

In vitro Cell Proliferation Assay of Demineralized Dentin Material Membrane in Osteoblastic MC3T3-E1 Cells

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Aim: Demineralized dentin material membrane (DDMM) is a novel bioresorbable guided bone regeneration (GBR) which is derived from the demineralization process of bovine dentin. This material/process could be an alternative to resolve musculoskeletal dysfunction that harms the quality of human life.

Purpose: To evaluate the cytotoxic effect of DDMM as GBR membrane on MC3T3-E1 osteoblast cell line. 10

Methods: Cytotoxic effect was evaluated using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay. Osteoblast MC3T3-E1 cell culture was used as a parameter of cell viability after reacting with GBR materials. The absorbance values were examined at each treatment to determine the percentage of cell viability. There were four groups created in the present study: two treatment groups and two control groups. The treatment groups consisted of a DDMM group and a bovine pericardium collagen membrane (BPCM) group. The control groups comprised a group containing cell culture medium as a negative control group and another positive control group that contained cell cultures. 15

Results: The results revealed no significant difference in MC3T3-E1 cell viability between the treatment and control groups ($p < 0.05$). Moreover, as observed in the DDMM group, there was an increase in the number of osteoblast cells. 20

Conclusion: DDMM is a suitable alternative biomaterial for GBR as it is non-cytotoxic and could potentially increase the rate of repair of craniofacial defects.

Keywords: cells, biomedical and dental materials, oral surgical procedures, materials testing, wound, injuries, cytotoxicity test, pre-prosthetic 25

Introduction

The scope of oral and craniofacial surgery includes various surgical procedures, such as tooth extraction, implants, removal of pathological infections in the oral cavity, cleft lip, palatal surgery, orthognathic surgery, and head-and-neck reconstructive surgery. The etiology of craniofacial defects is trauma, malignancy, and hereditary defects. In general, craniofacial defects will disrupt the function of chewing function, aesthetic function, and musculoskeletal dysfunction, which negatively affects the quality of human life. Rehabilitation of patients with craniofacial defects is necessary to restore lost functions.¹ 30

Scientists have found several techniques to reconstruct craniofacial defects, including bone grafting, bone splitting, and guided bone regeneration (GBR). Among the reconstruction techniques mentioned before, GBR is the chosen 35

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technique by many operators in dentistry. The medical community often chooses the GBR technique because it is considered more biocompatible and has been proven in almost all studies and medical procedures to reconstruct craniofacial defects, which proves increased tissue regeneration.²

Guided Bone Regeneration has been widely circulating with various brands, including Bovine Pericardium Collagen Membrane (BPCM) and Demineralised Freeze-Dried Bovine Cortical Bone Membrane (DFDBCMB). The membranes have been tested for biocompatibility and shown promising results when applied to bone.³

Previous research using BPCM and DFDBCMB, on the other hand, revealed a number of flaws. After it observed, such as an increase in inflammatory cell invasion following the implantation of chemically cross-linked collagen and a high operational cost.^{3,4} Based on the reasons above, it is necessary to find an alternative type of biomedical dental material with similar structures and functions to significantly treat bone defects. This study attempted to discover the possibility of Bovine demineralized Dentine Material Membrane (DDMM). So, it can be used as a guided bone regeneration membrane. The organic matter in bovine dentine is similar to human dentine. Human teeth are the most mineralized tooth of all living creatures, and bovine enamel and dentine have the closest similarity to that of human.²⁷ Dentine contains extracellular Type I collagen and various growth factors, such as bone morphogenetic proteins (BMP), Osteocalcin, Osteonectin, and Phosphoprotein, recognized to play a role in the mineralization of bones.¹ Novel Bovine DDMM has several advantages because this material has growth factors like FGF, IGF-I, IGF-II, bFGF, and TGF- β .⁴ Therefore, the use of DDMM as GBR is expected to improve bone healing. The first principle of any biomaterial is that it must be compatible with the host body. Biocompatibility is the ability of a material to adapt to the host not to harm the body. To examine the biocompatibility of a material, a cytotoxicity test is required to determine the effect of a substance on cells directly by MTT assay.⁶

Materials and Methods

Materials Examined

The sample population of experimental groups were novel DDMM and BPCM (Jason Membrane[®], Botiss, Germany). This study used six replications from each group. In addition to the experimental groups, a control

group was set under the same experimental condition, consisting of MC3T3-E1 osteoblast cell culture as a positive control group and a media as a negative control group.^{6,7}

The first procedure was to prepare DDMM by collecting ten intact bovine teeth with an open apex that were then washed in tap water. The teeth were then refrigerated in 70% ethyl alcohol.

Next, they were rinsed before the removal of attached soft tissue and pulp. Bovine dentine was soaked in 3% hydrogen peroxide, which was replaced every day until the tissue became white. It was soaked in sterile distilled water for 5–6 days. Then, refluxed for 2 hours in isopropanol to eliminate any leftover of soft tissue or fat.

The dentine was cut and administered in a freeze-drying process at -80°C in a freeze dryer container under vacuum pressure (<20 Pa) and dried for 18–24 hours until the remaining water content was 5%. The demineralization process used 0.6 N HCl. The tissue was cut into a 5×5 mm square with a thickness of 300 μm . The double packing and sterilisation process were conducted using gamma irradiation.^{26,28} The present research was approved by Health Research Ethical Clearance Commission, Universitas Airlangga, Faculty of Dental, Number 780/HERCC.FODM/X/2020, and was conducted at the Research Center, Faculty of Dental Medicine, Universitas Airlangga, Surabaya, Indonesia.

Cell Culture

The MC3T3-E1 cells (ATCC, USA) were incubated in a humidified incubator with 95% humidity and 5% of CO₂, immersed in the α -Minimal Essential Medium (Gibco, Grand Island, USA), 1% penicillin/streptomycin (Sigma Aldrich, P0781), and 10% Fetal Bovine Serum (Gibco, Grand Island, USA). The whole medium was changed every three days until the cells reached an 80% confluency.¹⁵

Osteoblast cell culture MC3T3-E1 was treated with the following steps. First, the osteoblast culture was prepared in a 15 mL centrifuge tube. The culture was centrifuged for 5 minutes at 4900 RPM. The media was eliminated afterwards and the cell sediment was deposited. Subsequently, the incubation process was administered for three days at 37°C and 5% of CO₂.⁸

After a three-day incubation process, the cells were ready to be harvested. The first step was preparing the petri dish tissue culture containing the confluence cells

plus 1mL EDTA trypsin. Then, it was stored in the incubator at 39° C for 5 minutes. After the incubation was completed, approximately 2 mL of media was taken and put into a Petri dish of tissue culture. The following procedure would be inserting it into a 15 mL centrifuge tube. Then, a centrifugation process was conducted at 4500rpm for 5 minutes which resulted in the cells settling at the bottom of the tube. Osteoblasts that had been successfully cultured were used as an MTT test indicator to determine the cytotoxicity of novel DDMM.

For morphological analysis, MC3T3-E1 cells were positioned into a 96-well plate. The viable cells were monitored using a fluorescence imaging. The CellID software from the photomicroscope (PX71, Olympus) was operated to verify the morphology of the cells throughout the culture.

MTT Assay

MTT test was carried out with the following stages. A pipette of 100µL cell in the media control wells, cell control and experimental groups. Then, a pipette of 10µL cells with a total of 5000–10,000 cells was put into a cell control well and treatment group/well. Afterwards, they were moved into a 24-hour incubation process with a temperature of 37°C and 5% of CO₂. After the incubation was completed, the media were disposed of and new media were added with approximately 100 µL in the cell control and media control wells. The pipette in the treatment well contained approximately 50 µL extract. On the other hand, the pipette in the treatment contained approximately 50 µL media. After that, it was incubated for 24 hours at 37°C and 5% of CO₂. The pipette of 10 µL MTT reagents in all wells, then it was incubated for 4 hours at 37°C and 5% of CO₂ pipette 50µL DMSO in all wells. Mitochondria in osteoblast cells would reduce MTT formazan salt and produce a purple colour. The last process was the incubation of ... ? For 10 minutes at 37°C which was followed with a process of determining the absorbance value with a wavelength of 540nm using a microplate reader.^{6,8}

After the test was conducted, the optical density (OD) value would appear from the microplate reader, which was used to determine the percentage of the cell's viability to proliferate in the treatment. This was calculated using a viability cell formula.

$$\% \text{ viability cell} = \frac{\text{OD treatment} - \text{OD media}}{\text{OD cell control} - \text{OD media}} \times 100\%$$

The % viability cell was the percentage of viable cells after treatment, OD treatment was Formazan OD value

in each test sample, OD media was Formazan OD value in media control, and OD cell was Formazan OD value in cell culture as a positive control. The result for using the formula described above, OD value was used to compute the percentage of cells.

Statistical Analysis

After the percentage of viable cells was obtained, a normality test was applied with the Shapiro–Wilk method, a homogeneity test with the Levene's test method and a significance test with the independent *T*-test method to determine whether the difference in cell percentage between samples was significant ($p < 0.05$).

Results

Morphology of Cell Culture

After a three-day incubation, the morphology of MC3T3-E1 cells, was examined with a photomicroscope. The MC3T3-E1 cells exhibited a rounded morphology. The densities began forming a monolayer with spindle-shaped cells and without distinct abnormal changes (Figure 1).

MTT Assay

The results of the MTT assay reveal the absorbance value of the experimental sample group, positive control group and negative control group. The mean absorbance of the positive control group is 0.536, while the negative control group is zero. Furthermore the absorbance of the DDMM sample group is 0.464, and the BPCM group mean is 0.516. Based on the results, it is known that the DDMM group has a higher percentage than the negative control group. The BPCM group has a higher percentage than the novel DDMM group. The positive control group has the highest percentage of absorbance values (Figure 2).

Statistical Analysis

The percentage of live cells was statistically examined to prove the research premise. The Shapiro–Wilk test was employed to determine normality in this study ($p > 0.05$ = average distribution data). The Shapiro–Wilk test revealed that the data were normally distributed, with a *p*-value of 0.653 for the DDMM group and a *p*-value of 0.068 for the BPCM group.

The results of this study indicated the value of absorbance of different materials. In DDMM, the average percentage of living cells was 84.4%, whereas, in the BPCM

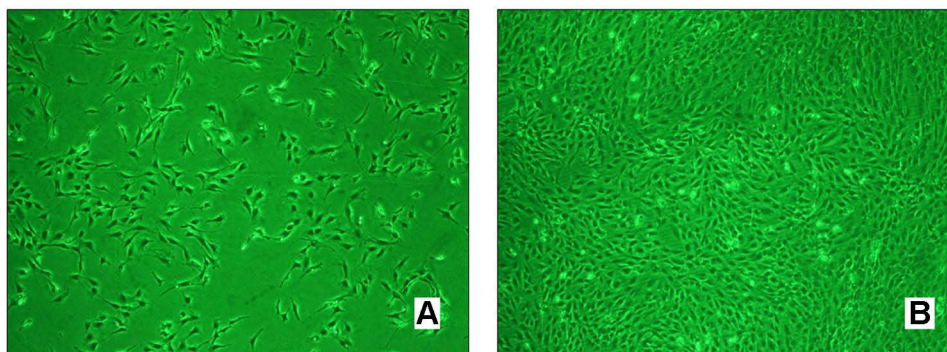


Figure 1 (A) Fluorescent images of the MC3T3-E1 cells, most cells exhibited a rounded form, achieved confluency of around 50% (B) well-spread cells displaying a polygonal morphology, achieved confluency of around 80%.

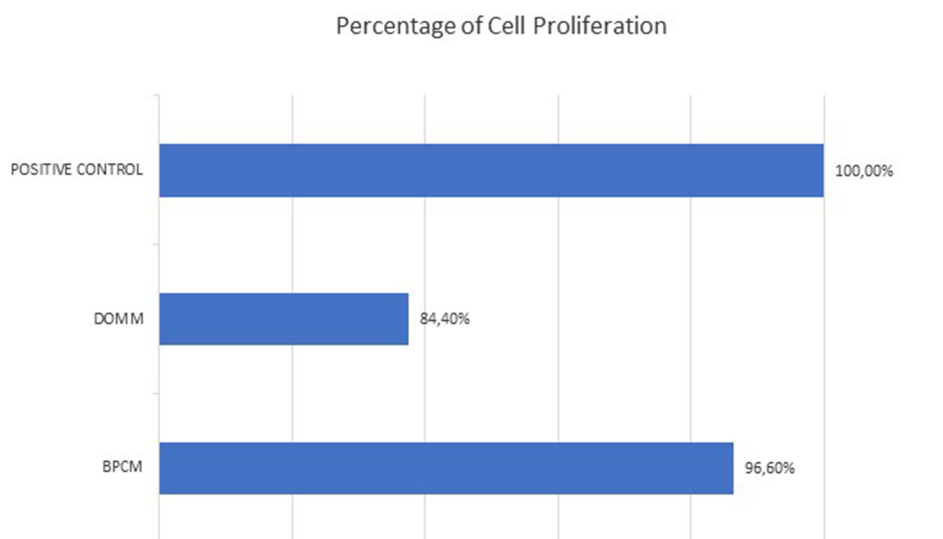


Figure 2 Table of absorbance value percentage of cell proliferation on each sample group.

membrane, it was 96.6% and the percentage of living cell was 100%.

The average percentage of live cells in both membranes, DDMM and BPCM, was greater than 50%. Therefore, both of membranes can be concluded to be non-toxic because they had a cell viability percentage of above 50% using the CD50 parameter.¹¹

In order to determine the homogeneity of the data, a homogeneity test is performed. The data were homogeneous according to Levene's test ($p = 0.091$).

The data were regularly distributed and homogeneous, according to the normality and homogeneity tests. The Independent Sample *T*-test can be used to perform a variety of tests. The Independent Sample *T*-test revealed that no statistically significant difference existed between the DDMM and BPCM groups ($p < 0.05$).

Discussion

DDMM is a bio-resorbed membrane derived from bovine dentine that has been demineralized. DDMM is a material with osteoinductive properties that are better than mineralised membranes because of the release of type 1 collagen matrix. Apart from humans, DDMM material can also be obtained from animals. Bovine dentine is easy to obtain, morphologically similar to human dentine, and possesses protein derivatives to support bone growth, namely BMP and TGF- β .^{1,5,9,10}

Performance measurement of mitochondrial dehydrogenase, also known as 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (methyl thiazolyl tetrazolium; MTT), is a rapid assessment of cell proliferation and a colorimetric cytotoxicity test that can lead into the cell metabolism or cell function measurement. The

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250 basic idea is that the tetrazole ring would be cut by
mitochondrial dehydrogenase in the cytochrome b and
c sites of live cells. Purple formazan crystals are formed
when the yellow water-soluble MTT is reduced. This sub-
255 stance dissolves in dimethyl sulfoxide and other organic
solvents. However, it does not dissolve in water. The
proportion of live cells and their activity are positively
correlated with the quantity of crystals that formed. The
percentage of viable cells and metabolic activity are
260 reflected in the measurement of colorimetric
absorbance.^{12,13} The higher the value of optical density,
the higher the cell viability will be. The optical density
value of each treatment is used in the formula for calculat-
ing the percentage of viable cells.

In vivo cell attachment is influenced not only by the
265 membrane material, but also by the implantation site and
time, as well as the patient's age and condition.²³ The
rough surface structure of the membrane encourages
attachment and proliferation of osteoblasts, whereas fibro-
blasts prefer to attach to smooth surfaces more easily. The
270 overlapping fibrils or the poor protein binding capability
cause the surface roughness.^{7,14} In addition, the incubation
period of osteoblast cells did not achieve the desired target
of the study because within 24 hours of incubation, this
cell is still alive, while more than that time, the cell will
275 lysis.^{12,16,24}

The primary material of BPCM is pericardium that is
composed of thick and dense fibrous tissue.^{14,17} Whereas
DDMM is based on dentin which has tubules with a high
level of porosity. DDMM has a more porous and rougher
280 surface structure than BPCM. Thus, the osteoblast cell
culture will be more attached to the surface of DDMM,
and when the membrane is removed from the culture,
many osteoblast cells are carried by DDMM, resulting in
a lower absorbance value. Previous studies proved that the
285 collagen content on BPCM was a factor that increased the
initial attachment of cells, so the cell viability of BPCM
was higher than novel DDMM.¹⁸⁻²⁰

Collagen membranes has been demonstrated in several
previous studies. It designed to reduce immunologic
290 responses, including macrophage control.^{25,29}

Based on this ability and the control of pro-
inflammatory and anti-inflammatory characters, the col-
lagen membrane is an innovative strategy to promote
bone regeneration.

295 GBR's capacity to produce space might provide an
impressive advantage in its ability in terms of maintaining
and creating space for the new bone to spread. The

intricate interfaces of cells in vivo cannot be recreated
using an in-vitro experimental model. Although MC3T3-
E1 osteoblast cells are a mouse osteoprogenitor cell line, 300
well considered and well raised in tissue culture. Some
possible changes might occur between these cells, which
could be generated orally from human osteoblasts. More
research is needed to clarify the clinical validity of the
305 results reproduced in this study and space preservation
ability in the clinical setting.^{21,22}

Although there are certain disadvantages to using
resorbable membranes, such as the blockage of periosteal
blood supply by the ingrowth of angiogenic cells with
310 slow healing, resorbable membranes have more advan-
tages than non-resorbable membranes. They provide for
an outstanding handling, dramatically change in the surgi-
cal methods, boost regenerative capacity, and improve
surgical results.

315 Conclusion

The present research showed that no significant difference
between DDMM and BPCM as GBR materials ($p < 0.05$)
by MTT assays on osteogenic MC3T3-E1 cells viability.
An increase in osteoblast cells was observed in the
320 DDMM group cell viability of the BPCM group. From
this study, it could be concluded that the DDMM mem-
brane is non-toxic. These results indicate that DDMM may
be a potential material to increase the number of osteo-
blasts. The study's implication will assist further related
325 research in the path of DDMM application for bone regen-
eration through in vivo research.

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Disclosure

The authors report no conflicts of interest in this work. The
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335 or indirectly, in the products used in the study.

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