SOME BIOCHEMICAL CHANGES IN PATIENTS WITH ACUTE ISCHEMIC STROKE

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The aim was to determine the alterations in levels of vasculair endothelial growth factor (VEGF), homocysteine (HC) and lipid profile in patients with acute ischemic stroke (AIS), to establish the relationship between them. A clinical and neurological examination of 120 patients aged 56 to 75 years with AIS was done. Diagnosis was established taking into account results of spiral computer and magnetic resonance tomography. The control group included 20 people with no signs of acute cerebrovascular and severe somatic pathology aged 59.35±1.8 years. Assessment of disability was performed at admission and during the treatment using the scale NIHSS. In all relevant patients in 1-2 days after stroke, the concentration of HC and VEGF levels was determined by ELISA. Blood lipid profile was determined by enzymatic colorimetric method. The study found that increasing concentration of plasma HC is associated with intensity of hyperlipidemia, and hyperhomocysteinemia severity prevails over the changes in the lipid profile components. During AIS a significant increase in the level of VEGF in the blood was observed, indicating the presence of vascular accident and activation of angiogenesis. Severity of vascular endothelial lesions is associated with the degree of increasing concentration VEGF that appears most significant in patients with severe AIS.

Key words: acute ischemic stroke, vasculair endothelial growth factor (VEGF), homocysteine, blood lipid profile.

Annually in Ukraine 100-120 thousand of strokes are registered, and more than 30% – in people of working age. About 50% of patients die within the first year after a stroke, a third of those who survived, is severely disabled.

These statistics motivate scientists to more detailed study biochemical cascade of brain damage in order to identify new trends and strategies in the diagnosis and treatment of ischemic stroke.

The results of one of the largest study MONICA, which covered 21 countries, found that known risk factors (smoking, hypertension, overweight and atherosclerosis) can not fully explain the occurrence of cardiovascular and cerebrovascular diseases, since their prevalence reaches 15% in women and 40% in men [1-4]. In this regard, the search for new risk factors is necessary, identification of which would affect the level of mortality from cerebrovascular disease [5, 6]. Among the potential causes of vascular brain accidents, which are of great interest in experts, a special place belongs to hyperhomocysteinemia (HHC). Homocysteine (HC) – a sulfur natural aminoacid that is not found in proteins, and is the product of the degradation of methionine – one of the 8 essential aminoacids in the body [7, 8]. Among the factors affecting the level of HC and increase the likelihood of its content in the blood include vitamine deficiency, smoking, alcohol abuse, diet with high content of meat food and food with high cholesterol level. We know also that the HHC can be hereditary disease caused by deficiency of enzymes involved in the metabolism of HC [3, 5, 9, 10]. In the case of HC excess in the body it accumulates in the blood, and the main place of the harmful effects of this substance is the inner surface of blood vessels. There are evidence that HHC causes damage and activation of endothelial cells, which significantly increases the risk of thrombosis, causes “oxidative stress”, increases platelet aggregation. Activation of the coagulation cascade leads to disruption of endothelium dependent vasodilatation.

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and stimulate proliferation of smooth muscle cells. So HHC has a negative effect on the mechanisms regulating vascular tone, lipid metabolism and coagulation cascade, contributing to the development of various vascular diseases [1, 4, 6, 7]. Given the significant negative impact of HHC on the vascular wall and insufficient justification of its role in acute ischemic stroke (AIS), the object of our interest was the study of the prevalence HHC in healthy subjects and in patients with AIS and clarify the interaction of HHC with risk factors as hyperlipidemia. The most influential physiological inhibitor of programmed cell death among other are growth factors. They reduce the concentration of apoptotic effectors or their activity to a safe level and stimulate the process of angiogenesis, which is necessary for long-term adaptation of tissues in state of damage [11-13]. The subject of our interest was just vascular endothelial growth factor (VEGF), whose role remains uncertain for the formation of brain damage. Proved that VEGF – is angiogenic protein, which is released in response to hypoxia with marked damage microvasculature and is a major inducer of angiogenesis [14, 15]. In the blood of healthy individuals free VEGF content is low, it is usually expressed on platelets and appears in a free state in large numbers at their excessive activation or damage, however, increased plasma VEGF may reflect a process of rising activity of compensatory and protective mechanisms (angiogenesis) and increasing the permeability of blood-brain barrier due to hypoxia, reflecting the deepening of pathological changes [12, 14]. Given discrepancy of versions and research results, it is reasonable to study changes in VEGF concentration in acute cerebral ischemia and establish the presence of interrelation HHC and hypercholesterolemia.

The aim – to determine the concentration changes of vascular endothelial growth factor, homocysteine and lipide profile in patients with acute ischemic stroke and to establish the relationship between them.

Materials and Methods

A clinical - laboratory analysis of 120 patients with AIS aged 56 to 75 years. All patients were admitted in the first 24-48 hours from the onset of clinical manifestations of disease. The diagnosis is verified using spiral computer and magnetic resonance tomography. The control group consisted of 20 people with no signs of acute cerebrovascular accident and severe somatic pathology age (59.35 ± 1.80) years. Stroke severity was determined according to the scale NIHSS. (National Institutes of Health Stroke Scale, USA). The level of VEGF in the blood was determined using ELISA kits Biosource (USA) on 1-2 day illness. HC serum levels were determined using ELISA kits Axis Homocysteine EL, production Axis-Shield Diagnostics Ltd the Tehnology ParkDundee DD2, XA United Kingdom on 1-2-nd day after the onset of disease. Blood lipid spectrum was determined by using enzymatic colorimetric method for biochemical analyzer Screen master lab production Hospitex diagnostik (Germany) with a certain determination of total cholesterol level (TC), triglycerides (TG), high density lipoprotein cholesterol (HDL). The content of low-density lipoprotein cholesterol (LDL) was calculated using the equation W. Freedwald: LDL cholesterol = TC - (HDL cholesterol + TG/2.2); cholesterol lipoprotein very low density (VLDL): VLDL = TG/2.2. Atherogenic factor – the equation of Klimov A. N : AF = (HDL-C)/HDL.

Statistical analysis of the data was performed using universal statistical software “Excel 5.0” to a personal computer for the definition of the mean value. For authenticity difference between groups was used Student test (t). The difference between groups acknowledged significant at (P < 0.05).

Results and Discussion

HC level in control group was 8.45 ± 0.30 mmol/l. In all surveyed patients with AIS on 1-2 day of illness HC content corresponded 17.57 ± 1.1 mmol/l, respectively which was significantly higher compared to control (P < 0.001). In the study, we have observed widely oscillations in severity of HHC and dependence of HC level from the severity of the neurological deficit. (14.1%) patients with mild stroke level HC was 10.8 ± 0.12 mmol/l (P < 0.05); in patients with moderate stroke (45.8%) – 15.9 ± 0.32 mmol/l (P < 0.001). In patients with severe stroke (40%) had the most high levels HHC and grew more than two time compared the control group and were 19.5 ± 0.5 mmol/l (P < 0.001). In patients with severe AIS and premorbid background which included hypertension combined with cerebral atherosclerosis and diabetes with a history of multiple thrombotic episodes (transient ischemic attack, ischemic stroke, myocardial infarction) significantly higher values of serum HC were recorded which reached 25.3 ± 0.3 mmol/l. It should be noted that patients with AIS in vertebrobasilar pool, which arose as a result external vascu-
lar compression on the background of degenerative disc disease of the cervical spine, level HC was at the upper limit of normal, indicating lack of impact of HC on destructive changes in cervical spine. In all patients with AIS was found increased level of VEGF (378.5 ± 44.6) pg/ml (P < 0.001) compared to (105.5 ± 8.74) pg/ml in the control group, indicating the presence of “vascular accident” and activation of angiogenesis. VEGF, providing growth of new capillaries and arterial collaterals, the recovery of perfusion in the area penumbry, creating a positive effect and can be used as a method of therapy. Severity of vascular endothelial lesions was associated with the degree of expression of VEGF. This pattern is the most often revealing in the examined patients with severe AIS (453.5 ± 44.6) (P < 0.001), in mild cases increase in VEGF concentration in blood was lowest (190.8 ± 12.52) pg/ml, (P < 0.05). It should be noted that in older patients with severe stroke VEGF concentration was lower compared to younger patients, which probably indicates exhaustion of angiogenesis with age and less activity of reparative processes. In assessing the state of lipid metabolism in 1-2 day illness changes of all parameters of lipide profile were compared with the control group (P < 0.05). Levels of HDL, LDL, VLDL cholesterol and triglycerides were increased significantly compared with control (P < 0.05) and HDL cholesterol level decreased (P < 0.05), which explains the presence of atherosclerotic lesions of cerebral vessels (Table).

Attention is drawn to the fact that in patients with AIS hyperhomocysteinemia growth and VEGF are expressed to a greater extent compared to the severity of their hyperlipidemia. In patients with acute ischemic stroke a direct strong correlation connection between plasma levels of HC content and TC (r = 0.71; P < 0.01), LDL cholesterol (r = 0.72; P < 0.01), and direct medium strength connection between HC and TG (r = 0.31; P < 0.05), VLDL (r = 0.59; P < 0.01) and AF (r = 0.69; P < 0.01). Reliable reverse connection of average force is established between the level of plasma HDL and HC (r = -0.43; P < 0.05). This confirmed the impact of hyperhomocysteinemia on the activation process of atherogenesis and lowering blood levels of proatherogenic factors (Fig. 1).

Conducting correlation analysis between the lipid spectrum of serum and VEGF in patients with AIS we identified a direct strong correlation between the level of VEGF and content TC (r = 0.72; P < 0.01), LDL cholesterol (r = 0.71, P <0.01), and a direct medium strength relationship between VEGF and TG (r = 0.50; P <0.05), VLDL (r = 0.52; P < 0.05) and AF (r = 0.47; P < 0.05). Also we found reverse medium strength connection between the level of VEGF and HDL (r = -0.45; P < 0.05) (Fig. 2).

The presence of interrelation between VEGF and blood lipid spectrum in patients with AIS confirms the hypothesis of activation of protective mechanisms such as angiogenesis process in the atherosclerotic blood vessels, which contributes to the restoration of collaterals and perfusion in the area penumbry. After analysis of the correlation between the concentration of VEGF and plasma HC was found direct strong correlation (r = 0.79; P < 0.001). Thus, the process autooxidation of HC in plasma occurs with the formation of free radicals,

<table>
<thead>
<tr>
<th>Lipide profile, mM/l</th>
<th>Control group (M ± m, n = 20)</th>
<th>AIS patients (n = 120, P &lt; 0.05)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TC</td>
<td>4.62 ± 0.11</td>
<td>5.98 ± 0.21</td>
</tr>
<tr>
<td>HDL</td>
<td>1.53 ± 0.06</td>
<td>1.06 ± 0.08</td>
</tr>
<tr>
<td>LDL</td>
<td>2.53 ± 0.14</td>
<td>3.55 ± 0.33</td>
</tr>
<tr>
<td>VLDL</td>
<td>0.57 ± 0.04</td>
<td>0.95 ± 0.17</td>
</tr>
<tr>
<td>TG</td>
<td>1.36 ± 0.15</td>
<td>3.05 ± 0.15</td>
</tr>
</tbody>
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Fig. 1. Graphic representation of power correlation between plasma homocysteine and lipid spectrum of the blood in patients with acute ischemic stroke
Fig. 2. Graphic representation of power correlation between vascular endothelial growth factor level and lipid spectrum of the blood in patients with acute ischemic stroke

which have a toxic influence on the vascular wall endothelial cells, so the body reacts to this by a compensatory increase in the production of VEGF. Thus, in patients with AIS the increase in HC and VEGF in the blood can act as markers destruction of vascular endothelium, but they have a different genesis and oppositely acting on the vascular wall.

**DEЯKИЙ БIОХIМІЧНИЙ ЗMIНИ У ПАЦIЄНТІВ IЗ ГОСТРИМ IШЕМІЧНИМ IНСУЛЬТОМ**

O. Я. Михалойко, И. Я. Михалойко

Целью работы было определить концентрационные изменения васкулоэндотелиального фактора роста (ВЭРФ), гомоцистеина (ГЦ) и показателей липидограммы у пациентов с острым ишемическим инсультом (ОИИ), установить наличие взаимосвязи между ними. При проведении клинико-неврологического обследования пациентов (56–75 лет) с ОИИ диагноз верифицировали по данным спиральной компьютерной и магнитно-резонансной томографии; контрольная группа – пациенты (59,35 ± 1,8 лет) без признаков острых нарушений мозгового кровообращения и тяжелой соматической патологии. Оценку неврологического дефицита проводили за госпитализации пациентов в процессе лечения по шкале NIHSS. Всем тематическим
больным в 1–2-й день после возникновения инсульта методом иммуноэнзимного анализа были определены концентрации ГЦ и ВЕФР в крови. Исследование липидного спектра крови проводили энзиматическим колориметрическим методом. В результате исследования установлено, что увеличение концентрации ГЦ ассоциируется с проявлениями гиперлипидемии, причём степень выраженности гипергомоцистеинемии превалирует над характером изменений составляющих липидного профиля. При ОИИ отмечается достоверное повышение концентрации васкулоэндотелиального фактора роста в крови, что свидетельствует о наличии сосудистой катастрофы и активации процессов ангиогенеза. Выраженность поражения сосудистого эндоцеля ассоциируется с повышением концентрации васкулоэндотелиального фактора роста, что наиболее показательно проявляется у больных с тяжёлым ОИИ.

**Ключевые слова:** острый ишемический инсульт, васкулоэндотелиальный фактор роста, гомоцистеин, липидный спектр крови.

**References**


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