Cellular viability and genetic expression of human gingival fibroblasts to zirconia with enamel matrix derivative (Emdogain®)

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PURPOSE. The objective of this study was to investigate the biologic effects of enamel matrix derivative (EMD) with different concentrations on cell viability and the genetic expression of human gingival fibroblasts (HGF) to zirconia surfaces. MATERIALS AND METHODS. Immortalized human gingival fibroblasts (HGF) were cultured (1) without EMD, (2) with EMD 25 μg/mL, and (3) with EMD 100 μg/mL on zirconia discs. MTT assay was performed to evaluate the cell proliferation activity and SEM was carried out to examine the cellular morphology and attachment. The mRNA expression of collagen type I, osteopontin, fibronectin, and TGF-ß1 was evaluated with the real-time polymerase chain reaction (RT-PCR). RESULTS. From MTT assay, HGF showed more proliferation in EMD 25 μg/mL group than control and EMD 100 μg/mL group (P<.05). HGFs showed more flattened cellular morphology on the experimental groups than on the control group after 4h culture and more cellular attachments were observed on EMD 25 μg/mL group and EMD 100 μg/mL group after 24h culture. After 48h of culture, cellular attachment was similar in all groups. The mRNA expression of type I collagen increased in a concentration dependent manner. The genetic expression of osteopontin, fibronectin, and TGF-ß1 was increased at EMD 100 μg/mL. However, the mRNA expression of proteins associated with cellular attachment was decreased at EMD 25 μg/mL. CONCLUSION. Through this short term culture of HGF on zirconium discs, we conclude that EMD affects the proliferation, attachment, and cell morphology of HGF cells. Also, EMD stimulates production of extracellular matrix collagen, osteopontin, and TGF-ß1 in high concentration levels. CLINICAL RELEVANCE. With the use of EMD, protective barrier between attached gingiva and transmucosal zirconia abutment may be enhanced leading to final esthetic results with implants. [J Adv Prosthodont 2014;6:406-14]

KEY WORDS: Enamel matrix derivative (Emdogain®); Human gingival fibroblast; Zirconia; Cell proliferation; Cell attachment

INTRODUCTION

Zirconia has adequate mechanical properties similar to those of stainless steel, such as high flexural and tensile strength, high resistance to abrasion and corrosion, low thermal conductivity.¹² Survival rate and fracture load after chewing simulation of zirconia abutments showed to be similar to that of titanium abutments.³

The biocompatibility of zirconia is as comparable to that of titanium. Accumulation of bacteria on Y-TZP proved to be satisfactorily lower than titanium, which is an important characteristic of zirconia in maintaining prosthodontic restorations free of periodontal problems.⁴⁵ Stable
soft tissue attachment which has influence on the periodontal health is affected by the material composition of the implant abutments.32-34 To obtain the most esthetic results as in maxillary anterior implant restorations, soft tissue regeneration may be needed in cases with even zirconia abutments. In this study, we discuss positive effects of enamel matrix derivatives on soft tissue regeneration around zirconia abutments with which periodontal tissue formation can be promoted.

Enamel matrix derivative (EMD, Emdogain®, Straumann, Malmö, Sweden), is a commercially available extract of enamel matrix proteins, introduced for periodontal regeneration. EMD is predominantly composed of amelogenins and part of peptides such as enamelin, tuftelin, ameloblastin, and proteases. Enamel matrix proteins secreted by cells of Hertwig's epithelial root sheath are deposited on the developing root surface and start the formation of cellular cementum, a process known as cementogenesis. Development of periodontal ligaments and alveolar bone follows subsequently.9-12 EMD can be employed on denuded root surfaces to stimulate regenerative responses and restore periodontal attachment successfully.13 Therefore, EMD facilitates the adhesion, proliferation, mineralization, and differentiation of osteoblasts and contribute to enhanced osteogenesis and periodontal regeneration.14-19 Practically, the use of EMD may offer positive benefits in treatment of periodontal defects and guided bone regeneration.20-24 Thus, beneficial outcomes of EMD on periodontal regeneration have been proven through clinical and histological data. However, effects of EMD on human gingival fibroblasts need more evaluation. Gingival fibroblasts are major cells of the gingival connective tissue and play a major role in the soft tissue sealing in the trans-mucosal part of implants. Based on the previous reports of EMD on the treatment of gingival recession and the ability to improve clinical parameters,25-29 we can speculate that EMD may have direct benefits on the gingival fibroblasts. Studies on EMD have reported that EMD has concentration dependent stimulatory effects on human gingival fibroblasts and the amount of matrix produced.30,31

Studies on proliferation and attachment characteristics of human gingival fibroblasts on titanium surfaces have been reported.32-34 However, promoted attachment of human gingival fibroblasts may be achieved on zirconia surfaces also.35,36 With the additional use of EMD, protective barrier between attached gingiva and transmucosal abutment with zirconia surface may be enhanced leading to final esthetic results with implants.

This study was designed to characterize the proliferation, attachment, and gene expression of immortalized human gingival fibroblasts on zirconia by means of MTT assay, SEM, and real-time polymerase chain reaction (RT-PCR).

MATERIALS AND METHODS

The sixty zirconium discs (Ø 12 mm × 4 mm height) were sectioned from a cylinder block of zirconia (LAVA, 3M ESPE, St. Paul, MN, USA). Zirconia discs were washed twice in absolute alcohol and once in demineralised water ultrasonically, and sterilized in an autoclave.

The immortalized human gingival fibroblasts (HGF; Applied Biological Materials Inc., Richmond, BC, Canada) were grown at 37°C in an atmosphere of 5% CO₂ using Dulbecco's modified Eagle's medium (DMEM, Gibco BRL, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (FBS) and 1% antibiotics. The cells with typical fibroblastic morphology from the 4-8th passage were used.

The cell proliferation of cells cultured under different concentrations of EMD on cell proliferation was examined with an MTT test assay (Sigma, St. Lois, Mo., USA). In the control group, cells were cultured with DMEM, 10% FBS, and 1% antibiotics only; in the EMD 25 group, Emdogain® (25 µg/mL) was added to the control group media; in the EMD 100 group, Emdogain® (100 µg/mL) was added to the control group media.

The substance used for the MTT test was a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-tetrazolium salt, which turns into a blue formazan product due to the viable mitochondria in active cells. HGF cells were seeded at a density of 2×10⁴ cells/ml on the discs. Cells were cultured at 37°C in 5 % CO₂ for 4, 24, and 48 hours. The discs were moved to well plates after incubation, and new media were added. Diluted MTT (5 mg/ml) solution was added, and the incubation was continued at 37°C in 5% CO₂ for 4 hours. After the incubation medium removed, 400 µL of isopropanol with 0.04 N HCl was added to each well. The resulting formazan crystal products, which reflect the level of cell metabolism, were dissolved. The absorbance of the formazan product at 490 nm was measured in triplicate with a microplate reader (Bio-kineletics reader, EL312e, Winooski, VT, USA).

The cellular adhesion and morphology of the cells were analyzed using SEM. HGF cells were seeded at a density of 2×10⁴ cells/ml and grown for 4, 24 and 48 hours in an incubator of 37°C in 5% CO₂. Experiment wells were rinsed twice with a 0.1 M Phosphate Buffered Saline (PBS) buffer (pH 7.4) after loosely bound cells were aspirated and removed. The bound cells were fixed with 4% glutaraldehyde for 6 hours. The excess glutaraldehyde solution was removed from the cells by rinsing with PBS one more time. Dehydration in ethanol baths of progressively higher concentrations (50, 60, 70, 80, 90, 95, and 100%) for 10 min in each bath was performed. The cells were dried to a critical point and the discs were gold sputter coated with a thickness of 100 nm using an ion coater (Eiko IB-3, Eiko Engineering Ltd., Tokyo, Japan). The discs with the cultured cells were evaluated and the images were recorded by SEM (Tescan VEGA II LMU, Brno, Czech Republic) at 100× and 500× magnification.
To quantify the expression levels of collagen type I, osteopontin, fibronectin, and TGF-β1, real-time polymerase chain reaction (RT-PCR) was done. The cells were incubated for 24 hours. A thermal cycler and a laser with fiber optics were used with the 384-well plates (AB gene, Hamburg, Germany) for intensification of the fluorescent signals on this system. The fluorescence emission from each sample is collected by a charge-coupled device-camera and the quantitative data were analyzed using the ABI Prism 7900 Sequence Detection System (Applied Biosystems, San Francisco, CA, USA) and the Sequence Detection System software (SDS version 2.2, PE Applied Biosystems). Reaction regimes were composed of mixtures 10 pmol/µL of the primers, 2x SYBR Green PCR Master Mix (Applied Biosystems) with HotStarTaqt DNA-Polymerase included, the dNTP mix with dUTP added, the SYBRs Green I fluorescent dye, and ROX dye as a passive reference. To produce the relative standard curves of each gene, serial dilutions (1, 1/2, 1/4, 1/8, and 1/16) of cDNA were included in the real-time quantitative PCR plates. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as the housekeeping gene. The amplification program consisted of 50°C for 2 min and 95°C for 10 min followed by 40 cycles of initial denaturation at 95°C for 30 seconds, followed by annealing at 60°C for 30 seconds and extension at 72°C for 30 seconds. All primers were amplified using the same conditions. The amplifications were quantified using ABI 7900HT (Applied Biosystems). Real-time PCR cycle numbers were modified to molecular size (ng) according to the equation. Experiments were repeated independently in triplicate. The name and primer sequences of the genes used are listed (Table 1).

| Gene Name   | Forward primer sequence | Reverse primer sequence | Molecular size |
|-------------|--------------------------|-------------------------|----------------|
| Type 1 collagen | TCC GAC CTC TCT CCT CTG AA | AAA CTG CTT TGT GCT TTG GG | 125 |
| Osteopontin  | CAT CAC CTG TGC CAT ACC AG | GCC ACA GCA TCT GGG TAT TT | 87  |
| Fibronectin  | AAG GTT CCG GAA GAG GTT GT | CGA GTC ATG CGT AGG TGT GT | 72  |
| TGF-β1       | GAG CCT GAG GCC GAC TAC TA | CGG AGC TCT GAT GTG TTG AA | 131 |

RESULTS

The cell proliferation of control, EMD 25, and EMD 100 group were improved as the culture time increased. After 4 hours of cell culture, HGF proliferation of EMD 100 group decreased in comparison with the control group. After 24 hours of culture, cell proliferation of EMD 100 group increased to be comparable with the control group. After 48 hours of cell culture, cell proliferation of EMD 100 group considerably increased compared to that of the control group. Cellular viability of EMD 25 group showed to be the greatest of all groups at 4, 24, and 48 hours of cell culture. Cell proliferation was not affected by the concentration of the EMD. Cell proliferation of the control group and EMD 25 group was found to be statistically significant (Fig. 1).

After 4 hours of culture, EMD 25 group showed more cellular attachments with lamellipodia and spreading of the cellular matrix (Fig. 2). The control group and the EMD 100 µg/mL group showed stretching of the cells. Both experimental groups showed more flattening of the cells in a concentration-dependent manner compared to the control group (Fig. 3).
At 24 hours of culture, EMD 25 group showed the greatest amount of cellular attachments. However, EMD 100 group showed more flattened cells compared to the control and EMD 25 group with coherent vesicles of EMD with extracellular matrix covering the cells and the surface of the matrix (Fig. 4 and Fig. 5).

At 48 hours of culture, cells of all control and experimental groups showed higher adhesion properties. In the EMD 25 group, cellular processes showed to be longer and thinner (Fig. 6). Under higher magnification, HGF of experimental groups showed to be more flat and more cell to cell contact with each other was shown in the EMD 100 group (Fig. 7).

The amount of type I collagen expression was remarkably increased in a concentration-dependent manner and was found to be statistically significant ($P<.05$). Expression of osteopontin decreased with EMD 25 group. However, it was increased with significance in the EMD 100 group. Fibronectin and integrin both decreased with EMD 25 group and then increased with EMD 100 group, but did not show any significant differences. TGF-β1 showed to be decreased with EMD 25 group and then increased with EMD 100 group with statistical significance (Fig. 8).

**DISCUSSION**

For long-term stability of dental implants, biological seal of soft tissue and implant interface is very important. Transmucosal part of dental implant requires sufficient connective tissue attachment and inhibition of bacterial invasion and proliferation. Titanium has been used successfully as transmucosal part of dental implants. Titanium is the predominant metal used because of its biocompatibility and stability of the attachment of surrounding hard and soft
Fig. 4. SEM photomicrographs of HGF on zirconia discs at 24h (×100). (A) control group, (B) EMD 25 μg/mL group, (C) EMD 100 μg/mL group. SEM picture shows more cellular attachment observed on EMD 25 μg/mL group than control and EMD 100 μg/mL group.

Fig. 5. SEM photomicrographs of HGF on zirconia discs at 24h (×500). (A) control group, (B) EMD 25 μg/mL group, (C) EMD 100 μg/mL group. SEM picture shows more flattened cellular morphology in EMD 100 μg/mL group than control and EMD 25 μg/mL group and aggregation of extracellular matrix and EMD vehicles in EMD 100 μg/mL group.

Fig. 6. SEM photomicrographs of HGF on zirconia discs at 48h (×100). (A) control group, (B) EMD 25 μg/mL group, (C) EMD 100 μg/mL group. More cellular attachments were similar in EMD groups. Cellular processes in EMD 25 μg/mL group were thin and long.
tissues. However, titanium abutments may sometimes exhibit esthetic problems, especially in the anterior region. To improve the esthetics of dental implants, zirconia abutments with the biocompatibility and mechanical strength of titanium can be used. Previous investigations support this and thus, zirconia abutments are now used successfully in clinical use. Raffaeli suggested that zirconia is a more biocompatible material compared to feldspathic porcelain and can strongly increase the adhesion competence and the proliferation rate of fibroblasts. Teté proposed that zirconia showed collagen fiber orientation that was similar to that of machined titanium surface and demonstrated less plaque formation resulting in periodontal health and improved esthetics. Recent studies on zirconia surfaces with HGF show good attachment and cell proliferation and support the clinical significance of zirconia abutments. However, soft-tissue regeneration procedures for further esthetics and treatment of peri-implantitis may lead to need for increased attachment and growth of HGF on zirconia surfaces. Thus, enamel matrix proteins, which have been found to have positive effects on regeneration of periodontal defects, may be applied to enhance fibroblast attachment.
and proliferation.

Previous research has reported that enamel matrix derivative may enhance periodontal wound regeneration which has direct effect on the gingival fibroblasts and human periodontal ligament cells. In this study, expression of osteopontin was suppressed in the EMD 25 group but increased significantly in the EMD 100 group. These data comply with previous studies where EMD had stimulatory effect on expression of TGF-β1 of HGF.

EMD had stimulatory effect on type I collagen in a concentration dependent manner. This indicates the possibility of HGF cell differentiation to osteoblast-like phenotype. Overall, more cellular attachment occurred with lower concentration of EMD. However, as gene expression of proteins investigated in this study was promoted with higher concentration of EMD, the adequate concentration of EMD effective on attachment and proliferation of gingival fibroblasts on zirconia surface needs further investigation. Also, further study must be directed on whether high concentration of EMD would possibly transfer human gingival fibroblasts into cells with osteogenic potential and possibility of bone and tissue regeneration with gingival fibroblasts.

CONCLUSION

Through this short term culture of HGF on zirconium discs, we conclude that enamel matrix derivative contribute to proliferation and attachment of HGF cells. Also, enamel matrix derivative stimulates production of extracellular matrix collagen, osteopontin, and TGF-β1 in high dose. But more investigation is required to determine appropriate concentration of enamel matrix derivative for utmost cell proliferation and attachment and further investigation on differentiation potential of HGF to osteoblast-like cell would be necessary.

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