Serum Vascular Endothelial Growth Factor and Insulin-Like Growth Factor-1 in Liver Cirrhosis: Relation to Disease Severity and Development of Portal Hypertension

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ABSTRACT

Aims: To assess the level of Vascular Endothelial Growth factor (VEGF) and Insulin-like Growth Factor -1(IGF-1) in serum of patients with liver cirrhosis and correlate them to Child-Pugh classes and to correlate Vascular Endothelial Growth factor to Color Doppler indices of portal and splenic veins. Patients and Methods: fifty-five patients with liver cirrhosis and ten healthy controls were chosen. They underwent a thorough history, physical examination, abdominal ultrasonography and Color Doppler examination of portal vein and portal pressure. The serum levels of VEGF and IGF-1 were measured using commercial ELISA kits. Results: The median (and interquartile range) of serum VEGF was significantly lower in patients than controls (120ng/L [110-330] and 341ng/L [258-990] respectively, p value < 0.001), also there was a significant decrease in IGF-1 in patients than controls (34ng/ml [23-48.3] and 147.2ng/ml[125.8-220.2] respectively, p value < 0.000). There was a significant difference in median serum VEGF and IGF-1 levels among the different Child-Pugh classes (class A: 110ng/L [109-120], class B: 120.5ng/L [120-462], and class C 126.5ng/L [110-286], p value < 0.005 for VEGF and class A: 48.3ng/ml [42.8-49.4], class B: 23ng/ml [20.8-34], and class C 36.6ng/ml [32.3-49.7], p value < 0.000 for IGF-1). A significant positive correlation was noted between serum VEGF and maximum portal vein velocity and maximum splenic vein velocity (Spearman’s r = 0.780, r = 0.693 respectively, p value < 0.000). A significant negative correlation was noted between serum VEGF and the hepatic artery resistance index and splenic hilar diameter (Spearman’s r = -0.462, r = - 0.695 respectively, p value < 0.000). Significant positive correlation was found between IGF-1 serum levels and serum albumin (Spearman’s r = 0.310, p value = 0.012). No correlation was found between VEGF serum levels and serum albumin. Conclusion: Circulating VEGF level in patients with liver cirrhosis could not serve as an indicator of the progression of liver cirrhosis but rather it may reflect development of complication in the form of portal hypertension. Also, liver cirrhosis is associated with changes in serum IGF-1 that is related to the degree of liver dysfunction. Key words: vascular endothelial growth factor, insulin like growth factor-1, liver cirrhosis, portal hypertension.
INTRODUCTION

Liver cirrhosis constitutes a major health problem in Egypt. Portal hypertension is a complication of liver cirrhosis. Portal hypertension is associated with the development of a porto-systemic collateral circulation which decompresses the portal vascular system.

Vascular endothelial growth factor (VEGF), a 46kDa dimeric protein, is a direct and specific mitogen for endothelial cells. It induces migration, proliferation and survival of endothelial cells (1). At the liver level, VEGF is significantly expressed by sinusoidal endothelial cells and hepatocytes (2). VEGF has been extensively investigated recently in various hepatic diseases such as hepatitis, liver cirrhosis, primary and secondary hepatocellular carcinoma. VEGF serum concentration was found to increase in acute hepatitis and decrease in chronic hepatitis and liver cirrhosis (3), which suggests that VEGF levels might correlate with disease severity. Moreover, VEGF circulating level was found to be closely related to Child-Pugh classification in cirrhotic liver (4).

Portal hypertension is a complication of liver cirrhosis and is associated with the development of a porto-systemic collateral circulation. Until recently, it was thought that the development of collateral circulation was due to the passive opening of vascular channels in response to increased portal pressure. However, recent studies showed that the formation of collateral circulation may be, at least in part, due to angiogenesis driven by VEGF (5).

Insulin-like growth factor-I (IGF-I) is a polypeptide hormone secreted by multiple tissues in response to growth hormone (GH). It is partly responsible for GH activity, and also has glucose lowering and anabolizing effects (6).

Insulin-like growth factor-1 (IGF-1) also called somatostatin, low molecular peptide, is considered an important anabolic hormone inducing anabolic metabolism and stimulating DNA synthesis, cell proliferation and meiotic division in a variety of tissues (7). The liver is the main source of circulating IGF-1 and its synthesis is regulated by the growth hormone. Serum level of IGF-1 was found to decrease in liver cirrhosis (8).

The present study aims to assess the serum level of VEGF and IGF-1 in patients with liver cirrhosis and correlate them to Child-Pugh classes and to correlate VEGF to Color Doppler indices of portal vein and splenic vein.

PATIENTS & METHODS

Patients
A total of 70 patients with liver cirrhosis were recruited from the Inpatient Department of Tropical Medicine and Gastroenterology, Assiut University Hospital. All patients were diagnosed by clinical, biochemical and radiological findings as liver cirrhosis. The severity of liver disease was evaluated by the Child Pugh's score. Portal hypertension was clinically implied by the presence of ascites, splenomegaly, encephalopathy and/or history of bleeding varices that occurs in the presence of known chronic liver disease. Using the ultrasound and Doppler ultrasound portal hypertension was, also, suggested by the presence of hepatic architectural changes suggesting
chronic liver disease, splenomegaly (spleen hilar diameter >4.5 cm, longitudinal diameter >11 cm) \(^{(9,10)}\), portal vein diameter greater than 1.25 cm, portal vein flow velocity <21 cm/sec, and congestion index exceeded 0.1 \(^{(11)}\).

In addition to ten healthy controls were included in the study. The control group underwent the same clinical, biochemical, and radiological examination as patients group.

Abdominal ultrasonography/Doppler examination:

**Equipment**

The ultrasound equipment used for Doppler studies was Siemens Sonoline Sienna Ultrasound Imaging System, (Siemens, Germany), software version 1.5, 2.0 and 3 with frequency range (2.5-5MHz).

**B-mode Examination**

Spleen hilar and longitudinal diameter, portal vein diameter in mm, the portal vein cross-sectional area in cm\(^2\) and splenic vein diameter in mm were measured using B-mode.

**Doppler Examination**

All patients were studied after fasting overnight or for at least 6 hours and a resting period of 15 minutes in the supine position prior to the examination. The examination was carried by a single experienced examiner who was unaware of the grade of cirrhosis so as to prevent bias. The following parameters were recorded during the examination: 1) portal vein maximum velocity \(V_{\text{max}}\) in cm/sec, 2) hepatic artery resistance index (RI), and 3) Splenic vein maximum velocity \(V_{\text{max}}\) in cm/sec.

**Blood samples** were collected from overnight fasted patients and controls. Specimens were allowed to clot, the sera were separated and stored frozen at -70°C until assayed.

**Determination of serum level of VEGF:**

Concentration of serum VEGF was measured using ELISA kit that contains all components required for the quantitative measurements which was provided by Koma Biotech Inc.

**Determination of serum level of IGF-1:**

The concentration of IGF-1 in serum samples was done using ELISA technique that uses antibodies with high affinity and specificity for two different epitopes on IGF-1. The ELISA kit was provided by Biosource, Europe.

**Determination serum level of alanine transferase (ALT), aspartate transferase (AST) and albumin by using colorimetric method provided by Stanbio Laboratory and serum level of bilirubin by Quimica Clinical Aplicada.**

**Statistical analysis:**

Data was entered and analyzed using SPSS version 13 for Windows. Nonparametric statistics were used throughout the analysis. Quantitative variables were summarized as median (interquartile range). Nominal and ordinal variables were summarized as ratios. The significance of the difference between two groups was assessed using Mann-Whitney's U statistic and Kruskall-Wallis test was used to test the difference between more than two groups. Bivariate correlations were analyzed using Spearman's rank correlation coefficient.

**RESULTS**

Out of 70 patients who were eligible, we were unable to obtain
reliable Doppler indices in 8 patients. Seven patients were, also, excluded from the study after ultrasonographic examination (4 had hepatic focal lesions and 3 had portal vein thrombosis). That left 55 patients in the patient's group for analysis.

Baseline characteristics of the study population are presented in table (1). Thirteen patients were Child Pugh's class A, 22 were Child class B, and 20 were Child class C.

The median of VEGF in patients with liver cirrhosis was significantly less than controls (120ng/L (110-330) versus 34ng/L (258-990)) respectively, P value< 0.001). Insulin-like growth factor was, also, significantly less in patients than controls (34 ng/ml (23-48.3) versus 147.2ng/ml (125.8-220.2) respectively, P value < 0.000).

The serum levels of VEGF and IGF-1 in different Child-Pugh classes are presented in table (2). There was a significant difference in median serum VEGF levels among the different Child-Pugh classes (p value< 0.005). Vascular endothelial growth factor was positively correlated to Child-Pugh classes (Spearman's $r = 0.120$, p value= 0.381), and IGF-1 was negatively correlated to Child-Pugh classes (Spearman's $r = -0.030$, p value = 0.826).

A statistically significant positive correlation was found between VEGF and maximum portal vein velocity (figure 1) and maximum splenic vein velocity (Spearman's $r = 0.780$, $r = 0.693$ respectively, p value< 0.000). A significant negative correlation was found between VEGF and hepatic arterial resistance index and spleen hilar diameter (Spearman's $r = -0.462$, $r = -0.695$ respectively, p value< 0.000). No significant correlation was noted between VEGF and both portal vein diameter and splenic vein diameter (Spearman's $r = 0.049$, p value .696 and $r = -0.132$, p value 0.295 respectively). A significant positive correlation was found between IGF-1 and albumin (Spearman's $r = 0.310$, p value 0.012). However, no correlation was found between VEGF and serum albumin (Spearman's $r = 0.082$, p value= 0.554).

### Table (1): Baseline clinical, biochemical and radiological characteristics in patients and controls (median (interquartile range))

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Patients (n=55)</th>
<th>Control (n=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>51 (25-57)</td>
<td>43.5 (30.3-54)</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>31/24</td>
<td>6/4</td>
</tr>
<tr>
<td>Albumin (g/l)</td>
<td>22 (19-26.6)</td>
<td>43 (40.8-44.5)</td>
</tr>
<tr>
<td>Bilirubin (μmol/l)</td>
<td>37 (22-53)</td>
<td>4.8 (3.7-5.9)</td>
</tr>
<tr>
<td>AST (IU/L)</td>
<td>68 (47-85)</td>
<td>20 (14.5-27.8)</td>
</tr>
<tr>
<td>ALT (IU/L)</td>
<td>48 (33-69)</td>
<td>18.5 (14.8-25.8)</td>
</tr>
<tr>
<td>Portal vein diameter (mm)</td>
<td>13.1 (12.1-14.5)</td>
<td>9.9 (9.2-10.1)</td>
</tr>
<tr>
<td>Portal vein velocity (cm/sec)</td>
<td>18 (16-21)</td>
<td>26.5 (42.5-30.5)</td>
</tr>
<tr>
<td>Splenic hilar diameter (cm)</td>
<td>5.5 (4.9-6.6)</td>
<td>3.9 (3.5-4)</td>
</tr>
<tr>
<td>Splenic vein diameter (mm)</td>
<td>10 (8.9-11.9)</td>
<td>8.1 (7.9-8.5)</td>
</tr>
<tr>
<td>Splenic vein velocity (cm/sec)</td>
<td>21.4(18-27)</td>
<td>29(27-30.3)</td>
</tr>
</tbody>
</table>
Table (2): The median (interquartile range) serum levels of VEGF and IGF-1 in different Child-Pugh classes

<table>
<thead>
<tr>
<th>Child Classes</th>
<th>VEGF (ng/L)</th>
<th>IGF (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Child A (n=13)</td>
<td>110 (109-120)</td>
<td>48.3 (42.8-49.4)</td>
</tr>
<tr>
<td>Child B (n=22)</td>
<td>120 (120-462)</td>
<td>23 (20.8-34)</td>
</tr>
<tr>
<td>Child C (n=20)</td>
<td>126 (110-286)</td>
<td>36.6 (32.3-49.7)</td>
</tr>
<tr>
<td>p-value</td>
<td>0.005</td>
<td>0.000</td>
</tr>
</tbody>
</table>

![Graph showing correlation between VEGF levels and portal vein blood velocity in patients with liver cirrhosis.](image)

**Figure (1): Correlation between VEGF levels and portal vein blood velocity in patients with liver cirrhosis (r = 0.780) (p value< 0.000).**

**DISCUSSION**

The present study clearly demonstrates the reduction of VEGF in patients with cirrhosis than the control group and a positive correlation to Child-Pugh classes. There is, also, a significant relation between VEGF serum levels and three accurate radiological parameters of portal hypertension, portal vein velocity, hepatic artery resistance index and splenic hilar diameter.

The findings of VEGF changes with cirrhosis agree with others\(^4,12,13\). This could be attributed to increased activity of several inhibitor substances including angiostatin and endostatin in liver cirrhosis\(^13\). The low circulating VEGF levels in patients with liver cirrhosis, also, indicate that VEGF was derived neither from a large burden of tumor cells as hepatocellular
carcinoma was excluded from the study nor from platelet activation by portal vein thrombosis which was, also, excluded.

The increase in the median serum levels of VEGF with Child-Pugh stage was, also, reported by Assy et al.\(^\text{12}\) Such increase could be attributed to the positive regulators such as the acute phase proteins, tumor necrosis factor-α (TNF-α) and transforming growth factor-β (TGF-β), and the pro-inflammatory cytokines such as interleukin-1 (IL-1) and interleukin-6 (IL-6) which are unregulated in liver cirrhosis\(^\text{14}\). However, Fuyuhiko et al.,\(^\text{3}\) reported a decrease in VEGF serum levels with the progression in Child-Pugh classes which may suggest that serum VEGF may be associated with hepatocyte regeneration grade.

VEGF has been shown to induce the proliferation of hepatic sinusoidal cells by virtue of increased expression of its receptors flt-1 and flk-1 following partial hepatectomy.\(^\text{2, 15}\) Furthermore, Assy et al.\(^\text{12}\) noted that exogenous VEGF isoform 165 administration to partially hepatectomized rats stimulate liver cell proliferation. It is likely that hepatocyte proliferation requires growth factors such as hepatocyte growth factor (HGF) and transforming growth factor-α (TGF-α) in the early phase that directly stimulate hepatocytes randomly at the periportal areas but is rarely accompanied by reconstruction of the architecture of hepatic sinusoids, and growth factors that indirectly stimulate hepatocytes, such as VEGF, in the late phase where proliferative hepatocytes express VEGF to obtain sufficient blood flow by accompanying reconstruction of the architecture of hepatic sinusoids.

The relation to portal hypertension was, also, reported by others\(^\text{12,16}\). This confirms the contention that VEGF plays a role in the neovascularization process through angiogenesis and vasculogenesis in liver cirrhosis. Fernandez et al.\(^\text{5}\) demonstrated that the development of hyperdynamic circulation and formation of porto-systemic collateral vessels in portal hypertensive rats are angiogenesis-dependent process that could be markedly inhibited by blockage of VEGF signaling pathway.

Insulin like growth factor-1 (IGF-1) is a peptide hormone with metabolic and trophic actions particularly on skeletal muscle and bone\(^\text{17}\). The adult liver is considered to represent the major source of circulating IGF-1\(^\text{18,19}\) and decreased serum IGF-1 concentrations have been reported in patients with liver disease\(^\text{20}\), this agree with the present results that demonstrates a reduction of IGF-1 in patients with cirrhosis than the control group and a negative correlation to Child-Pugh classes. The decreased serum IGF-1 concentration is considered to be the result of reduced hepatic synthesis.

Scharf et al.\(^\text{21}\) reported that serum IGF-1 concentrations were correlated with serum albumin. This finding further indicates that the reduction in hepatocellular function provides an explanation for low IGF-1 concentrations in patients with cirrhosis.

In conclusion, circulating VEGF level in patients with liver cirrhosis could not serve as an indicator of the
progression of liver cirrhosis but rather, it may reflect development of complication in the form of portal hypertension. Also, liver cirrhosis is associated with decrease in serum IGF-1 that is related to the degree of liver dysfunction.

REFERENCES


عامل النمو للطبقة الطالبية المبطنة للرئوية الدموية وعامل النمو الشبيه للإنسولين-1 في مرضى تليف الكبد: وعلاقتهما بحالة المرض وحصيلة ارتفاع الضغط البائي

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بهدف هذا البحث الى قياس مستوى عامل النمو للطبقة الطالبية المبطنة للرئوية الدموية، وعامل النمو الشبيه للإنسولين-1 في مرضى تليف الكبد، ثم علاقتهما بتقسيم الـ Child-Pugh، وعلاقة عامل النمو للطبقة الطالبية المبطنة للرئوية الدموية بتصوير دوبلر الملون للوريد البائي والوريد الطحالب.

وبعد اجراء هذا البحث على 55 مريضاً يعانون من مرض تليف الكبد مع مقارناتهم بمجموعة ضابطة مكونة من 10 أشخاص اصحاء متوازنين لهم في السن والجنس وتم تشخيصهم جميعاً بواسطة التاريخ المرضي وفحص الطبي المتكامل، وكذلك عمل فحص بالجوهرات الفوق الصوتية على البطن ودوبلر الملون للوريد البائي وقياس ضغط الدم للوريد البائي، وتم قياس مستوى عامل النمو المبطن للرئوية الدموية وعامل النمو الشبيه للإنسولين-1 بطريقة الـ Child-Pugh.

أظهرت نتائج البحث أن هناك فجوة في دالة إحصائية لعامل النمو للطبقة الطالبية المبطنة للرئوية الدموية وعامل النمو الشبيه للإنسولين-1 عند مقارنتهم بالمجموعة الضابطة، كذلك وجدت اختلافات ذات دالة إحصائية في مستوى عامل النمو المبطن للرئوية الدموية وعامل النمو الشبيه للإنسولين-1 في مرضى تليف الكبد حسب تقسيم Child-Pugh للإنسولين-1 وحيد، وجد أن هناك علاقة ارتباط إيجابية بين عامل النمو المبطن للرئوية الدموية، وسرعته القصوى للوريد الطحالب وتيني أيضاً أن هناك علاقة ارتباط سلبي بين عامل النمو المبطن للرئوية الدموية، ومقاومة شريان الكبد وقطر الحلقات الفقارية. هناك علاقة ارتباط إيجابية بين عامل النمو الشبيه للإنسولين-1 ومستوى الأمين واتوجد هذه العلاقة مع عامل النمو المبطن للرئوية الدموية.
ومن ذلك نستنتج أن مستوى عامل النمو المتباطئ للأوعية الدموية في مرضى تليف الكبد لا
يمكن أن يكون مؤشراً لявление حالة تليف الكبد، ولكنه ربما يعكس حدوث مضاعفات في صورة
ارتفاع ضغط الدم البابي، ويمكن الاستنتاج أيضاً أن تليف الكبد قد يكون متأثراً بالتغيرات التي
تتطور لعامل النمو الشبيه للانسولين-1 التي لها علاقة بدرجة الخلل الوظيفي للكلد.