Does recurrent catatonia manifest in a similar fashion in all the episodes of mood disorder? A case series with literature review

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ABSTRACT
Catatonia, originally conceptualised by Kahlbaum in 1868, is a neuropsychiatric condition that has been found to occur concomitantly with several organic and psychiatric conditions. Starting from the era of Kraepelin and Bleuler, this condition was faultily linked with schizophrenia alone; however, over time, greater associations have been found between catatonia and mood disorders. Despite the availability of several reports supporting this finding, there is a relative paucity of studies that specifically focus on catatonia to be the first symptom manifestation heralding a subsequent mood episode. In addition, there is scant literature to determine whether there are specific presentations of catatonia that show greater associations with mood disorders and whether these signs and symptoms recur in a stereotypical fashion in the subsequent mood episodes in the lifetime of an individual. We hereby report two cases with a diagnosis of mood disorders (bipolar disorder and recurrent depressive disorder) who had catatonia as the initial symptom not only at presentation but also at subsequent episodes. The report emphasises that recurrent catatonia can be the initial clinical manifestation of an underlying mood episode, which appears otherwise masked behind the catatonic presentation. These catatonic symptoms can be interestingly similar in all the subsequent episodes. A detailed clinical evaluation is thus warranted after catatonia has been duly treated to provide a holistic management.

INTRODUCTION
Catatonia is a neuropsychiatric condition characterised by alterations in motor behaviour, thought, affect and vigilance. Originally conceptualised by Kahlbaum in 1868, catatonia was initially incorporated in dementia praecox in 1893 by Kraepelin and later in schizophrenia by Bleuler. Over the years, catatonia has been found to be associated with not only schizophrenia but also mood disorders and a multitude of organic brain conditions such as epilepsy, neurodegenerative disorders, tumours, haemorrhage and neuro-infections. Recent advances have led to the understanding that among psychiatric conditions, catatonia occurs more frequently with mood disorders than with non-affective psychosis such as schizophrenia. There are examples in the literature that point to the recurrent nature of catatonia that occurs in the setting of the relapsing-remitting course of mood disorders: the catatonia understood as a phenotypical manifestation within the particular mood episode. When a patient presents with recurrent catatonia, clinicians should attempt to identify the underlying conditions that otherwise remain masked under the catatonic presentation. A detailed assessment after catatonia management is always needed to diagnose the underlying psychiatric disorder, which will entail a holistic management. There is a paucity of literature that suggests catatonia to be the first phenotypical manifestation of a mood episode. We hereby report a series of two cases in which catatonia was the first symptom manifestation of all the mood episodes in both along with its persistence in a similar fashion in all the episodes. We will also briefly review the existing literature about recurrent catatonia and its association with episodes of mood disorders.

CASE HISTORY
Case 1
A 25-year-old Hindu unmarried man with unremarkable family and personal history but with past two episodes of suggestive catatonia presented with a 3-day history of remaining mute, maintaining a posture for long and refusing to interact with others along with refusal to eat and drink. His initial mental status examination (MSE) revealed catatonia (mutism, negativism and posturing) (Bush-Francis Catatonia Rating Scale (BFCRS) score=16), and it precluded further examination of his affect, thought and perception.
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He was admitted, and on investigations, his haematological and biochemical parameters (complete haemogram; renal, hepatic and thyroid function tests; serum electrolytes; creatinine phosphokinase) and MRI of his brain were within the normal range. He was initially managed with the injection of lorazepam 4 to 8 mg/day, which led to remission of catatonia in 3 days (BFCRS score=0).³ On remission of catatonia, he started to show goal-directed hyperactivity and talkativeness with thought content revealing grandiosity, which gradually increased for the next few days. During this time, his MSE revealed increased speech output with prolixity, increased psychomotor activity, delusion of grandiosity and no perceptual abnormalities (Young Mania Rating Scale (YMRS) score=26).⁴ Further clarification of the past two catatonia episodes revealed that in each episode, he would first present with the similar type of symptoms such as mutism, posturing and negativism (and stupor as an additional symptom in the first episode), which would be the initial presentation and have an abrupt onset with a dramatic response to treatments prescribed by a psychiatrist that would ultimately manifest in a similar mental state similar to the index episode (ie, mania). However, in none of these episodes, he continued with the psychiatrist’s advice. Considering the past episodes and the current index presentation, he received a provisional diagnosis of bipolar affective disorder, current episode mania with psychosis according to ICD-10,⁵ and he was started on tablet sodium valproate 1 g/day, quetiapine 100 mg/day and lorazepam 2 mg/day, with which he showed significant improvement in the next 3 weeks (YMRS score=11).¹ He was discharged on this regimen and on a subsequent follow-up after a month, he was found to maintain well.

Case 2

A 42-year-old Hindu married man with no significant family and personal history but with a history of catatonia thrice in the last 4 years presented with a 5-day history of remaining mute, not taking food and water, not interacting with family members and moving away from any interventions. On admission, his vital parameters were within the normal range, and his initial MSE revealed catatonia (posturing, mutism, active negativism) (BFCRS score=18).³ His blood investigations (complete haemogram; renal, hepatic and thyroid function tests; serum electrolytes) and his MRI of the brain were within the normal range. His catatonia was not responding to lorazepam (given up to 12 mg/day) even after the third day, and therefore, he was started on modified electroconvulsive therapy (ECT). After the second ECT, his catatonia resolved significantly (BFCRS score=0).³ He however got a call from his wife 4 months later after a month later he was found to be well. We however got a call from his wife 4 months later that there was another similar episode for which he was taken to a tertiary care psychiatric hospital and was managed in similar lines. We however planned to follow him prospectively having a suspicion whether he may land up in a bipolar disorder diagnosis in future.

DISCUSSION

Catatonia has long been thought to be associated more with schizophrenia, starting from the era of Kraepelin and Bleuler. However, Kahlbaum, who initially gave the descriptions of catatonia along with its types (table 1) in 1868, conceptualised it to be associated mostly with mood disorders.¹ Over the years, its association with mood disorders has been strengthened. Catatonic signs have consistently been reported with mood disorders in 13% to 31% cases.⁷ Various systematic studies reported that, among hospitalised manic patients, more than 25% meet the criteria for catatonia, whereas among patients with catatonia, more than half meet the criteria for a manic-depressive illness.⁸ The seminal paper of Barnes et al also favoured this association. Thirty-six per cent of 25 patients with catatonia whom they described over a span of 12 years had affective disorders, whereas only 4% had a diagnosis of schizophrenia. The remaining (60%) had either an organic brain

Table 1  Kahlbaum’s catatonia types¹⁵

| Kahlbaum’s catatonia types | Clinical descriptions |
|---------------------------|----------------------|
| Catatonia mitis           | Mild form, rapid development, associated with depression, good prognosis. |
| Catatonia gravis          | Severe form, relatively rapid development, associated with pure mania and mixed states, good prognosis. |
| Catatonia protracta       | Severe form, insidious onset, bizarre movements and behaviours, chronic, poor prognosis. |
condition or no concomitant diagnosis. The affective disorders were all recurrent depressive disorders having mostly recurrent catatonia in the episodes. What was also striking was the fact that such a presentation (recurrent catatonia with depression) was also present in their family members, which points to its high genetic loading. Usman et al. did a retrospective chart review of 13968 patients who presented with catatonia in their tertiary care neuropsychiatry hospital in Nigeria and found an increase in the diagnosis of mood disorders (from 10% to 20.7%) in a span of 20 years alongside a decrement of a schizophrenia diagnosis (from 82.5% to 53.4%) during the same period. The patients with mood disorders were mostly females, but there was no significant difference in the sociodemographic profile in these two diagnostic groups. Parker et al. also held the same opinion in their review, reporting most inpatients with catatonia having a comorbid mood disorder (36% to 63.2%) than a psychotic disorder (29.8% to 32%). Within bipolar disorder, catatonia has been reported to be more associated with mixed states in comparison with a manic episode as reported in two studies that found 61% and 73% of patients with catatonia have an index mixed episode, respectively. Cata-tonia has also been reported with atypical presentations of catatonia in mood disorders, especially manic. According to them, catatonia is mostly associated with mixed states, they are transient and are markers of severity of mood disorders but still have a better prognosis than catatonic schizophrenia. The two cases discussed above had mood disorders (bipolar effective disorder in case 1 and recurrent depressive disorder in case 2), and for both of them, the initial symptoms were those of catatonia, the treatment of which ultimately led to the emergence of the underlying mood symptoms. What was also strikingly noticeable was the recurrence of catatonia in a similar fashion (their initial most presentation and the same catatonic signs and symptoms) in all their subsequent mood episodes. Some important observations were pointed out: (1) catatonia can mask a mood episode, which is consistent in subsequent episodes whether it is a bipolar illness or a recurrent depressive disorder; and (2) the similarity in its pattern and typology in the later episodes, for example, mutism, agitation and withdrawal in our cases. The current report thus points to the need for looking beyond such type of catatonic presentations, more so when they are the first presenting symptoms and occurring recurrently in future episodes. A thorough assessment after treating the initial catatonia can unmask an affective disorder that warrants a holistic management. Usman et al. in their retrospective chart review reported that among many catatonic symptoms, mutism (87.5%), staring (50%), withdrawal (43.8%) and posturing/catalepsy (56.3%) were the most frequent and most consistent symptoms in patients with mood disorder when compared with schizophrenia. Francis et al. however, reported a consistency of motor symptoms across all the catatonic episodes in the five patients they reported, although only one received a diagnosis of mood disorder among them. There are pieces of literature evidence to support that the ‘retarded’ type of catatonia is highly associated with mood disorders. Our cases as described above both had ‘retarded’ type of catatonia with mutism, posturing and withdrawal being their manifested signs. Peralta et al. endeavoured to dissect the catatonic phenotype in psychosis and mood disorders and came to the conclusion that ‘psychomotor retardation’ was the most frequent catatonic sign in their cohort followed by rigidity and mutism, which had the highest familiarity. Lin et al. in their 15-year retrospective record review of 30 patients with a history of recurrent catatonia, found that 6.7% and 16.7% of patients had a diagnosis of bipolar disorder and depressive disorder, respectively. All had ‘retarded’ type of catatonic symptoms and signs viz stupor and mutism, the Kahlbaum syndrome. The higher prevalence of mutism, stupor, and withdrawal was also reported in a recent study on a naturalistic setup involving 26 patients with catatonia having bipolar disorder. All these evidences point to a possible association of retarded catatonia with mood disorders and a consistency of their manifestation across future mood episodes (whether manic or depressive). However, further research is warranted that will longitudinally follow up patients with mood disorders, who had catatonic presentation during the episodes, to systematically look into the consistency of the catatonic signs and symptoms in their lifetime mood episodes.

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