Identification of Reduced Fibronectin 1 Gene Expression in Osteoblasts following Hydrogen Peroxide Treatment using Subtractive Gene Cloning

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Abstract: Bone formation steadily declines with age resulting in a significant loss of bone mass, while reactive oxygen species (ROS) are thought to be major contributors to the aging process. In the present study, we attempted to identify which gene had an altered transcription level when osteoblasts were treated with ROS, H2O2, using a subtraction gene cloning technique. A gene clone was obtained from cDNA of cells from the osteoblastic cell line MC3T3-E1 after subtraction of the cDNA of H2O2-treated cells and designated as MC-ROS-7. To confirm the gene product, the DNA sequence of MC-ROS-7 was determined and homology was assessed using a nucleotide-nucleotide BLAST (BLASTN) search of the NCBI database, which revealed a 99.7% homology with fibronectin 1 (FN) gene. A significant lower level of FN mRNA was found in H2O2-treated cells when compared to that in non-treated cells using endpoint RT-PCR and real-time PCR assays. Our results suggest that a reduction of FN gene expression by ROS is involved in the decline of bone formation in osteoblasts during the ageing process.

Key words: Ageing, Fibronectin, Gene Expression, Osteoblasts

Introduction

Bone remodeling requires a delicate balance between bone formation and bone resorption, in which bone-forming osteoblasts and bone-resorbing osteoclasts play central roles1). It is well known that bone formation steadily declines with age, resulting in a significant loss of bone mass, and may be due to a decrease in osteoblast proliferating precursors or the number of pre-osteoblasts that differentiate, or possibly decreased synthesis and secretion of essential bone matrix proteins2). However, the nature of the mechanisms that underlie the aging process and their effects on bone metabolism are not well understood. A currently popular hypothesis states that reactive oxygen species (ROS), produced during normal cellular metabolism, are major contributors to the aging process1). ROS include superoxides anions, hydroxyl radicals, and H2O2, which can cause severe damage to DNA, protein, and lipids. High levels of oxidants produced during normal cellular metabolism or from environmental stimuli disturb the normal redox balance and shift cells into a state of oxidative stress1). Recent evidence has shown that ROS may be involved in the pathogenesis of bone loss-related diseases, a marked decrease in plasma antioxidants was found in aged osteoporotic women3). Osteoporosis was noted in 2 different mouse models of premature aging associated with oxidative damage, in which osteopenia was speculated to be the consequence of oxidative damage4, 5). However, evidence specifically linking ROS-induced damage to bone formation as a part of the aging process is quite sparse and the function of aged osteoblasts in molecular cell biology has not been well characterized.

We previously studied the effects of ROS on bone nodule forming activity using cells from the osteoblastic cell line MC3T3-E1 and found that bone nodule formation was significantly reduced by H2O2 -treatment6). However, the regulatory mechanisms leading to that observation could not be elucidated, as it is necessary to use a molecular biological approach to comprehensively study the effects of ROS on bone forming activity by osteoblasts. Recently, subtractive gene cloning, a polymerase chain reaction (PCR)-based DNA subtraction method, has been shown to have a number of advantages for cloning
differentially expressed genes. This method has been utilized to identify novel genes and may be useful in studies of the mechanisms involved with the decline of bone formation with aging.

In the present study, we used subtractive gene cloning to analysis genes in MC3T3-E1 cells whose expression was altered by H₂O₂. Herein, we describe our subtractive gene cloning strategy and focus on identification of the fibronectin gene, and also show results confirming the effects of H₂O₂ on mRNA expression using a reverse transcription-polymerase chain reaction (RT-PCR) methods.

**Materials and Methods**

**Cell culture and H₂O₂ treatment**

MC3T3-E1 cells were purchased from Riken Cell Bank (Ibaraki, Japan). They were seeded at 1 × 10⁵ cells/well in a 24-well culture plate (Corning, Grand Island, NY, USA) and incubated for 24 hours in α-MEM medium containing 5 mM of HEPES supplemented with 50 µg/ml of gentamycin sulfate (Sigma, St. Louis, MO, USA), to which was added 0.3 µg/ml of amphotericin B (Flow Laboratories, VA, USA), 100 units/ml of penicillin G potassium (Sigma, St. Louis, MO, USA), and 10% fetal bovine serum in a humidified atmosphere containing 95% air and 5% CO₂ at 37°C. For RNA isolation, 100-mm diameter dishes were used. The cells were treated with 400 µM of H₂O₂ for 3 hours, after which they were washed and incubated with the differentiation medium, α-MEM medium described above supplemented with 50 µg/ml ascorbic acid and 10 mM β-glycerol phosphate.

**Quantitation of bone-nodule formation**

For quantitation of bone nodule formation, MC3T3-E1 cells were cultured for 40 and 50 days in the medium, after which the contents of each well were fixed for 15 minutes in 10% neutral buffered formalin and stained using a von Kossa technique.

**RNA preparation and construction of subtractive cDNA library**

Total cellular RNA was isolated from MC3T3-E1 cells at 6 hours after treatment with or without H₂O₂ using an Rneasy Fibrous Tissue Mini Kit (Qiagen, GmbH, Hilden, Germany) according to the manufacturer’s instructions. The cDNA was synthesized by an Omniscript Reverse Transcription kit (Qiagen) according to the manufacture’s instructions and utilized a PCR-Select cDNA Subtraction Kit (Clontech) for construction of the subtractive cDNA library, according to the manufacturers’ protocols. The resulting cDNA was directly inserted into a pGEM-T Easy Vector (Promega, Madison, WI) and transformed into competent Escherichia coli JM109 cells.

**Dot-blot differential screening**

The cloned cDNA inserts were initially evaluated for differential expression using DNA dot-blot analysis. White colonies from a subtractive cDNA library were picked randomly and the inserts amplified by PCR using the universal vector primers M13 and M13R. The PCR products were then purified with a QIAquick PCR purification kit (Qiagen) and dot-blotted on Hybond-N+ nylon membranes (Amersham Pharmacia Biotech, Buckinghamshire, United Kingdom), after which the membranes were hybridized with horseradish peroxidase (HRP)-labeled tester cDNA as a positive probe and HRP-labeled driver cDNA as a negative probe.

**Nucleotide sequencing and homology search**

Sequencing of the cDNA inserts in the vectors was performed using a Big Dye Terminator Cycle Sequencing Kit with T7 primer with an ABI PRIZM 310 Genetic Analyzer (Applied Biosystems, USA). The DNA sequence was analyzed using DNASIS sequence analysis software (Hitachi Software Engineering, Yokohama, Japan). Database homology searches for the cDNA sequence and deduced protein sequences were performed using BLASTN programs with the DNA and protein databases at the National Center for Biotechnology Information (NCBI).

**Endpoint RT-PCR and real-time PCR**

For the endpoint RT-PCR, PCR amplification was performed on QuantiTect SYBR Green PCR Kit (Qiagen) as instructed by the manufacturer under the following conditions: initial denaturation step of 95°C for 15 mins followed by 20 cycles 94°C for 15 sec, 56°C for 30 sec and 72°C for 30 sec. DNA products by endpoint RT-PCR were electrophoresed with DNA size standards on 1.5% agarose gel, followed by staining with ethidium bromide and confirmed the expected size of DNA.

**PCR DNA primers used were as follows; 5'-GCCGACTCTGACTGGCCTTAC-3' and 5'-C C G T G T A A G G TCAAAGCAT-3' (Gene bank accession no. NM010233) for FN, 5'-ATCACCATCTTCCAGGAG-3' and 5'-A T G G A C T G A C T G G TCAAAGCAT-3'
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TGGTCTAGG-3' (Gene bank accession no. M33197) for glyceraldehydes-3-phosphate dehydrogenase (GAPDH). The FN primers were amplified as a product of 166 base pair (bp), the GAPDH primers were amplified as a product of 318 bp.

The real-time PCR was carried out using a QuantiTect SYBR Green PCR Kit (Qiagen). The PCR mixture, which contained 20 pmol of forward and reverse primers and 2 µl of cDNA, was subjected to amplification with a DNA Engine Opticon 1 (MJ Research, San Francisco, CA, USA). The cycles were set at 95°C for 15 min for preheating, followed by 36 cycles at 94°C for 15 sec, 56°C for 30 sec and 72°C for 30 sec. The amplicons were detected directly by measuring the increase in fluorescence caused by the binding of the SYBR Green I dye to gene-specific, amplified, double-stranded DNA using a DNA Engine Opticon 1. Following the completion of the PCR reaction, the temperature was raised from the annealing temperature to 95°C for a melting curve analysis.

### Table 1

<table>
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<tr>
<th>Gene ID</th>
<th>Description</th>
<th>Identity (%)</th>
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<td>Mus musculus cDNA, fibronectin 1</td>
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<td>AK133868</td>
<td>Mus musculus 8 days embryo, fibronectin 1</td>
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<td>AK147701</td>
<td>Mus musculus 14 days female placenta, fibronectin 1</td>
<td>99.7</td>
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<td>AK147683</td>
<td>Mus musculus 14 days female placenta, fibronectin 1</td>
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<td>Mus musculus dendritic cells, fibronectin 1</td>
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<td>Mus musculus fibronectin 1, mRNA</td>
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<td>Mus musculus bone marrow stroma cell, fibronectin 1</td>
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<td>L29191</td>
<td>Rat fibronectin</td>
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Figure 4. Endpoint RT-PCR analysis of FN mRNA levels. An ethidium bromide staining pattern of the amplified PCR products using agarose gel electrophoresis is shown. FN mRNA was expressed at a lower level in the \( \text{H}_2\text{O}_2 \)-treated MC3T3-E1 cells as compared to untreated cells.
Results

As shown in Figure 1, bone nodule formation was significantly reduced in MC3T3-E1 cells following treatment with H$_2$O$_2$ as compared with the untreated cells. Next, we isolated RNA from both H$_2$O$_2$-treated and untreated cells, and examined differential gene expression by the subtractive gene cloning.

A triple dot-blot analysis of subtracted DNA to screen the expressed genes is shown in Figure 2. Several of the subtracted gene clones were differentially expressed in the initial tester and driver samples. Among those, a gene clone designated as MC-ROS-7, which showed an intensity of 3 dots, was significantly reduced in H$_2$O$_2$-treated cells as compared to the control cells.

Next, nucleotide sequencing of MC-ROS-7 and a homology search were carried out. The BLASTN search resulted in a number of BLAST hits in the query sequence with a variety of homologies. Figure 3A shows the distribution of BLASTN search hits in the query sequence. Among the genes found, MC-ROS-7 DNA exhibited a 99.7% homology to the Mus musculus fibronectin 1 (FN) gene (Figure 3B). Genes with high homology sequences with MC-ROS-7 in the DNA database are summarized in Table 1.

To confirm the reduced mRNA level in the FN gene from H$_2$O$_2$-treated cells, endpoint RT-PCR analysis was performed. A significantly lower level of FN mRNA was found in H$_2$O$_2$-treated cells as compared to untreated cells, while the mRNA levels of GAPDH, used as a housekeeping control gene, were the same with or without H$_2$O$_2$-treatment (Fig. 4).

Additional experiments to determine the precise rates of enhancement of the FN genes were performed using real-time PCR. PCR exponential curves of the FN and GAPDH genes are shown in Figure 5A, with data obtained following conversion to relative mRNA amounts and the ratio of FN/GAPDH calculated for standardization shown in Figure 5B. A significant reduction of FN gene expression was observed in H$_2$O$_2$-treated cells.

Discussion

The extracellular matrix (ECM) is known to act as a substratum in a variety of tissues and plays an important role in cellular functions. FN is a heterodimeric ECM glycoprotein that has been shown to regulate the adhesion, migration, and differentiation of various mesenchymal cells$^9$. Interestingly, FN deposition on the cell surface has been reported to be decreased in human aged fibroblasts$^{10}$, while FN isolated from late passage fibroblasts has been described as unable to support normal cell substrate interactions$^{11}$.

Previously, we investigated hydroxy radicals generated by H$_2$O$_2$-Cu ion on degraded and/or modified FN proteins. Bone nodule formation by rat osteoblasts in FN-coated culture dishes was significantly greater than in the non-coated dishes, while that in FN-coated dishes following hydroxy radicals treatment was significantly less than without such treatment$^{12}$. Further, alkaline phosphatase activity in and the secretion of type I collagen from those osteoblasts were reduced by FN-treated hydroxy radicals. RT-PCR analysis also showed that mRNA levels of alkaline phosphatase and type I collagen genes were diminished by treatment with hydroxy radicals$^{13}$. Thus, tissue aging may be directly related to altered interactions between cells and the ECM. However, the direct effect of ROS on the gene expression of FN in osteoblasts is not well documented.

In the present study, we attempted to identify the effects of H$_2$O$_2$ on gene expression in MC3T3-E1 cells using subtractive gene cloning technology. Based on our results, we selected MC-ROS-7 cells, which showed reduced mRNA expression following H$_2$O$_2$ treatment. To confirm the gene product of MC-ROS-7, we used nucleotide sequencing of MC-ROS-7 DNA and conducted a homology search of the NCBI BLAST nucleic acid databases. We found that the DNA sequence of MC-ROS-7 exhibited a high homology to that of FN. We also analyzed mRNA levels in H$_2$O$_2$-treated cells.
-treated and untreated MC3T3-E1 cells using endpoint PCR and real-time PCR assays. As expected, a significant lower level of FN mRNA was observed in the H₂O₂-treated cells as compared to that in untreated cells. These findings suggest that transcription of the FN gene in MC3T3-E1 cells is reduced following challenge with ROS.

Based on our results, we concluded that ROS may cause a decline in bone formation through a reduction of FN gene expression and a dysfunction of the FN protein by degradation and/or modification during the ageing process.

Acknowledgement

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References