Effect of cold adaptation on activities of relevant enzymes and antioxidant system in rats

Ji-Qing Xing¹, Yang Zhou², Jian-Feng Chen³, Shang-Bin Li¹, Wei Fang¹, Jun Yang¹

¹Department of Physical Education, Harbin Engineering University, Harbin 150001, China; ²Department of Radiology, Harbin Medical University Cancer Hospital, Harbin 150081, China; ³Animal Experimental Center, Harbin Medical University, Harbin 150081, China

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Abstract: Exercise in cold environments can cause significant metabolic regulation and antioxidant behavior. For discussing enzymatic responses towards cold adaptation, we investigated enzyme activities of adenylate cyclase (AC) and phosphodiesterase (PDE) in liver, skeletal muscle, and brown adipose tissue (BAT), as well as Na⁺-K⁺ ATPase and Na⁺/K⁺ ratio in blood. Malondialdehyde (MDA) and superoxide dismutase (SOD) activity in blood were also studied to address the effect of cold adaptation on oxidative damage and antioxidant system. Experimental results indicated that enzyme activities in liver, skeletal muscle and BAT maintained relatively constant for the control group. For the cold adaptation group, enzyme activities in liver and skeletal muscle were in high levels at the beginning, and then gradually decreased to similar values with the control group. However, enzyme activities in BAT performed an increasing trend and significantly higher than the control at the end. In addition, decreased oxidative damage and activated antioxidant system was observed along with the cold adaptation process.

Keywords: Cold adaptation, adenylate cyclase, phosphodiesterase, Na⁺-K⁺ ATPase, malondialdehyde, superoxide dismutase

Introduction

Cold exposure is a common issue in several sporting events (winter swimming, etc.) and special work in both indoor and outdoor conditions (fishing, forestry, dairy industries, etc.) [1]. In recent years, exercise in cold environments has attracted much attention on its potential benefits for athletes [2, 3]. Exercise in cold environments means that human body has to produce more heat for maintaining normal body core temperature, involving a series of physiological response within human body to change the utilization patterns of energy substances [4-6]. These metabolic processes lead to the enhancement of blood pressure, heart rate, and oxygen consumption [7]. It is well known that the increase of oxygen consumption (or energy consumption) can be mainly due to the thermogenesis effect and elimination of lactic acid [8]. Although some studies indicated that the local hypothermia can delay the release of lactic acid, the metabolic block of lactic acid from reduced blood supply resulted in lactic acid accumulations to some extent and obviously increased the respiratory quotient [9, 10].

Exercise in cold environments results in significant variation to body metabolisms. Sink et al. carried out the training project under different temperatures with the training strength of 50% VO₂max and 60% VO₂max, and the concentrations of plasma fatty acids and glycerol (as the lipid metabolic markers) significantly increased at 0°C environmental condition [11]. Fink et al. made the excise project with the training strength of 70%~80% VO₂max, and they found that the concentrations of triglyceride in muscle decreased for 23% under cold environments, and only 11% under warm condition. Parkin et al. found that after the exhausting excise of 70% VO₂max, muscle glycogen concentration in vastus lateralis was 30% higher under cold environments (3°C) than that under normal temperature (20°C) [12]. Febbraio et al. attributed the decrease of glycogen decomposition rate to the decrease of local muscle temperature and the sympathetic nervous system excit-
ability [13]. They concluded that high lipid utilization and low glycogen utilization occurred under cold environments. As a special physiological response towards cold exposure, the phenomenon can be explained as a protection mechanism for maintaining body core temperature [14]. Since the demand of glycogen is significantly increased under cold environments, skeletal muscle has to obtain more energy from lipid. In this way, more energy can be saved for making sure the organ function to sustain life [15].

The variation of body metabolisms under cold environments can be fundamentally due to metabolic regulations by relative important enzymes and compounds. For further understanding the stress/response metabolic mechanisms towards exercise in cold environments, in this study, we investigated enzyme activities of adenylate cyclase (AC) and phosphodiesterase (PDE) in liver, skeletal muscle, and brown adipose tissue (BAT) of rats, as well as Na⁺·K⁺ ATPase, K⁺ concentration, and Na⁺ concentration in blood. In addition, malondialdehyde (MDA) concentration and superoxide dismutase (SOD) activity were measured to discuss oxidative damage and free radical metabolism during cold adaptation process.

Materials and methods

Samples

One hundred Sprague-Dawley (SD) rats (50 males and 50 females) were chosen as the samples in this study, which was approved by the Ethics Committee in our institution. The rats were fed in the same conditions and all performed perfect physical status. While the rats were about 8 weeks old and had a weight range from 156 g to 203 g, they were prepared for the following experiment.

Experimental procedures

The rats were randomly divided into the control group and the cold adaption group with 50 individuals in each group. All the experiments were carried out in a temperature-controlled swimming chamber (Western Environmental Chamber, Napa, CA). For each day during the experimental process, the rats in the control group swam in the chamber at 20°C for 30 min, and the rats in the cold adaption group experienced a cold swimming exercise at 4°C for 30 min. At the end of the 1, 2, 3, 4, and 6 weeks, ten rats in each group were randomly chosen and killed respectively by decapitation. Blood sample, left liver lobule, intestinal muscle of right calf, and BAT in scapular region were taken for analyzing enzyme activities and other items.

Analytical methods

Enzyme activities of AC were measured by the unlabeled substrate method described in a previous study [16]. Enzyme activities of PDE were investigated by using cytochemical procedures described by Okruhlicova et al. [17]. K⁺ concentration and Na⁺ concentration in blood were examined by a Beckman CX-3 ion selective electrode. Na⁺·K⁺ ATPase activities, MDA concentration, and SOD activities were analyzed by using kits from Nanjing Jiancheng Biotechnology Institute, including the Na⁺·K⁺ ATPase assay kit, the MDA assay kit (TBA method), and the SOD assay kit (hydroxylamine method), respectively.

Statistical analysis

The experimental results were shown as mean ± standard deviation. Differences in experimental data between groups were evaluated by statistical analysis, which was performed with the use of the commercially available SPSS 16.0 software. The independent samples t tests were used to characterize the anthropometric and baseline data between groups. The Kolmogorov-Smirnov test was used to ensure normality. The analysis of variance (ANOVA) was conducted to locate the differences with the level of significance set at $P < 0.05$.

Results

Effects of cold adaptation on AC enzyme activities

Table 1 represents time-course profiles of AC enzyme activities in liver, skeletal muscle and BAT. AC enzyme activities in the control group maintained relatively constant during the whole period. In the cold adaptation group, AC enzyme activities performed a decrease trend in liver and skeletal muscle, and gradually increased in BAT. After six weeks of cold adaption, AC enzyme activities were almost the same in liver (control: 2.49 umol/mg-prot·h; cold adaption: 2.66 umol/mg-prot·h), and skeletal muscle (control: 2.19 umol/mg-prot·h; cold adaption: 2.32 umol/mg-prot·h) for the two groups. However, AC enzyme activities in BAT were sig-
Enzyme/antioxidant responses for cold adaptation

Table 1. Time-course profiles of AC enzyme activities in liver, skeletal muscle, and BAT

<table>
<thead>
<tr>
<th>Time</th>
<th>Liver (umol/mg-prot·h)</th>
<th>Skeletal muscle (umol/mgh)</th>
<th>BAT (umol/mg-prot·h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Cold adaption</td>
<td>Control</td>
</tr>
<tr>
<td>1 week</td>
<td>2.22 ± 0.23</td>
<td>4.26 ± 0.40*</td>
<td>2.07 ± 0.33</td>
</tr>
<tr>
<td>2 weeks</td>
<td>2.31 ± 0.26</td>
<td>3.70 ± 0.32*</td>
<td>2.14 ± 0.35</td>
</tr>
<tr>
<td>3 weeks</td>
<td>2.42 ± 0.27</td>
<td>3.24 ± 0.27*</td>
<td>1.98 ± 0.26</td>
</tr>
<tr>
<td>4 weeks</td>
<td>2.58 ± 0.29</td>
<td>2.86 ± 0.22</td>
<td>2.22 ± 0.23</td>
</tr>
<tr>
<td>6 weeks</td>
<td>2.49 ± 0.25</td>
<td>2.66 ± 0.30</td>
<td>2.19 ± 0.27</td>
</tr>
</tbody>
</table>

*Represents the difference between two groups is significant (P < 0.05).

Table 2. Time-course profiles of PDE enzyme activities in liver, skeletal muscle, and BAT

<table>
<thead>
<tr>
<th>Time</th>
<th>Liver (umol/mg-prot·h)</th>
<th>Skeletal muscle (umol/mgh)</th>
<th>BAT (umol/mg-prot·h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Cold adaption</td>
<td>Control</td>
</tr>
<tr>
<td>1 week</td>
<td>2.18 ± 0.31</td>
<td>3.18 ± 0.31*</td>
<td>2.39 ± 0.19</td>
</tr>
<tr>
<td>2 weeks</td>
<td>2.24 ± 0.27</td>
<td>3.06 ± 0.28*</td>
<td>2.55 ± 0.26</td>
</tr>
<tr>
<td>3 weeks</td>
<td>1.95 ± 0.22</td>
<td>2.68 ± 0.32*</td>
<td>2.33 ± 0.29</td>
</tr>
<tr>
<td>4 weeks</td>
<td>2.08 ± 0.26</td>
<td>2.25 ± 0.19</td>
<td>2.19 ± 0.32</td>
</tr>
<tr>
<td>6 weeks</td>
<td>2.12 ± 0.20</td>
<td>2.19 ± 0.21</td>
<td>2.07 ± 0.31</td>
</tr>
</tbody>
</table>

*Represents the difference between two groups is significant (P < 0.05).

Figure 1. Enzyme activities of Na⁺K⁺ ATPase at different time for the control group and the cold adaptation group.

![Figure 1](image1.png)

Figure 1. Enzyme activities of Na⁺K⁺ ATPase at different time for the control group and the cold adaptation group.

Significantly higher (7.35 umol/mg-prot·h) in the cold adaption group than those in the control (2.26 umol/mg-prot·h).

Effects of cold adaptation on PDE enzyme activities

Enzyme activities of PDE in liver, skeletal muscle and BAT are listed in Table 2. For the control group, the enzyme activities of PDE were relatively stable, with the values ranged from 1.95-2.14 umol/mg-prot·h (liver), 2.07-2.55 umol/mg-prot·h (skeletal muscle), and 2.72-2.92 umol/mg-prot·h (BAT), respectively. For the cold adaption group, PDE enzyme activities performed a decrease trend in liver and skeletal muscle, and finally kept at 2.19 umol/mg-prot·h and 2.37 umol/mg-prot·h, respectively. Impressively, PDE enzyme activities in BAT dramatically increased for the cold adaption group and finally reached 8.07 umol/mg-prot·h.

Effects of cold adaptation on enzyme activities of Na⁺K⁺ ATPase

Figure 1 showed the variation trend of enzyme activities of Na⁺K⁺ ATPase for the two groups. For the control, the enzyme activities of Na⁺K⁺ ATPase was almost the same all the time with little fluctuation (0.21-0.25 umol/mg-prot·h). For the cold adaption group, the enzyme activities of Na⁺K⁺ ATPase experienced a continuous drop from 0.46 ± 0.11 umol/mg-prot·h to 0.30 ± 0.08 umol/mg-prot·h. At the end of the experiment, the cold adaption group showed relatively higher Na⁺K⁺ ATPase activity than the control group.

Effects of cold adaptation on K⁺ concentration, Na⁺ concentration and Na⁺/K⁺ ratio

Table 3 illustrates the variation of K⁺ concentration, Na⁺ concentration and Na⁺/K⁺ ratio at different times for the two groups. K⁺ concentration, Na⁺ concentration and Na⁺/K⁺ ratio for the control group were relatively steady all the time.
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Figure 3. SOD activities at different time for the control group and the cold adaptation group.

However, for the cold adaptation group, K⁺ concentration indicated an increasing trend and Na⁺ concentration decreased along with the adaptation time, leading to a remarkable decrease of Na⁺/K⁺ ratio (from 0.38 ± 0.08 to 0.18 ± 0.05). At the end of the cold adaptation process, we can not find significant differences of K⁺ concentration, Na⁺ concentration and Na⁺/K⁺ ratio for the two groups.

**Effects of cold adaptation on MDA concentration and SOD activity**

The change of MDA concentrations for the two groups is shown in Figure 2. For all the time, MDA concentration for the control group was almost the constant around 1.6 nmol/mg-prot. For the cold adaptation group, the MDA concentration was extremely high at the beginning (3.68 nmol/mg-prot.), followed by a rapid decrease to 1.62 nmol/mg-prot. At the end, MDA concentrations were almost the same for the two groups. Figure 3 illustrated the variation trend of SOD activities for the two groups. For the control group, SOD activities performed less fluctuation ranged from 221.08 U/mg-prot to 256.78 U/mg-prot. For the cold adaptation group, SOD activities slowly increased from 157.81 U/mg-prot to 267.11 U/mg-prot.

**Discussion**

Recently, exercise in cold environments became a hot topic for enhancing exercise efforts of athletes. Under the cold conditions, human body can alter the neuroendocrine regulation and metabolic levels to maintain core temperature and normal life activities [19]. After a long period of cold stimulation, a series of biochemical and physiological regulations are induced to increase cold resistance capability, which finally leads to the status of cold adaptation. Generally, the cold adaptation process can be divided into two phase: cold stimulation and cold adaptation, characterized by the variations of important enzymes and compounds.

In this study, for the beginning three weeks, AC and PDE activities in liver and skeletal muscle
of the cold adaptation group were significantly larger than those of the control group (P < 0.05). For the following three weeks, AC and PDE activities in BAT significantly increased and played an important role in heat production. It can be concluded that the beginning three weeks was cold stimulation period, which was characterized by main heat production by liver and skeletal muscle. In this stage, enhanced catecholamine hormones had regulatory effects on tissue cells, resulting in the reinforcement of AC and PDE activities. By catalyzing cytoplasm ATP into cAMP, Fat kinase was activated to strengthen lipolysis. More free fatty acids were produced to supply abundant substrates for oxidative phosphorylation of tissues, leading to the enhancement of tissue heat production [20].

For the following three weeks, it can be recognized as the cold adaptation period. In this stage, heat production in liver and skeletal muscle decreased and enzyme activities in BAT performed higher values. In previous studies, it has been proved that cold adaptation was mainly achieved by increased heat production from BAT and regulated by sympathetic nervous system [21]. During the process, expression of some important components in BAT (uncoupling proteins, thyroxine- 5’ - iodine enzyme, lipoprotein lipase, etc.) was increased to accelerate the decomposition of triglyceride in plasma lipoprotein, resulting in strengthened heat production and decrease of body fat [22]. Since lower body fat concentration might be more susceptible towards sympathetic stimulation, heat production process can be better operated to maintain core temperature, and further strengthen athletes’ physique [23].

In addition, cold adaptation process led to the increase of Na⁺-K⁺ ATPase activity and Na⁺/K⁺ ratio, which act vital efforts on maintain normal physiological function of cells. It has been assumed that exposure in cold environments induced the production of more catecholamine hormones, which act on sodium ion channel of cell membrane and increase the sodium ion concentration. Sodium pump was activated to uptake sodium and release potassium. At the same time, ATP was largely decomposed to release much energy for meeting metabolic requirements of body [24].

Exercises in cold environments were accompanied by enhanced metabolic processes and energy consumption. Moreover, more oxygen free radicals were simultaneously produced to make damage towards lipids, proteins and DNA. It was reported that cold stimulation led to the decrease of SOD activity in rat hypothalamus and adrenal medulla [25]. Kaushik et al. found that cold exposure led to the change of oxidative stress and antioxidant system of rat tissues [26]. In this study, we investigate the effect of cold adaptation on oxidative damage and antioxidant system. MDA and SOD were chosen as the important indexes to evaluate the contents of oxidative stress injury and antioxidant repair. We concluded that cold adaptation process can strengthen the oxidative stress level and decrease the antioxidant ability at the beginning. However, systemic adaptation was successfully established after a long period, including the activation of antioxidant system, intervention of repair/elimination system, and change of redox sensitive transcription. Based on effective removal of oxygen free radicals, virtuous circle can be made to strengthen the adaptation capability of body antioxidant defense system.

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Disclosure of conflict of interest

None.

Address correspondence to: Shang-Bin Li, Department of Physical Education, Harbin Engineering University, Harbin 150001, China. Tel: 86-451-82519915; Fax: 86-451-82519915; E-mail: lishangbinheu@163.com

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