Anthropometric and Biochemical Characteristics of Patients with Nonalcoholic Fatty Liver Diagnosed by Non-Invasive Diagnostic Methods

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ABSTRACT

Introduction: Non-alcoholic (NAFLD) encompasses a spectrum of disease states, from steatosis (fatty liver) to non-alcoholic steatohepatitis (also called NASH steatosis with inflammatory changes) followed by progression to fibrosis and cirrhosis and hepatocellular carcinoma. Excess liver fat is believed to be a manifestation of the metabolic syndrome and not surprisingly, NASH is associated with obesity, insulin resistance, dyslipidemia and type 2 diabetes in humans. Aim of the study: is to establish anthropometric and biochemical specificities in patients with non-alcoholic steatohepatitis diagnosed with non-invasive diagnostic methods. Material and methods: Study enrolled 170 participants, 130 with NASH steatosis. The non-alcoholic group (control), consisted of 40 normal weight patients without metabolic syndrome. Alcohol intake was estimated with established protocol. Routine biochemistry analysis were performed by standard laboratory procedures; serum levels of serum levels of fasting cholesterol and triglycerides, fasting glucose and insulin, insulin resistance estimated by HOMA index (Homeostasis model assessment), biochemistry tests and a liver ultrasound examination. Results: In study participants group, patients were more obese comparing with controls p < 0.01, waist line extent also was of greater statistical significance in the non-alcoholic group fatty liver (p < 0.01). Comparing biochemical parameter values, significant statistical deference has been noted in glaucosis and insulin levels, total cholesterol and gama-glutamil transferase levels, between groups (p<0.01). Fasting glucose and insulin levels, HOMA-IR were significantly greater in study cohort group patients, as was significantly positive correlation between BMI and waist line extent. Conclusion: Patients with non-alcoholic fatty liver are excessively obese, have greater waist line extent, consequently insulin resistance and impaired glucose metabolism, insulin resistance, dyslipidemia, risk factors known to be associated with the development of cardiovascular disease.

Kay words: non-alcoholic fatty liver, obesity, body mass index, waist line extent, insulin resistance.
(5, 6). Being almost always present in NAFLD insulin resistance is considered hepatic manifestation of the metabolic syndrome (7, 8).

Investigating pathophysiology background of the metabolic syndrome, insulin resistance and visceral obesity are recognized as main factors of risk. It could be pointed out that insulin resistance is core and obesity most frequent clinical manifestation (9). Early recognition of the metabolic syndrome is of great importance for atherogenesis which starts on the very beginning of the process, jointly with other multiple factors proceeds towards diabetes mellitus (DM) type 2, arteriosclerotic complications (10). Patients with non alcoholic fatty liver disease, adults and children usually meet diagnostic criteria for metabolic syndrome (abdominal fat deposition, hypertension, atherogenic dyslipidemia and hyperglycemia) respectively have factors of risk for cardiovascular disease development (5, 7, 9).

In estimating cardiovascular risk, fat distribution, is of greater importance, than amount of body fat and adipocyte dysfunction. Visceral fatty tissue correlates with insulin resistance, which contrary to subcutaneous fatty tissue is metabolic active organ and exhibits numerous local and systemic effects through secretion of whole lot of variety peptides, some of them with altered function in obesity. Enlarged and inflamed visceral fatty tissue releases wide spectrum of the molecules potentially involved in development of the insulin resistance and atherosclerosis, including free fatty acids, interleukin 6, tumor necrosis factor α (TNF-α), monocytes chemotactic protein 1 and other proinflammatory cytokines (7). Therefore recently modified definitions of the metabolic syndrome include waist line measurements as insulin resistance marker (11). Zhu and associates pointed out that waist line better points out towards NAFLD (2). Body mass measurements (BMI index), waist line measurements (VC) waist-hip ratio (VHR), bio electrical impedance analysis (BIA) are widely recommended in patients with NFLD as diagnostic tools, for being comfortable and relatively at low cost.

A group of authors from our surrounding area, also support the attitude that waist line extent is a good indicator of the visceral obesity, that can independently point out health risk (12). Those values that indicate sudden rise of the risk factors for cardiovascular diseases for men are 96 cm, and for women 86 cm in waist line (13).

Notification of either of the disorders in patients with established disrupted AST/ALT (ratio- aspartate aminotransferase/alanine aminotransferase), ratio, obesity, especially morbid obesity (BMI>35), DM type 2 diagnosis, diagnosis of the metabolic syndrome, obstructive apnea, point out towards NAFLD (2). Body mass measurements (BMI index), waist line measurements (VC) waist-hip ratio (VHR), bio electrical impedance analysis (BIA) are widely recommended in patients with NFLD as diagnostic tools, for being comfortable and relatively at low cost (2). In raised suspicion for NAFLD ultrasonographic imaging is diagnostic method first to be used in liver steatosis detection, in order to avoid more invasive methods as liver biopsy (1, 2). Existing dogma posits that liver biopsy is the most reliable approach for identifying the presence of steato- hepatitis and fibrosis in patients with NAFLD, but it is generally acknowledged that biopsy is limited by cost, sampling error, and procedure-related morbidity and mortality. So, liver imaging is the common test used in liver steatosis screening (2). Imaging reveals lever of the homogenic structure, light with increased echogenicity (glittering liver) (15).

Aim of this study was to establish anthropometric and biochemical specificities for patients with non alcohol fatty liver, diagnosed by noninvasive diagnostic procedures.

2. MATERIAL AND METHODS

Prospective study conducted during 24 months (2009-2011) covered and analyzed 170 participants through clinical monitoring. Diagnosis of "non alcoholic fatty liver" confirmed by noninvasive imagining liver ultrasound procedure and by the AST/ALT ratio (1, 2).

Respondents were distributed in two groups:

- Study group (consisted of responders with imaging confirmed fatty liver and AST/ALT pathological values. Group numbered 130 participants.
- Controls (consisted from responders of the same age, slim with imagining that match normal liver findings, 40 of them.

Patients have been monitored in internists ambulances of the Clinical-Hospital Center of Prishtina, and through Health Centre of Gracanica, with already estimated protocols that included patient’s name, generalities and valid anamnestic data. By those protocols anthropometric measurements have been carried out in order to estimate the degree of the nutritional status, mass, and the fatty tissue distribution.

Building up databases and processing was done in SPSS program version 10.0. Also standard descriptive methods were performed, as were analytical methods, t-test for related samples, t-test for independent samples and χ2 test. Correlation between biochemical and anthropometric characteristics tested with Pearson’s correlation differences of p<0.05 taken as statistically significant.

3. RESULTS

Basic data on the respondents are listed on table 1. There were no big difference among respondents considering age, the average age of the study group was 56, 26 ± 4, 2 years and 56, 40 ± 4, 22 years in control.

In group with nonalcoholic fatty liver there were 78% of man or 60%, and 52 women or 44%, while controls consisted of 24 men or 60% and 16 women or 40%. Between study group and control group no statistical important difference between gender representation could be noted.

Table 1. shows the anthropometric characteristics of the subjects with NAFLD and control group also clinical and biochemical characteristics of the tested groups. Respondents of the study group were more obese at the level of significance compared to the controls, p < 0.01 (BMI (kg/m2)) (32.22±5.50 vs 24.67±3.83), waist line circumdactio also showed statistically significant higher values in the group with NAFLD (p < 0.01). Waist/hip ratio was higher in the group with NAFLD, but there were no statistical significance among the groups (p =0.502).

Comparing biochemical parameters between groups significant statistical importance could be established among the glucose and insulin values, total cholesterol values and glutamyltransfere (γ-GT) (p<0, 01) values. No statistically significant difference between AST i ALT...
was noted. Anthropometric characteristics, and their frequency among the tested groups are shown on the Table 2. and 58.46 % tested subjects of the group with NAFLD had BMI ≥35, what according to the WHO guidelines fall in obesity of second degree. Waist circumference > 88 cm had 88.64 % of the responders. These was followed by increased fat deposition, fat body value were detected in 32 % responders, between 20 and 25, while values >25, had 64.02 % of them. Responders with NAFLD had a significantly higher incidence of disorders compared with those in controls.

Comparing participants to the input of alcohol abuse and the factor of risk for the cardiovascular disease presence, we came out with results pointing on study group, which had significantly higher incidence of hypertension, diabetes, and hypercholesteremia (p<0.001), comparing with controls, also significantly higher incidence for obesity and diabetes was noted among close relatives (Table 3).

Fasting glucose and fasting insulin values, insulin resistance (HOMA-IR) were significantly higher in responders with NAFLD, also significantly positive correlation with BMI and waist circumference (Tables 4 and 5) established. Correlation between WHR relationship and insulin resistance was not confirmed (r =-0.08782; p=0.3167).

### 4. DISCUSSION

There is all the more evidence that NAFLD presents hepatic component of the "metabolic syndrome", defined with obesity, hypercholesteremia, peripheral insulin resistance, diabetes mellitus, hypertriglyceridemia and hypertension (5,6,26,27). There are not that many studies that dwelt with anthropometric measurements in subjects with non-alcoholic fatty liver so assessed data are conflicting. Obesity is well documented factor of risk for NAFLD. Excessive obesity defined as BMI ≥25 and visceral obesity, waist circumference of ≥88 cm in women and ≥102 cm in men have been recognized as factors of risk for NAFLD (21). Our results confirmed that responders in study group were more obese (BMI 32.22±5.50kg/m² vs 24.67±3.83± 3, 51 kg/m²) than responders in controls which were, statistically highly significant (p <0, 001). In group of steatosis also were assessed statistically higher values for waist circumference 122.6±11.7 cm vs 86.0±5.8 cm (p<0, 001). Comparing data from available literature, studies from the region confirm facts that subjects with liver steatosis are significantly more obese than responders without steatosis (26, 27). The impact of the excessive body mass on liver disorders appearance confirmed studies that based diagnosis on liver biopsy. Ratziu V and associates confirmed in their studies, that obesity is the factor of risk in development of severe fibrosis (28). Cheung O and a group of authors based on collected data, that the degree of abdominal obesity correlates with necroinflammatory liver activity (29). Uslusoy HS and associates on the contrary did not establish statistically significant correlation between body mass index, waist circumference and severe necroinflammatory inflammation, respectively fibrosis (30).

Obesity is an independent risk factor for DMT2 de-
Accumulation of triglyceride in hepatocytes is considered the main pathogenic trigger in the development of NAFLD (44). Cholesterol metabolism in patients with NAFLD (45) is different from healthy individuals. Lipids, including triglycerides, are accumulated in the liver. This is often associated with insulin resistance and metabolic syndrome, diabetes and NAFLD (39). Also studies with conducted type 2, from 40 up to 200 % comparing to those without NAFLD. New American guidelines, unfortunately does not support non invasive biomarkers for identification NAFLD. New American guidelines, unfortunately does not support non invasive biomarkers for identification NAFLD (30). In the preceding study biochemical analysis of the 116 patients with NAFLD biopsy verified diagnosis, were compared with the results of the healthy subjects. NAFLD patients had higher levels of insulin, glycemia and HOMA-IR values, compared to controls, even when they did not have diabetes or glucose intolerance.

In addition to related disorders of glucose metabolism and insulin resistance, we found a large frequency of the hypertension in patients with NAFLD. Although amino-transferase activities (a surrogate marker for NAFLD) increase long term risk for diabetes, metabolic syndrome and cardiovascular incidents (36). Degree of enzyme is not determined, it is usually between 1 and 4 times over upper limit. Although amino- transferase (ALT) values are higher than aspartate aminotransferase (AST) levels, in most cases (37) the AST levels temporarily could be higher than levels of AST, especially in cirrhosis (38). All the same, the fact is that AST / ALT ratio never exceeds 2.

Kotronen and associates in their research pointed out that AST and ALT values were higher in patients with DM type 2, from 40 up to 200 % comparing to those without diabetes and fatty liver (39). Also studies with conducted by Duvnjak et al. Most of the patients had ALT levels higher than the requirements for inclusion in this study was disrupted AST/ALT ratio. Data from our study implicate higher γ-GT values, statistically high significant in patients with NAFLD (35, 36, 37, 38, 39). Investigating the importance of the liver enzymes disorders Goessling and associ. state that in general population high aminotransferase (a surrogate marker for NAFLD) increase long term risk for diabetes, metabolic syndrome and cardiovascular incidents (36). Although amino-transferase activity of liver enzymes and their impact for the diagnosis of nonalcoholic fatty liver, one of those is study conducted by Noguchi and ass (34), while several cross sectional studies found firm connection with elevated ALT, γ-GT values, and metabolic syndrome, diabetes and NAFLD (35, 36, 37, 38, 39).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycemia (mmol/L)</td>
<td>r = -0.2265; p &lt; 0.01</td>
</tr>
<tr>
<td>Triglycerides (mmol/L)</td>
<td>r = -0.1085; p = 0.2154</td>
</tr>
<tr>
<td>Total cholesterol (mmol/L)</td>
<td>r = -0.04339; p = 0.6213</td>
</tr>
<tr>
<td>γ GT(U/L)</td>
<td>r = 0.1567; p &lt; 0.05</td>
</tr>
<tr>
<td>Insulinemia (mIU/L)</td>
<td>r = 0.2981; p &lt; 0.01</td>
</tr>
<tr>
<td>HOMA-IR</td>
<td>r = 0.1502; p = 0.0858</td>
</tr>
</tbody>
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Table 5. Correlation between waist circumference, glycemia values, triglicerides, total cholesterol values, insulinemia and HOMA-IR index. Pearson’s correlation coefficient p-level of the correlation significance.

development. Risk of developing the disease increases exponentially when body mass index crosses 25 kg/m² (1, 2, 3). Our data indicate significant correlation between cardiovascular risk factors and risks for the development of diabetes with almost all anthropometric experimental procedures. Fasting glucose and fasting insulin values, insulin resistency (HOMA-IR) were significantly higher in responders with NAFLD, as was significantly positive correlation with BMI, waist circumference with other risk factors for the cardiovascular diseases development. The correlation between WHR relations and insulin resistency (r= 0.1502; p = 0.0858) have not been confirmed.

HOMA index is a simple method used to assess the function of pancreatic beta-cells, and the degree of metabolic insulin resistance from the basal (fasting) glucose values and insulin concentration. HOMA index was introduced in 1985 (9). Nowadays is widely recognized and useful assessment tool in clinical and epidemiological studies, revealing very valuable information on IR and beta-cell function (24). Our study data confirmed the existence of IR in subjects with NAFLD. The study group had a significantly higher values of fasting insulin 15.7 ± 7.6 mU/L compared to controls were the average values of this parameter were 6.00 ± 2.8 mU/L (p < 0.01). Beta-cell dysfunction was assessed as HOMA-IR index > 3, established in the group with NAFLD, with average 3.9 ± 2.8 vs 1.2 ±0.6 values, which was statistically significant. Results very similar to ours obtained Ana Lúcia Farias de Azevedo Salgado (32). In the preceding study biochemical analysis of the 116 patients with NAFLD biopsy verified diagnosis, were compared with the results of the healthy subjects. NAFLD patients had higher levels of insulin, glycemia and HOMA-IR values, compared to controls, even when they did not have diabetes or glucose intolerance.

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<tbody>
<tr>
<td>Glycemia (mmol/L)</td>
<td>r = 0.1322; p = 0.07</td>
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<tr>
<td>Triglycerides (mmol/L)</td>
<td>r = -0.06499; p = 0.4591</td>
</tr>
<tr>
<td>Total cholesterol (mmol/L)</td>
<td>r = -0.0922; p = 0.2929</td>
</tr>
<tr>
<td>γ GT(U/L)</td>
<td>r = 0.1567; p = 0.0727</td>
</tr>
<tr>
<td>Insulinemia (mIU/L)</td>
<td>r = 0.2028; p = 0.01</td>
</tr>
<tr>
<td>HOMA-IR</td>
<td>r = 0.151; p = 0.05</td>
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Table 4. Correlation between BMI, glycemia values, triglicerides, total cholesterol values insulinemia and HOMA-IR index. Pearson’s correlation coefficient p-level of the correlation significance.

Finally, by analyzing the above mentioned facts, it can be observed that despite the presence of disturbed anthropometric parameters, responders with nutrition fat-ty liver have also disrupted specific biochemical findings. Presented data confirmed significant deviation from the normal values in levels of glucose, liver enzymes, γ GT, insulin but lipids too. Patients with NAFLD have disrupt-ed lipid status, concluded by several studies (41, 42, 43) as for our results they are in accordance to the cited studies. Accumulation of triglyceride in hepatocytes is considered the main pathogenic trigger in the development of NAFLD (44). Cholesterol metabolism in patients with NA-
FLD is still insufficiently explored, but it is well known that fact that insulin resistance is associated with increased cholesterol synthesis (45). In these patients in order to maintain lipid homeostasis, synthesis of VLDL (very-low density lipoproteins) increases, increased levels of VLDL leads to rise of LDL cholesterol levels (46). The results of our study go along with the last statement in the line, raised levels of the cholesterol of LDL cholesterol was established in patients with NAFLD (47). In order to assess the size of the total fat mass, we used reference values by Bray, so referent values FAT% for men were 12-20%, and for women were 20-30%. This method confirmed that participants with nonalcoholic fatty liver have significantly greater amount of the fat mass, than the controls.

5. CONCLUSION

Our results revile that patients with nonalcoholic fatty liver excessively obese, have enlarged waist circumference, consequential insulin resistance, and impaired glucose metabolism, impaired lipid profile, and thus the risk of developing cardiovascular diseases.

CONFLICT OF INTEREST: NONE DECLARED

REFERENCES