



genetic testing for this person was classified as a variant of uncertain significance.

Uniquely, this case report describes an adult who has been tested, as opposed to paediatric cases which comprise much of the literature in this area. The notable case described by Verhoven of a patient who presented with a late onset of seizures, similar symptoms, progression and prior failed trials of medication as the case presented here, also benefited from genetic testing in later life. In addition, this patient responded well to sodium valproate which is known to be beneficial in cases with SLC6A1. This is most likely related to the GABA mediation. The findings of this case demonstrate the SLC6A1 gene-related disorder provides a unifying explanation for this diverse clinical phenotype, previously thought to be a constellation of syndromes and co-morbid symptoms.

The individual in this case study failed to respond to three adequate trials of medication for ADHD, all which have a robust evidence base. This suggests a possible alternate pathway for the development of ADHD features in cases such as this, as the majority of individuals with moderate-severe ADHD symptoms achieve symptomatic relief with pharmacological intervention.

Conclusion: This case highlights the relevance of genetic testing in adults and the reporting of variants of uncertain significance that could possibly lead to reclassification of a variant.

Abstracts were reviewed by the RCPsych Academic Faculty rather than by the standard *BJPsych Open* peer review process and should not be quoted as peer-reviewed by *BJPsych Open* in any subsequent publication.

Parkinson's Disease With Psychosis: A Case Report of Neuropsychiatric Manifestations

Dr Aung Sein, Dr Mobolape Olajide, Dr Rachel Rajaratnam,
Dr Lior Mevorach and Dr Sakshi Alewad

Essex Partnership University NHS Trust, Colchester, United Kingdom

doi: [10.1192/bjo.2025.10758](https://doi.org/10.1192/bjo.2025.10758)

Aims: Parkinson's disease (PD) is mainly a movement disorder, although 30% of PD patients may also suffer from psychosis, which may impair quality of life. PD psychosis (PDPsy) may result following approximately 10 years of treatment using dopaminergic agonists. PDPsy is characterized by recurrent and continuous hallucinations and delusions lasting for at least 1 month.

Methods: A 59-year-old female was admitted under Section 2 of the Mental Health Act (MHA) due to concerns raised by the police and her family regarding her mental health. The police were particularly alarmed after she repeatedly contacted them, expressing paranoid delusions. Her confusion and memory issues further contributed to the concerns leading to her admission. She was under the Early Intervention Psychosis team at the time of admission but her engagement with them was very erratic. She was started on quetiapine 50 mg ON but was non-compliant. She was diagnosis with PD 12 years prior to admission, and was still under the neurology team, and being regularly reviewed. The medication prescribed for PD were Sinemet 12.5/25 2 tabs QDS and ropinirole 8 mg BD.

Upon admission, the patient reported feeling monitored via 16 satellites connected to her television, broadcasting signals worldwide. She denied calling the police and instead suggested that her phone had been hacked. The police reported that she alleged that she had been sexually assaulted by her ex-partner or individuals organized by him while being drugged, though she had no memory of these events when questioned. She believed she was being followed, a notion she first experienced on a train to Cornwall a year prior. She also alleged that her ex-partner frequently entered her home at night to steal from

her. She, also, exhibited delusional beliefs regarding her YouTube presence, asserting that her ex-partner manipulated satellites to influence her views online. Risk factors identified included medication non-compliance, poor insight, risk of falls due to Parkinson's disease, potential financial exploitation due to engagement with strangers on social media, and vulnerability.

Results: The differential diagnosis considered for this patient:

Delusional Disorder.

Late-Onset Psychosis.

Parkinson's Disease Psychosis.

Alzheimer's Disease with Psychosis.

Brief Reactive Psychosis.

A literature search showed that management of PDPsy involves a balance between reducing PD medication and introducing an antipsychotic for symptom management.

Conclusion: This case highlights the diagnostic challenges of psychosis in Parkinson's disease and underscores the importance of a multidisciplinary approach in managing psychiatric symptoms in neurodegenerative conditions.

Abstracts were reviewed by the RCPsych Academic Faculty rather than by the standard *BJPsych Open* peer review process and should not be quoted as peer-reviewed by *BJPsych Open* in any subsequent publication.

Case Report: The Forgotten Functions of the Hindbrain

Dr Shalina Ramsewak and Dr Shruti Lodhi

Surrey and Borders Partnership NHS Trust, Camberley, United Kingdom

doi: [10.1192/bjo.2025.10759](https://doi.org/10.1192/bjo.2025.10759)

Aims: This case highlights the cognitive functions of the cerebellum which is often forgotten about.

Methods: An 80-year-old lady with a previous mental health diagnosis of Bipolar Affective disorder (BIPAD). She presented to the general hospital in July 2024 with a sudden onset of confusion and aggressive behaviour. She was referred to the Psychiatric Liaison Service (PLS). On assessments, she presented with no clear mood or psychotic disorder suggestive of relapse in her BIPAD. She scored 8 out of 30 on the Mini-Addenbrooke's Cognitive Examination (M-ACE) – losing marks on attention, memory – registration/recall, verbal fluency and the clock drawing test showed neglect. She lacked insight into her presentation and thus was in hospital under Deprivation of Liberty Safeguards (DoLS). Collateral history from her son corroborated that this was not a relapse in BIPAD. He enquired about an MRI head which revealed a small old infarct in the left cerebellar hemisphere.

Results: Cerebellar cognitive affective syndrome (CCAS; Schmahmann's syndrome) would explain the timeline of symptoms, assessment findings and collateral history. CCAS is characterized by deficits in executive function, linguistic processing, spatial cognition, and affect regulation. Neuropsychiatric features include impairments in attentional control, emotional control, psychosis spectrum disorders and social skill set. The deficits suggest a disruption of the cerebellar modulation of neural circuits that link frontal, parietal, temporal, and limbic cortices with the cerebellum which is known as Cerebellum-Cerebrum cortex loop. Movement, co-ordination, and cognition are intricately connected within the brain, however the role of the cerebellum in cognition has often been ignored.

Conclusion: Clinicians tend to focus on the motor-coordination functions of the hindbrain, and this case report highlights cognitive functions of the hindbrain that are often forgotten. CCAS may be underdiagnosed due to the lack of awareness of the intricate