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Decreased integrity of exercise-induced plasma cell free nuclear DNA – negative association with the increased oxidants production by circulating phagocytes

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Strenuous exercise increases circulating cell free DNA (cfDNA) and stimulates blood phagocytes to generate reactive oxygen species (ROS) which may induce DNA strand breaks. We tested whether: (A) elevated cfDNA in response to three repeated bouts of exhaustive exercise has decreased integrity; (B) each bout of exercise increases luminol enhanced whole blood chemiluminescence (LBCL) as a measure of ROS production by polymorphonuclear leukocytes. Eleven men performed three treadmill exercise tests to exhaustion separated by 72 hours of resting. Pre- and post-exercise concentrations and integrity of cf nuclear and mitochondrial DNA (cf n-DNA, cf mt-DNA) and resting (r) and fMLP (n-formyl-methionyl-leucyl-phenylalanine)-stimulated LBCL were determined. Each bout increased concentrations of cf n-DNA by more than 10-times which was accompanied by about 2-times elevated post-exercise rLBCL and fMLP-LBCL. Post-exercise cf n-DNA integrity (integrity index, $I_{229/97}$) decreased after the first (0.59 ± 0.19 vs. 0.48 ± 0.18) and second (0.53 ± 0.14 vs. 0.44 ± 0.17) bout of exercise. There were negative correlations between $I_{229/97}$ and rLBCL ($q = -0.37$), and $I_{229/97}$ and fMLP-LBCL ($q = -0.40$) – analysis of pooled pre- and post-exercise data ($n = 66$). cf mt-DNA integrity ($I_{218/78}$) did not alter in response to exercise. This suggests an involvement of phagocyte ROS in cf n-DNA strand breaks in response to exhaustive exercise.

There are rising evidences suggesting that besides, health promoting effects, exhaustive exercises might have potential adverse effects on immune system^{1,2}. Therefore, explanation of this issue is essential for training recommendations or customization of individual training load. Interestingly, bouts of strenuous exercise caused rapid increase in cell free DNA (cfDNA) concentration in plasma and, moreover, it was suggested to be a promising marker of acute exercise induced-metabolic changes in human body³. Exercise resulted in the surge of plasma cfDNA independently of workout associated with various sport disciplines and types of training such as weight-lifting⁴, running⁵, soccer⁶, cycling⁷, rowing⁸, strength training⁹ or repeated sprint⁶. Concentration of post-exercise cfDNA positively correlated with the duration and the intensity of aerobic running¹⁰ as well as with selected markers of muscle damage¹¹. Moreover, the ratio of post- to pre-exercise cfDNA associated with the distance covered by soccer players during the game⁶. In another study, increased cfDNA in response to heavy resistance exercise was described to have possible predictive value of muscle-performance decrease within 2 days after the bout¹². Therefore, measurements of exercise-related variations of cfDNA can be used for monitoring of training load or diagnosis and prevention of overtraining syndrome¹¹.

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DNA Integrity	1 st bout (day 7)		2 nd bout (day 10)		3 rd bout (day 13)	
	Before	After	Before	After	Before	After
cf n-DNA (I _{229/97})	0.59 ± 0.19 (0.57; 0.36)	0.48 ± 0.18* (0.46; 0.36)	0.53 ± 0.14 (0.55; 0.13)	0.44 ± 0.17* (0.46; 0.33)	0.50 ± 0.22 (0.38; 0.36)	0.45 ± 0.17 (0.41; 0.22)
cf mt-DNA (I _{218/78})	0.54 ± 0.16 (0.59; 0.28)	0.60 ± 0.26 (0.69; 0.39)	0.63 ± 0.18 (0.53; 0.33)	0.67 ± 0.16 (0.63; 0.30)	0.65 ± 0.18 (0.67; 0.16)	0.66 ± 0.20 (0.79; 0.40)

Table 1. Cell free nuclear (cf n-DNA) and mitochondrial (cf mt-DNA) DNA integrity in eleven average-trained men before and after each of three bouts of exhaustive treadmill exercise. Eleven average-trained men completed the study composed of four visits at days 1, 7, 10 and 13. After determination of VO₂max at day 1, three repeated treadmill exercise tests to exhaustion at speed corresponding to 70% of personal VO₂max were executed at days 7, 10 and 13. Results are expressed as mean ± SD (median; IQR). * vs corresponding value before the bout, $p < 0.05$ (for exact p-values please see Table 9, supplementary file).

The common process following exhaustive exercises is leukocyte demargination from the vascular, hepatic, pulmonary or spleen reservoirs in parallel with exercise-induced increase in cardiac output and blood flow. This elevates the white blood cell count, however, it seems to be not the main contributor to the exercise-induced increase in cfDNA because the rise in the latter is many times higher than post-exercise leukocytosis³. Therefore, mechanisms responsible for this phenomenon could include at least the elevated number of circulating leukocytes accompanied by the increase in their activity leading to the release of a variety of biomolecules including DNA³. This is in line with the recent observation of increased formation of intravascular neutrophil extracellular traps (NETs) that could be responsible for the rise in cfDNA in subjects after exhaustive exercise^{13,14}.

NETs formation involves disintegration of nuclear and granular membranes, diffusion of decondensed chromatin into the cytoplasm and its mixing with various proteins. After rupture of neutrophil plasma membrane, this chromatin associated with granular proteins such as myeloperoxidase, neutrophil elastase or cathepsin G is released into the extracellular space^{15,16} leading to the increase in plasma cf n-DNA.

Respiratory burst of circulating phagocytes (mainly polymorphonuclear – PMNs) in combination with phagocytosis and NETs formation is essential for innate immunity^{17,18}. Increased generation of reactive oxygen species (ROS) related to activation of circulating PMNs NADPH oxidase complex was reported in healthy subjects after exhaustive exercise^{19,20}.

Luminol-enhanced whole blood chemiluminescence technique (LBCL), which reflects mainly superoxide radicals production by circulating PMNs^{21,22}, was used for evaluation of pre- and post-exercise ROS generation^{23–25}. This is a rapid test which due to its low day-to-day variability and relative simplicity is suitable in epidemiological and clinical studies²⁶. It can be executed in two settings: (a) – blood samples without stimulation – resting LBCL (rLBCL) and (b) – blood samples after stimulation with n-formyl-methionyl-leucyl-phenylalanine (fMLP), a chemotactic peptide which induces PMNs respiratory burst via activation of NADPH oxidase²⁶ – fMLP-stimulated LBCL (fMLP-LBCL). Both rLBCL and fMLP-LBCL were increased in patients suffering from diseases accompanied by systemic inflammatory response such as systemic sclerosis, infective endocarditis and chronic renal failure^{27–29}. Moreover, LBCL was attenuated in healthy subjects in response to regular consumption of fruits rich in anti-oxidant and anti-inflammatory polyphenols^{30,31}.

ROS can induce damage to a variety of biomolecules including modification of DNA bases and DNA strand breaks. It is possible that simultaneous activation of respiratory burst of circulating PMNs and NETs formation in response to exhaustive exercise lead to strand breaks of released DNA and thus formation of cfDNA with decreased integrity. cfDNA can be divided in two pools: cell free nuclear DNA (cf n-DNA) and cell free mitochondrial DNA (cf mt-DNA), deriving from nucleus or cytoplasmic mitochondria's, respectively. Because of lack of nucleosomal structure, cf mt-DNA fragments present different pattern of integrity than cf n-DNA³². Moreover, mt-DNA seems to be more sensitive to oxidative stress and other genotoxic damages due to the lack of protective proteins and efficient DNA repair mechanism³³. Therefore, it seems that integrity of cf mt-DNA could be more affected by ROS increase following exhaustive exercise than that of cf n-DNA. To solve this question, we compared the pre- and post-exercise plasma concentrations and integrity of cf n-DNA and cf mt-DNA in relation to rLBCL and fMLP-LBCL (reflecting ROS production by circulating PMNs) in healthy physically active men subjected to three repeated exhaustive treadmill runs.

Results

All included men successfully completed the study protocol of three repeated exhaustive treadmill exercises. Mean concentration of plasma cf n-DNA increased 11.3-, 11.8- and 17.3- times ($p < 0.05$) after the 1st, 2nd, and 3rd exercise, while cf mt-DNA increased significantly by about 2- and 2.4-times only after the 2nd and 3rd exhaustive treadmill run, respectively. Data of plasma concentrations of cf nDNA and cf mt DNA, other parameters related to exercise tests, changes of selected markers of muscle damage as well as metabolic and cardiovascular responses to exercise have been described elsewhere³.

Changes of integrity of circulating cell free nuclear and mitochondrial DNA in response to three repeated bouts of exhaustive treadmill exercise. Pre-exercise integrity of cf n-DNA expressed as I_{229/97} did not alter significantly over the study period. However, the tendency to a gradual decrease in I_{229/97} was noted e.g. I_{229/97} = 0.59 ± 0.19 before the 1st bout in comparison to I_{229/97} = 0.50 ± 0.22 before the 3rd one (Table 1). Exhaustive exercise resulted in the decrease in cf n-DNA integrity especially after the 1st (pre-exercise I_{229/97} = 0.59 ± 0.19 vs post-exercise I_{229/97} = 0.48 ± 0.18, $p < 0.05$) and the 2nd bout (pre-exercise

Blood Chemiluminescence	1 st bout (day 7)		2 nd bout (day 10)		3 rd bout (day 13)	
	Before	After	Before	After	Before	After
rLBCL	326 ± 269 (196; 479)	697 ± 302* (786; 431)	335 ± 244 (280; 551)	870 ± 663* (728; 1056)	319 ± 261 (274; 528)	765 ± 610* (577; 1125)
fMLP-LBCL	892 ± 435 (941; 561)	1531 ± 607* (1575; 1014)	909 ± 375 (885; 692)	1598 ± 737* (1613; 1192)	960 ± 525 (802; 746)	1470 ± 907* (951; 1442)

Table 2. Resting (rLBCL) and fMLP-stimulated luminol enhanced whole blood chemiluminescence (fMLP-LBCL) in eleven average-trained men before and after each of three bouts of exhaustive treadmill exercise. Results expressed in relative light units per 10^4 phagocytes present in the assayed sample are shown as mean ± SD (median; IQR). * vs corresponding value before the bout, $p < 0.05$ (for exact p-values please see Table 10, supplementary file). Other details as for Table 1.

Correlated variables	rLBCL	fMLP-LBCL	Explanation
Integrity of cf n-DNA ($I_{229/97}$)	-0.29	-0.36*	Pooled individual pre-exercise data from three bouts (n = 33)
Integrity of cf n-DNA ($I_{229/97}$)	-0.31 [#]	-0.32 [†]	Pooled individual post-exercise data from three bouts (n = 33)
Integrity of cf n-DNA ($I_{229/97}$)	-0.37*	-0.40*	Pooled individual pre- and post-exercise data from three bouts (n = 66)

Table 3. Spearman's (ρ) correlations between integrity of cell free nuclear DNA expressed as $I_{229/97}$ and resting and fMLP-induced luminol enhanced whole blood chemiluminescence in eleven average trained healthy men before and after three repeated bouts of exhaustive treadmill exercise. cf n-DNA – cell free nuclear DNA, rLBCL – resting luminol enhanced whole blood chemiluminescence, fMLP-LBCL – stimulated luminol enhanced whole blood chemiluminescence. Other details as for Table 1. * - $p < 0.05$, # – borderline significance – $p = 0.08$, † – borderline significance – $p = 0.06$. Further details are shown in Figures 2, 3, 4, 5, 6 and 7 of supplementary file.

$I_{229/97} = 0.53 \pm 0.14$ vs post-exercise $I_{229/97} = 0.44 \pm 0.17$, $p < 0.05$). In contrast to cf n-DNA integrity, the pre- and post-exercise-integrity of cf mt-DNA ($I_{218/78}$) was relatively stable over the study period and did not alter in response to exercise (Table 1).

Changes of resting and fMLP-stimulated luminol enhanced whole blood chemiluminescence in response to three repeated bouts of exhaustive treadmill exercise. Mean rLBCL increased by about 2.1-, 2.6- and 2.4-times ($p < 0.05$) in response to the 1st, 2nd and the 3rd bout of exhaustive exercise (Table 2). The values of pre-exercise rLBCL were relatively stable over the study period and the same behavior was noted for the post-rLBCL. Similarly, behaved mean fMLP-LBCL: 1.7-, 1.8- and 1.5-fold increase ($p < 0.05$) after each exhaustive treadmill run (Table 2). Mutual comparisons of pre-exercise fMLP-LBCL values at consecutive days did not reveal any significant differences, as well for post-exercise fMLP-LBCL values. Consequently, the mean ratio of pre-exercise fMLP-LBCL to pre-exercise rLBCL values was similar and ranged from 2.7 at the 1st bout to 3.0 at the 3rd bout. The mean ratio of post-exercise fMLP-LBCL to post-exercise rLBCL was also stable and reached 2.2, 1.8 and 1.9 after the 1st, 2nd and 3rd bout of exercise, respectively.

Correlations between integrity of cf n-DNA and whole blood chemiluminescence before and after three repeated bouts of exhaustive exercise. Because we studied a relatively low group of men ($n = 11$), the direct analysis of correlations between variables obtained during one bout was inconclusive. Therefore, to overcome (at least partially) this problem, we analyzed Spearman's (ρ) correlations between integrity of cf n-DNA ($I_{229/97}$) and LBCL by using the following datasets from three bouts of exhaustive exercise: (A)- pooled individual pre-exercise data ($n = 33$); (B)- pooled individual post-exercise data ($n = 33$); and (C)- pooled pre- and post-exercise data ($n = 66$) (Table 3). There was a negative correlation of moderate strength between pre-exercise $I_{229/97}$ and pre-exercise fMLP-LBCL ($\rho = -0.36$, $p < 0.05$), while correlation between post-exercise $I_{229/97}$ and post-exercise rLBCL or fMLP-LBCL reached borderline significance ($\rho = -0.31$, $p = 0.08$ and $\rho = -0.32$, $p = 0.06$). In the case of analysis of dataset involving pre- and post-exercise data together, $I_{229/97}$ negatively correlated with rLBCL ($\rho = -0.37$, $p < 0.05$) and fMLP-LBCL ($\rho = -0.40$, $p < 0.05$) (Table 3). The same analyses performed between integrity of cf mt-DNA ($I_{218/78}$) and whole blood chemiluminescence (rLBCL, fMLP-LBCL) revealed no significant correlations (ρ ranged between 0.11 and 0.38).

Discussion

It is generally believed that regular physical activity has important health promoting effect. However, long term, repeated exhaustive exercises can evoke pathological reactions leading to the development of overtraining syndrome^{34–36}. High intensity exercise related to various sports disciplines^{4,5} was reported to cause fast, transient increases in circulating cfDNA reaching concentrations comparable to those observed in trauma or sepsis^{37,38}. Some studies suggest that elevated plasma cfDNA levels may be associated with the increased risk of occurrence of overtraining syndrome in athletes^{39,40}. Recently, we found that three repeated bouts of exhaustive treadmill exercise induced increase in cf n-DNA and cf mt-DNA in average trained healthy men without development of tolerance³. In the present study (which is the continuation of the afore-mentioned one), we found that the increase in cf n-DNA in response to each of three repeated bouts of exhaustive treadmill exercise is accompanied

by the decrease in cf n-DNA integrity. In parallel to this phenomenon, each bout of exercise caused significant increase in rLBCL and fMLP-LBCL and that reflects an increased post-exercise ROS production by circulating phagocytes, namely PMNs. Moreover, analysis of pooled data from all bouts revealed a few significant negative correlations between integrity index of cf n-DNA and LBCL. These considerations suggest the involvement of ROS generated from circulating phagocytes in fragmentation of post-exercise cf n-DNA. Although, cf mt-DNA raised in response to the 2nd and 3rd bout of exercise, very surprisingly, no changes of the integrity index of cf mt-DNA were noted over the study period.

Effect of three repeated bouts of exhaustive treadmill exercise on luminol enhanced whole blood chemiluminescence.

Each bout of exercise increased mean rLBCL and fMLP-LBCL in healthy men. Moreover, individual results revealed substantial increase in post-exercise rLBCL and fMLP-LBCL in each volunteer at any occasion. These observations indicate that exhaustive run increased basic generation of ROS by circulating PMNs and primed these cells for increased ROS production after stimulation with fMLP. Our results are consistent with the previous studies showing increased zymosan-induced luminol enhanced chemiluminescence of neutrophils isolated from blood collected after intensive aerobic exercise in healthy men^{23–25}.

Moreover, neutrophils of subjects who completed 100 km marathon run revealed increased ROS production as reflected by the afore-mentioned method⁴¹. On the other hand, other studies did not confirm these observations and even reported suppression of ROS production by neutrophils in the group of well-trained men after marathon run or after acute bout of moderate intensity running and cycling using the same stimulator of phagocyte oxidative burst^{42,43}. These differences may result from various study protocols, including exercise load and duration, time of blood collection, usage of whole blood or isolated cells for determination of ROS generation after stimulation with various cell activators. For instance, a continuous 90 min. exercise at the intensity of 50% VO₂max caused an increase of phorbol 12-myristate 13-acetate (PMA)-induced neutrophils chemiluminescence while zymosan-induced chemiluminescence remained unchanged in healthy men⁴⁴.

PMA is a direct activator of protein kinase C with following activation of NADPH oxidase and neutrophils oxidative burst while stimulation with opsonized zymosan (insoluble cell-wall preparation from the fungi *Saccharomyces cerevisiae*) involves phagocytosis of these particles, cell degranulation and massive production of ROS⁴⁵. In our study, we used fMLP (an analog of N-formylated bacterial chemotactic peptides). In our study, we used fMLP (an analog of N-formylated bacterial chemotactic peptides). This peptide activates signal transduction pathways (phospholipase C-dependent generation of diacyl glycerol and inositol 1,4,5-triphosphate, rise in the intracellular Ca²⁺ concentration, protein kinase C activation) upon binding to specific G protein-coupled receptors on the PMNs plasma membrane. This, in turn, leads to the formation of the active NADPH oxidase complex and ROS production⁴⁵. Thus, the peak time – time from agonist addition to appearance of maximal chemiluminescence – is about 7-times shorter for fMLP than for opsonized zymosan^{23,46}. This remark may explain the discrepancies between results of our study and some previous investigations^{42,43} about the effect of exercise on ROS production by isolated neutrophils after stimulation with opsonized zymosan.

Moreover, we studied the effect of exercise on LBCL, thus neutrophils and monocytes were not subjected to any process of isolation that could change the cell ability to respond to stimulations with various agonists. Elimination of the risk of cell priming or de-priming during the isolation procedure is the important advantage of this method^{21,31}. Such technique allows monitoring of oxidants release by circulating phagocytes under conditions that more closely resemble the *in vivo* situation than the analysis of isolated cells^{31,47}. Severe exercise resulted in essential increase in plasma pro-inflammatory cytokines, namely IL-6, IL-8, granulocyte colony-stimulating factor (G-CSF), macrophage colony-stimulating factor (M-CSF), and granulocyte macrophage colony stimulating factor (GM-CSF)^{48,49}. Although these pro-inflammatory cytokines alone induced rather weak oxidative response of PMNs, they strongly enhanced ROS generation in response to secondary stimulation with fMLP^{50–53}, and that could be an explanation of increased rLBCL and fMLP-LBCL in healthy men after exhaustive exercise.

Effect of repeated bouts of exhaustive treadmill exercise on the integrity of plasma cell free DNA.

Each bout of exhaustive exercise increased several times the concentration of circulating cf n-DNA, accompanied by the decrease in I_{229/97} of post-exercise cf n-DNA. This indicates that apool of cf n-DNA released in response to exhaustive exercise is subjected to factors leading to its enhanced fragmentation. Conversely, no changes in I_{218/78} were noted over the study period, although cf mt-DNA raised significantly after the 2nd and 3rd bout. Thus, one may conclude that exercise-induced release of cf mt-DNA was not accompanied by parallel processes resulting in its additional fragmentation. It is believed that the source of exercise-induced circulating cf n-DNA are NETs¹⁵. Various factors related to strenuous exercise including heat stress, catecholamines, pro-inflammatory cytokines (e.g IL-6, IL-8) can induce formation of NETs¹⁵. Intracellular generation of ROS by NADPH oxidase together with their elaboration by myeloperoxidase in intracellular granules are involved in triggering NETs formation⁵⁴. Thus, nuclear DNA mixed with some components of granules before expulsion of NET fibers from PMNs can be exposed to and damaged by ROS, including breaking of DNA strands into shorter pieces. This could be a plausible explanation of increased post-exercise plasma levels of cf n-DNA with decreased integrity and is in line with observed moderate negative correlations between I_{229/97} and rLBCL and fMLP-LBCL. Moreover, previous studies showing increased concentration of 8-oxo-7,8-dihydro-2-deoxyguanosine (an oxidized derivative of deoxyguanosine - marker of cellular oxidative stress and oxidative damage to DNA) in leukocytes and urine of athletes after intense exercise⁵⁵ support this explanation.

The exercise-induced rise in plasma cf n-DNA is transient^{8,56,57} probably due to simultaneous increase in the activity of circulating deoxyribonuclease I⁸. Thus, the post-exercise cf n-DNA decreased back to baseline within 0.5 to 2 hours of recovery⁸.

The bouts of exhaustive treadmill exercise were separated by 3 days of rest, and all healthy volunteers did not perform any additional strenuous exercise during the whole 13-day study period. Therefore, one may conclude

that the contribution of NETs in maintaining the pre-exercise cf n-DNA levels was low (and even negligible) and other processes with the unchanging intensities (e.g. necrosis, apoptosis) were the source of abaseline pool of circulating cf n-DNA. This may explain the relatively stable concentrations of pre-exercise cf n-DNA and its integrity over the study period. Circulating cf mt-DNA increased in response to exhaustive exercise at the same time as cf n-DNA. However, pre- and post-exercise $I_{218/78}$ of cf mt-DNA did not differ at any occasion. This is quite surprising considering the release of cf mt-DNA from NETs⁵⁸ and formation of NETs from pure mt-DNA in response to skeletal injuries and orthopedic surgery in humans⁵⁹. On the other hand, neutrophils infected with bacteria formed NETs with extracellular fibers containing n-DNA as the main structural component *in vitro*¹⁸. It cannot be excluded that neutrophils in response to exhaustive exercise extrude cf n-DNA alone or with very low admixture of cf mt-DNA. Thus, the exercise-induced increase in cf mt-DNA was about 5- to 6-times lower than that of cf n-DNA and no changes of cf mt-DNA integrity were noted.

Acute severe exercise on cycle ergometer until exhaustion increased mitochondrial ROS production in neutrophils of sedentary young males while the same physical exertion had no stimulatory effect on neutrophils mitochondrial ROS after 2 months of regular exercise training program in these subjects⁶⁰.

Aerobic strenuous exercise resulted in an increase in superoxide radical activity in contracting muscles. However, the contribution of muscle mitochondria to this increase was small^{61,62}, and even mitochondria generated less ROS during exercise than at rest⁶¹. We studied average trained healthy men who regularly performed recreational training where some of them participated in sport disciplines in the past. Therefore, it seems that exhaustive treadmill run did not induce increased generation of mitochondrial ROS and subsequent oxidative damage to mt-DNA.

Platelets are able to release microparticles containing functional mitochondria as well as free mitochondria^{63,64}. Circulating phospholipase A2 can digest mitochondrial membranes with subsequent release of mt-DNA into extracellular space⁶³. Exhaustive exercise (e.g. marathon run) can induce platelets activation and their degranulation⁶⁵. Additionally, short term lifestyle intervention (moderate energy restriction and aerobic training for 6 weeks) increased serum phospholipase A2 activity in overweight or obese subjects⁶⁶.

Consequently, it is possible that platelets could be another source of cf mt-DNA and encapsulating membranes can protect mt-DNA from oxidative attack. On the other hand, even mt-DNA released in response to exercise could be damaged by ROS originated from other sources (e.g. NADPH oxidase), and its amount seems to be too little to decrease the integrity of total post-exercise pool of circulating cf mt-DNA. These considerations may also explain no differences between pre- and post-exercise integrity of plasma cf mt-DNA.

There are very scant and unconvincing data on the integrity of exercise-induced cf DNA in humans. Incremental treadmill test until volitional exhaustion induced significant increase in circulating cf n-DNA but without changes of its integrity compared to pre-exercise cf n-DNA in male athletes⁶⁷. On the other hand, 10 km relay race resulted in a significant decrease in the integrity of post-exercise cf n-DNA in a group of recreational runners⁶⁸. Moreover, neither of these studies evaluated circulating concentrations and integrity of pre- and post-exercise cf mt-DNA. Our results showed that repeated exhaustive exercise decreased integrity of circulating cf n-DNA but did not change the integrity of cf mt-DNA.

Correlations between integrity of cf n-DNA and luminol enhanced whole blood chemiluminescence.

Because of the low number of studied volunteers, we pooled pre- and post-exercise data from three consecutive bouts of exhaustive treadmill run and calculated correlations between cf n-DNA integrity and rLBCL and fMLP-LBCL. Such approach has some limitations and can increase the risk of bias. Nonetheless, for instance, we found significant negative correlations between $I_{229/97}$ and rLBCL, and $I_{229/97}$ and fMLP-LBCL for pooled individual pre- and post-exercise data from three bouts. Moreover, all values of Spearman's ρ ($n = 6$) ranged between -0.40 to -0.29 and that suggests that there is some negative relationship between integrity of cf n-DNA and generation of ROS by circulating phagocytes.

However, according to values of ρ , these correlations can be interpreted as weak or moderate. This information suggests indirectly that other factors could be responsible for decreased integrity of post-exercise cf n-DNA, or post-exercise LBCL did not reflect precisely the generation of ROS by PMNs during the exhaustive run when NETs were formed and cf n-DNA released. Circulating cf n-DNA increased already after 9 min. from the onset of incremental treadmill run test (at the end of 10 km/h stage corresponding to 63% $\dot{V}O_{2,max}$)⁶⁸. In our study, mean run time was 47 min., 57 min. and 56 min. at the 1st, 2nd and 3rd bout, respectively³. Thus, pre- and post-exercise LBCL (rLBCL and fMLP-LBCL) reflected the activity of PMNs just before and after the exercise but not during the last 30 min. of run accompanied by the formation of NETs. Moreover, luminol crosses the cellular membrane of phagocytes (PMNs and monocytes); in this way, LBCL can mirror the extra- and intracellular production of ROS⁶⁹. Hence, additional blood sampling during exercise followed by measurements of LBCL and cf n-DNA integrity (to estimate the dynamics of changes) would give more valuable results of the analysis of associations between cf n-DNA fragmentation and ROS production by blood phagocytes.

It is possible that numerous plasma antioxidants can inactivate extracellular ROS (in the close vicinity of PMNs) before their reaction with n-DNA. In consequence, intracellular activity of ROS during formation of NETs seems to be crucial for the decrease of cf n-DNA integrity. These aforementioned reasons may explain the low strength of observed correlations between pooled data of cf n-DNA integrity and LBCL.

Acute exercise of different duration and intensity increased plasma antioxidant activity in men⁷⁰. Exhaustive cycling raised the activity of antioxidant enzymes e.g. superoxide dismutase and glutathione peroxidase in lymphocytes from peripheral blood⁷¹. Moreover, young athletes had higher Trolox-equivalent antioxidant capacity of plasma than their sedentary counterparts⁷². Thus, exercise-induced increase in circulating antioxidants (as adaptive response to muscle workout) may explain no effect of the third bout of exercise on cf n-DNA integrity and be an additional elucidation of weak correlations between post-exercise cf n-DNA integrity and LBCL.

It should be mentioned that one subject had no decrease in cf n-DNA integrity in response to each of the three bouts of exhaustive exercise (Table 5, supplementary file). Although, he presented an increase in post-exercise rLBCL and fMLP-LBCL (Tables 7 and 8, supplementary file) along with many-fold increase in post-exercise cf n-DNA³, the integrity indexes of pre- and post-exercise cf n-DNA were similar over the study period. It is possible that this volunteer had high activity of circulating and intracellular antioxidants which inactivated ROS before their reaction with n-DNA released from NETs. Circulating neutrophils are functionally a heterogeneous cell population^{16,73}. Certain neutrophil subpopulations could have increased capacity to form NETs and release cf n-DNA^{16,74}, and some of them can also release more ROS spontaneously or upon stimulation by various mediators^{75,76}. It cannot be excluded that in some persons these neutrophil subpopulations do not overlap, thus released cf n-DNA in response to exhaustive exercise would not be exposed to ROS and undergo subsequent fragmentation. Moreover, NETosis can occur without activation of NADPH oxidase and ROS generation¹⁶. These processes may also be a possible explanation of the afore-mentioned observations. Nevertheless, it is interesting that some subjects can increase cf n-DNA in response to exhaustive exercise without changes of its integrity.

Concomitant increase in the serum activity of DNase I was recognized as a factor responsible for transient nature of cf n-DNA response to strenuous exercise⁷⁷. Therefore, decreased integrity of post-exercise cf n-DNA could be the result of cleaving NETs DNA by this endonuclease. However, we did not observe any changes of post-exercise cf mt-DNA integrity. For that reason, the contribution of DNase I to the decreased integrity of post-exercise cf n-DNA seems to be not important under the conditions of our study. Increased post-exercise cf n-DNA and its integrity index as well as LBCL returned to baseline levels within 72 hour interval between bouts of exercise. Therefore, it is difficult to conclude about the clinical significance of these phenomena. Perhaps, in athletes with very high training- and competition-load, these changes may persist for longer time and be an additional marker of the risk of injury or over-training syndrome. However, it requires further more extensive studies.

Limitations of the study. Relatively low number of studied subjects ($n = 11$) and exclusion of female volunteers from the trial could be recognized as a weakness of the present study. Because this report is the extension of our previous one, all these afore-mentioned limitations were discussed in details elsewhere³. In addition, the inhibitory effect of progesterone and estradiol on ROS production by human PMNs *in vitro*⁷⁸ supported our decision on participation of only male volunteers in the study. The low number of studied subjects was counterbalanced by three repeated bouts of exercise. We found that all bouts induced a significant increase in cf n-DNA, rLBCL and fMLP-LBCL, while two bouts were accompanied by a decrease in cf n-DNA integrity and increase in circulating cf mt-DNA without changes of its integrity. This indicates the repeatability of these phenomena despite the small studied group of volunteers.

Conclusions

We found that repeated bouts of exhaustive exercise separated by three days of resting caused an increase in luminol enhanced whole blood chemiluminescence and in concentrations of circulating cf n-DNA. Post exercise cf n-DNA revealed decreased integrity which negatively correlated with LBCL. Because whole blood chemiluminescence reflects ROS production by circulating phagocytes, one may conclude that oxidants may be involved in the release of cf n-DNA and cf n-DNA strand breaks in response to exhaustive exercise. Although exercise caused moderate increase in plasma levels of cf mt-DNA, its integrity was stable and did not associate with blood chemiluminescence. This suggests a minor role of ROS in exercise-induced changes of cf mt-DNA. Further studies involving larger groups of male and female volunteers should be performed, especially bearing in mind the analysis of correlation and determination of causality between decreased of post-exercise cf n-DNA integrity and enhanced LBCL.

Methods

Studied group. The studied group of eleven, average-trained, non-smoking healthy men as well as the inclusion/exclusion criteria were the same as in our previous article³. Volunteers had mean age 34.0 ± 5.2 years, mean body mass index 26.2 ± 3.1 kg/m² and mean maximal oxygen consumption $VO_{2max} 49.6 \pm 4.5$ ml/kg x min. They were free of any pharmacological treatment including vitamins and food supplements for 3 months preceding the study. They signed informed consent and did not change their dietary habits during the study period.

The study design. The study design was described in details in our previous report³. Briefly, the study consisted of 4 visits at the 1st, 7th, 10th and 13th day of observation. At the first visit (day 1st) 11 average-trained men underwent treadmill VO_{2max} test and afterwards, at the three consecutive visits (day 7th, 10th and 13th), participants performed treadmill exercise to volitional exhaustion at speed corresponding to 70% of their personal VO_{2max} (Fig. 1). Venous blood (15 ml) was collected into vacutainer tubes with EDTA (Becton Dickinson, Franklin Lakes, NJ) within 5 min. before and after each bout of exercise. One milliliter thereof was placed into a separate tube for resting (rLBCL) and n-formyl-methionyl-leucyl-phenylalanine (fMLP) – stimulated LBCL as well as blood cell count (Micros Analyzer OT 45, ABX, Montpellier, France). The rest was used for determination of concentration and integrity of cell free nuclear (cf n-DNA) and mitochondrial DNA (cf mt-DNA). During the whole study period (13 days) volunteers did not perform any exhaustive exercise besides those related to the study protocol. The study was conducted according to the Declaration of Helsinki. The protocol was reviewed and approved by The Medical University of Lodz Ethics Committee (RNN/95/14/KB), and all volunteers provided a written informed consent.

Blood processing and measurement of cfDNA. EDTA blood samples were centrifuged ($1600 \times g$, 4 °C, 10 min.) immediately after collection. The obtained plasma samples were subjected to the second centrifugation ($16\,000 \times g$, 4 °C, 5 min.) to remove the cell debris and were stored at -80 °C for no longer than 4 weeks until

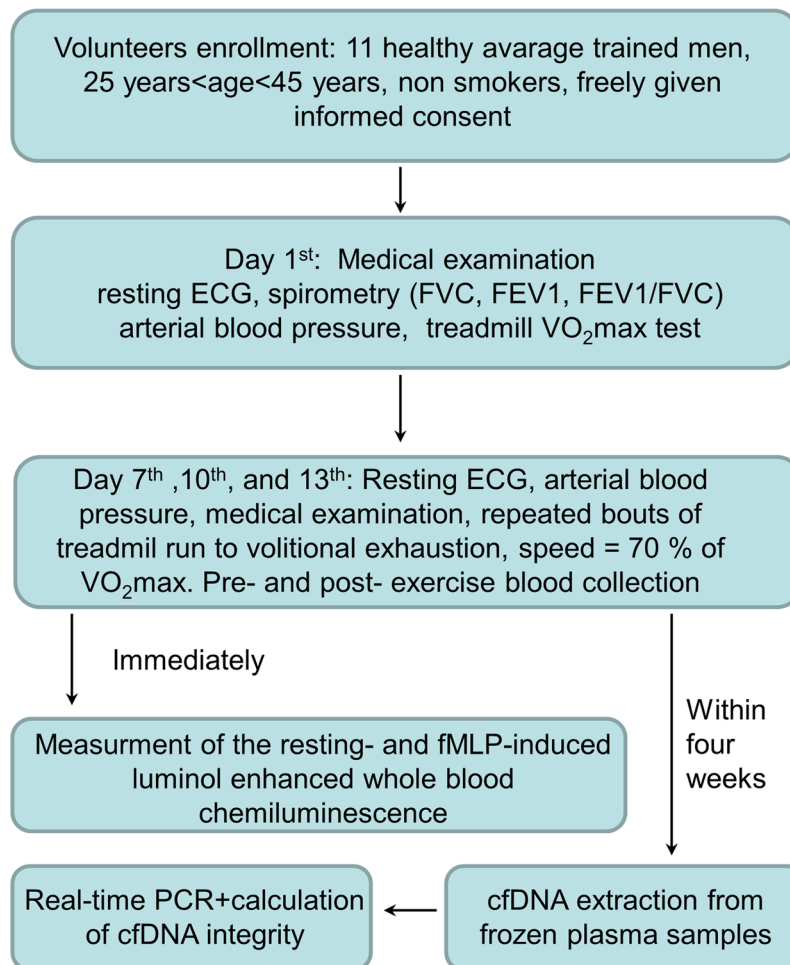


Figure 1. Overview of the study design. ECG -electrocardiography, FVC – forced vital capacity, FEV1 – forced exhaled volume in the first second, VO₂max – maximal oxygen consumption, fMLP – n-formyl-methionyl-leucyl-phenylalanine, cfDNA – cell free DNA, PCR – polymerase chain reaction. VO₂max was measured by a continuous incremental maximal exercise test using programmable treadmill with gas exchange analysis system and 12-lead wireless ECG at day 1st. Afterwards volunteers performed treadmill run to volitional exhaustion at speed corresponding to 70% of their personal VO₂ max at days 7th, 10th and 13th. Volitional exhaustion was defined as volunteer inability to maintain the required speed of running (exercise intensity) or its wish to stop the test.

isolation of cfDNA with QIAamp DNA Blood Mini Kit (Qiagen GmbH, Hilden, Germany) and measurement of plasma concentrations of cf n-DNA and cf m-DNA with real time quantitative PCR as described previously³. Individual results were obtained as a mean from two separate runs and expressed in ng/mL for cf n-DNA and as genome equivalents (GE)/ml plasma (1 GE = 6.6 pg DNA) for cf mt-DNA⁷⁹.

Determination of plasma cell free DNA integrity. The integrity of circulating cf n-DNA was evaluated by a quantitative real-time PCR (qPCR) targeting the human GAPDH (glyceraldehyde 3-phosphate dehydrogenase) gene (gene ID 2597). The length of the amplicons selected for this assay was 97 and 229 bp, respectively. The ratio of the concentration of the longer amplicon (GAPDH₂₂₉) to the concentration of the shorter one (GAPDH₉₇) (ranging from 0 to 1) defined the integrity index 229/97 ($I_{229/97}$), which was used to estimate the fragmentation of cf n-DNA. Higher values of $I_{229/97}$ (e.g. $I_{229/97} = 1$) indicate that all the cf n-DNA molecules are at least 229 bp in length in the GAPDH gene while lower values show that cf n-DNA contains fragments below 229 bp in the same target gene sequence. Similarly two amplicons, one of 218 bp length (encoding part of mitochondrial ATPase 6 gene, ID 4508, and mitochondrial ATPase 8 gene, ID 4509) and the second one of 78 bp length (encoding part of mitochondrial ATPase 8 gene) for calculation of cf mt-DNA integrity index 218/78 ($I_{218/78}$) were chosen. The assay was designed in a way that the forward primer and the probe were the same for each pair of amplicons, whereas two different reverse primers were used. Both, GAPDH₉₇ and MTATP₇₈ primers and corresponding probes were described and successfully used in previous studies⁷⁹. The primers for longer sequences (GAPDH₂₂₉ primer and MTATP₂₁₈ primer) were planned using *Primer3* software (Table 4). qPCR was carried out in 20 μ L of total reaction volume containing 6.5 μ L H₂O, 10 μ L TaqMan[®] Universal PCR Master Mix (Applied Biosystems, Branchburg, New Jersey, USA), 0.25 μ L of each of the two matched primers (one forward and one reverse primer for longer

Primer/probe	Integrity index of plasma cell free DNA	
	Nuclear I _{229/97}	Mitochondrial I _{218/78}
Forward primer	CCCCACACACATGCACTTACC	AATATTAAACACAAACTACC
Reverse primer	ATCAAACCTCAAAGGGCAGGA [229 bp]	TGGGTGGTGATTAGTCGGTTG [218 bp]
Reverse primer	CCTAGTCCCAGGGCTTTGATT [97 bp]	TGGTTCTCAGGGTTTGTAT [78 bp]
MGB probe	FAM-TAGGAAGGACAGGCAAC-MGB	VIC-CCTCACCAAAGCCATA-MGB

Table 4. The sequences of primers and probes used for determination of integrity index of circulating cell free nuclear DNA (cf n-DNA) and cell free mitochondrial DNA (cf mt-DNA). The forward primer and the probe were the same for each pair of amplicons, whereas the sequences of reverse primer were different. In brackets the amplicon length.

amplicon or one forward and one reverse primer for shorter amplicon) (Sigma-Aldrich), 1 μ L of a FAM-labeled MT-ATP 8-probe or 1 μ L of a MVIC-labeled GAPDH-probe (both probes from Applied Biosystems, Branchburg, New Jersey, USA), and 2 μ L of TE buffer containing cfDNA isolated from plasma. The final concentrations of primers and probes were 0.6 μ mol/L and 0.4 μ mol/L, respectively. Negative control samples received 2 μ L of TE buffer without cfDNA from plasma. Reaction was done in duplicate and performed using the 7900 HT Real-time PCR System (Applied Biosystems) under the following conditions: an initiation step at 50 °C for 2 min., followed by a first denaturation at 95 °C for 10 min., then 40 cycles at 95 °C for 15 s. and annealing at 60 °C for 1 min. Serial dilutions of human genomic DNA (Roche) (final concentrations from 0.5 ng/mL to 5000 ng/mL) were used to construct the calibration curve ($r^2 = 0.9996$) for measurement of PCR products.

Resting and fMLP-induced luminol enhanced whole blood chemiluminescence. The resting (rLBCL) and fMLP-induced LBCL (fMLP-LBCL) reflecting ROS production by circulating phagocytes were measured according to the method of Kukovetz *et al.*²¹ with some modifications³¹.

Briefly, 30 μ L of venous blood (collected into vacutainer tubes with EDTA) was initially diluted with 1 mL of mixture solution prepared fresh before the assay and containing 127.5 μ g/mL of luminol³¹. Then 103 μ L of diluted blood was added to 797 μ L of mixture solution (to obtain a final blood dilution of 300 times), placed in a multi-tube luminometer (AutoLumat Plus LB 953, Berthold, Germany) equipped with a Peltier-cooled detector to ensure high sensitivity, and a low and stable background noise signal, and incubated for 15 min. at 37 °C in the dark. Then 100 μ L of the control solution (solvent for fMLP) or 100 μ L of fMLP solution (final agonist concentration in the sample of 2×10^{-5} mol/L) for measurement of rLBCL and fMLP-LBCL was added by automatic dispensers and the total light emission was automatically measured for 120 seconds. Individual results were obtained as the mean of triplicate experiments and rLBCL and fMLP-LBCL was expressed in relative light units (RLU) per 10^4 phagocytes (PMNs and monocytes) present in the assayed sample³¹.

Statistical analysis. Results are expressed as a mean (SD) and median (interquartile range). Analysis of variance (ANOVA) for repeated observations (parametric test) or Friedman's ANOVA (non parametric test) was applied for the assessment of changes in variables over time (before and after three consecutive exercise bouts) depending on data distribution which was tested with Shapiro-Wilk's W test. In case of statistically significant ANOVA, the post hoc analyses were done with Scheffé's test or post hoc analysis for Friedman's ANOVA (multiple comparisons at 2 different time points). Correlations between variables were determined using Spearman's ρ . A p value < 0.05 was considered significant.

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Author contributions

R.S. D.N. K.W. were involved in conceptualization, study design and methodology, R.S. P.M. collected the data, R.S. E.P. A.S. A.W. conducted experiments, P.K. T.B. K.W. were responsible for participant safety during the execution of treadmill exhaustive exercise, D.N. G.P. supply resources, R.S. performed data analysis, statistics and visualization, D.N. was responsible for project administration and supervision, R.S. D.N. wrote the main text, D.N. R.S. E.P. G.P. review and edit the manuscript. All authors approved final submission.

Competing interests

The authors declare no competing interests.

Additional information

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