Plasma irisin in runners and nonrunners: no favorable metabolic associations in humans

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Abstract
Irisin is a hormone which mimics the favorable metabolic effects associated with regular exercise, by converting subcutaneous white fat into brownish fat, in rodents. Thirty-three human subjects (16 runners, 17 nonrunners) were measured for: resting energy expenditure (REE), body composition, VO2 Peak test, [irisin], and plasma metabolic profile. Nine female nonrunners then participated in a 10-week supervised 5 km training program and tested after the race. Two runners underwent 18F-FDG-PET scans to quantify brown fat. No gender or age (28 ± 10 years) differences noted between matched cohorts. Runners averaged 58 ± 26 miles/week for 13 ± 6 years and had lower body-weight (63 vs. 88 kg; \( P < 0.001 \)), BMI (21 vs. 30 kg/m²; \( P < 0.0001 \)), triglycerides (58 vs. 123 mg/dL; \( P < 0.01 \)), total (white) fat (14 vs. 32%; \( P < 0.0001 \)), and had higher VO2 Peak (63 vs. 34 mL/kg-min; \( P < 0.0001 \)) and HDL (65 vs. 48 mg/dL; \( P < 0.01 \)) compared with nonrunners. [Irisin] was lower in runners versus nonrunners both before (179 vs. 197 ng/mL; NS) and after (207 vs. 226 ng/mL; NS) the VO2 Peak test. Significant \( (P < 0.05) \) positive correlations were noted between [irisin] versus BMI \( (r^2 = 0.15) \), triglycerides \( (r^2 = 0.40) \), and total body fat \( (r^2 = 0.24) \) with a significant negative correlation between [irisin] versus respiratory quotient \( (r^2 = 0.33) \). Total lean mass significantly correlated with REE \( (r^2 = 0.58) \) while total fat mass inversely correlated with VO2 Peak \( (r^2 = 0.64) \). Nonrunners had lower [irisin] after completion of the training program (194 vs.197 ng/mL; pre- to post-training; \( P > 0.05 \)). Neither runner selected for 18F-FDG-PET scans had brown fat. Runners demonstrated significantly healthier metabolic and body composition profiles compared with nonrunners. None of these favorable exercise effects were positively associated with [irisin].

Introduction
Habitual running reduces disability compared with nonrunning controls, as demonstrated by a longitudinal study that tracked runners over two decades (Chakravarty et al. 2008). Furthermore, runners demonstrated 30% lower adjusted all-cause mortality and 45% lower cardiovascular mortality rates compared with nonrunning controls, as documented in a prospective study involving 55,137 adults (Lee et al. 2014a). Unfortunately, half of all participants who commence regular exercise programs drop out within the first 6 months and never attain the health benefits associated with sustained aerobic fitness (Herring et al. 2014). Therefore, with the current (2011–2012) National Health and Nutrition Evaluation Survey data estimating the prevalence of obesity for US adults at 34.9% (68.5% are overweight and obese) (Ogden et al. 2014), recent discovery of the elusive "exercise hormone", irisin, has generated avid scientific, medical and pharmaceutical interest.

Irisin is a circulating 112 amino-acid extracellular dimer fragment of fibronectin type III domain containing protein 5 (FNDC5) that signals the conversion of white subcutaneous fat into metabolically active beige fat (Bostrom...
et al. 2012). Exercising muscle activates peroxisome proliferator-activated receptor γ coactivator-1α (PGC-1α) that then triggers FND5C protein expression within muscle cell membranes (Bostrom et al. 2012). Subsequent proteolytic cleavage of the N-terminal extracellular FND5C dimer fragment (irisin) into the circulation then triggers the “browning” of subcutaneous fat via upregulation of uncoupling protein 1 (UCP1). It is hypothesized that this “browning” of storage fat explains the favorable body composition and metabolic profiles characteristic of regular exercisers, such as endurance runners, via augmentation of resting energy expenditure (REE) and improvement of diet-induced insulin resistance.

To date, the irisin messenger sequence between muscle and fat tissue appears valid in rodents, but equivocal in humans. Bostrom et al. (2012) documented a twofold increase in [irisin]p in eight healthy middle-aged men after 10 weeks of supervised aerobic training. However, this increase in basal [irisin]p has yet to be supported by independent investigators who conversely report either a decrease (Hecksteden et al. 2013; Norheim et al. 2014) or no change (Kurdiova et al. 2014) in [irisin]p following aerobic (Hecksteden et al. 2013; Kurdiova et al. 2014; Norheim et al. 2014) and strength (Hecksteden et al. 2013) training programs. These conflicting human data are thought to result from: lack of validation between commercially available assays used by independent laboratories (Sanchis-Gomar et al. 2014); mutation of the FND5C start codon in humans compared with rodents (Raschke et al. 2013); ambiguity with regards to quantitation of extracellular FND5C protein after cleavage of irisin (Erickson 2013); and/or evidence suggesting that [irisin]p is an adipokine (in addition to a myokine) (Roca-Rivada et al. 2013). Cross-sectional human studies also paradoxically demonstrate positive correlations between [irisin]p and fat mass (Stengel et al. 2013; Crujeiras et al. 2014), body mass index (BMI) (Huh et al. 2012; Park et al. 2013; Stengel et al. 2013; Crujeiras et al. 2014), insulin resistance (Park et al. 2013), and triglycerides (Park et al. 2013). Thus, the favorable body composition and metabolic adjustments from exercise-induced irisin secretion and augmentation of REE have yet to be convincingly validated in humans.

We wished to extend the current spectrum of understanding from unhealthy (Park et al. 2013; Stengel et al. 2013; Wen et al. 2013), middle-aged (Bostrom et al. 2012; Hecksteden et al. 2013; Park et al. 2013; Crujeiras et al. 2014), populations toward a healthier extreme: endurance runners. Thus, the purpose of this investigation was to: (1) Assess plasma irisin concentrations ([irisin]p) in well-trained runners and nonrunners before and after a VO2 peak test; (2) Examine relationships between [irisin]p versus REE, body composition, blood glucose, and lipid parameters; and (3) Assess changes in [irisin]p in nonrunners before and after a 5 km run/walk training program. We hypothesized that highly trained runners would epitomize the classic irisin phenotype, which would be characterized by: elevated [irisin]p, high REE, low fat mass, favorable lipid panel, and robust depots of metabolically active beige fat (detected through positron emission tomography scans). We further hypothesized that nonrunners would demonstrate improvements in body composition, metabolic parameters and REE following 10 weeks of supervised aerobic training that were associated with favorable changes in [irisin]p.

Materials and Methods

Subjects

Thirty-three healthy (no chronic medical condition requiring regular prescription medication) habitual runners (>50 km/week running for >3 months) and nonrunners (<60 min endurance activity for >3 months) between the ages of 18–50 years were recruited to participate in this trial. Sixteen runners (eight male, eight females) were age and gender-matched with nonrunners (eight males, nine females). All female runners were tested during the follicular phase of their menstrual cycle. Informed written consent was obtained prior to participation. Phase I and II protocols were approved by Oakland University’s Institutional Review Board (IRBNet #394700 and #471164, Phase I and II, respectively). Two runners (one male, one female) then underwent pilot cold-stress18F-fluorodeoxyglucose positron emission/computed tomography (18F-FDG-PET/CT) scans as part of a separate research study conducted at the Children’s Hospital of Michigan PET Center and approved by Wayne State University’s IRB (Brown Fat PET study).

Experimental protocols

Phase I: runners versus nonrunners

All 33 participants presented to the laboratory after a minimum 4-h fast. Weight and height were measured using a weight beam eye-level physicians scale (Detecto, Webb City, MO). REE was measured via indirect calorimetry, using a metabolic canopy (VIASYS Vmax Encore, CareFusion, Loma Linda, CA). For the REE test, subjects were allowed 5–10 min to achieve steady-state while resting comfortably in a recumbent position in a dimly lit room. REE and respiratory quotient (RQ) were averaged over a 15-min assessment period. Next, in this recumbent position, 10 mL of venous blood was withdrawn from an antecubital vein for pre-exercise measurement of plasma
lipids, glucose, [irisin]p, and protein. After venipuncture, body composition was assessed using a dual energy X-ray absorptiometry scan (Hologic Discover A, Boston, MA). A treadmill running test (VO2 Peak) was then conducted to determine aerobic fitness (TrueOne 2400, Parvo Medics, Sandy, UT). For the VO2 Peak test, runners started at an easy jog while nonrunners started at a comfortable walking speed. After 1 min, the treadmill speed increased 0.5 mph every minute until subjects could no longer keep pace with the treadmill (volitional exhaustion). After completion of the VO2 Peak treadmill test, all subjects returned to the recumbent position to allow re-equilibration of body fluid compartments. After 15 min, another 10 mL of venous blood was withdrawn for postexercise assessment of plasma lipids, glucose, [irisin]p, and protein. Room temperature was maintained between 22–24°C.

Phase II: training trial
Within 5 days of baseline Phase I testing (above), nine female nonrunners participated in a 10-week supervised run/walk 5 km training program. Phase II participants met three times per week and followed a freely available online training program designed to prepare beginners to finish a 5 km race comfortably (http://www.active.com/running/articles/how_to_run_your_first_5k?page=2). With in 5 days of completion of the target 5 km race, all Phase II runners repeated the Phase I testing protocol.

Biochemical measures
Total cholesterol, high density lipoproteins (HDL), triglycerides, blood glucose, aspartate transaminase (AST), and alanine transaminase (ALT) were measured in whole blood (low density lipoproteins [LDL] and very low density lipoproteins [VLDL] were calculated) within 10 min of venipuncture, using a Lipid Panel Plus cartridge/PicoXpress analyzer (Abaxis, Union City, CA). Irisin was measured in aprotinin-treated plasma, using a commercially available enzyme immunoassay kit (EK-067-29, Phoenix Pharmaceuticals, Burlingame, CA). The Phoenix Pharmaceutical ELISA kit has been previously validated against western blotting (Wen et al. 2013) and mass spectrometry (Lee et al. 2014b), with a detection range between 0.1 and 1000 ng/mL. All plasma was stored at −80°C until analysis could be performed as a single batch, within 9 months of collection. Changes in plasma volume (PV) were estimated by comparing pre- and post-VO2Peak measurements of plasma protein using a clinical refractometer (Schuco Clinical Refractometer 5711-2020, Tokyo, Japan) according to a method described previously (Stricker 1968).

Cold-stress fludeoxyglucose (18F) positron emission tomography/computed tomography (18F-FDG-PET/CT) scans
Thermoregulatory challenge was applied using a specialized whole-body garment through which subjects were exposed to a cold temperature stimulus. To achieve maximal stimulation, each subject was cooled until he or she was close to shivering and then the skin temperature was raised by 0.5°C and held steady thereafter. The tube suit cooling garment incorporated a network of small-diameter plastic tubing (Allen Vangard, Inc., Ottawa, CA) through which temperature-controlled cold water (2–4°C) was circulated. The effects of these exogenous temperature stressors on body temperature was monitored using a GaAs crystal sensor located at the tip of an optical fiber cable (OpSense, Inc., Quebec City, CA). This approach relies on the temperature dependence of the energy band gap of GaAs semiconductor crystal. The GaAs sensor is opaque for wavelengths below the bandgap and transparent for wavelengths above the energy band gap. The sensor was taped to the skin at the location of the left rib cage. This location was selected on the basis of proximity to important anatomical features (close to the pulmonary blood vessels which are possibly the most representative sites for body core temperature) and the ability to consistently place the sensors based on those anatomical landmarks.

Activated brown adipose tissue (BAT) in supraventricular fat depots was considered present if there were areas of tissue that were more than 5 mm in diameter, had the CT density of adipose tissue (−250 to −50 Hounsfield units [HU]), and had a maximal standardized uptake value (SUV) of FDG of at least 2.0. This cutoff represented more than 2 SD above the maximal SUV seen in typical depots of white adipose tissue. BAT volume was determined by thresholding both the CT image volume (−250 < HU < −50) and the FDG volume (SUV > 2.0) and then applying the logical AND operation to the two masks, followed by removal of all areas that were smaller than 0.125 cm³. Using this definition of activated BAT, no voxel survived the thresholding operation. The PET protocol and quantification of brown adipose tissue has been described previously (Muzik et al. 2013).

Statistical analyses
Unpaired t-tests were chosen to express differences between group (runners vs. nonrunners) and gender (male vs. female) within each group, after 2-way ANOVA tests revealed no significant interaction effects between group x gender for any measured variable (Phase I testing). Paired t-tests were utilized to evaluate pretraining
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versus post-training changes in those nonrunners who completed the 10-week run/walk training program (Phase II). Linear regression analyses (Pearson’s) were performed to assess relationships between variables. Statistically significant alpha level was set a priori at $P < 0.05$. All data presented as mean ± SD.

Results

Phase I: runners versus nonrunners

Eight male (109 ± 42 km/week; 11 ± 5 years running) and eight female (76 ± 38 km/week; 14 ± 9 years running) runners were age and gender-matched with nonrunners. When males and females were combined, runners demonstrated significantly lower body weight and total body fat along with higher aerobic fitness (VO2 Peak) compared to nonrunners (Table 2). There were gender differences in both the runner and nonrunner groups with respect to total lean tissue, % total body fat, bone mineral content (BMC), and measured REE (Table 1).

For the biochemical measurements, runners as a combined group (males and females) demonstrated significantly higher pre-VO2Peak HDL, AST and post-VO2 Peak blood glucose levels as well as lower pre-VO2 Peak triglycerides and VLDL when compared to nonrunners (Table 2). There were no significant differences in pre-VO2 Peak (resting) or Post-VO2 Peak (stimulated) [irisin]p, with a trend for higher [irisin]p in nonrunners compared to runners. Nonstatistically significant exercise-induced increases in [irisin]p were shown in both runners and nonrunners following the VO2 Peak test (Δ: postexercise minus pre-exercise), even when corrected for plasma volume change (Table 2).

When all data were combined ($N = 33$), resting [irisin]p was positively correlated with BMI, total cholesterol, VLDL3, triglycerides, trunk fat (Table 3), and total fat mass (Fig. 1A), while negatively correlated with RQ (Fig. 1C). Total lean mass was positively correlated with measured REE (Fig. 1D). Total fat mass was positively correlated with REE (Fig. 1D), BMI, VLDL3, and triglycerides while negatively correlated with HDL (Table 3) and VO2 Peak (Fig. 1B).

Phase II: training trial

Nine female nonrunners successfully completed the supervised 10-week run/walk 5 km training program. Two male nonrunners started the training program but dropped out within the first 4 weeks (82% completion rate overall). Although the average trend favored a net ~2 kg weight loss over the training period, six females lost weight (Losers) while three females either gained or maintained weight (Gainers). [Irisin]p tended to decrease following the training program, when compared to pretraining levels, despite (nonsignificant) gains in aerobic capacity. This pre- to post-training decrease in [irisin]p was noted both before and after the VO2 Peak test, regardless of weight loss or gain. There were no statistically significant correlations noted between resting [irisin]p versus the change in either lean mass or fat mass immediately post-training (Fig. 2A) or when expressed as a change (Fig. 2B). There was a significant difference in post-training pre-VO2 Peak [irisin]p between Losers versus Gainers (169.7 vs. 204.9 ng/mL; $P < 0.001$, respectively) with the Gainers demonstrating higher [irisin]p levels before and after the training program. The only other statistically significant difference between pretraining versus post-training variables or between Losers versus Gainers was seen in ALT, which decreased more in Losers compared to Gainers after the training program (Table 4).

18F-FDG-PET/CT scans

One female and one male runner underwent cold-stimulated 18F-FDG-PET/CT scans, to assess brown adipose tissue activation as part of a separate project. Possible activation of brown adipose tissue was assessed based on an FDG standard uptake value (SUV) threshold (SUV > 2.0) and presence of adipose tissue as determined by CT (HU < −50). No detectable activated brown fat was noted in either runner 1 (Fig. 3A: 48 year old female, 30 years running, 50 km/week running, SUVmax = 1.5, VO2 Peak 49 mL/kg-min, BMI 17 kg/m2, body fat 14%, [irisin]p 217 ng/mL pre and 229 ng/mL postexercise) or runner 2 (Fig. 3B: 24 year old male, 10 years running, 174 km/week running, SUVmax = 0.7, VO2 Peak 84 mL/kg-min, BMI 22 kg/m2, body fat 10%, [irisin]p 166 ng/mL pre-exercise and 192 ng/mL postexercise). The reported SUVs correspond to the location of supraclavicular fat depots defined based on HU values of (−250 < HU < −50).

Discussion

These data do not support any positive association between [irisin]p with favorable body composition profiles, healthy blood lipid parameters, or cold-stimulated brown fat activation in fit and unfit humans. Furthermore, there was a tendency for [irisin]p to decrease following a 10-week supervised run/walk training program, despite increases in aerobic fitness in our cohort of healthy, young, and overweight females. This lack of a positive [irisin]p response has been similarly reported in other prospective cohort training studies conducted in healthy humans (Hecksteden et al. 2013; Kurdiøva et al. 2014; Norheim et al. 2014). To date, only the original
Table 1. Demographic, body composition, aerobic fitness, and resting metabolism measurements obtained for both runners and nonrunners.

| Variable (units) | Runners | Nonrunners |
|------------------|---------|------------|
|                  | Male (min–max) | Female (min–max) | Combined (min–max) | Male (min–max) | Female (min–max) | Combined (min–max) |
|                  | n=8     | n=8        | n=16          | n=9     | n=9        | n=17          |
| Age (years)      | 30.1±11.0 (19–45) | 30.4±11.6 (20–48) | 30.3±10.9 (19–48) | 29.8±9.9 (20–48) | 23.4±6.9 (18–41) | 26.4±8.8 (18–48) |
| BMI (kg/m²)      | 22.4±2.4 (20–26)  | 20.3±1.7 (17–23)  | 21.4±2.3 (17–26)  | 29.8±7.5 (20–43) | 30.6±4.9 (25–41) | 30.2±6.0 (20–43)  |
| Weight (kg)      | 69.9±8.4 (61–85)   | 56.6±8.3 (43–68)   | 63.2±10.6 (43–85) | 95.5±23.5 (63–126) | 81.6±17.7 (67–125) | 88.1±21.2 (63–126) |
| Total Fat (%)    | 6.9±2.4 (4–11)    | 10.8±3.6 (6–15)    | 8.8±6.6 (4–15)    | 26.1±14.2 (12–47) | 30.0±9.9 (22–54) | 28.1±11.9 (12–54) |
| Tissue (kg)      | 59.5±6.5 (54–72)   | 43.3±5.4 (35–50)   | 51.4±10.2 (35–72) | 65.2±10.0 (47–81) | 47.8±7.3 (41–65) | 56.0±12.3 (41–65) |
| Bone Mineral     | 2.5±0.2 (2–3)     | 1.97±0.23 (1.8–2.4)| 2.26±0.39 (1.8–3.0)| 2.6±0.37 (2.2–3.4) | 2.14±0.31 (1.7–2.6)| 2.36±0.41 (1.7–3.4) |
| Content (kg)     | 9.8±2.7 (7–15)    | 18.9±4.0 (12–24)   | 14.4±5.0 (7–24)   | 26.1±8.2 (18–39)  | 36.9±3.8 (32–44)  | 31.8±8.2 (18–44)  |
| VO₂ Peak (ml/kg-min) | 70.5±9.6 (57–83)  | 53.6±5.1 (47–59)   | 62.6±11.5 (47–83) | 37.3±8.9 (25–48)  | 30.8±5.2 (24–39)  | 33.9±7.7 (24–48)  |
| Peak Treadmill   | 13.2±1.8 (11–16)  | 10.4±1.4 (9–12)    | 11.8±2.0 (9–16)   | 7.6±1.8 (5–10)    | 6.7±0.9 (6–8)     | 7.1±1.4 (5–10)    |
| Speed (mph)      | 184.9±10.1 (170–197) | 188.6±8.1 (176–198) | 186.9±9.0 (170–198) | 191.1±19.0 (149–211) | 189.2±0.3 (172–205) | 190.1±14.1 (149–211) |
| Maximum Heart    | 1781.8±107.1 (1466–2027) | 1436.4±107.1 (1312–1556) | 1609.1±235.6 (1312–2027) | 2059.9±542 (1405–3129) | 1570.9±241.6 (1438–2065) | 1810.0±470.3 (1405–3129) |
| Rate (bpm)       | 0.80±0.2 (0.7–0.9) | 0.81±0.07 (0.7–0.9) | 0.80±0.06 (0.7–0.9) | 0.76±0.04 (0.7–0.8) | 0.78±0.09 (0.7–1.0) | 0.77±0.07 (0.7–1.0) |

**Note:** $^a$P < 0.05; $^{aa}$P < 0.01; $^{aaa}$P < 0.001 between runners versus nonrunners.

$^b$P < 0.05; $^{bb}$P < 0.01; $^{bbb}$P < 0.001 between males versus females within cohort.
Table 2. Blood chemistry variables measured pre-VO2 Peak test (resting), unless otherwise specified as post-VO2 Peak test (exercise stimulated) or the change (Δ: post-VO2 Peak test minus pre-VO2 Peak test value).

| Variable (units) | Runners Mean ± SD | Nonrunners Mean ± SD |
|-----------------|------------------|----------------------|
|                 | Male (min to max) | Female (min to max)  | Combined (min to max) |
|                 | n=8              | n=8                  | N=16                 |
| Cholesterol (mg/dL) | 166.0 ± 24.8 (137 to 217) | 176.8 ± 32.0 (144 to 227) | 171.4 ± 28.2 (137 to 227) |
| HDL (mg/dL)     | 59.4 ± 15.8 (38 to 76)      | 70.8 ± 12.5 (49 to 87)      | 65.1 ± 15.0 (38 to 87)  |
| LDL (mg/dL)     | 95.3 ± 22.4 (54 to 130)     | 94.3 ± 25.8 (62 to 137)     | 94.8 ± 23.4 (54 to 137) |
| VLDL (mg/dL)    | 11.3 ± 3.9 (7 to 19)        | 12.0 ± 3.0 (8 to 16)        | 11.6 ± 3.4 (7 to 19)    |
| Triglycerides   | 56.0 ± 19.5 (36 to 95)      | 60.4 ± 14.5 (39 to 79)      | 58.2 ± 16.8 (36 to 95)  |
| ALT (U/L)       | 31.9 ± 7.8 (25 to 44)       | 21.5 ± 9.6 (7 to 40)        | 26.7 ± 10.0 (7 to 44)   |
| AST (U/L)       | 41.5 ± 10.6 (25 to 59)      | 32.5 ± 5.8 (23 to 39)       | 37.0 ± 9.5 (23 to 59)   |
| Pre-VO2 peak glucose (mg/dL) | 96.0 ± 4.0 (89 to 101) | 96.4 ± 5.8 (87 to 105) | 96.2 ± 4.8 (87 to 105) |
| Post-VO2 peak glucose (mg/dL) | 133.8 ± 11.1 (117 to 153) | 125.8 ± 23.0 (96 to 193) | 129.8 ± 17.9 (96 to 193) |
| Pre-VO2 peak irisin (ng/mL) | 177.8 ± 33.1 (128 to 228) | 180.0 ± 41.0 (103 to 223) | 178.9 ± 36.0 (103 to 228) |
| Post-VO2 peak irisin (ng/mL) | 191.8 ± 13.9 (174 to 217) | 221.9 ± 30.9 (161 to 270) | 206.9 ± 27.9 (161 to 270) |
| Δ irisin (ng/mL) | 14.0 ± 23.5 (−13 to 53)    | 6.2 ± 6.0 (0 to 19)        | 3.4 ± 6.4 (−6 to 6)     |
| Δ Plasma volume (%) | 2.4 ± 17.2 (−19 to 27)      | 36.0 ± 57.0 (−21 to 154)    | 19.2 ± 44.2 (−21 to 154) |
| Δ irisin PV corrected (ng/mL) | 2.4 ± 17.2 (−19 to 27) | 36.0 ± 57.0 (−21 to 154) | 19.2 ± 44.2 (−21 to 154) |

*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001 between runners versus nonrunners.

bP < 0.05; b**P < 0.01; b***P < 0.001; b****P < 0.0001 between males versus females within cohort.
Bostrom et al. (2012) study has demonstrated a significant (two-fold) increase in circulating irisin concentrations following an aerobic training program in humans. Investigations targeting FNDC5 gene expression within human muscle following both a 10-week endurance training and an 11-week strength training program also failed to demonstrate increases in FNDC5 gene expression (Raschke et al. 2013). Cross-sectional studies revealed significant increases in muscle FNDC5 gene expression only in older (~55 years) active versus sedentary subjects (Timmons et al. 2012) and in heart failure patients (~67 years) with higher aerobic capacity, when subdivided into more fit (VO2 Peak ~17 mL/kg-min) versus less fit (VO2 Peak ~11 mL/kg-min) groups for statistical comparison (Lecker et al. 2012). Because circulating irisin is the cleaved, extracellular, polypeptide fragment of FNDC5, these upstream results strengthen a growing body of evidence suggesting that [irisin]p is unrelated to the favorable metabolic or body composition effects associated with regular endurance exercise in healthy humans.

The runners in this study were well-trained and highly fit, as demonstrated by mean peak oxygen uptakes (VO2 Peak) above the 90th percentile for both males (90th percentile: 55 mL/kg-min; our cohort: 71 mL/kg-min) (ACSM’s Guidelines for Exercise Testing and Prescription 2014a) and females (90th percentile 46 mL/kg-min; our cohort: 54 mL/kg-min) (ACSM’s Guidelines for Exercise Testing and Prescription 2014b). Of note, our male cohort included elite, professional, runners, with mean peak aerobic capacities representative of international competitors, such as the Swiss National Team (73 mL/kg-min) (Marti and Howald 1985). Conversely, the nonrunners in this study had peak oxygen uptakes below the 20th percentile for both males (20th percentile: 38 mL/kg-min; our cohort 20th percentile: 38 mL/kg-min; our cohort

| Variable                  | BMI (kg/m²) | Cholesterol (mg/dL) | HDL (mg/dL) | VLDL (mg/dL) | Triglycerides (mg/dL) | ALT (U/L) | Trunk fat (kg) |
|---------------------------|-------------|---------------------|-------------|--------------|-----------------------|-----------|----------------|
| Pre-irisin (ng/mL)        | 0.15*       | 0.26**              | 0.00        | 0.40****     | 0.40****              | 0.07      | 0.16*          |
| Total lean tissue (kg)    | 0.26**      | 0.00                | (−0.05)     | 0.06         | 0.06                  | 0.28**    | 0.15*          |
| Total fat tissue (kg)     | 0.88*****   | (−0.01)             | (−0.20**)   | 0.28**       | 0.28**                | 0.12      | 0.98****       |

*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001. (−) designates that the correlation is negatively directed (inverse).
37 mL/kg-min) (ACSM’s Guidelines for Exercise Testing and Prescription 2014a) and females (20th percentile: 31 mL/kg-min; our cohort: 31 mL/kg-min) (ACSM’s Guidelines for Exercise Testing and Prescription 2014b). However, despite the wide disparity in aerobic fitness levels between runners and nonrunners, all of our subjects were healthy by design, with fasting blood lipid and glucose levels within the normal range. The only exception was pre-training HDL levels in the female nonrunners (49 mg/dL), with an overall mean value just below the biochemical range of “poor” (<50 mg/dL) (Table 2).

Even amongst “healthy” individuals, the runners demonstrated superior metabolic health with significantly higher HDL, lower triglycerides, and VLDL compared to the nonrunners. The male and female runners were also exceptionally lean, with significantly lower total body fat mass when compared to nonrunners. With regards to our main outcome measure, [irisin]p, none of the favorable aerobic fitness, metabolic health, or body composition profile measurements were positively correlated with either resting (pre-VO2 Peak) or exercise stimulated (post-VO2 Peak) [irisin]p. In fact, our mean [irisin]p values tended to be lower in runners compared to nonrunners, both before and after the VO2 Peak test. Also in direct contrast with Bostrom et al. (2012) results, [irisin]p decreased in our cohort of female nonrunners immediately following 10-weeks of endurance training. Neither post-training [irisin]p or the pre- to post-training change in [irisin]p were linearly related to changes in total lean mass. This decrease in resting and stimulated [irisin]p occurred regardless of body weight loss or gain and despite nonsignificant increases in aerobic fitness and improvements in metabolic health (increased HDL and decreased triglycerides). Thus, our collective findings from Phase I and II trials were opposite of what was expected, assuming that irisin receptor number, sensitivity, and clearance rate were equivalent in runners and nonrunners before and after the training program. Additionally, resting [irisin]p was positively correlated with BMI, total cholesterol, triglycerides, VLDL, trunk, and total fat mass while negatively correlated with resting RQ. Similar relationships between [irisin]p versus increasing adiposity and poorer metabolic health have been verified previously (Huh et al. 2012; Park et al. 2013; Stengel et al. 2013; Crujeiras et al. 2014) and thereby strengthen growing support that [irisin]p neither mimics nor augments the beneficial effects of regular endurance exercise in humans.

Resting energy expenditure was not related to [irisin]p in our cohort of runners and nonrunners, as originally expected. Total lean mass explained 58% of the variance in REE. Nonrunners who gained weight after the 10-week training program (Gainers) demonstrated gains in lean tissue, which were accompanied by increases in REE. Although total lean mass was not linearly related to [irisin]p, on average, nonrunners tended to have greater lean mass, REE, and [irisin]p compared to runners, but these differences were not statistically significant. These trends in magnitude support irisin as a myokine, stimulated in both runners and nonrunners by high intensity exercise even after correcting for plasma volume change. Both submaximal exercise and shivering have also been shown to induce transient increases in [irisin]p, further supporting its myokine lineage (Huh et al. 2012; Kraemer et al. 2014; Lee et al. 2014b). However, although it appears that irisin is released by muscular contraction, plasma irisin levels do not seem to be linearly related with increases in REE, total lean mass or any beneficial metabolic or body composition effects associated with aerobic fitness.

Unlike total lean mass, significant positive relationships were noted between resting [irisin]p versus total fat mass and resting RQ. Total fat mass explained 21% of the variance in REE and positively correlated with BMI, triglycerides and VLDL, while negatively correlated with HDL.
and VO$_2$ Peak. For the nonrunners who completed the 10-week training program, there was a nonstatistically significant trend for post-training [irisin]$_p$ to be positively correlated with changes in total fat mass and not lean mass as originally hypothesized. Thus, our data from both Phase I and Phase II trials suggest that [irisin]$_p$ better reflects a white, not brown, adipose tissue lineage.

Lastly, because the metabolic and body composition benefits of irisin are related to the “browning” of subcutaneous white adipose tissue, we performed cold-stress $^{18}$F-FDG-PET/CT scans (Muzik et al. 2013) in a pair of runners (one female and one male) deemed most likely to exhibit robust depots of beige or brown fat. The female was lean (BMI = 17 kg/m$^2$), had been running for >30 years and had higher than average [irisin]$_p$ levels (217–229 ng/mL). The male was a professional runner, averaging >160 km/week running, and had the highest VO$_2$ peak of the cohort (84 mL/kg-min). Unexpectedly, neither runner had any detectable beige or brown fat activity within the regions of interest. Because cold-induced postganglionic sympathetic nervous system (SNS) release of norepinephrine activates UCP1 in brown fat, we hypothesized that athletes would have an augmented capacity to secrete adrenaline (“sports adrenal medulla”) (Zouhal et al. 2008). We then hypothesized that chronic, repetitive, SNS stimulation from daily running would enhance the browning of subcutaneous fat and lead to greater cold-stimulated brown fat activation in more seasoned runners, regardless of circulating plasma irisin levels. Circulating norepinephrine concentrations have

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**Table 4.** Changes (Δ: post-training minus pretraining) in body composition, aerobic fitness, resting energy expenditure, and blood chemistry variables for the nine female nonrunners who completed the 5 km training program (Phase II). “Losers” refer to those females who lost weight after the 10-week walk/run program (n = 6). “Gainers” refer to those females who maintained or gained weight after the 10-week walk/run program (n = 3).

| Variable (units) | Total (min to max) | Losers n=6 | Gainers n=3 |
|-----------------|-------------------|------------|------------|
| Δ Weight (kg)   | –1.8 ± 3.9 (–11 to 2) | –3.5 ± 3.9 | 1.4 ± 1.2  |
| Δ Total fat tissue (kg) | –1.9 ± 3.0 (–9 to 2) | –3.0 ± 3.0 | 0.3 ± 1.5  |
| Δ Total lean tissue (kg) | 0.2 ± 1.0 (–0.8 to 1.9) | –0.1 ± 1.0 | 0.9 ± 0.9  |
| Δ Bone mineral content (g) | –16.6 ± 26.2 (–55 to 39) | –12.2 ± 30.8 | –25.7 ± 13.8 |
| Δ VO$_2$ peak (ml/kg-min) | 2.2 ± 3.0 (–1 to 8) | 2.6 ± 3.2 | 1.3 ± 2.8  |
| Δ Measured REE (kcal/day) | 8.9 ± 175.6 (–182 to 254) | –14.8 ± 169.9 | 56.3 ± 214.8 |
| Δ Cholesterol (mg/dL) | 6.9 ± 15.3 (–14 to 31) | 17.7 ± 12.2 | 1.5 ± 14.6  |
| Δ HDL (mg/dL) | 3.3 ± 8.2 (–12 to 16) | 6.3 ± 9.5 | 1.8 ± 8.0  |
| Δ LDL (mg/dL) | 6.4 ± 11.2 (–16 to 20) | 9.7 ± 4.6 | 4.8 ± 13.5  |
| Δ VLDL (mg/dL) | –2.8 ± 6.6 (115 to 3) | –4.7 ± 7.5 | 1.0 ± 2.0  |
| Δ Triglycerides (mg/dL) | –14.6 ± 33.6 (–77 to 15) | –24.8 ± 37.5 | 6.0 ± 7.6  |
| Δ ALT (U/L) | –4.0 ± 6.8 (–18 to 4) | –7.2 ± 6.0* | 2.3 ± 2.1  |
| Δ AST (U/L) | –3.3 ± 8.7 (–18 to 8) | –4.2 ± 10.8 | –1.7 ± 2.1  |
| Δ Glucose (mg/dL) | 3.0 ± 4.4 (–4 to 12) | 3.0 ± 8.2 | 3.0 ± 1.9  |
| Δ Pre-VO$_2$ peak irisin (ng/mL) | –12.5 ± 36.2 (–29 to 61) | –17.0 ± 42.2 | –3.6 ± 24.3 |
| Δ Post-VO$_2$ peak irisin (ng/mL) | –44.4 ± 46.9 (–120 to 9) | –47.3 ± 55.7 | –39.7 ± 37.9 |

*P < 0.05 between responders and nonresponders.

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**Figure 3.** FDG PET/CT images of a female (A) and a male (B) runner show low FDG uptake in supraclavicular adipose tissue depots (white arrows). The maximal FDG SUV in these depots was determined as 1.5 in the female and 0.7 in the male runner.
also been shown to increase >20 times basal levels after a 400 m race and rise exponentially with increases in exercise intensity (Zouhal et al. 2008). Although we did not measure catecholamine's in this study, the maximum heart rates of 185–191 bpm seen in runners and nonrunners after the VO₂ Peak test indirectly imply strong SNS activation, well above previous administration of ephedrine in another study which failed to induce brown fat activation, with pharmacologically driven heart rate increases of only ~10 bpm (Cypess et al. 2012). Previous studies using cold-stress ¹⁸F-FDG-PET/CT also document high brown fat mass (>50 g) and activation in 96% of healthy males (van Marken Lichtenbelt et al. 2009) and 53% of females (Muzik et al. 2013). This high prevalence of brown fat in healthy humans makes our negative PET findings all the more curious, given the potential metabolic benefits of both running and brown fat (Ouellet et al. 2012; Chondronikola et al. 2014) which did not coexist in our two carefully chosen lean subjects.

**Limitations**

The wide variability in circulating irisin levels reported in this study - and in other published reports (Sanchis-Gomar et al. 2014) - represents the biggest limitation in the interpretation of [irisin]ₙ. Although the Phoenix Pharma-
ceutical ELISA kit has been previously validated against immunoblotting and mass spectrometry, there have been three generations of commercial assay kits (EK-067-16, EK-067-52, and EK-067-29). Our first samples were measured using two separate EK-067-52 kits, which differed in lot number. When a twofold difference was noted between the mean values from the separate kits, all samples were subsequently reanalyzed using the newest generation assay (EK-067-29). Linear regression (Fig. 4A) and Bland–Altman analyses (Fig. 4B and C) highlight the wide variability between the mean values (90 to ~89) for the different ELISA kits and underscores the inconsistencies in measurement and reporting of [irisin]p (Erickson 2013; Sanchis-Gomar et al. 2014). The coefficient of variation (CV) for each kit was fairly similar and seemed to increase with increasing subject numbers: (EK-067-29; N = 84; CV = 0.24. EK-067-52A; n = 68; CV = 0.22; EK-067-52B; n = 16; CV = 0.18).

Another potential limitation was our 4-h fasting period, which may not have been adequate to obtain truly fasted lipid and REE values. We choose the 4-h fasting interval from a previous investigation (Haugen et al. 2003) showing that RMR differed by ~100 kcal/day when compared to an overnight fast. We also analyzed the REE data using the kcal/day “correction” factor and found that these data were not significantly altered between the subjects who participated in an overnight versus 4-h fast (n = 7; 21%; data not shown).

Perspectives and significance

The lack of a positive relationship between exercise, aerobic fitness, circulating irisin and the burning of fat highlights the potential difficulty translating evidence obtained in murine models directly to humans. Exercise training of healthy male Yucatan miniature swine pigs similarly did not elicit significant changes in muscle FNDC5 or circulating irisin levels after ~16–20 weeks of regular treadmill running (Fain et al. 2013). Genomic sequencing reveals that humans contain an alternative start codon for FNDC5 (ATA) while rodents bear the highly conserved ATG start codon (Raschke et al. 2013). As such, annotated noncanonical ATA start sites demonstrate low-translation efficiency. Only three human genes with an ATA start codon have been previously identified to translate full length proteins. In vitro data verify that only 1% of the full length FNDC5 protein is translated with an ATA start, which may explain the discrepancies between human versus rodent [irisin]p data (Raschke et al. 2013). Unfortunately, swine were not included in these multispecies FNDC5 sequence alignments to corroborate these negative human and pig exercise training data.

In addition to the potential inconsistencies when translating animal data to human models, the vast majority of human exercise interventions target disease (inactive) and/or normal (active) populations. Investigations targeting exceptionally healthy populations, such as endurance athletes, are often overlooked when evaluating the favorable physiological adaptations associated with regular physical activity. While we typically construct our physiological understanding of dysregulation starting with disease (chronic inactivity), a simultaneous deconstruct of highly fit individuals (chronic physical activity) would anchor both sides of the spectrum. The athletes profiled in this investigation have maximized their physiological capacity to effectively respond to daily bouts of intense homeostatic challenge. As such, we feel that our Phase I and Phase II exercise testing sequence provides a useful “reality check” in the evolution of molecular pathways and targets which serve to explain and - ultimately mimic – the health benefits of exercise as derived from in vitro and animal models.

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Conflict of Interest

None declared.

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