Evaluation of TGF β1, IL-8 and Nitric Oxide in the Serum of Diffuse Axonal Injury Patients and Its Association with Clinical Status and Outcome

Difüz Aksonal Yaralanma Hastalarının Serumunda TGF β1, IL-8 ve Nitrik Oksit Değerlendirilmesi ve Klinik Durum ve Sonuç ile İlişkileri

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ABSTRACT

AIM: The aim was to evaluate the level of interleukin 8 (IL-8), transforming growth factor β1 (TGF β1) and Nitric oxide (NO) in diffuse axonal injury (DAI) and its association to the outcome and clinical status.

MATERIAL and METHODS: This cross-sectional study was conducted on 20 patients with DAI and 20 patients with multiple traumas without head injury and 20 healthy subjects as controls. Blood levels of IL-8, TGF β1 and nitric oxide in the 1st, 2nd, 3rd and 7th days of injury were measured. Glasgow coma scale (GCS) of patients was recorded. The patients' outcome was evaluated by Glasgow Outcome Scale (GOS).

RESULTS: The level of TGF β1 was increasing during the admission and had the maximum level at the 7th day. In the DAI group, there was significant correlation between GOS score and serum IL-8 at 7th day of admission (r=-0.68, p= 0.002). In this group the GCS was found to be significantly correlated with the IL-8 concentration at 7th day of admission (p= 0.026, r=-0.55).

CONCLUSION: IL-8 has negative correlation with GCS and GOS. TGF β1 could protect the brain from cytotoxics, hypoxia and acidosis so its level comes down in brain injuries as a result of its overuse.

KEYWORDS: Diffuse axonal injury, NO, IL-8, TGF β1

ÖZ

AMAÇ: Burada amaç difüz aksonal yaralanmada (DAY) interlökin 8 (IL-8), transforme edici büyüme faktörü β1 (TGF β1) ve Nitrik oksit (NO) düzeyini ve sonuç ve klinik durumla ilişkisini değerlendirilmekti.


BULGULAR: TGF β1 düzeyi hastaneleye yatışla birlikte artış göstermiş ve maksimum düzeyde 7. günde ulaştı. DAY grubunda GOS skoru ile yattıktan sonra 7. gün serum IL-8 değeri arasında anlamlı bir korrelasyon vardı (r=-0.68, p= 0.002). Bu grupta GCS’nin yattıktan sonra 7. günde IL-8 konsantrasyonu % olumlu oranda korrelasyon gösterdiği bulundu (p= 0.026, r=-0.55).

SONUÇ: IL-8’in GCS ve GOS ile negatif korrelasyonu vardır. TGF β1 beyni sitotoksik maddeler, hipoksi ve asidozdan koruyabilir ve bu nedenle beynin yaralanmalarında fazla kullanılmasıyla düzeyi düşer.

ANAHTAR SÖZCÜKLER: Difüz aksonal yaralanma, NO, IL-8, TGF β1
INTRODUCTION

Neurodegeneration is a clinical statement in which the neurons degenerate and die after a severe sudden insult to the brain. This could happen after some acute insults including stroke, subarachnoid hemorrhage, head injury and other cerebrovascular accidents (CVA) (15).

Inflammatory processes have been shown as effective factor in the degeneration and acute injury-induced inflammation causing neurodegeneration (1, 3). An increase in the level of interleukin (IL) 8, IL-6, IL-10 and peripheral monocytes count after intracranial hemorrhage and traumatic brain injury shows the effect of these inflammatory mediators in the degeneration process and secondary insult. These mediators also have effect on multiple organ dysfunctions after severe brain injury (5,11,13,14). In addition to these inflammatory degenerative factors, increased release of growth factors, neurotrophins and anti-inflammatory mediators including transforming growth factor β1 (TGF β1) and IL-10 protects the neurons against the excitotoxocities, acidosis, hypoxia and peroxidants after acute brain injuries (12).

There are few investigations about the clinical state and outcome of these patients and their association to the blood and CSF level of protective and degenerative mediators. This study was designed to evaluate the level of IL-8, TGF β1 and nitric oxide (NO) in diffuse axonal injury (DAI) and its association to the outcome and disease severity.

MATERIAL and METHODS

This study was a prospective study on 20 patients with diffuse axonal injury in the neurosurgery department of Bahonar hospital, Kerman, Iran from March to December 2010.

Inclusion criteria were male sex, age over 18, admission to the hospital within 24 hours of the injury and Glasgow Coma Scale (GCS) less than 10. Patients with infectious disease, myocardial infarction, autoimmune, kidney and liver disease and multiple trauma were not enrolled to the study.

As control groups, 20 patients with multiple trauma without head injury (MT), and 20 healthy men were enrolled in the study.

Evaluation

The patients and controls were evaluated for blood levels of IL-8, TGF β1 and nitric oxide (NO) in the 1st, 2nd, 3rd and 7th days after injury. The blood samples were centrifuged and the serum was collected and kept at -70°C until assessment. The concentration of IL-8 and TGF β1 was measured using enzyme linked immunosorbent assay (ELIZA) method with Biosource Europe Belgium and DRG instruments GmbH Germany kits, respectively, and the NO level was measured using Griess method with Molecular Probes Europe company kit (Netherlands). The body temperature, platelet and leukocyte count and GCS of patients were recorded in every evaluation. The patients’ outcome was evaluated on the discharge day by Glasgow Outcome Scale (GOS).

RESULTS

There were 20 men with DAI and 20 MT and 20 healthy men in this study. The mean age of DAI group was 33.6±12.6 years which had no significant difference with MT (36.8±18.13) and healthy groups (25.45±4.68).

In DAI group, mean GCS at first day of injury was 7±1.29. Mean GCS in patients with DAI increased in 2nd, 3rd, and 7th days after injury to 7.85±1.81, 8.7±2.55 and 11.7 ±3.75, respectively. Mean GOS in the study group (DAI) was 4.6 ±0.68.

IL-8

The blood level of IL-8 was 76.22±31.34 at the first day and it decreased to 70.13±37 at 2nd, 66.02±33.36 at 3rd and 49.07±23.54 at 7th day of admission in the DAI group. The level of IL-8 showed increase from 1st to 2nd day of admission in the MT group but it began to decrease after the 2nd day to the least level at the 7th day (Table I). The mean level of IL-8 in 4 measurements in DAI group was higher than the healthy group (65.3±11.6 vs. 21.54±5.16, p=0.04) but less than MT group (65.3±11.6 vs. 279±124.5, p=0.014).

TGF β1

The level of TGF β1 was increasing during the admission and had the maximum level at the 7th day (Table I). The mean level of TGF β1 in the DAI group was significantly lower than the healthy group (117.8±9.4 vs. 215.1±44.9, p =0.003) but it did not have significant difference with MT group (117.8±9.4 vs. 127.5±43.7, p=0.67).

NO

The serum level of NO metabolites had the highest level at the first day and it was decreasing during the admission to the lowest level at the 7th day of admission in the DAI group. The NO concentration in the DAI group was significantly higher than the healthy group (37.9±3.4 vs. 25.2±3.37, p =0.43) but lower than MT group (37.9±3.4 vs. 49.5±7.3, p=0.28).

Correlation

In the DAI group, there was significant correlation between GOS score and serum IL-8 at 7th day of admission (r=−0.68, p= 0.002). This correlation was not significant in the other days. In this group the GCS was found to be significantly correlated.
with the IL-8 concentration at 7th day of admission ($p = 0.026$, $r = -0.55$) but not in the other days. The IL-8 level had significant relation to WBC count at 7th day of admission ($p = 0.002$, $r = 0.71$), this relation was not significant in the other days. There was no significant correlation between platelet count and IL-8 level. Also the association of IL-8 and the age of patients were not statistically significant.

In the DAI group, there was no significant correlation between TGF β1 at 1st, 2nd, 3rd and 7th day and GOS and GCS. The TGF β1 did not have significant association with WBC and platelet count and the patients’ age. The correlation of NO with GOS and GCS was not statistically significant within this group.

The serum NO level did not have association with WBC and platelet count and the patients’ age.

**DISCUSSION**

There are many studies that show increased cytokine release in acute ischemic stroke, intracerebral hemorrhage and traumatic brain injury in the CNS and systemic flow (5,11,13).

IL-8 is a potent chemoattractant that could activate PMN and elongate its half-life in vivo (11). Polymorphonuclear leukocyte (PMN) infiltration has been shown to be associated with the disability after ischemic cerebrovascular attack (1). Therefore the IL-8 activate and elongate the effect of PMNs in the inflammation-induced brain disability process. Based on the results of the present study, the level of IL-8 has significant positive correlation with the WBC count at the 7th day of admission in the DAI patients. This finding may confirm the mentioned effect of IL-8 in the PMN activation and number. However, this correlation could be an accidental finding considering loss or the association of IL-8 and WBC count in the previous days and in the MT group. In a study on acute stroke patients the level of IL-6 and WBC count increased significantly. The level of TGF β1 and IL-6 did not have correlation with WBC count in that study (10). The IL-8 level of the MT patients was found to be significantly higher than DAI patients in the present study whereas the IL-8 level in the DAI group was not different with healthy group. These findings could be due to the source of IL-8 in the body that is mainly peripheral organs.

The level of IL-6 in the cerebrospinal fluid has been reported to be related to the lesion extension in the ischemic stroke (17). The serum concentration of IL-6 and IL-8 has been shown to be associated with disease severity and patients’ GCS in traumatic brain injury (7). There is no report of this association in DAI patients. We found that there is no correlation between the patients GCS and GOS with the inflammatory mediators (IL-8).

It has been shown that hypothermia induces anti-inflammatory cytokine profile. Hypothermia has various effects on inflammatory cytokines, depending on site, time of measurement, and the presence of infection (4,19). Previous studies have reported increased levels of IL-6 and IL-8 in cerebrospinal fluid of hypothermic pediatric traumatic brain injury and patients with cardiac arrest compared with normothermic patients (2,6). This was not confirmed in DAI patients in our study.

The IL-8 level had association with the GOS and GCS at 7th day of admission and higher IL-8 caused lower GOS score. In a study by Kostulas et. al. administration of antibody against IL-8 in cerebral perfusion injured rabbits was shown to protect the brain from polymorphonuclear infiltration and reduces the brain edema within 6 hours and infarction within 12 hours (11). Considering the results of the present study and the conclusion of the previous studies, IL-8 has important exacerbating effect on the brain lesion and its level could predict the outcome and clinical status of patients. TGF β1 could protect the brain from cytotoxics, hypoxia and acidosis so its level comes down in brain injuries as a result of its overuse. Significant difference of TGF β1 between DAI
and MT group in one hand and control healthy group in the other hand confirmed this theory. In a study by Slevin et al. patients with ischemic stroke had significantly lower level of TGF β1 in comparison to healthy group (16). TGF β1 had no significant association with the platelet and WBC count that is due to different type of cells releasing this factor.

NO begins to elevate in the ischemic stroke patients by IL-1 mediating rule which is released to CSF and blood soon after the attack (17). In the traumatic brain injury, the NO level of CSF and extracellular fluid has been found to elevate within the first day after injury and then decreases gradually within the next 5 days (8, 9). The results of the present study showed that NO level increased in the first day after injury and began to decrease until 7th day of admission. This finding was similar to the previous studies. NO concentration has been shown to be correlated with the GCS and GOS in some investigations that are in contrast to the results of the present study (8,9,18). This controversy may be due to different way of sampling. They measured the NO level in the CSF but we evaluated its serum level.

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REFERENCES