Effect of Acute High-Intensity Exercise on Rat Myocardium metabolic Profiles. An LC-MS Based Metabolomics Study

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Abstract

Acute high-intensity exercise is a harmful manner associated with a series of myocardial injuries. Metabolism disorder of myocardium is one of the most serious conditions. However, few metabolomics-based studies provide data on the effect of exercise along with myocardial metabolism. Our study aimed to identify metabolic signatures in rat myocardium during acute high-intensity exercise and evaluate their diagnostic potential to sports injuries. SD rats were divided into control group and acute high-intensity exercise group and their myocardium samples were analyzed by LC-MS to explore metabolic alterations of rats’ myocardium. This study showed myocardium metabolism clearly differed between the two groups. there were 6 target metabolic pathways and 12 potential metabolic markers for acute high-intensity exercise. Our findings provide an insight that myocardium metabolism during acute high-intensity exercise have distinct disorders in complex lipids and fatty acids. Moreover, an increase of purine degradation products as well as signs of impaired glucose metabolism were observed. However, the amino acid was enhanced, which had a certain protective effect on the myocardium.

Background

Acute high-intensity exercise refers to the increase of cellular energy consumption and changes of substrate energy supply, which leads to a decrease in ATP synthesis efficiency and disorders of material metabolism. This moment, the oxygen supply of tissue decreases, associated with an electron leakage of the mitochondrial respiratory chain, finally inducing oxidative stress damage[1].

Heart is sensitive to exercise intensity. Strenuous sports make myocardium ischemia and hypoxia with systemic metabolic abnormalities and damages. Overtraining has been able to increase the content of angiotensin II and other hormones sharply, which aggravates cardiomyocyte apoptosis and inflammation, causing cardiac dysfunction[2]. Fine material metabolism provides the organism with ATP to ensure the normal operation of cardiomyocytes[3]. Therefore, improving the material metabolism of myocardium is critical to prevent exercise-induced myocardial injury.

Metabolomics is a new approach following genomics, proteomics and transcriptomics, which has the advantages of high throughput, high specificity and high sensitivity. It can capture metabolic states under the specific external stimuli and scans panoramic metabolites to reveal the transition between health and disease[4]. While, the application of metabolomics technology on sports remains unexplored. Here, we analyzed the changes of metabolomics in rat myocardium during acute high-intensity exercise to explain the effect of acute high-intensity exercise on cardiac function from the perspective of material metabolism.

Experimental

Subjects and groups
16 male SD rats at 7 weeks of age (weighing 190–200g) purchased from Beijing vital river laboratory animal technology biotech (Beijing, China), animal license number: SCXK (Beijing) 2016-0006) were randomly divided into a control group (C) and an acute high-intensity exercise group (E), 8 in each group. Raising condition: Light and dark cycled every 12h, free access to tap water and food, the temperature was 26–28°C, the humidity was 50–60%.

**Exercise program**

The rats exercised on a three-track rat-special treadmill. On the first day after grouping, group E performed an adaptive exercise, the intensity was 3m/min, the time was 5min, 1 week in total; On the 8th day, Exercise intensity pre-adaptation was conducted with the initial speed of 3m/min, took a break every 3 minutes of exercise and the speed increased by 3m/min after rest until the rat was exhausted (exhaustion standard: rats stayed at the last end of the treadmill, run on the abdomen after an electric shock and returned to the original place within three seconds), repeated 5 times; On the 20th day, the exercise capacity was tested, which based on the Bedford's article, the average exhaustion speed was 21m/min (75% VO2max); On the 28th day, did the final acute high-intensity treadmill exercise at a speed of 24m/min, a slope of -16°. The exercise time of each group was 5min, the interval time was 1min, a total of 8 groups.

**Samples**

After exercise (within 10 minutes), decapitated all rats immediately and stripped the myocardium. The myocardium was washed with ice physiological saline, absorbed dry with filter paper and placed in liquid nitrogen for rapid freezing. After that, stored it at -80°C.

**Detection**

**Reagents and instruments**

Acetonitrile (1499230-935, Merckg); Ammonium acetate (70221, Sigma); Ammonia (Fluka). Triple TOF 6600+ mass spectrometer (AB SCIEX); 1290 Infinity LC ultra-high pressure liquid chromatograph (Agilent); ACQUITY UPLC BEH Amide (1.7µm, 2.1 mm×100mm column) column (Waters); 5430R low temperature high speed centrifuge (Eppendorf).

**Sample pretreatment**

Thawed each sample at 4°C, weighed 80mg into 2ml EP tubes. added 200ul water to homogenize and vortexed for 1min. Put the pre-cooled 800ul acetonitrile/methanol mixture (1:1, V/V) into the EP tube and put the sample solution in the refrigerator for 1h to remove the sample protein after two low-temperature ultrasonic treatments (30min/time). Later, centrifugated at 4°C and 14000 rcf for 20 minutes, the supernatant was vacuum freeze-dried and stored it at -80°C.
Added 100ul acetonitrile/water (1:1/V: V) mixture and blended thoroughly to reconstitute sample. After centrifugation at 4°C and 14000rcf for 15 minutes, the supernatant was taken for mass spectrometry injection analysis.

The samples were prepared as quality control samples (QC) for testing the instrument and system status. The QC sample was analyzed 4 times at random.

**Chromatographic conditions**

Put the sample into the autosampler (temperature is 4°C) and added the HILIC chromatographic column. The injection volume was 2ul, the column temperature was 25°C and the flow rate was 0.3ml/min. Phase A: water+25mmol/L ammonium acetate+25mmol/L ammonia; Phase B: acetonitrile. Gradient elution procedure: 0~0.5min, 95%B; 0.5~7min, 95~65%B; 7~8min, 65~40%B; 8~9min, 40%B; 9~9.1min, 40~95%B; 9.1~12min, 95%B. QC were randomly added to the samples to reduce the signal fluctuations and ensure the data reliability.

**Mass spectrometry conditions**

Electrospray ionization source setting parameters: ion source gas 1: 60; ion source gas 2: 60; curtain gas: 30; source temperature: 600°C; ionsapary voltage floating: ±5500V. The secondary mass spectrum was obtained through data correlation acquisition in high-sensitivity mode. Related setting parameters: collision energy: 35±15ev; declustering potential: ±60V; candidate ions to monitor per cycle: 6; exclude isotopes: 4Da.

**Data processing**

Collected data from LC-MS. Used SPSS 24.0 for univariate statistical analysis and SIMCA 14.1 for multivariate statistical analysis based on peak areas data of the detected metabolites. Statistically significant compounds were evaluated by using ROC curve analysis. Made the volcano map and cluster map and Searched Met-PA databases to screen out target metabolic pathways and potential metabolic markers.

**Results**

**QC analysis**

The UHPLC-Q-TOF MS ion chromatograms of the QC sample were overlapped and compared (Figure 1, a). the QC sample chromatograms overlapped well, indicating the instrument was in good condition and the experimental data was reliable.

**Overall sample Hotellings T2**

Overall sample Hotellings T2 analysis was used to detect outliers in this experiment. Here, all samples were within the 99% confidence interval and there were no outliers (Figure 1, b).
**Typical metabolic profile**

Myocardial samples were detected by LC-MS to obtain typical metabolic spectrums (Figure 2). The contours of the myocardial metabolites between the two groups had changed to different degrees. The differences can be found by further analysis.

**PLS-DA analysis results**

PLS-DA uses to identify experimental data and predict differences between the two groups. This experiment supervised the data of the two groups after myocardial preconditioning and established a regression model. The model parameters (7 cycles of verification) showed the model establishment was stable and reliable (Table 1). Constructed a PSL-DA model score chart (Figure 3). The positive and negative ion points between group C and group E had a significant separation trend but within the group were more concentrated, indicating the metabolites of the two groups had differences.

**Table 1 PLS-DA analysis model parameter table**

| Grouping | Negative ion mode | Positive ion mode |
|----------|-------------------|-------------------|
|          | A                 | R²X(cum) | R²Y(cum) | Q²(cum) | A | R²X(cum) | R²Y(cum) | Q²(cum) |
| E/C      | 3                 | 0.477    | 0.999    | 0.897   | 2  | 0.242    | 0.995    | 0.754    |

Note: A is the number of main components; R² is the explanatory rate of model variables to X or Y; Q² is the predictive ability of the model; the closer R²Y and Q² are to 1, the more stable and reliable the model is; the model is stable and reliable when Q² is greater than 0.5

**OPLS-DA analysis results**

OPLS-DA statistical analysis revises the PLS-DA experimental data and enhances the significance of the differences between groups. The model parameters were shown in table 2. In this study, R²Y and Q² were both greater than 0.5, indicating the two groups had significant differences and the OPLS-DA model is stable and reliable. Constructed the OPLS-DA model score chart (Figure 4). The separation trend of positive and negative ions between the two groups was obvious but the tendency of aggregation within the group was obvious, indicating the metabolites of the two groups had significant differences.

Created 200 models on the basis of OPLS-DA to perform permutation tests on the random sorting of categorical variables Y and determined the R² and Q² values of the random model (Figure 5). On the same abscissa, the R² value was greater than the Q² value and can be well separated. The rightmost points of R² and Q² were both larger than the other points, the leftmost value of Q² was less than 0, indicating the model verification of this research had passed, the metabolites of the two groups existed differences and the analysis of PLS-DA, OPLS-DA results was meaningful.
Table 2 OPLS-DA analysis model parameter table

| Grouping | Negative ion mode | Positive ion mode |
|----------|-------------------|-------------------|
|          | A                 | R²ₓ(cum)          | R²ᵧ(cum)  | Q²(cum)  | A                 | R²ₓ(cum)          | R²ᵧ(cum)  | Q²(cum)  |
| E/C      | 1+2               | 0.477             | 0.999     | 0.817    | 1+1               | 0.242             | 0.995     | 0.796    |

Note: A is the number of main components; R² is the explanatory rate of model variables to X or Y; Q² is the predictive ability of the model; the closer R²ᵧ and Q² are to 1, the more stable and reliable the model is; the model is stable and reliable when Q² is greater than 0.5

**Univariate statistical analysis results**

Combined T test and FC analysis to make Volcano Plot (Figure 6). Visually displayed the significantly changed metabolites between the two groups and speed up the screening of potential metabolic markers involved in the pathway. On the basis of p<0.05, FC>1.5 or FC<0.67, the substances represented by the red dots in the upper left and upper right corners of the coordinates were the difference metabolites.

**Comparison results of different metabolites**

The VIP value obtained from OPLS-DA analysis screened the difference metabolites. The T test and FC analysis judged the significance of difference metabolites between the two groups and whether the difference metabolites increased or decreased. VIP>1.0, p<0.05, FC>1.5 represented the differential metabolites were significantly increased; VIP>1.0, p<0.05, FC<0.67 represented the differential metabolites were significantly decreased. The metabolites of the two groups were screened. It was found there were 32 different metabolites between group C and group E in the positive and negative ion mode (Table 3).

Table 3 Different metabolites and change trend in positive and negative ion mode of E/C group (n=8)
| Mode | Quantity | Mass-to-charge ratio | Retention time (s) | Differential metabolites | Variation tendency |
|------|----------|----------------------|--------------------|--------------------------|-------------------|
| ESI+ | 1        | 120.079              | 150.693            | Tyramine                 | ↓**               |
| ESI+ | 2        | 810.599              | 50.791             | 1-Stearoyl-2-oleoyl Lecithin (SOPC) | ↑**               |
| ESI+ | 3        | 496.336              | 216.480            | 1-palmitoyl-sn-glyceryl-3-phosphocholine | ↑**               |
| ESI+ | 4        | 171.004              | 471.382            | Glyceraldehyde 3-phosphate | ↑*               |
| ESI+ | 5        | 145.049              | 335.610            | L- (-) Sorbitose         | ↑*               |
| ESI+ | 6        | 426.318              | 225.515            | cholic acid              | ↓*               |
| ESI+ | 7        | 123.054              | 456.870            | Nicotinamide             | ↓*               |
| ESI+ | 8        | 400.339              | 169.317            | L-palmitoyl carnitine    | ↑*               |
| ESI+ | 9        | 134.044              | 449.359            | L-Aspartic Acid          | ↑*               |
| ESI+ | 10       | 734.564              | 175.564            | Phosphatidylcholine      | ↑*               |
| ESI+ | 11       | 278.061              | 472.020            | D-glucose 6-phosphate    | ↑*               |
| ESI+ | 12       | 175.119              | 579.330            | L-Arginine               | ↑*               |
| ESI+ | 13       | 127.038              | 471.296            | Larric acid              | ↑*               |
| ESI+ | 14       | 109.027              | 452.725            | Quinone                  | ↑*               |
| ESI- | 15       | 175.024              | 370.209            | D-galacturonic acid      | ↑**              |
| ESI- | 16       | 267.195              | 68.536             | Thapsic acid             | ↑**              |
| ESI- | 17       | 187.133              | 102.738            | 3-hydroxydecanoic acid   | ↑**              |
| ESI- | 18       | 180.033              | 124.399            | Acamprosate              | ↑**              |
| ESI- | 19       | 613.137              | 441.629            | Cytidine monophosphate N-acetylneuraminic acid | ↑** |
| ESI- | 20       | 241.082              | 98.070             | Thymidine                | ↑**              |
| ESI- | 21       | 103.039              | 188.660            | D (-)-β-hydroxybutyric acid | ↑** |
| ESI- | 22       | 125.035              | 73.774             | Thymine                  | ↑**              |
| ESI- | 23       | 111.020              | 85.016             | Uracil                   | ↑**              |
| ESI- | 24       | 295.226              | 62.648             | L-arabinose 1,4-lactone  | ↑**              |
| ESI- | 25       | 147.029              | 76.925             | D-arabin-1,4-lactone     | ↑*               |
| ESI- | 26       | 191.016              | 99.294             | D-galactate              | ↑*               |
Hierarchical clustering analysis of differential metabolites

The difference metabolites of myocardial samples between the two groups were analyzed by hierarchical cluster analysis (Figure 7). The red was a significant increase in metabolites, The blue was a significant decrease in metabolites. Here, the color changes of the samples in the same group were relatively concentrated and that of different groups were sharply contrasted, indicating the differences of the myocardial metabolites within the group were small but the differences between the groups were obvious. The selected different metabolites were reliable.

Pathway analysis of target metabolites

MetaboAnalyst 4.0 was used to analyze the differential metabolites between the two groups by Met-PA approach. Imported the data of 32 different metabolites into Pathway Analysis to explore the weight of the metabolic pathways involved (Figure 8). There were 26 metabolic pathways involved during high-intensity exercise (Table 4). Here, Raw p<0.05 and Pathway Impact>0.05 were used as the critical point to screen the above-mentioned metabolic pathways. We found there were 6 potential target metabolic pathways that affect the myocardial metabolism of rats during acute high-intensity exercise, namely fructose and mannose metabolism, Linoleic acid metabolism, pyrimidine metabolism, niacin and nicotinamide metabolism, arginine metabolism, amino sugar and nucleotide sugar metabolism (Figure 9).

Table 4 Differential metabolite pathway analysis results obtained through Met-PA
| Pathway                                           | Total | Expected | Hits | Raw p   | FDR     | Impact |
|--------------------------------------------------|-------|----------|------|---------|---------|--------|
| Linoleic acid metabolism                         | 5     | 0.08284  | 2    | 0.00282 | 0.2148  | 1      |
| Arginine metabolism                              | 14    | 0.23194  | 2    | 0.01365 | 0.39832 | 0.07614|
| Amino sugar and nucleotide sugar metabolism      | 37    | 0.61299  | 3    | 0.02228 | 0.39832 | 0.07043|
| Niacin and Niacinamide metabolism                | 15    | 0.24851  | 2    | 0.02473 | 0.39832 | 0.1943 |
| Pyrimidine metabolism                            | 39    | 0.64612  | 3    | 0.02963 | 0.39832 | 0.17143|
| Biosynthesis of neomycin, kanamycin and gentamicin|2      | 0.03314  | 1    | 0.02963 | 0.39832 | 0      |
| Fructose and mannose metabolism                  | 18    | 0.29821  | 2    | 0.03329 | 0.39832 | 0.12422|
| Biosynthesis of Pantothenic Acid and CoA          | 19    | 0.31478  | 2    | 0.03937 | 0.39832 | 0      |
| Beta-alanine metabolism                           | 21    | 0.34791  | 2    | 0.04664 | 0.42614 | 0      |
| Biosynthesis of unsaturated fatty acids           | 36    | 0.59642  | 2    | 0.1179  | 0.82529 | 0      |
| Glycerophospholipid metabolism                   | 36    | 0.59642  | 2    | 0.1179  | 0.82529 | 0.11182|
| Arachidonic acid metabolism                      | 36    | 0.59642  | 2    | 0.1179  | 0.82529 | 0.33292|
| Aminoacyl-tRNA biosynthesis                      | 48    | 0.79523  | 2    | 0.18753 | 1       | 0      |
| Alpha-linolenic acid metabolism                  | 13    | 0.21537  | 1    | 0.19592 | 1       | 0      |
| Histidine metabolism                             | 16    | 0.26508  | 1    | 0.23559 | 1       | 0      |
| Starch and sucrose metabolism                    | 18    | 0.29821  | 1    | 0.26098 | 1       | 0.13851|
| Pentose phosphate pathway                        | 21    | 0.34791  | 1    | 0.29756 | 1       | 0.18501|
| Galactose metabolism                             | 27    | 0.44732  | 1    | 0.36557 | 1       | 0      |
| Alanine aspartate and glutamic acid              | 28    | 0.46388  | 1    | 0.37627 | 1       | 0.22356|
| Inositol phosphate metabolism                    | 30    | 0.49702  | 1    | 0.39716 | 1       | 0      |
| Arginine and proline metabolism                  | 38    | 0.62956  | 1    | 0.47419 | 1       | 0.05786|
| Biosynthesis of primary bile acids               | 46    | 0.76209  | 1    | 0.54172 | 1       | 0      |
| Metabolic degradation of fatty acids             | 39    | 0.64612  | 1    | 0.48312 | 1       | 0      |
| Tyrosine metabolism                              | 42    | 0.69583  | 1    | 0.50907 | 1       | 0.02463|
| Biosynthesis of fatty acids                      | 47    | 0.77866  | 1    | 0.54955 | 1       | 0      |
| Purine metabolism                                | 66    | 1.09342  | 1    | 0.67608 | 1       | 0.01334|
Note: Total is the total number of compounds in the pathway; Expected is the expected value; His is the number of accurate matches in the uploaded marker data; Raw P is the original P value obtained through the analysis of the pathway score map; FDR is the error trigger rate; Impact is obtained through topological analysis Out-of-path influence value

**Metabolic markers of rat myocardium**

The receiver operating characteristic curve (ROC) evaluated the diagnostic ability of differential metabolites during acute high-intensity exercise. Combined the area value (AUC) and P value (P<0.05) under the ROC curve to screen out the potential metabolism of the above 6 acute high-intensity exercise metabolic pathways. It was Thymine (AUC=1.0), linoleic acid (AUC=0.84), cytidine-phosphate-N-acetylneuraminic acid (AUC=1.0), L-aspartic acid (AUC=0.85), 1-Stearoyl-2-oleoyl lecithin (AUC=1.0), thymidine (AUC=0.93), uracil (AUC=0.89), D-mannose (AUC=0.87), lecithin (AUC=0.95), L-arginine (AUC=0.94), nicotinamide (AUC=0.86), D-mannose, 1-alanine phosphate (AUC=0.91). In this study, these 12 metabolites were regarded as potential markers affecting fatigue metabolism.

| Metabolites                          | KEGG ID | Metabolites                              | KEGG ID |
|--------------------------------------|---------|------------------------------------------|---------|
| Linoleic acid                        | C01595  | D-mannose-1-phosphate                    | C00636  |
| 1-stearoyl-2-oleoyl lecithin         | C00157  | D-Mannose                                | C00159  |
| Nicotinamide                         | C00153  | Uracil                                   | C00106  |
| L-Aspartic Acid                      | C00049  | L-Arginine                               | C00062  |
| Thymidine                            | C00214  | Cytidine-phosphate-N-acetylneuraminic acid | C00128  |
| Thymine                              | C00178  | Phosphatidylcholine                      | C00157  |

**Discussion**

Metabolomics studies the metabolic mechanism from the overall metabolite profile. This study used LC-MS to explore the influence of acute high-intensity exercise on rat myocardial metabolism. We found there were 32 different metabolites, participating in 26 metabolic pathways during acute high-intensity exercise. Among them, fructose and mannose metabolism, linoleic acid metabolism, pyrimidine metabolism, niacin and nicotinamide metabolism, arginine metabolism, amino sugar and nucleotide sugar metabolism were the 6 target metabolic pathways during acute high-intensity exercise involved in 12 potential metabolic markers.

Phosphatidyl choline (PC), 1-stearoyl-2-oleoyl lecithin (SOPC) and linoleic acid (LA) participated in the metabolic pathway of linoleic acid (Figure 10.a). After acute exercise, the content of PC in subjects’ plasma increased, leading to impaired utilization of cardiac fatty acids and inflammation-mediated
metabolic disorders, which induced heart failure\[7\]. The reason may be that under the high oxidative stress conditions, PCs were easily transformed into lysophosphatidylcholine (LPC) catalyzed by phospholipase A2 (PLA2). LPC induces the production of inflammatory factors such as TGF-β1, IL-1β, accelerates the apoptosis of cardiomyocytes and promotes the development of coronary heart disease\[8\]. Lu’s study showed the myocardium produced a large amount of free radicals during high-intensity exercise and led to the overexpression of PLA2, furthermore, accelerated the production of LPC, which finally induced heart damage\[9\]. The metabolic process and biological effects of SOPC are consistent with those of PC. Research showed after excessive consumption of red meat, the content of SOPC was high and trimethylamine oxide (TMAO) was produced, which was a risk factor to induce coronary atherosclerosis and cardiovascular diseases\[10\]. In our study, SOPC and PC in group E increased significantly, indicating acute high-intensity exercise increased the cardiac oxidative stress sharply, possibly producing the lipid peroxidation and toxic substances, damaging the health of myocardium. LA were essential nutrients for organisms, which has the functions of lowering blood pressure and promoting microcirculation\[11\]. However, patients with diastolic dysfunction were found a significant increase of LA in the neointimal part of the myocardium and atherosclerotic plaques which improved lipid metabolism to provide energy for the heart\[12\]. Study also showed LA increased when the myocardium is in a pathological state, inducing myocardial hypertrophy through the calcineurin-activated T cell nuclear factor signaling pathway. In addition, the oxidation products of LA can be easily produced because of oxidative stress during acute high-intensity exercise, which causes macrophage apoptosis and induced coronary plaque rupture, thrombus formation or myocardial infarction\[13\]. In this study, LA in group E was significantly increased, indicating the organism accordingly improved the utilization of myocardium fatty acids during acute high-intensity exercise. However, myocardial ischemia and hypoxia is prone to generate linoleic acid oxidation products or its derivatives, causing heart damage.

D-mannose and D-mannose-1-phosphate participated in the metabolic pathway of fructose and mannose (Figure10.c). D-Mannose exists as a component of mannan in the body, which will be phosphorylated into D-mannose-6-phosphate by hexokinase. Later, a small part is isomerized to form D-mannose-1-phosphate\[14\]. D-mannose is structurally similar to glucose. When its content in organism is high, glucose transporter will be snatched by D-mannose to produce high levels of D-mannose-6-Phosphoric acid, which disrupts the aerobic oxidation of glucose, accelerates glycolysis and causes abnormal energy supply to the myocardium. These changes will hinder succinate-mediated activation of hypoxia-inducible factors, thereby inhibiting the expression of vascular endothelial growth factor and heme oxygenase 1, reducing angiogenesis and cardiac antioxidant capacity and ultimately leading to heart disease\[15\]. Study showed D-mannose in type 2 diabetic rats was significantly increased with the risk of myocardial infarction\[16\]. Ultramarathon runners also been detected high levels of D-mannose in their urine\[17\]. D-mannose-1-phosphate easily reacts with proteins or lipids to participates in the glycosylation process and generate glycosylation end products (AGEs)\[14\]. AGEs cause myocardial lipid metabolism disorders, induce atherosclerosis and mediate myocardial chronic inflammation or cell apoptosis through the myeloid differentiation receptor 2/toll-like receptor 4 pathway, which leads to chronic heart failure\[18\].
Myocardial pressure, corresponding to heart failure, overloaded during high-intensity exercise, which will make material metabolism abnormal and produced more AGEs\textsuperscript{[19]}. In this study, D-mannose and D-mannose 1-phosphate (p<0.05) in group E were significantly increased, indicating the aerobic oxidation of glucose in the rat myocardium during acute high-intensity exercise was blocked and the glucose metabolism was disturbed. At this time, cardiovascular regeneration and antioxidant capacity decreased, promoting the production of AGEs and inducing heart failure.

Cytidine-phosphate-N-acetylneuraminic acid (CMP-Neu5Ac), D-mannose-1-phosphate and D-mannose participated in the metabolic pathway of amino sugar and nucleotide sugar (Figure 10.e). CMP-Neu5Ac is the activated form of Neu5Ac, existing as the component of glycolipids and glycoproteins. The two is positively correlated. Neu5Ac is synthesized in the cytoplasm of eukaryotic cells and transferred to the nucleus. It is activated by CMP-Neu5Ac synthase to transfer cytidine monophosphate (CMP) residues from cytidine triphosphate (CTP) and generate CMP-Neu5Ac\textsuperscript{[20]}. Study showed the increase of CMP-Neu5Ac caused cardiomyocyte apoptosis and inflammatory response through Rho/ROCK-JNK/ERK signaling pathway, interfered with lipid metabolism and accelerated the occurrence of atherosclerosis, resulting in myocardial injury\textsuperscript{[21]}. Neu5Ac can also cause myosin light chain phosphorylation and integrin aggregation, increase the permeability of endothelial cells and promote the release of oxidized low-density lipoproteins and inflammatory factors, which destroyed the intravascular microenvironment and apoptosis of arterial smooth muscles, finally causing a cardiovascular disease\textsuperscript{[22]}. During high-intensity exercise, myocardium ischemic necrosis increased, which promoted the movement of Neu5Ac in serum to the conjugate in plasma, inducing myocardial injury\textsuperscript{[23]}, so the concentration of Neu5AC can represent the level of inflammatory response and be served as a marker for heart diseases. In this study, CMP-Neu5Ac (p<0.01) increased significantly in group E, indicating the heart damage during acute high-intensity exercise may be related to the inflammatory reaction of the heart.

Niacinamide and L-aspartic acid (Asp) participated in the metabolic pathway of niacin and niacinamide (Figure 10.b). Niacinamide is the precursor of Coenzyme I (NAD+) and has a significant antioxidant effect\textsuperscript{[24]}. Nicotinamide can increase the bioavailable NO content and up-regulate the expression of forkhead box protein 1 (Foxo1) by activating Silent Information Regulator 1 (SIRT1), thereby enhancing angiogenesis activity, inhibiting cardiomyocyte apoptosis and maintaining cardiovascular health\textsuperscript{[25]}. After excessive exercise, more nutrients were consumed, resulting in myocardial ischemia or hypoxia and a significant decrease in Niacinamide\textsuperscript{[26]}. However, supplementing nicotinamide during exercise will increase the antioxidant enzyme activity of cardiomyocytes and mitochondrial protein, activate autophagy to degrade damaged cell components in a timely manner, maintain the homeostasis of cardiomyocytes and improve the exercise endurance of rats\textsuperscript{[27]}. In this study, nicotinamide in group E (p<0.05) decreased significantly, showing acute high-intensity exercise reduced myocardial niacinamide, which easily induces myocardial pathological damage. Asp is an important substrate of gluconeogenesis with the effects of protecting cardiovascular health and promoting fatigue recovery\textsuperscript{[28]}. Study showed Asp can reduce hyperammonemia caused by high-intensity exercise and prolong exercise exhaustion time by promoting muscle glycogen retention, free fatty acid oxidation and gluconeogenesis\textsuperscript{[29]}.
Supplementing Asp in repeated cycling sprints can increase the concentrations of glutamic acid, alanine, phenylalanine and total amino acid in blood, reduce the body's lactic acid production by generating carbonates and maintain the normal PH value of the blood, thereby alleviating exercise fatigue and enhancing the output power in bicycle sprinting. In this study, Asp (p<0.05) in group E increased significantly, indicating during acute high-intensity exercise, the rat myocardium will produce Asp to resist the damage of myocardium by improving the oxidative stress, glucose metabolism and lipid metabolism, thereby protecting heart health to a certain extent.

L-arginine (L-Arg) and L-aspartic acid participated in the metabolic pathway of arginine (Figure 10.f). L-Arg is an essential amino acid in human's body with the functions of detoxification and alleviating fatigue. Research showed L-Arg can produced NO, which dilated blood vessels and increased cardiac blood flow to enhance lung ventilation and maximum oxygen uptake, thereby improving coronary perfusion or cardiomyocyte death during strenuous exercise and maintaining the normal cardiopulmonary function. L-Arg can also promote the phosphorylation of PI3K/Akt and then accelerated the secretion of insulin to regulate the glucose transport process, improved the utilization of glucose and alleviated secondary heart damage in diabetic patients. Exogenous supplementation of L-Arg can reduce the MDA of myocardium, lipid peroxides and free radicals, enhance the activity of antioxidant enzymes and ATPase, thereby reducing heart oxidative stress damage after a one-time continuous downhill running. In this study, L-Arg and Asp (p<0.05) in group E increased significantly, indicating during acute high-intensity exercise, the metabolism of amino acid was enhanced, which provided energy for the myocardium and protected the heart.

Uracil, thymidine and thymine participated in the metabolic pathway of pyrimidine (Figure 10.d). The three play an important role in the regulatory functions and energy metabolism. Uracil nucleotides in the venous plasma of patients with myocardial ischemia increased significantly to improve cardiac output, protect the ischemic heart and reduce TNF-a-mediated myocardial apoptosis, which were the important positive inotropic factors and alleviated chronic heart failure. Laitano's research showed during the recovery period after high-intensity running wheel training, the level of uracil in the myocardium of mice was significantly increased, which was essential for myocardial repair after strenuous exercise. Study showed the plasma thymine in mice with acute myocardial ischemia increased significantly and the expansion of thymine was closely related to the risk of heart disease. Peng's research showed a significant decrease in thymidine in rats' serum can be used as a biomarker of early acute myocardial infarction. After strenuous exercise, the increase of thymine in rats' serum had been confirmed and it will induce cardiac ischemia damage. Thymidine supplementation can promote the regeneration of rat myocardial cells, provide energy for the heart and improve anti-fatigue ability. In this study, thymine (p<0.01) in group E increased significantly, indicating during acute high-intensity exercise, the myocardium was in an ischemic state and the heart may have pathological changes. At this time, uracil and thymidine (p<0.01) in myocardium increased which can improve cardiac function.

Conclusion
In this study, we found compared with the control, rat myocardium acute high-intensity exercise had 32 different metabolites and 12 potential metabolic markers which participated in 6 target metabolic pathways by LC-MS and metabolic pathway analysis. Cardiac dysfunction caused by acute high-intensity exercise may be related to myocardial lipid peroxidation, lipid and glucose metabolism disorders. At this moment, the increase of amino acid and nucleotide metabolism in organism can speed up the repair of damaged myocardium and provide energy for myocardium to maintain the normal physiological function of heart. Therefore, improving myocardial material metabolism may be an important target for the treatment of heart disease.

Declarations

Author contributions

L-JW: First author, Providing the funding to this research, Revising the manuscript and confirmation of final version to be published and Submitted manuscript. J-YW: Corresponding author, Substantial contributions to conception and design, performed sample collection and the data analysis, revising the manuscript and confirmation of final version to be published. X-HC: Revising the manuscript. YT: Substantial contributions to conception and design. JL: data analysis.

Ethics approval and consent to participate

This experiment was approved by the ethics committee of Shanxi University (No. SXULL20200064).

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Conflict of interest

The authors declare no conflict of interest.

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Figures
Figure 1

(a) Overlapping TIC spectra of negative and positive ion modes of QC samples; (b) Hotellings T2 diagram in negative and positive ion mode of the sample

Figure 2

The total ion current diagram of typical metabolites in a myocardial sample. (a) Group C negative ion current diagram; (b) Group C positive ion current diagram; (c) Group E negative ion current diagram; (d) Group E positive ion current diagram
Figure 3

PLS-DA score chart of myocardial samples in positive and negative ion mode. (a) negative ions in group C/E; (b) positive ions in group C/E.

Figure 4

OPLS-DA diagram of myocardial sample positive and negative ions. (a) negative ions in C/E group; (b) positive ions in C/E group.
Figure 5

OPLS-DA model replacement test diagram of myocardial sample. (a) E/C group negative ions; (b) E/C group positive ions

Figure 6

The volcano diagram of the positive and negative ion mode of the myocardial sample. (a) C/E group negative ions; (b) C/E group positive ions
Figure 7

Metabolite hierarchical clustering of significant differences in positive and negative ion patterns in myocardial samples. (a) E/C group negative ions; (b) E/C group positive ions

Figure 8

Acute high-intensity exercise metabolic pathways constructed by MetPA database.

Note: The abscissa pathway impact is the importance value of the metabolic pathway obtained by topological analysis, and the ordinate -logP is the significance level of the metabolic pathway enrichment analysis; the greater the pathway impact and -logP value are, the higher the correlation of the metabolic differences between different groups is, the bigger the circle is.
Figure 9

Metabolic pathways involved in differential metabolites. Red is the potential marker of the pathway involved in this study; Blue is not in the metabolites of this study