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Tim Hofer
University of Florida

Luigi Fontana
Washington University School of Medicine

Stephen D. Anton
University of Florida

Edward P. Weiss
Washington University School of Medicine

Dennis Villareal
Washington University School of Medicine

See next page for additional authors

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Authors
Tim Hofer, Luigi Fontana, Stephen D. Anton, Edward P. Weiss, Dennis Villareal, Bhaskar Malayappan, and Christiaan Leeuwenburgh

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Long-Term Effects of Caloric Restriction or Exercise on DNA and RNA Oxidation Levels in White Blood Cells and Urine in Humans

Tim Hofer,1* Luigi Fontana,2,3* Stephen D. Anton,1 Edward P. Weiss,2,4 Dennis Villareal,2 Bhaskar Malayappan,1 and Christiaan Leeuwenburgh1

Abstract

Excessive adiposity is associated with increased oxidative stress and accelerated aging. Weight loss induced by negative energy balance reduces markers of oxidation in experimental animals and humans. The long-term effects of weight loss induced by calorie restriction or increased energy expenditure induced by exercise on measures of oxidative stress and damage have not been studied in humans. The objective of the present study was to compare the effects of 20% caloric restriction or 20% exercise alone over 1 year on oxidative damage to DNA and RNA, as assessed through white blood cell and urine analyses. Eighteen men and women aged 50 to 60 years with a body mass index (BMI) between 23.5 to 29.9 kg/m² were assigned to one of two conditions — 20% CR (n = 9) or 20% EX (n = 9) — which was designed to produce an identical energy deficit through increased energy expenditure. Compared to baseline, both interventions significantly reduced oxidative damage to both DNA (48.5% and 49.6% reduction for the CR and EX groups, respectively) and RNA (35.7% and 52.1% reduction for the CR and EX groups, respectively) measured in white blood cells. However, urinary levels of DNA and RNA oxidation products did not differ from baseline values following either 12-month intervention program. Data from the present study provide evidence that negative energy balances induced through either CR or EX result in substantial and similar improvements in markers of DNA and RNA damage to white blood cells, potentially by reducing systemic oxidative stress.

Introduction

Oxidative damage to DNA, proteins, lipids, and other cellular components accumulates over time and has been hypothesized to be a major cause of aging and age-associated diseases.1,2 In support of this hypothesis, accelerated aging and cellular oxidative damage have been linked in rodents.3 Experimental evidence indicates that excessive adiposity is associated with increased oxidative stress.4 In contrast, weight loss in obese patients causes a significant reduction in markers of oxidation, such as urinary 8-iso-PGF2α and protein carbonylation.5,6

Negative energy balance can be achieved by reducing energy intake or increasing energy expenditure. In rodents, both caloric restriction (CR) and exercise training have been found to reduce oxidative damage to lipids, protein, and DNA in many tissues.7–9 Calorie restriction has consistently been shown to extend lifespan and reduce age-related diseases in numerous species.10 Exercise training, however, has only been found to increase average lifespan and does not affect maximal lifespan.11 One explanation for these disparate effects is that exercise training may increase oxidative damage in some instances.12,13 Recent reports, however, indicate that exercise training does not increase oxidative damage in weight-matched rodents.14 Thus, the literature is currently mixed regarding the effect of exercise on oxidative damage. Moreover, the effects of long-term negative energy balance induced by either caloric restriction or exercise alone...
for reducing markers of oxidative stress in humans are not known.

We conducted a one-year randomized controlled trial in middle-aged lean and overweight men and women to evaluate the effect of body fat reduction induced by a 20% decrease in energy intake alone or 20% increase in energy expenditure on markers of DNA and RNA damage. We hypothesized that both calorie restriction (CR) and exercise (EX) would reduce DNA and RNA oxidative damage, but that these changes would be more pronounced when the energy deficit was achieved through CR than through EX.

Method

The current study represents an ancillary project of the main study entitled Comprehensive Assessment of Long-term Effects of Reducing Intake of Energy (CALERIE, phase 1). This study was approved by the Human Studies Committee and the General Clinical Research Center Scientific Advisory Committee of Washington University School of Medicine. All subjects gave their informed consent before their participation.

Participants

An extensive screening process was employed in the CALERIE study to ensure participants were healthy and suitable for participation in this trial. Details of the screening process have been previously reported. Briefly, men and women aged 50 to 60 years with a body mass index (BMI) between 23.5 to 29.9 kg/m² were recruited. Potential participants were excluded for the following reasons: (1) a history or clinical evidence of coronary artery disease, stroke, or lung disease, (3) a resting blood pressure (BP) of diabetes or a fasting blood glucose value ≥126 mg/dl, (2) a history or clinical evidence of coronary artery disease, stroke, or lung disease, (3) a resting blood pressure (BP) ≥170 mmHg systolic or ≥100 mmHg diastolic, or (4) a recent history or evidence of malignancy. Furthermore, all candidates had to be nonsmokers and sedentary (defined as exercising <40 min per week during the 6 months before baseline testing). Women had to be postmenopausal.

Study design

For the larger study, eligible participants were randomized, with stratification for sex, to one of three groups in a 2:2:1 sequence: caloric restriction (CR) group \( n = 19 \), exercise (EX) group \( n = 19 \), or healthy lifestyle (HL) control group \( n = 10 \) for 1 year. For the purpose of the current study, data were analyzed only for participants assigned to the CR and EX groups. The group samples were randomized and batch analyses were performed under the same conditions (see DNA and RNA oxidation section). Because one subject dropped out of each group before completing the study and because biological specimens were not available for all subjects, sample sizes for this ancillary study are smaller than those reported previously for the Washington University CALERIE study (sample sizes for each outcome are provided in the table and figures).

CR intervention

The goal of the CR intervention was to create a 20% energy deficit through a reduction in energy intake (without changing physical activity levels) for the duration of the 1 year intervention. Participants were instructed to decrease energy intake by 16% during the first 3 months and by 20% during the remaining 9 months. Diet prescriptions were based on participants’ baseline energy intake, which was assumed to be equal to total daily energy expenditure as determined by the doubly labeled water (DLW) method. To enhance compliance to the intervention, participants met with registered dietitians on a weekly basis to discuss and review strategies for reducing energy intake, as well as attended weekly group meetings led by a dietitian and a behavioral psychologist. Further details about the CR intervention, including compliance data, have been reported previously.

Exercise intervention

The goal of the EX intervention was to induce an energy deficit identical to the CR intervention by increasing daily energy expenditure through physical activity without changing caloric intake. Participants were instructed to increase energy expenditure by 16% of baseline total daily energy expenditure for the first 3 months and by 20% for the subsequent 9 months. They were informed that exercise sessions could be completed in one or several daily bouts, and exercise trainers worked closely with participants to monitor their energy expenditure goals. The participants exercised while using heart rate monitors (Polar S610, Polar Electro Oy, Kempele, Finland), which provided estimates of energy expenditure during exercise. To ensure participants’ energy intake remained stable, the study dietitians periodically monitored energy intake using 7-day food diaries and provided consultation as needed. Additional details about the EX intervention, including compliance data, have been reported previously.

Weight measurements

Body weight was measured twice in the morning following a 12 h fast. At baseline, weight was calculated from the mean of four weights measured over a 4-week baseline period. Twelve-month body weights represent the mean of three weekly weights obtained at the beginning, middle, and end of the 2-week assessment periods.

DNA and RNA oxidation

Urine analysis. Urine samples were collected for 12 h overnight from the participants using standardized procedures, aliquoted and frozen under argon at −80°C until analysis. The urinary RNA and DNA oxidative damage products 2,6-diamino-4-hydroxy-5-formamidopyrimidine (FapyGua), 8-oxo-7,8-dihydroguanine (8-oxoGua), 8-oxo-7,8-dihydroguanosine (8-oxoGuo), and 8-oxo-7,8-dihydro-2′-deoxyguanosine (8-oxodGuo) (chemical structures are shown in Fig. 1) were simultaneously measured for participants on CR and exercise employing electrospray tandem mass spectrometry detection (MS/MS) in multiple reaction monitoring (MRM) mode on a Finnigan TSQ 7000 triple quadrupole mass spectrometer (Thermo Electron Corporation, San Jose, CA). This method does not require any sample preparation except for the addition of buffer and 13C- and 15N-labeled internal standards (isotope dilution) to the urine prior to sample injection into the HPLC-MS/MS system, as
described in detail elsewhere. The urinary biomarkers were normalized to the concentration of creatinine, which was assessed using a creatinine kit (Cayman Chemical, Ann Arbor, MI). The manufacturer’s instructions were followed and the formation of acid-sensitive chromogen after reduction of the sample with picrate was measured spectrophotometrically at 500 nm.

**Blood analyses.** A venous blood sample was taken after participants had fasted for at least 12 h at baseline and at 12 months. In the EX group, blood samples were obtained at least 48 h after the last exercise session. Blood was collected in 10 mL EDTA-Vacutainer tubes from Becton Dickinson (Franklin Lakes, NJ), and white blood cell (WBC) “buffy coats” were collected following centrifugation (800 g, 20 min, 4°C) using a large orifice pipette tip and placed into 1.5 mL Eppendorf tubes and immediately frozen at –80°C. RNA and DNA oxidative damage levels to WBC were measured in the CR (\(n = 9\)) and EX (\(n = 9\)) subjects. Buffy coats of WBC were thawed from –80°C and placed on ice. Working on slush ice (0°C) during all steps, cells were lysed in 4.5 mL of 3 M GTC buffer (0.2 wt.% N-L-Sarcosine, 20 mM tris [pH 7.5]) containing 10 mM of the freshly dissolved metal chelator deferoxamine meylate (DFOM) during homogenization using a Potter-Elvehjem homogenizer. All chemicals and supplies used to extract and analyze nucleic acids were as previously described. After transferring the homogenates to 15 mL Phase-Lock Gel (PLG) tubes, an equal amount of a phenol-
chloroform mixture (pH 6.7) was added, and proteins and lipids extracted into the phenol phase. After vortexing in intervals of 10 min keeping the tubes on ice, samples were centrifuged (2000 g, 30 min, 0°C), and the upper aqueous phase containing nucleic acids was transferred into a new PLG tube. The procedure was repeated once. After transferring into a new PLG tube, an equal amount of chloroform-isoamylalcohol (24:1) was added to remove any remaining phenol by hand mixing followed by centrifugation. The procedure was repeated once, and the upper aqueous phase was collected and nucleic acids precipitated by adding 1:1 isopropanol, mixing and incubating at –80°C overnight. Total nucleic acids were collected by centrifugation at 10,000 g, 0°C for 10 min. Nucleic acids were washed in 70% ethanol, spun down at 3000 g (10 min, 0°C), and air-dried at room temperature for 10 min. RNA/DNA hydrolysis was performed using Nuclease P1 and alkaline phosphatase, and 8-oxoGuo/creatinine was performed using HPLC-ECD with a Coulochem III electrochemical detector (ESA Inc., Chelmsford, MA), as described previously.25

**Statistical analysis**

Baseline characteristics of participants were compared between groups using Fisher’s exact test for categorical data and independent t-tests for quantitative data. One-way ANOVA and paired t-test were performed to assess within-group changes, and analysis of covariance (ANCOVA) was used for between group comparisons after adjustment for initial values with subsequent Kruskal-Wallis test for post-hoc comparisons. All statistical tests were two-tailed, and significance was accepted at p < 0.05. Data are presented as means ± standard error (SE) at each time point, and for the change between baseline and 12 months. All analyses were performed using Prism 4 from GraphPad software (San Diego, CA).

**Results**

**Participants**

Subject characteristics data represent all participants who were included in the WBC- or urine-based analyses (N = 34). Male/female representation in the CR (7 men, 10 women) and EX (5 men, 12 women) groups did not differ significantly (p = 0.72). The participants in the CR group were slightly younger than those in the EX group (54.6 ± 3.1 vs. 58.6 ± 2.7 years, p = 0.0004), although all participants were within the 50 to 60 year range required for the study. Average BMI was in the overweight range (CR, 26.8 ± 2.4 kg/m²; EX, 27.0 ± 1.8 kg/m²) and did not differ between groups (p = 0.89). Results from statistical analyses performed on each subgroup alone (i.e., participants included in the WBC-based analyses (n = 18) or those included in the urine-based analyses (n = 29) were not different from analyses conducted on the entire sample.

**Body weight**

Baseline body weight was similar in the CR (79.1 ± 9.8 kg) and exercise (75.6 ± 10.3 kg) groups and did not differ between groups (p = 0.32). Body weight decreased in both groups (CR, −10.2 ± 1.5%, p < 0.0001; EX, −8.0 ± 1.5%, p < 0.0001) and these decreases did not differ between groups (p = 0.31). These results are based on subjects who were included in either the WBC- or urine-based data analyses (n = 34); statistical results from analyses on either subgroup alone were not different from the results from all subjects combined (data not shown).

**RNA and DNA oxidative damage**

Analyses of white blood cells. The two groups did not differ in levels of RNA or DNA oxidative damage at baseline. As presented in Figure 2, both interventions significantly reduced oxidative damage to both DNA and RNA. Levels of WBC DNA oxidation (8-oxoGuo/106 dGuo) decreased by 48.5% from baseline (4.24 ± 0.39) to 1 year (2.19 ± 0.34) for the CR group, and by 49.6% from baseline (3.03 ± 0.56) to 1 year (1.66 ± 0.22) for the EX group. Similarly, levels of total WBC RNA oxidation expressed as 8-oxoGuo/106 Guo decreased by 35.7% from baseline (4.15 ± 0.49) to 1 year (2.67 ± 0.28) for the CR group, and by 52.1% from baseline (4.15 ± 0.63) to 1 year (1.99 ± 0.43) for the EX group.

**Urine levels of nucleoside oxidation products.** Urinary levels for each nucleic acid oxidation product at baseline and at one-year are presented in Table 1. In contrast to the WBC data, nucleic acid oxidation products were not found to be significantly different from baseline following either 12-month intervention program. We did, however, find differences in the amount of baseline levels between the different excreted oxidized nucleic acid bases. Comparison of urinary excretion levels of the four nucleic acid products at baseline before interventions in all groups were different: baseline

<table>
<thead>
<tr>
<th>Table 1. Urinary Levels of RNA and DNA Oxidation Products</th>
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<tr>
<td><strong>EX (n = 15)</strong></td>
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<td><strong>Baseline</strong></td>
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<td>Creatinine</td>
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<td>FapyGua/creatinine</td>
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<td><strong>One year</strong></td>
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<tr>
<td>Creatinine</td>
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<tr>
<td>FapyGua/creatinine</td>
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<td>8-oxoGua/creatinine</td>
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<td><strong>CR (n = 14)</strong></td>
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<td><strong>Baseline</strong></td>
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Levels of FapyGua, 8-oxoGua, 8-oxoGua, and 8-oxoGua are expressed as nmol/mmol creatinine, creatinine alone as mM. Values are given as mean ± SE.
levels of 8-oxoGua (162 ± 33 nmol 8-oxoGua/mmol creatinine) was significantly higher ($p < 0.001$) than the levels of FapyGua (4.19 ± 1.4), 8-oxoGuo (6.68 ± 1.5), and 8-oxodGuo (2.57 ± 0.50) (Fig. 3).

**Discussion**

In this 1 year randomized trial, we compared the effects of weight loss induced by CR or EX, both producing a 20% energy deficit, on DNA and RNA oxidative damage in healthy normal weight and overweight middle-aged men and women. Our results provide evidence that negative energy balance induced through both CR and EX decrease levels of oxidative RNA and DNA damage in WBC to a similar extent. No significant changes in urinary nucleic acid oxidation levels were found. The reason for these disparate findings is not entirely clear but may be related to build-up of oxidative damages in WBC over longer time periods. Additionally, the large individual differences we observed in urinary levels of nucleic acid oxidation products makes determining statistical differences difficult. The variability was very apparent when comparing the four different nucleic acid oxidation products, with 8-oxoGua/creatinine having the highest levels and greatest variability.

This study represents a first attempt to examine the effects of energy deficits created from either prolonged CR or EX over 1 year on both oxidative RNA and DNA damage. In line with our findings, a previous 6-month study found that both a CR only (25% CR) and a CR plus EX intervention (12.5% CR plus 12.5% increase in energy expenditure through EX) reduced DNA strand breaks in human WBC, assessed through the comet assay. Additionally, other studies have found that oxidative DNA damage to leukocytes is reduced in physically active individuals, as compared to sedentary persons, following an acute bout of exercise. The reduction in oxidative DNA and RNA damage to WBC observed may be due to a reduction in systemic inflammation and related production of reactive oxygen species (ROS); both CR and EX have been found to reduce circulating markers of inflammation, such as C-reactive protein (CRP) and related cytokines (e.g., interleukins). In addition to affecting systemic hormone and metabolic regulatory parameters, CR and EX may affect behavioral patterns (e.g., sleep duration and quality). Moreover, a recent theory suggest that metabolic syndromes (obesity and type 2 diabetes) are related to changes in gut microbial composition and mass, where uptake of metabolites and gram-negative bacteria released lipopolysacharide (LPS) affect systemic metabolic parameters and can increase the systemic inflam-
mation related to ROS production.\textsuperscript{31} Both CR and EX can affect the gut microbial mass as well as composition. Further, it has been unknown which RNA and DNA oxidation product is mainly excreted in urine as it has been debated if the initial 8-OH-8-H-Gua\textsuperscript{*} radical intermediate (Fig. 1) formed after oxidant attack undergoes an oxidation (forms 8-oxoGua) or a reduction (ring opening into FapyGua).\textsuperscript{20,32} The clearly higher concentrations of 8-oxoGua than FapyGua found in human urine (Fig. 3) strongly supports an oxidation of the 8-OH-8-H-Gua\textsuperscript{*} radical intermediate as the major mechanism. As the base guanine is the moiety having the lowest oxidation potential in both RNA and DNA,\textsuperscript{32} the urine suggest that 8-oxoGua is the major nucleic acid lesion formed and excreted in humans as result of oxidative damage. However, we cannot rule out that urinary 8-oxoGua stems from RNA or DNA as it could stem from both, after being recognized, removed, and excreted. Excetration of the specific RNA oxidation product 8-oxodGuo was higher than the specific DNA oxidation product 8-oxodGuo. It should also be noted that oxidative base lesions in urine may to a certain extent also stem from absorption of digested oxidized food products, from gut microbial end products, or from dead cells which could affect the results.\textsuperscript{22–24} In DNA, oxidized bases are mainly removed by base excision repair (BER), whereas the processes for removal of damaged RNA (rRNA, tRNA, mRNA, and siRNA) are largely unknown. In (BER), whereas the processes for removal of damaged RNA oxidized bases are mainly removed by base excision repair (BER), whereas the processes for removal of damaged RNA (rRNA, tRNA, mRNA, and siRNA) are largely unknown. In agreement with previous observations,\textsuperscript{33,34} we found that the urinary level of the RNA-specific oxidation product 8-oxoGuo was higher than that of DNA (8-oxodGuo) for both CR and EX groups at baseline. The results of the present study should be interpreted in the context of its limitations. First, the generalizability of these findings is limited by our small sample size, as well as restricted body mass index range. Thus, these findings need to be replicated in future studies that utilize larger sample sizes and more diverse populations. Another weakness is the inability to completely control for food intake during the week prior to the collection of the urine samples, which may explain the variability of the oxidized nucleic acids in the urine, specifically for 8-oxoGua. The present study also had a number of strengths. This is the first study to test the effects of energy deficits created through prolonged CR versus EX on DNA and RNA oxidative damage, and the first report of RNA oxidative damage levels in WBC. Other strengths include the use of a randomized controlled trial design, comprehensive assessments of energy intake and expenditure using DLW, and the high rate of adherence to both interventions, as evidenced by the significant weight loss in both groups. In conclusion, prolonged CR and EX interventions, both producing a 20% energy deficit, were found to decrease oxidative DNA and RNA damage in human WBC. However, significant changes in DNA and RNA oxidation products were not observed through urinary analyses, potentially due to greater individual variability on this measure. Overall, our findings suggest energy deficits created through both CR and EX reduce DNA and RNA damage to WBC, potentially by reducing systemic oxidative stress.

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Address reprint requests to:
Steve Anton, Ph.D.
Christiaan Leeuwenburgh, Ph.D.
Department of Aging and Geriatrics
University of Florida
210 East Mowry Road
P.O. Box 112610
Gainesville, FL 32611
E-mail: santon@aging.ufl.edu; cleeuwen@aging.ufl.edu
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