Label-free, microfluidic separation and enrichment of human breast cancer cells by adhesion difference†

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Received 3rd July 2007, Accepted 9th July 2007
First published as an Advance Article on the web 1st August 2007
DOI: 10.1039/b710054j

A label-free microfluidic method for separation and enrichment of human breast cancer cells is presented using cell adhesion as a physical marker. To maximize the adhesion difference between normal epithelial and cancer cells, flat or nanostructured polymer surfaces (400 nm pillars, 400 nm perpendicular, or 400 nm parallel lines) were constructed on the bottom of polydimethylsiloxane (PDMS) microfluidic channels in a parallel fashion using a UV-assisted capillary moulding technique. The adhesion of human breast epithelial cells (MCF10A) and cancer cells (MCF7) on each channel was independently measured based on detachment assays where the adherent cells were counted with increasing flow rate after a pre-culture for a period of time (e.g., one, two, and four hours). It was found that MCF10A cells showed higher adhesion than MCF7 cells regardless of culture time and surface nanotopography at all flow rates, resulting in label-free separation and enrichment of cancer cells. For the cell types used in our study, an optimum separation was found for 2 hours pre-culture on the 400 nm perpendicular line pattern followed by flow-induced detachment at a flow rate of 200 μl min⁻¹. The fraction of MCF7 cells was increased from 0.36 ± 0.04 to 0.83 ± 0.04 under these optimized conditions.

Introduction

The separation and enrichment of cancer cells from a mixed phase of cancer and normal cells are the fundamental steps of diagnosis and further biological analysis of cancer. Current methods for detecting and separating cancer cells rely on observing morphological features or labeling with specific biomarkers.¹⁻⁵ Use of a biomarker is highly specific and accurate but requires laborious, time-consuming preparation for staining prior to separation, and a post process such as removal of labeling. Also, a limited number of effective biomarkers (e.g., aptamers⁶) have been reported for cancer probing and diagnosis. Most widely used labeling-assisted methods are fluorescence activated cell sorting (FACS) and magnetic activated cell sorting (MACS) because of their high separation speed and specificity.⁷ These cell sorting systems have been miniaturized to microfluidic devices, demonstrating the possibility of being integrated into lab-on-a-chip devices.⁸,⁹

In contrast, label-free methods typically utilize differences in physical properties such as cell size,¹⁰ density,¹¹ cell adhesion,¹²⁻¹⁵ and dielectric properties.¹⁶⁻¹⁸ One potential drawback in label-free methods is that the physical difference is not high enough for efficient separation in many cases, which limits the widespread use of the label-free methods. Nonetheless, the efficiency of separation could be enhanced under controlled micro/nanoenvironments by amplifying the physical differences.

Cell adhesion is one of the physical markers that can be used to detect and separate cells of interest. Currently, most cell separation and enrichment methods such as cell affinity chromatography (CAC) utilize immunoreactions to enhance the adhesion of one cell type than those of the other cell types.¹²⁻¹⁵ It is noted in this regard that the cell adhesion is not high enough for efficient separation in many cases, which limits the widespread use of the label-free methods. Nonetheless, the efficiency of separation could be enhanced under controlled micro/nanoenvironments by amplifying the physical differences. Cell adhesion is one of the physical markers that can be used to detect and separate cells of interest. Currently, most cell separation and enrichment methods such as cell affinity chromatography (CAC) utilize immunoreactions to enhance the adhesion of one cell type than those of the other cell types.¹²⁻¹⁵ It is noted in this regard that the cell adhesion is not high enough for efficient separation in many cases, which limits the widespread use of the label-free methods. Nonetheless, the efficiency of separation could be enhanced under controlled micro/nanoenvironments by amplifying the physical differences.

A microfluidic system is advantageous for analysis of cell adhesion in various aspects because of small sample consumption, use of multiple parameters, and generation of large shear stress.²⁶,²⁷ By integrating nanostructured surfaces with a microfluidic system, we herein devise a simple, label-free cell separation and enrichment scheme based on standard soft lithography and microfluidics protocol. The PDMS microfluidic device consists of four branch channels, each of which contains flat or polymeric nanostructures on the bottom (400 nm pillars, 400 nm perpendicular, and 400 nm parallel lines, respectively) to increase the difference in cell adhesion. Subsequently, the cells bound to the floor of a microfluidic
channel were detached and the adhered cells were counted with increasing flow rate of medium in a stepwise fashion. As a consequence, an optimum condition for efficient separation and enrichment of cancer cells was established for the cell types used in this study, which is described below.

**Methods and materials**

**Fabrication of PUA mould**

The ultraviolet (UV)-curable polyurethane acrylate (PUA) mould material consists of a functionalized precursor with an acrylate group for cross-linking, a monomeric modulator, a photoinitiator and a radiation-curable releasing agent for surface activity. Details of the synthesis and characterization of the material have been published elsewhere. To fabricate a sheet-type mould, the liquid precursor was drop-dispersed onto a silicon master and then a flexible, transparent polyethylene terephthalate (PET) film was brought into contact with the precursor surface. Subsequently, the mould was exposed to UV light (λ = 200–400 nm) for 20 s through the transparent backplane (dose = 100 mJ cm⁻²). After UV curing, the mould was peeled off from the master and additionally cured overnight to terminate the remaining active acrylate groups prior to use as a first replica. The resulting PUA mould used in the experiment was a thin sheet with a thickness of ~50 μm.

**Fabrication of microfluidic channel with PUA nanopatterned substrate**

Glass substrate was rinsed with isopropyl alcohol (IPA) in an ultrasonic bath for 30 min and washed in a flow of distilled water and dried in a stream of nitrogen. Three types of nanostructures were fabricated onto glass substrate using UV-assisted capillary moulding as shown in Fig. 1(a). A small amount of the PUA precursor (~0.1–0.5 mL) was drop-dispersed on the substrate and a first-replicated PUA mould (same material but without active acrylate groups) was directly placed onto the surface. The PUA precursor spontaneously moved into the cavity of the mould by means of capillary action and was cured by exposure to UV for ~20 s. After curing, the mould was peeled off from the substrate using a sharp tweezer. To increase the adhesion at the PUA nanostructure/glass interface, an adhesive promoter (phosphoric acrylate: propylene glycol monomethyl ether acetate = 1 : 10, volume ratio) was coated onto glass substrate.

PDMS microfluidic moulds were fabricated by curing PDMS pre-polymer (Sylgard 184 Silicon elastomer, Dow Corning) on silicon masters prepared by photolithography. The masters used for microfluidic moulds had protruding (positive) features with the impression of the microfluidic channel. The microfluidic channel consists of four branch channels (600 μm in width and 50 μm in height), which merged into two branches (1200 μm in width and 50 μm in height) and finally into one big channel (2400 μm in width and 50 μm in height) that was connected to a syringe pump (Fig. 1b). Rulers with 500 μm spacing were included in the four branch channels to easily locate the cells. To cure the PDMS pre-polymer, a mixture of 10 : 1 silicon elastomer and the curing agent was poured on the master and cured at 70 °C for 1 h. The PDMS replicas were then peeled from the silicon masters and cut prior to use. For the PDMS microfluidic moulds, holes were punched through the inlet and the outlet as a reservoir. For channel bonding, the PDMS microfluidic mould and glass substrate were treated in oxygen plasma for 60 s (60 W, PDC-32G, Harrick Scientific, Ossining, NY). After plasma treatment, the channel mould was brought in conformal contact with the substrate with careful alignment and firmly pressed to form an irreversible seal. The device was heated on a hot plate for several tens of minutes to further strengthen the sealing.

**Scanning electron microscopy (SEM)**

High-resolution SEM images of PUA nanostructures and cell morphologies were obtained using a HITACHI S-4800 microscope (Hitachi, Japan) operating at an accelerating voltage of 5 kV. To avoid charging effects, substrates were sputter-coated with Au to a thickness of 20 nm prior to measurements.

**Cell culture and measurement of cell adhesion**

Both MCF10A and MCF7 cells were obtained from the American Type Culture Collection (ATCC). The human normal breast epithelial cell line, MCF10A, was cultured in DMEM/F-12 medium containing 5% horse serum, 100 U ml⁻¹ penicillin, 100 μg ml⁻¹ streptomycin, 100 ng ml⁻¹ chola toxin, 10 ng ml⁻¹ epidermal growth factor, 0.5 mg ml⁻¹ hydrocortisone, 10 mg ml⁻¹ insulin, and 1% (w/v) L-glutamine at 37 °C under an atmosphere of 5% CO₂ (all from Sigma Chemical, St Louis, MO). The human breast cancer cell line, MCF7, was similarly cultured in DMEM containing 10% FBS, 100 U ml⁻¹ penicillin, 100 μg ml⁻¹ streptomycin, 10 mg ml⁻¹ insulin, and 1% (w/v) L-glutamine. Both cells were grown at a concentration of 1.0 × 10⁵ ml⁻¹ and sub-cultured when the cells were confluent at about 70%. For cell culture in a microfluidic chamber, cells were harvested by treating with 0.025% trypsin–EDTA, washed three times to remove the trace of trypsin and incubated for 30 min before injection into the channel to stabilize cells. Cell density was controlled at ~2.5 × 10⁶ ml⁻¹ to provide consistent experimental conditions. When the mixed cells of MCF10A and MCF7 lines were used, the medium was made by mixing two different culture media at a ratio of 1 : 1.

To measure cell adhesion, each cell line was injected inside a microfluidic channel (coated with 0.1% (w/v) gelatin) by surface tension driven capillary flow and the cells were left adhered to the floor of the channel in an incubator for different culture times of 1, 2 and 4 h, respectively. Flat or PUA nanostructures (400 nm pillar, 400 nm perpendicular, and 400 nm parallel line patterns) were constructed on the bottom of four branch channels. To apply a uniform shear stress, cells were distributed uniformly with a sufficient spacing between adhered cells. After cell culture, the microfluidic channel was connected to a syringe pump and then the Hank’s balanced salt solution (HBSS) was flowed into the channel with increasing flow rate in a stepwise manner. For each step, the medium was flowed for 1 min and the adherent cells were counted under an optical microscope (Nikon Eclipse TE2000, Tokyo, Japan).
Separation and enrichment of human breast cancer cells (MCF7)
Approximately 30% of MCF7 cells were mixed with MCF10A cells at a total density of $2.5 \times 10^6$ ml$^{-1}$ as described above. To distinguish MCF7 and MCF10A cells, the nuclei of MCF7 cells were stained with 10 mg ml$^{-1}$ Hoechst 33342 before mixing and both cell types were introduced inside a microfluidic channel having a patterned substrate with 400 nm perpendicular lines after culture for 2 h. After the microfluidic device was installed in an optical microscope equipped with fluorescence filter blocks, the non-viable cells were initially washed by the HBSS at a flow rate of 10 ml min$^{-1}$ for 1 min and the locations of MCF7 cells were recorded under the fluorescent microscope. Subsequently, the HBSS was flowed into the channel at a flow rate of 200 ml min$^{-1}$ for 3 min and adherent cells were counted under the microscope.

Results and discussion
Integration of a microfluidic device with nanopatterned substrate
Fig. 1(c)–(f) shows SEM images of flat PUA and three types of PUA nanostructures included inside a microfluidic channel: (c) PUA flat surface, (d) 400 nm pillars (e) 400 nm perpendicular lines, and (f) 400 nm parallel lines.

Adhesions of MCF10A and MCF7 cells
Difference in cell adhesion is a key parameter to enable separation and enrichment of cancer cells for the microfluidic approach presented here. According to recent studies, cell adhesions are significantly affected by surface chemical composition and nanotopography. More specifically, the cell...
adhesion could be increased or decreased depending on the material and geometry used to construct the surface structure. In this regard, the adhesion strength was evaluated for normal and cancer cells by increasing flow rate of the HBSS in a stepwise fashion. The step increase was adjusted to each cell line and flow rate, i.e., 50 μl min⁻¹ for low flow rates and increased to 100 μl min⁻¹ for high flow rates for MCF10A cells and 50 μl min⁻¹ for MCF7 cells (fixed regardless of flow rate). The duration of each step was 1 min throughout the experiment.

In Fig. 2, the fraction of adherent cells was plotted for (a)–(c) normal and (d)–(f) cancer cells as a function of flow rate for a given culture time. A number of notable findings were derived from the figure. First, the fraction of MCF10A cells was higher than that of MCF7 cells regardless of culture time and surface nanotopography at all flow rates. In particular, the fraction of MCF7 cells decreased drastically at a relatively low flow rate compared to that of MCF10A cells. Second, the fraction of MCF10A cells was much lower on glass and PUA flat substrate than that on nanopatterned substrate, suggesting that the PUA nanostructure could create a favorable environment for adhesion and spreading of MCF10A cells. Third, MCF7 cells were not sensitive to the patterned substrate as judged by a negligible difference in cell adhesion with different nanostructures. It is noted that the values in Fig. 2 were averaged over at least 3 repetitions using 80–100 cells per each experiment, thus representing statistical significance.

Shown in Fig. 3 is the plot for the fractional difference of adherent cells as a function of culture time for the same flow rate of 200 μl min⁻¹. With the exception of glass control the...
fractional difference showed a peak around 2 h of culture and decreased as the culture time increased. This fact reveals that the adhesion strength of MCF7 as well as MCF10A cells increases with increasing culture time. On glass control, it seems that the cells are not able to recognize the surface within a short period of time (1 h). For all the conditions tested, 1 h culture was not sufficient for proper separation of cancer cells, indicating that MCF10A cells would not spread over the surface up to 1 h. If the culture time is increased to 4 h with a higher flow rate of 400 μl min⁻¹, the fractional difference could be as high as 0.84 (see Fig. 2). Such a long time culture, however, is not suitable for high speed cell analysis and this high flow rate can exert more shear damage to the adhered cells, thus compromising cell viability. Based on these observations, the optimum condition for separation and enrichment of MCF7 cells from a mixed phase of MCF7 and MCF10A cells could be given at a flow rate of 200 μl min⁻¹ following 2 h culture on the perpendicular line pattern. Under these conditions, the maximum fractional difference was 0.78 ± 0.04. The reason why the perpendicular line pattern showed the best performance can be attributed to the largest shear stress generated on the cell surface than the other two patterns. This increased shear stress appears to facilitate detachment of MCF7 cells while suppressing detachment of MCF10A cells without compromising cell-substrate interactions.

**Morphology and hydrodynamic force of MCF10A and MCF7 cells**

Fig. 4 shows representative SEM images of (a)–(d) MCF10A and (e)–(h) MCF7 cells cultured for 2 h on various surfaces used in the experiment. As shown in the figure, the morphology of MCF10A cells on the 400 nm pillar pattern was flat with a spherical nucleus inside, similar to those on bare PUA and glass surfaces. Also, the morphology of MCF10A cells on the 400 nm line pattern showed contact guidance along the line direction presumably with more clusters of focal adhesions at the leading edges, corresponding to earlier results. By contrast, MCF7 cells showed star-shaped morphology (typical characteristics of cancer cell) on the 400 nm pillar and line patterns. It is noted that the elongation of MCF7 cells was not pronounced on the line pattern compared to MCF10A cells, suggesting that the adhesion of cancer cells would be relatively weak. For cell spreading, lamellipodia of MCF10A cells spread widely on the surfaces regardless of the material and geometry and partially migrated to the bottom of the substrate between nanostructures. By contrast, lamellipodia of MCF7 cells stretched radially on the pillar pattern whereas those on the line pattern showed more irregular and asymmetric shapes. Based on these observations, the adhesion strength of MCF7 cells would be much lower than that of MCF10A cells.

In order to examine whether the observed differences in the flow rate that are required to remove cells from surfaces are due to a difference in cell adhesion, not to the other factors, it would be beneficial to calculate hydrodynamic force ($F_\text{H}$) that is applied to the adherent cell surfaces. $F_\text{H}$ is related to the adhesive force ($F_\text{A}$), which might be equal to $F_\text{H}$ when 50% of the adherent cells detach from the cultured surfaces. According to a previous study, $F_\text{H}$ can be given by the summation of two components of hydrodynamic shear force and torque. Additionally, $F_\text{H}$ should be modified as a function of the ratio of cell to channel height ($\gamma$) in a microchannel geometry and needs to be implemented with shape evolution during cell spreading. For the cell shape elongated on the 400 nm line pattern as shown in Fig. 4(b), (i) with small $\gamma$ (<0.25), $F_\text{H}$ is given by the following form after algebraic manipulation:

$$F_\text{H} = \tau_{\text{avg}} A_{\text{surf}} \sqrt{1 + \left(\frac{c}{b}\right)^2} \geq 6\sqrt{c^2 + a^2} b \tau_{\text{wall}} \sqrt{1 + \left(\frac{c}{b}\right)^2} \quad (1)$$

where $\tau_{\text{avg}}$ is the average shear stress, $A_{\text{surf}}$ is the cell surface area, $\tau_{\text{wall}}$ the wall shear stress, and $a$, $b$, and $c$ are the length, width, and height of the adhered cell, respectively.
The geometrical parameters of the adhered cell were measured from the SEM images in Fig. 4 and given on the bottom along with the graphical definition. Here, $t_{\text{wall}}$ is given by

$$
\tau = \frac{3\mu Q}{2h^2w} \tag{2}
$$

where $Q$ is the volumetric flow rate, $h$ is the half the channel height, $w$ is the channel width, and $\mu$ is the fluid viscosity. In deriving eqn (1), it was assumed that $\tau_{\text{avg}} = \tau_{\text{max}}/2$ where $\tau_{\text{max}} = 3t_{\text{wall}}$. Accordingly, the hydrodynamic forces were calculated to be 9.15 nN and 7.37 nN for MCF10A and MCF7 cells, respectively. This result demonstrated that the detachment of MCF7 cells is mediated by weak adhesions to the substrate since the hydrodynamic force applied to the cancer cell surface is smaller.

To demonstrate potential applications to label-free separation and enrichment of cancer cells, a simple separation scheme is presented here. As shown in Fig. 5, the fraction of adherent MCF7 cells to the total adherent cells was 0.36 $\pm$ 0.04 prior to flow-induced detachment. After flowing the HBSS under optimized conditions (flow rate of 200 $\mu$m min$^{-1}$ for 3 min), the fraction of detached MCF7 cells to the total detached cells was increased to 0.83 $\pm$ 0.04 $\mu$m min$^{-1}$, indicating that the cancer cells can be readily separated and enriched by utilizing the difference in cell adhesion. These values were averaged over 13 independent experiments using $\sim$1000 cells in total, thus representing statistical significance. To check the damage on cells caused by this removal process, viability of the

Fig. 4 SEM images of (a)–(d) MCF10A and (e)–(h) MCF7 cells on three different PUA nanostructures, PUA flat surface, and glass control after 2 h pre-culture: (a), (e) 400 nm pillars, (b), (f) 400 nm lines, (c), (g) PUA flat surface, and (d), (h) glass control. On the bottom, the parameters of cell size are shown along with the definition. These values were averaged over 18 repetitions and error bars represent the standard errors of the mean (standard deviation/$\sqrt{n}$, where $n = 18$).

Fig. 5 Separation of MCF7 cells from a mixed phase of MCF10A and MCF7 cells

Separation and enrichment of MCF7 cells from a mixed phase of MCF10A and MCF7 cells

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detached MCF7 cells was tested. In this experiment, the MCF7 cells were cultured on the 400 nm perpendicular line pattern for 2 h and detached under the same conditions. It was found that the percentage of viable cells was >95% (see ESI†).

**Potential limitations and challenges**

The separation efficiency in our method turned out to be as high as that of other affinity-based cell separation methods (~0.8) including cell affinity chromatography (CAC).\(^1\)\(^2\)\(^3\)\(^4\)\(^5\)\(^6\)\(^7\)\(^8\)\(^9\) It seems that the efficiency could be further enhanced by better control over surface nanotopography and patterned material properties and by better understanding of cell-substrate interactions. In addition to this labeling free characteristic, our method is an intrinsically parallel process such that it takes merely 3 min to separate and enrich MCF7 cells from a mixture while recognizing the fact that a pre-culture step is required for several hours. This feature is contrasted with sequential DEP and FACS systems in that cells are separated one by one. Also, an increase in throughput can be readily implemented by multiplying channel number or by increasing channel length.

Our method also poses a number of potential limitations and challenges. First, the laminar flow inside a microfluidic channel would represent different velocities at different locations, thus preventing cell detachment at the edge of the channel would represent different velocities at different channel length.

Also, an increase in throughput can be readily implemented by multiplying channel number or by increasing channel length.

Second, it was found that the fraction of remaining MCF7 cells was increased when both cell lines were mixed and incubated together inside the microchannel. For example, the fraction of MCF7 cells after detachment assay was 0.13 ± 0.04 in the absence of MCF10A cells, which was increased to 0.23 ± 0.07 in the presence of MCF10A cells. It appears that some factors including epidermal growth factor in the medium of MCF10A cells enhance the adhesion of MCF7 cells. Third, the separation efficiency is currently 78%, which is not sufficient for accurate functioning of the device. Although the diversity in cell size and cell–surface interactions would be difficult to control, the efficiency could be further improved by a modified cell culture protocol or geometry of the microfluidic channel.

**Conclusions**

We have presented a simple microfluidic device for label-free separation and enrichment of cancer cells using difference in cell adhesion as a physical index. Our results demonstrated that the adhesion strength of MCF10A cells is higher than that of MCF7 cells regardless of pre-culture time and surface nanotopography at all flow rates. Also, the fractional difference between the adhered MCF10A and MCF7 cells varied with pre-culture time and showed a maximum around 2 hours with the exception of glass control. Of the patterns tested, the 400 nm perpendicular line pattern exhibited the best performance under the optimum conditions of 2 hours culture at a flow rate of 200 μl min\(^{-1}\) for 3 min. In a simple experimental scheme of enriching cancer cells, the fraction of MCF7 cells was increased from 0.36 ± 0.04 to 0.83 ± 0.04. The cell viability was also checked for the detached MCF7 cells, revealing that most of the cells were viable (>95%). It is hoped that this microfluidic device could serve as an alternative for separation and enrichment of cancer cells with minimal volume of body fluid containing a mixed population of normal and cancer cells. A future study would be directed towards analyzing the captured cancer cells for drug screening or other biological studies.

**Acknowledgements**

This work was supported by the Ministry of Science and Technology through Bio Tool R&D Project for Cell Research and in part by the Micro Thermal System Research Center of Seoul National University.

**References**