Expression analysis of selected classes of circulating exosomal miRNAs in soccer players as an indicator of adaptation to physical activity

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ABSTRACT: Recently studies have shown that, depending on the type of training and its duration, the expression levels of selected circulating myomiRNAs (c-miR-27a,b, c-miR-29a,b,c, c-miR-133a) differ and correlate with the physiological indicators of adaptation to physical activity. To analyse the expression of selected classes of miRNAs in soccer players during different periods of their training cycle. The study involved 22 soccer players aged 17-18 years. The multi-stage 20-m shuttle run test was used to estimate VO2 max among the soccer players. Samples serum were collected at baseline (time point I), after one week (time point II), and after 2 months of training (time point III). The analysis of the relative quantification (RQ) level of three exosomal myomiRNAs, c-miRNA-27b, c-miR-29a, and c-miR-133, was performed by quantitative polymerase chain reaction (qPCR) at three time points – before the training, after 1 week of training and after the completion of two months of competition season training. The expression analysis showed low expression levels (according to references) of all evaluated myomiRNAs before the training cycle. Analysis performed after a week of the training cycle and after completion of the entire training cycle showed elevated expression of all tested myomiRNAs. Statistical analysis revealed significant differences between the first and the second time point in soccer players for c-miR-27b and c-miR-29a; between the first and the third time point for c-miR-27b and c-miR-29a; and between the second and the third time point for c-miR-29b. Statistical analysis showed a positive correlation between the levels of c-miR-29a and VO2 max. Two months of training affected the expression of c-miR-27b and c-miR-29a in soccer players. The increased expression of c-miR-27b and c-miR-29 with training could indicate their probable role in the adaptation process that takes place in the muscular system. Possibly, the expression of c-miR-29a will be found to be involved in cardiorespiratory fitness in future research.

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INTRODUCTION

miRNAs have been identified as small regulators of gene expression by repressing specific target genes at the post-transcriptional level [1, 21]. A single gene can be regulated by multiple miRNAs, and, likewise, a single miRNA may regulate several target genes that are often grouped in a specific biological pathway [3]. Important roles of miRNAs have emerged in the control of metabolic pathways involved in lipid and glucose metabolism [4], energy homeostasis and nutrition [5]. MiRNAs have the ability to control cell proliferation and apoptosis and are located at fragile sites in the genome regions [6].

An interesting feature of miRNA activity is that while miRNAs are often moderate regulators under homeostatic conditions, their function becomes more amplified in response to injury or excessive stress [7]. Moreover, miRNAs have been identified as intracellular modulators of mitochondrial metabolism, inflammation, muscle recovery and hypertrophy. These findings attracted the attention of sports scientists and started the research on miRNA regulation in exercise physiology [8]. The identification of miRNA expression patterns characteristic for physical exercise could be used to monitor physical fatigue and recovery, and even to evaluate physical performance capacity [9]. Moreover, some miRNAs (c-miRNAs) are potential markers of doping manipulations [10].

Physical activity and exercise training induce changes in extracellular and intracellular signalling that influence the expression of genes controlling inflammation, angiogenesis, mitochondrial synthesis, myocardial and skeletal muscle metabolism, regeneration and remodelling [11-14].
The level of soccer ability depends on many factors. Technical drills/technical skills and physical capacity seem to be very important components of a player’s competence. The monitoring and analysis of the training process is an indispensable part of coaching the young players. The volume and intensity of the training are the primary determinants of the training load. Training volume in sports such as soccer is defined as the total duration in minutes or as the number of repetitions of an exercise [15]. The second component of the training load is usually described using heart rates and lactate concentration measures or ratings of perceived exertion (RPE) [16-20]. Moreover, some biomolecular markers, e.g. miRNAs and c-miRNAs, have been identified as intracellular modulators of physical exercises, suggesting their role in measuring the training load or as mediators of exercise training-induced adaptations [21-22].

Long-term endurance training induces many physiological adaptations in both the central and peripheral systems. The physiological adaptations in the central cardiovascular system mainly involve decreased heart rate, increased stroke volume of the heart [23], and increased blood plasma volume without any major changes in red blood cell count. The last item reduces blood viscosity and increases cardiac output. Moreover, this type of training leads to an increase in the total mitochondrial volume and an increase in the maximal oxygen consumption (VO₂ max) [23].

Circulating miRNAs (c-miRNAs) are also small non-coding RNA molecules that are secreted by cells in membrane-enclosed particles that include exosomes or microvesicles, bound to proteins. C-miRNAs have recently been identified extracellularly in body fluids, and differences in peripheral c-miRNA profiles have been observed in several diseases [24]. Furthermore, c-miRNAs have been found to be stable through digestion by ribonucleases (RNases), repeated freeze-thaw cycles, and prolonged storage [24]. These properties mean that extracellular miRNAs (e.g. circulating, exosomal miRNAs) may have potential for use in diagnostic and prognostic testing, as well as in identifying novel pathways and molecular mechanisms of sports. This is why we decided to investigate the effects of training on exosomal miRNA levels. So far, only one prior study has examined the effect of exercise on microRNAs specifically isolated from extracellular vesicles [25].

Expression levels of all the miRNAs we study may fluctuate (increase or decrease) depending on the regulation of particular biological processes involved in physical training. Among these miRNAs, miR-29a, miR-133a-1/-2-3p, let-7a-1/-2-5p, miR-27b-3p, miR-26a-5p, miR-1-3p, and let-7i-1/-2-5p are closely related to myogenesis, cell growth, myocyte proliferation, and cell apoptosis [26].

MiR-27a/b, a potential regulator of myogenesis, could induce skeletal muscle hypertrophy by down-regulating myostatin, an inhibitor of myogenesis [8, 27]. MiR-27b inhibition leads to increased proliferation and delays the onset of differentiation [28].

The miRNA-29 family (29a, 29b, and 29c) targets miRNAs that encode collagens and other proteins involved in fibrosis. Van Rooij et al. [29] validated target genes of the miRNA-29 family involved in cardiac fibrosis that play an important role in fibrosis during cardiac remodelling after myocardial infarction. MIr-133a belongs to the class of muscle-specific miRNAs (myomiRNA) [30] that play a central role in the regulation of myogenesis [31]. This type of miRNA is important in skeletal muscle plasticity because it causes changes in fibre type I/II synthesis and muscle mass regulation in response to activity [26]. Predominantly, miR-133a is a key factor of proliferation and differentiation of cultured myoblasts in vitro [32].

We aimed to examine whether expression levels of three myomiRNAs differed at three time points (I – first, II – second, and III – third time point) during a two-month training cycle. We also investigated the association between myomiRNA levels and VO₂ max (VO₂ max was measured at the first and third time points). Potential expression level differences may support the occurrence of physiological adaptations among individuals during soccer training.

**MATERIALS AND METHODS**

**Material**

The study was approved by the Medical University of Lodz Ethics Committee (RNN/157/16/KE). All participants gave full written informed consent prior to study commencement.

Twenty-two young soccer players entered the experiment (age 17.5±0.70 years, height 178±0.70 cm, weight 68.05±9.18 kg). They showed high sport competence and participated in the junior Regional League in Poland. The experiment was performed in spring during the two-month training cycle of the competition season from the middle of April to the middle of June.

**Methods**

All the players were subjected to the same football training that involved endurance, speed and strength drills (Table 1). Small-sided games and interval runs at anaerobic threshold (AnT) were performed on a natural grass field. The intensity of the effort yielded was determined by measuring the heart rate (HR) that was equal to or higher than the AnT value, but did not exceed 90% HRmax. Individual maximal intensity and lactate threshold of the players running were determined as previously described [33] on a synthetic field at the beginning of the experiment. The test protocol included 3.5–5 min running stages separated by a 1-min rest, during which a capillary blood sample was taken from the fingertip. The initial speed was set at 2.8 m-s-1 and increased by 0.4 m-s-1 after each stage until exhaustion. The Dmax method [34] was used to determine the lactate threshold (V/LT) running velocity and HR/LT. Blood samples were conducted using the Random Access Automatic Biochemical Analyzer for Clinical Chemistry and Turbidimetry A15 (BIO-SYSTEMS S.A.). Lactate concentration was measured using the Rx Monza LC 2389 kit. The manufacturer stated that the method’s intra-assay coefficient of variation (CV) is 3.62%. Moreover, the maximal heart rate (HRmax) was determined during the test. If a
higher HR value was observed during the small-sided games, the higher value was used as the HRmax. An overview of the typical weekly training load during the experiment is presented in Table 1.

**Beep test**

The multi-stage 20-m shuttle run test was used to estimate the VO$_2$max among soccer players. The test involved running continuously between two points 20 m apart. These runs were synchronized with a pre-recorded audio tape, CD or laptop software, that played beeps at set intervals. As the test proceeded, the interval between each successive beep was reduced, forcing the soccer players to increase their speed over the course of the test, until it was impossible to keep in sync with the recording (or, in rare occasions, if the player completed the test). The recording was typically structured into 21 ‘levels’, each of which lasted around 62 seconds. The interval of beeps was calculated to require a speed of 8.5 km/h at the start, which then had to increase by 0.5 km/h with each level thereafter. The progression from one level to the next was signalled by 3 rapid beeps. The highest level attained before failing to keep up was recorded as the score for that test [35-36].

**Serum collection**

Participants visited the laboratory in a fasted state and at least 12 h after exercise. After reporting to the laboratory at a standardized time (between 8 and 10 a.m., intra-individually the same hour for all tests) athletes rested in the supine position for 10 min prior to blood collection. A winged cannula was inserted into the antecubital vein during a short stasis (max. 30 s).

Participation of the players in training sessions was 90%. Before, during, and after the experiment, the tested subjects lived in the school dormitory and were nourished in the same way (sports diet). All the subjects had valid medical cards. All subjects and their parents or guardians were provided with detailed information about the research procedures and gave their written consent.

Samples were collected at baseline (time point I), after one week (time point II), and after 2 months of training (time point III). Blood was collected in Eppendorf tubes and left for about 30-45 minutes at 37°C (until clot formation). Then it was placed in a refrigerator at a temperature of 4°C for several hours (0.5-4 h) until the total organization of the clot. Next, the tube was centrifuged (1200 x g 10 min, 4°C), and serum was separated from the clot carefully into a new sterile tube, frozen and stored at -80°C.

**RNA extraction and complementary DNA (cDNA) synthesis**

All molecular and statistical analyses were conducted at the Medical University of Lodz. RNA was extracted from serum exosomes using the Total Exosome Isolation Reagent (from serum) and Total Exosome RNA & Protein Isolation Kit (Applied Biosystems, USA), according to the manufacturer’s instructions, with a starting volume of 200 μL. From the resulting RNA eluate, cDNA was synthesized using the TaqMan MicroRNA Reverse Transcription Kit (Applied Biosystems, USA) using 5 μl of template RNA. cDNA products were diluted up to 250 μL with double-distilled water and loaded on plates for storage at -20°C until further analysis.

**miRNA analysis**

Commercially available primer assays (Applied Biosystems, USA) were used for quantitative polymerase chain reaction (qPCR) in miRNA analysis (target miRNAs: c-miR-27b, c-miR-29a, c-miR-133; control miRNAs: RNU6B). The candidate miRNAs measured in the

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**TABLE 1.** Overview of typical weekly training load during experiment.

| Day of week | Morning                     | Afternoon                   |
|-------------|-----------------------------|-----------------------------|
|             | Training drills | Intensity (%HRmax) | Duration (min) | Training drills | Intensity (%HRmax) | Duration (min) |
| Monday      | Free                       | -                           | -              | aerobic exercises, coordination, technical | 70             | 90             |
| Thursday    | plyometric and speed, technical | 85             | 80             | small-sided games, tactical | 90             | 90             |
| Wednesday   | Free                       | -                           | -              | coordination, technical, tactical | 70             | 100            |
| Thursday    | Free                       | -                           | -              | technical, tactical     | 75             | 90             |
| Friday      | coordination, technical     | 70             | 75             | interval run, tactical  | 80             | 90             |
| Saturday    | Competition game            |                             |                |                            |                |                |
| Sunday      | Free day                   |                             |                |                            |                |                |
present study were selected due to their reported upregulation at rest in response to endurance rowing training and for their status as myomiRNAs [37-39]. MiRNA levels were measured in triplicate, using the synthesized cDNA with the TaqMan Universal PCR Master Mix (Applied Biosystems, USA). The qPCR mixture contained: cDNA (1 to 100 ng), 20× TaqMan miRNA Expression Assay, TaqMan Universal PCR Master Mix, and RNase-free water in a total volume of 20 μl. The qPCR reactions were performed in the Applied Biosystems 7900HT Fast Real-Time PCR System for 39 cycles, with an annealing temperature of 60°C, repeated 3 times for each sample. The relative expression of the studied samples was assessed using the comparative delta-delta CT method (TaqMan Relative Quantification Assay software) and presented as RQ values, adjusted to the RNU6B expression level.

Five control assays (let-7a-5p, miR-9, miR-21, miR-122, RNU6B) were selected based on previous reports showing expression stability within plasma. Norm finder software was used to determine which single gene or group of genes was at the most stable level within the samples. RNU6B alone was found to be most stable across the three groups and was therefore used to determine the relative expression of the target genes, calculated using the 2-ΔΔCt method. Consistent with RNU6B being the most stable miRNA between groups, data from other reports indicate that the other 4 miRNAs may not have been good candidates for controlling genes due to their association with other stimuli including cancer.

**Statistical analysis**

The Wilcoxon signed-rank test was performed in order to evaluate the differences between the expression levels (RQ values) of the studied miRNAs (c-miRNA-27b, c-miRNA-29a, c-miRNA-133) at the three time points. Spearman’s rank correlation coefficient was used to find relationships between the expression levels of the studied miRNAs and VO₂ max. In this study all statistical analyses were performed using the Statistica 12.0 program.

**RESULTS**

The average values of VO₂ max at the studied time points were calculated. At the first time point, i.e., before the cycle training, the average value was 51.28 (ml/kg/min); at the third time point, i.e., after the training cycle, the average value was 54.93 (ml/kg/min). General characteristics of the subjects are shown in Table 2.

The obtained results showed lower expression levels of all studied myomiRNAs in relation to the calibrator at the first time point: c-miRNA-27b (RQ = 0.34), c-miRNA-29a (RQ = 0.56) and c-miRNA-133 (RQ = 0.64). Analysis of the expression levels of the studied miRNAs at the second time point showed elevations in each case: c-miRNA-27b (RQ = 1.22), c-miRNA-29a (RQ = 1.75) and c-miRNA-133 (RQ = 1.32). After completion of the entire training cycle (third time point), analysis showed once again an increase in expression levels for all studied myomiRNAs: c-miRNA-27b (RQ = 2.53), c-miRNA-29a (RQ = 4.12) and c-miRNA-133 (RQ = 1.49).

**TABLE 2.** Demographic and physiological characteristics of subjects.

|                | Soccer players’ first time point | Soccer players’ second time point | Soccer players’ third time point |
|----------------|----------------------------------|----------------------------------|----------------------------------|
| Age            | 17±0.68                          | 17±0.68                          | 17±0.68                          |
| Weight         | 69±9.55                          | -----                            | 69±9.15                          |
| Height         | 178±6.96                         | 178±6.96                         | 178±6.96                         |
| VO₂ max        | 51.28±3.71                       | -----                            | 54.93±3.00                       |
Statistical significance was obtained between the first and the second time point for c-miRNA-27b (P = 0.0098; Wilcoxon signed-rank test) and c-miRNA-29a (P = 0.0220; Wilcoxon signed-rank test); between the first and third time point for c-miRNA-27b (P = 0.0003; Wilcoxon signed-rank test) and c-miRNA-29a (P = 0.0241; Wilcoxon signed-rank test); and between the second and the third time point for c-miRNA-27b (P = 0.0016; Wilcoxon signed-rank test), as presented in Figures 1, 2, and 3.

We did not find any significant correlations between the second and the third time point for c-miRNA-29a and at any time points for c-miRNA-133 (P>0.05; Wilcoxon signed-rank test).

Finally, we assessed the reciprocal relationship between the expression levels of the studied miRNAs and VO\(_2\) max.

Statistical analysis showed a positive correlation between the expression level of c-miRNA-29a and VO\(_2\) max \((R = 0.54, P =0.008;\) Spearman’s rank correlation coefficient) at the third time point (Figure 4). There was no statistically significant correlation between other studied myomiRNAs and VO\(_2\) max at any of the time points.

DISCUSSION

MiRNAs can be considered as molecular biomarkers related to the type of training – some of them correlate with exercise capacity and/or anthropometric characteristics and/or biochemical markers. We believe that molecular analysis would allow optimization of the training process and would contribute to the players’ harmonious development.

The identification of miRNA expression patterns characterizing physical exercise could be applied for monitoring physical fatigue and recovery, and even for evaluating physical performance capacity [9]. Moreover, as circulating miRNAs (c-miRNAs) are of tissue or cellular origin, they seem to be potential markers of adaptation to physical activity. So far, there has been only one published study regarding the effects of exercise on microRNAs specifically isolated from extracellular vesicles [32]. Our study is the second such study to assess the effects of training on exosomal miRNAs.

In our study we found significantly increased expression of c-miR-27b between all three measured points (the first and the second; the second and the third; the first and the third). There are
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