Acute and chronic watercress supplementation attenuates exercise-induced peripheral mononuclear cell DNA damage and lipid peroxidation

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Abstract
Pharmacological antioxidant vitamins have previously been investigated for a prophylactic effect against exercise-induced oxidative stress. However, large doses are often required and may lead to a state of pro-oxidation and oxidative damage. Watercress contains an array of nutritional compounds such as β-carotene and α-tocopherol which may increase protection against exercise-induced oxidative stress. The present randomised controlled intervention was designed to test the hypothesis that acute (consumption 2 h before exercise) and chronic (8 weeks consumption) watercress supplementation can attenuate exercise-induced oxidative stress. A total of ten apparently healthy male subjects (age 23 (SD 4) years, stature 179 (SD 10) cm and body mass 74 (SD 15) kg) were recruited to complete the 8-week chronic watercress intervention period (and then 8 weeks of control, with no ingestion) before crossing over in order to compete the single-dose acute phase (with control, no ingestion). Blood samples were taken at baseline (pre-supplementation), at rest (pre-exercise) and following exercise. Each subject completed an incremental exercise test to volitional exhaustion following chronic and acute watercress supplementation or control. The main findings show an exercise-induced increase in DNA damage and lipid peroxidation over both acute and chronic control supplementation phases (P<0·05 v. supplementation), while acute and chronic watercress attenuated DNA damage and lipid peroxidation and decreased H₂O₂ accumulation following exhaustive exercise (P<0·05 v. control). A marked increase in the main lipid-soluble antioxidants (α-tocopherol, γ-tocopherol and xanthophyll) was observed following watercress supplementation (P<0·05 v. control) in both experimental phases. These findings suggest that short- and long-term watercress ingestion has potential antioxidant effects against exercise-induced DNA damage and lipid peroxidation.

Key words: Exhaustive exercise; Watercress prophylaxis; DNA damage; Reactive oxygen species

Free radicals are produced in mammalian cells during a complex array of physiological processes, which, if left uncontrolled, have the potential to cause metabolic damage (3). Free radical activity is moderated by a complex antioxidant network (2), which has the capacity to temper potentially negative cellular effects (5). The human body depends on the intake and recycling of dietary antioxidant vitamins such as vitamins C, E and β-carotene in order to support the enzymatic systems (4). However, a systemic increase in free radical production may overwhelm antioxidant defences leading to a state of oxidative stress (5) in the form of molecular damage to DNA, cellular lipid membranes and proteins (60).

It has been established that various modes and intensities of exercise can lead to an exacerbated state of oxidative stress, although the exact mechanism of free radical production remains elusive (7). Despite this, pharmacological interventions have frequently been investigated for a prophylactic effect against oxidative stress induced by exercise (59–11), and although research has shown a prophylactic effect with exogenous antioxidant consumption, in some cases, large doses of antioxidants are required. As it has been shown that an overconsumption of oral antioxidants may lead to a pro-oxidant state, causing a disturbance in redox biochemistry (12), it is therefore imperative that food sources naturally high in antioxidant vitamins are considered, due to their capacity to provide increased systemic and cellular protection without excessively elevating in vivo antioxidant vitamin concentrations (13).

Cruciferous vegetables, in particular cauliflower, cabbage, broccoli and watercress, are known to reduce oxidative DNA experimentation in human cells (14).

Abbreviations: LOOH, lipid hydroperoxides; PC, protein carbonyls.

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More specifically, Gill et al.\(^{(15)}\) demonstrated that watercress, which contains per gram weight one of the highest concentrations of glucosinolates and carotenoids of any vegetable, protects lymphocyte DNA against damage in smokers, while Boyd et al.\(^{(16)}\) suggested that crude watercress extracts are also known to provide increased protection against DNA damage associated with human colon cancer cells and H\(_2\)O\(_2\) exposure.

Given that exhaustive exercise promotes oxidative stress, causing severe perturbations to cell structure and function, there is a need to ascertain an antioxidant vitamin-rich vegetable that has the potential to counteract exercise-induced oxidative stress. Although recent research has indicated that watercress may play a role in lymphocyte cell protection\(^{(15)}\), the efficacy of short- and long-term watercress ingestion on exercise-induced oxidative stress, to the best of our knowledge, is not yet known.

We therefore hypothesise that acute and chronic watercress consumption will provide effective prophylaxis against exercise-induced oxidative stress. Thus, the primary aim of the present investigation is to examine the efficacy of acute and chronic watercress ingestion against exercise-induced oxidative stress. A randomised controlled experimental design incorporating a comprehensive assessment of oxidative stress indices was used to test this hypothesis.

**Methods**

**Human subjects and experimental design**

A total of ten apparently healthy male subjects were recruited to participate in the chronic phase of the experiment first. The study was conducted according to the guidelines laid down in the Declaration of Helsinki and all procedures involving human subjects were approved by the University of Ulster Research Ethics Committee. It is noted that randomised assignment of participants to a supplemented or control group and incorporating a cross-over element is the most robust method of testing food supplements. However, due to the availability of this naturally occurring product and the quantities required to complete the experiment, delivery of product could only be guaranteed during the optimum growing months. For this reason, the participants completed 8 weeks of chronic watercress ingestion before washing out for an additional 8 weeks and then subsequently completing 8 weeks of no ingestion (control). Upon the completion of the chronic phase (lasting in total 24 weeks), eight out of the original ten participants volunteered to take part in the acute phase with an additional two participants recruited. A randomised cross-over design was utilised in the acute phase with participants washing out for a 2-week period between the intervention and the control. Please see Table 1 for participant characteristics.

<table>
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<tr>
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<th>Acute Mean</th>
<th>SD</th>
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*bpm, Beats per min.*

Table 1. Participant characteristics (Mean values and standard deviations, n 10)

Participants with a family history of any known haematological or cardiovascular-related condition were excluded from the study. Subjects were also excluded if they smoked or were taking any form of antioxidant supplement. All subjects were assigned to complete two phases of experimental testing, of which one was a chronic interventional phase lasting 24 weeks and the other an acute experimental phase, consisting of a randomised cross-over assignment to either watercress supplementation 2 h before exhaustive exercise or the control group. All commercially available watercress was supplied by Vitacress (Vitacress Limited). All participants were instructed to refrain from altering their usual dietary habits throughout the experimental phases of testing. Written informed consent was obtained from all subjects before participation. A schematic overview of each experimental phase is provided in Figs. 1 and 2.

**Chronic interventional phase**

Before chronic supplementation and control, a baseline venous blood sample was collected for quantification of lipid-soluble antioxidants. Participants were then required to consume one 85 g portion\(^{(15)}\) of raw watercress on a daily basis for a total of 8 weeks. Watercress was ingested in a single sitting between the hours of 12.00 and 14.00 daily in combination with other foods. Following 8 weeks of supplementation, all participants completed an exercise test to volitional exhaustion while venous blood was drawn at rest and after exercise to determine lipid-soluble antioxidant status and other indices of oxidative stress. Following an 8-week washout period\(^{(15)}\), participants entered an 8-week control phase with no watercress ingestion and then repeated the exercise test to exhaustion.

**Acute experimental phase**

During the acute experimental phase, participants were randomly assigned using a simple lottery system to either a supplementation or control group. For the supplementation group, participants were asked to consume a single portion of...
85 g of raw watercress and 500 ml of water within 30 min between 10.00 and 10.30 hours. Following ingestion, participants rested for 2 h before completing an exercise test to volitional exhaustion. The control group also completed the exercise test to volitional exhaustion, and did not ingest watercress but did consume 500 ml of water. Following a 2-week washout period (17), all participants crossed over to consume either watercress or nothing and were retested as above.

Exhaustive acute exercise protocol

To avoid the potential effects of diurnal variation, each participant was required to consistently attend the laboratory testing sessions at the same time of day (09.00 hours) for all phases. On each arrival of participants at the laboratory, body mass and stature were measured using standard methods (see Table 1). All participants were required to complete a treadmill test to volitional exhaustion in which the protocol was specifically designed to be progressive and incremental in order to elicit VO2max. Treadmill speed was set at 11 km/h with a 1% gradient rise at each 1 min interval until volitional fatigue. Validation of VO2max was confirmed when (1) the RER was above 1·15 arbitrary units, (2) a plateau in the oxygen uptake/exercise intensity relationship (>2 ml/kg per min) and (3) a heart rate of within 10 beats per min of age-predicted maximum (220 – age). Oxygen uptake was measured using standard laboratory gas analysis (Cosmed Quark b2), while heart rate was measured via a portable short-angle telemetry device (Polar Sports Tester). All subjects achieved maximum oxygen uptake.

Supplementation compliance

Subjects were asked to complete a weekly journal detailing total amounts of watercress ingested over the chronic interventional phase.

Haematology

Following a standardised 12 h overnight fast, whole blood was drawn from a prominent forearm ante-cubital vein using the vacutainer method (Becton-Dickinson). Blood was collected at baseline (pre-supplementation), at rest (post-supplementation/pre-exercise) and exercise (immediately post-exhaustive exercise) for the quantification of a range of lipid-soluble antioxidants, and at rest and exercise for various indices of oxidative stress. Blood was drawn into either di-potassium EDTA or serum separation tubes. EDTA and serum separation tubes (once clotted) were centrifuged (MSE Centuar 2; Labcare) at 3000 rpm for 10 min at –4°C. Aliquots of plasma and serum were stored at –80°C for a maximum of 3 months. All samples from the same participant were analysed within the same batch. Hb (g/dl) was measured using a β-Hb photometer (Hemocure Limited), and packed cell volume (%) was measured using the standard microcapillary reader technique to aid in the correction of all biochemical markers for acute exercise-induced plasma volume shifts using the equations of Dill & Costill (18).

Biochemical analysis

DNA. Lymphocytes were isolated from the whole blood (EDTA) by layering 3 ml onto Histopaque-1077 (3 ml) and centrifuged (MSE Centuar 2, Labcare) at 3500 rpm (relative centrifugal force 1356 g) for 30 min at room temperature. The opaque mononuclear layer was aspirated and washed three times using 15 ml PBS (one tablet added to 200 ml of double-distilled water, 0·2 M, pH 7·2). The comet assay was subsequently carried out on prepared cells using the protocol of Singh et al. (19). Dakin fully frosted slides (3 inch × 1 inch; 1·2 mm thick; Richards Supply Company Limited) were covered with 100 μl of 0·5% normal-melting point agarose and allowed to solidify under a coverslip (22 × 40 mm, no. 1 thickness; GBH, Laboratory Supplies). Then, 50 μl of cells (1 × 10⁴) were mixed with 50 μl of low-melting point agarose layered on top of the
normal-melting point agarose and allowed to solidify under a coverslip. The coverslip was then removed and the slides were placed in lysis buffer (2·5 M-NaCl, 100 mM-Na-EDTA, 10 mM-Tris, 1 % Triton X and 10 % dimethyl sulfoxide, pH 10) for 1 h at 4°C to break down the cell membranes. The slides were removed and placed in a horizontal electrophoresis unit containing electrophoresis buffer (300 mM-NaOH, 1 mM-EDTA, pH 12·5) for 20 min to allow the DNA double helix to relax and unwind. Electrophoresis (25 V, 300 mA, 0·15 V/cm) was performed for 20 min at room temperature. Following electrophoresis, the slides were rinsed with neutralising solution (0·4 M-Tris, pH 7), stained with 50 μl ethidium bromide and a coverslip was applied. A random sample of fifty cells from each slide was analysed using a Hewlett-Packard VGA monitor and Fenestra Comet software program (version 2.22) at magnification of 400 × using an epifluorescent microscope (Olympus BH2). Tail length was selected to report DNA damage, as it acts as a sensitive measure of low levels of DNA damage(20). This assay was performed on each experimental day using fresh lymphocytes. The intra/inter-assay CV were <9 and <11 %, respectively.

**Lipid hydroperoxides.** Serum lipid hydroperoxides (LOOH) were measured spectrophotometrically using the method of Wolff(23). This ferrous iron/xylene orange assay quantifies the susceptibility to Fe-induced LOOH formation in the blood. The presence of Fe ions in the assay protocol might therefore yield slightly higher LOOH values compared with other methods. Briefly, 90 μl serum were incubated with 10 μl catalase for 30 min at room temperature. To this solution, 900 μl ferrous iron/xylene orange reagent 1 (250 μM-ammonium ferrous sulphate, 100 μM-xylene orange, 100 μM-sorbitol and 25 μM-H2SO4) were added and incubated for a further 30 min at room temperature in the dark. Standard solutions were prepared from H2O2 in the range of 0–5 μMol/l and also incubated for 30 min with the ferrous iron/xylene orange 1 reagent, after which the samples were centrifuged in a Beckman microfuge for 5 min to remove any flocculated material. The absorbance of the supernatant was read spectrophotometrically (U-2001; Hitachi) at 560 nm against the standard curve that was linear in the range of 0–5 μMol/l. The intra/inter-assay CV were <2 % and <4 %, respectively.

**Protein carbonyls.** Protein carbonyls (PC) are a reliable measure of protein oxidation and were quantified with an assay kit purchased from Caymen Chemical. After thawing, 20 μl of plasma from each sample were pipetted into 1 ml plastic eppendorfs. The volume was brought up to 200 μl with distilled water and the protein precipitated with 200 μl of 20 % TCA and centrifuged for 1 min at 13 000 g at room temperature to remove the supernatant. Then, 500 μl 2,4-dinitrophenyl hydrazine (in 2 M-HCD) were added to each sample, which was subsequently vortexed to resuspend the sample in the solution. The reaction was allowed to proceed for 15 min with occasional vortexing. At the end of the reaction, each sample was re-precipitated with 500 μl TCA vortexed and centrifuged for 1 min at 13000 g. The supernatant was then discarded and the sample was drained by touching the lip of the inverted tube to a clean Kim-wipe. Each sample was washed three times with 1 ml of 1:1 ethanol–ethyl acetate, vortexing each time and centrifuging for 5 min at 16 000 g between washes. After the final wash, the samples were left to briefly air dry. Then, 1 ml of 6 M-guanidine, 500 mM-KPO4 (pH 2·5), was used to redissolve/resuspend each sample with vortexing. The samples were then added to a ninety-six-well plate and read at 370 nm for carboxyl and 276 nm for total protein estimation. The intra/inter-assay CV were <5 and <9 %, respectively.

**Hydrogen peroxide.** H2O2 was measured spectrophotometrically using a commercially available kit (Sigma Chemical Company). After thawing, the serum sample was diluted 1:64 by adding 5 μl of serum to 315 μl of the sample diluent. Then, 50 μl of this diluted sample were added to a ninety-six-well plate in triplicate along with an appropriate standard curve (part no. H2163) and blanks in addition to 100 μl of H2O2 colour reagent (part no. H2288). After 30 min of incubation at room temperature, the plate was read on a Dynatech MRX 650 plate reader with Revelation software (American Instrument Exchange) using 550 nm as the primary wavelength. The intra/inter-assay CV were <3 and <6 %, respectively.

**Lipid-soluble antioxidants.** Plasma retinol, α-tocopherol, γ-tocopherol, β-carotene, lycopene and xanthophyll were analysed by simultaneous determination using the HPLC method of Thurnham et al.(22). Tocopherol acetate was used as the internal standard during this procedure. Briefly, 200 μl of the internal standard were added to each extraction while recovery was measured by injecting 50 μl tocopherol acetate directly on HPLC with each batch of samples. The internal standard was used to monitor sample recovery and the results were adjusted accordingly. Then, 1000 μg of tocopherol acetate were added to 100 ml of heptane and magnetically stirred. The stock solution (1 ml) was then diluted in 249 ml ethanol (95 %) and stored at 4°C until use. Thereafter, 100 μl of 10 mM-SDS were added to 1 ml of the plasma sample and mixed. Then, 200 μl of the internal standard were added to each sample along with two blanks (for recovery). Those were then vortexed for 1 min and 1 ml heptane (containing 50 mg butylated hydroxytoluene) was added to each sample and vortexed for 4 min. The samples were centrifuged for 10 min at 10 000 g at 10°C. Thereafter, 700 μl of the supernatant were discarded and the samples were left to dry under N2 in a fume cupboard. Each sample was reconstituted with 100 μl of the mobile phase (750 ml acetonitrile, 200 ml methanol, 50 ml dichloromethane, 500 μg butylated hydroxytoluene and sonicated for 30 min to de-gas) and vortexed. The samples were dispensed into HPLC vials and placed on sampling carousel for analysis. For the analysis, the samples were combined with the mobile phase at a flow rate of 1·5 ml/min at 21°C. The HPLC system comprised a Waters 510, Waters 2996 photodiode array detector, connected to a Waters 717 autosampler and a Waters SunFire™ C18, 3·5 μm, 4·6 × 100 mm column. The temperature of the column was maintained using a Jones chromatography 7900 series HPLC column oven. Samples were analysed using a range of wavelengths (325–450 nm) to detect retinol, tocopherol and carotenoid peaks. The concentration of retinol, α-tocopherol, γ-tocopherol, β-carotene, lycopene and xanthophyll in each sample was corrected in relation to the percentage of the internal standard recovered for each sample. The intra/inter-assay CV were <6 and <10 %, respectively.
Statistical analysis

Statistical analysis was performed using the SPSS statistics package (version 15.0; SPSS). A prospective calculation of power was performed using the equations of Altman(23) and based on experimental DNA data by Gill et al(19). Data were analysed using parametric statistics following mathematical confirmation of a normal distribution using Shapiro–Wilk tests. Baseline, pre- and post-intervention data were analysed using a two-way (A X B) mixed ANOVA which incorporated one between-subject (group: watercress v. control) and one within-subject factor (state: rest v. exercise). When a significant interaction effect was detected, within-participant factors were analysed using Bonferroni-corrected paired-sample t tests. Between-participant differences were analysed using a one-way ANOVA with an a posteriori Tukey’s honestly significant difference test.

The α level was established at P<0.05 and all values are reported as means and standard deviations, unless otherwise stated.

Results

Compliance

A total of ten participants (100%) completed the chronic phase of the study, and compliance to supplementation throughout was 98% (548 consumed doses from 560 possible doses). The ten participants also completed the acute phase (100%) and compliance with the supplementation protocol was 100% (ten consumed from ten possible doses).

Indices of oxidative stress

DNA damage increased following exercise in both control conditions (chronic-phase increase of 61% from rest, P<0.05 v. supplementation; acute-phase increase of 63% from rest, P<0.05 v. supplementation; Table 2), however, watercress consumption attenuated DNA integrity following exercise in both experimental phases (chronic phase, P<0.05 v. control; acute phase, P<0.05 v. control; Table 2). Similarly, there was a rise in LOOH from rest to exercise in both control conditions (chronic-phase increase of 15% from rest, P<0.05 v. supplementation; acute-phase increase of 16% from rest, P<0.05 v. supplementation; Table 2), which was conversely decreased as a function of watercress supplementation (chronic phase, P<0.05 v. control; acute phase, P<0.05 v. control; Table 2). PC concentration remained unchanged following exhaustive exercise in both chronic and acute supplementation phases (P>0.05; Table 2).

Reactive oxygen species

There was a comparatively higher production of H2O2 as a function of exhaustive exercise in both control conditions (chronic phase, P<0.05 v. supplementation; acute phase, P<0.05 v. supplementation; Fig. 3), while watercress ingestion decreased H2O2 concentration in both experimental groups (chronic phase, P<0.05 v. control; acute phase, P<0.05 v. control; Fig. 3).

Lipid-soluble antioxidants

Table 3 shows the effect of watercress and exercise on plasma antioxidant concentration. There was an increase in α-tocopherol concentration following chronic watercress ingestion (P<0.05 v. control), while α-tocopherol, γ-tocopherol and xanthophyll all increased in the acute phase (P<0.05 v. control). Following exhaustive exercise, α-tocopherol increased in both acute watercress and control states (P<0.05 v. rest); however, exercise decreased α-tocopherol in the chronic watercress group (P<0.05 v. control). Exercise also lowered γ-tocopherol concentration across the watercress and control groups for both phases (P<0.05 v. rest) and β-carotene decreased in the acute and chronic watercress groups (P<0.05 v. control), with no change observed in the control state (P>0.05).

Discussion

Strong experimental evidence indicates that exhaustive exercise may activate the production of free radical species and cause damage to important biological molecules such as DNA, lipid...
Exercise-induced oxidative stress

Previous work from our laboratory and, indeed, others demonstrates that intense exhaustive exercise can increase damage to circulating lymphocyte DNA and lipid membranes \(^{(10,24)}\). The comet assay is a sensitive, valid and versatile tool for the measurement of single- and double-strand breaks in DNA \(^{(20)}\) and has been previously used to detect DNA damage following exhaustive exercise \(^{(8,10,25–27)}\). Moreover, many of these reports attribute an increase in free radical production as the main cause of exercise-induced DNA damage; however, the exact mechanism(s) for the production of primary free radical species during exercise remains elusive, although free radicals generated within the skeletal muscle are often proposed as the main protagonist \(^{(20)}\). Evidence from the present investigation portrays a potential primary mechanism of cellular exercise-induced DNA damage which may be mediated by \(\text{H}_2\text{O}_2\). The up-regulation of leucocytes and neutrophil enzyme activation is known to cause an increased release of superoxide \(^{(30,31)}\), and superoxide has the capacity to directly damage DNA \(^{(32)}\) and/or be dismutated to \(\text{H}_2\text{O}_2\) by superoxide dismutase \(^{(33)}\). \(\text{H}_2\text{O}_2\) can readily diffuse across cell membranes \(^{(20)}\), and depending on the availability of transition metal ions, hydroxyl radicals may be generated which can cause single- and double-strand breaks in DNA and base-pair modification \(^{(34)}\). The change in \(\text{H}_2\text{O}_2\) as a function of exercise in both control groups provides a tentative explanation for DNA damage observed. An elevated \(\text{H}_2\text{O}_2\) concentration may also be a direct mediator of DNA damage through the disruption of DNA repair pathways \(^{(36)}\). Furthermore, \(\text{H}_2\text{O}_2\) can be generated from a number of extracellular pathways during exercise and may also be independent of superoxide formation. The auto-oxidation of neurological chemicals such as catecholamine has been shown to increase following exercise resulting in \(\text{H}_2\text{O}_2\) formation \(^{(37)}\). In addition, the oxidation of dopamine by monoamine oxidase releases \(\text{H}_2\text{O}_2\) as a metabolic by-product \(^{(38)}\).

Superoxide and hydroxyl radical production may also initiate vascular cell membrane lipid peroxidation, which may generate LOOH and facilitate further DNA damage. PUFA decomposition can produce an array of metabolic by-products including mutagenic compounds and lipid peroxidation intermediates \(^{(37)}\), including alkoxyl free radicals \(^{(39)}\) and malondialdehyde \(^{(39)}\). Both molecules are potentially capable of directly damaging DNA, while alkoxyl free radicals, in particular, may perpetuate the process of lipid peroxidation \(^{(37)}\). Unsaturated alkenals are also by-products of lipid peroxidation, which can lead to the production of PC \(^{(39)}\). PC are normally produced following the oxidation of amino acids and have previously been used as a sensitive measure of protein damage during high-intensity exercise \(^{(40)}\). Exhaustive exercise has been found to have no effect on protein oxidation throughout any condition, which perhaps suggests that protein oxidation has a greater affinity with exercise duration rather than intensity \(^{(41,42)}\).

Prophylactic effect of watercress

Watercress is a cruciferous vegetable of the \textit{Brassica} variety and contains, per gram weight, one of the highest concentrations of glucosinolates and carotenoids of any vegetable \(^{(43)}\). Glucosinolates can be hydrolysed by the enzyme myrosinase (released from plant cells during the chewing process) to form isothiocyanates which have been shown to possess anticarcinogenic properties and reduce DNA damage properties in both animal models and humans \(^{(44)}\). Cellular exposure to isothiocyanates

### Table 3

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<th>H(_2)O(_2) concentration (mmol/l)</th>
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*Within-group difference \((P< 0.05 \text{ v. rest})\); †between-group difference \((P< 0.05 \text{ v. rest})\).
Table 3. Plasma lipid-soluble antioxidants at baseline, rest and exercise for the watercress-supplemented and control groups for both experimental phases

(Mean values and standard deviations)

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<tr>
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<td>Baseline (mmol/l)</td>
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* Between-group differences (P<0·05).
† Within-group difference (P<0·05).
‡ Change from baseline to rest.
§ Change from rest to exercise.

Watercress ingestion attenuates DNA damage.
and antioxidants may represent a possible mechanism for the observed protection of cell DNA following both chronic and acute supplementation phases. LOOH also reduced following exercise in both supplemented conditions, providing further evidence of the additional cellular protection provided by the constitutes of watercress. In the present investigation, serum H2O2 decreased following exhaustive exercise under both supplemented conditions but increased in the control groups. These data would lend support to the suggestion that watercress may provide effective in vivo protection against H2O2 production as a function of exercise. It is also plausible that the elevated lipid-soluble antioxidants (under both supplemented protocols) are directly scavenging superoxide and therefore result in a net decrease in H2O2 production \(^ {45} \). The observed increase in lipid-soluble antioxidants, as that demonstrated following exercise, may also play a key role in the protection against cell membrane lipid peroxidation.

Watercress has been shown to increase important lipid-soluble antioxidants and, in particular, the tocopherols and carotenoids. Gill et al. \(^ {15} \) identified a 33% increase in β-carotene, a 26% increase in α-tocopherol and a 35% increase in ascorbic acid, thus highlighting watercress’s potential to elevate lipid- and aqueous-soluble systemic antioxidants. In contrast, we observed no change in β-carotene or retinol under either experimental condition. However, the lipid-soluble antioxidants α-tocopherol, γ-tocopherol and xanthophyll were all elevated following supplementation with watercress, and, in doing so, indicates the potential for watercress to act as a source of blood-rich antioxidants. Xanthophyll is an effective scavenger of singlet oxygen \(^ {46} \) and has shown antioxidant activity that is particularly effective in protecting DNA \(^ {47} \). The increased concentration of xanthophyll following the acute dose of watercress may therefore have played a contributory role in the increased protection of lymphocyte DNA in this supplemented group. Moreover, an increase in lipid-soluble antioxidants occurred as a function of exercise and is perhaps as a result of membrane-bound antioxidants being released into the vascular medium during the lipolysis of adipose tissue \(^ {48} \). The notion that exhaustive exercise may increase systemic levels of lipid-soluble antioxidants comes from a plethora of evidence suggesting that an increased mobilisation of fatty acids may give rise to α-tocopherol \(^ {24,49,50} \) in addition to other fat-soluble antioxidants such as retinal and lycopene \(^ {51} \).

PC are generated following the oxidation of amino acids and have previously been used as a sensitive measure of protein damage during high-intensity exercise \(^ {40} \). However, we observed no change in PC following exhaustive exercise with or without watercress ingestion, and this is in contrast to the observed DNA damage and lipid peroxidation. As previously reported, protein oxidation may have a greater affinity with exercise duration rather than intensity \(^ {41,42} \), and it is also conceivable that muscle damage may be a potential mediating factor in protein oxidation following exercise.

**Conclusion**

The present study suggests that dietary watercress intervention provides effective protection against exercise-induced oxidative stress, with no additional benefits gained from long-term watercress consumption. The prophylactic effect of watercress may be due in part to the high concentration of antioxidants or other cytoprotective compounds.

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**References**

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