Effects of Exercise Training on Behavior and Brain Function after High Dose Isoproterenol-Induced Cardiac Damage

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Abstract

Acute sympathetic stress can result in cardiac fibrosis, but may also lead to mental dysfunction. Exercise training after isoproterenol (ISO)-induced acute sympathetic stress was investigated regarding cardiac damage, neuroinflammation, brain function and behavior.

Male Wistar rats (12 months) received ISO or saline. One week later, treadmill running or control handling (sedentary) started. After four weeks, cognitive- and exploratory behavior were evaluated, and heart and brain tissues were analyzed regarding cardiac damage, hippocampal neuroinflammation and neuronal function.

ISO did not affect cognitive performance nor hippocampal function. However, ISO reduced anxiety, coinciding with locally reduced microglia (processes) size in the hippocampus. Exercise in ISO rats reversed anxiety, did not affect microglia morphology, but increased brain function.

Thus, exercise after ISO did not affect cardiac damage, cognition or hippocampal neuroinflammation, but normalized anxiety. Increased localized BDNF expression may indicate improved brain function.

Introduction

Acute sympathetic stress is characterized by overactivation of beta-adrenergic receptors, resulting in cardiac fibrosis and ultimately cardiac dysfunction [1]. Acute administration of high dose of beta-receptor agonist isoproterenol (ISO) is commonly used to mimic this clinical condition [2–4]. ISO, administered twice with 24h in between, evoked inflammation and increased cytokine production resulting in cardiac fibrosis [1, 3], that over a period of weeks developed into left ventricular hypertrophy and dilatation, and ultimately heart failure [4]. This was associated with reduced exploratory behavior [5] and cognitive decline [6].

Numerous studies have been performed to investigate potential therapeutic approaches in the ISO model, including physical exercise [1, 7]. However, usually these interventions were provided before, during or immediately after the ISO application, in order to prevent deleterious consequences. However, as Alemasi and coworkers [1] already indicated, acute stress is often caused by unexpected events and is difficult to predict. Therefore, it seems worthwhile to explore interventions that could be applied after ISO had exerted its effect on the heart, and cardiac fibrosis was eminent.

Exercise training is generally acknowledged for having positive effects on physical as well as mental conditions [8]. Evidence is accumulating for an association between mental dysfunction and neuroinflammation [9]. Exercise is known to have anti-inflammatory effects [10]. Although effects of potential treatment on cardiovascular aspects are commonly investigated in the ISO model [1, 7], effect on brain and behavior are only sparsely studied [5, 6]. Exploratory behavior six days after ISO (+ pituitrin) could be improved by Corvitin and 2-Oxoglutarate [5], while cognition, tested 2-8 days after ISO, could be
improved by sodium thiosulfate [6]. However, also in these studies interventions started at the time of ISO application, suggesting prevention rather than treatment of consequences.

Aim of the present study was to investigate effects of exercise training, starting after cardiac damage induced by high dose of ISO has been established, with focus on the brain. For that, effects on neuroinflammation, neuronal function and behavior were studied.

**Material And Methods**

**Animals and experimental protocols**

The study has been separated into two experiments. First, to validate the model of ISO in our hands, the short-term effects of ISO on mortality, cardiac function and damage, and neuroinflammation were studied in young rats. Because of availability of ECHO-cardiography equipment for rats necessary to measure cardiac function, this part of the study was performed in Groningen. For that, male 3 months old Wistar rats (Harlan), housed 2-3 per cage with food and tap water available at lib, were used. All methods were performed in accordance with the ARRIVE guidelines. All experiments were performed in accordance with relevant guidelines and regulations/legislations. Experimental animals and procedures were approved by the local animal committee of the University Groningen, the Netherlands. Rats were randomized to receive either ISO or saline treatment. One week later, rats were anesthetized for cardiac function assessment. Subsequently, rats were sacrificed and heart- and brain tissues were dissected and processed for (immuno)histochemistry.

The actual study, long term effects of physical training after ISO, was performed in Budapest, for the local availability of adult rats, and exercise equipment (treadmill). For that, 12 months old Wistar rats (breeding colony of University of Physical Education, Budapest, Hungary) were used. All methods were performed in accordance with the ARRIVE guidelines. All experiments were performed in accordance with relevant guidelines and regulations/legislations. Experimental animals and procedures were approved by the local animal committee of the University of Physical Education, Budapest, Hungary. Rats were randomized to receive either ISO or saline treatment. One week later, rats were randomized to physical exercise by treadmill running, or were handled but remained sedentary for five weeks. During the sixth week following the injections, behavioral testing was performed, and rats were subsequently sacrificed. Heart and brain tissues were dissected and processed for (immuno)histochemistry.

**ISO treatment**

Animals were randomly assigned to the two experimental interventions. The experimental group received 70 mg/kg ISO dissolved in 1 ml/kg saline, control rats received 1 ml/kg saline. Administrations were performed on two consecutive days via intraperitoneal injections.
Cardiac function

Cardiac function was estimated with transthoracic echocardiography (Sonos 5500, Philips, the Netherlands). For that, isoflurane anesthetized (±2% in air/oxygen= 2/1) rats were placed in supine position on a heating pad to maintain body temperature at 37°C. Standard two-dimensional and M-mode long- and short-axis images at the midpapillary level were acquired using a 12-13 MHz transducer. Cardiac function was determined as left ventricular ejection fraction (EF).

Exercise

Exercise training (running) was performed on a six-lane rat treadmill (Tartonik Elektronika, Italy) with individual lanes of 12*54*13 cm. The training program lasted for 5 weeks, 5 times per week on each weekday. On the first week of the training program rats were habituated to running: on the first day 10 minutes of running with a maximal speed of 10 m/min, which was gradually raised to 30 minutes and maximal speed of 18 m/min by the fifth day. For the following four weeks each running session lasted 30 minutes: starting with a 5-minutes warm up to reach the desired speed. Running at a speed of 18 m/minutes was considered a moderate intensity of approximately 65% of VO₂max (pilot study).

Behavior

Different aspects of behaviour were assessed. Effects on anxiety were assessed by exploratory behavior in an open field (OF) test. Cognition was measured as short-term memory in the novel object recognition test (NOR) and the novel location recognition test (NLR). All tests were recorded with a digital video camera and stored on a memory card. Tests were carried during at the end of the five weeks's intervention, and encompassed 10 days.

Open field exploration
Rats were placed in a round shaped arena and were given 5 minutes to freely explore it, while behavior was recorded. The arena was divided into wall and center areas, delineated by concentric circles. Time spent in the areas were measured from the recordings by Eline (University of Groningen), regarding time spent in center and wall areas. Percentage of time spent in the area along the wall was taken as a measure for anxiety/depressive-like behavior. After removal of animals, the arena was cleaned with 70% ethanol and paper tissue.

Novel object and novel location recognition
The two memory tests were combined in one protocol [11]. The rat was placed in a box shaped arena and was let 3 minutes to get accustomed to the settings, then two identical objects were placed into the arena. After again 3 minutes the objects were removed and cleaned, and 1 minute later, one familiar and
one novel object were placed back on the same locations. Again, after 3 minutes objects were removed, cleaned, and after 1 minute placed back but one of the objects was moved to a new location. Test was ended 3 minutes after exploration of the last setting. Arena and objects were cleaned with 70% ethanol after each animal. Time spent exploring the objects was measured (Eline). Preference for the novel object (NOR) or novel location (NLR) was calculated as percentage time spent exploring the novel/relocated object divided by time spent exploring both objects.

(Immunohistochemistry)

Cardiac collagen
Since ISO was anticipated to cause focal myocardial infarcts, percentage collagen was used to measure cardiac damage. For that, 20 µm thick transverse slices at mid-ventricular and apex level of the heart were stained with Sirius red (Sigma, Aldrich) and fast green as counter staining [12]. Colour pictures were taken. Image analysis (Image Pro plus, USA) was used to measure the collagen positive (red) area and was expressed as percentage of total left ventricular tissue area.

Neuroinflammation
Since microglia are regarded the immune cells of the brain, and change shape when activated, microglia morphology was used to measure microglia activity. In IBA-1(Wako, USA) stained brain slices (20µm thick), different hippocampal areas, Cornu Ammonis (CA)1 and Hilus [11] were photographed (200X), and microglia were analysed, according to altered morphology, including density, coverage, cell size, cell body size and processes size. Microglia cell body to cell size ratio was used as a measure for microglia activity; neuroinflammation [13].

Brain function
For brain function, brain slices were stained with Brain Derived Neurotrophic Factor (BDNF) antibody (Alomone Labs, Israel). In the different areas of the dorsal hippocampus, CA1, CA3, Dendate Gyrus and Hilus, BDNF expression was obtained as corrected optical density (Image-J) compared to an underlying reference area [11].

Data analyses
The study has been reported in accordance with ARRIVE guidelines. All reports were performed in accordance with relevant guidelines and regulations/legislations. Data are presented as mean and standard error of mean (SEM), unless indicated otherwise. Results outside twice the standard deviation of its group were considered outliers and were excluded before analyses (maximally 1-2 per experimental group). Data of the first study, to establish the ISO model, were compared with a student's T-test for independent samples, for ISO versus saline treatment. Data of the actual study, effects of long-term exercise in the ISO model, were compared using two-way analysis of variance (ANOVA) with least square difference (LSD) post-hoc test, with saline/ISO and sedentary/runner as factors. Association between selected parameters were measured with Pearson linear correlation. For the Novel Object / Novel Location
recognition test, outcomes were also tested against change level (=50%), using a single sample t-test. A p-value of <0.05 was considered statistically significant and presented as *. Potentially relevant tendencies (p<0.1) were mentioned as well.

Results

The ISO model

Although rats exhibit symptoms described by Wexler and Kittinger [14], including prostrate and stuporous behavior, with irregular breathing, they recovered well and mortality was 14%. One week after the injections, no decline in cardiac function was observed (Table 1). Heart and lung weights were not affected by ISO (Table 1). However, cardiac collagen percentage, representing damage, was significantly higher after ISO, both at the mid-ventricular level and at the apex (Figure 1). Absence of increased tissue area in microscopical slices supported the lack of increased heart weight. Apart from a significantly decreased relative brain weight, no effects of ISO were observed on organ weights.

Table 1
Body- and organ weights, as well as heart rate and left ventricular function (ejection fraction) obtained from ECHO-cardiography, 1 week after isoproterenol (ISO) or saline treatment. *: significant effect of ISO

|                    | saline   | ISO       |
|--------------------|----------|-----------|
| Body weight (g)    | 360±6    | 391±7*    |
| Heart weight (%)   | 0.34±0.02| 0.34±0.01 |
| Lung weight (%)    | 0.40±0.02| 0.39±0.01 |
| Liver weight (%)   | 3.54±0.12| 3.34±0.31 |
| Spleen weight (%)  | 0.22±0.01| 0.22±0.02 |
| Left adrenal gland (%) | 0.007±0.001 | 0.006±0.0004 |
| Brain weight (%)   | 0.56±0.01| 0.52±0.01*|
| Left ventricular ejection fraction (%) | 89±1 | 90±1 |
| Heart rate (beats/min) | 396±21 | 379±9 |

Neuroinflammation in the hippocampus, measured as microglia activity in the CA1 and hilus, was not significantly affected one week after ISO treatment (CA1 area: saline 5.3±0.5%, ISO 5.3±0.3%; Hilus: saline 14.7±1.9%, ISO 11.3±0.7%). Neither did any of the other microglia morphology parameters, including density, coverage, cell size, cell body size and processes size, change.
Effects Of Exercise In Iso Rats

General

From 35 male rats injected with ISO, 15 rats died within the first days. No mortality was observed in 20 saline treated rats, resulting in 10 rats in each of the experimental groups; saline sedentary; saline runner; ISO sedentary; ISO runner. General characteristics of the experimental groups are summarized in Table 2. Two-way ANOVA revealed a significant effect of ISO versus saline for heart weight, but no effect of running, nor interaction between saline/ISO and sedentary/runner.

|                       | saline sedentary | saline runner | ISO sedentary | ISO runner |
|-----------------------|------------------|---------------|---------------|------------|
| N                     | N=10             | N=10          | N=10          | N=10       |
| Body weight start (g) | 448±19           | 442±11        | 430±12        | 426±8      |
| Body weight end (g)   | 438±22           | 437±15        | 430±13        | 420±7      |
| Heart weight (% of body weight) | 0.30±0.01 | 0.28±0.01 | 0.32±0.01* | 0.32±0.01* |
| Brain weight (% of body weight) | 0.50±0.02 | 0.49±0.02 | 0.49±0.01 | 0.49±0.01 |

Cardiac collagen levels, as measure for ISO-induced focal infarcts, were significantly increased at the apex of the heart (Figure 2). Two-way ANOVA revealed a significant effect of ISO, which appear most pronounced in runners. No effect of sedentary versus running, nor interaction effects were observed. Apical collagen, but not mid-level collagen, was significantly correlated to heart weight (r=0.55, p<0.000).

Neuroinflammation

Neuroinflammation was obtained from morphologic changes in microglia in the CA1 and Hilus area of the hippocampus. No significant differences were observed in microglia parameters in the CA1 area (Table 3). In the Hilus (see figure 3), ISO decreased the size of the microglia by decreasing the area covered by processes. This did not result in increased microglia activity measures, although cell body size seemed unaffected. The reduced microglia size was not compensated for by increased density, resulting in reduced coverage (Figure 3). No correlations were observed between open field behavior and Hilus microglia parameters. Running in saline treated rats, had similar effects on microglia morphology as had ISO. However, running in isoproterenol-treated rats, may partly reverse the ISO-induced declined microglia coverage.
Table 3
Parameters of microglia morphology in the CA1 area of the hippocampus in saline and isoproterenol (ISO) treated male rats, and effects of exercise (runner) rats versus sedentary conditions.

|                          | saline sedentary | saline runner | ISO sedentary | ISO runner |
|--------------------------|------------------|--------------|--------------|------------|
| N=10                     | N=10             | N=10         | N=10         | N=10       |
| Density (# cells/area)   | 4.86±0.21        | 4.79±0.10    | 4.74±0.19    | 4.86±0.21  |
| Coverage (% area)        | 9.9±0.6          | 9.9±0.7      | 8.4±0.4      | 10.6±0.9   |
| Cell size (pixel)        | 2102±151         | 2146±187     | 1825±71      | 2268±255   |
| Cell body size (pixel)   | 182±7            | 177±5        | 181±10       | 184±8      |
| Processes size (pixel)   | 1919±154         | 1969±188     | 1644±73      | 2085±255   |
| Activity (cell body/cell size; %) | 9.6±0.9 | 9.0±0.7     | 10.7±0.8    | 9.3±1.0    |

**Brain function**

Since the hippocampus is mostly involved in learning and memory, function of the brain was estimated by Brain Derived Neurotrophic Factor (BDNF) expression in the hippocampus. In sedentary animals, no effects of ISO were observed in any of the areas. Running exercise was not affecting BDNF in saline treated rats. However, in all areas BDNF expression in ISO treated rats with exercise was slightly higher than in saline treated rats, reaching statistical significance in the CA1 and Hilus areas (Figure 4).

**Behavior**

General exploratory behavior was obtained from the open field test. Anxiety/depression levels were considered as more time spent at the wall areas rather than in the center of the open field. Two-way ANOVA revealed no significant effect of either saline/ISO or sedentary/running alone, but a significant interaction between both factors, resulting in post-hoc analysis of significantly decreased time spent at the wall area after ISO, which was reversed by running (Figure 5).

Cognitive behavior was tested in the Novel Object Recognition (NOR) and the Novel Location Recognition (NLR) tests. Figure 6 shows the results of these tests. In the NOR test, all groups performed significantly above chance level (=50%). Running may slightly improve performance in saline treated rats, but not in ISO treated rats. Although in the NLR test only saline runners performed above chance level, similar effects as seen in the NOR test were observed.

**Discussion**

**General**

Aim of the present study was to investigate the effects of five weeks exercise training on hippocampal neuroinflammation, neuronal function and behavior after established ISO-induced cardiac damage. At the start of exercise training, one week after ISO, cardiac damage seemed eminent, without declined cardiac
function or neuroinflammation (microglia activity). Although long-term cardiac damage after ISO appeared substantial, it did not induce cognitive impairment nor associated changes in microglia or BDNF expression in hippocampal CA1. However, long-term effects of ISO suggested altered microglia morphology in the Hilus and reduced anxiety in the OF. Five weeks of running reversed the reduced anxiety, but did not affect cognition or neuroinflammation in ISO rats. However, running treatment after ISO-induced cardiac damage may increase brain function, measured by BDNF expression.

The model

Acute sympathetic stress is associated with overactivation of beta-adrenergic receptors, leading to cardiac damage and ultimately cardiac dysfunction and heart failure [1,2,15]. However, acute stress is often caused by unexpected events and therefore difficult to predict [1]. Acute administration of high dose of the beta-receptor agonist isoproterenol (ISO) is commonly used to mimic this clinical condition [15]. Although interventions in this model were mostly aimed at prevention of the cardiac damage, regarding the unpredictable character of the clinical condition, it seemed worthwhile to explore interventions that could be applied after ISO had exerted its effect on the heart, and cardiac fibrosis was eminent.

Interventions that target consequences of cardiac damage were mostly investigated in the more labor-intensive and technical skills requiring model of coronary artery ligation [16]. Moreover, our previous study [12] indicated a mixed effect of the surgical procedure for coronary artery ligation and the effect of the subsequent introduced cardiac damage, on cognition and (neuro)inflammation. The model of ISO-induced cardiac damage circumvented these additional surgery-induced effects.

Effects of ISO

ISO, administered twice with 24h in between, evoked inflammation and cytokine production and cardiac fibrosis [1], that over a period of weeks developed into left ventricular hypertrophy and dilatation, and ultimately heart failure [4]. Effects of ISO appeared age dependent [17,18]. Accordingly, in the present study mortality in the adult (12 months old) animals was substantially higher than in the young (3 months old) rats. Cardiac damage and scar formation one week after ISO administration in young rats indeed implicated activation of the innate immune response necessary for wound healing. However, this had not led to declined cardiac function yet. Up to 1 week after injections, blood pressure, cardiac output and work were in the normal range under basal conditions but maximum cardiac output and work were well below normal [2]. Normal heart weight and tissue coverage (microscopical sections) suggested cardiac myocyte hypertrophy to compensate for the loss of viable myocardium; compensated cardiac hypertrophy. Absence of increased lung weight supported absence of overt heart failure. Moreover, microglia morphology was not altered, indicating no signs of neuroinflammation yet. Six weeks after ISO in 12 months old rats, cardiac fibrosis was still present, though less pronounced than in 3 months old rats one week after ISO. This would be in agreement with the old study of Beznak et al. [2]; during the first few weeks following ISO injection, histological repair took place, and the cardiovascular parameters measured were not different from those found in naive rats [2]. Although direct effects of beta-adrenergic stimulation on behavior have been studied; long-term effects
of this acute beta stimulation on neuroinflammation and behavior are less well known [5]. Still, ISO seemed to reduce exploratory behavior and the autonomic emotional state [5] and induced cognitive decline [6]. In the present study, no cognitive effects of ISO were observed. Since the used cognitive tests were mainly aimed at hippocampal function, accordingly microglia morphology was analyzed in this brain structure. In accordance with the lack of cognitive effects, no microglia activity was observed in the CA1 area of the hippocampus, neither was brain function, as measured by hippocampal BDNF expression, affected by ISO. Surprisingly, ISO seemed to have reduced anxiety, as measured by less time spent at the wall area in the open field. This would be in contrast to our previous findings in coronary artery ligation rats [19]. The effect on open field behavior coincided with altered microglia morphology in the hippocampal Hilus area. This would be in agreement with our study in coronary artery ischemia-reperfusion induced cardiac damage, showing microglia activation in the Hilus, but not in the CA1 area [12]. However, in the previously mentioned study declined cognitive performance in the NLR test was observed, which was not seen in the current investigation. An explanation could be that for the present study at 12 months of age even (saline treated sedentary) control rats did not recognize the relocated object. Therefore, the effects of ISO to mimic the consequences of acute sympathetic stress-induced cardiac damage, seemed in general agreement with literature, and showed similarities with the consequences of myocardial infarction-induced by coronary artery ligation in our hands.

Effects of exercise

Exercise training is generally acknowledged for having positive effect on physical as well as mental conditions. Moreover, exercise is known to have anti-inflammatory effects [10]. The potentially anti-inflammatory effect of exercise could counteract (neuro)inflammation and thereby improve cognition and mood [20]. ISO induced an inflammatory response for healing and focal cardiac scar formation. Therefore, in the present study, the effects of exercise training in the ISO model were investigated. Exercise did not interfere with cardiac fibrosis in ISO-treated rats. Heart weight was significantly increased in the combined ISO and exercise rats. Exercise reversed the ISO effect in open field behavior by normalizing anxiety. Exercise tended to improve cognitive performance in saline treated rats. However, exercise could not improve cognitive performance in ISO rats. In fact, only saline rats that had exercise training performed above chance level in the NLR test. Exercise-induced stimulation of brain BDNF is well-known [21]. Accordingly, in the present study, exercise increased BDNF expression in both the CA1 as well as hilus area of the hippocampus, but only in ISO-treated rats. Morphological changes indicated a reduction of microglia size, mainly due to lower processes area. Although not reflected in increased microglia activity, measured as cell body size/cell size [13], a lower processes area would indicate de-ramification of the microglia, which is usually associated with activation [22]. Exercise may partly reverse this microglia activation in ISO rats. Although exercise is often performed as prevention rather than reversal of effects of ISO [1,7], or coronary ligation-induced MI [23], effects are generally going in the same direction as seen in the present study. Interestingly, exercise training is reported to affect glycogen metabolism in skeletal muscles [24], and in the brain glycogen metabolism was associated with neuroplasticity [25]. The Hilus appeared to be the most glycogen-rich subregion of the hippocampus [26].
Accordingly, in the present study, most pronounced effects of exercise in the brain were observed in the Hilus area of the hippocampus.

Limitations

Each study has its limitations. The first part of the study was aimed at developing the ISO model in our lab. For that, young male Wistar rats were used. Based on the outcome of established cardiac damage one week after ISO treatment, in the subsequent study, the intervention of exercise training was started from one week onwards. This latter study was performed in 12 months old male Wistars as adult rats. Since the effects of ISO seemed age dependent, we chose for adult rats, to mimic the population of patients experiencing acute sympathetic stress. Indeed, mortality in these adult rats appeared higher than in the young rats. However, whereas the young rats were studied in Groningen and the adult rats in Budapest, differences in facility condition may have contributed as well.

Moreover, timing of the exercise intervention, from 1-6 weeks after ISO treatment, was chosen carefully regarding the aim of exercise intervention; treatment rather than prevention of an unpredictable event. Other starting time as well as duration could have provided different results.

No blood samples have been collected from these rats to evaluate levels of circulating inflammatory markers, which could have helped to elucidate on the underlying mechanisms.

Finally, since no echocardiography equipment was available in the lab in Budapest, effects of exercise training on cardiac function could not be obtained. Although cardiac damage, as percentage fibrosis was not altered by exercise, the increase in heart weight in the ISO+exercise group could have pointed at changes in cardiac function.

Conclusion

Aim of the present study was to investigate the effects of 5 weeks exercise training on neuroinflammation, neuronal function and behavior after established ISO-induced cardiac damage. Although cardiac damage after ISO appeared substantial, it did not induce long-term cognitive impairment nor associated changes in the hippocampal CA1 area. Accordingly, exercise training did not affect these aspects. Surprisingly, ISO seemed to reduce anxiety, which was reversed by exercise. ISO-induced microglia activation in the Hilus was not significantly affected by exercise, but exercise in ISO-treated rats showed increased brain function. In conclusion, five weeks of exercise, starting one week after ISO treatment, did not affect ISO-associated cardiac damage, cognition or neuroinflammation, but normalized the reduced anxiety shown in the open field. Increased localized BDNF expression may indicate improved brain function. Additional behavioral tests, and measurement of circulating inflammatory markers in a follow-up study may further elucidate on the underlying mechanisms.

Abbreviations

BDNF
brain-derived neurotrophic factor
CA
Cornu Ammonis
EF
ejection fraction
ISO
isoproterenol
Iba-1
ionized-binding adaptor protein 1
LSD
Least square difference
NLR
novel location recognition test
NOR
novel object recognition test
OF
open field
SEM
standard error of the mean
VO$_2$ max
maximal oxygen uptake capacity

Declarations

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Funding information
No external funding was obtained.

Compliance with ethical standards

Human subjects/Informed consent
No experiments on humans were carried out

Ethical approval of animal studies
The animal experiments were approved by the animal committee of the University Groningen, the Netherlands and the animal committee of the University of Physical Education, Budapest, Hungary.

Conflict of interest
All authors declare that they have no conflict of interest
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**Figures**
Figure 1

Photographs of Sirius Red/Fast green stained apical heart slices from rats treated with saline or isoproterenol (ISO), showing more collagen (red) in ISO treated hearts. Lower panel shows actual measurement of percentage collagen 1 week after ISO or saline treatment. *: significant effect of ISO
Figure 2

Collagen positive area in the left ventricle of the heart, at mid ventricular level and apex, in saline and isoproterenol (ISO) treated rats, and effects of exercise (runner) versus sedentary conditions. *: significant difference between saline and ISO.
Figure 3

Microglia morphology parameters obtained from the Hilus of the hippocampus in saline and isoproterenol (ISO) treated rats, and effects of exercise (runner) rats versus sedentary conditions. ISO: isoproterenol treatment; *: significant difference between indicated groups (n=9-10 per group)
Figure 4

Brain Derived Neurotrophic Factor (BDNF) expression in the different regions of the hippocampus in saline and isoproterenol (ISO) treated rats, and effects of exercise (runner) rats versus sedentary conditions. *: significant difference between saline runners and ISO runners.
Figure 5

Time spent in the wall area of the open field in saline and isoproterenol (ISO) treated rat, and effects of exercise (runner) versus sedentary conditions. *: significant difference between indicated groups.

Figure 6
Cognitive effects of isoproterenol (ISO) versus saline treatment, and the effects of running versus sedentary conditions. *: significant difference between indicated groups; # significantly different from chance level (=50%; dashed line).