CARDIOVASCULAR EFFECTS OF ORAL ANTIVIRAL TREATMENT FOR NON-CARDIAC HCV PATIENTS

By


*Department of Cardiology, **Department of Tropical Medicine, ***Department of Clinical Pathology

Faculty of Medicine, AL-Azhar University (Cairo)

ABSTRACT

Background: The prevalence of hepatitis C virus (HCV) in Egypt is quite high, and the combined oral direct-acting antiviral agents (DAAs) may have impressive results.

Objectives: To assess the cardiovascular effects of oral antiviral treatment in non-cardiac HCV patients.

Patients and Methods: The study enrolled 100 patients with positive HCV who were enrolled for the new oral antiviral therapy in the form of Sofosbuvir and daclatasvir with or without ribavirin for 3 or 6 months. All patients on our study were subjected to follow up pre-and post-oral antiviral course, and the end point of the study was the development of a major cardiovascular event, e.g. congestive heart failure, echocardiographic evidence of left ventricular dysfunction, occurrence of significant arrhythmias, or acute coronary syndrome. The following parameters were accomplished: medical history and clinical examination, electrocardiogram, Echo-Doppler study, and laboratory investigations.

Results: No significant differences were found between the patients regarding demographic criteria. None of the patients had developed any major cardiac event. No significant changes were observed regarding ST segment and T wave abnormalities, arrhythmias, or QT interval. None of patients developed echocardiographic regional wall motion abnormalities at baseline or at study end. Systolic function parameters showed minute non-significant changes over study visits. Diastolic function parameters showed non-significant changes between baseline and post oral antiviral course visit.

Conclusion: The DAAs used in combination regimen didn’t significantly affect the cardiovascular system.

Key words: Hepatitis C virus, oral direct antiviral agents, echocardiography.

INTRODUCTION

Hepatitis C virus (HCV) is a significant public health problem and the leading cause of liver transplantation and hepatocellular carcinoma (Moyer, 2013). More recent surveillance data suggest that HCV infection has increased to 5.2 million people in the U.S. (Chak et al. 2015).

Sofosbuvir represents the first key step towards the new era in the management of chronic hepatitis C, since it is the first approved directly acting antiviral agents with excellent tolerability and favorable pharmacokinetic profile, limited potential...
for drug interactions, potent antiviral activity and high genetic barrier against all HCV genotypes. Sofosbuvir has recently become commercially available in combination with ribavirin, achieving high sustained virologic response (SVR) rates after 12-24 weeks of therapy (Keating, 2014 and Pawlotsky, 2014).

On the other hand, oral antiviral treatment for HCV patients is known to induce cardiac adverse effects, three cases of severe bradyarrhythmia that occurred during treatment with Sofosbuvir plus Daclatasvir, Simeprevir, or Ribavirin occurred among 415 patients treated (Fontaine et al. 2015). Recent reports had drawn the attention to the possible cardiotoxic effect of Sofosbuvir when combined with amiodarone. Occurrence of syncopal attacks was reported in two cases few days after administration of Sofosbuvir (Lashen and Sanhoury 2016).

Svere pulmonary hypertension was newly discovered or exacerbated after using sofosbuvir (Renard et al. 2016).

The present work aimed to assess the cardiovascular effects of oral antiviral treatment in non-cardiac HCV patients.

**PATIENTS AND METHODS**

**I- Patients:**

**Inclusion criteria:** This study was carried out on 100 Egyptian cases, who had chronic HCV infection. They attended at Health Insurance clinic in the Luxor governorate during the period from July 1st 2017 to February 28th 2018 they were candidated for treatment by oral antiviral for a period 3 or 6 months.

**Exclusion criteria:**

1- Decompensated liver cirrhosis (ascites, encephalopathy and bleeding varices).
2- Auto-immune hepatitis.
3- Chronic hepatitis B.
4- Combined chronic hepatitis B and C.
5- Patients with uncontrolled psychiatric disorders.
6- Cardiac disease (cardiomyopathy, arrhythmias, ischemia, myocarditis and valvular heart diseases).
7- Advanced renal impairment.
8- Uncontrolled thyroid dysfunction.
9- Poorly echogenic patients.

**II- Methods:**

All patients provided written informed consent. All patients included in the study were subjected to the following:

**I- Full history** taking which included name, age, sex, occupation, residence, shortness of breath, cough, hemoptysis, palpitation, chest pain, and paroxysmal nocturnal dyspnea.

**II- Clinical examination** including determination of blood pressure, pulse, jugular venous pressure, pallor, jaundice, cyanosis, ascites, edema of the lower limbs, S3, S4 and any detected murmurs.

**III- Laboratory investigations:**

a- Liver function tests: SGOT, SGPT, serum bilirubin, serum albumin, alkaline phosphatase and PT.

b- Serum Creatinine and serum urea.

c- Blood glucose (fasting and post-prandial).

d- Complete blood picture.
e- Viral markers specially for HBV, HCV (Screened by ELIZA technique).

f- TSH.

IV- Twelve lead ECG: Standard 12 lead ECG was recorded pre-and post-oral antiviral course to document the presence of significant ST changes suggestive of ischemic heart disease with assessment of rhythm, QT/ QTc, PR interval, P wave size, QRS duration and axis and voltage, abnormal Q, T and U waves.

V-Echo-Doppler study: Full 2-D, M-mode, Doppler and color flow mapping echocardiography study (by Phillips HD7 XE machine, 2-4probe) was done pre-and post-oral antiviral course.

2D and M-mode dimensions:
- Left ventricular end diastolic diameter (LVEDD).
- Left ventricular end systolic diameter (LVESD).
- Inter-ventricular septum thickness (IVS).
- Posterior wall thickness (PWT).
- Left atrial and aortic root diameters
- Right ventricle internal dimensions.

Left ventricular systolic function (Mitchell et al, 2018):
- Ejection fraction (EF)% by M-mode in parasternal short axis view at level of papillary muscle guided by 2D and long axis parasternal at level mitral valve leaflets tips or apical four chambers(A4C) by (modified Simpson's single plane method).

Left ventricular diastolic function assessed by (Mitchell et al, 2018):
- E/A ratio and tissue Doppler study (e-velocity &E/ e).
- Isovolumetric relaxation time (IVRT).
- Deceleration time (DT).

Right ventricular (RV) function (Rudski et al, 2010):
- RV function both systolic and diastolic by myocardial performance index (MPI).
- RV systolic function by TDI (systolic signal of the lateral tricuspid valve-TV-annulus) and tricuspid annular plane systolic excursion (TAPSE).
- RV pressure using tricuspid regurgitations (TR) jet and the RV outflow (RVOT) pattern i.e. parabolic or triangular.

Pulmonary hemodynamics:
- Pulmonary artery systolic pressure by modified Bernoulli equation from peak TR jet and mean right atrial pressure (Rudski et al, 2010).
- Pulmonary vascular resistance (PVR) estimation by the following equation:

\[
\{(TRV)/VTI_{RVOT}\} \times 10^{+0.16} (Abbas et al. 2013).
\]

(Where TRV, TR velocity. VTI_{RVOT}, right ventricular outflow tract velocity time integral).

Statistical analysis:
All results were obtained pre and post oral antiviral course and statistically analyzed using t-test. P value ≤ 0.05 were considered significant.
RESULTS

Baseline Characteristics (Table 1):

Patient’s baseline demographics are shown in (Table 1). The mean age of the recruited patients was 46.2 ±12.0 years. The most frequent age group is from 35-40 years presenting 23%, followed by the age group (30-35 years) presenting 18%. It is followed by the age group (45-50 years) presenting 13%. Then it is followed by the age groups (40-45 years) and (65-70 years), each presenting 12%. This is followed by age groups 50-55, 55-60 and 60-65 years representing 6%, 8% and 7% respectively. The least frequent age group is from 70-75 years representing only 1 %. The normal curve reveals that age distribution does not follow the normal distribution as the data are skewed to the left.

Forty-eight were males (48%) and 52 females (52%). No statistical significant difference was noted as regard patient’s gender (p-value = 0.16).

Table (1): Baseline demographics in the studied group

<table>
<thead>
<tr>
<th>Baseline characteristic</th>
<th>N=100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>46.2 ±12.0</td>
</tr>
<tr>
<td>Gender (Male)</td>
<td>48%</td>
</tr>
<tr>
<td>Diabetes</td>
<td>28%</td>
</tr>
<tr>
<td>Hypertension</td>
<td>43%</td>
</tr>
<tr>
<td>Smoking</td>
<td>38%</td>
</tr>
<tr>
<td>Family history of cardiac disease</td>
<td>23%</td>
</tr>
<tr>
<td>Treatment regimen</td>
<td></td>
</tr>
<tr>
<td>3 Drugs for 3 months</td>
<td>17%</td>
</tr>
<tr>
<td>3 Drugs for 6 months</td>
<td>2%</td>
</tr>
<tr>
<td>2 Drugs for 3 months</td>
<td>78%</td>
</tr>
<tr>
<td>2 Drugs for 3 months</td>
<td>3%</td>
</tr>
</tbody>
</table>

The most prevalent risk factors were hypertension (HTN) in 43 patients (43%), diabetes mellitus in 28 patients (28%), smoking in 38 patients (38%), and positive family history of cardiac disease in 23 patients (23%).

According to the treatment regimen, the patients enrolled in this study were distributed into 4 groups as the following (Figure 1):

- Group 1: 3 Drugs for 3 months (sofosbuvir, daclatasvir and ribavirin).
- Group 2: 3 Drugs for 6 months (sofosbuvir, daclatasvir and ribavirin).
- Group 3: 2 Drugs for 3 months (sofosbuvir and daclatasvir).
- Group 4: 2 Drugs for 6 months (sofosbuvir and daclatasvir).

No significance difference between regimen groups when compared by repeated measures ANOVA (p-value=0.43).
In the present study, baseline clinical examination was done as regard shortness of breath, cough, palpitation and chest pain and then reassessed by the end of treatment.

As regard blood pressure, systolic blood pressure showed statistically significant difference with p-value 0.00, however this difference was clinically insignificant (3 mmHg), while diastolic blood pressure showed no statistically significant difference after treatment when compared to their values before treatment (78.30±7.66 vs 78.90±9.31 mmHg respectively) (Table 2).

Table (2): Blood pressure before and after treatment

<table>
<thead>
<tr>
<th>Blood pressure</th>
<th>Before Treatment</th>
<th>After Treatment</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic(mmHg)</td>
<td>125 ±14</td>
<td>122±12.3</td>
<td>0.0</td>
</tr>
<tr>
<td>Diastolic(mmHg)</td>
<td>78.9±9.3</td>
<td>78.3±7.6</td>
<td>0.29</td>
</tr>
</tbody>
</table>
Heart rate showed no statistically significant difference after treatment when compared to their values before treatment (83.43±11.36 vs 83.60±12.06 beat/minute respectively). The PR interval showed no statistically significant difference after treatment when compared to their values before treatment (143.85±16.83 vs 142.23±17.67 msec respectively). The QRS duration showed no statistically significant difference after treatment (96.86±8.13 msec) when compared to their values before treatment (97.09±7.80 m/s). The QT interval showed no statistically significant difference after treatment when compared to their values before treatment (357.97±26.35 vs 358.67±26.42 msec respectively) and corrected QT interval also showed no statistically significant difference after treatment (422.10±18.81) when compared to their values before treatment (421.82±27.23). No significant alterations regarding ST segment and T wave or arrhythmias between the start and the end of treatment course (Table 3).

Table (3): ECG parameters before and after treatment

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Time</th>
<th>Before Treatment</th>
<th>After Treatment</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate( b/m)</td>
<td></td>
<td>83.60 ±12.06</td>
<td>83.43 ±11.36</td>
<td>0.59</td>
</tr>
<tr>
<td>P-R interval (ms)</td>
<td></td>
<td>142.23 ±17.67</td>
<td>143.85 ±16.83</td>
<td>0.44</td>
</tr>
<tr>
<td>QRS duration (ms)</td>
<td></td>
<td>97.09 ±7.8</td>
<td>96.86 ±8.13</td>
<td>0.79</td>
</tr>
<tr>
<td>QT (ms)</td>
<td></td>
<td>358.67 ±26.42</td>
<td>357.97 ±26.35</td>
<td>0.86</td>
</tr>
<tr>
<td>QTC</td>
<td></td>
<td>421.82 ±27.23</td>
<td>422.10 ±18.81</td>
<td>0.38</td>
</tr>
</tbody>
</table>

There was no statistically significant difference between any of echocardiography parameters before and after treatment. Only three measurements showed statistically significant difference; LVESD mm (by 2D), IVSd mm (by 2D) and EF by 2D% with p-values less than 0.05. However, these differences were not clinically significant (Table 4).
Table (4): Echo-Doppler parameters before and after treatment

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Time</th>
<th>Before Treatment</th>
<th>After Treatment</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>LVEDD mm(by MM)</td>
<td></td>
<td>44.92 ±4.26</td>
<td>45.82 ±4.68</td>
<td>0.09</td>
</tr>
<tr>
<td>LVESD mm(by MM)</td>
<td></td>
<td>29.18 ±3.08</td>
<td>29.60 ±3.41</td>
<td>0.16</td>
</tr>
<tr>
<td>IVSd mm(by MM)</td>
<td></td>
<td>9.48 ±0.71</td>
<td>9.44 ±0.76</td>
<td>0.64</td>
</tr>
<tr>
<td>PWd mm(by MM)</td>
<td></td>
<td>9.63 ±0.92</td>
<td>9.54 ±0.94</td>
<td>0.31</td>
</tr>
<tr>
<td>LA mm(by MM)</td>
<td></td>
<td>33.47 ±3.81</td>
<td>33.73 ±3.22</td>
<td>0.46</td>
</tr>
<tr>
<td>AO mm(by MM)</td>
<td></td>
<td>23.72 ±3.56</td>
<td>23.69 ±3.5</td>
<td>0.96</td>
</tr>
<tr>
<td>RVd mm(by MM)</td>
<td></td>
<td>19.00 ±2.44</td>
<td>19.37 ±2.36</td>
<td>0.13</td>
</tr>
<tr>
<td>LVEDD mm(by 2D)</td>
<td></td>
<td>42.29 ±5.1</td>
<td>45.82±4.68</td>
<td>0.09</td>
</tr>
<tr>
<td>LVESD mm(by 2D)</td>
<td></td>
<td>27.53±3.58</td>
<td>29.60±3.41</td>
<td>0.0*</td>
</tr>
<tr>
<td>IVSd mm(by 2D)</td>
<td></td>
<td>9.10±0.44</td>
<td>9.35±0.67</td>
<td>0.0*</td>
</tr>
<tr>
<td>PWd mm(by 2D)</td>
<td></td>
<td>9.39±0.88</td>
<td>9.44±0.79</td>
<td>0.38</td>
</tr>
<tr>
<td>LVEDV cm3(by 2D)</td>
<td></td>
<td>103.73±23.5</td>
<td>102.42±25.1</td>
<td>0.36</td>
</tr>
<tr>
<td>LVESV cm3(by 2D)</td>
<td></td>
<td>40.82±11</td>
<td>41.10±12.43</td>
<td>0.92</td>
</tr>
<tr>
<td>EF by MM %</td>
<td></td>
<td>63.23±0.34</td>
<td>62.73±3.17</td>
<td>0.35</td>
</tr>
<tr>
<td>EF by 2D%</td>
<td></td>
<td>62.87±3.83</td>
<td>64.25±3.2</td>
<td>0.01*</td>
</tr>
<tr>
<td>EF by Simpson%</td>
<td></td>
<td>63.61±3.81</td>
<td>63.59±3.51</td>
<td>0.72</td>
</tr>
<tr>
<td>E/A</td>
<td></td>
<td>2.21±8.56</td>
<td>1.29±1.0</td>
<td>0.15</td>
</tr>
<tr>
<td>e-(cm/s)</td>
<td></td>
<td>10.36±2.67</td>
<td>10.31±3.12</td>
<td>0.61</td>
</tr>
<tr>
<td>E/e-</td>
<td></td>
<td>8.25±2.31</td>
<td>8.28±2.07</td>
<td>0.78</td>
</tr>
<tr>
<td>IVRT(ms)</td>
<td></td>
<td>86.86±15</td>
<td>87.59±17</td>
<td>0.79</td>
</tr>
<tr>
<td>DT(ms)</td>
<td></td>
<td>161.62±43.57</td>
<td>159.77±39</td>
<td>0.65</td>
</tr>
<tr>
<td>RV MPI</td>
<td></td>
<td>.16±0.1</td>
<td>.15±0.09</td>
<td>0.42</td>
</tr>
<tr>
<td>RV S-(mm)</td>
<td></td>
<td>14.64±2.4</td>
<td>14.86±2.52</td>
<td>0.47</td>
</tr>
<tr>
<td>TAPSE(mm)</td>
<td></td>
<td>24.33±4.5</td>
<td>24.23±3.77</td>
<td>1.00</td>
</tr>
<tr>
<td>PASP(mmHg)</td>
<td></td>
<td>21.48±4.58</td>
<td>21.50±4.87</td>
<td>0.89</td>
</tr>
<tr>
<td>PVR(WU)</td>
<td></td>
<td>1.53±0.33</td>
<td>1.47±0.36</td>
<td>0.17</td>
</tr>
</tbody>
</table>

*Statistically significant difference.

2D=2-dimensional echocardiography, AO=aortic root, EF=ejection fraction, IVRT= isovolumic relaxation time. DT=deceleration time, IVSd= inter-ventricular septum in diastole, LA =left atrium, LVEDD =left ventricular end diastolic diameter, LVEDV=left ventricle end diastolic volume, LVESD= left ventricular end systolic diameter, LVESV=left ventricle systolic volume, Mm=millimeters, MM= M-mode, mmHg= millimeter of mercury, PASP=pulmonary artery systolic pressure, PVR=pulmonary vascular resistance, PWd =posterior wall in diastole, RV MPI=right ventricle myocardial performance index, TAPSE=tricuspid annular plane systolic excursion, RVd=right ventricle in diastole, WU=wood units.
DISCUSSION

Regarding the new DAAs, Food and Drug Administration (FDA) recently announced a change in labeling for the hepatitis C antiviral LDV/SOF (Harvoni) and SOF (Sovaldi) after the manufacturers reported bradycardia, pacemaker invention, and even death in patients who took the medications along with amiodarone (FDA hepatitis update, 2015).

As the prevalence of HCV in Egypt is high, and the new treatment combined with therapy involving oral DAAs either with or without PEG-IFN is extensively used.

The present study was designed to investigate the cardiovascular effect of DAAs on non-cardiac HCV patients. It included (100) Egyptian cases with HCV infection, who attended Luxor Insurance clinic for treatment by DAAs.

All included subjects were submitted to full history taking, clinical examination, laboratory investigations, twelve lead resting ECG, transthoracic Echo-Doppler study pre & post antiviral course.

Several studies from different parts of the world have reported that HCV infection may also contribute to the development of diabetes mellitus (DM), and higher prevalence of type 2 DM has been observed in patients with HCV infection than in those with other forms of chronic hepatitis (Bernsmeier & Heim 2009, and Lonardo et al. 2009).

(Rouabhia et al., 2010) reported that the prevalence of diabetes in HCV Algerian patients was reported to be 39.1%.

In the present work, the laboratory tests showed the fitness of the studied patients to receive DAAs therapy; unfortunately, no values at the end of the course of treatment were obtained. In the literature, it was reported that, DAAs resulted in suppression of baseline levels of hemoglobin, hematocrit, white blood cells, and pancytopenia.

In the present study, baseline clinical examination was done as regard determination of blood pressure, pulse, jugular venous pressure, pallor, jaundice, cyanosis, ascites, edema of the lower limbs, S3, S4 and any detected murmurs &shortness of breath, cough, hemoptysis, palpitation and chest pain, paroxysmal nocturnal dyspnea and then reassessed by the end of treatment. No statistically significant difference was found between after treatment and baseline findings before treatment.

Wong. (2009) reported that, several electrophysiological abnormalities have been observed in cirrhosis and these include of chronotropic incompetence, electromechanical uncoupling, and prolonged QT interval.

These results are quite different that those reported in the present work, as it was reported with liver cirrhosis but in the present study the included patients are in early stages of hepatitis C infection with no evidence of cirrhosis. In addition, there was no data in the literature as regards the effect of DAAs QT and QTc.

Fontaine et al. (2015) reported three cases of severe bradyarrhythmia that occurred during treatment with Sofosbuvir plus Daclatasvir, Simeprevir, or ribavirin.
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among 415 patients treated in their unit from January 2 to December 31, 2014.

But in the present study, there was no statistically significant difference was in rhythm found between after treatment and baseline findings before treatment.

Ahmad et al. (2015) Reported in their study on Thirty-four patients received Interferone-free BMS (Bristol-Myers Squibb) regimens (Sofosbuvir included regimen); Six patients had left ventricular ejection fractions (LVEFs) <30%, 8 had LVEFs 30%-50%, and 11 required hospitalization for suspected cardiotoxicity. Of the patients with LVEF <50%, 6 had normalization of systolic function after a median of 20 days. T-wave inversions were the most sensitive predictor of LVEF dysfunction. These results are quite different that those reported by Biomy et al. (2015) on their Study findings which showed no significant alterations in patient symptoms (shortness of breath, palpitations, and chest pain), signs (heart rate and blood pressure), or ECG recordings (arrhythmias, QT interval, or ST-T wave changes) before and after the oral antiviral treatment. In addition, there were no significant alterations in echocardiographic parameters regarding the systolic and diastolic functions before and after the oral antiviral treatment.

Renard et al. (2016) reported three cases of newly diagnosed or exacerbated severe pulmonary arterial hypertension (PAH) in patients treated with Sofosbuvir.

Our study results indicated that DAAs therapy did not induce any significant change in pulmonary artery pressure. This could be due to that PAH have been related to multiple associated factors besides DAA treatment.

CONCLUSION

The present findings suggested that this therapy seemed that it has no cardiovascular effects.

This study has some limitations:
- The small sample size of included cases.
- Incomplete follow up of DAAs therapy course.
- Follow up for the possible electrophysiological abnormalities were not used, e.g. Holter monitoring.
- Myocardial perfusion imaging for the detection of possible ischemia were not used.
REFERENCES


الآثار القلبية الوعائية من العلاج عن طريق الفم المضاد للفيروسات في أصحاب القلب مرضي التهاب الكبد الفيروسي سي

سامح رفعت حسن علام* - جمال محمد سليمان** - أشرف عبد المنعم سيد***
أحمد عبد الروؤف مهدي* - محمود محمد أبو الحجاج*

قسم القلب والأوعية الدموية **الأمراض المزمنة ***الباثولوجيا الإكلينيكية
كلية الطب جامعة الأزهر

خلفية البحث: إنشار فيروس الالتهاب الكبدى الوبائى (سي) في مصر مرتفع، وقد تكون نتائج العلاج عن طريق الفم المضاد للفيروسات مثيرة للاعجاب.

الهدف من البحث: تقييم الآثار القلبية الوعائية للعلاج عن طريق الفم في أصحاب القلب المرضي بالتاريخ الوبائى سي.

المريض وطرق البحث: التحق بالدراسة (100) مريض مصاب بفيروس التهاب الكبد الوبائي سي الذين تم تسجيلهم في العلاج المضاد للفيروسات عن طريق الفم على شكل سوفوسوفر و داكاتاسفير مع أو بدون الربافيرين لمدة 3 أو 6 أشهر. وقد تم أخذ التاريخ المرضي بصورة مفصلة، كما تم عمل الفحوصات العملية الضرورية لمثل هذه الحالات علاوة على عمل فحص إكلينيكى شامل قبل وبعد تناول العلاج كذلك تم عمل رسم قلب وفحص بالموجات فوق الصوتية على القلب قبل وبعد العلاج.

النتائج: لم تكن هناك تغييرات في العلامات الحيوية ولا في الفحص الإكلينيكي للمرضي بعد العلاج عن مثيلها قبل العلاج كما أن التغيرات في تخطيط القلب الكهربائي وفي الموجات فوق الصوتية على القلب كانت غير دالة إحصائيا بعد العلاج مقارنة بثانيتها قبل العلاج.

الاستنتاج: نتائج الدراسة الحالية تشير إلى أن العلاج عن طريق الفم المضاد للفيروس (سي) لا يوجد لديه تأثيرات على القلب والأوعية الدموية.