Sorting Inactivated Cells Using Cell-Imprinted Polymer Thin Films

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ABSTRACT

Previous work showed that cell imprinting in a poly(dimethylsiloxane) film produced artificial receptors to cells by template-assisted rearrangement of functional groups on the surface of the polymer thin film which facilitated cell capture in the polymer surface indentations by size, shape, and, most importantly, chemical recognition. We report here that inactivation of cells by treatment with formaldehyde (4%), glutaraldehyde (2%), or a combination of the two leads to markedly improved capture selectivity (a factor of 3) when cells to be analyzed are inactivated in the same manner. The enhanced capture efficiency compared to living cells results from two factors: (1) rigidification of the cell surface through cross-linking of amine groups by the aldehyde; and (2) elimination of chemicals excreted from living cells which interfere with the fidelity of the cell-imprinting process. Moreover, cell inactivation has the advantage of removing biohazard risks associated with working with virulent bacteria. These results are demonstrated using different strains of Mycobacterium tuberculosis.

KEYWORDS: cell imprinting · cell inactivation · cell sorting · diagnosis · tuberculosis

Cell imprinting is a recently developed technology that captures the structural and chemical information of cells on a polymer surface through template-assisted assembly of functional groups.1–5 A polymer is cured around template cells that are removed subsequently, leaving complementary cavities that not only spatially fit but also chemically recognize the target cells.5 The cell-imprinted materials exhibit specific chemical affinity to the original template cells, thereby functioning as artificial receptors. Such receptors are much less expensive to produce and more durable than natural receptors; consequently, these cell-imprinted polymer films could potentially be broadly used in cell-sorting assays. One promising potential application is the detection of pathogens causing infectious diseases, in particular, as a diagnostic tool for tuberculosis.

Mycobacterium tuberculosis is one of the three leading causes of morbidity and mortality in humans.6 Quick and accurate diagnoses of tuberculosis are the key to choosing proper antimicrobial treatment and preventing further spread. However, one major challenge is that the tubercle bacilli in patient sputum samples are at variable concentrations and embedded in a complex mixture of oropharyngeal contaminating microorganisms. To distinguish the suspected pathogen from the other constituents of the sample, microscopic examination combined with differential staining of the acid-fast bacilli on a glass slide is most commonly performed worldwide. However, this strategy suffers from low sensitivity and therefore misses about 50% of the cases. To achieve a conclusive diagnosis, a selective microbiological cell culture is performed, which requires sophisticated

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biosafety facilities and is limited by the slow growth rate of microorganisms.\textsuperscript{7} We are working on a potential solution based on cell-imprinted polymer thin films, which may overcome the limitations mentioned above. This strategy selectively captures the target cells in a sample onto a cell-imprinted polymer thin film. There have been attempts to use bioactive molecules like antibodies as a stationary phase for extraction to realize a similar strategy.\textsuperscript{8–10} However, the bioactive molecules are expensive to prepare, fragile to handle, and not easy to store. Alternatively, on the basis of our recent findings,\textsuperscript{7} we believe cell imprinting could be a promising strategy to achieve this goal.

When live cells are used as the template for imprinting, several major challenges arise.\textsuperscript{7} First, live cells are fragile during imprinting, and effective means are needed to ensure the conformational integrity of the template cells. Second, living cells can actively adapt to the environment by secreting chemicals, which thereby could detrimentally affect recognition by the cell-imprinted polymer film, causing the capturing selectivity to be decreased. Last, but not the least, the involvement of pathogens in the production of the imprinted material as well as the cell-capturing process brings occupational risk of infection to those working with them.

We describe here a method that can effectively overcome the challenges mentioned above. Recently, we discovered that inactivated bacteria can be selectively captured by a polymer film imprinted with the bacteria after having been treated with the same inactivation process, avoiding the use of live virulent bacteria; moreover, the inactivation strategies, especially those that utilize chemical reagents, resulted in better selectivity of capture than when living cells were used. In this study, we present the results we obtained on the effect of inactivation methods on enriching for identification of \textit{Mycobacterium smegmatis}, a surrogate of \textit{M. tuberculosis}, compared with other bacteria, including \textit{Escherichia coli} and \textit{Staphylococcus aureus}. In addition, we propose two mechanisms that help to explain why cell imprinting with inactivated bacteria can be superior to that of live bacteria. Finally, we demonstrate optimization of the inactivation method based on the mechanisms we have proposed.

\textbf{RESULTS AND DISCUSSION}

The bacterial model system used in these experiments was chosen to mimic the identification of \textit{M. tuberculosis} bacilli (\textit{MTB}), an extremely slow-growing bacterium (one division every 20 h) that causes millions of infections per year worldwide.\textsuperscript{12} A major challenge for \textit{TB} control programs is the lack of a rapid and low-cost diagnostic strategy. \textit{TB} infections are most commonly diagnosed by acid-fast staining smears of sputum samples.\textsuperscript{13} The work described here is a preliminary exploration of a potential strategy that could selectively capture \textit{MTB} cells from a patient sample to a small region of a test slide, thereby increasing the sensitivity of a smear-based \textit{TB} diagnosis.

In our preliminary experiment, poly(dimethylsiloxane) (PDMS) films imprinted with \textit{living} \textit{M. smegmatis} showed preferential binding of living \textit{M. smegmatis} as compared to \textit{E. coli} and \textit{S. epidermidis}. However, the use of virulent bacteria during the production of the cell-imprinted polymer thin films and the cell-capture process may bring risk of infection, which could be a major hurdle for implementation of this method. The biohazard risk could be fully removed if the imprinting—capture process is based on inactivated bacteria, which have lost their biological activity but still retain their shape and surface characteristics.

We screened both physical (UV) and chemical inactivation methods to determine the most appropriate method for our cell-imprinting strategy and compared these procedures to imprinting with live bacteria (Figure 1A). In our experiments, we carefully chose chemicals that are already confirmed by literature to be effective in killing \textit{MTB}. The chemicals we have tested include 75\% v/v ethanol, 5 wt \% bleach, 1 wt \% hydrogen peroxide, and 4 wt \% formaldehyde; the concentrations were chosen according to literature reports on sufficient dosages to kill \textit{MTB}.\textsuperscript{14} Also, as UV has been widely used for sterilization,\textsuperscript{15} including killing \textit{Mycobacteria} in milk,\textsuperscript{16} UV was also tested to inactivate the bacteria in our experiments. Every reagent effectively inactivated \textit{M. smegmatis}, \textit{E. coli}, and \textit{S. aureus} at the tested concentration.

After inactivation, the morphology of bacteria was inspected under the microscope. It was found that ethanol (not shown), bleach (not shown), and hydrogen peroxide (Figure 1B) dissolved or thinned the cell wall of \textit{M. smegmatis}, while formaldehyde kept the cell morphology largely intact (Figure 1C). Also, we found UV light with a 254 nm peak emission to be effective in inactivating \textit{M. smegmatis} and led to no detectable change to the cell morphology (Figure 1D).

Figure 1 shows that the morphology of the cells is maintained upon exposure to UV or formaldehyde but not hydrogen peroxide. Using inactivated cells, the capturing performance of the cell-imprinted PDMS films was tested. Fluorescently tagged bacteria in PBS were flowed over the imprinted region with the assistance of an array of microfluidic channels,\textsuperscript{3} and the captured bacteria were inspected under a fluorescence microscope. Figure 2 presents the number of counted cells in each inspected area for each different procedure. It was found that the imprinted area captured significantly more bacteria than the nonimprinted area.

Furthermore, we found that the selectivity of capturing, defined as the ratio of captured target bacteria to other bacteria, was higher when inactivated
bacteria were used in the process, compared to living bacteria. In addition, formaldehyde-treated bacteria led to better selectivity than when UV-treated bacteria were used. Table 1 summarizes the ratios of selectivity in capturing, calculated from the data shown in Figure 2.

Additional experiments were conducted to understand better the effect of inactivation on imprinting and capture performance. A common function of inactivation methods is to terminate the biological activity of cells. We observed that this function effectively eliminated the formation of a thin film on the template cells, as shown in Figure 3. The template for imprinting was prepared by sedimentation of cells to form a single layer on a substrate. When living bacteria were used in the template preparation, formation of a thin film was observed (Figure 3A,C). These thin films, which are thought to be formed by extracellular matrix material secreted by cells in response to their

Table 1. Selectivity of Capture, Calculated by Dividing the Number of Captured M. smegmatis by the Number of Captured E. coli, on the Imprints of M. smegmatis

<table>
<thead>
<tr>
<th>Inactivation method</th>
<th>without inactivation</th>
<th>4% formaldehyde</th>
<th>UV exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td>selectivity ratio</td>
<td>7.4</td>
<td>13.8</td>
<td>9.1</td>
</tr>
</tbody>
</table>
environment, may reduce the recognition effect by blocking the surface of the bacteria. When inactivated cells were used, no thin film was observed on the template (Figure 3B,D). The absence of this thin film ensures that the surfaces of templated bacteria are exposed to the imprinting polymer during the imprinting step. This was confirmed using atomic force microscopy (AFM), which showed that the imprinted surface obtained from inactivated bacteria had more distinguishable features than those from living bacteria (Figure 4). Quantitative measurement showed that the average depth of the cavities from inactivated M. smegmatis was 285 ± 6 nm, while that from living M. smegmatis was 261 ± 7 nm. However, the absence of a thin film is not the only reason for the observed improvement in selectivity when using inactivated cells. We observed that better selectivity was demonstrated by formaldehyde-inactivated cells as compared to UV-inactivated cells. To interpret the underlying mechanism, we looked into the mechanisms of their inactivation effects. Surface-maintaining inactivation methods can be divided into two categories based on their effects on the cell wall. Physical inactivation methods, such as UV and heating treatment, terminate the biological activity by damaging the DNA molecules and/or denaturing the proteins while generally leaving the surface morphology of the cells unchanged. Chemical inactivation methods, particularly, those based on chemical fixatives, involve
The selectivity is calculated by dividing the number of template cells (M. smegmatis) with the number of other cells (E. coli), captured on the imprints of templated cell on PDMS. For each test, the template cells and the sample were prepared with the same method.

Figure 5 shows that a combination of 4% formaldehyde and 2% glutaraldehyde, was further evaluated with various MTB strains (MTB H37Ra, MTB H37Rv, and MTB ΔlprG) to verify its general applicability to virulent cells. Cell culture experiments confirmed effective inactivation of all the strains tested. Table 2 presents the selectivity of cell capture, calculated by dividing the number of template cells with the number of other cells captured on the imprints of templated cells on PDMS. The results confirmed that, by inactivation with aldehydes that removes biohazard risks associated with the procedure, the virulent strains of MTB cells, from a mixture containing other cells, could be selectively captured on a cell-imprinted polymer thin film.

**CONCLUSIONS**

We found that if the cells used in creating the template for cell imprinting and the cells present in the sample are inactivated in the same way, inactivated cells could be used with an advantage in the imprinting – capturing strategy we proposed previously with living cells. When inactivated bacteria are employed in the entire process, the concern about the occupational risk of using virulent bacteria in preparation and utilization of the imprinted polymer is eliminated. In our previous work, we have found that the mechanism of cell capture with cell-imprinted polymer involves both shape and chemical recognition. The inactivation process employed in our current strategy may have played two roles: (1) to terminate the biological activity of the cells and therefore eliminating secretion of extracellular matrix, which helps expose the surface of the inactivated cells to the imprinting polymer; (2) to fix the morphology and surface structure of the inactivated cells, which helps to preserve the structural and chemical information on the cell surface. In general, physical inactivation methods only realize the first, whereas chemical fixatives can achieve both of these functions. This work suggests that when cells, especially, pathogenic cells, are used for imprinting, the inactivated form may be a better choice not only for better selectivity but also for increased biosafety.

**TABLE 2. Selectivity of Cell Capture, Calculated by Dividing the Number of Template Cells with the Number of Other Cells (E. coli) Captured on the Imprints of Templated Cell on PDMS**

<table>
<thead>
<tr>
<th>templated cell strain</th>
<th>MTB H37Ra</th>
<th>MTB H37Rv</th>
<th>MTB ΔlprG</th>
</tr>
</thead>
<tbody>
<tr>
<td>selectivity ratio</td>
<td>15.8</td>
<td>16.2</td>
<td>12.6</td>
</tr>
</tbody>
</table>

**EXPERIMENTAL SECTION**

**Materials.** M. smegmatis mc^2^155, MTB H37Ra, MTB H37 Rv, MTB ΔlprG, E. coli (ATCC25922), and S. aureus (ATCC 29213) were obtained from the Stanford University medical center clinical microbiology laboratory strain collection. Poly(dimethylsiloxane) was obtained from GE Silicone. CellTracker Orange, auramine-rodamine stain, and Kinyoun’s acid-fast stain kit were purchased from Invitrogen and Sigma Aldrich, respectively, and
used following the protocols provided by the vendors. All other chemicals were purchased in analytical or higher grade from Sigma Aldrich or VWR.

Cell Handling. Each bacterial strain was cultured on an LB agar plate at 37 °C. Before the experiment, fresh culture was harvested and rinsed using 1× PBS (pH 7.4) by centrifuging for 10 min at 1200g and 4 °C. The rinsed cells were subjected to a mild centrifugation for 3 min at 200g to remove clumps. OD600 was used to measure the cell density in a suspension. Unless specified, the cell suspension of each strain was diluted to be 10⁸ cell/mL for making templates and 10⁴ cell/mL for preparing the sample. For visualization, we stained the sample with CellTracker Orange (excitation wavelength 541 nm; fluorescence emission wavelength 565 nm) following the protocol suggested by the vendor.

Inactivation of the Cells. Cells in suspension were inactivated with physical (UV) or chemical means (75% v/v ethanol, 5 wt % bleach, 1 wt % hydrogen peroxide, 4 wt % formaldehyde, and 2 wt % glutaraldehyde in PBS). Inactivation with UV was carried out in microcentrifuge tubes under 6 W 254 nm UV (UVG11, UVP) for 1 min. Treatment with reagents was approached by suspending the cells in a buffer solution with the reagents and kept at room temperature for 10 min, followed by resuspension in PBS. Inactivation was confirmed by testing for colony formation.

Template Preparation. A 10 μL cell suspension (approximately 10⁶ cells/mL) was spread on the surfaces of precleaved microscope slides and kept in a covered Petri dish at 4 °C for 8 h. After the excess water was removed by centrifugation at 1500g for 1 min, the substrate was dried at 60 °C for 2 h and rinsed with water before use as the template.

Stamp Fabrication. Optimization of the imprinting protocol was conducted and discussed in our previous work. Briefly, we diluted a PDMS curing mixture (monomer/cross-linker = 10:1) using cyclohexane to a volume ratio of 2:1 and spin-coated this solution onto a microscope slide (30 s at 1500 rpm). After precurring the PDMS at 80 °C for 4 min, we pressed the template stamp into the prepolymer and kept it at 37 °C for 8 h. Then we peeled off the template slide and cleaned the imprinted polymer film by immersing it in a Petri dish filled with distilled water and sonicating it for 5 min. The substrates were then inspected with a scanning probe microscope (XE-70, Park Systems) under non-scan (tip radius <10 nm, with medium-low tip oscillation damping ~15%) and a scan rate of 0.2 Hz.

Cell Capture. After being cleaned with water, the imprinted surface is used to selectively capture the template cells in each of the cell suspensions, which have been treated with the same method for making the inactivated cells used in the template. A microfluidic channel was employed to assist passing the cell suspension over the imprinted surface. PDMS chips containing an array of microchannels were fabricated via standard soft lithography. Each channel was 30 μm high and 100 μm wide. The total volume of the channels was about 1 μL. The chip was reversibly bonded to the imprinted substrate by the adhesion between PDMS surfaces without heat or plasma treatment. A pipet tip was inserted into the inlet of the channel as a reservoir and was filled with cell suspension. A syringe was connected to the outlet of the channel to draw the cell suspension through the channel via negative pressure. For each test, a 50 μL cell suspension (OD600 = 0.2) was infused at a flow velocity of 0.2 mm/s. Then 50 μL of PBS was used to rinse the channel at the same flow rate. The imprinted area of the chip was inspected under a confocal microscope (TCS SP2, Leica).

Conflict of Interest: The authors declare no competing financial interest.

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REFERENCES AND NOTES


