Excess Linoleic Acid Increases Collagen I/III Ratio and “Stiffens” the Heart Muscle Following High Fat Diets*

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Background: Dietary n-6 polyunsaturated fats (n-6 PUFA) like linoleic acid (LA) may worsen cardiac remodeling after injury.

Results: Excess LA increased cardiac collagen I/III ratio and lysyl oxidase causing early diastolic dysfunction. In vitro experiments in fibroblasts with genetic manipulation confirmed such mechanisms.

Conclusion: LA promotes noncompliant collagen and cardiac stiffening.

Significance: This study demonstrates a novel, cardiac-specific lipotoxic pathway of n-6 PUFAs.

Controversy exists on the benefits versus harms of n-6 polyunsaturated fatty acids (n-6 PUFA). Although n-6 PUFA demonstrates anti-atherosclerotic properties, survival following cardiac remodeling may be compromised. We hypothesized that n-6 PUFA like linoleic acid (LA) or other downstream PUFAs like γ-linolenic acid or arachidonic acid alter the transforming growth factor-β (TGFβ)-collagen axis in the heart. Excess dietary LA increased the collagen I/III ratio in the mouse myocardium, leading to cardiac “stiffening” characterized by impaired transmitral flow indicative of early diastolic dysfunction within 5 weeks. In vitro, LA under TGFβ1 stimulation increased collagen I and lysyl oxidase (LOX), the enzyme that cross-links soluble collagen resulting in deposited collagen. Overexpression of fatty acid desaturase 2 (fads2), which metabolizes LA to downstream PUFAs, reduced collagen deposits, LOX maturation, and activity with LA, whereas overexpressing fads1, unrelated to LA desaturation, did not. Furthermore, fads2 knockdown by RNAi elevated LOX activity and collagen deposits in fibroblasts with LA but not oleic acid, implying a buildup of LA for aggravating such pro-fibrotic effects. As direct incubation with γ-linolenic acid or arachidonic acid also attenuated collagen deposits and LOX activity, we concluded that LA itself, independent of other downstream PUFAs, promotes the pro-fibrotic effects of n-6 PUFA. Overall, these results attempt to reconcile opposing views of n-6 PUFA on the cardiovascular system and present evidence supporting a cardiac muscle-specific effect of n-6 PUFAs. Therefore, aggravation of the collagen I/III ratio and cardiac stiffening by excess n-6 PUFA represent a novel pathway of cardiac lipotoxicity caused by high n-6 PUFA diets.

Our understanding of cardiopotoxicity is undergoing a paradigm shift (1). Instead of saturated fats, recent analyses point toward a possible detrimental effect of n-6 polyunsaturated fatty acids (PUFAs) on cardiovascular health (2, 3). This is surprising, as n-6 PUFA prevents atherosclerosis and attenuates mortality from vascular disorders by lowering cholesterol (4). Thus, potential cardiotoxicity, if present, may exist within the heart muscle itself. Work from our group and others has demonstrated inflammation and oxidative stress with excess n-6 PUFA in the myocardium in animal models of diet-induced obesity and diabetes (5–7). More recently, Galvão et al. (8) demonstrated earlier death with dietary n-6 PUFA in a hamster model of cardiomyopathy, which was independent of changes in heart function. Although adverse cardiac remodeling was suspected, the mechanisms remained elusive.

The role of individual n-6 PUFA on cardiac remodeling remains unresolved. Under in vivo conditions, linoleic acid (LA, C18:2), the parent dietary n-6 PUFA, undergoes desaturation/elongation to longer, more unsaturated n-6 PUFAs like γ-linolenic acid (GLA, C18:3) and arachidonic acid (AA, C20:4) in mammals (9); the role of individual n-6 PUFAs on fibrotic processes remains unclear. The aims of this study were as follows: (i) to identify whether excess n-6 PUFA altered TGFβ and col-

* This work was supported by operating grants from Canadian Diabetes Association (to S. G. and K. M. M.). The authors declare that they have no conflicts of interest with the contents of this article.
1 Both authors contributed equally to this work.
2 Supported by an Natural Sciences and Engineering Research Council of Canada Masters award.
3 Supported by a Canadian Institutes of Health Research doctoral award.
4 Supported by the China Scholarship Council.
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6 The abbreviations used are: PUFA, polyunsaturated fat; GLA, γ-linolenic acid; ARA, arachidonic acid; LOX, lysyl oxidase; LA, linoleic acid; OA, oleic acid; HF, high fat; CO, corn oil; OO, olive oil; MUFA, monounsaturated fatty acid; AA, arachidonic acid; ANOVA, analysis of variance; LV, left ventricle; FADS, fatty acid desaturase; SNP, single nucleotide polymorphism.
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TABLE 1

| Detailed composition of high fat diets | Ingredients | Olive oil diet | Corn oil diet |
|--------------------------------------|-------------|---------------|---------------|
| Casein                               | 240 g/kg    | 240           |               |
| d-Methionine                         | 3.6         | 3.6           |               |
| Corn starch                          | 150         | 150           |               |
| Sucrose                              | 298.8       | 298.8         |               |
| Cellulose                            | 50          | 50            |               |
| Calcium carbonate                    | 3.6         | 3.6           |               |
| Mineral mix                          | 42          | 42            |               |
| Vitamin mix                          | 12          | 12            |               |
| **Oils**                             |             |               |               |
| Soybean oil                          | 10          | 10            |               |
| Corn oil                             | 0           | 190           |               |
| Olive oil                            | 190         | 0             |               |
| **Total**                            | 1000        | 1000          |               |

| a Mineral mix (mg/g) contains the following: di-calcium phosphate 500; magnesium oxide 24; potassium citrate 220; potassium sulfate 52; sodium chloride 74; potassium KSO, 12H₂O 0.55; cupric carbonate 0.3; potassium iodate 0.01; ferric citrate 6; manganous carbonate 3.5; sodium selenite 0.01; zinc carbonate 1.6; sucrose 118.03.
| b Vitamin Mix (mg/g) contains the following: vitamin A 0.8; vitamin D₃ 1; vitamin E 10; menadione sodium bisulfite 0.08; nicotinic acid 3; calcium pantothenate 1.6; pyridoxine HCl 0.7; riboflavin 0.6; thiamin 0.6; sucrose 978.42.

Materials and Methods

Animal Protocols and Diet—All protocols were completed in accordance with the University of British Columbia Animal Care Committee Guidelines. Eight-week-old male C57BL/6 mice were purchased from The Jackson Laboratory (Bar Harbor, ME) and housed in a temperature-controlled room (26 °C) on a 12-h light/dark cycle with free access to food and water. These mice were randomly assigned into two high fat (HF) groups composed of olive oil (OO; monounsaturated fatty acid (MUFA)) or corn oil (CO; n-6 PUFA). Both HF diets were commercially prepared. Sucrose was also added to the HF diets to mimic the current dietary trend of high simple sugar intakes. However, sucrose content was similar between the two HF diets, and both the HF diets were isocaloric and isonitrogenous (Harlan Teklad, XD88232). The HF diets were supplemented with 20% w/w of either olive oil or corn oil (40% energy from fats) as described previously (10). The carbohydrate and protein contents were 39 and 19% energy, respectively. The final diet composition is given in Tables 1 and 2. The chow diet (14% energy from fat; Lab Diet-5P76) was utilized to compare the effects of HF diets per se. All diets were fed ad libitum for 5 weeks. At the end of all experiments, mice were euthanized with isoflurane-CO₂ and hearts excised. A section of the left ventricle (LV) was put in RNALater™ and formalin for mRNA quantification and histology, respectively. The remainder was flash-frozen and stored at −80 °C for biochemical assays.

Echocardiography—At the end of the dietary protocols, Vevo® 2100 system (Fujifilm Visualsonics, Ontario, Canada) was used to examine LV function in mice by noninvasive transthoracic echocardiography. Mice were anesthetized with 2% isoflurane and maintained under anesthesia with 1% isoflurane. Body temperature was kept at 37 ± 0.5 °C by placing mice on a warming platform. Two dimensional M-mode and parasternal short and long axis scans were made to assess changes in left ventricle fractional shortening, ejection fraction, and cardiac output. In addition, transmitral pulsed wave Doppler echocardiography was performed. An apical four-chamber view was acquired by positioning the transducer as parallel to the mitral inflow as possible. From this view, mitral valve early (E) and late (A) flow velocities, E/A ratio, isovolumetric contraction, and relaxation times were determined (11). Deceleration time of early left ventricular filling peak (EDT) was calculated as a surrogate of LV wall stiffness, where LV stiffness was inversely proportional to EDT (12).

TABLE 2

| Macronutrient composition of experimental diets | % w/w | % energy |
|-----------------------------------------------|-------|---------|
| **High fat diet**                             |       |         |
| Protein                                       | 21.2  | 19      |
| Carbohydrate                                  | 44.4  | 39      |
| Fat                                           | 20.0  | 40      |
| **Total**                                     |       | 4.53 kcal/g diet |
| **Normal chow**                               |       |         |
| Protein                                       | 22.6  | 26.4    |
| Carbohydrate                                  | 51.2  | 60.1    |
| Fat                                           | 5.2   | 13.7    |
| **Total**                                     |       | 3.41 kcal/g diet |

Note: Normal chow ingredients are variable as with any semi-purified diet and have not been listed.

Gas Chromatographic Analysis—Composition of high fat diets was analyzed using gas chromatography (GC) by NP Analytical Laboratories (St. Louis, MO) on behalf of Harlan Teklad. For GC analysis of fatty acids in heart muscle or cell culture studies, evaluation of fatty acids was done using a combined extraction and methylation protocol (13). Briefly, 400 µl of media was added to 1.5 ml of hexane and 1.5 ml of BF₃/MeOH. Samples were heated between 90 and 110 °C for 1 h. Next, 3 ml of water was added to the samples and centrifuged for 1 min, and the top hexane layer was removed. Fatty acid methyl esters were analyzed on a Trace 1300 gas chromatograph with flame ionization detector (FID) using a Supelco Fawemux column (30 m × 0.32 mm inner diameter × 0.5 µm). For the analysis of relative ratios of 16:0, 16:1n-9, and 16:1n-10, a shorter, more efficient column from Agilent (DB-WAX, part number 123-7011; 15 m × 0.32 mm inner diameter × 0.15 µm) was utilized with authentic standards of 16:0, 16:1n-9 (Sigma), and 16:1n-10 (Matreya LLC). For all methods, the GC peak area percentage of each fatty acid methyl ester was calculated as a percent of the total of all fatty acid esters.

Western Blotting—Western blotting was performed as described previously (14). In brief, protein was extracted from flash-frozen mouse hearts. 25 µg of protein was separated using SDS-PAGE (10%). Following transfer to a nitrocellulose membrane and blocking, membranes were incubated with primary antibodies for collagen I and III (Santa Cruz Biotechnology, Dallas, TX) at a 1:250 dilution or with LOX...
TABLE 3
Primer sequences used for quantification of mRNA levels by real time PCR
Primer sequences were used to determine gene expression. Both the forward (F) and reverse (R) primers are indicated.

| Gene    | Primer sequences (5′–3′)                     | GenBank accession no. |
|---------|---------------------------------------------|-----------------------|
| tgfβ1   | F: GTCAGCTGAGTGTACCACA                    | NM_011577.1           |
|         | R: AACCTCTATGTTCTGACCC                     |                       |
| loxl1   | F: ACTGTGTGCTCTCCAATCA                  | NM_007742.3           |
|         | R: GACTTGGCTCTGCCCTTGG                   |                       |
| fads2   | F: CGCTACACATCCAAGCAA                    | NR_003278             |
|         | R: GCCTGAAATTACCGCGCT                    |                       |
| 18S rRNA| F: GCTGAAATTACCCGGCT                     | NR_003278             |

(Novus Biological) at a 1:1000 dilution. Membranes were then probed with horseradish peroxidase-conjugated secondary antibodies against rabbit and goat IgGs (Applied Biological Materials Inc., British Columbia, Canada) at a 1:500 dilution. Signals were detected using C-DiGit Blot Scanner (LI-COR) with Image Studio DiGit software (version 3.1). Band density was quantified and expressed relative to total protein loaded, as determined via Ponceau stain, in arbitrary units for collagen.

Masson Trichrome Stain and Immunolocalization of Collagens I and III—Masson trichrome stain was used to evaluate total collagen in the heart as described previously (15). In brief, paraffinized heart sections were rehydrated and mordanted in Bouin’s solution for 1 h at 56 °C. Then they were stained with Weigert’s iron hematoxylin and Biebrich Scarlet-Acid Fuchsian followed by incubation in phosphomolybdic-phosphotungstic acid solution. Finally, sections were stained with aniline blue, differentiated in acetic acid, mounted, and visualized using light microscopy. Double immunolabeling was performed as described previously (14). Briefly, paraffinized sections of mouse hearts were blocked and incubated overnight at a 1:50 dilution with a rabbit polyclonal antibody against collagen I (Santa Cruz Biotechnology) and then with a goat polyclonal antibody against collagen III (Santa Cruz Biotechnology). Secondary anti-rabbit DyLight 488 (BioLegend) and anti-goat DyLight 594 (Santa Cruz Biotechnology) antibodies were used to detect the stain, 250 μl of 0.1 n NaOH and transferred to a 48-well plate. Absorbance was read at 540 nm. Next, cells were stained with Sirius Red and centrifuged at 10,000 × g for 5 min to pellet the collagen, and the supernatant was carefully discarded. To the pellet, 1 ml of HCl was gently added to wash off any excess/unbound dye. To dissolve the stain, 250 μl of 0.1 n NaOH was added. The dye was transferred to a new 48-well plate, from which the absorbance intensity was read at 540 nm using the GloMax Multi+ detection system. Treated cells were compared with control untreated cells, and values were expressed in relative absorbance units. Deposed collagen within cells was visualized using bright field microscopy.

LOX Activity—LOX activity was assessed using the Amplite™ fluorimetric lysyl oxidase assay kit (AAT Bioquest, CA) according to the manufacturer’s instructions and expressed in relative fluorescence units per mg of protein as determined by Bradford assay (Bio-Rad).

Overexpression of fads Isoforms in Fibroblasts—Fatty acid desaturase 2 (fads2) is a gene involved in the desaturation pathways of n-6 and n-3 fatty acids (17). fads2 was overexpressed in NIH/3T3 cells using the mammalian expression vector pTarget (Promega). Briefly, DH10β Escherichia coli containing pCMVSPORT6:FADS2 was obtained from ATCC (ATCC catalog no. 10470049). The 1508-bp fads2 cDNA fragment was isolated from the vector by digestion with MluI (Invitrogen), gel-purified using the QIAquick gel extraction kit (Qiagen), and ligated

VA). NIH/3T3 cells were cultured in DMEM-LG media supplemented with 2 mM sodium pyruvate, 4 mM glut-a-GRO, 10% FBS, and 2% penicillin/streptomycin at 37 °C in a humidified atmosphere of 5% CO₂. Cells were passaged at 70–80% confluency, and media were replaced every 3–4 days. Fibroblasts were seeded in 48-well plates at a density of 15,000 cells/well. Wells were treated with various fatty acids for 24 h and were treated with chemokines such as TGFβ1 and TGFβ3 (10 ng/ml).

Soluble and Deposited Collagen Assay—Both soluble and deposited collagen were measured from cell media extracts using Sirius Red staining, adapted from Tullberg-Reinert and Jundt (16). Briefly, supernatants from cell cultures were incubated with Sirius Red for 30 min, and soluble collagen was collected by centrifugation. Following a wash with 0.01 n HCl, soluble collagen was resuspended in 0.1 n NaOH and transferred to a 48-well plate. Absorbance was read at 540 nm. Next, cells were stained with Sirius Red and centrifuged at 10,000 × g for 5 min to pellet the collagen, and the supernatant was carefully discarded. To the pellet, 1 ml of HCl was gently added to wash off any excess/unbound dye. To dissolve the stain, 250 μl of 0.1 n NaOH was added. The dye was transferred to a new 48-well plate, from which the absorbance intensity was read at 540 nm using the GloMax Multi+ detection system. Treated cells were compared with control untreated cells, and values were expressed in relative absorbance units. Deposited collagen within cells was visualized using bright field microscopy.

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| loxl1 | F: ACTGTGTGCTCTCCAATCA | NM_007742.3           |
|       | R: GACTTGGCTCTGCCCTTGG |                       |
| fads2 | F: CGCTACACATCCAAGCAA | NR_003278             |
|       | R: GCCTGAAATTACCGCGCT |                       |
| 18S rRNA | F: GCTGAAATTACCCGGCT | NR_003278             |
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into pTARGET-fads2 using Attractene transfection reagent (Qiagen), according to the manufacturer’s instructions. Scrambled RNA was used for control transfections (ABM-Good, catalog no. LV015). After 48 h, cells were treated with various fatty acids as indicated for respective experiments for an additional 24 h. At 72 h post-transfection, cells were collected for analysis.

Knockdown of fads2 in Fibroblasts—NIH/3T3 fibroblasts were plated and incubated overnight in complete DMEM. Knockdown of fads2 was achieved by using siRNA (ABM-Good, Richmond, Canada; catalog no. i048001) and DharmaFECT 1 transfection reagent (ThermoScientific, catalog no. T-2001-01) following the manufacturer’s instructions. Scrambled RNA was used for control transfections (ABM-Good, catalog no. LV015). After 48 h, cells were treated with various fatty acids as indicated for respective experiments.

Statistical Analysis—Results are expressed as mean ± S.E. Data were analyzed by either one-way or two-way ANOVA with Tukey post-tests as appropriate, to determine the level of significance. Level of significance was set at p < 0.05. GraphPad Prism (version 5) software was used to conduct statistical analyses.

Results

Short Term Fat Feeding Does Not Alter Body Weight in Mice—Fatty acid analysis of experimental OO and CO diets revealed that CO had higher LA (18:2n-6), whereas the OO diet had higher oleic acid (OA, 18:1n-9) (Fig. 1a). Other major classes of fatty acids like saturated (palmitic (16:0) and stearic (18:0) or n-3 PUFA (α-linolenic acid (18:3n-3)) were similar between both diets (Fig. 1a). Long chain n-3 PUFA like docosahexaenoic acid or eicosapentaenoic acid were low in both HF diets. Despite having sucrose, with a higher energy density (4.53 kcal/g versus 3.41 kcal/g), mice on both HF diets consumed fewer grams of food compared with the chow diet (Fig. 1b). However, there was no difference in food intake among mice eating the two HF diets (Fig. 1b). This adjustment of caloric intake among rodents on a HF diet is known (18) and could have led to similar weight gain compared with chow-fed mice over 5 weeks. Mice on high fat diets consumed less food on a per g basis than chow diets. c, body weight of mice over 5 weeks of diet regimens. Body weight was not significantly different among any group of mice. Bottom panel, Masson trichrome stains of LV depicting perivascular and interstitial regions of LV at 400 and 200 magnifications, respectively. No difference in total collagen staining (blue) in either areas were observed with any diet. Data were analyzed using one-way ANOVA with Tukey post hoc tests (n = 6 mice). *, p < 0.05 versus the chow-fed group.

Statistical Analysis—Results are expressed as mean ± S.E. Data were analyzed by either one-way or two-way ANOVA with Tukey post-tests as appropriate, to determine the level of significance. Level of significance was set at p < 0.05. GraphPad Prism (version 5) software was used to conduct statistical analyses.
TABLE 4
Proportion of total cardiac fatty acids in mice fed various diets over 5 weeks

Data are means ± S.E. (n = 4 mice in each group). Differences between CO- and OO-fed hearts were analyzed by Mann-Whitney test.

| Fatty acid | Chow | OO | CO |
|------------|------|----|----|
| C14:0      | 0.66 ± 0.13 | 0.62 ± 0.05 | 0.87 ± 0.05 |
| C14:1      | 0.73 ± 0.12 | 0.40 ± 0.04 | 0.29 ± 0.03 |
| C16:0      | 19.26 ± 0.73 | 17.19 ± 2.39 | 14.99 ± 0.27 |
| C16:1      | 0.93 ± 0.19 | 0.77 ± 0.11 | 0.32 ± 0.17 |
| C18:0      | 0.27 ± 0.17 | 0.22 ± 0.13 | 0.35 ± 0.04 |
| C18:1t9h12 | 34.30 ± 1.24 | 45.63 ± 3.00 | 31.79 ± 1.92 |
| C18:2n6    | 21.88 ± 1.19 | 10.84 ± 1.71 | 30.12 ± 0.86 |
| C18:3n6    | 0.18 ± 0.08 | 0.08 ± 0.06 | 0.12 ± 0.01 |
| C18:4n6    | 0.34 ± 0.07 | 0.75 ± 0.17 | 0.33 ± 0.03 |
| C20:0n6    | 0.10 ± 0.09 | 0.36 ± 0.41 | 0.90 ± 0.91 |
| C20:4n6    | 0.42 ± 0.29 | 1.20 ± 1.15 | ND |
| C20:5n6    | 0.15 ± 0.11 | 0.28 ± 0.02 | 0.13 ± 0.07 |
| C22:0n6    | 0.97 ± 0.36 | 3.64 ± 0.14 | 1.99 ± 0.16 |
| C22:4n6    | 0.26 ± 0.09 | 0.63 ± 0.53 | 0.12 ± 0.01 |
| C22:6n6    | 10.76 ± 1.55 | 7.07 ± 0.84 | 6.45 ± 0.65 |

a p < 0.05 versus CO-fed mice.

b ND means not detectable.

TABLE 5
Cardiac function as estimated by echocardiography in mice fed various diets

Data are means ± S.E. (n = 6). Differences between CO and OO groups were analyzed by two-tailed Mann-Whitney tests. NC, normal chow; MV, mitral valve; MV A, A wave; MV E, E wave; DT, deceleration time; IVCT, isovolumetric contraction time; IVRT, isovolumetric relaxation time; NFT, no flow time; ms, millisecond. Significant p values are italicized.

| Parameter | Chow | OO | CO | P value |
|-----------|------|----|----|---------|
| LV mass (mg) | 96.97 ± 6.1 | 113.78 ± 12.6 | 107.10 ± 11.7 | 0.59 |
| Diameter (mm), systole | 1.57 ± 0.1 | 1.58 ± 0.1 | 1.60 ± 0.2 | 0.91 |
| Diameter (mm), diastole | 3.59 ± 0.1 | 3.70 ± 0.1 | 3.66 ± 0.2 | 0.88 |
| Ejection fraction (%) | 85.9 ± 0.9 | 1.581 ± 0.121 | 1.596 ± 0.145 | 0.70 |
| Fractional shortening (%) | 54.74 ± 1.1 | 57.51 ± 2.7 | 56.59 ± 2.8 | 0.69 |
| Cardiac output (ml/min) | 24.32 ± 1.3 | 26.78 ± 1.3 | 26.25 ± 2.7 | 1.00 |
| MV A (mm/s) | 536.72 ± 28.93 | 462.50 ± 21.11 | 526.46 ± 44.10 | 0.13 |
| MV E (mm/s) | 714.23 ± 29.4 | 762.72 ± 33.89 | 740.68 ± 63.05 | 0.73 |
| MV A/E | 1.53 ± 0.06 | 1.68 ± 0.14 | 1.40 ± 0.07 | 0.041 |
| DT (ms) | 29.44 ± 1.02 | 31.45 ± 1.7 | 26.56 ± 1.3 | 0.030 |
| IVCT (ms) | 14.17 ± 1.28 | 14.81 ± 1.53 | 16.30 ± 0.53 | 0.39 |
| IVRT (ms) | 16.88 ± 1.61 | 20.55 ± 1.44 | 16.14 ± 0.53 | 0.16 |
| NFT (ms) | 81.32 ± 4.93 | 83.92 ± 3.53 | 84.12 ± 3.37 | 1.00 |

a p < 0.05 versus CO-fed mice.

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FIGURE 2. Corn oil alters collagen subtypes in the mouse heart. a, distribution and localization of collagen I and III in the heart across diet groups via double immunolabeling. Chow mouse hearts depicted low collagen I (fine arrow) with negligible collagen III positivity. OO increased diffuse collagen III across the myocardium (block arrow) and reduced punctate collagen I stains. CO hearts demonstrated the highest collagen I (fine arrows) with negligible collagen III stains. Magnification ×200. b, cardiac mRNA expression of tgfβ1 and tgfβ3 across diet groups. c, cardiac mRNA expression of col1a1 and col3a1. d, representative bands from Western blots for collagen (Col) I and III with total protein depicted by Ponceau stain. e, Western blot analysis of collagen I and III in mouse hearts normalized to total protein under various diets. Data were analyzed using one-way ANOVA with Tukey post hoc tests (n = 6 mice). *, p < 0.05 versus chow fed group. t, p < 0.05 versus corresponding olive oil fed group. A.U., arbitrary units; tgfβ3, gene of tissue growth factor β.

diastolic function and flow characteristics from mice fed various diets. No differences in either LV mass or parameters of systolic function like ejection fraction, fractional shortening, and cardiac output were observed among any group after 5 weeks of HF feeding. However, changes in mitral flow characteristics were observed, which are known markers of abnormal ventricular filling (19). Ratio of E/A waves was significantly reduced in CO-fed compared with OO-fed hearts, which suggests diastolic dysfunction and increased LV "stiffness," especially that E DT was also significantly reduced in CO-fed mice (12). Impaired LV filling without systolic dysfunction indicative of early diastolic dysfunction is associated with altered collagen subtypes (20).

Corn Oil Increases Cardiac Collagen I/III Ratio without Altering Total Fibrosis—Double immunolabeling for collagen I and III was performed in LV sections (Fig. 2a). Out of the two HF diets, OO- and CO-fed hearts demonstrated less and more punctate collagen I immunopositivity, respectively. In contrast, compared with chow and CO, OO-fed hearts demonstrated increased collagen III, at least across some parts of the myocardium (Fig. 2a, red block, arrow). Collagen production is under the direct control of TGFβ in the heart. With respect to tgfβ subtypes, gene expression of tgfβ1 was decreased, whereas tgfβ3 levels was increased in OO-fed hearts (Fig. 2b). tgfβ subtypes in CO-fed hearts remained similar to chow-fed hearts (Fig. 2b). To quantify changes in collagen subtypes, immunoblotting revealed a global decrease of both collagen I and III in both HF diet groups compared with chow-fed mice. However, between the two HF diets, CO-fed mouse hearts demonstrated increased collagen I and decreased collagen III compared with OO-fed mouse hearts (Fig. 2, c and d). In contrast, col-
lagen III was attenuated in the CO-fed mouse hearts but not in OO-fed mouse hearts (Fig. 2, c and d). Taken together, such differences between the two HF diets led to a 3-fold increase in collagen I to III ratio in CO compared with OO-fed hearts (Fig. 2 e).

Linoleic Acid Increases Collagen in Response to TGFβ1 but Not TGFβ3 in Fibroblasts—To decipher underlying mechanisms of differential cardiac collagen subtypes induced by various diets in vivo, mouse NIH/3T3 fibroblasts were incubated with LA or OA, which represent the major fatty acids in CO- and OO-fed diets, respectively (Fig. 1 a). In a high fat (40% energy) fed C57/Bl6 mouse, plasma levels of nonesterified LA and OA are ~0.5 and 1.2 mM/liter, respectively (21). In preliminary studies, 0.5 mM fatty acids induced toxicity in NIH/3T3 fibroblasts (data not shown), and only 0.1 mM fatty acid was used instead in all incubation experiments. 0.1 mM LA caused an increase in total col1a1 and a decrease in col3a1 within 24 h in fibroblasts (Fig. 3 a). As a consequence, akin to in vivo, the ratio of collagen I to III gene expression was increased specifically in LA-treated fibroblasts within 24 h (Fig. 3 b). Sirius Red staining can differentiate between soluble and deposited collagen and assess their interconversion in culture (22). With TGFβ1 stimulation, fibroblasts deposited more collagen with LA than OA-treated cells (Fig. 3 c). To confirm these results, light microscopy revealed increased intracellular collagen deposits within fibroblasts treated with LA but not OA in response to TGFβ1 (Fig. 3 d). In contrast, TGFβ3 stimulated collagen production in OA-treated cells only (Fig. 3 c). Interestingly, with TGFβ3, LA induced even lower collagen deposition compared with untreated fibroblasts (Fig. 3 c).

Overexpressing fads2 but Not fads1 Abolishes Collagen Deposition in LA-treated Fibroblasts—Having identified specific increases in collagen I and deposited collagen with LA both in vivo and in vitro so far, our next goal was to identify the n-6 PUFA moiety that drives such responses. In this regard, LA begins to undergo bioconversion to GLA and ultimately ARA using Δ6-desaturase encoded by the fatty acid desaturase 2 (fads2) gene (17). The ratio of LA to AA is used to study clinical outcomes associated with FADS activities (9). Table 6 depicts fatty acid composition of cells following fads2 overexpression. fads2 overexpression did not alter major fatty acids in fibroblasts either in BSA- or OA-treated fibroblasts compared with nontransfected controls. As expected, overexpression of fads2 significantly lowered LA and GLA, while increasing ARA within fibroblasts treated with LA only. This lowered the ratio of LA/ARA by ~50% in fibroblasts treated with LA.

Some earlier studies had suggested that Δ6-desaturase, the enzyme encoded by fads2, can lead to increased bioconversion of palmitate to sapienic acid (C16:1n10) in the human sebaceous gland (23) and even in COS-7 cells (24). However, in this
TABLE 6
Fibroblast fatty acid composition with fads2 overexpression

Either nontransfected or stably transfected cells were incubated with BSA, 0.1 mM OA, or LA for 24 h. Data represent mean ± S.E. of relative peak area in gas chromatograms from three different experiments. Differences between cells with or without fads2 overexpression were analyzed by t tests.

|                | BSA          | BSA + fads2 | OA            | OA + fads2 | LA            | LA + fads2 |
|----------------|--------------|-------------|---------------|------------|---------------|------------|
| C16:0          | 27.8 ± 1.6   | 34.0 ± 0.7  | 22.4 ± 1.1    | 20.1 ± 6.6 | 29.2 ± 1.3    | 26.5 ± 0.8 |
| C16:1:10       | 3.3 ± 0.8    | 4.4 ± 0.1   | 4.0 ± 0.1     | 3.9 ± 1.8  | 3.1 ± 0.3     | 4.1 ± 0.6  |
| C16:1:n9       | 2.8 ± 1.1    | 3.4 ± 0.1   | 2.9 ± 0.1     | 2.5 ± 0.9  | 3.1 ± 0.1     | 4.4 ± 0.5  |
| C18:1:n9       | 23.9 ± 4.7   | 22.1 ± 1.3  | 49.2 ± 2.1    | 59.3 ± 9.8 | 17.2 ± 1.1    | 16.2 ± 0.2 |
| C18:2:n6       | 2.2 ± 0.7    | 1.7 ± 0.0   | 0.9 ± 0.1     | 1.0 ± 0.2  | 18.2 ± 0.9    | 12.6 ± 0.2 |
| C18:3:n6       | ND           | 0.6 ± 0.1   | 1.0 ± 0.1     | 1.6 ± 0.4  | 1.3 ± 0.2     | 0.8 ± 0.1  |
| C20:4:n6       | 12.9 ± 1.5   | 10.9 ± 0.4  | 5.9 ± 0.2     | 4.4 ± 1.6  | 10.3 ± 1.2    | 16.2 ± 0.4 |
| C18:2/C20:4 ratio | 0.17 ± 0.1 | 0.16 ± 0.0  | 0.16 ± 0.1    | 0.23 ± 0.2 | 1.29 ± 0.4    | 0.70 ± 0.1* |

* p < 0.05 versus nontransfected cells with the same fatty acid treatment. ND, not detectable.

FIGURE 4. Overexpression of fads2 does not alter 16:1:n-10 in fibroblasts. a, characterization of gas chromatographic separations for 16:1:n9 and 16:1:n10. Panel i, representative chromatogram of a spiked fibroblast cell extract with 1 μg of 16:1:n9. Panel ii, representative chromatogram of a spiked fibroblast cell extract with 1 μg of 16:1:n10. Palmitate (C16:0) is also depicted in panels i and ii for positional reference. Representative chromatograms of C16:0, C16:1:n9, and C16:1:n10 in untransfected NIH/3T3 cell extracts treated with BSA (b), OA (d) and LA (f). Impact of fads2 overexpression in NIH/3T3 cells treated with BSA (c), OA (e) and LA (g). Overexpression of fads2 did not affect either C16:0, C16:1:n9, or 16:1:n10 content in any group. Experiments were repeated at least twice in triplicate. Table 6 depicts numerical values for these data. fads2, fatty acid desaturase 2.

study, cells treated with BSA (Fig. 4, b and c), OA (Fig. 4, d and e), or LA (Fig. 4, f and g) had identical levels of both palmitate and total 16:1 (both 16:1:n-9 and 16:1:n-10) with or without fads2 overexpression (also Table 6). Therefore, any effects observed with fads2 overexpression is unlikely to be due to either a loss of palmitate or an increase in sapienic acids in fibroblasts.

Regarding collagen, overexpression of fads2 prevented collagen deposition with and without TGFβ1 stimulation in LA- but not with OA-treated fibroblasts (Fig. 5a). Interestingly, fads2 overexpression lowered collagen deposition with both OA and LA under TGFβ3 stimulation as well (Fig. 5b). To negate nonspecific effects of fads2 overexpression, we overexpressed fads1, which is not directly involved in LA desaturation, in fibroblasts (17). Overexpression of fads1 did not affect collagen deposition in fibroblasts with either fatty acid (Fig. 5c), thus indicating that the increased collagen deposition with LA was due to LA itself and not to other downstream n-6 PUFAs.

LOX Activity Is Augmented with LA Both in Vivo and in Vitro—LOX is a copper-dependent amine oxidase that promotes cross-linking of collagen fibrils and therefore the deposition of insoluble collagen fibers from soluble collagen (25). First, we sought to determine LOX activity in the hearts of various fat-fed mice. In vivo, LOX activity was higher in CO-fed than in OO-fed hearts (Fig. 6a). To identify the mechanism of LOX induction in vivo, gene expression of loxl1 in various fat-fed mice. In vitro, LOX activity was higher in CO-fed than in OO-fed hearts (Fig. 6a). To identify the mechanism of LOX induction in vivo, gene expression of loxl1, the major genes contributing to LOX activity in the myocardium (26), were measured in mouse hearts. Gene expression of either protein was unaltered with dietary intervention in any group of mice (Fig. 6b). In vivo, LOX activity was higher in CO-fed than in OO-fed hearts (Fig. 6a). To identify the mechanism of LOX induction in vivo, gene expression of loxl1, the major genes contributing to LOX activity in the myocardium (26), were measured in mouse hearts. Gene expression of either protein was unaltered with dietary intervention in any group of mice (Fig. 6b). As LOX proteins can also undergo post-translational modifications whereby prolyl oxidase (50 kDa) is cleaved to the active, mature LOX (30 kDa), Western blotting was per-
In vitro, to identify LOX activity, its substrate, i.e. soluble collagen, was assessed in fibroblasts incubated with fatty acids. Soluble collagen increased similarly with both OA and LA under TGFβ1 stimulation (Fig. 6d). As described earlier with deposited collagen (Fig. 3c), TGFβ3 also failed to increase soluble collagen (Fig. 6d) in response to LA, and it was not included in further studies to assess collagen dynamics with LA or other n-6 PUFAs. With regard to LOX, as LA increases collagen deposition under TGFβ1 (Fig. 3c), we hypothesized an increase in LOX activity in response to LA and TGFβ1, which would generate the insoluble (i.e. deposited) collagen. Without stimulation, LOX activity was increased with both OA and LA incubation (Fig. 6e). However, as predicted, TGFβ1 led to a further increase in LOX activity in LA-treated fibroblasts only (Fig. 6e).

To verify the specific impact of LA and its relationship to LOX, fads2-overexpressing fibroblasts were again utilized. OA-treated cells did not demonstrate any change in LOX activity under increasing doses of this fatty acid or with fads2 overexpression (Fig. 7a). In contrast, increasing doses of LA from 0.05 to 0.2 mM led to a dose-dependent increase in LOX activity in LA-treated cells (Fig. 7b). In contrast, cells overexpressing fads2 lowered and completely abolished LOX activities when incubated with 0.2 mM for 24 h (Fig. 7b). Western blotting of prolyl oxidase and mature LOX in nontransfected and fads2-transfected fibroblasts revealed higher mature LOX in nontransfected fibroblasts treated with LA (Fig. 7c), followed by a significant down-regulation of mature LOX with fads2 overexpression. In contrast, OA-treated cells did not demonstrate any change in either prolyl oxidase or mature LOX with fads2 overexpression. In conjunction with Table 6, which demonstrates a specific down-regulation of LA and an increase in ARA with fads2 overexpression, these data suggest that LOX activity and maturation is directly related to cellular LA levels in fibroblasts.

Knockdown of fads2 Exacerbates LA-mediated Fibrogenesis—To confirm the role of fads2 in LA-mediated induction of collagen deposition and LOX activity, we performed RNAi-mediated knockdown of fads2 in fibroblasts. fads2 gene levels were reduced by ~70% with RNAi (Fig. 8a). Incubation with incremental doses of OA did not change collagen deposition when stimulated with TGFβ1 (Fig. 8b). In contrast, fads2 RNAi led to almost doubling of collagen deposition when incubated with either 0.1 or 0.2 mM LA (Fig. 8c). Similar effects were noted with LOX activity with OA in the presence of fads2 RNAi where only the lowest concentration of OA at 0.05 mM led to an increase in LOX activity (Fig. 8d). However, at higher concentrations of fatty acids (i.e. at 0.1 and 0.2 mM), LA demonstrated augmented LOX activity with fads2 RNAi (Fig. 8e). Western blotting of these fibroblasts revealed a significant up-regulation of the mature form when fads2 RNAi-treated fibroblasts were incubated with 0.2 mM LA but not with OA (Fig. 8f). These data suggest that inhibition of fads2 directly increases both maturation of LOX and its enzymatic activity and results in augmented collagen deposition with LA.

Downstream n-6 PUFA Reduces Deposited Collagen in the Presence of TGFβ1—To confirm our results with fads2 overexpression, fibroblasts were directly incubated with 0.1 mM

**FIGURE 5.** Overexpression of fads2 but not fads1 reduces fibroblast collagen deposition with LA. a, impact of fads2 overexpression in fatty acid-treated cells with or without 10 ng/ml TGFβ1 on collagen deposition. fads2 overexpression lowers collagen deposition with and without TGFβ1 with LA but not OA. b, impact of fads2 overexpression in NIH/3T3 cells with or without 10 ng/ml TGFβ3 on collagen deposition. fads2 overexpression lowers collagen deposition with both OA and LA, with or without TGFβ3. c, impact of fads2 overexpression in NIH/3T3 cells with or without 10 ng/ml TGFβ3 on collagen deposition. Unlike fads2, overexpression of fads1 does not affect collagen deposition in cells treated with either OA or LA, with or without TGFβ3. Experiments were repeated at least twice in triplicate. Data are presented relative to control. Data were analyzed using two-way ANOVA with Tukey post hoc tests, p < 0.05, *p < 0.05 versus control cells. †, p < 0.05 versus corresponding OA-treated cells. RAU, relative absorbance units; fads1, fatty acid desaturase 1; fads2, fatty acid desaturase 2.

formed to identify these two fractions (26). CO feeding led to the highest increase in mature LOX to prolyl oxidase ratio in mouse hearts (Fig. 6c).
GLA or ARA, which are the downstream n-6 PUFAs of LA. Compared with LA, GLA and ARA induced lower deposited collagen (Fig. 9a). Furthermore, unlike LA, TGFβ1 stimulation had no effect on collagen deposition in GLA- or ARA-treated cells (Fig. 9a). Overexpression of fads2 inhibited collagen following TGFβ1 stimulation in LA-treated cells but had no effect on GLA- or ARA-treated cells (Fig. 9b). Measurement of LOX activity revealed lower activity in GLA and an almost 3-fold reduction in activity in AA-treated fibroblasts compared with LA (Fig. 9c). Overexpression of fads2 also lowered LOX activity in LA-treated cells alone and had no effect on either GLA- or AA-treated fibroblasts (Fig. 9c). These results confirmed that downstream n-6 PUFAs like GLA or ARA reduce collagen deposition in fibroblasts compared with LA and reinforce the predominant role of LA, the parent n-6 PUFA, in driving collagen deposition under TGFβ1 stimulation.

Discussion

Effects of lipids on heart disease have been extensively studied in animal models. The usual mechanism for lipotoxicity involves generation of ceramide and reactive oxygen species by palmitic acid (or other saturated fatty acids) leading to ectopic lipid deposition and heart failure (27). However, these pathways, based on saturated fats, may only partially explain the current heart disease pandemic in the Western world (1). This is because dietary saturated fats are extensively substituted by n-6 PUFA in the Western diet, and the intake of saturated fats has been relatively constant in the Western population for some time (28). In this regard, the intake of n-6 PUFA has increased from 2.79 to 7.21% energy intake in North America over the last century (28). However, unlike saturated fats, the cardiotoxic potential of n-6 PUFA remains less well defined. We and others have demonstrated that n-6 PUFA overload induces oxidative stress (6, 29, 30), inflammation (7), and mitochondrial damage (31, 32) in endothelial cells and cardiomyocytes. Although it is conceivable that any of the above could alter cardiac remodeling, n-6 PUFA-mediated effects on cardiac fibrosis remain unclear. Here, we present evidence that LA, but not other downstream PUFAs, induce specific TGFβ isoforms and increase the collagen I/III ratio and LOX activity both in vivo and in vitro. Such changes in collagen subtypes induce cardiac stiff-
Excess n-6 PUFA Alters Collagen I/III Ratio in Heart

FIGURE 7. Effect of fads2 overexpression on LOX activity in fibroblasts. Control or stably fads2 transfected fibroblasts were treated with 0.05, 0.1, or 0.2 mM of OA (a) and LA (b) for 24 h. Experiments were repeated at least twice in triplicate. Overexpression of fads2 abolished LOX activity in LA-treated fibroblasts. c, representative Western blots for LOX across 0.2 mM fatty acid treatments. Mature 30-kDa fraction of LOX was profoundly inhibited by fads2 overexpression with LA. Data were analyzed using two-way ANOVA with Tukey post hoc tests, *p < 0.05 versus corresponding untransfected cells. †, p < 0.05 versus corresponding 0.05 mM cells. ††, p < 0.05 versus corresponding 0.1 mM cells. RFU, relative fluorescence units; fads2, fatty acid desaturase 2.

Some reports are in disagreement with other analyses showing that n-6 PUFA lowers coronary heart disease by lowering cholesterol, atherosclerosis progression, and coronary heart disease mortality (4). As shown in this study, LA-mediated cardiac changes may be mediated by alterations of the collagen I/III ratio in the heart muscle itself and not atherosclerotic properties. This is because the C57Bl6 mice as used in this study are resistant to atherosclerosis/fatty streaks with a short term HF diet (34). It is also curious that a negative impact of n-6 PUFA on cardiac function has not been reported earlier. This could be because mitral valve flow measurements are not the standard practice during echocardiography in animal studies (35), and they are difficult to assess using in situ working heart preparations. It has been known for some time that changes in the collagen I/III ratio could lead to early diastolic dysfunction in the absence of changes in commonly assessed systolic or diastolic parameters (20). Clinically, such occurrence is steadily increasing among heart failure patients. It is predicted that heart failure with preserved ejection fraction will be a leading type of heart failure within a decade (36).

Type I and III collagens include ~90% of all collagen within the myocardium (37). Physically, in a single fiber, collagen I is around 37% stiffer than collagen III (38). It is considered that collagen I provides rigidity, whereas collagen III promotes elasticity of the cardiac muscle (39). Therefore, instead of total collagen, collagen subtypes dictated by TGFβ isoforms dictate cardiac stiffness (40). Although TGF-β1 stimulates collagen I production (41), little is known about the role of TGF-β3 in the post-natal heart. Taking cues from its role in the skin, where TGF-β3 attenuates TGF-β1 and collagen I production (42–44) and appears up to 48 h after TGF-β1 expression (45), it is tempting to speculate that TGF-β3 might act to “balance” the production of the noncompliant collagen I expression with the flexible collagen III production or prevent further collagen deposition altogether. Indeed, in avian hearts, specific TGF-β3 stimulation reduces compaction of the extracellular matrix (46).

LOX is an extracellular amine oxidase that plays a crucial role in cross-linking collagen fibrils and deposition of insoluble collagen (25). Excess LOX, fibrillar collagen cross-linking, and fiber deposition are characteristics of heart failure (47). Moreover, TGFβ1 and LOX are known to cross-enhance each other’s activities (48). Soluble collagen is non-aggregated, whereas deposited collagen represents aggregated collagen fibers. LOX catalyzes the oxidation of the amino groups in lysine or hydroxylysine residues, resulting in the formation of aldehydes, which react with each other or with another amino group resulting in cross-linking two polypeptide chains.

In this study, both high fat diets did not induce a change in gene expression of either lox or loxl1 genes in mouse hearts, which are the primary proteins contributing to LOX activities in the adult heart muscle (49). In contrast, corn oil-based diets did induce augmented LOX maturation from the proenzyme form (50 kDa) to the mature form (30 kDa). Such results were then repeated in vitro in fibroblasts, where gene expressions of either LOX or LOXL1 did not change (data not shown), but increased maturation of LOX was noted with conditions that promoted LA (either direct incubation of LA or RNAi mediated inhibition of fads2 with LA in the incubation medium). These results are supported by an earlier study, where a high fat, high sucrose diet induced LOX activity in mouse hearts by promoting LOX maturation, independent of its mRNA levels, via T cell activation (26). Although unknown in the heart, we recently demonstrated T cell recruitment in the gut with an LA-rich diet (10). In an earlier study, the increased dietary n-6 to n-3 PUFA ratio was shown to aggravate cross-linking of collagen in the bone dur-
ing osteoarthritis (50). Despite such evidence, the direct effect of LA or other n-6 PUFA on cardiac LOX remains unclear. As LOX was also activated in fibroblasts with direct incubations with LA, this indicates that LA, rather than any other component of corn oil, might be directly responsible for up-regulation of LOX activity. In this regard, it was demonstrated earlier that oxidized LA promotes cross-linking of collagen in rat tail tendons (51). In contrast, ARA-derived prostaglandin E2 or prostacyclin can down-regulate lysyl oxidase expression or collagen production in lung (52) and cardiac fibroblasts (53), respectively. These opposing effects of LA and ARA on LOX activities are supported in our study as overexpression of FADS2 lowered LA, increased ARA, and simultaneously reduced LOX activity in fibroblasts. Interestingly, in an earlier study, ARA directly inhibited TGFβ activity as well (54).

Unlike PUFA, the role of OA in fibrosis is clearer. Similar to this study, olive oil reduces both TGFβ1 and collagen I in the liver after injury (55). Collagen III mRNA levels were also increased in olive oil-fed mouse hearts in this study. This result is in agreement with an earlier study describing the same in MUFA-fed BALB/c mouse skin following wounding (56). We also show that olive oil did not induce gene levels of LOX proteins in mouse hearts. Moreover, we found that OA up to 0.2 mM did not affect LOX processing in fibroblasts. In an earlier study, higher levels of oleate (0.8 mM) were required for LOX activity with collagen in vitro (57). Therefore, up-regulation of collagen III mRNA by olive oil/OA, reduction of collagen I/III ratio, and inhibition of LOX activities are novel cardioprotective pathways induced by the OO-rich Mediterranean diets (55, 56).

In this study, both fads1 and fads2 overexpression strategies were incorporated to clarify the role of various PUFAs on collagen dynamics in vitro. Although single nucleotide polymorphisms (SNPs) that alter fads activities do alter plasma free fatty acid profiles (58), the role of fads on cardiac events remain unclear. The reported effects with fads SNPs range from nil (59) to detrimental (9) to beneficial (60). Usually, increased desaturase activities that increase AA are blamed for coronary artery diseases and atherosclerosis through pro-inflammatory pathways (9). However, lower activity of fads2 can also induce higher oxidative stress markers in healthy nonobese middle aged men (61), which is in line with earlier studies showing pro-oxidant effects in the myocardium with an LA-rich diet (5–7). In this study, RNAi-mediated inhibition of fads2 also augmented LOX maturation and activity under incubation with LA. In a recent study, cardiac collagen production was diminished by increased activation of fads1/2 and elevated

![FIGURE 8. Effect of fads2 RNAi on collagen deposition and LOX activity in fibroblasts. a. fads2 gene levels after incubation with scrambled or fads2 RNAi in fibroblasts. b. Deposited collagen in fibroblasts treated with incremental doses of OA (b) or LA (c) in the presence of either scrambled or fads2 RNAi after 24 h. No effect of fads2 RNAi was noted with OA, but it increased collagen deposition with both 0.1 and 0.2 mM LA. LOX activity in fibroblasts treated with incremental doses of OA (d) or LA (e) in the presence of either scrambled or fads2 RNAi after 24 h. OA demonstrated increased LOX activity at 0.05 mM, whereas LA demonstrated increased LOX activity at 0.1 and 0.2 mM under fads2 RNAi. f. Western blots for LOX from fibroblasts with fads2 RNAi with 0.2 mM fatty acids. fads2 RNAi induced significant increase in mature/prolysyl oxidase levels with LA but not OA. Experiments were repeated at least twice in triplicate. Data were analyzed using two-way ANOVA with Tukey post hoc tests, *p < 0.05, **p < 0.05 versus corresponding scrambled controls. RFU, relative fluorescence units; fads2, fatty acid desaturase 2.]

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SEPTEMBER 18, 2015 • VOLUME 290 • NUMBER 38

JOURNAL OF BIOLOGICAL CHEMISTRY

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Alternatively, pharmacologically increasing the TGFβ1
preventative strategy to limit adverse cardiac remodeling.
(as in traditional Mediterranean diets) could be an effective
holds true in humans, decreasing dietary LA and increasing OA
5–10% energy intake. Thus, these findings represent a novel
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ARA accumulation in a rat model of hypertension. Not surpris-
ingly, such effects were also down-regulated by knockdown of
fads1 or fads2 in cultured cardiac fibroblasts in the same study
(53). Based on such results, investigations of fads SNPs in clinical
cellular populations may reveal significant associations with car-
diac fibrosis and remodeling.
In summary, we demonstrate that n-6 PUFA-rich corn oil
diets alter the collagen I to III ratio and LOX activity, which can
have a functional consequence in the heart within 5 weeks (Fig.
10). Although the nutritional requirement of n-6 PUFA is only
1% energy (62), current clinical recommendations propose
5–10% energy intake. Thus, these findings represent a novel
pathway of cardiolipotoxicity associated with LA, a fatty acid
increasingly present in Western diets (28). If this mechanism
holds true in humans, decreasing dietary LA and increasing OA
(as in traditional Mediterranean diets) could be an effective
preventative strategy to limit adverse cardiac remodeling.
Alternatively, pharmacologically increasing the TGFβ3/colla-
gen III axis represents an attractive therapeutic target to confer
a more “elastic” phenotype to the failing myocardium. Beyond the
heart, increased collagen I/III ratio can induce adverse clin-
ical outcomes in fibrotic diseases like cirrhosis of the liver (63),
pancreatitis (64), and respiratory distress in the lungs (65).
Whether soaring dietary LA influences the progression of such
diseases as well remains an attractive but unexplored area of
research.

Author Contributions—This project was primarily carried out by
J. B., who also wrote the first draft. Animal work and revisions were
conducted by A. B., who also contributed to the writing of the
revised version. J. Y. performed the RNAi experiments. H. S. per-
formed echocardiography and analyzed data. B. J. M. performed
gene expression and Western blotting. M. F. constructed the
fads2-overexpressing cell line. K. M. M. critically reviewed the data and
the manuscript. The core idea, funding, coordination, and writing of
the final draft of the manuscript was performed by S. G.

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