Dried mulberry fruit ameliorates cardiovascular and liver histopathological changes in high-fat diet-induced hyperlipidemic mice

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Abstract

Background and aim: Metabolic disease encompasses most contemporary non-communicable diseases, especially cardiovascular and fatty liver disease. Mulberry fruits of Morus alba L. are a favoured food and a traditional medicine. While they are anti-atherosclerotic and reduce hyperlipidemic risk factors, studies need wider scope that include ameliorating cardiovascular and liver pathologies if they are to become clinically effective treatments. Therefore, the present study sought to show that freshly dried mulberry fruits (dMF) might counteract the metabolic/cardiovascular pathologies in mice made hyperlipidemic by high-fat diet (HF).

Experimental procedure: C57BL/6J mice were fed for 3 months with either: i) control diet, ii) HF, iii) HF + 100 mg/kg dMF, or iv) HF + 300 mg/kg dMF. Body weight gain, food intake, visceral fat accumulation, fasting blood glucose, plasma lipids, and aortic, heart, and liver histopathologies were evaluated. Adipocyte lipid accumulation, autophagy, and bile acid binding were also investigated.

Results and conclusion: HF increased food intake, body weight gain, visceral fat accumulation, fasting blood glucose, plasma lipids, and aortic, heart, and liver histopathologies were evaluated. Adipocyte lipid accumulation, autophagy, and bile acid binding were also investigated. HF+100 mg/kg dMF, or iv) HF+300 mg/kg dMF. Body weight gain, food intake, visceral fat accumulation, fasting blood glucose, plasma lipids, and aortic, heart, and liver histopathologies were evaluated. Adipocyte lipid accumulation, autophagy, and bile acid binding were also investigated.

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1. Introduction

Life-style dominated by excess energy intake leads to wide range of disease processes collectively termed metabolic disease. These conditions are characterised by but not restricted to: hypertriglyceridemia, diabetes, hypercholesterolemia (especially very low- and low-density lipoprotein cholesterol (LDL)),12 hepatic steatosis12 and cardiovascular disease (atherosclerosis resulting in hypertension, myocardial infarction, and stroke).5 Vascular dysfunction is dominated by inflammatory beginning with endothelial damage and discontinuity; loss of vasodilatory signalling; oxidised LDL entering the arterial intima, then engulfed by invading macrophages that become lipid-laden foam cells manifest as fatty streaks. Vascular smooth muscle cells proliferate and migrate: the arterial intima becomes fibrotic; platelets adhere to the damaged vascular wall thus triggering coagulation and compromised blood flow.7,8

As fat accumulates, the adipocyte lipid capacity becomes limiting whereupon resident macrophages switch to inflammatory phenotypes, especially in visceral and perivascular adipose tissues.9 From the latter, inflammatory cytokines exacerbate vascular inflammation. Insulin and leptin resistance develops so excess circulating lipids are then removed by non-adipocyte lineages (ectopic fat: muscle, liver, etc.). However, the physiological mechanisms such as exercise or thermogenesis can no longer mobilise fat from refractory adipocytes and ectopic fat and fail to impact body weight. Autophagy recycles dysfunctional cell constituents but becomes less effective in metabolic syndrome10 and fatty liver disease.11 The related lipophagy mobilises fats from cell lipid droplets from a variety of cells including hepatocytes. Lipophagy also may stimulate transdifferentiation of white adipocytes into brown fat cells, absent from obese patients, that metabolise lipids by uncoupled oxidative thermogenesis.12

Most current anti-atherogenic treatment strategies focus on antihypercholesterolemic treatments: (a) diets low in cholesterol or its precursor; (b) intestinal cholesterol absorption inhibitors; (c) PCSK9 inhibitors that remove LDL cholesterol, and (d) statins, the mainstay treatment, that target hepatic cholesterol synthesis by inhibiting 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase.5,13 Statins are the most prescribed drugs but risk causing myopathy, diabetes, weakness by Q10 depletion, etc.13 However, these patients have a constellation of pathologies and targeting only one risk factor fails to treat the underlying ill health. Indeed, many of these patients drift into treatment polypharmacy that increases iatrogenic conditions. Traditional medicine with herbal medicines takes a holistic approach to improving overall efficient function body, well-being, and quality of life, a strategy well-suited to metabolic disease. Food derived medicines also provide levels of safety rarely achieved by allopathic drugs.

Mulberries from Morus alba L. are a source of nourishment and used as a traditional medicine to relieve fever, hepatic protectant, improve eyesight, and as an antihypertensive.14 Several pharmacological studies showed that the extracts of this fruit ameliorate diabetes, obesity, hyperlipidemia, and atherosclerosis.15–18 Water- or anthocyanin-rich mulberry extracts inhibited LDL oxidation and foam cell formation in vitro.19 Aqueous or etOH extracts of mulberries improved blood lipids and reduced aortic atherosclerotic lesions and liver lipid accumulation in cholesterol fed rabbits and high-fat fed rats.15,16 Thus, mulberries may well be an efficacious treatment in humans, but the predominant use of extracts may not replicate its traditional use by losing pharmacologically active ingredients. Furthermore, extractions are energy-wasteful. However, getting soft fruit, including mulberries, to market before spoilage is challenging, particularly as medicines that need reliable supplies. Freeze-drying mulberries circumvents these problems. Indeed, such preparations also improved lipid profiles.17,18

Nevertheless, data on various mulberry extracts so far described is fragmented and does not account for the diversity of pathologies manifest as metabolic disease. The current study tests how the unrefined product affects some important risk factors and goes some way define some mechanisms involved protecting against vascular disease and fatty liver that could be clinically applied.

2. Material and methods

2.1. Plant sample preparation

Dried mulberry fruit powder (dMF) was obtained from the Thailand Institute of Scientific and Technological Research, Bangkok, Thailand. Ripe mulberries (M. alba L.) from Nakhon Pathom province in Thailand were freshly harvested, freeze dried and roller ground to obtain dMF that was stored at −20 °C in darkness until use.

2.2. Chemical composition of dMF by LC-MS

dMF was dissolved in water (20 mg/mL), sonicated and 0.22 μm filtered. Separation used a Luna C18(2), 150 × 4.6 mm, 5 μm column (Phenomenex, USA). The analysis used an Agilent HPLC 1260 consisting of a vacuum degasser, a binary pump, an autosampler and a column thermostat equipped with QTOF 6540 UHD accurate mass spectrometer (Agilent Technologies, USA). A 20 μL sample (20 mg/mL) was injected into the LC system with a solvent flow rate of 500 μL/min. The mobile phase consisted of a gradient elution between water (solvent A) and acetonitrile (solvent B), both containing 0.1% v/v formic acid. The linear gradient elution was 5–95% for solvent B starting at 0–30 min and post-run for 5 min. The column temperature was controlled at 35 °C. The mass analysis was performed using a QTOF 6540 UHD accurate mass spectrometer. The conditions for the dual negative electrospray ionization source were drying gas (N2) at a flow rate of 10 l/min, a drying gas temperature of 350 °C, nebulizer 30 psig, fragmentor 100 V, capillary voltage 3500 V, skimmer, 65 V, octopole (Q1 1 RF Vpp), 750 V and scan spectra from m/z 100–1000 amu. The auto MS/MS for the fragmentation was set with collision energies of 10, 20 and 40 V. The positive mode was also set up with the same MS conditions as the negative mode. The data were processed by the Agilent MassHunter Qualitative Analysis software version 8.06.0, which provided a list of possible molecular formulas. The MS data, MS/MS fragmentation profiles and molecular formula proposed by the MassHunter were compared with the literature data and some
databases, such as Human Metabolome, ChemSpider, METLIN, and Lipidmap to annotate the phytochemicals analyzed from the extracts. The molecular formula proposed by MassHunter in MS experiments was compared with the literature data, and a maximum error of 10 ppm was accepted.

2.3. Anthocyanin content determination and DPPH assay

dMF total anthocyanins were measured by the pH differential method. In addition, the reducing ability of the dMF was evaluated by DPPH assay.

2.4. Animal and diet

Twenty four female C57BL/6J mice (aged 8 weeks; weight 18–20 g) were obtained from the Nomura Siam International Co. Ltd., Bangkok, Thailand. Experiments were approved by Naresuan University Animal Care and Use Committee (NUACUC, Naresuan University, Phitsanulok, Thailand; ethic protocol number NU-AE 610727) for the care and use of animals for scientific purposes. The mice were acclimatized for 1 week in their polycarbonate cages at 22 ± 1 °C, 12-12 h light-dark cycle, and allowed access to food and water ad libitum at the Center for Animal Research, Naresuan University, Phitsanulok, Thailand. The animals were randomly divided into 4 groups each of which were fed a different diet for 3 months;

i) Control: Animals were fed with normal diet (Teklad Global: “18% Protein Rodent Diet”, 3.1 kcal/g containing 6.5% fat, 44.2% carbohydrate, 18.6% protein from Harlan Teklad Laboratory, Madison, Wisconsin, USA), and gavaged with water.

ii) High fat diet (HF): Animals were fed with HF (Harlan Teklad Atherogenic Rodent Diet, 4.5 kcal/g composed of 21.2% fat, 0.2% cholesterol, 46.9% carbohydrate, 17.3% protein) and gavaged with water. Since the pellets were more friable than the standard diet, they were treated with polyvinylpyrrolidone to match their hardness and strength of the standard pellets.

iii) HF + 100 mg/kg dMF (HF + dMF100): Animals were fed with HF and dMF was administered by gastric tube once daily as an aqueous suspension.

iv) HF + 300 mg/kg dMF (HF + dMF300) Animals were fed with HF and dMF was administered by gastric tube once daily as an aqueous suspension.

All animals were weighed and food intake was measured daily throughout the 3 month test period. At the end of the experiment, mice were fasted (12–14 h), deeply anesthetized by 50–70 mg/kg sodium thiopental intraperitoneally. Then blood, aorta, heart, liver and visceral fat were removed as below.

2.5. Visceral fat accumulation

Visceral fat was manually separated from the abdominal cavity, then weighed and expressed as g/100 g body weight.
2.6. Blood lipid profile

Whole blood was collected by cardiac puncture and transferred to heparin-coated tubes, centrifugated at 3000 g, 4 °C for 10 min and the plasma retained. Plasma levels of total cholesterol (TC), triglyceride (TG), and high density lipoprotein (HDL) were measured using an enzymatic colorimetric test according to manufacturer protocols (Human diagnostic company, Wiesbaden, Germany). Plasma LDL was calculated by the following equation: LDL = TC - (HDL + TG/5).

2.7. Blood glucose analysis

Before euthanasia, the tip of tail was cut and ~10 μL of blood directly dropped on to test strips and blood glucose measured using an ACCU-Chek Performa blood glucose monitor (Roche Diabetes Care, Mannheim, Germany).

2.8. Determination of histological changes in aorta and heart tissue

Aorta and heart were quickly separated, rinsed with normal saline and then cleaned of fat and fixed with buffered 10% formalin, dehydrated, and embedded in paraffin. De-waxed sections of 3–5 μm were stained by Masson’s trichrome for collagen. Sections were photographed via light microscopy and analyzed by Image J (version 1.51J, National Institutes of Health, USA). Areas of collagen fibre accumulation in the tunica media was calculated as a percentage of the total area of tunica media bounded by the internal elastic lamina and the external elastic lamina.

2.9. Hepatic lipid accumulation

Livers were washed with normal saline and then preserved in 30% sucrose in paraformaldehyde at 4 °C. They were cut at 10 μm as frozen sections, stained with oil red O, mounted in Permount and observed under a microscope.
photographed at 40× magnification by light microscopy. Images were analyzed by Image J software (version 1.51j8, National Institutes of Health, USA) to quantify liver lipid area occupied by oil red O positive staining and expressed as the proportion of region of interest (%).

2.10. Autophagy immunohistochemistry of liver

Liver samples were collected and fixed in 10% neutral buffered formalin before paraffin embedding (FFPE). For immunohistochemistry (IHC) evaluations, 4 μm sections of FFPE liver tissue were

![Fig. 2.](image-url)
mounted onto commercially coated glass slides, dried at room temperature, and incubated at 65 °C for 1 h prior to use. Sections were deparaffinized/hydrated. For antigen unmasking, sections were submersed in 1x citrate unmasking solution, heated to boiling point and left for 10 min (at 95–98 °C) and then cooled for 30 min. Endogenous peroxidases were blocked by incubating with 3% hydrogen peroxide for 10 min. The sections were washed in deionized water and wash buffer, then each section blocked with 100–400 μL of TBST/5% normal goat serum for 1 h at room temperature. After removing the blocking solution, rabbit monoclonal anti-LC3A/B (1:500, Cell Signaling, USA) or rabbit monoclonal anti-SQSTM1/p62 (1:250, Cell Signaling, USA) were added to each section. These were incubated overnight at 4 °C. The sections were washed by wash buffer before covering with IHC detection reagent (UltraView Universal HRP multiplier, Ventana, USA) and incubated in a humidified chamber for 30 min at room temperature. SignalStain® DAB chromagen was applied on each section. Tissue sections were counterstained with hematoxylin and then dehydrated using 95% ethanol, 100% ethanol, and xylene and mounted in mounting medium. The slides stained for LC3A/B and SQSTM1/p62 were photographed with Olympus microscope using cell Sens Imaging Software. Numbers of hepatocytes positive for LC3 A/B or SQSTM1/p62 were counted using image J.

2.11. dMF on adipocyte function

dMF was tested on pre-adipocyte differentiation (adipogenesis) and fat loss from mature adipocytes (lipolysis). 3T3-L1 pre-adipocytes were cultured in high glucose Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% fetal bovine serum and 1%/mL penicillin/streptomycin as completed medium under a humidified 5% CO2 atmosphere at 37 °C in CO2 incubator. Inhibition of lipid accumulation by dMF was assessed as reported.24 After reaching confluence in flasks, cells (2 × 103 cells/well) were seeded into 96 well plates and cultured in complete media for 2 days.

Action of dMF on adipocyte differentiation: The medium was changed to adipocyte differentiation medium, complete medium containing with added: dexamethasone (1 μM DEX); isobutylmethylyanthine (0.5 mM, IBMX); and insulin (10 μg/mL), to initiate adipocyte differentiation (defined as day 1), also with dMF (20, 50, 100, 200, 500, or 1000 μg/mL). This medium was removed at days 3, 5, and 7 and replaced with complete medium with 10 μg/mL insulin and dMF only. At day 9, cells were tested for either (i) lipogenesis by cell content using oil red O staining, or (ii) cell viability using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazoium bromide (MTT). Cells without dMF served as negative controls.

Action of dMF on lipolysis: For lipolysis, a similar protocol was used for 9 days but without dMF using adipocyte differentiation medium and supplemented with 10 μg/mL insulin only. At day 9, dMF (20, 50, 100, 200, 500, or 1000 μg/mL) was then added to the medium and further incubated for 24 h. Finally, treated cells were assessed for either lipid content, or cell viability.24

For assay of adipocyte lipid content25, cells, were washed twice with of PBS (100 μL), fixed with 10% formalin/PBS (100 μL) for 8 min and 1 h, and then washed twice with 100 μL of 60% isopropanol. The cells were allowed to dry in room air. Then 50 μL of 0.5% oil Red O in distilled water was added and incubated for 45 min, then washed twice with 100 μL distilled water. The dye was extracted with 100 μL of isopropanol, shaken for 10 min, and optical density of the solution measured at 500 nm using microplate reader. Lipid contents were expressed as absorbance compared to control (untreated cells = 100%).

For cytotoxicity testing by the MTT assay,26 cells were incubated with MTT reagent (0.5 mg/mL) for 4 h at 37 °C in CO2 incubator. Then, the solution was discarded and DMSO (100 μL) was added and shaken for 15 min to dissolve formazan crystals. The absorbance was recorded at 595 nm using microplate reader. Cell survival expressed as absorbance compared to control (no treatment = 100%).

2.12. Bile acid binding assay

The bile acid binding to dMF was measured as previously reported.27 dMF was dissolved in distilled water, well mixed and diluted to 0.2, 2, 10, and 20 mg/mL of dMF. dMF solution (200 μL), (final concentrations 0.1, 1, 5, and 10 mg/mL) was incubated with 20 μL of 20 mM bile acids (taurocholic acid or taurodeoxycholic acid) with phosphate buffer saline (PBS) (180 μL) for 2 h at 37 °C. The mixtures were centrifuged at 10,000 rpm for 20 min and filtered through a 0.22 μm filter. The filtrate (10 μL) was reacted with 1 unit of 3α-hydroxysteroid dehydrogenase (10 μL) in 180 μL of reaction mixture (tris buffer pH 9.5; 1 M hydrazine hydrate; 7.7 mM nicotinamide adenine dinucleotide (NAD), ratio of 1:5:1:0.3) at 30 °C. Cholesterolamine and PBS were used as positive and negative controls, respectively. After being incubation for 90 min, the optical density of reactions was measured at 340 nm using microplate reader. The percentage of bile acid binding was compared with control (no treatment) which considered as 0%.

2.13. Statistical analysis

All data were expressed as mean ± standard error of mean (SEM) of n animals. The statistical significance between groups was evaluated using student t-test and/or ANOVA followed by the Tukey–Kramer post hoc test. Values of P < 0.05 were considered statistically significant.

| Table 2 | Plasma lipid profiles and blood glucose level at 3 months. |
|---|---|---|---|
| Parameters | Control | HF | HF + dMF100 mg/kg | HF + dMF 300 mg/kg |
| **Plasma lipid (mg/dL)** | | | | |
| TC | 60.9 ± 1.2 | 127.6 ± 4.6** | 118.4 ± 4.5** | 116.7 ± 9.4** |
| TGs | 70.5 ± 4.1 | 70.2 ± 2.7 | 49.3 ± 3.6**,### | 46.8 ± 19**,## |
| HDL | 37.2 ± 2.1 | 34.9 ± 4.3 | 44.2 ± 1.0**,## | 46.4 ± 0.6**,## |
| LDL | 19.4 ± 2.8 | 71.4 ± 5.0**,## | 64.5 ± 8.1* | 64.5 ± 12.0* |
| TC/HDL | 1:6 ± 0.1 | 3.6 ± 0.2** | 2.62 ± 0.1**,## | 2.59 ± 0.2**,## |
| **Blood glucose (mg/dL)** | | | | |
| Baseline | 95.5 ± 1.6 | 97.3 ± 5.2 | 94.8 ± 4.2 | 97.7 ± 2.9 |
| 3 months | 94.2 ± 2.5 | 113.7 ± 3.9** | 94.3 ± 0.7*** | 99.8 ± 3.0# |

Control, normal diet; HF, high-fat diet; HF + dMF 100, high fat diet + 100 mg/kg dried mulberry fruit powder; HF + dMF 300, high-fat diet + 300 mg/kg dried mulberry fruit powder; TC, total cholesterol; TGs, triglycerides; HDL, high density lipoprotein cholesterol; LDL, low density lipoprotein cholesterol. Values are means ± SEM (n = 6). **P < 0.01, ***P < 0.001 vs. control. #P < 0.05, ##P < 0.01, ###P < 0.001 vs. high fat diet.
3. Results

3.1. Composition of dMF

Contents by chemical analysis: Total anthocyanin content by the pH differential method was 1.46 g/100 g dMF.

Contents by LC-MS: Water soluble constituents of dMF yielded 37 peaks corresponding to tentatively identifiable compounds by their monoisotopic mass and fragmentation patterns (Fig. 1; Table 1). Hexoses and organic acids formed a large proportion of the content, while components having possible pharmacological actions formed 4 groups: (i) anthocyanins as cyanidins or pelargonidin glycosides (compounds 12, 15, 17, and 18); (ii) flavonols as rutin (27), and taxifolin (30), (iii) phenolic acids, as conjugates of caffeic acid and quinic acid (5, 6, 16, 19, 22); (iv) Amino acid compounds and other nitrogen contain compounds (1–3, 7, 10, 11, 20, 23–26, 28, 29, 36, 37).

3.2. Body weight gain, food intake, visceral fat and liver weight

Gains in body weight were consistently greater for mice on the HF diet than controls by ~100% at one month but less so for months 2 and 3 (Fig. 2A). Weight gains for the dMF-dosed groups (both 100 and 300 mg/kg) prevented almost all the weight gain ascribed to the HF at month 1, but less pronounced for the remaining 2 months (Fig. 2A). These weight changes mirrored changes in food intake which was doubled with the HF and likewise reduced by 100 and 300 mg/kg dMF (Fig. 2B).

The HF substantially increased the amount of visceral fat and the liver weight (Fig. 2C and D). Some of the increased visceral fat and liver weight was prevented by both dMF dose regimens, a ~60% reduction for visceral fat and ~30% for liver.

Table 3
Aortic wall thickness and collagen content, and hepatic lipid accumulation.

| Parameters                  | Control       | HF            | HF + dMF100    | HF + dMF300    |
|-----------------------------|---------------|---------------|----------------|----------------|
| Aortic wall thickness (μm)  | 275 ± 8       | 386 ± 11*     | 314 ± 6*,###  | 338 ± 6*,###   |
| Area of medial collagen (%) | 31.5 ± 1.3    | 48.9 ± 1.2*   | 39.5 ± 1.6*,###| 40.9 ± 1.4*,###|
| Proportion of hepatic lipid (%) | 33.6 ± 2.1 | 51.9 ± 1.2**  | 41.0 ± 2.6*, # | 42.0 ± 1.6*, # |

Control, normal diet; HF, high-fat diet; HF + dMF100, high-fat diet + 100 mg/kg dried mulberry fruit powder; HF + dMF300, high-fat diet + 300 mg/kg dried mulberry fruit powder. Aortic wall thickness was measured as the perpendicular distance between the inner wall of the tunica intima to the outer extremity of the tunica media. Hepatic lipid was the area occupied by oil red O positive staining and expressed as the proportion of region of interest area. Values are means ± SEM (n = 5 mice). *P < 0.01, **P < 0.001 vs. Control. #P < 0.05, ##P < 0.01 vs. HF.
3.3. Plasma lipids and glucose

Three months on the HF raised plasma TC, LDL cholesterol, and glucose as commonly seen with such nutrition (Table 2). TGs and HDL cholesterol were unchanged. In dMF treated mice, TC and LDL cholesterol were not substantially affected but TGs were substantially reduced even below the control while glucose matched the control concentration. HDL was raised even above the control level with dMF (Table 2).

3.4. Changes to aortic and cardiac histology

The major morbidities of metabolic syndrome arise from vascular insufficiencies including arterial stiffening and cardiac hypertrophy. Thus, the aortic wall became hypertrophied (Fig. 3A and B, Table 3) due in part to increased medial collagen due to increased dietary fat. Treatment with either 100 or 300 mg/kg dMF prevented around 50% of aortic wall-thickening ascribed to collagen accumulation (Fig. 3A and B, Table 3).

The HF also produced a hypertrophy of the left ventricular wall (Fig. 3C) and expansion of constituent cardiomyocyte (Fig. 3D). More collagen deposition was also observed, in particular around the blood vessels of the heart in the HF group (Fig. 3C and D), but this was less pronounced in the dMF-treated groups (Fig. 3D).

3.5. Liver lipid accumulation

Hepatic lipid content was measured directly using ‘oil red O’ that showed sparse staining in the control group, intense in the HF group, and substantial in the dMF-fed groups (Fig. 4). The incremented oil red O contents in the HF group was lowered by 49% in the two dMF groups as summarised in Table 3.

3.6. Autophagy by LC3 and p62 immunohistochemistry

The next experiments aimed to assess the role of autophagy in fatty liver disease by measuring expression of two marker proteins, LC3 (Fig. 5) and p62 (Fig. 6), involved in recycling dysfunctional proteins. The IHC using anti-LC3 antibody showed 4 types of distinct staining patterns of hepatocytes: no staining (0), feint (1+), intermediate (2+), and strong staining (3+). Endothelial and Kupffer cells lining the sinusoids stained intensely (Fig. 5). For a positive control, cerebral neurons also stained (Fig. 5E).

Cells identified as hepatocyte were counted according to staining category (Fig. 5) in 7 photomicrographs taken at random from livers of each of 4 mice. From all of 16 mice used, the average hepatocyte count was 2490 ± 386 (SD) of which 88 ± 56 were graded 1+, 44 ± 36, 2+, and 14 ± 14 3+ across all 112 frames. Stained hepatocytes tended to exist in groups which explained their highly variable count rates. Fig. 5F shows the proportions of all staining categories counted hepatocytes. Fig. 5E shows an IHC positive control using neocortex processed at the same time.

The ubiquitin-binding protein p62 is also involved in hepatic autophagy. The p62 positive cells tended to be confined to regions, and their numbers tended to increase in dMF treated group (Fig. 6E). There is no clear evidence that liver autophagy was dramatically changed for either marker protein.

3.7. Lipid accumulation and lipolysis in adipocytes

3T3-L1 cells were cultured in media that stimulated proliferation and promoted fat accumulation and inhibition of these processes could reduce cell fat content. dMF marginally reduced lipogenesis (Fig. 7A) while the cell viability was unaffected (Fig. 7B). Fat mass can also be lowered by mobilising accumulated TG stores through lipolysis to fatty acids and glycerol that are then metabolised or lost to the medium. Treatment of cells with the lipolysis protocol for just 24 h reduced fat content with the higher dMF concentrations (Fig. 7C) without adversely affected cell viability.
3.8. Bile acid binding activity of dMF

Sequestration of bile acids is used clinically to prevent intestinal fat absorption. While cholestyramine bound strongly to taurocholic acid and taurodeoxycholic acid, binding of dMF to these bile acids was generally undetectable except for very weak binding at 1 g/mL (Table 4) and thus unlikely to influence fat absorption.

4. Discussion

Increased dietary fat fed to mice induced many of the cardiovascular, biochemical risk factors, and histological changes associated with metabolic disease seen in humans. Specifically, HF consumption accelerated body weight gain, promoted visceral and hepatic fat accumulation, changes that were prevented or reduced by concomitant administration of 100 and 300 mg/kg dMF. HF increased fasting blood glucose, plasma TC, LDL and the TC/HDL ratio. Both dMF doses normalised blood glucose, reduced TC/HDL, and remarkably reduced TGs below the control level while elevating HDL above baseline level. The latter suggests that dMF might have benefits in the absence of an atherogenic diet. In contrast, dMF had little impact on TC and LDL. These observations generally reflect mouse studies using statins and mulberry extracts that reduce obesity, diabetes and hyperlipidemia. The hyperglycemia and dyslipidemia in HF mice was accompanied by deterioration in cardiovascular histology, i.e., fibrosis of the aortic wall that diminishes vascular compliance. With accompanying endothelial inflammation, the endothelial vasodilatory signalling is reduced leading to poorer tissue perfusion and feedback compensation that increases cardiac output and arterial
hypertension to maintain tissue perfusion. This sustained cardiac demand drives left ventricular myocyte hypertrophy that was clearly observed in our study. These observations accord characteristics of atherosclerosis.7,8

**dMF Mechanisms of action:** Inflammation underlies metabolic disease and all its cardiovascular ramifications. Anthocyanins are widely known for their beneficial cardiovascular supported by numerous investigations and convincing clinical trials.40 Of the dMF constituents, some polyphenols and especially the total cyanidin oral dose (as 3-glycosides) was ~1.5 mg/kg in 100 mg dMF. Bioavailabilities are low with oral anthocyanin doses higher than administered here, typically achieving blood concentrations of <1 μM and slightly higher for aglycone.41 Cyanidin conjugates are readily hydrolysed by enterocytes and numerous intestinal bacterial species.42 The gut microbiome also cleave the ‘C’ ring to yield two monocyclic polyphenols that are more bioavailable than cyanidins and achieve effective blood concentrations in humans.43 These metabolites have widespread anti-inflammatory effects, especially on vascular endothelial cells, some acting ~100 nM.44 Microbial metabolism of other polyphenols also generate the same monocyclic metabolites augmenting those from cyanidins.45 Polysaccharides may also contribute to dMF actions46 and α-glucosidase inhibition by its galactopyranosyl-1-deoxynojirimycin would help to reduce damaging hyperglycemic peaks in humans. Pharmacologies of other constituents await exploration.

The autophagy marker proteins LC3 and p62 were not convincingly affected by the HF diet nor by dMF as seen in other study.47 A related process, lipophagy may be more relevant to cells engorged with fat and provide a way of fat mobilisation. The doubling of HF food intake and commensurate weight gain is common yet overlooked by many studies but the cause of such hyperphagia is unclear.48 dMF prevented the hyperphagia

**Fig. 6.** Representative photomicrographs of immunohistochemical staining for p62 in brown for the control group (A), HF group (B), HF + dMF 100 (C), and HF + dMF 300 (D). Proportion of p62 positive cells was not different between groups (E). Values are mean ± SEM (n = 4). Control, normal diet; HF, high-fat diet; HF + dMF100, high-fat diet + 100 mg/kg dried mulberry fruit powder; HF + dMF300, high-fat diet + 300 mg/kg dried mulberry fruit powder.
accompanied by a reduced dietary fat load. This makes it difficult to unravel dMF actions that arise solely due to reduced fat load associated with the lower food intake from direct actions on cellular targets.

Clinical applications: Thus, compared to the single target allopathic medicines, dMF potentially acts on a range of cellular targets.

Table 4

Binding of dMF or cholestyramine to bile acids (taurocholic acid and taurodeoxycholic acid) in vitro.

| Concentrations of dMF or cholestyramine (mg/mL) | Bile acid binding             | Taurocholic acid (%) | Cholestyramine (%) |
|------------------------------------------------|-------------------------------|----------------------|--------------------|
|                                                 | dMF (± SEM)       | Cholestyramine (± SEM) | dMF (± SEM)       | Cholestyramine (± SEM) |
| 0                                               | 0.0 ± 0.0         | 0.0 ± 0.0           | 0.0 ± 0.0         | 0.0 ± 0.0             |
| 0.1                                             | 0.0 ± 0.0         | 4.9 ± 11.3          | 0.0 ± 0.0         | 8.2 ± 13.8            |
| 1                                               | 0.0 ± 0.0         | 47.2 ± 3.0**        | 0.0 ± 0.0         | 87.3 ± 3.7*           |
| 5                                               | 0.0 ± 0.0         | 83.6 ± 2.1**        | 0.0 ± 0.0         | 93.8 ± 2.1**          |
| 10                                              | 16.6 ± 5.9##      | 87.7 ± 1.7**        | 10.2 ± 25.5#      | 96.5 ± 2.6**          |

dMF, dried mulberry fruit powder. Values are means ± SEM (n = 3). *P < 0.05 vs. Control.

Fig. 7. Effect of dried mulberry fruit (dMF, 20–1000 μg/mL) in cultured 3T3-L1 adipocytes on (A) Lipogenesis (lipid accumulation) measured by oil red O staining; (B) Cell viability using the MTT assay; (C) Lipid loss by lipolysis; and (D) Cell viability using the MTT assay. Values are means ± SEM (n = 3), *P < 0.05 vs. Control.
pathologies typical of metabolic disease. At 100 mg/kg/day allo-
metrically translates to 1g of dMF for humans, a palatable and easily
consumed supplement. The safety of mulberries, being an
commonly consumed fruit, will always out-perform statin safety.40
Trials and medical advice focus on weight loss but instead need a
multi-pronged approach beginning with dMF supplements and a
profound change of diet. The former reduces inflammation and its
polyphenols assist microbial diversity,41 while the new diet shifts
microbial metabolism away from energy harvesting seen in obesity
to short-chain fatty acid production that improves enterocyte, im-
une, and hepatic function.51 Because metabolic syndrome man-
ifests as a constellation of interacting pathologies, treatment should be
holistic as practiced by traditional healers rather than a mono-
therapy using a single treatment outcome.

Study limitations: Rodents are poor models for human metabolic
disease.52 Commercial animal diets are formulated to optimise
health while human diets are designed to satisfy gustatory drive
causing calorie intake in excess of needs, which may create nutrient
deficiencies, and create intestinal dysbiosis. Here, we show that
mulberries prevent obese pathologies, but treatments for
established obesity in human fat mobilisation no longer operate
because of adipocyte inflammation, and leptin- and sex hormone-
resistance. Nevertheless, researchers have tight control their ani-
mal protocols.

Future studies: Human metabolic disease is best studied in hu-
man trials where the disease is already entrenched. While tight
control of the numerous lifestyle variables is impossible, strict and
detailed participant selection criteria greatly facilitates data inter-
pretation. Studies on dMF or any other herbal should be in the
context of a holistic treatment and include a quality of life outcome:
The trial should aim to: (i) reduce the excess energy input while
improving nutrition, and (ii) removing existing adipocyte or ectopic
fat through muscular utilisation or mitochondrial thermogenesis.
TOF/LC/MS measurement of circulating and fecal metabolites is
crucial for defining the actual agents generating the pharmaco-
logical efficacy.

5. Conclusion

This study suggests that the lyophilised mulberries can
ameliorate dyslipidemia, improve vascular and cardiac function
and promote weight loss in mice and the results suggest translation
to patients with metabolic syndrome. Animal models may be best
used to characterise mechanisms discovered in the human condi-
tion. Contemporary application of traditional medicines should
study the whole herbal product and its pharmacologically active
metabolism in clinical trials. However, to have a major impact on
the constellation of disease processes encompassing metabolic
syndrome, requires a paradigm shift in lifestyle where dMF serves
as an important supplement in the required holistic treatments.

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Declaration of competing interest

The authors declare that there are no conflicts of interest.

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