Evaporation-Induced Biomolecule Detection on Versatile Superhydrophilic Patterned Surfaces: Glucose and DNA Assay

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Supporting Information

ABSTRACT: We introduce a droplet-based biomolecular detection platform using robust, versatile, and low-cost superhydrophilic patterned superhydrophobic surfaces. Benefitting from confinement and evaporation-induced shrinkage of droplets on wetted patterns, we show enrichment-based biomolecular detection using very low sample volumes. First, we developed a glucose assay using fluorescent polydopamine (PDA) based on enhancement of PDA emission by hydrogen peroxide (H$_2$O$_2$) produced in enzyme-mediated glucose oxidation reaction. Incubation in evaporating droplets resulted in brighter fluorescence compared to that in bulk solutions. Droplet assay was highly sensitive toward increasing glucose concentration while that in milliliter-volume solutions resulted in no fluorescence enhancement at similar time scales. This is due to droplet evaporation that increased the reaction rate by causing enrichment of PDA and glucose/glucose oxidase as well as increased concentration of H$_2$O$_2$ generated in shrinking droplet. Second, we chemically functionalized wetted patterns with single-stranded DNA and developed fluorescence-based DNA detection to demonstrate the adaptability of the patterned surfaces for a different class of assay. We achieved detection of glucose and DNA with concentration down to 130 μM and 200 nM, respectively. Patterned superhydrophobic surfaces with their simple production, sensitive response, and versatility present potential for bioanalysis from low sample volumes.

INTRODUCTION

Superhydrophobic surfaces patterned with well-defined wetted patterns have been important for high-throughput biological and chemical screening inside droplets placed on top since they provide increased reaction rates and sensing signals. Microscale droplets are confined on superhydrophilic patterns due to wetting contrast between hydrophilic patterns and hydrophobic areas. Upon evaporation of droplets, substances dispersed in droplets enrich, which increases molecular interaction frequencies. Such an enrichment effect increases reaction rates, providing enhanced detection signals, whereas signals remain too low to be detected in low-concentration samples. A few reports have demonstrated detection of DNA, RNA, protein, and bacteria inside stationary droplets by exploiting high sensitivity advantage of evaporation-induced enrichment effect. Here, to obtain a practical platform for droplet-based assays, we present the use of superhydrophobic organically modified silica-coated surfaces with circular superhydrophilic patterns generated by converting surface-bound methyl groups into hydroxyl groups via UV/ozone treatment. Two-dimensional chemical patterning without physical decomposition resulted in isolation of aqueous droplets in small wetted spots (Figure 1). These surfaces provide a very strong contrast between wetting and nonwetting regions. Additionally, they can be generated with high precision and they are suitable for large-area processing and stable at a wide range of temperature. Here, we show the versatility of these surfaces using two types of assays, which are based on different chemistries, a novel glucose sensing assay and a DNA hybridization assay.

Hernandez-Perez et al. have recently developed a micro-pillar-based platform and have shown that 2.5 mM glucose can be detected in evaporating droplets placed on pillars. They used horseradish peroxidase (HRP)/glucose oxidase (GOx)/chromogen as the sensing probe. However, HRP has some drawbacks, such as high cost and instability. Therefore, various peroxidase-mimicking nanomaterials have been introduced, including quantum dots, metal nano-clusters, and dyes. Conjugated polymers have also been used as peroxidase-mimicking nanomaterials based on direct or indirect interactions between H$_2$O$_2$ and polymer backbones. Although such materials exhibit high sensitivity for glucose detection, they usually involve complicated synthesis steps and toxicity. Contrary to these materials, polydopamine (PDA) can be easily produced through oxidation and self-polymerization of dopamine in alkaline environments. Particularly, PDA fluorescence has recently found applications in bioimaging and dopamine sensing. Here, we used...
fluorescent PDA for droplet-based glucose sensing probe, which is another major contribution of this work. A sensitive glucose assay was prepared by combining the advantages of PDA, such as biocompatibility and ease of production with analyte/product-enriching capability of patterned surfaces.

PDA fluorescence is known to enhance due to decomposition of its stacked chain structure by $\text{H}_2\text{O}_2$, which is produced in glucose oxidation reaction. To the best of our knowledge, we have exploited PDA as glucose probe for the first time. We first produced PDA from dopamine and incubated with glucose/GOx inside microdroplets placed on wetted patterns. After droplet evaporation, aggregated PDA on the patterns radiated bright fluorescence, which was observed to increase with increasing glucose concentration. However, when incubated in milliliter-volume bulk solution, no significant fluorescence enhancement was obtained within the same time (~50 min). Fast fluorescence enhancement and stronger signals on patterns revealed increase of reaction rate and sensitivity in evaporating droplets.

As a second assay geared toward molecular diagnostics, we functionalized wetted patterns with single-stranded DNA (ss-DNA) and performed fluorescence-based DNA sensing in evaporating microdroplets. Hybridization of target and probe DNA strands inside droplets resulted in higher fluorescence intensities compared to milliliter-volume solution, where no enrichment occurred. Fluorescence of hybridized DNA with

**Figure 1.** Schematic representation of droplet pinning on superhydrophilic circular pattern of a superhydrophobic ormosil surface and evaporation-induced enrichment of low-concentration, low-volume fluorescent sample and resulting fluorescence enhancement.

**Figure 2.** (a) Scanning electron microscopy image of patterned superhydrophobic silica surface. (b) Water droplet dyed with Ponceau 4R, a food additive, on wetted spot and nonwetted region of patterned surface. Droplet was confined on the wetted area, whereas it was repelled from the nonwetted part and maintained its spherical shape due to low surface energy. (c) Time-dependent contact angle profile of a water droplet on superwetted pattern at room temperature.
concentration as low as 200 fM was detected thanks to the enrichment effect of patterned surfaces.

**RESULTS AND DISCUSSION**

Superhydrophilic–superhydrophobic patterned surfaces were produced using a one-pot sol–gel method, followed by UV/ozone treatment on the positions defined (see Materials and Methods section for details). Superhydrophobic coatings were composed of methyl (CH₃)-modified nanoparticles, which were organized in a porous and rough network (Figure 2a). The contact angle of superhydrophobic surfaces was 169° (Figure S1). Superwetted spots with diameter of 0.9 or 1.4 mm were generated on as-prepared superhydrophobic surfaces. To obtain these spots, surfaces were aligned under a shadow mask with holes formed by laser cutting system and treated with UV/ozone, which decomposed hydrophobic CH₃ groups, leaving silanol (Si–OH) groups. Increased π–π* interactions result in aggregation-induced quenching in the course of time (Figure S3). When treated with H₂O₂, PDA fluorescence is found to turn on due to decomposition of PDA polymer to nano-aggregates by H₂O₂. Accordingly, PDA fluorescence was increased with increasing glucose concentration due to H₂O₂ generated through glucose oxidation (Figure 3a,b). Further enhancement of PDA fluorescence was also observed by the naked eye under UV light (Figure 3c). Interestingly, fluorescence maximum was shifted to blue (470 nm) in the presence of glucose with concentration above 5 mM, revealing reduction of nanoaggregate size with gradual increase of H₂O₂. We investigated the effect of H₂O₂ on PDA fluorescence. Besides increasing fluorescence intensity, fluorescence maximum was observed to shift from 500 to 470 nm with increasing H₂O₂ (Figure S4). The limit of detection of fluorescence-based glucose assay was calculated to be 130 μM by plotting the linear calibration curve (inset of Figure 3b).

Fluorescence intensities were recorded after droplets were evaporated, which took approximately 50 min at 37 °C. PDA, which is formed through oxidation and polymerization of dopamine, yields weak green broad-band fluorescence in the wavelength range of 430–600 nm (Figure 3a). Dopamine oxidation produces oligomers, which self-assemble and form PDA through π–π* stacking interactions. Increased π–π* interactions result in aggregation-induced quenching in the course of time (Figure S3). When treated with H₂O₂, PDA fluorescence is found to turn on due to decomposition of PDA polymer to nano-aggregates by H₂O₂. Accordingly, PDA fluorescence was increased with increasing glucose concentration due to H₂O₂ generated through glucose oxidation (Figure 3a,b). Further enhancement of PDA fluorescence was also observed by the naked eye under UV light (Figure 3c). Interestingly, fluorescence maximum was shifted to blue (470 nm) in the presence of glucose with concentration above 5 mM, revealing reduction of nanoaggregate size with gradual increase of H₂O₂. We investigated the effect of H₂O₂ on PDA fluorescence. Besides increasing fluorescence intensity, fluorescence maximum was observed to shift from 500 to 470 nm with increasing H₂O₂ (Figure S4). The limit of detection of fluorescence-based glucose assay was calculated to be 130 μM by plotting the linear calibration curve (inset of Figure 3b).

Although fluorescence enhancement was also observed in bulk liquid, where insignificant evaporation occurs, the reaction was much slower. To investigate the same reaction in bulk, fluorescence intensities were recorded after incubating PDA with glucose/GOx in 4 mL of buffer solutions at 37 °C. Figure 4 shows time dependence of fluorescence intensity at 500 nm. During the initial stages of PDA growth, dopamine emission centered at 460 nm was observed besides PDA emission centered around 500 nm and then gradually disappeared (Figure S5). After the addition of dopamine to oxidized...
glucose, different fluorescence intensities were recorded for different glucose concentrations. However, enhancement response was observed with increasing glucose concentration only after 4 h incubation (Figure 4). Until 4 h incubation, there is no correlation between the glucose concentration and fluorescence intensity at 500 nm. This result can be explained by the suppression of PDA formation soon after dopamine addition to oxidized glucose when H$_2$O$_2$ amount is high. Reorganization of PDA nanoclusters through π-π* stacking interactions in the course of time resulted in fluorescence enhancement. Droplet-based assay accelerated reaction between PDA and H$_2$O$_2$, and reduced the required time to 50 min from 4 h to generate fluorescent enhancement for a wide range of glucose concentration from 1 to 50 mM. Dopamine emission was not observed in evaporating droplets due to high PDA growth rate. It is important to note that shift of fluorescence maximum was not observed in the bulk liquid, revealing the inefficient interaction with H$_2$O$_2$.

Besides glucose detection, patterned surfaces were also used for droplet-based DNA detection. First, probe DNA was covalently attached to wetted patterns using silane chemistry (see Materials and Methods section for details). Briefly, wetted patterns were first epoxylated with (3-glycidoxypropyl)-trimethoxysilane (GOPTS) and then amine-ended DNA molecules were anchored to epoxy groups in sodium carbonate buffer with pH 9 (Figure 5a). Surfaces were washed with bovine serum albumin (BSA) solution to remove the unbound DNA molecules. Binding of GOPTS and DNA to UV/ozone-treated patterns was analyzed by X-ray photoelectron spec-

Figure 4. Time dependence of fluorescence intensities of PDA at 500 nm after incubation with glucose with concentration ranging from 1 to 50 mM in bulk solutions. The complete spectrum for all incubation times is given in the Supporting Information.

Figure 5. (a) Schematic representation of epoxysilane binding onto UV/ozone-treated regions, followed by attachment of probe DNA, hybridization of fluorescence-tagged target DNA, evaporation-induced analyte enrichment, and fluorescence enhancement. (b) Confocal image of wetted pattern after the attachment of Cy5-tagged probe DNA. (c) Confocal image of wetted pattern after hybridization of probe DNA with 20 pM Cy3-tagged target DNA in 4 μL of evaporating droplet.
troscopy. C 1s spectrum of UV/ozone-treated areas showed the decomposition of CH₃ groups (Figure S6). Peaks at 286.7 and 287.7 eV confirmed epoxysilane binding, whereas peaks at 285.4 and 286.6 eV and increase of carbon intensities revealed DNA binding (Figure S7). Binding of probe DNA was also monitored with confocal microscopy using Cy5 tagging. Red emission of Cy5 collected from porous ormosil network shows DNA binding (Figure S7).

DNA was at the same level with buffer solution revealing the specificity of the biosensor for target DNA (Figure 6). Detection of DNA as low as 200 fM was achieved in the assay.

■ CONCLUSIONS

In conclusion, we fabricated a simple detection platform for glucose and DNA assays by using superhydrophilic–superhydrophobic patterned surfaces. A simple UV/ozone-based chemical treatment was used to generate wetted patterns on superhydrophobic coatings produced via sol–gel method. Aqueous droplets with very small volume can be confined on wetted patterns. Analytes dissolved in droplets can be enriched upon evaporation of microdroplets, which are confined on wetted patterns. Polydopamine (PDA) was used as a single fluorescent probe for glucose, for the first time in this study, by benefiting from H₂O₂-dependent fluorescent enhancement of PDA. A glucose assay with reduced response time and increased sensitivity was obtained due to enrichment-induced increase of reaction rates. Contrary to droplet-based assay, fluorescence enhancement was very slow in bulk solutions. Glucose detection with concentration ranging from 1 to 50 mM was shown using bright PDA fluorescence, which can also be observed by the naked eye. PDA with its biocompatibility and promising response curve shows potential for replacing the HRP/chromogen indicators in glucose sensors, especially for wearable and implantable sensors. Furthermore, wetted patterns were functionalized with DNA strands and were used for droplet-based DNA analysis using very low sample volumes. Target DNA with concentration as low as 200 fM can be detected by means of enrichment effect. The presented versatile patterned surfaces with their simple production scheme have potential for a wide range of biochemical analysis.

■ MATERIALS AND METHODS

Materials. Methyltrimethoxysilane (MTMS), oxalic acid, ammonium hydroxide (25%), sodium hydroxide, ethanol, glucose, and hydrochloric acid were purchased from Merck. Methanol was purchased from Carlo-Erba. Glucose oxidase (GOx), dopamine hydrochloride, tris(hydroxymethyl)-aminomethane, phosphate-buffered saline (PBS) tablets, (3-glycidoxypropyl)trimethoxysilane (GOPTS), sodium carbonate, and bovine serum albumin (BSA) were purchased from Sigma-Aldrich. Molecular biology (MB) grade water was purchased from Fisher Scientific. All chemicals were used as received. Amino-modified, Cy5/Cy3-tagged, and tag-free single-stranded DNA (ssDNA) sequences were purchased from Sentegen Biotech. Sequences were as follows:

- Probe DNA: 5′-C6-Amino-TTGGAACATTC-Cy5-3′ (fluorescence-tagged)
- 5′-C6-Amino-TTGGAACATTC-3′ (tag-free)
- Target DNA: 5′-Cy3-GAATGTTCCAA-3′
- Noncomplementary DNA: 5′-Cy3-TGCCGCTGCAC-3′

Preparation of Superhydrophilic–Superhydrophobic Patterned Surfaces. Superhydrophobic coatings were prepared using previously reported sol–gel method. MTMS (1 mL) was dissolved in 9.74 mL of methanol. Following 15 min stirring, 0.5 mL of 1 mM oxalic acid solution was added dropwise and the solution was gently stirred for 30 min. Then, the mixture was left to hydrolyze organosilane precursor completely for 24 h at room temperature. After the hydrolysis step, 0.19 mL of water and 0.42 mL of ammonia solution (25%) were added and gently stirred for 15 min. The resultant mixture was left for 2 days at room temperature to obtain a complete gel. After the aging step, 9 mL of methanol was added, followed by homogenization using an ultrasonic liquid processor for 45 s at 20 W. Portions of resultant colloidal silica solution (250 mL) were spin-coated onto clean quartz or glass substrates (1 × 2 cm²) at 3000 rpm for 45 s. The as-prepared surfaces were covered with cellulose acetate sheets, on which circular holes with 0.9 or 1.4 mm diameter were formed using a 30 W CO₂ laser cutter system (Epilog Zing). Covered surfaces were exposed to UV/ozone for 60 min using a surface cleaning system (PSD-UV, Novascan Technologies) to convert exposed
superhydrophobic regions to superhydrophilic. The power intensities of the UV/ozone cleaner were 8 and 30 mW/cm² for 185 and 253.7 nm wavelengths, respectively.

Characterizations. Static water contact angles of surfaces were measured with a contact angle meter (OCA 30, DataPhysics). Water droplets (20 μL) were used for the measurements. The Laplace—Young equation was used for contact angle calculations. Surface morphology was examined using a scanning electron microscope (Quanta 200F, FEI). Surface chemistry was characterized by X-ray photoelectron spectroscopy (K-Alpha, Thermo Scientific).

Preparation of Buffer Solutions. PBS (100 mM) was prepared by dissolving 10 PBS tablets in 200 mL of deionized water, and the solution pH was adjusted to 7.4 using 1 M sodium hydroxide solution. Tris solution (50 mM) was prepared and its pH was adjusted to 8.6 using 2 M hydrochloric acid. Sodium carbonate buffer (100 mM) was prepared and its pH was adjusted to 9.0 using 2 M hydrochloric acid.

Glucose Assay. PBS solutions (2 mL, pH 7.4), including glucose with concentrations varying from 1 to 50 mM, were mixed with 50 μL of 1 mg/mL GOx solution. Portions of the mixtures (20 μL) were drop-cast onto separate superwetted spots of patterned surfaces and incubated at 37 °C for 30 min. For bulk assay, the solutions were directly incubated at 37 °C for 30 min. Dopamine in Tris–HCl buffer (pH 8.6) with a final dopamine concentration of 0.5 mM was added to the glucose–GOx mixtures. For droplet assay, 20 μL portions of dopamine were added and incubated until droplets were completely evaporated, which took approximately 50 min. The assay time is a function of the hydrophilic spot size, droplet size, and evaporation rate, which should be optimized for any given assay. For bulk assay, dopamine and glucose–GOx mixtures were incubated at 37 °C up to 5 h. A control sample containing GOx enzyme without glucose was also prepared and incubated as all other samples.

DNA Assay. Patterned surfaces were immersed in ethanol solution containing 2% (w/v) GOPTS and incubated at 37 °C overnight. Then, the surfaces were washed with ethanol several times, followed by drying at 120 °C for 1 h. Portions (4 μL) of 1 μM probe DNA (Cy5-tagged or tag-free) in sodium carbonate buffer solutions (pH 9.0) were drop-cast onto superwetted spots. After incubation at 37 °C for 4 h in a closed chamber humidified around 80%, the surfaces were washed with PBS solutions (pH 7.4) to remove the unbound probe DNA strands. The surfaces were then washed with PBS, followed by washing with 1% BSA–PBS solution and MB grade water. Target DNA (4 μL; 200 fM, 20 pM, and 200 pM) in PBS solutions were drop-cast onto probe-bound spots. Microdroplets were left to incubate in the dark at 37 °C for 2 h in a closed chamber with 40% humidity. The surfaces were then washed with PBS (pH 7.4) and MB grade water and dried with nitrogen blow.

Fluorescence Measurements. Fluorescence intensities of droplets and bulk glucose assays were analyzed using a UV lamp (366 nm, 8W, CAMAG) and a fluorescence spectrophotometer (HORIBA, Fluorolog-3, 450 W) by exciting samples at 400 nm and collecting intensities within the 420–600 nm range. Fluorescent images of droplet-based DNA assay were taken using aconfocal microscope (LSM 510, Zeiss using 10× objective. Helium lasers at 543 and 633 nm were used for excitation. Emissions at around 570 and 650 nm were collected for Cy3 and Cy5, respectively.


