Application of “Tissueoid Cell Culture System” Using a Silicate Fiber Scaffold for Cancer Research

Shoko Murakami\textsuperscript{a,b} Ken-ichi Mukaisho\textsuperscript{a} Takuya Iwasa\textsuperscript{c} Masaaki Kawabe\textsuperscript{c}
Saori Yoshida\textsuperscript{a} Naoko Taniura\textsuperscript{a} Takahisa Nakayama\textsuperscript{a} Masaharu Noi\textsuperscript{a,b}
Gaku Yamamoto\textsuperscript{b} Hiroyuki Sugihara\textsuperscript{a}

\textsuperscript{a}Division of Molecular and Diagnostic Pathology, Department of Pathology, Shiga University of Medical Science, Otsu, Japan; \textsuperscript{b}Department of Oral and Maxillofacial Surgery, Shiga University of Medical Science, Otsu, Japan; \textsuperscript{c}Central Research Laboratory, Japan Vilene Company, Ltd, Koga, Japan

Keywords
Tissueoid · Three-dimensional culture system · Cell culture · Silica fiber · Cancer research

Abstract

\textbf{Background:} We developed a 3-dimensional (3D) culture system using a high-purity silica fiber scaffold of unwoven sheets called Cellbed\textsuperscript{TM}. \textbf{Methods:} We used adherent colon and esophagogastric junction adenocarcinoma cells, tongue squamous cell carcinoma (SqCC) cells, and nonadherent gastric cancer cells. These cells were subjected to staining with various substances and observed by electron microscopy. To evaluate the effects of extracellular matrix in carcinoma tissues, SqCC cells were cultured in Cellbed coated with collagens I, III, and IV. \textbf{Results:} Especially well-differentiated carcinoma cells cultured in this 3D system showed their own unique characteristics: luminal formation in adenocarcinoma cells and cell stratification and keratinization in SqCC cells. Scanning electron microscopy revealed the proliferation of cancer cells with cytoplasm entwined in Cellbed. Intercellular desmosomes in squamous epithelia were detected by transmission electron microscopy of vertical cross sections. SqCC cells cultured in Cellbed coated with collagen IV showed enhanced invasive and proliferative abilities. \textbf{Conclusion:} Because the morphology of cancer cells cultured in this 3D culture system is similar to that in living organisms, we called the system a “tissueoid cell culture system.” Coating with collagen IV enables the modification of cell-matrix interactions as well as recapitulation of the in vivo microenvironment.

Introduction

Conventional 2-dimensional (2D) monolayer cell culture reportedly does not recapitulate the in vivo cancer microenvironment where cancer cells grow in a 3D manner \cite{1}. 3D in vitro cell culturing is an innovative approach in cancer research to bridge the gap between conventional 2D culture and in vivo tumors \cite{2}. Thus, many tools are now available for the development of in vitro 3D cultures \cite{3}. The presence of a scaffold is important for the growth of cancer cells and the morphogenesis of a 3D structure \cite{3}. Various biomimetic materials including ex-
tracellular matrix (ECM) as well as synthetic materials are currently used as scaffolds in 3D culture systems [2, 4–7]. In this study, a 3D culture carrier known as Cellbed™ (Japan Vilene Co., Tokyo, Japan) was used as a scaffold for cancer cell culture. Cellbed is a high-purity silica glass fiber aggregate sheet which has ultrafine and continuous porosity (Fig. 1a) [8]. Its lattice-like structure is thought to resemble the loose connective tissue in living organisms (Fig. 1b) [8, 9]. It is difficult to observe cells during culture as Cellbed is opaque in the culture medium; however, cells can be visualized using a mounting medium with a refractive index of 1.46, similar to that of silica fiber, after staining with various substances [8]. Furthermore, horizontal and vertical cross-sectional specimens can be prepared from fixed Cellbed sections after culturing the cells in this system, thereby clarifying their behavior in 3D environments [8, 9]. Using this system, we cultured squamous cell carcinoma (SqCC) cells and other adherent adenocarcinoma cells as well as nonadherent gastric cancer cells. We performed morphological assessments using scanning electron microscopy (SEM) and transmission electron microscopy (TEM) as well as hematoxylin-eosin (HE) staining, immunostaining, and fluorescence immunostaining.

To reproduce the microenvironment found within living organisms in a 3D system, we focused not only on the layered framework of fibers found within loose connective tissue in vivo, but also on the presence of an abundant ECM between the fibers. The ECM not only serves as the scaffold upon which tissues are organized, but also provides critical biochemical and biomechanical cues that direct cell growth, survival, migration, and differentiation and modulate vascular development and immune function [10]. The purity and structure of Cellbed allows a matrix to be added. A previous study reported the effects of using laminin, collagen I, and fibronectin as matrices with Cellbed [11, 12]. In this study, we coated Cellbed with several collagens to clarify the roles of ECM.

### Materials and Methods

#### Cancer Cells and Cell Culture

We used various adherent cancer cells including the gastroesophageal junction adenocarcinoma (OE-19), colorectal adenocarcinoma (DLD-1), and tongue SqCC (HSC-3, HSC-4, and SCC-15) cell lines. Gastric cancer cell lines (SNU-1 and KATOIII) were also used as nonadherent lines. The characteristics of and suitable culture conditions for these cells are summarized in Table 1 [13–19]. Both 2D and 3D cultures were incubated at 37°C and 5% CO2. Initially, the cells were seeded in a 19-mm Cellbed in a 12-well plate (4 × 10^4 cells/mL) for 3D culture. Seven days later, cells were transferred to a larger container or flask with an adequate volume of medium. To prevent pooling of the medium consumed by cancer cells within the center of the Cellbed, it was necessary to shake the container even on days when the cell culture medium was not changed. This was done to prevent ischemia and subsequent necrotic death of the cells.

#### Various Stainings

HE staining of cells in the Cellbed was performed in accordance with a previously reported method [8]. Immunohistochemical staining was carried out using the Discovery XT Automated IHC Stainer with a Ventana DABMap Detection Kit (Ventana Medical System, AZ, USA). The primary antibodies were visualized by the addition of substrate and chromogen, 3,3′-diaminobenzidine (DAB), with positive cells staining brown. All antigens were retrieved by heating. Information regarding all antibodies is listed in Table 2. Fluorescence immunostaining was conducted in Cellbed for approximately 4 weeks, in accordance with a previously reported method [8]. The primary antibodies used are listed in Table 2.

#### Scanning Electron Microscopy

HSC-4 cells seeded on 19-mm-diameter Cellbed for 2 weeks were used for SEM. Samples were prepared with a Tokai Electron Microscope Analysis (Nagoya, Japan). The samples were fixed at

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Original region</th>
<th>Histologic type</th>
<th>Differentiation</th>
<th>Medium</th>
</tr>
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<tbody>
<tr>
<td>OE-19</td>
<td>gastroesophageal junction</td>
<td>adenocarcinoma</td>
<td>moderate</td>
<td>RPMI + 10% FBS + 1% ABAM</td>
</tr>
<tr>
<td>DLD-1</td>
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<td>adenocarcinoma</td>
<td>moderate to poor</td>
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<tr>
<td>HSC-3</td>
<td>tongue</td>
<td>squamous cell carcinoma</td>
<td>poor well-differentiated</td>
<td>DMEM + 10% FBS + 1% ABAM</td>
</tr>
<tr>
<td>HSC-4</td>
<td></td>
<td></td>
<td>well-differentiated</td>
<td>DMEM/Ham’s F-12 + 10% FBS + 1% ABAM + 0.4 µg/µL hydrocortisone</td>
</tr>
<tr>
<td>SCC-15</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SNU-1</td>
<td>stomach</td>
<td>adenocarcinoma</td>
<td>signet-ring cell carcinoma</td>
<td>RPMI + 10% FBS + 1% ABAM</td>
</tr>
<tr>
<td>KATOIII</td>
<td></td>
<td></td>
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</table>
4 °C for 1 h in 1% tannic acid in 0.1 M cacodylate buffer, pH 7.4, and then subjected to 4 washes, lasting 30 min each, in 0.1 M cacodylate buffer; they were then postfixed in 2% osmium tetroxide in 0.1 M cacodylate buffer at 4 °C. Next, the samples were dehydrated using alcohol. The alcohol was replaced with tert-butyl alcohol at room temperature, and the frozen samples were vacuum-dried. After drying, the samples were coated with osmium (30 nm) using an osmium plasma coater (NL-OPC80A; Nippon Laser & Electronics Laboratory, Nagoya, Japan), and viewed using a scanning electron microscope (JSM-7500F; Jeol, Tokyo, Japan).

**Transmission Electron Microscopy**

SCC-15 cells were seeded on a 19-mm-diameter collagen-IV-coated Cellbed and cultured for approximately 4 weeks. Sample preparation and imaging were performed by Tokai Electron Microscopy, in accordance with a previously reported method [8].

**Evaluation of the Effects of Collagen I, III, and IV Coating Collagen Coating**

Five hundred microliters of 10× diluted Cellmatrix I and Cellmatrix IV (Nitta Gelatin, Osaka, Japan) in dilute hydrochloric acid (pH 3.0), and 500 µL of 300× diluted collagen type III (3 mg/mL) (Nippi, Tokyo, Japan) in dilute acetic acid (5 mM) were added to each well of Cellbed in a 12-well plate. The plate was protected from light and immersed overnight at 4 °C. The solution was aspirated, after which the plate was air-dried on a clean bench for 30 min, followed by 2 washes with the medium used for the cell lines.

**Serial Dilution Experiment Using Collagen IV**

Five hundred microliters of 2x, 5x, 10x, and 100x diluted Cellmatrix IV in dilute hydrochloric acid (pH 3.0) were added to each well of Cellbed in a 12-well plate. The subsequent process for the preparation was as described above.

**Table 2. First antibodies**

<table>
<thead>
<tr>
<th>Isotype</th>
<th>Clone</th>
<th>Dilution rate</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immunohistochemistry</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>MUC1</td>
<td>mouse</td>
<td>Ma695</td>
<td>1/100</td>
</tr>
<tr>
<td>CK17</td>
<td>mouse</td>
<td>E3</td>
<td>1/40</td>
</tr>
<tr>
<td>CK (AE1/AE3)</td>
<td>mouse</td>
<td>AE1/AE3</td>
<td>diluted</td>
</tr>
<tr>
<td>Ki67</td>
<td>rabbit</td>
<td>Sp6</td>
<td>1/100</td>
</tr>
</tbody>
</table>

**Immunofluorescence**

<table>
<thead>
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<th>Isotype</th>
<th>Clone</th>
<th>Dilution rate</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ezrin</td>
<td>rabbit</td>
<td>EP1922Y</td>
<td>1/200</td>
</tr>
<tr>
<td>Cortactin</td>
<td>rabbit</td>
<td>1/1,000</td>
<td>Abcam</td>
</tr>
</tbody>
</table>

**Measurement of Depth of Tongue Cancer Cell Invasion**

After 2 weeks of 3D culture, vertical paraffin sections were stained with HE, the depth of invasion was measured using a light microscope at ×400 magnification and ×40 focus, and the average was calculated. The depths of invasion in collagen I-, III-, and IV-coated and uncoated Cellbeds were compared.

**Ki67 Index**

A light microscope at ×200 magnification and ×10 focus was used to determine the rate of Ki67 positivity. Immunostaining with Ki67 was performed using a sectioned sample of the vertical section of the collagen I-, III-, and IV-coated and uncoated Cellbeds after 2 weeks of 3D culture.

**Statistical Analysis**

Student’s t test was used for statistical analysis to evaluate the effects of collagens I, III, and IV on invasion depth and proliferative ability.
Results

Observation of Adherent Adenocarcinoma Cells

Clear ductal luminal formations were observed in horizontal and vertical cross sections of 3D cultures of OE-19 cells (Fig. 2a, b). Additionally, immunostaining was successfully performed using the first antibody of MUC1 (Fig. 2c). In experiments with colon cancer cell lines, luminal-like structures were observed in horizontal sections of 3D cultures of DLD-1 cells (Fig. 2d). In the vertical sections of 3D cultures, DLD-1 cells partly exhibited polarity and were regularly aligned on the surface of Cellbed (Fig. 2e).

Observation of Tongue SqCC Cells

Abnormal keratinization and cell stratification, which are characteristics of well-differentiated SqCC cells, were observed in horizontal and vertical cross-sections of HSC-4 and SCC15 cells grown in 3D culture (Fig. 3a–c). Staining positivity was confirmed upon immunostaining using CK17 (Fig. 3d) and fluorescence immunostaining using ezrin (green) and cortactin (red; Fig. 3e–h). HSC-4 scanning electron micrographs showed that cancer cells were present among the Cellbed fibers with cytoplasm (Fig. 4a, b). Desmosomes were observed between cells by TEM of vertical cross-sections of SCC-15 cells grown for 4 weeks in 3D culture (Fig. 4c, d).

Morphological Observation of Nonadherent Cells

SNU-1 and KATOIII cells proliferated in the 3D culture system (Fig. 5a, b). These cancer cells were partly clustered, but no luminal structure was detected. Immunostaining experiments were successfully performed using formaldehyde-fixed paraffin-embedded sections of
growing cells in Cellbed (Fig. 5c, d). Most SNU-1 cells were positive for Ki67 (nuclear staining; Fig. 5c) and all KATOIII cells were positive for CK (AE1/AE3; membranous and cytoplasmic staining; Fig. 5d).

**Collagen IV Coating Increases Invasion Depth**
We evaluated the invasion depth of cancer cells in the vertical section of Cellbed, the results of which are shown in Figure 6. We had reported previously that cell projec-
tions within HSC-3 and HSC-4 cells cultured in a Cellbed could be readily detected, in contrast to the results found using cells cultured in a 2D condition [8]. These cytological changes were also observed in this study, and we further show that HSC-3 and HSC-4 cells display similar cytological and histological characteristics regardless of the type of collagen (Fig. 6a). However, such invasion was significantly deeper for HSC-3 cells in the collagen IV-coated group than in the uncoated group (HSC-3: \( p < 0.01 \), HSC-4: \( p < 0.05 \); Fig. 6b, c).

Having found significant differences in the invasion depth on comparison of the uncoated group and the collagen-IV coated group, we evaluated the invasion depth of cancer cells in the collagen-IV coated group by means of serial dilution experiments. The invasion depth was significantly deeper for HSC-3 cells in the collagen IV-coated group, at all dilutions (×2, ×5, ×10, ×100), than for those in the uncoated group (\( p < 0.01 \); Fig. 6d). The invasion depth was significantly deeper for HSC-4 cells using collagen IV, coated at ×10 dilution, than for those in the uncoated group (\( p < 0.05 \); Fig. 6e). From these results, ×10 dilution of collagen-IV is the most suitable environment for the invasion of SqCC cells.

Fig. 4. Electron microscopy image of 3D culture of squamous cell carcinoma cells. Scanning electron micrograph of HSC-4 cells cultured for 2 weeks. a A proliferating cell entangled in Cellbed fibers. Scale bar, 20 µm. b Confirmation of cellular extension into Cellbed. Scale bar, 5 µm. c, d Transmission electron microscopy image of SCC-15 cells cultured for 4 weeks. d A magnified image of (c). Desmosomes were observed even in the vertical section (arrows) Scale bar, 1 µm.
Effects of Collagen I, III, and IV on Proliferative Ability

The Ki67 index was significantly higher in HSC-3 cells in the collagen I-, III-, and IV-coated group than those in the uncoated group (collagen I: \( p < 0.01 \), collagen III: \( p < 0.01 \), collagen IV: \( p < 0.05 \); Fig. 7a). The Ki67 index was significantly higher in HSC-4 cells in the collagen III- and IV-coated group than those in the uncoated group (collagen III: \( p < 0.05 \), collagen IV: \( p < 0.01 \); Fig. 7b). Figure 7c shows the representative figures of the immunostaining of Ki67 in each group.

Discussion

The morphology of 3D cultured cells using a Cellbed scaffold is very similar to that in cancer tissues. We termed this system “tissueoid cell culture system.” We were able to assess the significance of the in vivo microenvironment by coating Cellbed with various collagen fibers. We proved that collagen IV coating enhanced the invasive and proliferative abilities of SqCC cells.

“Tissueoid” is a recently coined term. It was initially used to refer to an artificial tissue using a decellularized part of living tissue [20]. We slightly modified it and used
Fig. 6. Effects of collagen I, III, and IV coating on invasion depth. 

a HSC-3 and HSC-4 cells display similar cytological and histological characteristics regardless of the type of collagen. Scale bar, 200 µm. 

b, c The invasion depth was significantly deeper for the collagen IV-coated HSC-3 and HSC-4 cells than for the uncoated group (* p < 0.05, ** p < 0.01). 

d The invasion depth was significantly deeper for HSC-3 cells in the collagen IV-coated group, at all dilutions (×2, ×5, ×10, ×100), than for those in the uncoated group (p < 0.01). 

e The invasion depth was significantly deeper for HSC-4 cells using collagen IV, coated at ×10 dilution, than for those in the uncoated group (p < 0.05).
the term “tissueoid cell culture system” to refer to our 3D cell culture system. The tissueoid cell culture system has several advantages over other 3D cell culture approaches, such as Matrigel coating and spheroid cultures for the following reasons [21, 22]: (1) The procedures undertaken for the culture system are very simple; (2) as the scaffold resists heat and organic solvents such as xylene or alcohol, cultured cells can be directly used for various types of staining; (3) pure cultured cells can be used for various experiments [8]; and (4) we can coat ECM on the silicate fibers of Cellbed.

One of the most important factors in 3D cultures is the scaffold, which serves as an ECM substrate for cell growth and the formation of in vivo 3D structures [3]. We were able to carry out the 3D culture of all of the cancer cells applied to the tissueoid cell culture system. Because the cells can change morphology into a spindle shape and move freely in a cavity through the pores formed by the silicate fiber aggregate, especially well-differentiated cancer cells could express the histological characteristics of the cells at each primary site. For example, luminal formation, which is a characteristic of differentiated adenocarcinoma, was observed when OE-19 and DLD-1 cells were cultured. However, Cellbed is not unique in facilitating the luminal structure formation in adenocarcinoma cells. For example, the physiological breast bilayer was reproduced by formation of luminal structures in collagen gels [23], and adenocarcinoma cells form a luminal structure in 3D Matrigel [24]. However, Cellbed is a 3D culture carrier that supports the formation of the unique characteristics of each cancer cell. On the other hand, neither signet-ring cell carcinoma SNU1 nor KATOIII formed glandular structures.
Cell stratification and abnormal keratinization, typical characteristics of well-differentiated SqCC, were observed when HSC-4 and SCC-15 cells were cultured. Furthermore, the presence of desmosomes in vertical sections of SCC-15 cell cultures was confirmed by TEM. In contrast, poorly differentiated SqCC HSC-3 cells did not form obvious stratification.

These results imply that the tissueoid cell culture system accurately recapitulates the 3D morphology of cancer cells in vivo.

Interactions and cross-talk between cancer cells and various stromal cells of tumor tissue might lead to the formation of a cancer-specific microenvironment in tumor invasion and metastasis [25]. Thus, some ECM plays an important role in cancer progression [26]. The ECM can be divided into 2 matrices: interstitial matrix (IM) and basement membrane (BM) [26]. IM makes up the main stroma and plays a major role in cell migration, cell adhesion, angiogenesis, tissue development, and repair. The BM is a well-structured membrane underlying epithelial and endothelial cells [27]. The IM is dominated by fibrillary-forming collagens such as collagen I, II, and III whereas the major components of the BM are network-forming collagens such as collagen IV [26]. Collagen II is the main collagen in cartilage, so we did not investigate it in this study. Collagens I and III are the most and second-most abundant collagens in the IM, respectively, where they play key structural roles [28]. Apart from its structural role, collagen I possesses important growth factor-binding potential, and regulates cell homeostasis by binding to a variety of proteins [28]. The remodeling of the BM leads to a complex array of pro- and antitumor signals from degradation products [29]. Previous studies have reported that the rate of collagen I and III degradation is associated with patient survival in head and neck SqCC [30, 31]. Collagen IV is usually found only in the BM, but during pathogenesis, it is associated with organ or tumor fibrosis, accumulates in the tumor stroma, and plays an important role in the adhesion, migration, differentiation, and growth of cells [32]. Moreover, collagen IV is a potent and essential regulator of angiogenesis [33–35]. Our results suggest that the ability of cancer cells to invade and proliferate was enhanced in microenvironments in which collagen IV was present.

ECM remodeling is primarily mediated by matrix metalloproteinases (MMPs), and MMP activity causes profound structural and mechanical changes in the developing ECM [36]. In this study, the invasion depths of HSC-3 and HSC-4 cells were significantly less in the collagen I-coated group than in the uncoated group. This suggests that collagen I-coated fibers may resemble the in vivo tumor microenvironment before collagen I is decomposed by MMPs, while uncoated fibers may reproduce the in vivo tumor microenvironment after collagen I is degraded by MMPs. The detailed role of collagen III-coated fibers was not revealed in this study, but our results upon coating Cellbed with various collagen fibers enabled the assessment of the invasive and proliferative abilities of SqCC cells, which were not previously characterized by in vivo study. In the near future, we would like to evaluate the effects of cancer progression using other ECM proteins known to be crucial for cancer cell biology, such as fibronectin and/or laminin [37, 38].

Taking all the obtained findings together, we consider the tissueoid cell culture system to be applicable for studies of SqCC cells as well as adherent and nonadherent adenocarcinoma cells. We showed that collagen IV promoted invasion by SqCC cells. The addition of a matrix component such as collagens facilitated the recapitulation of in vivo SqCC cells. The tissueoid cell culture system could be considered a suitable experimental tool for use before in vivo studies.

Acknowledgment

We thank Professor Wataru Yasui of Hiroshima University for naming the 3D culture method using Cellbed in this research the “tissueoid cell culture system.” Electron microscope images were provided by Tokai Electron Microscopy. We would like to express our gratitude to Shuji Kawamura (Tokai Electron Microscopy).

Conflict of Interest Statement

This research was done in collaboration with Japan Vilene Company, Ltd. The company provided Cellbed and information on Cellbed materials.

Author Contributions

S.M. performed almost all experiments and prepared the manuscript. K.M. planned the experiments and provided advice on the preparation of the manuscript. S.Y. and N.T. assisted in the experiment. T.I., M.K., T.N., M.N., G.Y., and H.S participated in discussions during the experiment and preparation of the manuscript and provided good advice.


References


