Short communication

Cell growth on 3D microstructured surfaces


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ABSTRACT

Chinese Hamster Ovary (CHO) cell cultures were grown on surfaces lithographed with periodic 3D hexagonal microcavity array morphology. The range of microcavity size (inscribed circle diameter) was from 12 μm to 560 μm. CHO cells were grown also on flat surfaces. The characterization was performed with respect to cell growth density (number of nuclei per unit area) by fluorescence optical microscopy and evaluated by correlation function analysis. We found that optimum microcavity radius was 80 μm, concerning to the maximum cell growth density, being even greater than the growth density on a flat (unstructured) substrate of the same material. This finding can be important for optimization of biotechnological processes and devices.

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1. Introduction

The chemical and structural similarities of cells lead us to search for systematics in their architecture and components [1]. It is known that cell growth is usually anchorage dependent and requires attachment to a solid surface [2–4], and cell adhesion to biomaterials with a suitable surface is fundamental to many biotechnological processes [5,6]. Numerous human and veterinary use pharmaceuticals, such as human tissue plasmid activator [7,8] and viral vaccines (hepatitis [9], rabies [10], influenza [11], polio, rotavirus, food engineering [12] and mouth disease [12,13]) are produced by anchorage dependent cells. Geometrical and mechanical properties of the cell microenvironment have great impact on cell morphogenesis and function, and can interfere with cell cytoskeleton architecture, polarity, migration, division, growth and differentiation. Cell behavior can be regulated by interaction between neighboring cells and by interaction with the extracellular matrix (ECM) [14,15]. These environmental conditions are vital for homeostasis maintenance, and their deregulation can lead to loss of integrity or even cell death [16]. Thus cell growth on microstructured surfaces is an important field of investigation.

The literature reports a number of studies of cell cultures grown on surfaces modified by nano/micro-patterning of different ECM receptors in selected areas [17–24]. These studies have explored cell growth on various kinds of patterning formed by a variety of techniques, and have looked at how the cell culture responds to these features.

However, cell growth on 3D microstructures that tend to isolate cells within 3D “microcavities” - a kind of microstructuring that differs fundamentally from the prior work - has been not reported previously. The work summarized here makes use of electron beam lithographed hexagonal microstructures as the surface morphology on which the cells are grown.

The objective of this work was investigating cell growth on chemically homogeneous microcavity patterns, monitoring the growth behavior of the cells due to morphology of the surface. The results of this work can contribute for scaffolds design [25–27] and for optimal conditions for cell growth in general sense.

2. Materials and methods

2.1. Surface microfabrication

Chinese Hamster Ovary (CHO) cells were cultured on 3D hexagonal-ly microstructured substrates. Fluorescence optical microscopy (FOM) and correlation function analysis were used to characterize the growth. Periodic microstructures were generated on SU-8 (2005 from Micho Chem, resin used for electron beam lithography – electron resist) surfaces by electron beam lithography. The morphology was composed of hexagonal cavities; see Fig. 1, where R is the radius of a circle inscribed within the hexagonal microcavity, h the cavity depth and β the wall thickness between cavities. The cavity aperture 2R was varied for different substrates, with values of 12, 30, 80, 280 and 560 μm, generating samples referred to here as HEX-12, HEX-30, HEX-80, HEX-280 and HEX-560, respectively. The cavity depth h was kept constant at h = 3 μm, and the wall thickness β varied according to the cavity cell size.
2R. The parameters used for electron beam lithography were 15 pA beam current, 0.07 nC/cm exposure dose, and 10 mm working distance.

2.2. CHO cell growth

The cells were stored in liquid nitrogen (freezing solution 10% DMSO) in aliquots of 1 x 10^6 cells/mL, and thawed and seeded in monolayers into flasks of 25 cm² (Corning), in HAM-F10 (Invitrogen) medium supplemented with 10% fetal bovine serum (Cultilab) and antibiotics (streptomycin, 1% and penicillin 1%). Cell cultures were incubated in humid 5% CO₂ atmosphere at 37 °C until reaching approximately 90% confluence, when they were subcultured. At that time, the medium was removed and 5 mL of Hanks 1X solution (0.4 g KCl, 0.06 g KH₂PO₄, 0.04 g Na₂HPO₄, 0.35 g NaHCO₃, 1 g glucose, 8 g NaCl, H₂O q.s.p. 1000 mL) were added for 2 min. The Hanks solution was removed and 2 mL of trypsin 0.25%/EDTA 1X (Invitrogen) were added for 5 min, until the cells detached. Subculture was performed splitting 1 confluent culture flask to 2 sub-confluent culture flasks (1:2 ratio) every 24 h in culture flasks of 25 cm². For the studies carried out, experimental design cells were always between the 3rd and 5th passages. Cell viability on the surfaces was evaluated beforehand by the trypan blue (Merck) exclusion test. The number of dead cells (apoptosis/necrosis) was about 0.7%, which is not significant.

2.3. Fluorescence optical microscopy

After 24 h incubation, cells were fixed and their nuclei labeled with DAPI fluorophore as follows. The attached cells were washed three times with PBS 1X 1 M at 37 °C and fixed using 4% paraformaldehyde for 15 min at room temperature (RT). Cells were permeabilized by washing three times with 0.2% Triton X100 (Sigma) in PBS for 5 min at RT. The cells were then washed again three times with PBS and stained with Prolong Gold DAPI (Life Technologies). A volume of 20–30 μL of this suspension was placed onto the substrate surface and covered with a coverslip. Samples were allowed to set in the dark, at room temperature, for 24 h before analysis. Nucleus morphology was observed using an epifluorescence microscope (Olympus BX 51 with U-RFL-T fluorescence module), with light source at 350 nm excitation wavelength and with a CCD camera (Qcolor 5) for image acquisition.

2.4. Image processing

The FOM images (Fig. 2(a)) were processed by extracting the blue channel using ImageJ software [29–31] or Fiji software [32], after which the resulting images were in grayscale, allowing correlation analysis. The color of the structures was first set to the color of the background, and binarization then done. The final image was evaluated by correlation function analysis, where the white cells were associated with a “1” value and the black background associated with “0” (Fig. 2(b)).

2.5. Statistical analysis

For each surface pattern size was performed three identical experiments, generating groups of three samples. Normality tests for

<table>
<thead>
<tr>
<th>Sample</th>
<th>Diameter, 2R (μm)</th>
<th>Wall thickness, β (μm)</th>
<th>Height, h (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HEX-12</td>
<td>12</td>
<td>1.3</td>
<td>3</td>
</tr>
<tr>
<td>HEX-30</td>
<td>30</td>
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</tr>
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</tr>
<tr>
<td>HEX-280</td>
<td>280</td>
<td>6.5</td>
<td>3</td>
</tr>
<tr>
<td>HEX-560</td>
<td>560</td>
<td>9.0</td>
<td>3</td>
</tr>
</tbody>
</table>
statistical analysis were applied for each group, using Shapiro-Wilk test \[33\], where any p-value below 0.1 was considered statistically significant. An unpaired t-test \[33\] was used for the statistical comparison method between the control and each samples group. The results are presented as histograms containing mean ± standard error. Asterisks indicate significant differences between the sample groups and the flat SU-8 substrate.

### 3. Results and discussions

The parameters of the hexagonal microstructures were confirmed by scanning electron microscopy (SEM) and by atomic force microscopy (AFM) measurements and are summarized in Table 1. Fig. 3 shows a typical SEM micrograph of the microstructured HEX-80 substrate. Previous works describe \[34-36\] more details about the lithographed surface characteristics.

Cell growth density results are shown in Fig. 4(a), indicating that the greatest growth density was on HEX-80 (t-test, p > 0.1, CI = 95), as confirmed in the box-plot of Fig. 4(b). Since the average size of CHO cells grown on flat SU-8 was 35 ± 1 μm (measured by conventional optical microscopy), we suggest that the reduced cell proliferation on surfaces containing periodic cavities of HEX-12 and HEX-30 can be problematic for cell adhesion because a cell can span more than a single microwavity. For HEX-80, HEX-280 and HEX-560 substrates, the cells primarily grew inside the cavities, suggesting that adherence was not adversely affected by the microstructures. We noted that the cell growth density on HEX-280 and HEX-560 was close to that for SU-8, as expected because the dimensions of the structures were much greater than the cell size, resembling flat substrates.

The results of the correlation function analysis using a method based on Gompper’s model \[37\] revealed that the distance between the first and second nearest-neighbor cells was lower for the HEX-80 substrate, i.e., the cells were closer to each other. For the other surface patterns, the analysis showed that cells were more separated. See Fig. 5. Table 2 summarizes the results for the correlation function analysis fitting parameters - the agglomerate radius rc, the distance between particle agglomerates dc, and the number of first nearest-neighbors Np. Table 2 includes statistic analysis using ANOVA; symbols, placed beside values, are indicating the statistical significance of similarity (\(p < 0.01\), **\(p < 0.05\), ***\(p < 0.001\) compare to the flat SU-8 control).

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the microcavity is greater than the cell size, agglomerates tend to separate, while maintaining the number of nearest-neighbors, as occurs in the case of flat SU-8 substrate.

The correlation analysis results showed clearly what was not evidenced by merely viewing the images - that the cells on flat SU-8 had a strong tendency to form agglomerates within a radius $r_c$ of 45 µm, close to the radius of the HEX-80 (2R = 80 µm) structures. Thus, the HEX-80 cavities showed the highest culture growth density performance. Supporting this superior performance of HEX-80 compared to flat SU-8 is Thery's study [14], which shows that cell cultures on patterned surfaces show similarity to physiological conditions in tissue what generates more comfortable environmental conditions. Thery argues that cells in vivo have control over the substrate properties of (1) space, which is related to the geometry of the substrate and the tissue to be formed, (2) chemical information, which is related to cell adhesion, and (3) mechanical information, which is related to the mechanical properties of the substrate. In cultures on flat substrates, cells lose control over the spatial property but still successfully culture because of the other substrate properties. By introducing a microarray with appropriate dimensions onto the surface, optimal conditions for cell culture growth are restored. These enhanced conditions result in the surfaces with microstructures having $2R = 80$ µm having the maximal cell growth density.

4. Summary and conclusions

We have grown cells on surfaces with 3D hexagonal microcavity array morphology, formed by electron beam lithography, over a range of microcavity sizes. These sizes ranged from much less than the cell dimension to much greater than the cell dimension. The cell growth was characterized by fluorescence optical microscopy and evaluated by correlation function analysis. The results indicated that for optimum microcavity dimension (HEX-80) - about equal to the flat surface cell agglomerate radius - the cell growth density was significantly greater than the growth density on a flat (unstructured) substrate of the same material. This finding can be important for optimization of biotechnological processes and devices.

Acknowledgements

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References


Table 2

<table>
<thead>
<tr>
<th>Sample</th>
<th>Agglomerate radius, $r_c$ (µm)</th>
<th>Distance between agglomerates, $d_c$ (µm)</th>
<th>Number of first nearest-neighbors, $N_p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flat SU-8</td>
<td>44.2 ± 1.5†</td>
<td>102.9 ± 1.3†</td>
<td>10 ± 2</td>
</tr>
<tr>
<td>HEX-12</td>
<td>51.6 ± 0.4</td>
<td>210.4 ± 8.1**</td>
<td>11 ± 1</td>
</tr>
<tr>
<td>HEX-30</td>
<td>50.8 ± 1.8</td>
<td>213.7 ± 1.4**</td>
<td>9 ± 1</td>
</tr>
<tr>
<td>HEX-80</td>
<td>45.9 ± 0.3†</td>
<td>105.7 ± 2.4*</td>
<td>15 ± 1</td>
</tr>
<tr>
<td>HEX-280</td>
<td>49.4 ± 0.9</td>
<td>146.4 ± 9.5**</td>
<td>10 ± 1</td>
</tr>
<tr>
<td>HEX-560</td>
<td>49.4 ± 1.7</td>
<td>150.8 ± 9.3**</td>
<td>9 ± 1</td>
</tr>
</tbody>
</table>

* Results for the correlation function analysis fitting parameters: agglomerate radius $r_c$, distance between particle agglomerates $d_c$, and number of nearest-neighbors $N_p$. ANOVA was used for the statistical analysis for each column. Symbols indicating the statistical significance of similarity are placed aside values ($p < 0.01$, *$p < 0.1$, **$p < 0.05$, †$p < 0.03$, ‡$p < 0.01$). The groups with the same symbol (or no symbol) showed no statistically significant differences between them.