Comparative Effects of Flavonoids from *Fructus Sophorae* on Rat Osteoblasts *in vitro*

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**Abstract:** *Fructus Sophorae* (FS), the dry mature fruit of *Sophora japonica* Leguminosae, is a valuable traditional Chinese medicine resource with flavonoids as the major active ingredients. To identify the plant-derived estrogen-like flavonoids serving as potential osteoporosis chemopreventive agents, we isolated and identified 27 flavonoids compounds, including 17 compounds obtained for the first time from FS. To screen out the flavonoids with estrogen-like biological behavior from the 11 high yield compounds, we set up an *in vitro* screening system in rat osteoblast MC3T3-E1 cells including MTT assay, alkaline phosphatase staining and Alizarin red S staining assay to examine the effects on cell proliferation, differentiation and mineralization of osteoblast cells respectively. Six flavonoids, including genistein, sophoricoside, sophorabioside, sophoraflavonoloside, nicotiflorin and rutin, significantly increased the cellular activity of MC3T3-E1 cells. Furthermore, blocking the estrogen receptor signaling by tamoxifen compromised the effects above significantly, suggesting the 6 compounds behave as estrogen-like reagents. Moreover, the interaction between the six flavonoids from FS and estrogen receptor was clarified by molecular docking method from Glide XP. Collectively, being used as food and medicine in China, FS is rich in flavonoids with estrogen-like effects, may be used as healthy supplementary in treating postmenopausal women osteoporosis.

**Keywords:** *Fructus Sophorae*; flavonoid; phytoestrogens; estrogen receptor; osteoporosis. © 2019 ACG Publications. All rights reserved.

1. Introduction

Osteoporosis is a bone disease featured with reduced quality of bone mass and micro-architecture, leading to increasing risk of fragility or bone fracture which would cause considerable morbidity and mortality [1,2]. Deficiency of ovarian estrogen production is a major reason for osteoporosis in over 200

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million postmenopausal women worldwide [3, 4]. Currently hormone replacement is the most effective therapy to ameliorate postmenopausal symptoms and lower or prevent osteoporosis, however the increased carcinogenic risk of hormones and synthetic hormone-like drugs [5,6] stimulates the demand of seeking for safe and health natural products for replacement, mostly phytoestrogens and novel estrogen-like compounds from plants [7]. Phytoestrogens (isoflavones, flavonoids, coumestanes, lignans, stilbenes and so on) are polyphenol compounds rich in several edible or medicinal plants (soybeans, flaxseeds and so on), mostly belonging to the Leguminosae family [7,8]. Owing to the structural and functional similarity with steroid hormone, phytoestrogens isolated from plants or herbs are verified to be potential hormone replacement reagent via cell line or ovariectomized rodent animal model [7]. Clinically, to avoid exposure to exogenous hormones and toxic drugs, large populations of postmenopausal women are shifting to phytoestrogens for relief [7,10,11]. However, more scientific evidences in the study on the protective properties of phytoestrogens on osteoporosis are required for broader clinical application.

Fructus Sophorae (FS) or Huaijiao, the dry mature fruit of Sophora japonica Leguminosae, is a well-established traditional Chinese medicine listed in the Chinese Pharmacopoeia, which has been used as a heat-clearing and fire-purging, cooling blood and hemostatic agent in China. Extracts from Sophora japonica were proved to prevent bone loss in ovariecotomized rat’s model [12]. Constituents studies of FS showed that it is a natural excellent source for flavonoids which account 20.2% of the total weight [13,14]. FS flavonoids mainly belong to isoflavonoid and flavonols along with their glycosides, serving as the therapeutic components of FS [15,16]. Further, genistein [17] and sophoricoside [18] isolated from FS show osteoporosis chemopreventive properties. Sophoricoside, genistin, genistein, rutin, quer cetin and kaempferol are the known major compounds of FS, occupying 11.4% of the total weight [13,14]. In our present study, we focused on isolating and identifying the unknown flavonoids in FS, and investigated their osteogenesis effects. Osteogenic effects of the compounds were tested in cultured osteoblasts MC3T3-E1 cells by alkaline phosphatase (ALP) staining. Alizarin red S (ARS) staining was used to evaluate the cellular mineralization. Tests of co-treatment with tamoxifen and molecular docking of estrogen receptor (ER) β protein were carried to verify if the flavonoids interacted with ER [19,20].

2. Materials and Methods

2.1. Materials

FS were purchased from Hebei, China, in November 2014 and authenticated by Professor Guangshu Wang in the department of Pharmacognosy, Jilin University, China. A voucher specimen (No. FSJ-20141012) was prepared and kept in the Jilin University.

MC3T3-E1 cells were gifts from Professor Hui Xu’s lab (Jilin University). Reagents for cell culture, including alpha-Minimum Essential Medium (α-MEM), Fetal bovine serum (FBS), and penicillin/streptomycin were all purchased from Gibco. 3-(4,5-Dimethyl-2-thiazolyl)-2,5-phenyl-2-H-tetrazolium bromide (MTT), 17β-estradiol and tamoxifen were obtained from Sigma. Silica gel (Tsingtao Ocean Chemical Co., Ltd, Shandong, China), Sephadex LH-20 (GE Healthcare Bio-Science AB, Sweden), and ODS (Silicycle, Quebec City, Canada) were used in column chromatography. All other analytical grade reagents were purchased from Beijing Chemical Factory (Beijing, China).

2.2. Extraction and Isolation

FS, the dried fruits of Sophora japonica L. (5.0 kg) were pulverized (120 mesh) and then extracted for 3 h with 70% EtOH (40 L, 3 times) at 50-60°C. The extract solution was removed to afford the EtOH extract, which was then suspended in distilled H2O and successively partitioned with n-Hexane, EtOAc and n-BuOH. The EtOAc soluble fraction (159.0 g) and n-BuOH soluble fraction (1320.0 g) were obtained respectively.
Effects of flavonoids from *Fructus Sophorae* on rat osteoblasts

The *n*-BuOH-soluble fraction (300.0 g) was subjected to silica gel column chromatography, gradiently eluted with CH₂Cl₂-MeOH by stepwise addition of MeOH (20:1-0:1 v/v) to yield 6 fractions. Fraction 1 (4.0 g) was subjected to silica gel (CH₂Cl₂-MeOH 22:1) to obtain compound 14 (40 mg). Fraction 2 (80.2 g) was subjected to silica gel (CH₂Cl₂-MeOH 15:1) and ODS (MeOH-H₂O 25:75:100:0) column chromatography to obtain compounds 3 (39.0 g), 17 (5.8 g). Fraction 3 (68 g) was subjected to ODS (MeOH-H₂O 20:80:100:0) column chromatography and recrystallization (MeOH-H₂O) to obtain compounds 4 (31.1 g), 16 (3.0 g). Fraction 4 (45.5 g) was subjected to silica gel (CH₂Cl₂-MeOH 8:1) to obtain compound 21 (2.0 g). Fraction 6 (11.0 g) was subjected to Sephadex LH-20 (MeOH) chromatography to obtain compound 22 (35.0 mg).

The EtOAc-soluble fraction (159.0 g) was subjected to silica gel column chromatography, gradiently eluted with CH₂Cl₂-MeOH by stepwise addition of MeOH (30:1-0:1, v/v) to yield 7 fractions. Fraction 2 (12.0 g) was subjected to silica gel (PE-EtOAc-acetone-H₂O, 10:4:1:1, upper phase), Sephadex LH-20 (MeOH) chromatography and HPLC (RP-18, MeOH-H₂O 60:40) chromatography to obtain compounds 1 (0.7 g), 5 (9 mg), 8 (12 mg), 9 (58 mg), 10 (5 mg), 11 (3 mg), 12 (26 mg), and 26 (11 mg). Fraction 3 (31.6 g) was subjected to silica gel (EtOAc-Acetone, 40:1-1:1), Sephadex LH-20 (MeOH) chromatography, HPLC (RP-18, MeOH-H₂O 58:42) chromatography and crystallization (MeOH-H₂O) to obtain compounds 6 (103.2 mg), 13 (0.5 g), 18 (8 mg), 25 (0.3 g) and 27 (0.2 g). Fraction 4 (28.0 g) was subjected to silica gel (CH₂Cl₂-MeOH 10:1) to obtain compound 15 (3.8 g). Fraction 5 (16 g) was subjected to Sephadex LH-20 (MeOH) chromatography to obtain compounds 7 (11.0 mg), 19 (6.5 mg), 23 (6.5 mg) and 24 (5.9 mg). Fraction 6 (14.3 g) was subjected to Sephadex LH-20 (MeOH) chromatography and HPLC (RP-18, MeOH-H₂O 55:45) to obtain compounds 2 (2.1 g) and 20 (37 mg).

### 2.3. Preparation of Test Samples

Compounds 1, 2, 3, 4, 13, 15, 16, 17, 21, 25 and 27 were dissolved into 0.1M stock solution by DMSO respectively, and further diluted by culture medium in different concentrations before use (0.1mM, 1.0μM, 0.01μM). 17β-estradiol, the positive control and tamoxifen, the estrogen-receptor inhibitor, are both solved into the final concentration of 5.0μM. The final concentration of DMSO was controlled within 0.1% (v/v).

### 2.4. Cell Culture

MC3T3-E1 cells were maintained at 37 °C with 5% CO₂ in α-MEM (10% FBS and 1% penicillin/streptomycin). To induce differentiation, 50 mg/L L-ascorbic acid and 10 mmol/L β-glycerophosphate were supplemented in the culture medium.

### 2.5. MTT Proliferation Assay

The cell viability of MC3T3-E1 cells treated by the test compounds was determined by MTT assay. Briefly, MC3T3-E1 cells were seeded in 96-well culture plates at a density of 5x10⁴ cells/well. After culture for 24 h, the cells were treated with varying concentrations of test compounds for another 2 days. 20 μl MTT solution (5 mg/ml) were added into each well and incubated for 4 h to allow the formation of purple formazan crystal. For the co-treatment with the estrogen-receptor inhibitor, tamoxifen was added 4 h before the drugs administration. Discarded the supernatant and dissolved the formazan crystals with 150 μl DMSO followed by 10 min incubation in a shaker. Measure the absorbance by Model 50 (Bio-Rad, USA) at 490 nm.
2.6. Alkaline Phosphatase Staining

MC3T3-E1 cells were seeded in 24-well plates at a density of $2 \times 10^4$ cells/well in differentiation medium. In the negative control, the estrogen-receptor inhibitor (tamoxifen) was added 4 h before the drugs pretreated the MC3T3-E1 cells. After 14 days, ALP staining was measured via modified calcium-cobalt (Ca-Co) method [21]. After 10 min fixation by 90 % alcohol and 4 h incubation at 37 °C, the cells were stained by 2% cobalt nitrate and 1% ammonium sulfide in turn [21]. Wash the cells and took photographs under the microscope. The ALP-positive cells exhibit dark-brown color in the bright field.

2.7. Mineralization Assay

Alizarin red S (ARS) staining was a method for the measure of mineralization [22]. Seeded the MC3T3-E1 cells in 24-well at a density of $2 \times 10^4$ cells/well in differentiation medium for 21 days. After washing with PBS twice, fixed the specimens with 95 % ethanol at room temperature for 10 min. Rinsed the plate with distilled H$_2$O, and stained the cells with 0.1% alizarin red S solution (pH 8.3) for 30 min at 37 °C. Rinsed the cells with distilled H$_2$O and took photographs under the microscope. Cells with pink color are the positive ARS staining suggesting the formation of bone nodule.

2.8. Molecular Docking within Estrogen Receptor

Molecular docking studies of anti-osteoporosis bioactive compounds were conducted using Schrödinger Maestro (2015-2, Schrödinger, USA). The structure information of Human ER β binding with genistein was downloaded from PDB websites (1QKM, www.rcsb.org/pdb/) [23]. Optimized the structure of ERβ using GLIDE software (v.6.7, Schrödinger, USA) with the default constraint set of 0.30 Å RSD [24]. The receptor grid box was restricted to the size of 2 nm at the active site [25]. The graphs of docking results were plotted with PyMol (Schrödinger, USA).

2.9. Statistical Analysis

All of the experiments were performed at least for three independent times. The values in Figure 2 and 5 were presented as means ± standard deviation (SD). One-way ANOVA analysis and graph plotting was perform via GraphPad Prism 5 software package (GraphPad Inc., San Diego, USA). P-value under 0.05 was considered to be statistically significant.

3. Results and Discussion

3.1. Extraction and Isolation

The 70% EtOH extract of FS was partitioned with n-Hexane, EtOAc and n-BuOH. The EtOAc and n-BuOH soluble fractions were separated and purified on silica gel, ODS, Sephadex LH-20 columns and semi-preparative HPLC to produce 27 compounds. The structures of compounds 1-27, shown in Figure 1, were identified by physicochemical properties, NMR spectral analyses and comparison with known samples as isoflavones, genistein (1), genistin (2), sophoricoside (3), sophorabioside (4), biochanin A (5), sissotrin (6), orobside (7), pratensein (8), prunetin (9), 3'-methylorobol (10), dehydroferreirin (11), cajanin (12), tectoridin (13), ononin (14); flavonols, kaempferol (15), sophoraflavonolioside (16), nicotiflorin (17), α-rhamnoisorobin (18), populnin (19), astragalalin (20), rutin (21), quercetin-3-O-β-D-sophoroside (22), quercitrin (23), isoquercitrin (24); flavone, thevetiaflavone (25), genkwanin (26) and dihydroflavone, choerospondin (27).
Effects of flavonoids from *Fructus Sophorae* on rat osteoblasts

Figure 1. Structures of the 27 compounds isolated from FS in this study. Including 14 isoflavones (1-14), 10 flavonols (15-24), 2 flavones (25 and 26) and 1 dihydroflavone (27).
Among them seventeen compounds 5-12, 13, 17, 19, and 22-27 were first reported from FS, and six compounds 10, 11, 19, 22, 25, 26 and 27 were obtained from genus *Sophora* Linn. for the first time (https://scifinder.cas.org). Details about the structure analysis of the chemicals would be described in supplemental data 1.

3.2. Primary Screening of High Yield Compounds on the Proliferation of MC3T3-E1 Cells by MTT

As shown in Figure 2 (A-B), each compound from 11 high yield flavonoids (compounds 1, 2, 3, 4, 13, 15, 16, 17, 21, 25 and 27) was diluted into 3 concentrations, 0.01 μM, 1.0 μM and 100 μM. For each compound, there are also a blank group as negative control and 17β-estradiol (5.0 μM) as positive control. After cell culture for 48 h, we performed MTT assay to examine the effects of the compounds on the proliferation of MC3T3-E1 cells. Compounds 1, 3 and 15 promoted proliferation of MC3T3-E1 cells at the concentration of 0.01 and 1.0 μM, but 3 and 15 showed no significant effects at 100 μM, and 1 even inhibited proliferation at 100 μM.

![Figure 2](image-url) Primary screening of 11 high yield compounds on the proliferation of MC3T3-E1 cells by MTT assay. (A) Effects of 1, 2, 3, 4, 13 and 15 on osteoblast proliferation of MC3T3-E1 cells. (B) Effects of 16, 17, 21, 25 and 27 on osteoblast proliferation of MC3T3-E1 cells. For each compound, there are one negative control (blank group), one positive control (5.0 μM 17β-estradiol), and 3 different concentrations of each chemical (0.01 μM, 1.0 μM and 100 μM). MTT assay of each compound has been analyzed by cell viability (%), which is calculated using following formula. Cell viability (%) = (OD_{490} of test compound wells/OD_{490} of control wells) ×100%. Each assay was repeated for three times independently. (*p < 0.05, **p < 0.01.).
Other compounds showed a dose-dependent effect on the MC3T3-E1 cells proliferation. Among them, cells treated with compounds 4 and 13 have a significant promoting effect at the concentration of 0.01 μM, while compounds 25 and 27 treatment have this effect only at the high concentration (100 μM) comparing to the control group. At the concentration of 1.0 μM, compounds 1, 2, 3 and 15 exhibit the maximal cell viability of 133.8%, 127.2%, 138.7% and 118.8% respectively (p<0.01 for all). While at the concentration of 100.0 μM, compounds 4, 13, 16, 17, 21, 25 and 27 reach the maximal cell viability of 140.4%, 129.6%, 131.1%, 122.6%, 126.8%, 117.0% and 116.0% respectively (p<0.05 for 25 and 27, and p<0.01 for the rest). For the following experiments, the concentration of each chemical is set as 1.0μM (9 compounds excepting 25 and 27) or 100μM (compound 25 and 27) according to the MTT assay results.

Results of genistein (1) and sophoricoside (3) are consistent with the previous work in FS [17, 18]. Other 4 compounds were first reported as the chemoprotective osteoporosis compounds in osteoblast cells (rat MC3T3-E1 cells), especially nicotiflorin (compound 17) were obtained from FS for the first time. The test of proliferative activity on MC3T3-E1 cells showed that 11 compounds promoted cells proliferation in different degrees. But the structure-activity relationship is not distinct due to the insufficient of sample quantity.

![Compounds Only (-Tamoxifen)](image1)

![Compounds +Inhibitor (+Tamoxifen)](image2)

**Figure 3.** Effects of 11 compounds with or without tamoxifen on ALP staining in MC3T3-E1cells. Group A is the blank group as negative control. The concentration of each chemical is set as 1.0μM ( 9 compounds excepting 25 and 27) or 100μM (compound 25 and 27) according to the MTT assay results of Figure 2. After ALP staining, 6 compounds from the 11 high yield compounds have obvious stronger signal than blank group, including 1(B), 3(C), 4(D), 16(E), 17(F) and 21(G). Other 5 groups have similar intensity signal with blank groups are not shown in Figure 3. From A’ to G’ groups, the inhibitor of ER signaling, tamoxifen (shorten as +T), was added in each group with a final concentration of 5.0μM comparing to A to G groups respectively. ALP signaling was significantly reduced to the blank level after tamoxifen administration as shown in group B’ to group G’. Arrows stand for the representative ALP staining positive cells. Each assay was repeated for three times independently.

### 3.3. Effect of Compounds on the Differentiation of MC3T3-E1 Cells

ALP enzyme participated in the regulation of calcium deposition and was generally used as a key marker in early differentiation process of osteoblast [26,27]. To investigate effects of compounds on the differentiation of MC3T3-E1 cells, we have performed ALP staining. As shown in Figure 3, treatment with compounds 1, 3, 4, 16, 17 and 21 at concentrations of 1.0 μM showed significant ALP positive staining with several black granules deposition in MC3T3-E1 cells comparing with blank control (Figure 3 A-G). The other compounds fail to have the obvious difference ALP positive signal comparing to the
The results suggested that these six compounds (compounds 1, 3, 4, 16, 17 and 21) could enhance ALP activity in MC3T3-E1 cells groups.

3.4. Effect of Compounds on the Mineralization of MC3T3-E1 Cells

In the process of osteoblast differentiation, a large number of calcium ions could deposit and form mineralized nodules, which could be stained by alizarin red S (ARS) with pink or pale purple color. We examined the mineralization effects of 6 ALP positive chemicals (compounds 1, 3, 4, 16, 17 and 21) of MC3T3-E1 cells through ARS staining. Highly consistent with the ALP staining results above, compounds 1, 3, 4, 16, 17, and 21 exhibited increased ARS positive area at concentrations of 1.0 μM comparing to the control group. As shown in Figure 4, the pink positive staining cells treated with compounds 1, 3, 4, 16, and 21 were significantly increased, and the mineralized nodules were obviously formed compared with control. These results indicated that these five compounds could enhance mineralization of MC3T3-E1 cells. The positive staining cells were increased and clustered treated with 17, but no obvious mineralized nodules were formed, indicating that the mineralization of 17 was weak.

![Figure 4](image)

**Figure 4.** Effects of compounds on Alizarin red S staining in MC3T3-E1 cells. Only 6 chemicals with ALP positive signaling (Compounds 1, 3, 4, 16, 17 and 21) were tested by Alizarin red S staining. Arrows indicated the Alizarin red S staining positive cells. Each assay was repeated for three times independently. a: Concentration 1.0μM (B-G); b: Concentration 100μM(B’-G’).

Positive results in ALP staining and calcification tests confirmed the cultured cells have the typical characteristics of mature osteoblast. In this study, 11 compounds with the capable of promoting osteoblast proliferation, among them, 6 compounds (1, 3, 4, 16, 17 and 21) could enhance ALP activity and promoting calcium salt deposition of osteoblast compared with control, suggesting that the compounds could improve the osteogenic properties of osteoblast, enhancing bone density.

3.5. Compounds Promote Proliferation and Differentiation of MC3T3-E1 Cells via Estrogen Receptor Pathway

To investigate whether the 6 compounds isolated from FS could interact with the endogenous ER signaling of MC3T3-E1 cells, we treated the rat cells with the compounds together with tamoxifen, inhibitor of ER pathway, and examined by ALP (Figure 3 A’-G’), and MTT assay (Figure 5) respectively. Compounds 1, 3, 4, 16, 17 and 21 used in the above assay were all diluted into the concentration of 1.0 μM according to the results in Figure 2. As shown in Figure 3, compounds co-treated with tamoxifen (Figure 3, B’-G’) compromise the ALP signal by compounds only groups (Figure 3, B-G). Consistently, all the chemicals would increase the OD value remarkably at 490 nm, while decreased to the blank group.
level by in the tamoxifen added groups (Figure 5). From the two independent assays, the 6 compounds have the estrogen-like biological activities to promote proliferation and differentiation of MC3T3-E1 cells.

Results showed that, co-treatment with tamoxifen and compounds significantly decreased MC3T3-E1 cells proliferation and differentiation comparing to exposure to compounds alone. These suggested that ER signaling influence the anti-osteoporosis properties of the compounds.

![Figure 5. Effects of compounds with or without tamoxifen on MTT assay in MC3T3-E1 cells. The groups with ALP and Alizarin red S signal positive (Compounds 1, 3, 4, 16, 17 and 21, at concentration of 1.0μM) were verified by MTT assay. Different with assay in Figure 2, co-treatment with tamoxifen was also one group for each compound. Consistent with the results in Figure 2, the cell viability of the these groups has remarkably increased, while decreased to the blank group level by in the tamoxifen added groups. Each assay was repeated for three times independently. (*p < 0.05, **p < 0.01.).](image)

### 3.6. Compounds Could be Docked into the Active Pocket of ER Protein via Molecular Docking

Molecular docking in the active pocket on ERβ protein were performed to clarify the mode of compounds 1, 3, 4, 16, 17 and 21 interaction with ER, and to localize their accurate binding sites. As shown in Figure 6, all compounds bind to ERβ protein, some similar binding sites of the observed compounds were revealed. The binding amino acid residues, His475, Arg346, Leu339, Glu305 and Met336 were commonly shared between ligands and ERβ protein. Compound 1 binds to ERβ by the A-ring phenol interacting with His-475 at the right, and the C-ring phenol interacting with hydrogen bond network of Glu-Arg-H2O triad at the left. Compounds 3, 16, 17 and 21 bind to ERβ by the A-ring phenol interacting with Glu-Arg-H2O triad, and hydroxyl of the sugar ligand interact with His-475. Compound 4 binds to ERβ by the sugar hydroxyl interacting with the Glu-Arg-H2O triad. The C-ring phenol of 4, 16, 17 and 21, interacting with the Leu 476.
Molecular docking by Schrödinger software showed that compounds 1, 3, 4, 16, 17, and 21 could form stable hydrogen bonds with the amino acid residues in the active pocket of ERβ respectively, which is consistent with in vitro experiments results. Hydroxy groups of the compounds could play an important role in achieving high binding affinities by interacting with the Glu-Arg-water triad (Glu305/Arg346) and His 475 in ERβ. Above results indicated that one of the anti-osteoporosis mechanisms for these six compounds was by activating ERβ receptor.

Hormone replacement is considered to be a popular regime for osteoporosis prevention and treatment. However, recent evidence suggests that hormone used increases the risk of endometrial and breast cancers. Therefore, phytoestrogens therapy as an alternative treatment option is becoming more popular throughout the world [7]. Flavonoids with diverse structures, but most are similar to estrogen [28], is a natural product library of potential phytoestrogens. From our present study, these compounds are more prone to serve as potential chemopreventive agents in the therapy of osteoporosis, particularly in postmenopausal. Being used as food and medicine in China, FS is rich in flavonoids with estrogen-like effects, may be used as healthy supplementary in treating postmenopausal women osteoporosis.
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