Platelet amino acid spectrum and gut microbiota, their links in patients with coronary artery disease and atrial fibrillation

Abstract. Background. The aim of our work was to identify the links between platelet amino acid (AA) spectrum and gut microbiota composition in patients with coronary artery disease (CAD) and atrial fibrillation (AF) and to evaluate them. Materials and methods. Three hundred patients were enrolled in the study. They were divided into 3 groups: I (CAD) — 149 patients with CAD but without arrhythmias, II (CAD + AF) — 124 people with CAD and AF paroxysm, and control group (CG) — 27 individuals without CAD and arrhythmias. Platelet AA level was assessed by method of ion-exchange liquid column chromatography. Gut microbiota composition was studied by 16-S rRNA sequencing. Results. A significant increase in isoleucine (10.73 %), leucine (12.63 %) and a decrease in threonine (23.05 %), serine (5.06 %), glycine (32.21 %) and valine (30.83 %) platelets levels was found in patients with CAD and AF (P < 0.05). In addition, they had a significant increase in Bacteroides spp., Faecalibacterium prausnitzii, Actinobacter spp., Streptococcus spp., Ruminococcus spp. and a decrease in Lactobacillus spp., Bifidobacterium spp., Eubacterium rectale (P < 0.05). Platelet glutamine acid, valine, glycine, asparagine acid, threonine had the highest number of significant correlations with gut microbiota species (P < 0.05). Actinobacter spp., Blautia spp., Streptococcus spp., Akkermansia muciniphila and Roseburia inulinivorans had the highest number of significant correlations with platelet amino acids (P < 0.05). Conclusions. Platelet amino acid spectrum and gut microbiota composition in patients with coronary artery disease and atrial fibrillation are closely linked. Keywords: coronary artery disease; atrial fibrillation; amino acids; blood platelets; gut microbiota composition

Introduction

Atrial fibrillation (AF) is the most common arrhythmia in the world, which prevalence increased each year. While coronary artery disease (CAD) is the most common cardiovascular disease and one of the known risk factors of AF [1, 2]. Both diseases share associated risk factors, as dyslipidemia, inflammatory diseases, diabetes mellitus, arterial hypertension etc. CAD leads to atrial fibrosis development, which produce the reentry morphological substrate of AF. Near the half of patients with AF have CAD [3].

AF presence increased mortality 1.5—3.5 times, mostly due to stroke [1]. Because, AF is strongly associated with prothrombotic tendency, which pathogenesis is highly intricate and multifactorial. Also increased platelets activity is common for CAD patients. Activated platelets have a lot of prothrombotic and vasoactive factors. One of the important sine of platelets activation is an increasing of mean platelets volume, that shows us the morphological platelets changes. So, platelets are an important part of hemostatic balance and they directly affect prothrombotic state [4]. Each AF paroxysm is associated with platelets activation [5]. Repeated platelets activation is contributing thrombosis formation [6]. Moreover, antithrombotic therapy is essential for patients with AF and CAD because of the high risk of thrombosis, whereas a combination of antiplatelets and anticoagulants is associated with a high risk of bleeding [7].

Due to the recent data, special microbiota signature is common for AF occurrence. In large population-based study AF is characterized by positive associations with genera Eisenbergiella, Enorma, Enterobacter, Kluyvera and negative with genera Bacteroides, Bifidobacterium, Holdemanna, Parabacteroides, Turicibacter [8]. Also, some case-control
studies have shown genus and species changes in gut microbiota in AF patients. As enrichment of *Ruminococcus*, *Streptococcus*, *Enterococcus* and depletion of *Faecalibacterium*, *Alistipes*, *Oscillibacter*, *Bilophila* in AF patients. Or overgrowth of *Parabacteroides*, *Lachnospiraceae*, *Streptococcus*, *Alistipes* and reduction of *Enterobacter* was observed in AF patients in another study [9]. But these results are controversial.

Gut microbiota acts at the heart health through its metabolites toxicity: increasing plasma lipo polysaccharides (LPS), trimethylamine (TMA), trimethylamine-N-oxide (TMAO), bile acids, indole sulfate and decreasing fecal short chain fatty acids (SCFA) due to impaired intestinal barrier function [9]. That leads for autonomic remodeling (increasing sympathetic activity in heart innervation), structural remodeling (cardiac fibrosis, sell apoptosis, increasing conduction velocity) and electrical remodeling (reduction effective refractory period and increase after depolarizations) [10].

Platelets protein composition is an important component of their morphological and functional state. Proteins changes occurs during platelets activation and highly connected with prothrombotic conditions [11]. Also, platelets hyperactivity is connected with gut microbiota metabolites, including plasma amino acids (AA) composition, TMA, TMAO levels [12].

**The aim:** to estimate the links between platelet AA spectrum and gut microbiota composition in patients with CAD and AF and evaluate their connections.

### Materials and methods

300 patients were enrolled in the study. They were divided into 3 groups: I (CAD) — 149 patients with CAD but without arrhythmias, II (CAD + AF) — 124 patients with CAD and AF paroxysm and control group (CG) — 27 patients without CAD and arrhythmias. CAD and AF diagnosis were based on latest ESC guidelines [1, 2]. All patients were treated in the Kyiv City Clinical Hospital 12, cardiological and therapeutic departments. Diagnosis CAD was confirmed by history of coronary arteries stenotic changes during invasive coronarography. AF paroxysm was checked by resting 12 leads electrocardiography. Exclusion criteria were: reported malignancies, chronic kidney disease (glomerular filtration rate (GFR) < 60 mL/min), valcular AF, heart failure class III to IV (by New York Heart Association), thyroid pathology, gout and hyperbilirubinemia are known risk factors of AF paroxysm development [1]. That’s why this baseline characteristics were analyzed and compared, because it can help us to exclude their influence on obtained results.

Platelets AA level was detected by method of ion exchange liquid column chromatography — such AA were identified: lysine, histidine, arginine, ornithine, taurine, asparagine acid, threonine, serine, glutamine acid, proline, glycine, alanine, cysteine, valine, methionine, isoleucine, leucine, tyrosine, phenylalanine, glutamate. Automatic amino acids analyzer T 339 (Mikrotechna, Czech Republic) were used in our study. Study was done in Bogomolets National Medical University. Blood sampling from patients was performed on an empty stomach from the cubital vein on the first day of hospitalization, before treatment. Citrated blood is centrifuged for 10 minutes at a speed of 1500 revolutions per minute. The middle layer was selected with a Pasteur pipette — the plasma is saturated with platelets. The obtained material is again centrifuged for 20 minutes at a speed of 3000 revolutions per minute. The upper supernatant liquid was collected with a Pasteur pipette, and the lower layer was washed with buffer (phosphate buffer solution pH 6.2). Washed platelets are resuspended in buffer (pH 7.4).

Determination of the gut microbiota composition was carried out using quantitative PCR qRT-PCR using primers for the 16S rRNA gene and taxon-specific primers. Such domains were checked: bacteria — *Firmicutes* (*Lactobacillus* spp., *Faecalibacterium prausnitzii*, *Enteroccous* spp., *Blautia* spp., *Streptococcus* spp., *Eubacterium rectale*, *Roseburia inulinivorans*, *Ruminococcus* spp.), *Bacteroides* (*Bacteroides* spp., *Bacteroides thetaiotaomicron*, *Prevotella* spp.) and other (*Bifidobacterium* spp., *Escherichia coli*, *Akkermansia muciniphila*, *Acinetobacter* spp.), and archaea (*Methanobrevibacter smithii* and *Methanosphaera stadtmanae*).

Results were presented as mean ± standard error or [95% confidence interval (CI)] for continuous variables or as a number for categorical variables. Data were compared using Wilcoxon signed-rank test or Student t-test with two critical

### Table 1 — Baseline characteristics of study sample, mean ± standard error

<table>
<thead>
<tr>
<th>Characteristic/group</th>
<th>I group</th>
<th>II group</th>
<th>CG</th>
<th>P I–II</th>
<th>P II-CG</th>
<th>P I-CG</th>
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<tr>
<td>Age (years)</td>
<td>67.71 ± 3.90</td>
<td>67.96 ± 0.94</td>
<td>56.25 ± 2.18</td>
<td>&gt; 0.05</td>
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<td>BMI (kg/m²)</td>
<td>27.02 ± 0.33</td>
<td>26.93 ± 0.43</td>
<td>28.12 ± 2.10</td>
<td>&gt; 0.05</td>
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<td>Total bilirubin (mmol/l)</td>
<td>11.30 ± 0.09</td>
<td>12.40 ± 0.08</td>
<td>11.70 ± 0.11</td>
<td>&gt; 0.05</td>
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<td>Uric acid (mmol/l)</td>
<td>380.50 ± 28.16</td>
<td>404.90 ± 36.11</td>
<td>310.20 ± 29.12</td>
<td>&gt; 0.05</td>
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<td>GFR (ml/min)</td>
<td>62.03 ± 2.31</td>
<td>67.73 ± 1.98</td>
<td>84.01 ± 5.48</td>
<td>&gt; 0.05</td>
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<td>TC (mmol/l)</td>
<td>5.73 ± 0.37</td>
<td>6.18 ± 0.31</td>
<td>4.32 ± 0.21</td>
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Notes (here and in Fig. 2): * — P < 0.05 I–II groups; ** — I group vs CG; *** — II group vs CG.

Figure 1 — Platelets AA spectrum in the investigated groups, μmol/l: A — platelets A spectrum; B — platelets taurine level

Figure 2 — Gut microbiota composition in the investigated groups, mean [95% CI], lg/CFU/ml

Results
At first baseline characteristics of investigated groups were analyzed. All investigated groups were similar by age, BMI and total bilirubin values. I and II groups patients were characterized by increase of TC (by 32.64 and 43.06 %, respectively), uric acid (by 22.66 and 30.53 %) and decrease of GFR (by 26.16 and 19.38 %) in comparison with CG. Data are shown in Table 1.
In this study platelets AA spectrum in I and II groups was compared with CG. During data analysis in the I group in comparison with CG was found significant increasing isoleucine (by 12.41 %), levels and decreasing taurine (by 20.26 %), serine (by 9.31 %) and glycine (by 19.73 %) levels. In II group in comparison with CG significant increasing isoleucine (by 24.47 %), leucine (by 10.20 %) and decreasing taurine (by 19.84 %), threonine by (29.37 %), serine (by 13.90 %), glycine (by 45.59 %) and valine (by 27.87 %) levels. Also, in II group in comparison with the I group significant increasing isoleucine (by 10.73 %), leucine (by 12.63 %) and decreasing threonine (by 23.05 %), serine (by 5.06 %), glycine (by 32.21 %) and valine (by 30.83 %) levels were detected. A general overview of investigated groups plasma AA levels is provided in Fig. 1.

Gut microbiota composition was estimate in investigated groups. By the species analysis results in the II group comparing with I group is significant increasing Actinobacter spp. and decreasing Blautia spp., Roseburia inulinivorans, Bacteroides thetaiotaomicron; in the II group comparing with CG is significant increasing Bacteroides spp., Faecalibacterium prausnitzii, Actinobacter spp., Streptococcus spp. and decreasing Lactobacillus spp., Bifidobacterium spp., Akkermansia muciniphila, Blautia spp., Eubacterium rectale; in the I group comparing with CG is significant increasing Bacteroides spp., Faecalibacterium prausnitzii, Actinobacter spp., Streptococcus spp., Ruminococcus spp. and decreasing Lactobacillus spp., Bifidobacterium spp., Eubacterium rectale. Results are presented in the Fig. 2.

The correlation analysis between platelets AA spectrum and the clinical and laboratory characteristics, gut microbiota composition of the examined groups was done. Gut microbiota composition and platelets AA spectrum had the largest amount of correlations in such species as Actinobacter spp. (total number = 9), Blautia spp. (total number = 8), Streptococcus spp. (total number = 7), Akkermansia muciniphila (total number = 7) and Roseburia inulinivorans (total number = 7); also, such AA as glutamine acid (total number = 11), valine (total amount = 10), glycine (total number = 9), asparagine acid (total number = 9) and threonine (total number = 8). At the same time, the highest amount of correlations was between TC level and platelets AA (total

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<th>Gut microbiota, risk factors/platelets AA</th>
<th>Lysine</th>
<th>Histidine</th>
<th>Arginine</th>
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<th>Taurine</th>
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Figure 3

Moderate positive correlation, 0.3 < r < 0.7
Strong positive correlation, r > 0.7
No significant correlations
Moderate negative correlation, –0.3 > r > –0.7
Strong negative correlation, r < –0.7
number = 7). Strong correlations ($P < 0.05$) were found between: taurine and *Escherichia coli* ($r = 0.714$), asparaginate and *Methanosaeta stadtmanae* ($r = 0.716$), threonine and age ($r = -0.727$), threonine and *Akkermansia muciniphila* ($r = 0.703$), glutamate and *Actinobacter* spp. ($r = 0.744$), glycine and total cholesterol ($r = -0.715$), glycine and *Blautia* spp. ($r = 0.708$), glycine and *Actinobacter* spp. ($r = -0.769$), valine and *Actinobacter* spp. ($r = -0.736$). All correlations are shown in the Fig. 3.

**Discussion**

Unfortunately, it is no evidence about platelets AA spectrum in patients with AF and CAD. But, due to literature data, platelets AA profile changes are present for their risk factors, as diabetes mellitus [15], etc. So, platelets AA spectrum in patients with AF and CAD was checked first time in our work.

The role of platelets in AF pathogenesis is undoubted. Such platelets morphological and functional characteristics as platelet count, mean platelets volume, platelet distribution width, platelet factor 4, beta thromboglobulin and p-selectin are closely linked with AF paroxysm occurrence and consequent stroke. Platelets volume, distribution and factors of activity are significantly higher, but platelets count is lower in AF patients. But it is still no evidence about deep pathophysiologival plot of this changes [4]. Moreover, platelets condition plays an important role in myocardial inflammation and regulates myocardial reperfusion in during myocardial ischemia [16]. It means, platelets characteristics in patients with AF and CAD are interesting pathogenetic aim for investigations.

According to the latest data, BCAA (leucine, isoleucine, valine) takes part in platelets activation. Isoleucine, leucine and valine concurrently used BCAA aminotranspherase and dehydrogenase in their metabolism. In our work significant increasing isoleucine and leucine platelets levels in AF patients was present. At the same time valine was significantly decreased, that can be explained by highly increased valine metabolism in activated platelets. In the animal experiment α-ketoisovaleric acid, which is valine catabolite, is able to activate platelets. Also, BCAA involved in the integrin β3-mediated bidirectional signaling pathway which can modulate platelets activation. Moreover, BCAAs enhanced propionylation of tropomodulin-3 what can lead for platelets hyperactivity [17]. So, role of AA metabolism in AF paroxysm formation in CAD patients is undoubted.

Gut microbiota composition is an important part of human health. Its influence on the host life is made by its metabolites, including fecal SCFA. They are presented mainly by acetic, propionic, butyric, valeric acids. BCAA and SCFA are strongly correlated with each other [18]. SCFA, especially valerate and butyrate, are also able to increase platelets activity. On the other hand, high level of fecal butyrate increases permeability of intestinal cellular barrier. CAD is commonly associated with low fecal SCFA, especially butyrate [19]. Due to the animal studies, SCFA production is associated with *Bifidobacterium*, *Bacteroides*, *Actinobacter*, *Ruminococcus*, *Roseburia*, *Bilophila*, *Coprococcus* species [20]. In our study, also *Bifidobacterium*, *Bacteroides*, *Actinobacter*, *Ruminococcus*, *Roseburia*, *Blautia*, *Akkermansia* are significantly correlated with BCAAs.

On the other hand, glycine, serine, threonine, taurine platelets levels were significantly higher in patients with AF and CAD in our study. By the animal studies, glycine reduces platelets aggregation by activating calcium flux through glycine-gated chloride channels in white blood cells, macrophages and platelets [21]. Glycine deficiency leads to hyperlipidemia by some data. In our study also, total cholesterol plasma level and platelets glycine level were significantly correlated. *Clostridia, Actinobacteria, Bacteroides* are associated with glycine level reduction. In our study *Actinobacter* spp. and *Bacteroides thetaiotaomicron* have significantly negative correlations with platelets glycine levels. Glycine and serine exchange are closely linked. Threonine is also a known glycine resort in microbial and human metabolism [22].

So, gut microbiota and platelets AA profile have strong connections what approved by appearance of strong correlations between their components: taurine and *Escherichia coli* ($r = 0.714$), asparaginate and *Methanosaeta stadtmanae* ($r = 0.716$), threonine and *Akkermansia muciniphila* ($r = 0.703$), glutamate and *Actinobacter* spp. ($r = 0.744$), glycine and *Blautia* spp. ($r = 0.708$), glycine and *Actinobacter* spp. ($r = -0.769$), valine and *Actinobacter* spp. ($r = -0.736$).

By the animal studies, taurine assimilation is closely linked with *Escherichia coli* metabolism. Also, taurine assimilation by *Escherichia coli* depends by the iron presence and SCFA content [23]. Asparaginic acids exchange is closely connected with archaea activity and is crucial for their metabolism [24]. *Akkermansia muciniphila* is widely discussed as a new probiotic bacterium. Threonine supplementation promotes its proliferation in rats and in human feces [25]. *Blautia* spp. is an also promising discussed new probiotic. Its decrease is linked with glycine metabolism in animal studies. *Blautia* spp. and glycine are decrease in aging and low fibers diet, also in animal models [26]. In *Actinobacter* spp. glycine derived pathways are N-oxygenase and amino acid-carrier protein ligase, which regulates bacterial biosynthesis and genes activity [27]. On the one hand, increase of BCAA is commonly associated with metabolic disorders (diabetes mellitus, dyslipidemia) [12], also *Actinobacter* spp. have proatheroscle-rotic and proinflammatory properties [28]. But *Actinobacter* spp. includes variety of species, which influenced in BCAA metabolism in different way. Some of them use valine in the top of metabolites in connection with γ-amino butyric acid [29], what can explain obtained strong negative connections between valine and *Actinobacter* spp. obtained in our study.

In conclusion, gut microbiota is directly linked with platelets amino acids profile. Gut microbiota composition answer for intestinal amino acids exchange, what plays crucial role for all human organism. Probiotics administration can modulate not only gut microbiota composition, but also amino acids profile. Because they improved amino acids and minerals absorption from the nutrients, decries inflammation, normalize lipids exchange [30]. Of course, the type of used probiotic is important and should be performed individually [31]. Moreover, by the latest data some amino acids have strong probiotic properties. For example, in animal studies glycine can decrease endotoxin production, improve anti-inflammatory response, increase non-pathogenic *Escherichia coli* production, potentiate *Lactobacietum* and *Bifidobacterium* activity, etc. [32, 33]. In our study glycine also...
had the largest number of the strong correlations with gut microbiota composition. So, glycine can be used as a promising component in probiotic treatment strategy for patients with CAD and AF.

Conclusions
Platelet amino acids spectrum and gut microbiota composition in patients with coronary artery disease and atrial fibrillation are closely linked:
1. Significant increasing isoleucine (10.73 %), leucine (12.63 %) and decreasing threonine (23.05 %), serine (5.06 %), glycine (32.21 %) and valine (30.83 %) platelets levels in patients with coronary artery disease and atrial fibrillation was found (P < 0.05).
2. Significant increasing Bacteroides spp., Faecalibacterium prausnitzii, Actinobacter spp., Streptococcus spp., Ruminococcus spp. and decreasing Lactobacillus spp., Bifidobacterium spp., Eubacterium rectale in patients with coronary artery disease and atrial fibrillation was determined (P < 0.05).
3. Platelets glutamine acid, valine, glycine, asparagine acid, threonine had the highest number of significant correlations with gut microbiota species (P < 0.05).
4. Actinobacter spp., Blautia spp., Streptococcus spp., Akkermansia muciniphila and Roseburia inulinivorans had the most pronounced amount of significant correlations with platelets amino acids (P < 0.05).
5. Strong correlations were found between (P < 0.05): taurine and Escherichia coli (r = 0.714), asparaginate and Methanosphaera stadtmanae (r = 0.716), threonine and age (r = −0.727), threonine and Akkermansia muciniphila (r = 0.703), glutamate and Actinobacter spp. (r = 0.744), glycine and total cholesterol (r = −0.715), glycine and Blautia spp. (r = 0.708), glycine and Actinobacter spp. (r = −0.769), valine and Actinobacter spp. (r = −0.736).

References


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