行政院國家科學委員會補助專題研究計畫成果報告

雙極性情感疾病患者之躁症狀態的免疫調節作用 (II)

Immunological Regulation in Bipolar Patients with Manic Episode (II)

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一、中文摘要

有許多文獻證據顯示中樞神經系統與免疫 系統間有密切的交互作用,比較特別的是有 關精神壓力與精神疾病患者有免疫異常情 形的描述。目前學者提出精神裂症患者有單 核細胞與 T 細胞活化的假說。躁鬱症是一種 可能會出現躁狀精神狀態的情感性疾病,目 前有關躁症的免疫研究較少且結果仍有爭 議,因此本實驗的目的主要是研究躁症患者 與正常控制組之可溶性的 sCD4, sCD8, IL-1RA, CC16, sTfR 的表現情形。SCD4 躁症患 者與正常控制組相比有較高, sCD8, IL-1RA, sTfR 則有意義的(p<0.05)高於正常組。CC16 則低於緩解期。總而言之,這些結果顯示雙 極性情感疾病的免疫病生理機制可能受症 狀嚴重程度的影響,且調節機轉與正常人不 同。

關鍵字:

關鍵詞:躁症,可溶性 CD4,可溶性 CD8,

CC16, IL-1RA, 可溶性 TfR

Abstract

Several studies suggest CMI is altered in

bipolar patients with manic episode, although findings have been contradictory. The inconsistent results may due to the effect of age, sex, medications, and severity of manic symptoms. In this case-control study we investigate the bipolar patients in manic episode as well as in consequent remission regarding the plasma soluble CD4 (sCD4) and sCD8 the plasma IL-1 concentration, receptor anatagonist (IL-1RA) and clara cell protein 16 (CC16) level and the soluble transferrin receptor (sTfR) level. The plasma sCD4 level in manic episode is higher than in that of but did not reach controls statistical significance. Whereas, the sCD8 level in bipolar patients in manic episode as well as in remission is significantly higher (p<0.05) than The levels of IL-1RA both in in controls. bipolar patients in manic episode as well as in remission is significantly higher (p<0.05) as compared to controls. The mean levels of CC16 in bipolar patients in manic episode is significantly lower than that of the remissions The sTfR concentration in manic (p<0.05). episode as well as in remission is significantly higher than controls (p<0.05). Basing on literature reviews, the increased levels of sCD8. but not sCD4 in manic episode suggests activation of cell-mediated immunity. reduced CC16 production and increase production of IL-1RA of bipolar disorder in acute mania and consequent remission is similar to that of schizophrenia. An increase sTfR level is significantly higher than controls which was also observed in schizophrenia. Taken together, the activation of CMI in bipolar disorder is observed, the immune-inflammatory variables in acute manic is different than controls suggest there is a immune-modulation of bipolar disorder may vary from normal.

INTRODUCTION

Data suggested that activation of the monocytic and T-cell arms of cell-mediated immunity (CMI) play a role in the pathophysiology of schizophrenia. However, reports on the assessment of immunity in mania are limited and remain controversial.

Base on our previous study, increased lymphocyte proliferation and significantly increased plasma concentrations of sIL-2R but not sIL-6R in manic patients than in normal controls, suggested activation of CMI in mania

patients. Since age, gender, and symptomatic severity have effects on immunity in patients with mood disorder, it is probable neither evaluating the immune parameters after a well-defined remission period nor comparing them with age- and sex-matched controls led to the inconsistent results.

The aim of the present study was to examine the plasma sCD4, sCD8, TfR, CC16 and IL-1RA concentrations in patients with mania and in normal controls.

MATERIALS AND METHODS

Subjects

bipolar Acute in-patients diagnosed disorder, manic (DSM-IV) with YMRS scores \geq 30, aged \leq 45 years and free from acute infections or chronic diseases were recruited. During the index hospitalization, all the patients were treated with lithium and typical antipsychotics (haloperidol or chlorpromazine). Follow-up blood samples of patients were collected while in consequent remission (YMRS scores < 12) free from any sign of depression. Ageand gender-matched healthy control subjects without any Axis I psychiatric disorder were recruited.

Plasma sCD4, sCD8, TfR, CC16, and IL-1RA level analysis The plasma levels of sCD4, sCD8, TfR, CC16 and IL-1RA level were measured in duplicate using ELISA method.

RESULTS

The plasma sCD4 level in manic episode is higher than in that of controls but did reach statistical significance. Whereas, the sCD8 level in bipolar patients in manic episode as well as in remission is significantly higher (p<0.05) than in controls. The levels of IL-1RA both in bipolar patients in manic episode as well as in remission is significantly higher (p<0.05) as compared to The mean levels of CC16 in controls. bipolar patients in manic episode is significantly lower than that of the remissions (p<0.05). The sTfR concentration in manic episode as well as in remission is significantly higher than controls (p<0.05).

CONCLUSION

Based on comparison between acute mania and consequent remission of bipolar individuals, a major finding of this study is that manic episode is accompanied by significantly elevated plasma sCD8, TfR, and IL-1RA levels. Since, increased plasma sCD8 and TfR levels are an unequivocal index of T cell activation. These results indicated that activation of cell-mediated immunity

occurs in acute mania. Furthermore, the plasma sIL-1RA levels was significantly elevated in bipolar mania than in normal controls. These finding suggests that the activation of the monocytic arm of CMI may occur in this illness.

Unmedicated patients shown significantly increased sCD4, TfR and lower plasma CC16 than normal controls. Patients with medication shown significantly increased TfR and IL-1RA as compared to controls.

Taken together, in this study bipolar patients in acute mania are characterized by T lymphocyte and monocyte activation as evidenced by elevated plasma sCD8, TfR and IL-1RA levels. In patients with bipolar disorder, the immunomodulatory mechanism may vary from normals.

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