Yam Extracts Increase Cell Proliferation and Bone Matrix Protein
Collagen Synthesis of Murine Osteoblastic MC3T3-E1 Cells

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Abstract

Yam extracts (Dioscorea batatas) have been reported to possess a variety of functions. However, studies on
its osteogenic properties are limited. In this study, we investigated the effect of ethanol and water extracts on
osteoblast proliferation and bone matrix protein synthesis, type I collagen and alkaline phosphatase (ALP), using
osteoblastic MC3T3-E1 cell model. MC3T3-E1 cells were cultured with yam ethanol and water extracts (0~30
mg/L) within 39 days of osteoblast differentiation period. Cell proliferation was measured by MTT assay. Bone
matrix proteins were assessed by the accumulation of type I collagen and ALP activity by staining the cell layers
for matrix staining. Also, the secreted (media) matrix protein concentration (type I collagen) and enzyme activity
(ALP) were measured colorimetrically. Yam ethanol and water extracts stimulated cell proliferation within
the range of 15~30 mg/L at 15 day treatment. The accumulation of type I collagen in the extracellular matrix, as
well as secreted collagen in the media, increased with increasing doses of yam ethanol (3~15 mg/L) and water
(3~30 mg/L) extracts. ALP activity was not affected by yam ethanol extracts. Our results demonstrated that
yam extracts stimulated osteoblast proliferation and enhanced the accumulation of the collagenous bone matrix
protein type I collagen in the extracellular matrix. These results suggest that yam extracts may be a potential
activator for bone formation by increasing osteoblast proliferation and increasing bone matrix protein type I
collagen. Before confirming the osteogenic action of yam, further studies for clarifying how and whereby yam
extracts can stimulate this osteogenesis action are required.

Key words: yam (Dioscorea batatas), MC3T3-E1 cells, proliferation, type I collagen, alkaline phosphatase (ALP)

INTRODUCTION

Bone formation is characterized by the formation of a collagen-rich extracellular matrix after which, ex-
pression of genes associated with differentiated osteoblast follows; alkaline phosphatase (ALP) and para-
thyroid hormone protein receptor are early markers while osteopontin and osteocalcin are late markers (1). There-
fore, bone formation including extracellular matrix mineralization by differentiated osteoblasts is an exquisitely
complex process that follows a temporal sequence of gene expression that eventually culminates in the miner-
alization of the matrix secreted by osteoblasts. Unlike any other tissues, bones are continuously being re-
modelled to maintain constant bone volume via a delicate balance between bone formation by osteoblasts and
bone resorption by osteoclasts. An increase in bone resorption over bone formation can lead to most adult skel-
etal diseases including osteoporosis, the most prevalent metabolic bone disorder (2). Until recently, therapies
aimed to ameliorate skeletal diseases, most notably osteoporosis, are usually focused on diminishing resorption
rather than increasing bone formation. However, more concern is focused recently in the determination of ther-
apeutics that might aid in bone formation by osteoblast that could provide more useful approach in the treatment
of bone loss in osteoporosis.

There has been an increasing interest in the utilization of food-origin products in the treatment of diseases since
they are basically safe and inexpensive. Moreover, numerous edible food-origin products are showing tremen-
dous potential for the treatment of bone-related diseases. For example, it has been reported that a number of com-
mon vegetables including onion, garlic and parsley, can inhibit bone resorption in ovariectomized rats (3).
Curcumin, which is an active compound of yellow spicy turmeric, have been reported to suppress receptor activ-
tor of NF-κB ligand (RANKL) and consequent osteo-
clastogenesis (4).

Yams (Dioscorea batatas) are tuberous vines that are
widely distributed in East Asia including China, Korea,
and Japan. Plants of the genus Dioscorea have been used
for edible starch food sources and traditionally used in
orlent societies. In Korea, yam is consumed as medical and functional food. Recently, several studies have suggested the potential of several species of yam to inhibit bone resorption, as well as to promote bone formation. The phytochemicals from yam such as glycosides, lignans, diosgenin, and dioscin have all been reported to have anti-osteoporotic activity both in vitro and in vivo (5-8). In our previous study, we reported that diosgenin, a major component of yam, showed osteogenic action through upregulating bone-specific transcription factor RunX2 and bone matrix protein synthesis in osteoblastic MC3T3-E1 cells (9). However, the in vitro effect of yam extracts on osteoblastic MC3T3-E1 cell proliferation and their effect on the accumulation of type I collagen and ALP, the bone matrix proteins crucial for extracellular matrix mineralization, have not been reported.

This study aimed to assess the osteogenic properties of extracts of yams grown locally within the vicinity of Andong, Kyungbook Province, South Korea. Specifically, this study was undertaken to examine the effect of various concentrations of yam ethanol and water extracts on the cell proliferation and synthesis of the bone matrix proteins, type I collagen and alkaline phosphatase (ALP), using murine osteoblastic MC3T3-E1 cell model. Findings from this study might provide an insight into the potential role of yam extracts in enhancing bone formation.

MATERIALS AND METHODS

Reagents

Antibodies for target proteins (type I collagen) were obtained from Santa Cruz Biotechnology (Santa Cruz, CA, USA) while antibody for loading control (GAPDH) was obtained from Cell Signaling (Beverly, MA, USA). Cell culture reagents including α-minimum essential media (α-MEM), fetal bovine serum, and penicillin/streptomycin were obtained from Gibco® (Grand Island, NY, USA). Sodium pyruvate, β-glycerophosphate, L-ascorbic acid were all obtained from Sigma (St. Louis, MO, USA). The BCA Protein assay kit, RIPA buffer, and the Super Signal West Pico Chemiluminescence detection reagents were from Pierce Biotechnology (Rockford, IL, USA) while the PVDF membrane was obtained from Millipore (Billerica, MA, USA). All other reagents used were from Sigma.

Yam extracts preparation

Yam (Dioscorea batatas) grown locally within the vicinity of Andong, Kyungbook Province, South Korea was provided by Bookwho Nonghyup which is the major area for yam production in Andong. Yam was extracted with 2 liters of 80% methanol for 48 hours. The extracted yam solvent was filtered and concentrated by rotary evaporation. After concentration, the extracted yam was graduated in ethanol (EtOH), hexane (Hex), chloroform (CH3Cl), ethyl acetate (EtOAc), butanol (BuOH) and water (H2O).

Cell culture

Mouse osteoblastic MC3T3-E1 cells were seeded at a density of 1×10⁵ cells/mL and cultured in regular growth culture media containing α-minimum essential medium (α-MEM), supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate, and 100 units/L penicillin and 100 mg/L streptomycin in a humidified atmosphere of 5% CO₂ at 37°C. At 80% confluence, the cells were cultured in differentiation media as growth media supplemented with 10 mM β-glycerol phosphate and 50 μg/mL L-ascorbic acid as osteogenic differentiation medium. Cells were treated with various yam extracts (EtOH, Hex, CH₃Cl, EtOAc, BuOH, and H₂O) for cell proliferation assay. Also cells were treated with 1~30 mg/L of yam ethanol and water extracts for the assay for type I collagen and ALP. The medium was changed every 2~3 days. The cells cultured in normal osteogenic differentiation medium without yam extracts were used as osteogenic control (OSM).

Cell proliferation assay

Cell proliferation was determined by MTT assay. Briefly, cells (1×10⁵ cells/well in 96-well plate) were maintained in growth media for 24 hr at 5% CO₂, 37°C. At 90% confluence, the cells were treated with various concentrations (1, 3, 5, 10, 15 and 30 mg/L) of yam extracts for 6, 15 and 39 days. Following the respective time treatment, MTT was added to the cell cultures and samples were incubated at 37°C for 3 hr. The production of formazan by the reduction of MTT was measured for the indication of mitochondrial dehydrogenases enzyme activity of viable cells. Absorbance for enzyme products was measured at 570 nm using an optical 96-well microplate reader (Sunrise Absorbance Reader, Tecan, GmbH, Grödig, Austria).

Protein quantification

Osteoblastic MC3T3-E1 cells were cultured on 6-well plates and treated with 100 mg/L of yam water, butanol, ethanol, and hexane extracts for 6 days. Cells cultured using normal osteogenic medium (OSM) were used as control. After treatment, cells were harvested and lysed with RIPA buffer (150 mM NaCl, 1% Nonidet P-40, 1% sodium deoxycholate, 0.1% SDS, 25 mM Tris-HCl, pH 7.6, supplemented with 1% protease inhibitor). Protein
amounts were determined using Pierce BCA assay kit.

**Western blot analysis**

Whole cell lysates (30 µg) were separated by 10% SDS-PAGE and transferred to PVDF membrane by Immobilon-P. The membranes were blocked with 5% non-fat milk and incubated with the primary antibodies for type I collagen and GAPDH. The membranes were then incubated with secondary antibody conjugated to horseradish peroxidase and were developed using Super Signal West Pico Chemiluminescence detection reagents.

**Collagen measurement**

Medium collagen concentration measurement: The amount of medium collagen was measured using the Picro-Sirius red method. Fifty micro liters each of the collagen standard, protein and medium samples were added on each well of 96-well plate and were allowed to dry at 37°C in humidified atmosphere for 24 hours for denaturation of the proteins. After drying, the wells were washed with 200 µL distilled water and 100 µL of 0.1% Picro-Sirius red dye in picric acid was added on each well to combine with the connective protein collagen. The samples and standards were then incubated with the dye for 1 hour at 37°C. After incubation, the samples and standards were washed three times with 200 µL of 10 mM HCl to remove the unbound dye. Finally, the bound collagen was dissolved by adding 200 µL of 0.1 M NaOH for 5 minutes. The samples and standards were then transferred to a clean 96-well plate and absorbance was read at 540 nm.

Cell matrix collagen staining: The synthesis of collagen was assessed by staining with Van Gieson. Cells were washed with distilled water and were allowed to dry completely. The cells were fixed with 2% formaldehyde at 4°C for 15 minutes. The cells were then washed twice with distilled water and stained with Van Gieson reagent for 15 minutes. Excess dye was removed by washing with at least three changes of distilled water. Collagen in the extracellular matrix is stained red.

**ALP activity measurement**

Medium ALP activity: Medium ALP activity was measured in cells treated with 1, 3, 5, 10, 15 and 30 mg/L yam ethanol and water extracts for 39 days. Media were collected for the measurement of the secreted ALP. The ALP activity in media was measured using p-nitrophenyl phosphate as substrate and the product of enzyme activity, p-nitrophenol, was measured as optical absorbance at 405 nm as previously described (9). The activity of medium ALP was expressed as nmol PNP (p-nitrophenyl) /mL medium/minute, which showed ALP activity as enzyme products produced per sample amount (mL) per minute from the substrate.

Cell matrix ALP staining: Cultured cells were rinsed with phosphate buffer saline (PBS, pH 7.4) and fixed with 2% formaldehyde. The cells were stained using Naphthol As-Mx phosphate disodium salt as a substrate for enzyme activity, N,N-dimethyl formamide, and fast red salt as dye for 30 min at 37°C, or until yellow color appeared. After washing with PBS, the cells were photographed. The products of ALP activity were stained red as an indicator for the products of enzyme activity.

**Statistical analysis**

Data were analyzed using software SPSS 18.0. Values for cell proliferation, ALP activity and collagen concentration are presented as mean±SEM. The data analysis was performed using one way ANOVA and Tukey’s HSD test was used as post hoc test if significance was detected among the treatments at the level of p<0.05.

**RESULTS**

**Cell proliferation of various yam extracts and expression of type I collagen**

The effect of various yam extracts (100 mg/L) on the proliferation of osteoblastic MC3T3-E1 cells was determined using MTT assay at 1 and 5 days of treatment (Fig. 1A). Cell proliferation was significantly higher in ethyl acetate, ethanol, water, and hexane extracts, compared to chloroform and butanol after 1 day treatment. However, extending the treatment up to 5 days showed a decrease in cell proliferation of ethyl acetate extract, while ethanol and water extracts showed consistently higher proliferation hence they are used for succeeding experiments. The expression of type I collagen as affected by extracts was analyzed by Western blotting after 6 days treatment (Fig. 1B). Butanol extract interestingly showed the highest type I collagen expression albeit its lower proliferation compared with other extracts while ethanol and water extracts both showed higher type I collagen protein expression than the normal osteogenic control (OSM).

**Cell proliferation of yam ethanol and water extracts**

Using our proliferation and protein expression data described in Fig. 1 above, we decided to utilize yam ethanol and water extracts for further experiments. The effects of serial concentrations (1 ~ 30 mg/L) of both yam ethanol and water extracts on the proliferation of MC3T3-E1 cells for 6, 13, 15, and 39 days treatment were examined using MTT assay (Fig. 2A & B). At 6 days treatment, both yam ethanol and water extract slightly stimulated proliferation but the increase was not statistically significant. A significant and dose-dependent stimulation
of proliferation was observed at 15 days treatment for ethanol and at 13 days treatment for water extracts. No significant differences on proliferation were observed on both yam ethanol and water extracts after extending the treatment to 39 days. The protein synthesis by osteoblastic cells treated by yam ethanol and water extracts was measured by BCA protein assay after 6 days treatment (Fig. 2C). The protein synthesis showed the trend of higher protein content as observed with yam ethanol extract treatment, even without statistical significance.

**Collagen synthesis and secretion**

The effect of yam ethanol and water extracts on the accumulation of collagen in the extracellular matrix was measured by Van Gieson staining at 15 days (Fig. 3A). After 15 days treatment, yam ethanol extract dose-dependently increased collagen accumulation up to 15 mg/L, but increasing yam ethanol extract concentration to 30 mg/L decreased collagen accumulation. On the other hand, yam water extracts dose-dependently increased collagen accumulation from 3 to 30 mg/L (Fig. 3A). The amount of collagen secreted into the media was measured after 39 days treatment. Both yam ethanol and water extracts dose-dependently increased the secreted collagen, and this pattern is more prominent in yam ethanol.

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**Fig. 1.** Effect of various yam extracts on osteoblastic MC3T3-E1 cell proliferation (A) and type I collagen expression (B). (A) At 90% confluence, MC3T3-E1 cells were treated with various yam extracts (100 mg/L) for 1 and 5 days and cell proliferation was measured by MTT assay. The results are presented as absorbance at 570 nm which is the indirect indication of the intensity of live cells. The effect of the different yam extracts for cell proliferation was analyzed by one-way ANOVA at p<0.05 followed by Tukey as post hoc test. Different superscripts indicated significant differences among various yam extract treatments (n=6). (B) Type I collagen protein expression was analyzed by Western blot after 6 days yam extracts treatment (100 mg/L). EtOH: ethanol, Hex: hexane, CHCl3: chloroform, EtoAc: ethyl acetate, BuOH: butanol, H2O: water, OSM: normal osteogenic media.

**Fig. 2.** Dose-dependent effect of yam ethanol (A) and water (B) extracts on osteoblastic MC3T3-E1 cell proliferation and protein synthesis (C). (A, B) Cell proliferation was measured by MTT assay and presented as absorbance at 570 nm. The effect of the different yam extracts was analyzed by one-way ANOVA at p<0.05 followed by Tukey as post hoc test. Different superscripts indicated significant differences among treatments (n=6). V: vehicle. (C) Protein concentration was measured by BCA protein assay. No significant difference in protein contents with respect to yam extracts treatment was observed (n=3). EtOH: ethanol, H2O: water, OSM: normal osteogenic media.
extract treatment (Fig. 3B, right panel). Yam ethanol extracts increased collagen accumulation more prominently within the range of 5–30 mg/L, while yam water extracts was within 15–30 mg/L indicating that yam ethanol extract may be more potent in increasing collagen content for the collagen accumulation in cell matrix.

**ALP activity stimulation and medium ALP activity**

The influence of yam extracts on cell layer ALP activity in extracellular matrix was measured by ALP staining using Fast red salt as dye. ALP activity was more stimulated by yam ethanol extracts compared with yam water extracts (Fig. 4A). However, within the concentration range (0–30 mg/L) analyzed for both yam ethanol and water extracts, there was no observed dose-dependent ALP activity stimulation. The activity of ALP secreted in the medium was measured at 39 days treatment (Fig. 4B). Yam ethanol extracts slightly decreased media ALP activity while yam water extract did not show any significant change.

**DISCUSSION**

The balance between bone formation and resorption must be delicately maintained to ensure the integrity of the skeleton. So far, a number of studies show the potential of several dietary components and natural products in promoting bone formation and inhibiting bone resorption giving a net positive result on the skeleton. A wide range of dietary products with therapeutic effect on bone formation have been reported and several natural compounds have been shown to enhance osteogenic differentiation (9-13), such as red yeast rice (10), black cohosh (14), and green tea (13). The effect of yam extracts and their bioactive component on bone formation *in vitro* (5,9,15), *in vivo* (8,16-18), and in human population samples (19) have been reported and so far all showed potential in the treatment of bone-related diseases, particularly in osteoporosis.

However, *in vitro* studies on the effect of yam on bone formation via its effect on osteoblast proliferation and its effect on bone matrix proteins have been limited. Our previous study on Diosgenin, one of the bioactive components extracted from yam, showed a positive role on bone formation via increasing the bone-specific transcription factor Runx2 as well as the bone matrix proteins type I collagen, ALP, and osteopontin (9). Thus, in the present study, we investigated the effect of yam ethanol and water extracts on the proliferation and bone matrix synthesis (type I collagen and ALP) using osteoblastic MC3T3-E1 cell model.

Bone formation is largely dependent on the executive cells and increasing osteoblast number could ensure that an efficient extracellular matrix is properly laid out for mineral matrix deposition, since the osteoblasts synthe-
size and secrete the bone matrix proteins. Hence, dietary compounds that could enhance osteoblast cell proliferation and differentiation are beneficial to bone formation. In the present study, we showed that both yam ethanol and water extracts increased cell proliferation indicating the ability of yam extracts to increase osteoblast number. This was observed at about 6 days to 13~15 days of treatment which coincides with the osteoblast in vitro proliferation and differentiation stage, respectively. The increase in cell number in these stages may indicate that more cells can participate in the secretion bone matrix protein and synthesis of a collagen-rich extracellular matrix.

The formation of a collagen-rich extracellular matrix is crucial for mineralization since collagen serves as the structural scaffold to which minerals are later embedded in the matrix. About 90% of the extracellular matrix is made up of collagen (mainly type I collagen) and aside from its structural role, it also influences osteoblast structure, differentiation, and gene expression (20) and this may be likely mediated through the interaction of collagen with the heterodimeric integrin receptors (21). In the present study, we showed that both yam ethanol and water extracts could increase collagen accumulation in the extracellular matrix indicating a potential role of yam extracts in enhancing bone formation. Culturing of osteoblast cells without ascorbic acid, a cofactor for collagen synthesis, showed a fivefold reduction in ALP activity and suppression of mineralization (20) which further emphasizes the crucial role of collagen synthesis in mineralization. Ascorbic acid stimulates procollagen hydroxylation and fibril assembly which is followed by robust induction of osteoblast associated genes including ALP (22).

Likewise, yam ethanol, but not water extracts, was found to slightly enhance ALP activity in the extracellular matrix, although no dose-dependent stimulation was observed. This may indicate that higher doses of yam extracts might be necessary to enhance ALP activity. ALP, a homodimeric metalloenzyme, is the most commonly used biochemical marker of an active osteogenesis. The ALP action on extracellular matrix mineralization is mainly degrading the ubiquitous mineralization pyrophosphate (PPi) which releases inorganic phosphate (Pi) thus providing the necessary Pi for mineral nucleation process (23) and eventually inducing the formation of hydroxyapatite \( \text{Ca}_{10} \left( \text{PO}_4 \right)_6 \text{(OH)}_2 \) crystals within the collagen fibrils in the extracellular matrix. The expression of collagen and ALP activity are usually a coupled process, and an increase collagen accumulation may also indicate an increase in ALP activity since collagen scaffold is synthesized for mineralization and ALP can produce phosphate nucleator for Ca deposition into the collagen network. The crucial role of type I collagen and ALP on mineralization came from the works of Murshed et al. (24) who used several knock-out mouse models to
demonstrate that the co-expression of collagen and ALP are necessary and sufficient to induce mineralization of any extracellular matrix. Therefore, the ability of yam extracts to enhance collagen accumulation in the extracellular matrix may indicate their positive role in enhancing bone formation.

In summary, our results suggest that yam ethanol and water extracts may have a potential positive effect in enhancing bone formation by 1) increasing osteoblast cell proliferation and 2) by increasing extracellular matrix collagen accumulation crucial for matrix mineralization. This study results may aid in the development of nutraceutical and functional food with the approach of utilizing yam extracts for the enhancement of bone health and prevention of bone-related disorders such as osteoporosis. Further studies are also needed to elucidate the potential mechanism whereby yam extracts affects bone formation so that a more effective treatment scheme may be developed.

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