Weight Loss via Exercise with Controlled Dietary Intake May Affect Phospholipid Profile for Cancer Prevention in Murine Skin Tissues

Ping Ouyang, Yu Jiang, Hieu M. Doan, Linglin Xie, David Vasquez, Ruth Welti, Xiaoyu Su, Nanyan Lu, Betty Herndon, Shie-Shien Yang, Richard Jeannotte, and Weiqun Wang

Abstract

Exercise has been linked to a reduced cancer risk in animal models. However, the underlying mechanisms are unclear. This study assessed the effect of exercise with dietary consideration on the phospholipid profile in 12-O-tetradecanoylphorbol-13-acetate (TPA)-induced mouse skin tissues. CD-1 mice were randomly assigned to one of the three groups: ad libitum–fed sedentary control; ad libitum–fed treadmill exercise at 13.4 m/min for 60 min/d, 5 d/wk (Ex+AL); and treadmill-exercised but pair-fed with the same amount as the control (Ex+PF). After 14 weeks, Ex+PF but not Ex+AL mice showed ~25% decrease in both body weight and body fat when compared with the controls. Of the total 338 phospholipids determined by electrospray ionization–tandem mass spectrometry, 57 were significantly changed, and 25 species could distinguish effects of exercise and diet treatments in a stepwise discriminant analysis. A 36% to 75% decrease of phosphatidylinositol (PI) levels in Ex+PF mice occurred along with a significant reduction of PI 3-kinase in TPA-induced skin epidermis, as measured by both Western blotting and immunohistochemistry. In addition, ~2-fold increase of the long-chain polyunsaturated fatty acids, docosahexaenoic and docosapentaenoic acids, in phosphatidylcholines, phosphatidylethanolamines, and lysophosphatidylethanolamines was observed in the Ex+PF group. Microarray analysis indicated that the expression of fatty acid elongase-1 increased. Taken together, these data indicate that exercise with controlled dietary intake, but not exercise alone, significantly reduced body weight and body fat as well as modified the phospholipid profile, which may contribute to cancer prevention by reducing TPA-induced PI 3-kinase and by enhancing ω-3 fatty acid elongation. Cancer Prev Res; 3(4); OF1–12. ©2010 AACR.

Introduction

Overweight or obesity, which may be due to a lifestyle of overconsumption or less expenditure of energy, has become a major public health problem associated with increased risk of many chronic diseases, including cancer (1). Physical activity as a major factor in energy expenditure has been suggested for cancer prevention by both human and animal studies. A large cohort study including 42,672 U.S. postmenopausal women recently reported that light and moderate physical activities were associated with a decreased risk of many chronic diseases, including cancer (2). Another large cohort study including 79,771 Japanese men and women found that a decreased cancer risk was related to increased daily physical activity (3). A population-based study from the 2005 Canadian Community Health Survey showed that physical activity was associated with a high survival rate in the skin cancer patients (4).

The cancer-inhibitory activity of weight loss by calorie restriction and/or exercise has been well documented in many animal models, including the skin carcinogenic model (5). A relationship between reduced dietary energy intake and decreased tumor rates has been established in rodents (6). The molecular targets in response to energy balance for prevention of skin carcinogenesis have also been suggested (7). Michna et al. (8) showed that voluntary wheel running exercise inhibited UVB-induced skin tumorigenesis in mice. The exercise-induced skin cancer inhibition was further associated with enhanced apoptosis in the epidermis via a p53-independent mechanism (9). Most studies from animal models suggest that physical activity inhibits carcinogenesis; however, the exercise-induced inhibition seems less consistent or less potent than calorie restriction (10). Research by the Hursting group suggested that it is negative energy balance rather than exercise alone that inhibited the development of intestinal polyps in APCMin mice (11). By means of dietary regimens in both of high fat and calorie-restricted diet, the...
groups of DiGiovanni and Hursting recently concluded that the dietary energy balance might modulate insulin-like growth factor-I (IGF-I) and IGF-I receptor signaling (12). This reduction is due to both reduced total levels of IGF-I and a reduction in bioavailability due to an increase in levels of IGF-I binding proteins. Similarly, our previous studies found that exercise with pair feeding, but not exercise alone, was effective in controlling body weight and selectively abrogating the tumor promoter–induced phosphatidylinositol 3-kinase (PI3K)–Akt pathway in the skin epidermis, resulting in enhanced apoptosis and reduced proliferation (13).

Many studies have investigated the molecular mechanism of cancer prevention by physical activity. However, the underlying mechanisms of this phenomenon are not clear. Physical activity has been proposed to prevent cancer through deletion of reactive oxygen species and an increase in antioxidant status (14); reduction of sex hormone levels (15); reduction of the metabolic hormones insulin, IGF-I, and/or leptin (16–18); and improvement in immune function (19). Hormone-related signaling alterations, especially of IGF-I, seem to be an important factor in weight control for cancer prevention. PI3K is one of the targets activated by IGF-I. Activated PI3K usually phosphorolates PIs in the cell membrane to produce PI phosphates or phosphoinositides. PI-3,4,5-trisphosphate [PI(3,4,5)P_3], a major product of PI3K, is able to bind and activate Akt, therefore activating many downstream signaling proteins that regulate cell survival and cell cycle progress. Elevated levels of PI(3,4,5)P_3 and upregulation of Akt are found to be oncogenic and promote the transition to malignancy (20–22). Therefore, in addition to being membrane structural building blocks, PIs and their derivatives phosphoinositides have been found to play an important role in cellular signaling and intracellular trafficking as well as in the cancer disease process (23).

Among the phospholipids, some lysophosphatidylcholines (lysoPC) may activate phospholipase C, leading to the production of diacylglycerols and inositol triphosphate, activation of protein kinase C, release of and intracellular Ca^{2+}, and activation of mitogen-activated protein kinase signaling (24). In addition to the signaling processes involving specific lipid head group classes, alterations in the membrane lipid fatty acid composition may also be involved in cancer progression. For example, breast cancer patients were found to have a higher risk of early occurrence of visceral metastasis when they had a lower level of polyunsaturated fatty acids in phosphatidylethanolamines (PE; ref. 25). Decreased levels of stearic acid in PC were found in breast cancer patients with metastasis compared with those who remained metastasis-free (26). Recent studies by the Bougnoux group showed that a lipid profile, rather than a single fatty acid, could be important; they showed that decreased linolenic acid, increased cis-monounsaturated fatty acids, and a low ω-6/ω-3 fatty acid ratio were associated with lower risk of breast cancer (27). Very long chain ω-3 fatty acids, such as docosahexaenoic acid (DHA), are well known to have a preventive effect on cardiovascular disease and cancer and may be involved in alterations of membrane structure and function, eicosanoid metabolism, gene expression, and inhibition of lipid peroxidation (28).

Physical activity has been found to affect membrane phospholipids. For example, it is reported that particular types of exercise increase sphinganine and sphingosine in rat skeletal muscle (29) and reduce total content of ceramide in rat heart muscle (30). Regular exercise was also found to significantly increase oleic acid 18:1 (ω-9) and DHA 22:6 (ω-3) in human muscle (31). However, previous lipid composition studies related to cancer were usually focused on total cholesterol, lipoproteins, and triglyceride (32, 33). The overview of membrane phospholipid profile and the response to exercise-induced weight control has not been studied yet.

With the development of recent lipid analysis, phospholipid compositional profiles can be determined by electrospray ionization–tandem mass spectrometry (ESI/MS-MS). The technique is highly sensitive, accurate, and reproducible (34–36).

In this study, we measured 338 membrane phospholipid species in the skin tissues of mice whose body weights were controlled via moderate treadmill exercise. The effect of exercise-induced weight loss on the phospholipid profile was further evaluated for potential cancer prevention mechanisms. The significant changes in certain species of phospholipids, especially PI3K-related PIs and ω-3 fatty acid–containing PCs, PEs, and lysoPE, might suggest novel cancer-preventive mechanisms by exercise-induced weight control.

Materials and Methods

Animals and animal treatments

Eight-week-old female CD-1 mice (Charles River Laboratories) were housed individually at 24 ± 1°C and 80% relative humidity with 12-h light/12-h dark cycle. They were randomly assigned into one of three groups: ad libitum feeding sedentary control, exercise and ad libitum feeding (Ex+AL), and exercise but pair feeding (Ex+PF). Ad libitum feeding groups were allowed to freely access the basal diet (AIN-93), whereas the pair-fed group was fed daily the same amount as the sedentary control.

A zero-grade adjustable-speed rodent treadmill (Boston Gears) was used to exercise the mice. After a 2-wk training period, mice did treadmill exercise at 13.4 m/min for 60 min/d, 5 d/wk for 14 wk. This exercise was recognized as moderate based on the intensity at 65% to 70% of the maximal oxygen uptake calculated by the predicting regression equation of Fernando et al. (37). To take into account the biological clocks of nocturnal rodents, the light cycle was adjusted for mice to exercise at night. The mice were fed until the last day, but exercise was stopped 24 h after the last bout to measure the effect of exercise training rather than acute exercise. At the end of experiment, the dorsal skin of the mice was shaved and topically treated.
once with 12-O-tetradecanoylphorbol-13-acetate (TPA) at 3.2 nmol in 200 μL of acetone. Mice were sacrificed 2 h after TPA treatment. The dorsal skin samples were snap frozen in liquid nitrogen and kept at –70°C until further analyses.

**Body fat analysis**

In the last week, the body composition of the mice was determined by a dual-energy X-ray absorptiometer using the small animal software (v5.6, Prodigy GE, Lunar-General Electric).

**Phospholipid measurements and profiling**

Each skin sample was ground with liquid nitrogen. After 2 mL of solvent [chloroform/methanol (1:2) + 0.01% butylated hydroxytoluene] were mixed with 1 g of tissue, 1 mL of chloroform and 1 mL of water were added; the mixture was centrifuged at 1,000 rpm for 15 min, and the lower layer was collected. Then, 1 mL of chloroform was added to the tissue, the mixture was centrifuged, and the lower layer was removed and combined with the previously removed lower layer. The combined extract was analyzed for phospholipids using an automated ESI/MS-MS approach. Data acquisition and analysis for acyl group identification were carried out as described previously (35, 36). Twelve phospholipid classes or subclasses, including phosphatidic acid, PI, PC, lysoPC, alk(en)yl/acyl phosphocholine (ePC), PE, lysoPE, alk(en)yl/acyl phosphoethanolamine (ePE), phosphatidylserine (PS), alk(en)yl/acyl phosphoserine, sphingomyelin (SM), and ceramide PE, were determined. Identification of phospholipid molecular species was based on total mass/charge and the presence of a fragment of mass/charge consistent with [M-H]− ions of PE species were identified following collision-induced dissociation of the [M−H]− ions, and acyl ions of PC species were identified following collision-induced dissociation of the [M + OAc]− ions.

**PI3K expression detected by Western blotting**

As described in our previous publication (13), the skin tissue was homogenized in Mg2+ lysis/wash buffer and the lysate was collected after centrifugation at 12,000 × g at 4°C for 15 min. Protein concentration was measured by the Bio-Rad protein assay. After running on 12% SDS-PAGE gel, the proteins were transferred to a nitrocellulose membrane. PI3K at 110 kDa and the internal loading control β-actin at 43 kDa were bound to their monoclonal antibodies (Santa Cruz Biotechnology, Inc.). After incubation with horseradish peroxidase–conjugated secondary antibody (Santa Cruz Biotechnology), the blot was visualized by the FluorChem 8800 Advanced Imaging System (Alpha Innotech). The relative density of the target PI3K band was normalized to the loading control β-actin and then expressed as a percentage of the controls.

**PI3K expression in situ detected by immunohistochemistry**

As described in our previous publication (13), the frozen dorsal skin tissues of mice 2 h after TPA treatment were fixed in –70°C absolute ethanol and rinsed with PBS before adding 10% formaldehyde. The skin tissues were sectioned and the slides were exposed to steam in the target retrieval solution (DakoCytomation). The monoclonal antibody against mouse PI3K (Santa Cruz Biotechnology) was used as a primary antibody, and the secondary antibody was BioGenex QP900 SS multilink horseradish peroxidase kit. Slides were counterstained with Gills hematoxylin followed by dehydration in alcohol and xylene. Staining was developed with diaminobenzidine chromogen, and the density of the stain for each section was scored by a pathologist. Ten to 15 sections for each group were blindly graded using computer standards. The standards of staining intensity were established at 400× by grading up to 40 cells in 5-unit increments from three to five mice per group. Data were statistically analyzed and group scores with P ≤ 0.05 were considered significantly different.

**Microarray and data analyses**

As described in our previous publications (13, 38), the skin samples from four mice in each group were analyzed by a GeneChip Mouse Genome 430 2.0 Array (Affymetrix). The intensities of probe sets were quantified by GeneChip Operating Software 1.0 (Affymetrix). One-way ANOVA and false discovery rate (Benjamini, P < 0.1) were applied to identify the gene expression difference between the treatment and control group. Then, the data were filtered using 1.5-fold change as a cutoff. The intensities of various probe sets for each gene expression of six fatty acid elongases (Elovl) were averaged and reported in this study.

**Statistical analysis**

Phospholipid data outliers were detected using a Q test, and statistical analyses were done using SAS (SAS Institute, Inc.). Results of phospholipid profiling between mice to which acetone vehicle and TPA were applied were found not significantly different, and the data for these groups were thus pooled. Phospholipid levels among the three treatment groups were compared using a one-way ANOVA and an F test for significance followed by pairwise comparisons by least significant difference method. To determine variables that may discriminate the three treatment groups and generate linear combinations, allowing the classification of unknown samples, an automatic backward stepwise discriminant analysis was done using 57 phospholipid species that were significantly different among three treatment groups. Thus, these discriminating variables could be considered putative biomarkers for the effects of diet and exercise on skin polar lipidome.
Body weight and body fat change

Body weights of mice during the total 14 weeks of experimentation are shown in Fig. 1A. Adult CD-1 mice in the sedentary control group gained weight gradually. No significant difference was found between Ex+AL mice and sedentary controls. However, the body weight of Ex+PF mice was significantly lower when compared with either sedentary or Ex+AL mice, beginning in the fifth week of the experiment. As shown in Fig. 1B, when compared with either sedentary control mice or exercised mice with ad libitum feeding, body fat (% of body weight) was significantly lower in exercised and pair-fed mice in comparison with either sedentary control mice or exercised mice with ad libitum feeding. Results are means ± SE (n = 10-15). *, P ≤ 0.05 versus sedentary control or exercised but ad libitum-fed mice.

Effect of exercise with or without controlled diet intake on phospholipid profile

A total of 338 phospholipid species were measured, and 57 of them were found to be significantly different among the treatment groups.

Among the phospholipids, PC was the most abundant head group class in mouse skin, and it represented 45.5% of the total phospholipids analyzed (Table 1). Compared with other classes, PC has the largest amount of shorter-chain species, with species with ≤38 acyl carbons in the two chains making up 9.1% of diacyl PCs, whereas species with ≥38 acyl carbons made up ~26.7% of diacyl PC. Some PCs with short-chain fatty acids, including 30:0-PC, 32:0-PC, and 32:1-PC, were significantly lower in Ex+PF mice than in either group of ad libitum–fed or control mice. However, 40:4-PC, 40:5-PC, and 40:6-PC were significantly higher in Ex+PF mice up to 2-fold when compared with sedentary control as well as Ex+AL mice.

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The second most abundant head group of phospholipids in mouse skin samples is PE, which accounts for 31.5% of the total phospholipids. PE has less short-chain species, with only 0.5% of diacyl PEs having 28, 30, or 32 total acyl carbons and >68% of diacyl PE having ≥38 carbons. The levels of 40:5-PE, 40:6-PE, and 40:7-PE of Ex+PF mice tended to be higher than that of sedentary mice, although they were not significantly different (4.99 ± 0.63 versus 3.13 ± 0.51, 5.94 ± 0.82 versus 4.17 ± 0.90, and 1.48 ± 0.22 versus 1.10 ± 0.15, respectively).

PS diacyl species are 4.6% of the total phospholipids and only 40:6-PS was significantly higher in Ex+PF mice compared with Ex+AL mice (40.71 ± 0.09 versus 0.58 ± 0.11).

Acyl composition analysis was conducted to identify the fatty acid species in 40:5-PC and PE and 40:6-PC and PE. The data in Fig. 2A suggest that 40:5-PC is primarily 18:0-22:5 with small amounts of 18:1-22:4 and 20:1-20:4. The 40:6-PC includes primarily 18:0-22:6 and also 18:1-22:5 species (Fig. 2B). For 40:5-PE, the acyl composition was mainly 18:0-22:5 and also 18:1-22:4 (Fig. 2C), whereas for 40:6-PE the acyl composition was mainly 18:0-22:6 with small amounts of 18:1-22:5 and 18:2-22:4 (Fig. 2D).

The 18:0-22:5 and 18:0-22:6 pair species of PC and PE were similar between Ex+AL and control groups but significantly higher in Ex+PF mice (P < 0.001; Fig. 2E).

For PIs, only seven species were detected and they comprise ~4.6% of the total phospholipids. The major molecular specie is 38:4 PI, which represents 91.4% of the total PI. Most PI species (five of seven) showed a significant (approximately 36-75%) decrease in the Ex+PF mice compared with sedentary control or Ex+AL mice (Table 1).

Table 1 also shows the profile of ePC and ePE, which represent 3.0% and 2.3% of total phospholipids, respectively. Most of the ePCs in the Ex+PF mice significantly decreased compared with the two ad libitum–treated groups. Two ePEs (36:2 and 38:2) significantly increased in Ex+AL mice but...
not in Ex+PF mice. The 38:5-ePE was significantly lower in Ex+PF mice compared with the Ex+AL mice. LysoPC, lysoPE, and SM compromise only a small percentage of total phospholipids, with 1.2%, 0.4%, and 3%, respectively. The data on lysoPC and lysoPE are shown in Table 1. For lysoPC, 16- and 18-carbon species are the predominant species. The lysoPC in Ex+AL mice was not significantly different compared with the sedentary control. In Ex+PF mice, however, 16:0-lysoPC, 18:1-lysoPC, 18:2-lysoPC, and 20:4-lysoPC were decreased significantly.

**Table 1.** Effects of exercise with or without controlled dietary intake on the profiles of phospholipids in mouse skin tissues

<table>
<thead>
<tr>
<th>Phospholipid group</th>
<th>Species*</th>
<th>m/z</th>
<th>Mol% of total polar lipids†</th>
<th>Sedentary</th>
<th>Ex+AL</th>
<th>Ex+PF</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PC</strong></td>
<td>30:0</td>
<td>706.5</td>
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<td>0.28 ± 0.03a</td>
<td>0.19 ± 0.03b</td>
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<td></td>
<td>32:0</td>
<td>734.6</td>
<td>2.21 ± 0.34a</td>
<td>2.10 ± 0.27a</td>
<td>1.02 ± 0.13b</td>
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<td>32:1</td>
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<td>40:5</td>
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<td></td>
<td>40:6</td>
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<td>0.81 ± 0.13a</td>
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<tr>
<td></td>
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<td>3.10 ± 0.41ab</td>
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<td></td>
<td>38:5</td>
<td>883.5</td>
<td>0.14 ± 0.08ab</td>
<td>0.20 ± 0.07b</td>
<td>0.02 ± 0.01b</td>
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<td><strong>ePC</strong></td>
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<td>0.27 ± 0.04a</td>
<td>0.12 ± 0.03b</td>
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<td></td>
<td>32:1</td>
<td>718.6</td>
<td>0.11 ± 0.02a</td>
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<td>0.06 ± 0.01b</td>
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<td>802.7</td>
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<td>800.7</td>
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<td>Ether phospho-etherine</td>
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<td></td>
<td>18:2</td>
<td>520.3</td>
<td>0.17 ± 0.05ab</td>
<td>0.22 ± 0.04a</td>
<td>0.12 ± 0.02b</td>
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<td></td>
<td>20:4</td>
<td>544.3</td>
<td>0.06 ± 0.02a</td>
<td>0.06 ± 0.02a</td>
<td>0.04 ± 0.01b</td>
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<tr>
<td>Lysophospho-etherine</td>
<td>22:5</td>
<td>528.3</td>
<td>0.03 ± 0.01ab</td>
<td>0.03 ± 0.01b</td>
<td>0.07 ± 0.01a</td>
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<tr>
<td></td>
<td>22:6</td>
<td>526.3</td>
<td>0.05 ± 0.01b</td>
<td>0.05 ± 0.01b</td>
<td>0.11 ± 0.02a</td>
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<tr>
<td>Sphingomyelin</td>
<td>16:0</td>
<td>703.6</td>
<td>2.36 ± 0.51a</td>
<td>2.72 ± 0.45a</td>
<td>1.14 ± 0.25b</td>
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<td></td>
<td>24:1</td>
<td>813.7</td>
<td>2.31 ± 0.41ab</td>
<td>2.62 ± 0.35a</td>
<td>1.73 ± 0.26b</td>
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NOTE: CD-1 mice were exercised with or without controlled dietary intake for 14 wk. Quantitative profiling of detectable phospholipids in the mouse skin tissues was done by ESI/MS-MS. Only the significantly changed phospholipids are shown.

*Total acyl carbons:total double bonds.
†Results are means ± SE (n = 8-15) per group. Means with different letters differ significantly, P ≤ 0.05.
when compared with the control or the Ex+AL group. Nearly half (51%) of lysoPE was species with 20 or 22 carbons. The 22:6 and 22:5 lysoPE significantly increased in Ex+PF mice. Two major species that make up 70% of total SM, 16:0 and 24:1 SM, were reduced in Ex+PF (Table 1).

**Discriminant analysis**

An automatic backward stepwise discriminant analysis generated two discriminant functions using 25 phospholipid variables. The variables selected are listed in Fig. 3. The three treatments (sedentary, Ex+AL, and Ex+PF) were significantly distinguishable (Wilks’ Λ = 0.001 at P < 0.00001) using the two discriminant functions. DF1 represented the effect of diet, and DF2 the effect of exercise.

**PI3K protein expression detected by Western blotting and immunohistochemistry**

The Western blot showed that class I PI3K expression was significantly decreased in Ex+PF but not Ex+AL mice (Fig. 4). The image in Fig. 5 depicts the median expression score of class I PI3K as measured by immunohistochemistry. In sedentary control mice, the expression of PI3K in epidermal cells (arrows) was higher in TPA-treated mice (Fig. 5B, scored as 10) than in acetone vehicle-treated mice (Fig. 5A, scored as 5). As the PI3K expression increased in Ex+AL mice (Fig. 5C, scored as 15), the expression level of PI3K in Ex+PF mice decreased to the same level as that of the acetone vehicle sedentary mice (Fig. 5D, scored as 5).

**Gene expression of the Elovl family**

All six members of Elovl gene family are detectable in mouse skin tissues by microarray analysis with various probe sets (up to 11 for each Elovl gene). As Elovl2 is specifically expressed in the liver, it is not unexpected that the expression level was lowest in skin, as shown in Fig. 6. Among the other five Elovl members, only Elovl1 expression increased significantly in Ex+PF mice when compared with either the sedentary control or the Ex+AL group. Elovl5 expression seemed to be enhanced in Ex+PF mice with comparison with the Ex+AL group only but not the sedentary controls. One of four probe sets for Elovl6 indicated overexpression; this was confirmed by a real-time PCR analysis. However, the average of Elovl6 expression as measured by all the four probe sets was not significantly different among the groups.

**Discussion**

This study showed that exercise with controlled dietary intake successfully prevented body weight gain and reduced body fat in the CD-1 mice. It also modified the phospholipid profile significantly in the skin tissues by decreasing some lysoPCs, most PIs, and ePCs; decreasing PI3K protein expression in the skin epidermis; and increasing long-chain polyunsaturated PC, PE, and lysoPE species containing 22:6 and 22:5 that are likely to be ω-3 fatty acids.

The results showed that exercise with *ad libitum* feeding did not effectively decrease body weight and body fat. Similar results are also observed by Mehl et al. (39) in APC<sup>Min</sup> mice and Michna et al. (8) in SKH mice. A moderate increase of food intake from 3.43 ± 0.28 g/d for the control to 3.67 ± 0.18 g/d for Ex+AL mice may compensate, at least in part, the exercise-induced energy expenditure. Furthermore, the dietary energy increase in Ex+AL might not necessarily match the treadmill exercise-induced energy expenditure because the energy expenditure could have been altered due to the changed spontaneous activity in the cage and/or resting energy metabolism. In addition to physical activity, Badman and Flier (40) have suggested that the total energy expenditure should also consist of adaptive thermogenesis and obligatory energy expenditure. It is interesting to compare our results with Huffman et al.’s study (41). The exercise training protocol seems a little bit different between two studies. In our experiment, the mice run treadmill at zero grade for 13 weeks, but Huffman et al.’s mice were exercised at 8% grade for 24 weeks. The most significant difference is due to strain- and diet-related phenomena. We used a lean mouse model that fed a normal AIN-93 diet (5% of calories from fat), but Huffman et al. (41) used C57BL/6 strain, a classic high-fat diet-induced obese model. To induce weight gain, Huffman et al.’s mice were fed a moderately high-fat diet (35% of calories from fat). Therefore, the average of body weights for the control mice in Huffman et al.’s study was ∼30 g in the beginning of experiments and 46 g in week 14. In contrast, the average of body weight for our control mice is ∼26 g in the beginning and 32 g in week 14. It is no doubt that the moderately high-fat diet-induced overweight C57BL/6 strain in Huffman et al.’s study should be much more susceptible to exercise-induced weight loss than our lean CD-1 strain. Actually, our data from the lean CD-1 strain are consistent with what we observed in

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**Fig. 2.** Quantification of 22:5- and 22:6-containing acyl species in PC/PE 40:5 and PC/PE 40:6 phospholipids via production spectra for acyl group identification. CD-1 mice were exercised with controlled dietary intake for 14 wk. The acyl groups of PC/PE 40:5 and PC/PE 40:6 in a lipid extract were identified as acyl anions from the appropriate negative ion precursors by ESI/MS-MS. PC and PE were analyzed as [M + OAc]<sup>−</sup>. These product ion analyses were done on selected molecular ions, indicating three major pairs of acyl composition (18:0-22:5 > 18:1-22:4 and 20:1-20:4) for PC40:5 (A), two major pairs of acyl composition (18:0-22:6 > 18:1-22:5) for PC 40:6 (B), two major pairs of acyl composition (18:0-22:5 > 18:1-22:4) for PE 40:5 (C), and three major pairs of acyl composition (18:0-22:6 > 18:1-22:5 and 18:2-22:4) for PE 40:6 (D). The fractions of the 40:5 and 40:6 species (determined from the lipid profile) that corresponded to the indicated 22:5- and 22:6-containing species were determined from the production spectra analysis (E). The PC/PE 18:0-22:5 and PC/PE 18:0-22:6 were significantly higher in exercised and pair-fed mice but not in exercised and *ad libitum*-fed mice when compared with the sedentary control. Results are means ± SE (n = 4). Means with different letters differ significantly, P ≤ 0.001.
another lean SENCAR strain (13). When compared with Huffman et al.’s study, our contradictory data may provide a diverse phenomenon for a lean strain model with normal diet treatment.

When we adjusted the food consumption of the exercised mice to that of sedentary control mice, significant effects on body weight and body fat were observed. Studies in the Hursting lab found that voluntary wheel running mice with restricted food consumption, a pair-feeding strategy similar to ours, had a significantly lower body weight and less intestinal polyp development (12). Overall, this study indicates that a negative calorie balance via both increasing energy expenditure and limiting calorie intake seems most effective in preventing body weight gain. It should be noted that we may not be able to differentiate body weight control from caloric balance. Although exercise alone with ad libitum feeding was not sufficient to decrease body weight due to, at least in part, the corresponding increase in dietary intake, the lack of an exercise effect in Ex+AL mice on the body weight/fat and various phospholipids might be in part due to the insufficient magnitude of the calorie deficit. Because the average food intake for three treatment groups is comparable, we did not find any significant correlation of calorie intake with specific phospholipids, and the caloric deficit via exercise was not matched with the diet intake, and thus, the results should be interpreted accordingly.

Although it wonders whether the turnover rate of skin would make it a better indicator of subtle changes than other tissues, previous studies by ours and others have shown a cancer-inhibitory activity of weight loss by dietary calorie restriction and/or exercise in animal skin cancer model (5–8). Furthermore, exercise-induced skin cancer inhibition has been linked to apoptosis induction and antiproliferation in the skin epidermis (9, 13). To further evaluate the effect of weight control, lipidomics analysis for all the phospholipids in skin tissues 2 hours after TPA treatment was done. First, we did not find any significant differences of phospholipids between TPA and ace- tone vehicle control. The reason related to a lack of a significant effect of TPA on the phospholipid profile may be due to a short time exposure to TPA treatment in vivo. The selection of 2-hour period for TPA treatment is based on the previous observation that was adequate for a significant activation of Ras and extracellular signal-regulated kinase activities in skin epidermis (13).
Furthermore, the finding that the major molecular species of PI was 38:4 is consistent with the typical pattern of PIs found in mammals and, in particular, in mouse tissues, where this major species has shown to be 1-18:0, 2-20:4 PI (42). The observed decrease of the most PI species in the Ex+PF mice led us to measure expression of PI3K, a key kinase required for various signaling cascades for cancer-related cellular function, including cell growth, proliferation, differentiation, motility, survival, and intracellular trafficking (13). As expected, we found that the decreased PI levels corresponded to lowered levels of PI3K protein in the skin epidermis of Ex+PF mice. Skin cancer development is usually associated with uncontrolled proliferation of epidermal cells (43), so the lower protein expression of class I PI3K in epidermal cells as measured by immunohistochemistry may result in less proliferation as found in our previous report (13). Furthermore, the increased levels of PI3K staining were observed in TPA-treated sedentary control when compared with acetone vehicle control. When compared with TPA-treated sedentary control (Fig. 5B) but not acetone vehicle sedentary control (Fig. 5A), no significant difference was found in Ex+AL group. However, TPA-induced increase of PI3K protein expression was significantly suppressed by exercise with pair feeding, suggesting that the direct product of PI3K, the second messenger PI(3,4,5)P3, might be reduced. It is well recognized that PI(3,4,5)P3 recruits some signaling enzymes with pleckstrin homology domains, such as protein serine-threonine kinases, including Akt and adaptor proteins, to the membrane. The activation of these enzymes affects protein synthesis, cell cycle entry, cell survival function, etc. (20). Decreased PIs and downregulation of PI3K expression observed in this study may imply that body weight control through exercise with controlled dietary intake may prevent against TPA-induced cancer risk. In addition, many studies by us and others have found that weight control was associated with reduced levels of circulating growth hormones or factors such as IGF-I (44). Considering the requirement of PI3K activation by IGF-I–dependent signaling, the downregulation of PI3K protein expression, and the reduced PI3K-related PI substrates in the exercised but pair-fed mice might be caused by a decrease in plasma IGF-I levels. In our studies, IGF-I was restored in the exercised and pair-fed mice either by i.p.

![Fig. 4. Effects of exercise with or without controlled dietary intake on PI3K protein expression in mouse skin tissues. CD-1 mice were exercised with or without controlled diet intake for 14 wk. The level of PI3K protein in skin tissues was determined by Western blotting and quantified by the FluorChem 8800 Advanced Imaging System. Results are means ± SE (n = 10-15). Means with different letters differ significantly, P ≤ 0.01.](image1)

![Fig. 5. Effects of exercise with or without controlled dietary intake on PI3K protein expression in skin epidermis. CD-1 mice were exercised with or without controlled dietary intake for 14 wk. Representative histologic skin sections with immunohistochemical staining for p110-PI3K in epidermis in acetone-treated control (A) and TPA-treated control (B), exercise with ad libitum feeding (C), and exercise with pair feeding (D) are shown. The arrows indicate representative staining of the target protein in epidermis (n = 3-5).](image2)
injection at 10 μg/g body weight twice per week (13) or via osmotic minipumps. We have found that the reduction of PI3K protein expression and the PI species were partially reversed by IGF-I restoration. In addition to PIs, we also found that most of the ePCs and lysoPCs were significantly reduced in the exercised and pair-fed mice, whereas 22:6 lysoPE was increased in Ex+PF group. The lower levels of ePCs and lysoPCs may prevent cancer by reducing cellular damage and proliferation because ePCs are required for the formation of platelet-activating factor and lysoPCs are produced during LDL oxidation within atherosclerotic plaques for atherosclerotic lesion development (45–48).

The effect of weight control via exercise on the PCs and PEs is interesting. When compared with the sedentary control, exercise with ad libitum–fed mice did not change the profile of PCs and PEs. However, there are significant changes observed in the exercised and pair-fed mice. As some short-chain fatty acids of PCs were decreased, the long-chain polyunsaturated fatty acids (i.e., 40:5 and 40:6) increased significantly in Ex+PF mice in comparison with either control or Ex+AL group. A similar effect was found on PEs (data not shown). By means of product ion analysis, the increased polyunsaturated 40:5 and 40:6 fatty acids in PCs and PEs were further discovered to contain a combination of either 18:0-22:5 or 18:0-22:6. The 22:6 fatty acid is undoubtedly DHA. The 22:5 fatty acid (i.e., docosapentaenoic acid) could be either ω-6 adrenic acid or ω-3 clupanodonic acid. As one of the three major ω-3 long-chain polyunsaturated fatty acids, clupanodonic acid could be intermediary between eicosapentaenoic acid and DHA (49). It should be noted that ω-3 22:6 DHA was found to be elevated significantly in the exercised and pair-fed mice not only for PCs and PEs but also for lysoPEs. It is well known that the mammals can make DHA and eicosapentaenoic acid through desaturation and elongation of essential ω-3-linolenic acid (50, 51). Our microarray data further confirmed that elongation of (very) long-chain fatty acid–like elongase gene 1 (Elovl1) was expressed significantly more in Ex+PF compared with the Ex+AL group. Elevation of DHA by exercise has been reported by others in human studies (31, 52). Considering the general health benefits of ω-3 fatty acids (53–56) and a specific inhibitory role in TPA-induced signaling activation (28, 57), the increase of ω-3 fatty acids found in this study may provide a novel approach to understand the mechanisms by which exercise with controlled calorie intake may protect against cancer.

To overview the effects of exercise with or without consideration of diet intake on the phospholipid profiling, we applied a discriminant function analysis to the 57 significantly changed phospholipids. Twenty-five of the total 57 phospholipids were able to distinguish the treatment groups with 92% classification efficiency. These 25 phospholipids are possible candidates for biomarkers to distinguish the effects of diet regimens and exercise in mice. It should be noted that the most 25 phospholipids selected are PCs, ePC, or lysoPCs. The functional effect of these PC-related species changes and how such changes might afford protection from cancer warrant further studies.

Taken together, these data indicate, for the first time, that exercise with controlled diet interventions, but not exercise alone, significantly reduced body weight and body

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5 Unpublished data.
fat as well as modified the phospholipid profile. This modified profile might provide potential cancer prevention benefits, perhaps via reducing TPA-induced PIs and PI-related P13K expression and by enhancing α-3 PC, PE, and/or lysoPE elongation mechanisms.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

We thank Mary Roth for assistance in lipidomics analysis and Dr. Mark Haub for assistance in body fat analysis by dual-energy X-ray absorptiometry.

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Grant Support

National Cancer Institute grants R01 CA106397 and P20 RR15563 (W. Wang); a graduate student summer stipend from Terry Johnson Center for Basic Cancer Research, Kansas State University (P. Ouyang); and National Science Foundation Major Research Instrumentation grant DBI 0521587 and EPSCoR grant EPS-0236913 with matching support from the State of Kansas, NIH grant P20 RR016475 for the Kansas Lipidomics Research Center, Analytical Laboratory (R. Welti), and K-State Functional Genomics Consortium (KSU Targeted Excellence Program). This is journal contribution 06-200-I of the Agricultural Experiment Station, Kansas State University. The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked advertisement in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

Received 02/04/2009; revised 12/18/2009; accepted 01/21/2010; published OnlineFirst 03/16/2010.
Weight Loss via Exercise with Controlled Dietary Intake May Affect Phospholipid Profile for Cancer Prevention in Murine Skin Tissues

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