Bio-inert Properties of TEG Modified Dendrimer Interface

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The bioinert interfaces that prevent adhesion of proteins and cells are important for biomaterial applications. In order to design a bioinert interface, the immobilization of an appropriate functional group and the control of molecular density is required. Dendrimer was modified with triethylene glycol (TEG) to display a dense brush structure. TEG with different density and terminal groups were immobilized with a dendrimer template and thiol terminated molecules. The inhibitory effect on protein and bacteria binding was investigated. The physical property of the interface was measured by QCM-admittance to clarify the factor of the bioinert property.

Keywords Bioinert interface, dendrimer, triethylene glycol, QCM-A

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Introduction

The advancements in biotechnology are fueling demand for advanced biomaterials, for which control of material interfaces is important. Biomaterial interfaces are involved in many aspects of performance, including molecular recognition, cell adhesion, and inflammation. In particular, controlling the adhesion of bio-substances such as proteins and cells is important, and the design of bioinert materials has been intensively investigated for the development of biomaterials with biocompatibility for use in bio-devices and artificial organs.1-3 As a result of many studies, polymers such as polyethylene glycol (PEG)4,5 and zwitter ionic polymers6,7 have been reported.8 However, the current technology is still insufficient to control the interface properties as required.

It has been also reported that the structure of water at the material interface is important, and protein adsorption is changed by the molecular mobility of water.9 That is, it is significant to refine the design of interfaces with molecular level. The self-assembled monolayer10,11 and the polymer brush12 are considered to be excellent techniques for controlling the molecular structure of the interface.2 However, there are of course limitations to control the interface molecular structure. It is an interesting challenge to control the interface structure with the special molecules like dendrimer.

Dendrimers are uniform and branched polymers.12 The dendrimers have special structures that diverge from the center core to the molecular end. It has been reported that dendrimers with high generation have a dense molecular structure. We have reported the control of the interface of dendrimer immobilized material.13 The unique performance can be demonstrated by controlling the density and nano-structure with dendrimers.14 We have previously reported that a molecular interface with a sugar chain immobilized at the end is realized, and the adhesion of proteins is greatly different.15 In other words, even if the functional groups are the same, their functions differ greatly depending on the density and structure of the molecules immobilized on the interface.16 Previous studies have shown that the molecular density of sulfated sugar has a significant effect on protein amyloidosis.

In this study, we have investigated the effect of dense functional groups on the pseudo-brush with dendrimer template. Dense brush layer of triethyleneglycol (TEG) was prepared by TEG dendrimer conjugates. The dendrimer had a core of disulfide group that was reduced to thiols to form a self-assembled monolayer on the gold substrate. Proteins and bacteria were applied to densely packed TEG structures to study their adhesion. Both methoxy and hydroxyl TEG were prepared.

Experimental

Reagent

The following reagents were used without purification: chloroform, N,N-diisopropylethylamine (DIEA) pyridine, sodium borohydride (NaBH₄), sodium azide (NaN₃) toluene, (Kanto Pure Chemical, Tokyo Japan), N,N-Dimethylformamide (DMF), methanol, potassium thioacetate, succinic anhydride, 2,4,6-trinitrobenzenesulfonic acid sodium salt dihydrate (TNBS), (Fujifilm Wako, Tokyo, Japan), 1-[bis(dimethylamino)methylene]-1H-1,2,3-triazolo[4,5-b]pyridinium 3-oxide hexafluorophosphate (HATU), diethylene glycol 2-bromoethyl methyl ether, 2-[2-(2-chloroethyl)ethoxy]ethanol, octadecane thiol (ODT) (TCI, Tokyo Japan), albumin from bovine serum (BSA), fibrinogen, polyamideamine dendrimer with cysteamine core generation 4 (PAMAM, 10 wt.% in methanol), polystyrene (PS) (Sigma-Aldrich Co., St. Louis, MO), concanavalin A (ConA, J-Oil Mills, Tokyo, Japan), and green-fluorescence SYTO9 (SYTO9, Molecular Probes Inc., Eugene, OR).

Staphylococcus epidermidis (S. coccus, Strain JCM 2414) was provided by RIKEN, BRC. E.coli ORN 178 was provided from Prof. T. Ooya (Kobe University).
Preparation of TEG derivatives

TEG derivatives with hydroxyl group and methoxy group, and TEG modified dendrimer with cysteamine core were prepared (Fig. 1). The detail of the synthetic procedure is summarized in the Supporting Information.

Substrate preparation

Gold substrate (Kenis, Osaka, Japan) was cut into 1 × 1 cm. The substrate was sonicated in MilliQ water and methanol for each 10 min. The substrate was cleaned with UV/O₃ (UV253E, UV_MT, Tokyo, Japan) using NCHV-10V (Reflector 2, Harrick Scientific Products Inc., Pleasantville, NY). Water contact angle was measured with a DropMaster300 (Kyowa Interface Science Co., Ltd., Saitama, Japan) instrument, and the thicknesses of the SAMs were estimated using an ellipsometer (PZ2000, Royal Philips Electronics, Eindhoven, Netherlands). AFM was measured with D-AFM (D-3000, Bruker AXS K.K., Karlsruhe, Germany) using NCHV-10V probe (Bruker AXS K.K., Karlsruhe, Germany) and Au(111) mica (φ10 mm, PHASIS, Geneva, Switzerland)

Characterization

Proton nuclear resonance (¹H-NMR) spectra were recorded on a JEOL-ECX400 spectrometer (JEOL, Tokyo, Japan) using CDCl₃, MeOD, DMSO-d₄, or D₂O as a solvent. X-ray photoelectron spectroscopy (XPS) spectra were measured on AXIS-ultra (Shimadzu/Kratos, Kyoto, Japan), and peaks were analyzed with peakfit v4.12 (Systat Software, Inc., CA, USA). FTIR spectra were measured with FTIR spectrum 100 (Perkin-Elmer Inc., Waltham, MA) equipped with an RAS attachment (Reflector 2, Harrick Scientific Products Inc., Pleasantville, NY). Water contact angle was measured with a DropMaster300 (Kyowa Interface Science Co., Ltd., Saitama, Japan) instrument, and the thicknesses of the SAMs were estimated using an ellipsometer (PZ2000, Royal Philips Electronics, Eindhoven, Netherlands). AFM was measured with D-AFM (D-3000, Bruker AXS K.K., Karlsruhe, Germany) using NCHV-10V probe (Bruker AXS K.K., Karlsruhe, Germany) and Au(111) mica (φ10 mm, PHASIS, Geneva, Switzerland)

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Substrate preparation

Gold substrate (Kenis, Osaka, Japan) was cut into 1 × 1 cm. The substrate was sonicated in MilliQ water and methanol for each 10 min. The substrate was cleaned with UV/O₃ (UV253E, Filgen, Inc., Aichi, Japan) for 30 min. The substrate was cleaned with MilliQ water and methanol and dried under N₂.

D-TEG-OCH₃ and D-TEG-OH were dissolved in water to a concentration of 0.2 mM. NaBH₄ (2 mL) and 4 mL methanol were added to the dendrimer solution to obtain thiol-terminated dendrimer solution. Dendrimer solution was diluted with water to a concentration of 0.1 mM. The substrate was immersed in 0.1 mM dendrimer solution for 12 h and washed with MilliQ water and methanol. The substrate was dried under N₂.

ODT was dissolved in ethanol to a concentration of 0.1 mM. The substrate was immersed in the ODT solution for 12 h. The obtained TEG derivatives were immobilized on the gold substrate.

Bacteria adsorption on the substrate

E. coli (ORN178) and S. coccus were cultured in LB broth (Lennox) medium (Sigma-Aldrich), and nutrient medium (polypeptone 5 g/L, beef extract 3 g/L, and NaCl 5 g/L). A solution of E. coli and S. coccus solution was prepared with cell densities of 10⁸ and 10⁷ CFU/mL with 10 nM SYTO 9. The substrate was immersed into the bacteria solution for 2.5 h and washed with PBS 5 mL twice. The bacteria were observed with fluorescence microscope, and analyzed with ImageJ.

Admittance measurement of the substrate with QCM

The admittance of the substrate was measured with 27 MHz QCM (AFFINIX Qn pro, Initium Inc., Kanagawa, Japan). The admittance curve was converted into a conductance curve (conductance–frequency plot) based on an admittance analysis to obtain both the frequency change (ΔF) and energy dissipation change (ΔD). The frequency change in the air phase (ΔF(air)) and that in aqueous solution (PBS buffer) (ΔF(buffer)) can be measured to determine the amount of hydrodynamic water on the interface. The energy dissipation change in water (ΔD(buffer)) and the frequency change after drying (ΔF(air)) are used to evaluate the viscoelastic properties in the aqueous buffer solution.

Results and Discussion

Preparation of the modified substrate.

TEG modified dendrimers (D-TEG-OCH₃ and D-TEG-OH) and thiol-terminated TEG (L-TEG-OCH₃ and L-TEG-OH) were prepared (Fig. 1 and Supporting Information), and confirmed by ¹H-NMR. The terminal of dendrimer was converted to TEG, the modification ratio was calculated to be 75%, and 48 TEG units were introduced in one dendrimer.

The obtained TEG derivatives were immobilized on the gold substrate. The surface modification with derivatives were analyzed by XPS, and contact angle measurements (Fig. 2 and Fig. S10). TEG and dendrimer immobilized substrates showed the spectra in C(1s), O(1s), and N(1s). TEG modified dendrimer showed the peaks at C–C, C–H (284.9 eV), C–N (285.7 eV), C–O (286.5 eV), C=O (287.8 eV), TEG, C=C (285.8 eV), C=O (284.1 eV), and C-S (286 eV). The reference hydrophobic substrate of ODT self-assembled monolayer showed only the peak of C=C, C-H (286 eV). The amount of molecular bands was estimated by XPS peaks of C(1s) and decrease of Au(4f) (Fig. S10). The amount of the molecular immobilization of TEG derivatives were compared with ODT that forms densely packed self-assembled monolayers. The amounts of L-TEG-OH and L-TEG-OCH₃ were estimated to be about 25% of ODT-self-assembled monolayer (SAM) based on the C(1s) peak.

Protein adsorption

The protein adsorption on the surface was evaluated by QCM (AFFINIX Q4, Initium Inc., Kanagawa, Japan). Fibrinogen, and BSA and ConA was used in this experiment. The modified sensor cell was covered with PBS buffer (10 mM, pH 7.4, 134 mM NaCl, 2.68 mM KCl, 1.76 mM KH₂PO₄, 8.1 mM Na₂HPO₄) until the frequency reached a steady state. Each protein, at various concentrations, was injected into the sensor cell, and the frequency changes (ΔF) were recorded.

Preparation of the modified substrate.

TEG modified dendrimers (D-TEG-OCH₃ and D-TEG-OH) and thiol-terminated TEG (L-TEG-OCH₃ and L-TEG-OH) were prepared (Fig. 1 and Supporting Information). The modified sensor cell was covered with PBS buffer (10 mM, pH 7.4, 134 mM NaCl, 2.68 mM KCl, 1.76 mM KH₂PO₄, 8.1 mM Na₂HPO₄) until the frequency reached a steady state. Each protein, at various concentrations, was injected into the sensor cell, and the frequency changes (ΔF) were recorded.
and the decrease of Au(4f) peak. The amount of D-TEG-OH and D-TEG-OCH₃ was approximately the same as that of ODT-SAM.

Contact angles of the substrates of D-TEG-OCH₃, D-TEG-OH, L-TEG-OCH₃, and L-TEG-OH were 37.3, 30.9, 58.8 and 36.0, respectively. The TEG modified substrates showed hydrophilicity, and the TEG-dendrimer substrate showed stronger hydrophilicity than monomeric TEG due to the dense hydrophilic group. The TEG immobilization was also confirmed by AFM and FTIR (Figs. S12 and S13). AFM analysis indicates the immobilization of the spherical dendrimer on the substrate. The thickness of the dendrimer layer is 0.5 - 1.0 nm, showing the dendrimer monolayer formation on the substrate. In spite of the bulky structure of dendrimer, the thiol-terminated dendrimer formed a self-assembled monolayer on the substrate based on the strong Au-S interaction.

**Protein adsorption to the TEG modified substrate**

The amount of protein adsorption was varied with the substrates using fibrinogen, and in the order of PSi, L-TEG-OCH₃, > ODT > L-TEG-OH >> D-TEG-OCH₃ > D-TEG-OH (Fig. 3). The amount of protein bound on L-TEG-OCH₃ is about 8 times larger than that on D-TEG-OCH₃, and the amount of protein bound on L-TEG-OH is about 7 times larger than that on D-TEG-OH at C = 0.01 g/L. The protein bound on D-TEG-OCH₃ and L-TEG-OCH₃ is larger than that on D-TEG-OH and L-TEG-OH, respectively. The large amount of protein was bound to the hydrophobic surface, but the amount of protein bound was not determined only by the hydrophobicity. PSi and L-TEG-OCH₃ showed larger protein adsorption than the ODT-substrate.

Inhibition of protein adsorption depended on the TEG display. TEG-dendrimer conjugate could display the densely packed TEG brush on the substrate and it showed a stronger inhibitory effect on protein adsorption than linear TEG. Though the PAMAM dendrimer contains an amine group, the electrostatic interaction by the amine group inside dendrimer was not affected by the protein binding. The dense-TEG structure was effective in terms of the protein inhibitory effect, which was consistent with previous results. Interestingly L-TEG-OCH₃ modified substrate and PSi showed a similar affinity to protein adsorption. The hydrophobicity of the substrate was favorable to protein adsorption.
adsorption.

In protein adsorption, the TEG-dendrimer showed a better inhibitory effect than the linear. This inhibitory effect of densely packed TEG has been suggested from a report by the Whitesides group, where the SAMs with oligoethylene glycol terminal have a remarkable ability to inhibit protein adsorption. In this study, however, the ability to inhibit protein adsorption was not observed in L-TEG-OH and L-TEG-OCH due to non-dense TEG and hydrophobicity. It is suggested that dendrimer-templated TEG is not a long-chain polymer such as polyethylene glycol, but exhibits an excellent protein inhibitory effect for dense TEG. In view of protein adsorption, dendrimer-TEG substrate was bioinert.

**Bacteria adsorption to the substrate**

Bacteria adsorption on the substrate was measured with various substrates (Fig. 4). In the experiment with *E. coli*, the *E. coli* adsorption on PSi was suppressed to 12% or less on the TEG immobilized substrates (D-TEG-OCH, L-TEG-OCH, D-TEG-OH and L-TEG-OH). In particular, the amount of *E. coli* on L-TEG-OH was 0.002% of that on PSi, indicating resistance of the substrate to *E. coli*. When comparing the dendrimer (D-TEG-OCH, and D-TEG-OH) and the linear (L-TEG-OCH; and L-TEG-OH) with the same terminal group, the amount of adsorption on the dendrimer-TEG was larger than that on linear TEG at the methoxy terminal and the OH terminal, respectively. In the experiment with *S. coccus*, the *S. coccus* adsorption on TEG immobilized substrate was less than half of that on the PSi substrate. In particular, L-TEG-OCH showed almost no adsorption. When comparing the dendrimer-TEG and linear TEG immobilized substrate with the same terminal groups, the amount of adsorption on the dendrimer TEG was 72.6 times and 3.0 times larger than linear TEG at the methoxy terminal and the OH terminal, respectively.

The results of PSi and ODT indicated that the interface with large amounts of protein adsorption was also high for bacteria. However, the adsorption behavior TEG interface of dendrimers and linear showed no correlation between protein and bacteria adsorption. Linear TEG (L-TEG-OCH and L-TEG-OH) showed a strong inhibitory effect on bacteria adsorption, though the linear TEG substrate did not inhibit protein adsorption. In addition, in protein adsorption, the effect of the functional group at the molecular terminal was large, and the methyl ester and hydroxyl group at the terminal had a large effect, but in bacterial adsorption, the effect of the functional group at the terminal was small, and the shape (linear or dendrimer) of the molecules had the most effect. In view of bacteria adsorption, the linear TEG substrate is bioinert.

**Viscoelasticity of the substrates**

The viscoelastic properties of the interface with immobilized TEG derivatives were evaluated using QCM-A. QCM is used as a sensitive mass sensor, utilizing the Saureybre relationship between the resonant frequency and the mass per unit area. The mass changes can be measured not only in vacuum but also in the aqueous solution as a label-free biosensor. Since the mass is measured by the difference in resonant frequency of the device, the energy dissipation by viscosity of the device is reflected in the QCM. The rheological change due to the hydrodynamic water in the aqueous solution is well affected. It has been reported that the viscoelasticity at the device interface is related to the energy dissipation of the QCM.

The results are shown in Fig. 5. The viscoelasticity of the interface was evaluated by comparing the energy dissipation value (ΔD/PBS) divided by the polymer fixed amount (ΔD/(air)). The result of energy dissipation per unit mass shows a correlation between protein and bacteria adsorption. Linear TEG (L-TEG-OCH and L-TEG-OH) showed a strong inhibitory effect on bacteria adsorption, though the linear TEG substrate did not inhibit protein adsorption.

The value of the dendrimer was 2.5 times higher than that of the linear. In the present measurement, since the higher the value of ΔD/(air), the more the viscoelastic, it was found that the dendrimer TEG substrate was more elastic than the linear TEG substrate, and that the linear TEG substrate was more viscostic than the dendrimer TEG substrate.
Okahata et al. suggested that the current dendrimer interface value had similar viscoelasticity to those of proteins.²⁴

Figures 4 and 5 suggest that there is a correlation between viscoelasticity and the amount of bacteria bound, even though the molecular thickness is only several nanometers (Fig. S13). It is considered that bacteria are more likely to adsorb to the elastic substrate and less likely to adsorb to the viscous substrate. It has been reported that not only the inhibition effect of proteins at the interface but also the physical properties of the interface have a great influence on the adsorption of bacteria. It has been reported that viscoelasticity is strongly related to the adsorption of bacteria.²⁵ Several cells and bacteria were reported to have mechanical property sensitive proteins.²⁶ In this study, the terminal functional groups, the contact angle, and the amount of protein bound are not appropriate for the amount of bacterial adhesion, suggesting that the difference in the physicochemical properties of the interface is involved. However, this mechanism needs further investigation.

Conclusion

Dendrimer terminal was modified with triethylene glycol (TEG) to form dendrimer-TEG conjugates. Densely packed TEG was immobilized on the gold substrate as the dendrimer conjugate. The protein and bacteria adsorption on the TEG modified substrates were investigated. It was found that the dendrimer-and the linear- TEG immobilized substrate had an inhibitory effect of the adsorption of proteins and bacteria. However, when the dendrimer- and the linear-TEG were compared, it was found that the dendrimer had a higher inhibitory effect on protein adsorption but a lower inhibitory effect on bacterial adsorption. It was suggested that the physical property of dendrimer TEG interface influenced the adsorption of bacteria.

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Supporting Information

This material is available free of charge on the Web at http://www.jsac.or.jp/analsci/.

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