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Large-Scale, Lithography-Free Production of Transparent Nanostructured Surface for Dual-Functional Electrochemical and SERS Sensing

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ABSTRACT: In this work, we present a dual-functional sensor that can perform surface-enhanced Raman spectroscopy (SERS) based identification and electrochemical (EC) quantification of analytes in liquid samples. A lithography-free reactive ion etching process was utilized to obtain nanostructures of high aspect ratios distributed homogeneously on a 4 in. fused silica wafer. The sensor was made up of three-electrode array, obtained by subsequent e-beam evaporation of Au on nanostructures in selected areas through a shadow mask. The SERS performance was evaluated through surface-averaged enhancement factor (EF), which was $\sim 6.2 \times 10^5$, and spatial uniformity of EF, which was $\sim 13\%$ in terms of relative standard deviation. Excellent electrochemical performance and reproducibility were revealed by recording cyclic voltammograms. On nanostructured electrodes, paracetamol (PAR) showed an improved quasi-reversible behavior with decrease in peak potential separation ($\Delta E_p \sim 90$ mV) and higher peak currents ($I_{pa}/I_{pc} \sim 1$), compared to planar electrodes ($\Delta E_p \sim 560$ mV). The oxidation potential of PAR was also lowered by $\sim 80$ mV on nanostructured electrodes. To illustrate dual-functional sensing, quantitative evaluation of PAR ranging from 30 $\mu$M to 3 mM was realized through EC detection, and the presence of PAR was verified by its SERS fingerprint.

KEYWORDS: lithography-free, dual-functional, electrochemical, SERS, paracetamol

Electrochemical (EC) detection and Raman spectroscopy have been historically linked since the discovery of surface-enhanced Raman spectroscopy (SERS), where it was found that nanostructured electrode surfaces, typically achieved via electrochemical roughening, introduced nanoscale hot spots that significantly enhanced the Raman signal.1,2 In recent years, SERS has become an analytical tool, allowing detection of analytes at ultralow concentrations by providing their molecular vibrational fingerprints.3–5 It has been widely used in biomedical diagnostics, environmental sensing, food safety, and illicit drug control.6–8 On the other hand, extensive research has been carried out, using defined nanostructures as electrodes for electrochemical sensing. By using electrodes consisting of nanostructured surfaces, such as nanoporous metallic structures, nanoparticles,9 nanotubes,10 nanowires,11–13 and vertically aligned nanopillars,14,15 performance of electrochemical sensing has been improved dramatically, due to the unique electrical, chemical, and catalytic properties, and increased surface areas of the nanostructures.16

The combination of SERS with electrochemistry (EC-SERS) has led to research in various areas, including corrosion and surface science,17 fuel cells, electrocatalysis, and surface redox reactions,18–20 as well as adsorption and reaction of biomolecules.21,22 Particularly, EC and SERS have been used as complementary techniques to reveal and explain chemical phenomena, or as a way to improve sensing performance. For example, coupling EC with SERS has been proven beneficial for monitoring electrochemical reactions at ultralow concentrations. In addition, by applying a potential, the adhesion of target analytes to the Au surface can be promoted. This would increase the number of analytes in the electromagnetic hot spots, improving the SERS detection limits.23,24 A more advanced technique involves using fluctuation of the applied electrode potential to adsorb and desorb different sets of molecules while monitoring the Raman signatures and thereby avoiding spectral congestion. This is especially relevant with coadsorption of molecules in complex solutions, typically biological solutions.22

As application expands by combining EC with SERS, in recent years, focus has been addressed on developing substrates...
that are dual-functional, i.e., integrating EC and SERS sensing on a single chip. In recent studies, Chen et al. reported the deposition of Ag nanowires on a coffee filter by a simple dip-coating route to develop an enzyme-free sensor for hydrogen peroxide detection.25 Bailey et al. reported polystyrene encapsulated Au electrodes with the electrochemically roughened surface for studying surface chemistry of neurotransmitters.26 Furthermore, to study neural stem cell differentiation, Au nanostar arrays were electrochemically deposited on indium tin oxide substrates.27 Single cell monitoring was achieved on zinc oxide nanorods grown on planar microelectrodes with Au electroplating.28 For studying biotin–avidin interaction, silicon nanowires were grown on microscope slides by plasma enhanced chemical vapor deposition, and the structures were coated by Au, rendering simultaneously working electrode and SERS probes.29 In addition, dual sensing was achieved on commercial Au electrode and was applied for evaluation of drug intercalation into a DNA duplex. While the molecular identification profiles were provided by SERS, the concentration dependent modification of the DNA by the drug was evaluated by EC detection.30

The success of combining EC and SERS relies on the nanostructured enhancing surfaces. Fabrication techniques have typically involved chemical synthesis and electrodeposition, sometimes in combination with lithographic techniques.32 For solid substrates, they are usually patterned by lithography, with a subsequent metallization process.33–35 The main drawback of lithographic techniques is that they are time-consuming and costly. Lithography-free reactive ion etching (RIE), on the other hand, has been proven to be a good alternative, since it is much faster and significantly less expensive.36–38 For example, using maskless RIE, wafer-scale, high performance SERS substrates consisting of nanorods that are able to form clusters, can be manufactured.39,40 However, the semiconducting nature of silicon has prevented these structures to be used as dual-functional sensors for EC and SERS. Recent developments have shown that lithography-free nanostructuring of insulating and chemically stable materials, such as fused silica (glass), is possible.36,41,42 This opens up new possibilities for integrating EC and SERS on a single chip, in a cost-effective manner, which is also reproducible and suitable for mass production.

In this paper, we report an easy to implement, inexpensive, and fast lithography-free process, for manufacturing a dual-functional EC and SERS sensor system, and present its characterization and application. Dense and homogeneous nanostructures with high aspect ratios were obtained on a 4 in. fused silica wafer using mask-less RIE process. By e-beam evaporation of Au on these nanostructures and was applied for evaluation of drug intercalation into a DNA duplex. While the molecular identification profiles were provided by SERS, the concentration dependent modification of the DNA by the drug was evaluated by EC detection.30

### EXPERIMENTAL SECTION

**Materials.** All chemicals and reagents were used as received. Potassium hexacyanoferrate(II) (ferrocyanide), potassium hexacyanoferrate(III) (ferricyanide), phosphate buffered saline (PBS) containing 0.01 M phosphate buffer, 0.0027 M potassium chloride, and 0.137 M sodium chloride (pH 7.4), trans-1,2-bis (4-pyridyl) ethylene (BPE), methanol, and paracetamol were purchased from Sigma-Aldrich Co. (St. Louis, MO, USA). Absolute ethanol was from

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**Figure 1.** Illustration of the fabrication process of gold capped glass nanopillars. (A) Al layer deposition through a shadowmask, (B) RIE resulting in fused silica nanostructures, (C) electrode patterning using Au deposition through a shadowmask, (D) photograph of whole assembly with inset showing three-electrode arrays, SEM images showing areas of silica nanostructures with (bottom) and without (top) Au deposition, (E) SEM images (top view and side view) of fused silica nanostructures with an evaporated Au layer (All SEM images have scale bar = 1 μm).
Kemetyl A/S (Koge, Denmark). All aqueous solutions were prepared in ultrapure water obtained from a Milli-Q water purification system (Millipore Corporation, Billerica, MA, USA). A 4 in. double-side-polished fused silica wafer with a 525 μm thickness (Innergie, Corporation, Waltham, MA, USA) was used to cut the desired pattern on PMMA (Evonik Industries AG, Darmstadt, Germany) and double-sided pressure sensitive adhesive (PSA) tape (140 μm thick, ARcare 90106 from Adhesive Research Ltd., Limerick, Ireland) were used.

**Fabrication of Dual-Functional Sensor.** The nanostructures were obtained using RIE technique on 4 in. fused silica wafer. Prior to performing RIE process, AI mask (600 nm thick) was deposited through a shadow mask via e-beam evaporator (Wordentec QCL 800 E-beam and sputtering deposition system). The 4 min RIE procedure at an O₂ flow of 70 sccm, an SF₆ flow of 170 sccm, a platen power of 110 W, a coil power of 450 W, chuck temperature 5 °C and a chamber pressure of 75 mTorr was performed using advanced oxide etcher (STS MESC Multiplex ICP). Further, 170-nm-thick Au film in the form of three-electrode array was achieved over the nanostructures using the e-beam evaporator. The morphology of nanostructures was characterized using a scanning electron microscope (Zeiss Supra VP 40 SEM). The planar electrodes were patterned on the fused silica wafer using e-beam evaporation (17 nm Cr adhesion layer/170 nm Au) through a shadow mask that was previously used to evaporate Au on nanostructured surface. The working area of the electrodes and the wells (150 μL) to accommodate the sample over the electrodes were defined by utilizing 0.6-mm-thick PMMA and 140-μm-thick double-sided PSA tape. The pattern to cut PMMA and PSA was designed using the computer aided design (CAD) software SolidWorks (Dassault Systèmes SolidWorks Corporation, Waltham, MA, USA). A computer-controlled Epilog Mini 18 laser cutter equipped with a 30 W CO₂ laser (Epilog Laser, Corporation, Waltham, MA, USA) was used to cut the desired pattern on PMMA, whereas Cameo 3 blade cutter (Silhouette, Inc., Lindon, UT, USA) was utilized to cut the PSA. Finally, the laser-cut PMMA was bonded to the fused silica substrate, incorporating the sensors, using PSA (see Figure 2 for final assembly). Simultaneous electrical connections to all eight sets of three-electrode arrays were achieved using custom-made printed circuit board with magnetic clamping, as described in our previous work.

**SERS Measurements.** SERS measurements were carried out using a Raman microscope (DXRxi Raman Imaging microscope, Thermo Scientific, Waltham, MA, U.S.) equipped with a 780 nm laser and a 10× objective. The sample was placed on a motorized stage controlled by using a microscope controller (Shamrock Spectrograph SR-303i-A, equipped with an Andor Newton 970 EMCCD). The obtained scattering spectrum was then divided by the reference white light spectrum to obtain the scattering signal from the substrates.

**Electrochemical Measurements.** All electrochemical measurements were carried out at room temperature using an eight-channel CHI1030A potentiostat (CH Instruments, Inc, Austin, TX, USA) connected simultaneously to eight sets of three-electrode arrays fabricated on 4 in. fused-silica wafer. For the electrode characterization, cyclic voltammograms (CVs) were recorded from −400 mV to 400 mV vs Au pseudoreference electrode (RE) in 10 mM ferri/ferrocyanide using PBS as supporting electrolyte at scan rate 50 mV s⁻¹. Electrochemical fingerprint of PAR in PBS was recorded using CV from −400 mV to 600 mV vs Au pseudo-RE at scan rate 50 mV s⁻¹. The quantification was performed with batch amperometry by successive additions of PAR at an applied potential of 450 mV vs pseudo-RE in PBS.

#### RESULTS AND DISCUSSION

**Fabrication Process.** The fabrication process of nanostructured surface obtained on fused silica wafer using a lithography-free process (see Experimental Section) is illustrated in Figure 1A,B. A previously described masking technique was utilized during the RIE process. Accordingly, a seed layer of Al was used to sputter nanoscale masks onto the fused silica surface. The Al pattern consists of an outer ring that ensures reduction of the maximum distance of the exposed fused silica wafer to areas with Al so that sputtering of nanomasks occurs evenly across the wafer. In this way, the arrangement of Al seed layer promoted fabrication of uniformly distributed dense nanostructures (Figure S1). Subsequent evaporation of Au on these nanostructures through a shadow mask resulted in formation of isolated metal caps and defined the working, counter, and reference electrodes (Figure 1C). SEM images of the Au capped fused silica nanostructures are shown in Figure 1D,E.

**SERS Characterization.** SERS performance of the Au-capped nanostructures was evaluated using trans-1,2-bis(4-pyridyl)ethylene (BPE) dissolved in ethanol. Figure 3A shows the distribution of the Raman signal intensity, recorded from 5 μL of 1 mM BPE, which was evenly deposited on the sensor surface. In addition, we were able to measure BPE both from the front as well as through the backside of the substrate. As shown in Figure 3C, the measurement through the back of the substrate showed a 37% lower SERS signal compared to those recorded from the front, with background signal recorded using pure ethanol. These measurements showed a strong enhancement with minimal background signal and the ability to measure from the back of the sensor. Furthermore, we found a homogeneous signal enhancement with a relative standard deviation of 13.4%, as shown in Figure 3C.

The surface-averaged SERS enhancement factor (EF) of the substrate was calculated from the formula:

$$\text{EF} = \frac{I_{\text{SERS}}/N_{\text{SERS}}}{I_{\text{RS}}/N_{\text{RS}}}$$

where $I_{\text{SERS}}$ and $I_{\text{RS}}$ were the SERS and the Raman intensity, respectively. $N_{\text{SERS}}$ and $N_{\text{RS}}$ were the numbers of molecules probed in the SERS and the Raman measurements. To establish the value of $N_{\text{RS}}$, a 100 mM ethanol solution of BPE was poured into a Petri dish. 1.4 counts mW⁻¹s⁻¹ ($I_{\text{RS}}$) were measured for the 1200 cm⁻¹ peak under an excitation wavelength of 780 nm. The number of molecules excited ($N_{\text{RS}}$),
was estimated to be $3.60 \times 10^9$, using the interaction volume of 60 $\mu$m$^3$. To establish the values for $I_{\text{SERS}}$ and $N_{\text{SERS}}$, 1 $\mu$L of 1 mM ethanol solution of BPE was pipetted onto the substrate. Hereafter, the substrate was dried in air. The introduced BPE solution forms a quasi-circular pattern on the surface of the substrate with a radius of $\sim 3400$ $\mu$m. This leads to a density of BPE on the substrate to be $1.67 \times 10^7$ $\mu$m$^{-2}$. Using identical experimental conditions as for $I_{\text{RS}}$, $\sim 8000$ mW$^{-1}$s$^{-1}$ ($I_{\text{SERS}}$) were measured for the 1200 cm$^{-1}$ peak. By assuming uniform adsorption of BPE onto Au and a laser spot size of 0.8 $\mu$m in radius, $N_{\text{SERS}}$ is then estimated to be $3.33 \times 10^7$. Hence, the calculated surface-averaged EF becomes $6.2 \times 10^5$. Note that this EF is a surface-averaged value, obtained by assuming that all the adsorbed molecules contributed equally to the integrated SERS signal, and is thus not the maximum enhancement, which others have attempted to calculate by estimating the number of molecules in the hot spots.

Dark-field scattering spectrum of the substrate is obtained as shown in Figure 3B. A broad peak ranging from $\sim 600$ nm to $\sim 1000$ nm is observed. This is because of the tremendous randomness of the structures, which enables the substrate to support extensive localized plasmon resonance modes. A wide range of excitation wavelengths is thus selectable to perform efficient SERS analysis for different applications. In addition, the dip at $\sim 500$ nm is caused by the interband transition of Au.

**EC Characterization.** EC performance and reproducibility of the nanostructured electrodes were evaluated by acquiring CVs in 10 mM ferro/ferricyanide dissolved in PBS. The electrodes showed a quasi-reversible electrochemical response with a peak potential separation ($\Delta E_p$) of $99 \pm 2$ mV at 50 mV s$^{-1}$ and the ratio between the reduction and oxidation peaks was 1.09 $\pm$ 0.03 as shown in Figure 4A. The recorded CVs confirm an excellent reproducibility of the electrodes.

According to SEM images shown in Figure 1E, the majority of Au that defined the electrode area consists of Au nanoparticles with randomized morphologies. Therefore, the following equation can be established to estimate the 3D surface area of the nanostructured WE, $S_{3D}$:

$$S_{3D} = \overline{S}_{\text{NP}} \times D_{\text{NP}} \times S_{2D}$$

where $\overline{S}_{\text{NP}}$ is the average surface area of a Au nanoparticle, $D_{\text{NP}}$ is the average density of the Au nanoparticles, and $S_{2D}$ is the two-dimensional surface area of the WE. Approximating each Au nanoparticle by an ellipsoid with lengths of axes being 100, 100, and 200 nm, 54,000 nm$^2$ is estimated for $\overline{S}_{\text{NP}}$. From top-view SEM images, $D_{\text{NP}}$ is estimated to be 35 $\mu$m$^{-2}$. As per the electrode design shown in Figure 1D, $S_{2D}$ is 14.92 mm$^2$. These lead to an estimated $S_{3D}$ of 28.20 mm$^2$. Thus, the surface area of the nanostructured electrode is increased by a factor of 1.89 relative to the planar electrode. Figure 4B shows a comparison of CVs for nanostructured and planar electrode in PBS. Due to increase in surface area, nanostructured electrode exhibit higher currents for charging double layer capacitance at the electrode/electrolyte interface as compared to the planar electrode. However, the CVs recorded in the presence of redox-active species did not confirm the significant increase in surface area for nanostructured electrode (Figure 4C). The possible explanation could be that the wetting of the nanostructured electrode was not complete, i.e., the electroactive probe did not have access to the crevices of the nanostructured electrode.

**Dual Sensing Detection and Quantification of Paracetamol.** PAR was chosen as a model analyte to be identified...
and quantified using the developed dual-functional sensor. As shown in Figure 5A, the presence of PAR, based on its Raman fingerprint, was confirmed with SERS-sensing at different concentrations. For SERS detection, in order to reduce interferences caused by salts present in the PBS and to increase the wettability of the sensing surface, the samples were diluted with 10% methanol. In comparison with the control measurement, distinct fingerprint peaks were observed for paracetamol (e.g., vibrational mode at 1167 cm\(^{-1}\)).

The improved electrochemical behavior of PAR on surface modified glassy carbon electrodes have been already shown in the literature.\(^{44-46}\) However, the presented nanostructured electrode exhibited an excellent electrocatalytic activity to PAR without any surface modification as compared to the planar electrode with similar geometry. In the case of planar electrode, the PAR was oxidized at 380 mV, whereas the oxidation potential was lowered by 80 mV for a nanostructured electrode with higher peak currents (\(I_{\text{pa}}/I_{\text{pc}} \sim 1\)) (Figure 5B). This can explain an efficient catalytic role of the Au-coated nanostructured surface that provides enough reaction sites for the enhancement of the current response toward PAR. For 1 mM PAR, oxidation and reduction peak currents on nanostructured electrodes were 40 \(\mu\)A \(\pm\) 2 and 40 \(\mu\)A \(\pm\) 2, and on planar electrodes were 22 \(\mu\)A \(\pm\) 2 and 15 \(\mu\)A \(\pm\) 1, respectively. In addition, PAR showed an improved quasi-reversible behavior on the nanostructured electrode with peak potential separation (\(\Delta E_p\)) of 90 mV as compared to 560 mV for the planar electrode. The decrease in the \(\Delta E_p\) value of the nanostructured electrode confirms the effective electron transfer ability of the Au-coated nanostructured surface. The CV was also acquired in 10 mM PBS with or without PAR to confirm that the oxidation and reduction peaks solely belongs to the PAR (Figure S2). As shown in Figure 5C, the linear range, from 30 \(\mu\)M up to 3.0 mM, was obtained when quantifying PAR on the

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**Figure 4.** CVs for (A) 16 electrodes patterned on two different substrates (each carrying 8 electrodes) acquired in 10 mM ferri/ferrocyanide using PBS as supporting electrolyte, red shaded area shows the standard deviation among the electrodes. (B) and (C) planar Au electrode vs nanostructured gold electrode in PBS and in 10 mM ferri/ferrocyanide using PBS as supporting electrolyte, respectively. (scan rate 50 mVs\(^{-1}\)).

**Figure 5.** (A) SERS measurement of paracetamol in PBS showing clear peaks for different concentrations of paracetamol. (B) CV of 1 mM PAR on planar and nanostructured electrodes. (C) Calibration curve of paracetamol obtained by amperometry at an applied potential of 450 mV in PBS vs Au pseudo-RE.
nanostructured electrodes using amperometry at an applied potential of 450 mV. In addition, attributing to the slightly higher surface area, nanostructured electrodes showed improved sensitivity as compared to the planar electrodes during quantification (Figure S3).

■ CONCLUSION

A dual-functional sensor for EC and SERS-sensing consisting of nanostructured three-electrode array was developed. The sensor was successfully applied for quantitative detection of PAR, during which EC and SERS served, respectively, for quantification and identification, complementing each other. The developed lithography-free fabrication process is reproducible, high-throughput, and inexpensive, and is thus suitable for commercialization. The high optical transparency of the substrate allows excitation and collection of SERS spectra from the backside of the sensors. Thereby, detecting analytes in complex samples containing, e.g., cells, becomes possible. In addition, the sensors patterned on a circular wafer would be suitable for integration into centrifugal microfluidic platforms, where a certain part of the assay, such as sample filtration and analyte concentration, can be performed before sensing.

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Notes
The authors declare no competing financial interest.

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