Protein Patterning Based on Electrochemical Activation of Bioinactive Surfaces with Hydroquinone-Caged Biotin

Kyuwon Kim,† Haesik Yang,*‡ Sangyoung Jon,* Eunkyung Kim,* and Juhyoun Kwak†§

Korea Research Institute of Standards and Science, Daejeon 305-600, Korea, Department of Chemistry, Pusan National University, Pusan 609-735, Korea, Department of Life Science, Gwangju Institute of Science and Technology, Gwangju 500-712, Korea, and Department of Chemistry, Korea Advanced Institute of Science and Technology, Daejeon 305-701, Korea

Received July 7, 2004; E-mail: Juhyoun_Kwak@kaist.ac.kr (J.K.); hyang@pusan.ac.kr (H.Y.)

An efficient attachment and patterning method of proteins on surfaces is crucial to the construction of protein chips and bioelectronic devices. In the attachment step, the site-specific immobilization of proteins is required to retain their biological activity. As a platform satisfying the requirement, a biotin-modified surface has been widely used because either avidin or biotin-labeled proteins can be selectively immobilized in right orientation through avidin/biotin interaction with high affinity ($K_a$ $\approx$ $1 \times 10^{15}$ M$^{-1}$). On the other hand, the serial patterning of proteins is essential to the preparation of protein chips for multiple analysis. To prepare such patterns utilizing biotin/avidin interactions, two kinds of strategies have been developed. First is site-selective immobilization of biotin before protein attachment by using the soft lithography technique or scanning probe microscopy. Although these proved to be versatile protein patterning methods, several drawbacks restrict their application. For example, time consuming in the complex and serial patterning may lower the activity of previously attached proteins under ambient condition. Electrochemical approaches based on this strategy have been also reported. However, their reaction conditions are not compatible with the previously presented protein. They may suffer from ligand contamination bringing about cross adsorption of the protein. The second strategy is a site-selective bioactivation of initially deactivated biotin surface by photo irradiation. In the approach, a whole surface is precoated with deactivated biotin containing a photolabile protecting group, and then only selected sites are activated by the photo irradiation through the photo mask. When the surface is exposed to a solution of a target protein labeled with avidin, the protein is immobilized on the activated sites through avidin linkage. In some cases, however, longer photo irradiation to achieve complete deprotection can cause irreversible damage to the surface, which results in increased nonspecific adsorption of the protein. This should be overcome for the retention of protein activity as well as for higher contrast patterning.

Therefore, it is of great benefit to develop a fast, mild, and protein-friendly method for protein patterning. In this paper, we report a method for protein patterning based on an electrochemically active biotin derivative that generates a bioactive biotin surface by mild electrochemical perturbation. The electrochemical activation proceeds under the buffered aqueous environment at neutral pH. It also allows site-selective generation of bioactive biotin for the immobilization of the target protein by using prepatterned electrode arrays.

Scheme 1 displays the chemical structure of the electrochemically active biotin derivative (I) employed in this paper and a proposed mechanism for the generation of a bioactive biotin surface after electrochemical oxidation. Once the hydroquinone (HQ) moiety is oxidized, it is converted to a benzoquinonium cation. Subsequently, benzoquinone (BQ) and CO$_2$ are immediately released after nucleophilic acyl substitution by H$_2$O, and finally, the bioactive biotin surface is generated. To demonstrate the surface reaction proposed in Scheme 1, cyclic voltammetry (CV) was performed with the self-assembled monolayers (SAMs) of compound I, prepared by immersion of gold surfaces in 1 mM THF solution of I for at least 12 h. Figure 1A shows an irreversible anodic peak at 0.20 V on the first scan and a dramatic disappearance of the peak on the second cycle, no change in the charging current of the double-layer region, compared to that of the first cycle, indicates that the oxidation potential is mild enough to not destroy the remaining organic SAMs.

Figure 1B shows the grazing angle Fourier transform infrared (FT-IR) spectra of chemical environments on SAM surfaces, which support our view that the electrochemical bioactivation step is very

† Korea Research Institute of Standards and Science.
‡ Pusan National University.
§ Gwangju Institute of Science and Technology.
† Korea Advanced Institute of Science and Technology.
fast and complete. In the spectrum of I-based SAMs, we observed two strong bands at 1743 and 1796 cm$^{-1}$ corresponding to the C=O stretching of three esters and two urethane groups (a and b in Scheme 1), respectively. The peak at 1511 cm$^{-1}$ is characteristic of aromatic C=C stretching of the HQ moiety. The C–H stretching bands of the alkyl chain appear at 2929 and 2861 cm$^{-1}$. The spectrum of the monolayers after electrochemical reaction in a 0.1 M phosphate buffer solution showed distinct changes from original SAMs. While the peak at 1743 cm$^{-1}$ (C=O stretching in the ester moiety) revealed no significant change, the aromatic C=C stretching at 1511 cm$^{-1}$ disappeared, and the peak intensity at 1796 cm$^{-1}$ (C=O stretching in urethane) was reduced by half of the initial intensity. These results indicate that the HQ moiety was completely detached, and the C=O of “b” was also removed from the I-based SAMs. Moreover, the negligible changes in intensity, as well as the position of the C–H stretches, also imply that the other parts of I are kept intact. This implication also supports the view that the electrochemical reaction of HQ is mild.

To evaluate biological activity of the activated biotin surface after the electrochemical reaction, we tested a biospecific interaction with streptavidin (SA). Tri(ethylene glycol) ester of thiocetic acid (II) was used as a diluent of I-based SAMs not only to adjust proper density of surface biotin but also to prevent nonspecific adsorption of protein. Under the condition, we optimized the surface density of I to maximize SA binding. The density can be easily controlled by changing the mixing ratio of I and II in adsorbent solutions. The binding of streptavidin on mixed SAMs with various mole fractions of I in the solution ($\Gamma_I$) was monitored using SPR after the electrochemical oxidation (Figure 2A). When $\Gamma_I$ was 0.25 and the absolute surface density from CV was $5 \times 10^{-11}$ mol cm$^{-2}$, the maximum attachment of SA was observed (175 ng cm$^{-2}$, curve “a”). This density is consistent with that of the previous reports on the maximum binding condition of surface biotin. Curve “c” shows that nonspecific binding of SA was negligible in II-based SAMs (2 ng cm$^{-2}$). Curve “f” supports that these bindings are biospecific.

We constructed I-based SAMs on an interdigitated microelectrode array (IDA) for protein micropatterning. To visualize the pattern, we used poly-(horseradish peroxidase)-conjugated SA (pHRP-SA). Using SPR, we determined $\Gamma_I$ presenting maximum pattern contrast between deactivated and activated biotin surfaces, by comparing the amounts of pHRP-SA bound to the surfaces before and after electrochemical activation. Maximum contrast was observed at $\Gamma_I = 0.5$, and thus we employed this condition for micropatterning. Only one electrode of an IDA substrate was electrochemically treated at 0.23 V for 1 min. The resulting substrate was immersed in phosphate-buffered saline solution containing pHRP-SA. After pHRP-SA immobilization, the pattern was visualized by a color change that resulted from catalytic precipitation of 4-chloro-1-naphthol by HRP. Figure 2B shows an optical microscopic image of micropatterned pHRP-SA on IDA. Bright and dark regions correspond to the electrodes before and after electrochemical activation, respectively. The two electrodes highly contrast each other, which indicates that the present approach is very efficient for site-selective protein micropatterning.

In conclusion, we demonstrated that site-specific, as well as site-selective, protein patterning with high contrast could be achieved by means of an electrochemical bioactivation reaction of a rationally designed HQ-caged biotin surface under mild electrochemical conditions. Since the present patterning process, including surface activation and protein attachment, can be conducted under neutral buffer conditions in a short time period, it enables the serial patterning of multiple proteins while retaining the activity of previously attached proteins. Therefore, it might have applications for fabricating protein chips and biomolecular electronic devices.

Acknowledgment. J.K. gratefully acknowledges support from the National R&D project for Nano Science and Technology, as well as Brain Korea 21, MICROS, and IMT-2000 projects. H.Y. is grateful for Grant 02-PJ-3PG6-EV05-0001.

Supporting Information Available: Detailed synthetic procedure of compounds I and II. This material is available free of charge via the Internet at http://pubs.acs.org.

References

(9) All electrochemical treatments were conducted by holding a SAM-modified gold surface as a working electrode at 0.23 V for 1 min on the CV scan.
(11) BIACORE X SPR and SIA Kit Au surfaces were used for SA binding studies. The 1000 RU corresponds to an SA density of 1 ng mm$^{-2}$.
(12) The absolute surface density of I in the mixed monolayers can be easily obtained from integrating the oxidation peak of the HQ of I on the CV.
(13) Due to the nonlinear structure of thiocetic acid of compound I, $\Gamma_I$ presenting maximum binding of SA, is higher than the mole fraction of biotin/alkanethiols ($\sim 0.1$) of the linear methylene chain in other investigations. Nevertheless, the absolute surface density of biotin is similar to that expected in them. (a) Spanke, J.; Liley, M.; Schnitt, P. J.; Guder, H. J.; Angermueller, L.; Knoll, W. J. Chem. Phys. 1993, 99, 7012–7011. (b) Nelson, K. E.; Gamble, L.; Jung, L. S.; Boeckl, M. S.; Naemie, E.; Kolledge, S. L.; Sasaki, T.; Castner, D. G.; Campbell, C. T.; Stayton, P. S. Langmuir 2001, 17, 2807–2816.
(14) HRP mediates the catalytic oxidation of 4-chloro-1-naphthol by H$_2$O$_2$. Only the pHRP-SA bound region produces precipitates of the oxidized product in aqueous solution, which paints the region in dark color.

JA0459330