Whether Delirium Acts as ECT?
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Abstract

Delirium is defined by the acute onset of fluctuating cognitive impairment and a disturbance of consciousness with reduced ability to attend. It may occur at any age but is more common in older adults. By virtue of working in the psychogeriatric hospital (Department of Geriatric Mental Health, King George’s Medical University, Lucknow, India), authors experienced that many elderly patients with psychiatric illness develop delirium because of various reasons. Subsequently, we observed that psychiatric manifestations of the patients improved significantly following recovery from delirium. A series of such four cases (case 1- Depression with psychotic features, 2- Manic episode, 3- Bipolar affective disorder current episode mania and 4- Depressive episode) later developed delirium has been presented and discussed in this paper. ICD-10 criteria were used for the diagnosis of delirium and psychiatric illnesses. Average stay of these patients in the hospital was 10 days. We found that signs and symptoms of these psychiatric disorders disappeared almost completely following the recovery from delirium. Therefore, it can be assumed that delirium potentially acts similar to ECT. By reporting such case we open a new dimension of research for effective treatment of the psychiatric patients. Inducing delirium in a patient is unethical but understanding pathophysiology behind it will certainly open the door for better cure of the ailing psychiatric patients.

Keywords: Delirium; Older adults; ECT

Introduction

Delirium is defined by the acute onset of fluctuating cognitive impairment and a disturbance of consciousness with reduced ability to attend. According to ICD-10 Delirium is an etiologically nonspecific syndrome characterized by concurrent disturbances of consciousness and attention, perception, thinking, memory, psychomotor behaviour, emotion, and the sleep-wake cycle. It may occur at any age but is most common after the age of 60 years [1]. None of the age is immune to the delirium, it can occur in anyone at any age but it is more common in older adults. In a review, it is reported that 15% to 30% of elderly patients will have delirium on admission to hospital and up to 56% will develop delirium during their stay [2]. The point prevalence of delirium in the community is 1.1% amongst the general population aged over 55 years and up to 14% in those over 85 years [3]. Since delirium is an acute medical condition, its prevalence can only be studied in hospital settings. A meta-analysis of 42 studies found delirium to be prevalent on admission in 10-31% of medical inpatients, and to occur in the hospital in 3-29% [4]. Fang et al. [5] reported that the overall prevalence of delirium in the community is just 1-2%, but in the setting of general hospital admission this increases to 14-24%. The incidence of delirium arising during a hospital stay ranges from 6% to as high as 56%, and this incidence is even higher when more specialized populations are considered, including those in postoperative, intensive-care, subacute and palliative-care settings [4,6,7]. Postoperative delirium occurs in 15–53% of surgical patients over the age of 65 years [8] and among elderly patients admitted to an Intensive Care Unit (ICU) the delirium incidence can reach 70-87% [9]. Advanced age and cognitive impairment are major risk factors for the development of delirium. About 15% of elderly inpatients in acute medical-surgical services have delirium at any time, and the rate is even higher among nursing home patients. Approximately 30 to 40 percent of hospitalized patients older than age 65 have an episode of delirium, and another 10 to 15 percent of elderly persons exhibit delirium on admission to the hospital [10,11].

The patient with delirium presents with a wide range of psychiatric symptoms making it difficult to distinguish with other psychiatric disorders such as schizophrenia. Thought content disturbances of delirium entails themes from the immediate environment and circumstances, hallucinations used to be frequently visual rather than auditory and formal thought disorder typically comprised poverty of thinking and illogicality [12]. Clouding of consciousness and inattention found in delirium may be helpful in differentiating it from psychosis. The major causes of delirium are neurological disorders, systemic disease, metabolic disorders and either intoxication or withdrawal from pharmacological or toxic agents. Cause of delirium can also be multifactorial. The permeability of blood brain barrier is impaired by ageing and dementia, which may predispose the patient to delirium. The major neurotransmitter hypothesized to be involved in delirium is acetylcholine, which is decreased in the brain. One of the most common causes of delirium is toxicity from too many prescribed medications with anti-cholinergic activity. Noradrenaline, serotonin and glutamate have also been implicated in the causation of delirium. Neurotransmitters involve for the causation of the psychiatric disorders are same which are involved in the delirium [13-15]. It was observed that psycho-geriatric patients are at receiving end of development of delirium because of multiple drugs prescribed for the co-morbidities along with psychotropic medications. Development of delirium during course of treatment is frequent in psychogeriatric patients as compared to their younger counterparts. During course of treatment of the psychiatric patients who developed delirium in our hospital, we observed an interesting phenomenon. These patients had a psychiatric illness before the onset of delirium as per the ICD-10 criteria. Somehow after recovery from the delirium most of the psychiatric symptoms were disappeared. Written informed consent was taken from the patients and their family members hospital to present and publish the case for the benefit of the scientific community. Their names are written as Mr. A, B, C and D to maintain confidentiality. A series of 4 cases is being presented below and discussed.

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Case 1

Mr. A, 61 year old male presented to the Department of Geriatric Mental Health, (DGMH) OPD with chief complaints of low mood, restlessness, decreased interaction, decreased appetite, decreased sleep and suspiciousness for the last 1 month. On mental status examination he was depressed. He was having persecutory delusions and auditory hallucinations. On enquiry he complained that the voices are coming from the outside, they abuse him and threatened to kill him. The provisional diagnosis of depression with psychotic features was entertained and was prescribed escitalopram 10 mg, olanzapine 5 mg and lorazepam 2 mg per day. His routine blood investigations and CT Head were within normal limits. On follow-up after 15 days of treatment he was confused. He developed visual hallucinations that insects are crawling on the ground. His sleep cycle was reversed. Possibility of drug induced delirium was kept. His CBC, electrolytes, blood urea, serum creatinine and CT scan Head were within normal limits. Delirium was managed by withdrawing all the psychotropic medications prescribing multivitamins and electrolyl powder. On follow-up next week Mr. A, recovered from the delirium. Interestingly his all psychiatric symptoms were improved. Family members reported him as normal as he was earlier. On follow-up after three weeks, there was re-emergence of the symptoms but of very low intensity. So he was prescribed escitalopram 10 mg and olanzapine 5 mg and he has been maintaining well on this treatment.

Case 2

Mr. B, 64 year old retired male presented to the DGMH OPD with chief complaints of over talkativeness, aggressiveness, doing big claims and decreased need for the sleep for the last 1 month. He often suspicious about the son that he had taken out money from his account for which he used to become agitated. On mental status examination he was elated, having delusions of grandiosity and theft. On detailed evaluation it was his suspiciousness about the money being withdrawn from his account has started little earlier than the overtalkativeness. Provisional diagnosis of 1st episode mania was made. He was hospitalised and he was given haloperidol and promethazine in the form of injectibles. He was not accepting medicines and food orally so the injectibles were continued further for the 2 more days but on 3rd day of hospitalization he developed confusion. He was not oriented to time and place. His sleep cycle was reversed and used to pick something from his belongings. Possibility of drug induced delirium was kept and injectibles were stopped. His all blood parameters were within normal limits. No abnormality detected on CT scan head except cerebral atrophy. He was kept on the Multivitamins and electrolyl powder. Hydration was maintained. Finally he recovered from the delirium but interestingly all his symptoms for which he was hospitalized were gone. He was in the state of remission and was discharged. Later on he was discharged on Mood stabilizer (sodium valproate) to prevent further relapse.

Case 3

Mr. C, 63 year old male presented to DGMH OPD with chief complaints of overtalkativeness, aggressiveness, irritability, blaming his family members about the mistreatment for last 15 days. On mental state examination his psychomotor activity was increased and mood was irritable. He had similar episode 2 years back which got treated from the DGMH and completely recovered within 2 months and discontinued the treatment without consultation. Provisional diagnosis of Bipolar Affective Disorder current episode Mania was entertained. He was hospitalized. Combination of oral haloperidol 15 mg and combination of chlorpromazine (200 mg) and trihexiphenidyl 8 mg was started but instead of improving, his symptoms started worsening after initial glimpse of improvement. He was quite agitated, trying to jump from the bed, was not oriented to time, place and person. He often visualizes small creatures creeping on the ground. Diagnosis of drug induced delirium was made and all the drugs were stopped and promethazine 25 mg once daily at night, electrolyl powder and multivitamins were prescribed to him. CT scan Head and routine blood investigations were within normal limits. He recovered from the delirium but interestingly he reached into remission. Later on he was discharged on Mood stabilizer (sodium valproate) to prevent further relapse.

Discussion

It is well proven that a marked increase in slow wave activity in inter-period on EEG may last for weeks to months after completion of an ECT-course [16,17]. ECT cumulatively leads to Electroencephalogram (EEG) changes characterized by increased amplitude and reduced frequency (‘delta’ activity) somewhat similar to delirium [18]. It is likely that such EEG changes are correlated with improved clinical response rates and reduced likelihood of relapse [19]. EEG characteristics of delirium include slowing or dropout of the posterior dominant rhythm, generalized theta or delta slow-wave activity, poor organization of the background rhythm, and loss of reactivity of the EEG to eye opening and closing [20]. It might be possible on the above theory that ultimate effect of ECT is not because of seizure activity but because of the postictal confusion which causes delirious state in the patient. It means that if by any mean psychiatric patient reach to delirious state the psychiatric symptoms may be subsided. Delirium is a common and serious acute neuropsychiatric syndrome with core features of inattention and cognitive impairment, and associated features including changes in arousal, altered sleep-wake cycle, and other changes in mental status [21]. Hypotheses about the pathophysiology of delirium are speculative and largely based on animal research. According to the neurotransmitter hypothesis, decreased oxidative metabolism in the brain causes cerebral dysfunction due to abnormalities of various neurotransmitter systems. Reduced cholinergic function, excess release of dopamine,
norepinephrine, and glutamate, and both decreased and increased serotonergic and gamma-aminobutyric acid activity may underlie the different symptoms and clinical presentations of delirium. Furthermore, severe illness and physiologic stress may give rise to modification of blood-brain barrier permeability and increased activity of the hypothalamic-pituitary-adrenal axis. These circumstances possibly also contribute to changes in neurotransmitter synthesis and release of cytokines in the brain, and consequently to the occurrence of delirium. Elderly patients are more at risk for developing delirium, very likely due to age-related cerebral changes in stress-regulating neurotransmitter and intracellular signal transduction systems [14]. Psychiatric disorders are result of disturbances in above mentioned neurotransmitters either increased or decreased. If neurotransmitter system is disturbed possibly they get rearranged on recovery and ultimately result in remission. Three of the four patients developed drug induced delirium and on recovery the psychiatric symptoms were completely absent. One of the four patients hospitalized for delirium with background history suggestive of depression. On recovery from the delirium there were no symptoms suggestive of depression. One of the four patients relapsed again on follow-up suggestive that recovery is not lifelong and it has to be relapsed after few days if not given psychotropic medication as we see in the cases of ECT. On the basis of above discussion we can say that delirium acts as ECT. If a less invasive substitute for ECT is desired, the priority would be to explore underlying mechanism of delirium which has similar effect on psychiatric disorders. By reporting such cases we open a new dimension of research by virtue of which we can treat effectively the psychiatric patients. Inducing delirium in a patient is unethical but understanding pathophysiology behind it will certainly open the door for better cure of the ailing psychiatric patients.

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