented to the hospital’s emergency room by her relatives with a new episode of confusion, spatial disorientation, myoclonus seizures, and stereotypies (such as lip smacking), during which she remained unconscious for ten minutes. Afterwards, she recovered by herself, but drowsiness and spatial disorientation persisted. After recovering, when we tried to establish contact with Ms. A. and elucidate what had happened, she could not remember the episode. According to her relatives at that time, she was taking desvenlafaxine 50 mg/day, alprazolam sustained-release tablets 0.5 mg/day, and olanzapine 5 mg/day. In order to rule out any iatrogenic cause, we replaced alprazolam 0.5 mg/day with diazepam 5 mg/8 h (as a muscle relaxer) and quitted desvenlafaxine.

In April, 2017, Ms. A. started to display migraine-like symptoms, drowsiness, and being increasingly more excited, up to the point that she could barely breathe, suffering from trismus and emitting gibberish sounds. Furthermore, she also presented with muscle contractions and claw digits. In our physical examination, she was neither cooperative nor resistant, but she appeared to be indifferent. We requested additional tests in order to rule out other organic conditions. Brain CT scan, EEG, and pulmonary scan (in order to rule out a pulmonary thromboembolism), as well as laboratory test results, were unremarkable, again. A psychiatric orientation was then reassessed as a key component of this clinical profile. At that time, we added quetiapine 25 mg/8 h replacing olanzapine for the purpose of reducing a long-term anxious mental state, which was the patient’s most remarkable clinical manifestation during those episodes.

2.3. Clinical Worsening and Hospitalization. In May 2017, within a few weeks, the patient developed hyperventilation, muscle stiffness, lockjaw, walking difficulty, falls, and psychiatric symptoms consisting of visual hallucinations, bizarre behaviour, and some abnormal gestures such as trying to grab nonexistent objects in the air. This initial presentation was followed by an acute course towards a catatonic state, mutism, rapidly progressive vision and audition loss (initially, she could not recognize colours, and then, she lost visual and audition acuity), frequent seizures, muscle rigidity, and facial dyskinesia. The patient was eventually admitted to a psychiatric inpatient unit, as our initial presumptive diagnosis had been, until that point, a mental disorder. There, she was started on intravenous diazepam 5 mg/8 h and haloperidol 5 mg/8 h. We did not observe any response, so we went further with physical evaluation.

A new brain MRI scan was performed, showing linear hyperintensities in basal segments and the VIII, VII, V, and III cranial nerves in FLAIR sequence, in the walls of both lateral ventricles in T2 sequences, as well as subtle communicant hydrocephalus (see Figures 1 and 2). A lumbar puncture showed very high cerebrospinal fluid (CSF)
pressure (55 mmHg, normal value 10-15 mmHg), notable hyperproteinorrachy (470 mg/dL, normal value 15-15 mg/dL), with normal glucose levels and cell account. Gram stain cultures and C-reactive protein (CRP) test for bacteria and viruses were also negative.

Eventually, CSF cytology detected tumor cells. These cells were compatible with a metastatic breast adenocarcinoma, according to immunohistochemical stain. However, we never found any primary tumor after performing body CT scan, mammography, and positron emission tomography- (PET-) CT. The clinical course was very aggressive, with refractory catatonia and rigidity, and the patient died few weeks later.

2.4. Final Diagnosis. The final diagnosis was carcinomatous meningoencephalitis, probable breast cancer as primary tumor.

3. Discussion

Catatonia is a neuropsychiatric syndrome easily identifiable by the clinician. However, an etiological diagnosis is not always easy. The Diagnostic and Statistical Manual of Mental Disorders, 4th Edition, Text Revision (DSM-IV-TR) [5] recognizes that many cases are secondary to an organic disorder, under the heading of catatonic disorder due to a general medical condition (CD-GMC). The fifth DSM edition (DSM-5) recognizes that catatonia can occur in the context of another mental disorder or as a disorder due to another medical condition. Despite the amount of evidence, there are not many publications available regarding characteristics of organic catatonia.

Many acute cases are attended at emergency departments, where the catatonias are often attributed to psychiatric problems, instead of medical and neurological conditions [1, 6]. This means delays in diagnosis and initial management, which are associated with increased morbidity and complications; some of these are serious and life-threatening. Many of these complications are related to decreased dietary and liquid intake and others to hospitalization and immobilization, such as rhabdomyolysis, deep vein thrombosis, pulmonary thromboembolism, and aspiration pneumonia, among others [6].

3.1. Differential Diagnosis. Apart from psychiatric disorders, catatonia is secondary to numerous medical and neurological conditions. In a cohort of patients from a single
In this case, the patient underwent a remarkable great number of tests; unfortunately, most of them were inconclusive. In fact, a first MRI was completely normal, and no abnormalities were found until a second MRI was performed two years later. Although prognosis of metastatic infiltration of CNS is bleak, a more rapid diagnosis may have been made if CSF had been analysed in an earlier stage.

This highlights how psychiatric symptoms often lead to a psychiatric diagnosis straightaway, with other possibilities seldom being considered. Although catatonia and akinetic mutism are commonly related to psychiatric diseases, several medical conditions can mimic psychiatric disorders [1, 5, 6, 10]. Cancer is just one of them, which can induce neurological and behavioural disturbances. Encephalitis is also an important differential diagnosis [9].

The correct diagnostic subcategorization of catatonia is critical for proper prognostication, identification of disease-specific acute therapies, and long-term therapeutic management of the underlying disease state. Delays are linked to poorer outcomes [6]. We strongly suggest to always rule out an organic condition as cause. We also recommend the use of lumbar puncture and CSF analyses for patients with catatonic disorder of unclear etiology [7, 8]. Focus should be placed on patients with neurological symptoms and recurrent bizarre episodes of undetermined origin with partial response to the treatment, as these patients may be at higher risk of any organic condition, and a complete differential diagnosis should be carried out.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest.

**Authors’ Contributions**

The authors of this paper were responsible for conceiving and designing the evaluation, clinical data collection, clinical data interpretation, and drafting the manuscript.

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