Carboxymethyl Guar Gum Nanoscaffold as Matrix for Cell Growth In Vitro

Abstract

Scaffolds are an important part of tissue engineering, because they can mimic the functions of the extracellular matrix on which a cell grows. Scaffolds are used either as a matrix for the growth and proliferation of cells, or as a delivery system for growth factors to help cells grow. Polysaccharide scaffolds are practical, efficient and do not pose the problem of eliciting an immune response. Guar gum is a hydrated biopolymer obtained from the seeds of the Cyamopsis tetragonoloba, which is used in various fields. The guar gum can be modified easily to get a unique polymer which is structurally suitable for use in tissue engineering. The various structural characteristics of guar gum led us to attempt to design a scaffold using it. This scaffold was further functionalized with a carboxymethyl group, thus forming the carboxymethyl guar gum (CMGG) scaffold, with an average pore diameter of 10 µm. The aim of this paper is to assess the effects of the scaffold on the growth of RAW 264.7 macrophage cells, NIH3T3 fibroblast cells and bone marrow-derived mesenchymal stem cells. RAW 264.7 cells, NIH3T3 cells and BM-MSCs (untreated, treated with LPS to induce inflammation, and treated with CoCl$_2$, to induce hypoxia) were grown on the scaffolds in 24 well plates, and observed for at least 48 hours. MTT assay was done to assess their ability to proliferate on the scaffolds. All the cell types were found to have successfully grown and proliferated on the scaffold, even under inflammatory and hypoxic conditions (induced by treating the cells with 1 µg/ml lipopolysaccharide and 100 mM CoCl$_2$6H$_2$O, respectively). These preliminary studies have showed that the carboxymethyl guar gum scaffold may be used as a matrix in tissue engineering.

Keywords: Biomatrix; Scaffold; Biopolymers; Hydrogels; Cell migration; Inflammation; Cellular adherence and fibroblasts

Introduction

Polymeric scaffolds can be used in tissue engineering as a substitute to the extracellular matrix of the cell, as its porosity and large surface area can provide a support for the growth of the cells, and its surface properties and geometric structure direct the cells towards regeneration [1]. Typically, scaffolds are designed as meshes, fibres or sponges, which allow the cells and nutrients to be, distributed uniformly [1]. Polysaccharide hydrogels are very practicable tools for the design of scaffolds for tissue engineering, as they have macromolecular structures similar to native tissues [2-4]. In contrast to the protein matrices, most polysaccharide hydrogels are risk free and elicitors of low or no immunogenic reactions. Hydrogels can also be bioconjugated to improve their performance [4]. Biopolymers like alginates, hyaluronic acid and cellulose are amply available in nature and can easily form porous biocompatible gel networks [5]. Alginate bio-hybrids and covalently modified hyaluronates have been investigated earlier in tissue engineering arenas. Layered tissues can be formed using cell sheets. In this method, calcium alginate hydrogels can be used as a substrate for the growth of cells [6]. Once the cells reach confluence, the hydrogel scaffold can be removed by using a chelating agent on the calcium, leaving the cells to proliferate and structure themselves into the tissue [6]. Cellulose nano-formations and metal cross-linked hydrogels are other examples that are currently accepted for the design of engineered biocompatible materials [7].

Tissue engineering is a multidisciplinary field that applies the principles of engineering and life sciences to develop biological substitutes that repair, maintain, or improve the function of a tissue or whole organ [8]. Organ failure and loss of tissue, caused due to an injury or any other type of damage, is a major life threatening human health problem. The solution to these problems is tissue or organ transplantation. Other therapies, like drug therapy or the expensive procedure of replacement by medical devices are not able to restore all the functions of a damaged or lost organ or tissue. Thus engineering cells, in combination with scaffolds, are currently used as a promising alternative in the field of tissue engineering, which can be used to treat the loss of a tissue or organ without the limitations of current therapies.

Abbreviations: CMGG: Carboxymethyl Guar Gum; BM-Mscs: Bone Marrow Derived Mesenchymal Stem Cells; DMEM: Dulbecco’s Modified Eagle Medium; FBS: Foetal Bovine Serum; Pen-Strep: Penicillin-Streptomycin; LPS: Lipopolysaccharide; Cod$_6$H$_2$O: Cobalt Chloride Hexahydrate; PBS: Phosphate Buffered Saline; EDTA: Ethylene Diamine Tetra Acetic Acid; MTT: 3-(4, 5-Dimethylthiazol-2-Yl)-2, 5-Diphenyltetrazolium Bromide; UV: Ultraviolet Light; Hcl: Hydrochloric Acid

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The field of regenerative medicine seeks to heal, replace, or regenerate tissues and organs damaged by injury or disease. Stem cells have been found to be a promising source of cells to address the challenges of tissue engineering [9]. Stem cells can self-renew and differentiate into multiple lineages. This makes them an ideal source for cell therapy and regenerative medicine [10,11]. Bone marrow derived mesenchymal stem cells (BM-MSCs) can be grown easily in cell culture dishes. By varying the composition of the medium in which they are grown, MSCs can be selectively differentiated into fat cells (adipocytes), bone cells (osteocytes), and cartilage cells (chondrocytes) [12]. Thus bone marrow is a good source of cells for skeletal tissue engineering. Stem cells are also known to be distributed around the body in various other ‘niches’. However, stem cells cannot be grown easily in vitro and are thought to have a limited replicative capacity [12]. The standard in vivo culture system involves growing cells in a monolayer on a charged, flat, plastic surface, although this cannot replicate the complexity of the cells’ natural environment and rarely supports the transformation of the cells into a functional tissue [13]. This can be achieved by providing an appropriate scaffold that will lead to the development of a functional tissue. This process is certainly not a simple matter, but this approach may overcome the problem, using a variety of materials [13].

Scaffolds play an important role in tissue engineering with the goal of manipulating cell behaviour by creating a micro-environment to allow the cells to attach and subsequently develop into a tissue [14]. There are mainly two approaches to using scaffolds in regenerative medicine. First, scaffolding can be used as a structural support upon which cells are seeded in vitro. Here, the matrix provides the foundations tissue development. Secondly, using the scaffold as a delivery device for growth factors. This strategy involves combining the scaffold with growth factors, so that, upon implantation, cells from the body are recruited to the scaffold site to form a tissue throughout the matrix [14]. Scaffolds are typically made up of polymeric bio-materials such as poly lactic-co-glycolic acid (PLGA) and poly lactic acid (PLA). Scaffolds may also be of other types, like high-pressure CO₂ foamed scaffolds, injectable scaffolds, and novel custom scaffolds. These have been extensively used for tissue engineering [14].

Nanoscaffolds have been used in the field of tissue engineering and regenerative medicine for quite some time, with satisfactory results. Nanoscaffolds can provide a suitable microenvironment for the growth and proliferation of cells. Several types of scaffolds, including biodegradable ones, have been made. Nanoscaffolds have to be designed in such a way that they have small pore sizes, so the cells cannot escape into the empty spaces, and so that they have no immunological side-effects. Many scaffolds have been used in mesenchymal stem cell-based therapy where MSCs have been cultured on scaffolds and differentiated into different cell types. [15].

Tissue engineering has long been used in healing of skin wounds. Scaffolds can be used to mimic the skin cells, and provide a matrix for the tissue to grow. These scaffolds can be produced from natural materials, like collagen and gelatin, or synthetic/polymeric materials like polyglycolic acid, poly乳酸 acid, hydrogels or self-assembling peptides. These materials generally do not pose the problem of rejection, and are biocompatible [16]. Guar gum (GG) is a hydrated biopolymer obtained from the seeds of the bean Cyamopsis tetragonoloba, composed of a straight chain of β-1,4-mannose with a single galactose unit linked to alternate manose units by α-1-6 linkages [17]. The biopolymer is used regularly as functional food [17], as well as in cosmetics [17] and drug delivery [17-19]. Inter- and intra-molecular hydrogen bonding of the galactose units of GG allow for its hydration in a three chain coil formation. Despite being a non-ionic polymer, GG interacts and binds with proteins and lectins in a physiological environment due to its glycopolysaccharide structure [20,21]. This ability of GG to interact with proteins has been used to design artificial tear to treat dry eyes [22].

With these unique structural characteristics in mind, we used GG to design a scaffold to be used for tissue engineering. The GG was modified in such a way as to confer a high degree of functionality to it, to allow for better interaction with molecules. The scaffold used in this study was functionalised with carboxymethyl group, and thus, named carboxymethyl gum guar gum (CMGG) nanoscaffold. This form of the scaffold has an average pore diameter of 10μm. The present study aims to assess how cell-matrix interactions influence the in vitro behavior of different cell types (RAW 264.7 macrophage cell line, NIH3T3 fibroblast cell line and BM-MSCs) grown on CMGG nano-scaffold, and how they react under the effects of inflammation and hypoxia.

Materials and Methods

Materials

Dulbecco’s Modified Eagle Medium (DMEM), foetal bovine serum (FBS), and Penicillin- Streptomycin (Pen- Strep) were bought from Himedia, India. Bacterial lipopolysaccharide (LPS) was obtained from Sigma Aldrich; cobalt (II) chloride hexahydrate (CoCl₂·6H₂O) was purchased from Merck, India. Phosphate buffered saline (PBS; pH 7.4) was purchased from Himedia (India). 24 well plates were obtained from Nest Biotech Co. Ltd., China, and were incubated in a CO₂ incubator (Thermo Fisher) at 37°C, with 5% CO₂. Cells scrapers from Himeda, India were used to detach cells. All cell-culture work was done inside the biosafety cabinet. CMGG scaffold was a kind gift from Dr. Arup Mukherjee, Calcutta University (India).

Cell Lines: RAW 264.7 macrophage cells were cultured in our laboratory. NIH3T3 cells were obtained from NCCS, Pune (India) and cultured in our laboratory. Bone marrow mesenchymal stem cells (BM-MSCs) were isolated from adult male BALB/c mice, purchased from National Institute of Nutrition (Hyderabad, India) and cultured in our laboratory.

Methods

Ethical statement for animal experiments: All experiments using mice (for isolation of BM-MSCs) were performed according to rules laid down by the animal ethics committee of the Department of Zoology, University of Calcutta, and the animals housed under specific pathogen free conditions at the animal house of the Department of Zoology, University of Calcutta.
Culture of cells: RAW 264.7 macrophage cells were cultured in DMEM, with 10% FBS and 1% Pen-strep, and incubated at 37°C with 5% CO₂ till confluent. On reaching the confluence, the cells were scraped out, washed with 1X PBS and seeded in the wells. NIH3T3 cells were also cultured in complete DMEM (DMEM + 10% FBS + 1% Pen-Strep) and incubated at 37°C with 5% CO₂ till confluent. On reaching the confluence, the cells were trypsinized using 1X Trypsin EDTA solution (0.25% Trypsin and 0.2% EDTA in Dulbecco’s Buffered saline), washed with DMEM complete medium and seeded in respective wells. Bone marrow mesenchymal stem cells (BM-MSCs) were derived from BALB/c mice by gently flushing out the bone marrow with DMEM. The bone marrow cells were then cultured in complete DMEM (DMEM + 15% FBS + 1% Pen-Strep) with periodic media change to eliminate the hematopoietic cells. From day 3, spindle shaped MSCs appeared which was and incubated at 37°C with 5% CO₂ till 70% confluent. On reaching confluence, the cells were trypsinized, washed, and used for assays.

Culturing RAW 264.7 cells on scaffolds for 3 days: The CMGG scaffolds were UV sterilized for 15 min in the biosafety cabinet, cut into small pieces and plated in a 24 well plate which was labeled for day 1, 2 and day 3. The pieces were further washed with 1X PBS and then incubated overnight in DMEM, at 37°C in a humidified 5% CO₂ incubator prior to cell seeding. This was done to facilitate cell attachment onto the nanofibers. Next day, 5x10⁴ RAW 264.7 cells were seeded in the well labeled for day1, same number of cells in the well with scaffold as well as the control well which was without scaffolds for day 1. The seeding density was properly maintained for wells labeled for rest of the days and the culture medium was changed regularly. The cells were observed at 20X of an inverted microscope (Nikon Eclipse TiS) and photographed with Nikon DS Fi2 camera for cell quantification.

Culturing NIH3T3 cells on scaffolds for 3 days: The CMGG scaffolds were UV sterilized for 15 min in the biosafety cabinet, cut into small pieces and plated in a 24 well plate which was labeled for day 1, 2, and day 3. The pieces were further washed with 1X PBS and then incubated overnight in DMEM, at 37°C in a humidified 5% CO₂ incubator prior to cell seeding. This was done to facilitate cell attachment onto the nanofibers. Next day, the 5x10⁴ NIH3T3 cells were seeded in the well labeled for day1, same number of cells in the well with scaffold as well as the control well which was without scaffolds for day 1. The seeding density was properly maintained for wells labeled for rest of the days and the culture medium was changed regularly. The cells were observed at 20X of an inverted microscope (Nikon Eclipse Ti S) and photographed with Nikon DS Fi2 camera for cell quantification.

Culturing Bone marrow derived mesenchymal stem cells (BM-MSCs) on scaffolds for 3 days: The mesenchymal stem cells were obtained by harvesting the bone marrow from Balb/c mice. The whole bone marrow cells were cultured in DMEM medium supplemented with 15% FBS and 1% Pen-Strep. The media was changed and re-feeding of cells with fresh medium was followed for 3 days to exclude the haematopoietic cells. The MSCs shows plastic adherent property which helps them to remain adhered to the plate and proliferate. Microscopic visualization was done periodicaly for morphological examination. The cells were further maintained in the above mentioned medium and used for MTT assay as well as treated with different stress factors.

MITT Assay for RAW 264.7 cells, NIH3T3 cells and BM-MSCs grown on scaffold for 3 days: Seeding of cells were done as mentioned above and the respective wells were allowed to grow for 24 hours in CO₂ incubator. They were imaged microscopically to examine the colonization of cells with and without scaffold. The cells were treated with 40µl of 0.5 mg/ml MTT reagent (in 1X PBS) and incubated for 3 hours at 37°C, 5% CO₂. The media was discarded and the formazan dye was dissolved with acidic isopropanol (0.04M HCl in isopropanol). Then 100µl of dissolved dye was transferred in triplicates to fresh 96 well plate and the absorbance was measured at 540nm in Microplate Reader (Thermo-Fisher Scientific). The assay was done for all the 3 days.

Treatment of bone marrow derived MSCs: To assess the effects on inflammation and hypoxia on the growth of the cells in scaffold, and to determine whether the scaffold plays any protective role in stress conditions, such stress conditions were induced. The cells were divided into the following treatment groups:

- A. Only BM- MSCs
- B. BM- MSCs + CMGG scaffold
- C. BM- MSCs + CMGG scaffold + 1µg/ml LPS
- D. BM- MSCs + CMGG scaffold + 100µM CoCl₂·6H₂O
- E. Only CMGG Scaffold

For the CMGG groups, the wells were in cubated overnight with the CMGG scaffold. 0.1 x 10⁵ cells/ ml of the isolated BM- MSCs were seeded in the respective wells of the 24-well plates. The cells were treated with 1 µg/ml lipopolysaccharide for 18 hrs to induce inflammation [23,24]. The cells were treated with 100 µM cobalt chloride for 24 hrs to induce hypoxia [25-27]. The plates were then placed in a CO₂ incubator at 37°C with 5% CO₂, and the cells were observed for over a period of 24 hours and 48 hours. The cells were observed at 20X of an inverted microscope (Nikon Eclipse Ti-S) and photographed with Nikon DS-Fi2 camera for cell quantification.

Statistical analysis: All data have been presented as Mean ± Standard error of mean (SEM). Statistical significance has been calculated using t-test, considering all values of p<0.05 to be significant.

Results

Proliferation and viability of RAW 264.7 cells cultured on the CMGG scaffold

RAW 264.7 cells are the murine macrophage cells which are easy to propagate for inflammatory assay. We have cultured these cells both in absence and presence of scaffolds. The aim is to get an idea if the CMGG scaffolds are suitable for the growth of macrophages. The study was done for 72 hrs where it was found that the scaffold is providing matrix for adherence and growth of cells on them. The morphology of the cells was monitored microscopically where the cells showed proliferation on the scaffold as compared to the control cells without scaffold (Figure...
1). MTT assay was done to quantify the cellular viability of RAW 264.7 cells with and without scaffold as well as to check if the CMGG scaffolds shows any cytotoxic effect on RAW 264.7 cells. There was a 4.18 fold increase (p<0.05) in the cell proliferation of RAW cells cultured on scaffold for 24 hrs as compared to the cells without scaffold. After 48 hrs and 72 hrs, there was a 1.58 fold increase in the cell proliferation and the fold change was significant for 72 hrs (Figure 2 & Table 1). Thus it can be said that the CMGG scaffold is not affecting the viability of the cells and promoting growth and proliferation of RAW 264.7 cells in vitro.

Table 1: Fold change in the MTT assay of RAW 264.7 cells culture on scaffold with respect to cells cultured without scaffold. RAW cells cultured on scaffold shows significant increase in proliferation after 24 hrs and 72 hrs (*: p<0.05 with respect to cells without scaffold).

|                  | Absorbance at 540nm | Fold Change with Respect to Cells without Scaffold |
|------------------|---------------------|-----------------------------------------------|
|                  | Without Scaffold    | With Scaffold                                 |
| Day 1            | 0.019 ± 0.005       | 0.078 ± 0.009 (+) 4.18*                       |
| Day 2            | 0.054 ± 0.001       | 0.085 ± 0.011 (+) 1.58                         |
| Day 3            | 0.071 ± 0.001       | 0.112 ± 0.008 (+) 1.58*                        |

MTT: 3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium Bromide

Proliferation and viability of NIH3T3 cells cultured on the CMGG scaffold

Fibroblast cells are mainly concerned for tissue metabolism and the structural integrity of the tissue, by secreting precursors for extra cellular matrix. In this study, we have used a NIH3T3 cell line, which is a mouse embryonic fibroblast cell to check the effect of CMGG scaffolds on the growth and proliferation of fibroblasts. On day 0, 0.5x10^5 cells were seeded on scaffold as well as without scaffold. The cells were cultured for 72 hrs, and the growth and proliferation was monitored microscopically after 24 hrs, 48 hrs and 72 hrs (Figure 3). The MTT assay was done to quantify the viability of cells at the given interval of time. There was significant increase in the proliferation of cells when cultured on CMGG scaffold as compared to without scaffold. On day 1, there was a 1.5 fold increase (p<0.05) in the viability of NIH3T3 cells which was increased to 2.8 fold (p<0.05) on day 2 as compared to cells culture without scaffold. On day 3, there was a 2.8 fold increase (p<0.05) as compared to the cells cultured without scaffold (Figure 4 & Table 2). Thus it can be said that CMGG scaffold is not affecting the viability of cells and is providing a suitable matrix for the fibroblast and may be used for tissue engineering.
Table 2: Fold change in the MTT assay of NIH3T3 cells culture on scaffold with respect to cells cultured without scaffold. NIH3T3 cells cultured on scaffold shows significant increase in proliferation after 24 hrs, 48 hrs and 72 hrs (*: p<0.05 with respect to cells without scaffold).

| Absorbance at 540nm | Fold Change with Respect to Cells without Scaffold |
|---------------------|-----------------------------------------------|
| Without Scaffold    | With Scaffold                                 |
| Day 1               | Day 2                                         |
| 0.036 ± 0.001       | 0.056 ± 0.004                                | (+) 1.5*                              |
| 0.037± 0.001        | 0.143 ± 0.005                                | (+) 3.8*                              |
| 0.044 ± 0.002       | 0.124 ± 0.004                                | (+) 2.8*                              |

MTT: 3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium Bromide

Figure 5: Growth of BM-MSCs on tissue culture treated plate (A and C) without CMGG Scaffolds, (B and D) with CMGG Scaffolds on Day 1 and 2. Microscopic images captured at 20X of Nikon Eclipse TiS inverted microscope.

Figure 6: MTT assay of BM-MSCs cultured on CMGG scaffold. The error bars represent the Mean ± SEM (n=3; * denotes p < 0.05 with respect to MSCs without scaffold).

Effect of LPS and CoCl₂ on BMMSCs grown on CMGG scaffold

The BM-MSCs are a promising area for tissue engineering studies for regenerative therapies. In this study, BM-MSCs are derived from Balb/c mice and are cultured on CMGG scaffolds to check their proliferation to validate scaffolds in vivo for the revival of degenerated tissues through tissue engineering studies in tissue degenerative disorders. The two treatment groups are LPS (1μg/ml) for 18 hrs and CoCl₂·6H₂O for 24 hrs to induce inflammation and hypoxic condition respectively. The microscopic images are captured under 20X after the incubation period of cells with treatment groups (Figure 7). This was done to understand the ability of the scaffold to protect the cellular functions under stressed condition which are mainly found during tissue degeneration or injury. There was a 1.37 fold decrease in the cell proliferation as compared to cells grown without scaffold which was relevant with the time kinetics growth of BM-MSCs after 48 hrs (Figure 5) where the proliferation was decreased after 48 hrs of culture. In presence of LPS, there was a 1.86 fold increase in the cell proliferation and 2.36 fold (p<0.05) increase under hypoxia; the fold change was compared with cells grown without scaffold (Figure 8 & Table 3).

Table 3: Fold change in the MTT assay of BM-MSCs culture on scaffold with respect to cells cultured without scaffold. BM-MSCs cultured on scaffold shows significant increase in proliferation after 24 hrs (*: p<0.05 with respect to cells without scaffold).

| Absorbance at 540nm | Fold Change with Respect to Cells without Scaffold |
|---------------------|-----------------------------------------------|
| Without Scaffold    | With Scaffold                                 |
| Day 1               | Day 2                                         |
| 0.033 ± 0.002       | 0.087 ± 0.012                                 | (+) 2.6*                              |
| 0.042 ± 0.042       | 0.038 ± 0.004                                 | (-) 1.1                               |

MTT: 3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium Bromide; BM-MSCs: Bone Marrow Derived Mesenchymal Stem Cells.
In this study we have performed in vitro experiments to understand the possible combination of CMGG nanoscaffolds to support the growth and adherence of cells on it. We have checked the cytocompatibility of scaffolds with RAW 264.7 macrophages, NIH3T3 fibroblast and BM-MSCs in vitro in this study. As the CMGG scaffolds are mannose rich, we tried to check the growth and proliferation of macrophages which expresses mannose receptor on them. The RAW 264.7 cells are the accepted model for inflammatory and cytotoxicity studies of the scaffold [28]. In the first part of our experiment, we have performed microscopic studies for RAW 264.7 macrophage cells, followed by MTT assay to quantify the proliferation of cells on scaffold. The microscopic studies showed that they have adhered and proliferated on the scaffolds and the MTT assay showed significant increase in the fold change when cultured for 72 hrs. The NIH3T3 cells are the most suitable cells for the migratory, wound healing and proliferation studies [29]. The fibroblasts have a tendency to attain a 3D morphology in a living system to communicate with their neighboring cells which is difficult when grown on a plate in 2D [30]. The CMGG scaffolds provide that 3D biomatrix to ensure the colonization in vitro. We have checked the adherence and proliferation of NIH3T3 cells on CMGG scaffolds and a significant increase in the proliferation was observed when monitored for 24 hrs to 72 hrs. Also the morphology of the cells microscopically showed that they were compatible with the CMGG scaffold which is supporting their adherence and proliferation. The bone marrow MSCs are used for differentiation towards different lineages when supplemented with the established lineage specific factors, thus BM-MSCs are considered as the potent cells for tissue engineering studies [31]. We have performed an experiment with BM-MSCs, to check if CMGG scaffolds are supporting the growth of BM-MSCs. The BM-MSCs adhered on the scaffold and their proliferation is also increased significantly after 24 hrs as compared to cells grown without scaffold. Studies showed that during inflammatory state, there is metabolic imbalance at the mucosal surface of the tissues causing hypoxia characterized by over expression of hypoxia inducible factor (HIF) [32,33]. The hypoxia preconditioned BM-MSCs have been used for tissue injury in previous studies considering the MSCs which survives the hypoxic stressed condition are adapted to counteract and survive under the conditions prevailing during the injury for its recovery [34]. In our experiment we have checked the effect of hypoxia on cells as well as on scaffolds which was done after treating the cells grown on scaffold with CoCl$_2$. We have also checked the effect of LPS which is a widely accepted inducing agent for inflammation [35]. There was 1.86 fold increase in proliferation of BM-MSCs when treated with LPS and 2.36 fold increase in the fold change when treated with CoCl$_2$ as compared to the untreated cells. Thus in vitro studies showed that the pore size of scaffold is suitable for cell adherence for RAW 264.7, NIH3T3 cells as well as BM-MSCs. Our study also showed that the scaffolds are least affected by the inflammatory as well as the hypoxic stimulus and serve as a suitable biomatrix for the maintenance of cellular morphology. Thus, from our results we can say that CMGG scaffold can sustain the stress conditions prevailing during the tissue injury and degenerative processes and may be used as a tool for tissue regeneration studies.

**Conclusion**

The emerging field of tissue engineering has gained interest to understand the possible combination of CMGG nanoscaffolds...
with cells. In this study, we have performed in vitro experiments to check the adherence of cells on the scaffold as well as their growth and proliferation on the scaffold matrix. The first stage of our experiment showed increased proliferation of two different types of murine cells- the RAW 264.7 macrophage cells and the NIH3T3 fibroblast cells, when cultured on CMGG scaffold for 72 hrs. Since scaffolds are mainly used in tissue engineering, where the main component is often stem cells, we further checked the proliferation of BM-MSCs derived from Balb/c mice on scaffold which showed increased proliferation when cultured for 24 hrs and 48 hrs. From our studies we can conclude that, up to a certain time point, the CMGG scaffold is providing a functional 3D matrix supporting the adhesion, growth and proliferation of different types of cells. Thus our in vitro study confirms the cytocompatibility of CMGG scaffolds as a biomaterial. However in vivo study with CMGG scaffold in mice model will confirm whether it can be used for development of tissue engineering and regenerative medicines.

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Conflict of Interest

The authors have declared that no conflict of interest exists.

Contribution of Authors

SM & NG performed the experiments, analysed the data and wrote the manuscript. ERB initiated the project with her idea, designed the experiments, analyzed all data and wrote the manuscript.

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