Caution in Co-administration of Aripiprazole and Lithium in Patients with Learning Disability

Abstract
Bipolar disorder is a complicated syndrome. The first line of bipolar disorder treatment is pharmacotherapy. Aripiprazole is a safe and effective medication in acute phase of bipolar disorder. This report is about a patient who suffers from comorbidity of bipolar mood disorder and learning disability. The patient developed with extra pyramidal symptom in combination of aripiprazole and lithium in doses lower than therapeutic dosages. This finding can be explained by susceptibility of a traumatized brain to neurological side effects of antipsychotics and drug interactions.

Introduction
Bipolar disorder is a complicated syndrome with multifactorial etiology such as neuro-endocrine, genetic, and environmental factors [1]. The first line of bipolar disorder is pharmacotherapy [2]. In acute phase of mania, lithium, anticonvulsants [3] and antipsychotic agents are suitable for symptom management [4]. Aripiprazole as a third generation antipsychotic, is a partial agonist of the dopamine D2 receptor, a partial agonist of the serotonin 5-HT1a receptor, and antagonist of the 5HT2a receptor sites [5]. Aripiprazole has been defined as a dopamine system stabilizer [6]. Due to its unique psychopharmacological profile the hypo-dopaminergic state is limited by this medication. Aripiprazole was approved by the FDA in 2004 for treatment of bipolar disorder [7]. An important meta-analyses which was designed by Dian-Jeng Li and co-authors confirmed the efficacy or safety of aripiprazole in manic episodes [8].

There are several studies that show long-term administration of aripiprazole in combination to lithium or valproate in bipolar mania is well tolerated and safe and this improvement in functioning is maintained [9,10]. One of the side effects of aripiprazole is akathisia [11]. The frequency of occurrence of other extra pyramidal symptom-related events such as acute dystonia, parkinsonian syndrome, akathisia, akinesia, rabbit syndrome, tardive dyskinesia or neuroleptic malignant syndrome in short-term, and in long-term, and in patients with schizophrenia or bipolar mania is similar in both the placebo and the aripiprazole-treated groups [11].

Case Report
The patient was 30 year old single lady, she couldn't finish primary school. She lives with her mother and could only do simple tasks such as individual care. In the other word she was a case of learning disability. Who came with increased energy, increased activity level, risky behaviors such as involvement in sexual contact, paranoid delusion about her neighbor and severe agitation and aggression. These symptoms began about 2 month's prior admission. This was the first episode of illness. Till now she never experienced depressive, hypomanic or psychotic episodes. All of the work ups such as biochemistry evaluation, thyroid function test and brain imaging was done and no abnormality was detected. With impression of Bipolar disorder type I, pharmacotherapy began. She received lithium 150 mg QHS and Aripiprazole 5 mg QD. The therapeutic response in management of agitation and control of hyperactivity was dramatic. During first week of therapy, agitation and aggression improved and paranoid delusions became shaky.

8th day of admission she became febrile (38 C axillary) and she developed with tremor, sialorrhea and drowsiness. All of the medications became hold and IV hydration started. The most important differential diagnosis was neuroleptic malignant syndrome. In lab data, white blood cells and creatine phosphokinase were normal. Vital signs were stable and no change in blood pressure or pulse rate were detected. After hydration the patient became afebrile. So neuroleptic malignant syndrome ruled out. The other differential diagnosis was extrapyramidal symptom. By conservative management such as administration of Linderal 10 mg PO BID, hydration, and supportive care after 4 days the tremor and sialorrhea completely improved.

At this time the patient was oriented to time, place and person. But she was restless and elevated mood was obvious. She reported auditory and visual hallucination. So management of symptoms with antipsychotic was necessary. This time quetiapine with dosage of 12.5 mg PO BID began for the patient. After 10 days the psychotic symptoms disappeared and mood swing became partially controlled. Quetiapine 25 mg po BID were effective for control of symptoms and the patient discharged from the psychosomatic ward at the end of 38th day of admission.
Discussion

Although akathisia is not uncommon in patients receiving aripiprazole, other extrapyramidal side effects of this drug were equivalent to placebo [11]. Patients who suffer from learning disability, benefit from atypical antipsychotics for management of agitation and aggression [12]. In these patients, comparison of typical and atypical antipsychotics, show that akatasi is more prevalent with first generation antipsychotics, but no differences is reported in happening of dystonia and Parkinsonism/Dyskinesia [13]. It means that both groups of antipsychotics can cause parkinsonism. Among atypical antipsychotics, the least prevalence of Parkinsonism is reported with Aripiprazole [14].

When you search drug interactions between Aripiprazole and lithium, it is reported that in patients with normal learning function this combination can lead to dizziness, drowsiness, confusion, and difficulty concentrating. Elderly, may experience dysfunction in thinking, judgment, and motor coordination [15]. In the other word this combination is defined as a safe and effective treatment for management of acute mania in bipolar disorder [1-18]. In patients who suffers from schizophrenia or schizoaffective disorder, therapeutic doses of lithium had no clinically significant effects on the pharmacokinetics of aripiprazole [19].

Although animal studies present the effective role of antipsychotics on cognitive function in cases with brain injury [20]. There are case reports that show more severe neurologic side effects with aripiprazole in patients with brain injury and learning disability [21,22]. These similar findings can be explained by susceptibility of a traumatized brain to neurological side effects of antipsychotics and drug interactions. This problem even can present in cases who use medications with lesser than therapeutic dosages. Of course designing of cross sectional and clinical trial studies can help us to know more about the adverse effect of medications in patients with learning disorders.

References