Clinical Findings Showing That Non-Convulsive Electric Shock Administration for Patients Receiving Clozapine Therapy for Schizophrenia May Contribute to Stability of Granulocyte Number, the Most Critical Drug Side-Effect

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Abstract

Two treatment-resistant schizophrenic patients were administered clozapine, but showed a decrease in granulocytes. As an alternative treatment, we administered non-convulsive Electric Shock Therapy (ECT) (modified-ECT: m-ECT) to both and the number of granulocytes increased to a level considered safe for clozapine ingestion. Then, under continuation of m-ECT, we gradually increased the dose of the drug. After 10-14 m-ECT sessions in both patients, the number of granulocytes became stable at approximately 5000/mm3 and m-ECT was completed. Our findings show the possibility of treatment for therapy-resistant patients who are unable to receive clozapine because of reduced granulocyte number.

Keywords: Clozapine therapy; Electric shock; Schizophrenia; Granulocytes

Introduction

Schizophrenia is a common psychotic disorder that severely affects human personality, mainly in the manner of thinking; including delusions, inappropriate hostility, strange behaviour, and others, with no proven cure. Although side-effects have been shown, clozapine has been recently reintroduced as drug therapy for the disorder because of its remarkable effects on normalizing personality (improved rapport, control of hostility increased quality of life, etc.) [1]. However, an unfavourable effect of clozapine administration is a decrease in granulocytes, especially neutrocytes, which leads to a large portion of candidates not able to use the drug [2].

Non-convulsive Electroconvulsive Therapy (ECT) and modified ECT (m-ECT) have been introduced widely for daily psychiatric therapy. Although ECT is generally used for mood disorders, it has also been applied for controlling schizophrenic psychomotor excitement and other unexplainable actions by patients [3]. M-ECT is generally preferred as compared to ECT, because the electric shock can be applied without resultant convulsions.

In the present study, we report results of 2 patients who received m-ECT while being administered clozapine because of concern regarding the instability of granulocyte numbers. Both showed improvement in granulocytes, which enabled safe use of clozapine with favourable effects. In addition, we discuss this combined therapy as a means for more wide use of clozapine for therapy-resistant patients and the mechanism of its effects.

Case 1: Male, 31 Years Old, Hebephrenic Schizophrenia

Three years prior, the patient was admitted for deterioration of personality (difficult to communicate with others due to meaningless language, inappropriate responses, discontinuation of medication or nutrition, delusions including personal odor, etc.).

Moreover, unexpected hostility had led him to management in an isolated protection room for approximately 3 years regardless of additional therapy, we also began m-ECT (450 V sin implus, 0.2-0.45 × 504 mC) for relaxation of the cumbersome symptoms. Unexpectedly, while clozapine administration continued, blood tests conducted after starting m-ECT showed that granulocytes recovered to approximately 10,000/mm3.

Thus, we decided perform blood testing on the day following the m-ECT sessions, which continued for approximately 3 months. As a result, the patient was able to socialize with other patents outside of the protection room and was freed from his self-smell delusion. As compared to prior to beginning clozapine and m-ECT therapy, the Positive and Negative Symptom Scale (PANSS) score decreased from 121 to 92. Presently, m-ECT is not being given and clozapine administration is continuing without a decrease in granulocytes (Figure 1).
Case 2: Female, 51 Years Old, Delusional Schizophrenia

The patient had been suffering from delusional schizophrenia for more than 30 years. In the early period, personality deterioration was not apparent and with antipsychotic drugs she was able to function as a housewife. However, recently the patient began to show difficulty with maintaining regular daily life due to delusional thinking, such as the feeling that someone was intruding into her house, and was unable to maintain regular drug administration. In addition, she showed bizarre behaviour such as going out of the house without wearing clothes. After noting no satisfactory effects from therapy with more than 2 types of atypical antipsychotics, we decided to begin an administration of clozapine. In the early period of clozapine therapy, the patient was unable to get along with other patients and stayed in an isolated protection room. Moreover, granulocytes showed a decreasing trend to less than 4000/mm$^3$. 

However, we were concerned about discontinuation of drug therapy and decided to start m-ECT (450 V sin impulse, 0.2-0.45 × 504 mC) sessions while maintaining a minimum dose of clozapine. Thereafter, granulocytes increased and stabilized at greater than 5000/mm³ and her fear of other people gradually disappeared.

Following 10 sessions of m-ECT, the patient began to participate in regular daily life in the hospital without apparent side-effects from the drug, thus we stopped m-ECT and administration of antipsychotics other than clozapine. As compared to prior to beginning clozapine and m-ECT therapy, the PANSS score decreased from 109 to 79, while granulocytes have been maintained at a safe level (>5000/mm³) (Figure 2).

Discussion

The present 2 patients were treated with m-ECT as an alternative complementary therapy method. Although the combined use of m-ECT and clozapine therapy had been reported in our country, the report did not inform us of the benefit gain from the combination [4]. Our results do not provide concrete evidence for the effects of m-ECT in patients receiving clozapine. Nevertheless, it is possible that such combined therapy decreases stress induced by drug therapy shown by a reduction in number of granulocytes.

Previous studies have found that ECT enhances immunological ability in humans. For example, it was reported that the activity of the natural killer granulocytes was activated or the number of leukocytes slightly increased [5,6]. However, to the best of our knowledge, no reports have directly shown an increase in number of granulocytes or the mechanisms of that phenomenon. The capability for ECT to enhance immunological ability remains speculative, though cytokine family members show changes in amount in blood and are candidates for maintaining the number of granulocytes [7].

An increase in number of granulocytes by m-ECT remains to be proven. Furthermore, the mechanisms by which an electric shock affects the immunological system, how long the increase in number of granulocytes persists, and whether the phenomenon observed in our patients is universal are not apparent.

Should clozapine be considered as the last choice for therapy-resistant schizophrenic patients, safer use must be investigated. We used the present combined therapy in 2 therapy-resistant patients who could not continue the drug because of a decrease in granulocytes along with continuation of symptoms and requirement of complementary therapy. Although the number of subjects was small, a recovery of the number of granulocytes was observed in each. In a future report, we intend to present additional information regarding these cases. For that purpose, more several cases were given the therapy that combined m-ECT and clozapine administration. First of all, we intended to know how long does it take for the patients showing the decrease of granulocyte to stabilize its number by m-ECT therapies. Moreover, if m-ECT increases the granulocytes, it should be apparent when the effect emerges after the therapy is done. We now are planning the multiple measurements of the number under permission by the patients. Furthermore, influences on the amount of candidate substances including cytokine families by m-ECT will be studied. On the other hand, the genome risk factors for Clozapine-Induced Granulocytopenia (CIG) are gradually reported. Iwata and colleague recently presented the specific risk factor Japanese patients [8]. Therefore, in the future we’ll have to try research about the genome risk factors for CIG of patients who were done the combined therapy successfully to clarify the relation between the risk factor and our combined therapy.

References