Huntington’s Disease in a Patient Misdiagnosed as Conversion Disorder

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Huntington’s disease (HD) is an inherited, progressive, and neurodegenerative neuropsychiatric disorder caused by the expansion of cytosine-adenine-guanine (CAG) trinucleotide at the coding region of the IT15 gene on chromosome 4. This pathology typically presents in individuals aged between 30 and 50 years and the age of onset is inversely correlated with the length of the CAG repeat expansion. It is characterized by chorea, cognitive deficits, and psychiatric symptoms. Usually the psychiatric disorders precede motor and cognitive impairment, Major Depressive Disorder and anxiety disorders being the most common presentations. We present a clinical case of a 65-year-old woman admitted to our Psychiatric Acute Unit. During the 6 years preceding the admission, the patient had clinical assessments made several times by different specialties that focused only on isolated symptoms, disregarding the syndrome as a whole. In the course of her last admission, the patient was referred to our Neuropsychiatric Team, which made the provisional diagnosis of late-onset Huntington’s disease, later confirmed by genetic testing. This clinical vignette highlights the importance of a multidisciplinary approach to atypical clinical presentations and raises awareness for the relevance of investigating carefully motor symptoms in psychiatric patients.

1. Introduction

Huntington’s disease (HD) is an inherited, progressive, and neurodegenerative neuropsychiatric disorder characterized by abnormal movements, cognitive deficits, and neuropsychiatric impairment with a worldwide service-based prevalence of 2.71 per 100,000 (95% CI: 1.55–4.72), based on a meta-analysis [1].

HD is caused by the expansion of cytosine-adenine-guanine (CAG) trinucleotide at the coding region of the IT15 gene on chromosome 4. Patients with more than 36 repeats will develop the disease, and the age of onset is inversely correlated with the length of the CAG repeat expansion [2].

Individuals who are diagnosed with HD are typically aged between 30 and 50 years and have an average life expectancy of 15–30 years after diagnosis.

Regarding the clinical presentation, chorea is the major neurologic finding, consisting in brief, abrupt, irregular, and unpredictable involuntary movements, which potentially leads to disturbances in posture, gait, and balance, resulting in increased risk of falls. There is also cognitive decline, usually beginning as attentional deficits, followed by onset of dementia [3]. Psychiatric symptoms found in HD include isolated symptoms such as apathy, irritability, or incipient memory loss as well as full blown syndromes, such as Major Depressive Disorder or anxiety disorders in 33–76%
of patients. Psychotic disorders and mania-like symptoms are rarer, affecting 3–11% [4].

The onset of serious motor and cognitive symptoms is often late in HD and psychiatric symptomatology often precedes motor symptoms by many years. Longitudinal research in the PREDICT-HD [5] and other clinical programs [6] indicates that HD patients develop subtle changes decades before classical presentation. These include cognitive, functional, and psychiatric symptoms, and altered brain morphology and connectivity and even subtle motor deficits [7]. Cross-sectional studies [8, 9] reported that sadness and depressed mood are early symptoms of HD that peak during the early motor symptomatic phase whereas significantly lower rates of depression are present in advanced stages of the disease.

A large international study [10] confirmed that depression, irritability/aggression, obsessive compulsive behaviors (OCBs) and apathy are highly prevalent neuropsychiatric symptoms in HD population. By contrast, the prevalence of psychosis in the same sample was low.

Moreover, the presence of neuropsychiatric symptoms was associated with a positive psychiatric history; particularly a past episode of depression was associated with manifestation of neuropsychiatric symptoms in HD.

However, there is still a dearth of data regarding psychiatric manifestations and care in HD.

2. Material and Methods

Literature revision was made using PubMed database under the following specific terms: “Huntington, Depressive symptoms, Prodromal symptoms, Mood, Gait, and Treatment”, as well as studies of epidemiology, etiology, and clinical presentation.

3. Case Report

We present the case of a 65-year-old Caucasian woman with no previous history of psychiatric disease until 2012 (60 years of age). At this point, she complained to her family doctor about low mood, reduced energy, and anhedonia. She was prescribed Sertraline 50 mg once a day and maintained the treatment for about 3 years with partial improvement, mainly in the first year.

At 63 years of age, she started complaining to her family doctor about nonspecific limb weakness without movement difficulties. A few months later (April 2015), she was admitted to hospital with a myocardial infarction and was discharged 5 days later with full recovery of coronary perfusion. She was followed in cardiology outpatient clinic for about one year and was discharged because of progressive noncompliance with medical treatment and prescribed medical exams.

During the months that followed the myocardial infarction, her depressive and motor symptoms gradually worsened, leading to eleven visits to the Psychiatric Emergency Room Service (ERS), where she was assessed by Psychiatry and Neurology with complaints of anxiety, low mood, social isolation, deficits in memory retrieval, unspecific pain, and limb weakness with progressive development of abnormal gait. She left the ERS before full medical evaluation multiple times and never accurately followed medication changes that were suggested.

After several visits to the ERS, she was referred to Psychiatry and Neurology outpatient clinics. At Psychiatry consultation, she was, once again, medicated for anxiety and depressive mood, with Duloxetine 30 mg and Gabapentin 300 mg daily. One year later, her condition worsened, which motivated addition of Mirtazapine 30 mg and titration of Gabapentin up to 600 mg, again without proper response.

Simultaneously, she was evaluated in a Neurology appointment because of diminished force and limitation of limb movements. The neurological examination noted unspecific wide-based gait, enhanced bilateral reflexes, and abolished bilateral postural sensitivity. Blood samples showed low folic acid (3.9 ng/mL), which was considered as partial justification for neurological findings and reposition with folic acid was initiated. Additionally, more exams were requested.

One month before the admission at our Acute Inpatient Unit (AIU) that leads to the diagnosis of HD, the patient was, once again, evaluated in the Psychiatric ESR. On observation, the patient showed worsening of mood status, increased pressure of speech, pseudo hallucinations (patient described the following: “a TV host speaks with me, he guides me and tells me to do certain things”), and suicidal ideation. She was started on Valproic Acid 1000 mg and Quetiapine 400 mg and Mirtazapine was suspended. However, behavior changes were maintained and even aggravated (she took her clothes off at the window, broke some lamps at home and was aggressive to her sister), eventually leading to the admission to the psychiatric ward.

The mental state examination at our unit showed low mood and partial disorientation in time. She was unable to provide dates of important life events saying repeatedly “I don’t remember” and scored 14/30 at Mini-Mental State Examination (MMSE). Because of her depressive symptoms and cognitive changes, diagnosis of Major Depressive Disorder/Pseudodementia was assumed and she was medicated with Mirtazapine 30 mg and Quetiapine 200 mg. She was discharged 12 (twelve) days later with improved mood, but she still presented abnormal sustained gait, which was interpreted as a comorbid Conversion Disorder.

The Computed Tomography (CT) and Electromyography did not reveal any significant alterations. Therefore, at the subsequent Neurology appointment, patient was discharged with diagnosis of Conversion Disorder.

After hospitalization, she did not follow the prescribed medication and her abnormal behavior was maintained, such as not cleaning her house and throwing clothes away, so she was referred again to the Psychiatric ERS. At the mental state examination she endorsed soliloquies, depressive mood with catathymic delusions, and suicidal ideation and was disoriented to time and place. For that reason, she was readmitted to the Psychiatry ward on March 2017.

During hospitalization the abnormal gait and movements were still obvious; therefore the case was referred to the Neuropsychiatry Team (NT). The NT considered that the psychiatric symptoms of the patient, in addition to the
The patient presented manic symptoms, interpreted as drug during her second hospitalization, Venlafaxine was initiated with SSRIs and the SNRIs Venlafaxine [12]. For that reason, during her second hospitalization, Venlafaxine was initiated with reversion of depressed mood. However, later on, our patient presented manic symptoms, interpreted as drug induced mania, and the antidepressive was discontinued. However, despite the fact that depressive episodes are much more frequent than manic episodes, HD is also considered a neurologic cause of mania [13] which could enhance the probability of secondary mania induced with Venlafaxine.

Her progressive noncompliance with medical advice and exams and multiple episodes of leaving the ERS against medical advice were probably early signs of behavioral and cognitive impairment. Dementia in HD is subcortical including deficits in processing speed, attention, visuospatial impairment, and dysarthric speech [12]. During the first hospitalization, these changes were interpreted as Pseudode- mentia in a 65-year-old woman, who had persistent depressed mood that had not responded to multiple antidepressants (probably due to therapeutic noncompliance) and mild cognitive impairment.

Movement dysfunction was evaluated at Neurology Consultation and AIU with an inconclusive clinical evaluation, and it was labelled as Conversion Disorder. The wide-based gait observed is a classical presentation of chorea in HD, but it is also similar to the one caused by cerebellar ataxia [14]. CT and Electromyography did not reveal any hypothesized lesion location that could explain the symptomatology. The choreiform movements can be worsened by stress factors, such as anxiety [15] which is another frequent psychiatric symptom of HD which she had complained of. The late-onset presentation (over 60 years) was another confounding factor for the diagnosis, due to the perceived low likelihood of HD at this age. Moreover, 94.4% of reported cases of LoHD had CAG repeat lengths of ≤44 (as our case) and, despite motor manifestations being the commonest initial presentation, 29.2% of the patients presented with nonmotor manifestations as the first clinical feature [15].

Regarding treatment, in this patient haloperidol was used successfully to address chorea, but there are several other approaches [12]. There is no available pharmacological treatment to stop the progression of HD, so treatment is focused on improving daily functioning [16]. Although there was a reduction in the severity of mood and movement dysfunction, our patient still manifests residual symptoms such as apathy. The latter is more frequent in advanced stages of the disease and is, among psychiatric symptoms, the most robustly associated with disease progression and cognitive and motor dysfunction [17].

5. Conclusions
Neuropsychiatric disorders often have overlapping and ambiguous presentations, constituting a diagnostic challenge. These could be more noticeable in individuals with LoHD, in which cognitive impairment, rather than chorea, may be the major source of disability.

Thereby, this case enhances the importance of effective collaboration between medical specialties that are dedicated to the study of the brain. The diagnosis of HD was finally achieved, only after the involvement of NT, showing that a multidisciplinary approach is a key factor in this kind of cases.

This case report also alerts clinicians for the need of a careful investigation of motor abnormalities in psychiatric
patients, before attributing these symptoms to causes, such as iatrogenic effects or Conversion Disorder.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest regarding the publication of this article.

**Authors’ Contributions**

João Machado Nogueira and Ana Margarida Franco contributed equally to this work.

**References**


