Nanowell Array based Sensor and Its Packaging

JuKyung Lee1, Tsuda Akira2, Myung Yung Jeong3 and Hea Yeon Lee1,†

1Department of Mechanical and Industrial Engineering, College of Engineering, Northeastern University, Boston, MA02115, USA
2Department of Environmental Health, Harvard School of Public Health, Boston, MA 02115, USA
3Department of Congo-Mechatronics Engineering, Pusan National University, Busan 609-735, Korea

(Received September 5, 2014: Corrected September 18, 2014: Accepted September 19, 2014)

Abstract: This article reviews the recent progress in nanowell array biosensors that use the label-free detection protocol, and are detected in their natural forms. These nanowell array biosensors are fabricated by nanofabrication technologies that should be useful for developing highly sensitive and selective also reproducible biosensors. Moreover, electrochemical method was selected as analysis method that has high sensitivity compared with other analysis. Finally, highly sensitive nanobiosensor was achieved by combining nanofabrication technologies and classical electrochemical method. Many examples are mentioned about the sensing performance of nanowell array biosensors will be evaluated in terms of sensitivity and detection limit compared with other micro-sized electrode without nanowell array.

Keywords: Nanowell Arrays (NWA), Nanobiosensor, Electrochemical (EC) method, Chip-packaging

1. Introduction

Recently, biosensors of nanoscale dimension have attracted wide attention for sensitive, multi-targeting, and labeling-free biomolecular detection. Various affinity-binding assisted analytical tools have been developed based on nanowires, nanoparticles, nanotubes. The widespread use of these tools, however, has been limited by random configuration of binding molecule and the formation of protein-protein cluster on a solid surface, resulting in a loss of masking of binding sites. Since the binding event is highly specific and varies depending on the targeting strategy employed, the biosensing platform needs to be flexible and constructed with many functional groups. To overcome this limit, we have developed nanowell array structures for reducing non-specific binding that highly affect to biosensor’s sensitivity and selectivity. In this geometry, most of the area of the Au electrode is covered with a blocking layer, and only a nanosized gold surface is exposed to the open space above each nanowell array (NWA). The depth and width of the nanowells can be adjusted to allow only one or a few biomolecules to enter the well and become attached to the self-assembled modified gold surface. The number of wells required in a NWA to obtain the desired level of assay signal is easily estimated and an appropriated array can be fabricated.

Electrochemical (EC) detection was selected as analysis method that has high sensitivity compared with other classical analysis. We have reported that NWA can enhance EC detection of molecular binding events by controlling the binding sites of the captured molecules. Using NWA biosensor based direct current (DC) and Alternative current (AC) analysis, we have detected biological macromolecules such as DNA, protein as antigen or aptamers at low concentrations. Moreover, miniaturization can be achieved by combining this simple EC method and nanosized electrode. The packaged device was tested for various type of biosensor. This system permits a much simpler packaging technique to be utilized and possesses advantages for many of the medical applications.

In this article, we will review a well-oriented NWA that is easily fabricated using current lithography technology and has numerous applications of biosensors. First, we will examine how to make this NWA structure by using various typed lithography. After this, we will also examine various applications of EC biosensors based on this NWA electrode. The area has been researched about our work through the use of on-line databases and browsing of journal content.
2. Fabrication of Nanowell Array Electrode

2.1. Lithographic Molding

Lithographic molding is lithography for the fabrication of elastomeric stamps for microcontact printing (µCP). This technique provides elastomeric stamps with sub-micrometer features and does not require routine access to clean rooms; it is thus an alternative to photolithography for the fabrication of stamps for µCP. Lithographic molding combines several steps: i) microcontact printing on a thin gold film supported on a silicon (100) wafer to form a patterned SAM; ii) etching of the gold film, using the patterned self-assembled monolayers (SAMs) as resists iii) anisotropic etching of silicon which reduces the scale of certain features of the original pattern-in regions where removal of gold has exposed the underlying silicon wafer; iv) molding of a new elastomeric stamp using the etched silicon as a master, and v) µCP with this new elastomeric stamp. This demonstration is one of several that explore alternatives to the photolithographic techniques normally used for the production of sub-micrometer scale features.

A schematic procedure for the fabrication of polyethylene glycol (PEG) NWA is shown in fig. 1. In this technique, a nanopatterned polymeric mold is directly placed on a drop-dispersed PEG solution before solvent evaporation. For the mold material, a UV-curable polyurethane acrylate (PUA) was used, which was recently introduced for lithography of features smaller than 100 nm. Using this method, various nanopatterns of a PEG random copolymer were fabricated using positive PUA molds with different feature size (Fig. 1b). AFM image demonstrated that these patterns were neatly constructed with the same lateral dimension with that of the original mold. The height varied depending on the geometry of the mold and the initial concentration/amount of the polymer solution, ranging from 70 to 200 nm.

2.2. Wafer-scale NWA patterning using KrF stepper

In general, e-beam lithography (EBL), nanoimprint lithography (NIL) are capable of creating nanopatterns that can be used to fabricate NWA. But these methods have low throughput and are limited to small areas. We also used

Fig. 1. (a) A schematic illustration for the fabrication of PEG nanowells with exposure of the substrate. (b) Planar and cross-sectional AFM images of the fabricated PEG nanopatterns on gold substrate: i) Squares of 300 nm width and 70 nm height. ii) Lines of 200 nm width and 80 nm height. iii) Holes of 80 nm diameter and 70 nm height. Reproduced with permission from ref. number.

Fig. 2. Fabrication of NWA based electrochemical sensor. (a) Simple and high-throughput nano process for industrial application. (b) Photograph of as-fabricated samples (top) and SEM images of NWA electrode (bottom). Reproduced with permission from ref. Num.
krypton-fluoride (KrF) stepper semiconductor process with a wavelength of 248 nm for the fabrication of NWA electrodes on 6 inch wafer. This wafer-scale fabrication method of nanopatterns has rapid, high-throughput and is highly reliable for the fabrication of NWA biosensor. As shown fig. 2 (a), a 6 inch wafer was pre cleaned using SPM (sulfuric peroxide mixture, H2SO4 : H2O2 = 4:1) and diluted HF to remove the organic contaminants and native oxide layer on the substrate, respectively. A 300 nm thick SiO2 layer was deposited on the Si substrate by plasma-enhanced chemical vapor deposition (PECVD). Au layer was deposited by sputtering. After micro-patterning, the Ti/Au layers were etched in an ICP etcher. Finally, the exposed SiO2 areas were etched using an ICP etcher, and photoresist was removed. Also we designed nanowell array electrode with electronic pad for efficient packaging, Fig. 2 (b) shows the 57 uniform and well-fabricated NWA electrodes on 6-inch wafer fabricated by KrF stepper semiconductor process for industrial application. The size of a single NWA sensor was 21×10 mm². Each sensor consisted of two NWA with a size of 4×2 mm². Each NWA structure had a diameter of 500 nm with an interwell spacing of 200 nm.  

2.3. Chip-Level Packaging

There exist several challenges when developing the packaging process for the sensor chip. Some of these challenges are addressed in the sensor design stage such as sensor size and interconnections.  

The goal of the packaging process is to enable the sensor to its intended specifications but causing minimal effect to its environment and the sensor itself. Fig. 2 is also shown about our chip-design for packaging. NWA (wide) and electronic pad (narrow) region are interconnected by Au pad. In order to reduce noise (that is most important factor for EC biosensor), we covered other region by PR that is insulating layer and reduce interconnection channel. Also we designed the chip to enable multi-targeting. Figure 3 shows the process for fabrication of a replica mold from Methacrylate octa-functionalized silsequioxane (SSQMA)-based formulation on a flexible and transparent poly-ethylene terephthalate (PET) film via UV-embossing. The duplicated pattern of parallel lines with a half-pitch of 50 nm showed almost the same half-pitch as the Si master mold. Rigid replicas duplicated onto a flexible backplane are called “rigiflex” replica molds. These results demonstrate that the SSQMA-based formulations are suitable for large-area replication with sub-50-nm features.

Rigiflex replica molds exhibit both local rigidity and global flexibility, which are advantageous in many applications. In particular, even when the substrate is not flat, a flexible mold can provide large-area conformal contact with the substrate without requiring high pressures. This method demonstrated good reliability and performance. Also deposition, patterning and packaging are performed by simple process, this method has advantage of providing a continuous and high throughput. The cost of production can be dramatically reduced.

3. NWA based Biosensor

3.1. Single Liposome based Sensor

Nanoscale arrays of functional lipid vesicles (FLVs) are potentially useful when employed in electrochemical studies and applications. Construction of such arrays on gold electrode with controlled dimension and density could allow for quantitative analysis on the level of single lipid vesicle with more reaction sites and low signal-to-noise ratio. To achieve a nanowell-based electrochemical biosensor with FLVs, three of the key directions are to address how to (I) capture FLVs at predefined locations without non-specific binding, (II) make the captured FLVs stable and functional,
and (III) quantitatively interpret signals from affinity binding. Of these, non-specific binding has been a hurdle to most label-free detection methods since many proteins in a biological sample adhered to the surface non-specifically. Thus it would be great benefit to apply NWA electrode with controlled geometry and density that can capture individual liposomes without non-specific binding.

To evaluate potential applications of the site-selective deposition of single FLVs, we tested streptavidin binding to biotin on the FLVs surface. This coupling is widely studied for development of biosensor for its strong affinity. Fig. 4 (a) illustrates the streptavidin-biotin interactions using biotinylated FLVs containing thiol, ferrocence and PEG moieties on the surface of FLVs. Fig. 4 (b) shows SWV measurements for the change of current density on both electrodes (dash lines: before streptavidin-biotin binding, solid lines: after streptavidin-biotin binding). The SWV data demonstrated that the electrochemical responses (both magnitude and difference in peak current densities) were significantly enhanced for the NWA electrode: the difference in peak current densities (DIP) were 33.7 pA µm² and 0.187 pA µm² for the NWA gold and the bare electrodes, respectively. Based on the active area exposed to the electrolyte, this difference corresponds to approximately 220 times increase in signal amplification for the NWA electrode, which could ultimately lead to a substantial increase in sensitivity.

3.2. Electrochemical Immunosensor

EIS (Electrochemical Impedance Spectroscopy) based immunosensors typically utilize antigen-antibody immunolayers that are very thin and have low electric permittivity. Molecular binding is detected by interruption of the faradaic current at the electrode, which generate an impedance signal. We developed an highly sensitive immunosensor for

![Fig. 4](image-url)

**Fig. 4.** (a) a schematic illustration of the binding event between streptavidin and captured biotinylated FLVs on NWA electrode, (b) SWV measurements of current density before and after addition of streptavidin using a series of pulse input. The NWA electrode with non-biotinylated FLVs was used as the negative control. All the SWV data were obtained at a scan rate of 200 mV/s. Reproduced with permission from ref.num.

![Fig. 5](image-url)

**Fig. 5.** EIS binding of STIP-1 to the immunolayer on the electrodes. (a) Nyquist plot for the NWA electrode, (b) Nyquist plot for the bare electrode, (c) Quantitative analysis of STIP-1 and anti STIP-1 antibody binding. Biotinylated anti-STIP-1 antibody layer was labeled with streptavidin on the NWA and bare electrodes. Error bars represent the standard deviations of three measurements. The applied potential was set to 50 mV against an Ag/AgCl reference electrode. The frequency range was 0.1 Hz to 1 MHz. Reproduced with permission from ref.Num.
quantiﬁcative detection of speciﬁc antigen (stress-induced-
phosphoprotein-1) [STIP-1] using NW A electrodes. Fig. 5 (a) and (b) show the Nyquist plot for NW A electrodes and
bare electrodes without NW A using STIP-1 as a model
analyte. In a nyquist representation, the real component
of the complex impedance is shown on the x-coordinate, and
the imaginary component on the y-coordinate. To compare
the sensitivity of the NW A biosensor with that of other
electrode-based biosensors, the same treatments were
applied to a bare electrode without NW A. The LOD was
estimated to be 10 pg/mL or less for the NW A electrodes,
which suggests a 100-fold improvement in the LOD when
using EIS with NW A electrode as shown ﬁg. 5 (c). The
sensitivity of the NW A impedimetric immunosensor was
better for each analyte concentration tested when compared
the sensitivity of the bare electrode sensor.

Based on these results, the electrochemical impedimetric
immunosensors using NW A electrodes can be applied for
label-free detection, with low levels of non-speciﬁc binding.
Because NW A are optimal for the selective docking of
single molecules, they reduce non-speciﬁc binding and
enhance electrochemical responses. Therefore NW A have
high sensitivity and selectivity as well as very low LOD.

4. Conclusions

In this article, we overviewed recent progress about NW A
biosensors from its fabrication to its application. There are
several potential advantages of NW A in biosensor
applications. First, NW A can act as a digital assay by
increasing the S/N ratio. The nanowell geometry should
minimize unwanted nonspeciﬁc binding and decrease the
noise signal, since the resist layer can restrict many
biomaterials from becoming attached to the EC active gold
electrode surface. Second, the array can be used in
numerous other integrated biosensor systems, such as in
electronic signaling, ﬂuorescence, and bioluminescence.
Third, the fabrication of the NW A is highly compatible with
recent advanced nanotechnologies and is quite economical.
Also we designed the NW A by considering the packaging
process. Improvement in the packaging process has resulted
in a compact single-chip NW A electrode for reducing noise
when measure the signal.

This type of array can be adapted relatively easily and
used with a high throughput method. Finally, the NW A
geometry should enhance the signal sensitivity so much that
in the near future it might be feasible to detect a single
biomolecular reaction. This nanometric system can also be
applied to other integrated digital biosensors.

Acknowledgements

This research was supported by the Basic Science
Research Program of the National Research Foundation of
Korea (NRF), funded by the Ministry of Education, Science
and Technology (2011-0014610).

References

1. J. M. Kim, H. S. Jung, J. W. Park, H. Oka, T. Yukimasa, H.
Y. Lee and T. Kawai, “Spontaneous Immobilization of Lipo-
somes on Electron Beam Technique”, J. Am. Chem. Soc.,
127(7), 2358 (2005).
2. H. S. Jung, J. M. Kim, J. W. Park, H. Y. Lee and T. Kawai,
“Amperometric immunosensor for direct detection based
upon functional lipid vesicles immobilized on nanowell array
electrode”, Langmuir : the ACS journal of surfaces and col-
loids., 21(13), 6025 (2005).
3. J. L. Wilbur, E. Kim and Y. Xia. “Whitesides GM. Litho-
graphic molding: A convenient route to structures with sub-
micrometer dimensions**”, Advanced Materials., 7(7), 649
(1995).
4. P. Kim, B. K. Lee, H. Y. Lee, T. Kawai and K. Y. Suh.
“Molded Nanowell Electrodes for Site-Selective Single Lipo-
some Arrays”, Advanced Materials., 20(1), 31 (2008).
5. J. Lee, S. Cho, H. Ryu, J. Park, S. Lim, B. Oh, C. Lee, W.
Huang, A. Busnaina and H. Lee. “Wafer-scale nanowell array
pattening based electrochemical impedimetric immunosens-
sor”, Journal of biotechnology., 168(4), 584 (2013).
6. D. C. Ng, T. Nakagawa, T. Mizuno, T. Tokuda, M. Nunoshiba,
H. Tamura, Y. Ishikawa, S. Shiosaka and J. Ohta. “Integrated
in vivo neural imaging and interface CMOS devices: design,
packaging, and implementation” Sensors Journal, IEEE., 8(1),
121 (2008).
7. B. K. Lee, N. G. Choa, H. Tanaka, N. Y. Hong, D. P. Kim,
H. Y. Lee, T. Kawai. “Photocurable Polyhedral Oligomeric
Silsequioxane-based Resists for Nanoimprint Lithography:
Fabrication of High-aspect Ratio Structures and Replica
Molds”, Langmuir : the ACS journal of surfaces and colloids.,
26(18), 14915 (2010).
JuKyung Lee, Tsuda Akira, Myung Yung Jeong and Hea Yeon Lee

- Ju Kyung
  - Northeastern University, department of mechanical and industrial engineering
  - Electrochemical biosensor
  - e-mail: chejueyes@gmail.com

- Tsuda Akira
  - Harvard school of public health, department of environmental health
  - Nanoparticle-lung interactions
  - e-mail: atsuda@hsph.harvard.edu

- Myung Yung Jeong
  - Pusan national university, department of congo-mechatronics engineering
  - Nano imprint, surface engineering
  - e-mail: myjeong@pusan.ac.kr

- Hea Yeon Lee
  - Northeastern university, department of mechanical and industrial engineering
  - Nano biosensor, nanotoxicology, drug discovery
  - e-mail: he.lee@neu.edu