Nano-visualization of oriented-immobilized IgGs on immunosensors by high-speed atomic force microscopy

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Oriented immobilization of sensing molecules on solid phases is an important issue in biosensing. In case of immunosensors, it is essential to scrutinize not only the direction and shape of immunoglobulin G (IgG) in solution but also the real-time movement of IgGs, which cannot be achieved by conventional techniques. Recently, we developed bio-nanocapsules (BNCs) displaying a tandem form of the IgG Fc-binding Z domain derived from Staphylococcus aureus protein A (ZZ-BNC) to enhance the sensitivity and antigen-binding capacity of IgG via oriented-immobilization. Here, we used high-speed atomic force microscopy (HS-AFM) to reveal the fine surface structure of ZZ-BNC and observe the movement of mouse IgG3 molecules tethered onto ZZ-BNC in solution. ZZ-BNC was shown to act as a scaffold for oriented immobilization of IgG, enabling its Fv regions to undergo rotational Brownian motion. Thus, HS-AFM could decipher real-time movement of sensing molecules on biosensors at the single molecule level.

Due to the high specificity and affinity of biological molecules (e.g. immunoglobulin G (IgG), ligands, receptors, aptamers, sugar chains, lectins), biosensors are expected to be a promising technology for sensing various biological materials. The sensor surface (e.g., glass or metal) is commonly functionalized with chemical crosslinking or self-assembled monolayers (SAMs). Especially, long-chain (number of methylene groups n > 10) alkanethiols assemble in a crystalline-like way and can introduce uniform functional groups onto gold surface. While biological molecules were subsequently immobilized on the sensor surface, the control over the orientation of these molecules (i.e., oriented-immobilization of IgGs) has not so far been fully achieved. In case of immunosensors, the crosslinking molecules should neither increase steric hindrance around the antigen-recognition Fv regions nor reduce the antigen-recognition activity of IgGs and, furthermore, should align the Fv regions for efficient antigen-recognition. Specific sites within IgG molecules have been used to achieve oriented immobilization, such as the Fc region via an Fc-binding protein A or G, aldehyde groups introduced into the carbohydrate moiety of the CH2 domain via an hydrazide-containing crosslinker, and the thiol group of monovalent Fab’ fragments via a thiol-containing solid phase. However, the orientation of protein A or G itself cannot be fully controlled on the solid phase. Chemical and enzymatic treatments may affect the antigen-recognition activity of IgGs. These situations led us to develop rigid and self-assembled scaffolds for aligning IgGs at the nanoscale level, without modification.

We previously developed ZZ-BNCs of ~30 nm diameter by expressing the gene encoding hepatitis B virus (HBV) surface antigen (HBsAg) L protein with a tandem form of Fc-binding Z domain (Fig. 1a) in yeast. ZZ-BNC contains about 120 molecules of ZZ-L protein (N-terminally ZZ-fused L protein) embedded in a liposome and has ability for capturing ~60 mouse IgG molecules, as well as displaying all the IgG Fv regions outwardly. Furthermore, ZZ-BNCs can enhance the sensitivity of immunosensors and immunoassays not only through the oriented immobilization of antibodies but also the clustering of antibodies and labelling molecules. Thus, ZZ-BNC is a promising scaffold for a variety of conventional immunosensors and immunoassays.

To evaluate immunosensors, it is necessary to elucidate not only the direction and shape of IgGs on the solid phase in solution but also the real-time movement of IgGs. However, conventional techniques cannot fully analyze the real-time movement of IgGs. Particularly, it is difficult for AFM to distinguish IgGs from surrounding molecules using these static observations (snapshots). Recently, HS-AFM (high-speed atomic force microscopy) equipped with a highly sensitive, ultra-fast cantilever and efficient AFM scanning capabilities, have

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calculated to be 21,300 nm$^3$ and 5,170 nm$^2$, respectively, corre-
cand c). The volume and surface area of an average ZZ-BNC were
between CH$_2$ and CH$_3$ of the Fc subunit$^{32}$. (b) (upper), (c) HS-AFM images
in a lipid bilayer. The dimeric forms of ZZ-L proteins are described in this
BNC. One ZZ-BNC particle consists of about 120 ZZ-L proteins embedded
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(10 particles from 48 frames) revealed protrusions on the surface
in diameter; 17.6 $\pm$ 1.0 nm in height (mean $\pm$ SD, n = 19)) (Fig. 1b
and c). The volume and surface area of an average ZZ-BNC were
calculated to be 21,300 nm$^3$ and 5,170 nm$^2$, respectively, corre-
sponding to those of a spherical $\sim$40.6-nm particle. Because this
guess agreed well with the diameter obtained by dynamic light
scattering (45.4 $\pm$ 2.2 nm), ZZ-BNCs were considered to adsorb
onto mica surface without disrupting their capsule structure. Semi-
automated, single-particle reconstructions using the EMAN software$^{16}$
(10 particles from 48 frames) revealed protrusions on the surface
of ZZ-BNC (Fig. 1d). The number and area of protrusions were
estimated by ImageJ software to be 27 $\pm$ 3 and 28.9 $\pm$ 10.1 nm$^2$,
respectively. The diameter was 6.1 $\pm$ 3.6 nm. The distance between
adjacent protrusions (centre to centre) was 8.2 $\pm$ 1.7 nm, which
agrees well with the space ($\sim$6 nm) between two HBsAg proteins
on the HBV virion$^{17}$. One ZZ-BNC was estimated to possess 54
protrusions, while reported to contain about 120 molecules of ZZ-L
protein$^2$. Meanwhile, the HBsAg protein of the HBV virion$^{18}$, as
well as the HBsAg L protein of BNC$^5$, is a transmembrane protein
containing three membrane-spanning domains as dimeric$^{20}$ and
multimeric$^6$ forms. ZZ-BNCs were then reduced with 0–1,000 mM
2-mercaptoethanol (2ME), separated by SDS-PAGE, and stained with
silver (Fig. 1e). By increasing the concentration of 2ME, two major
bands of 96 and $>$260 kDa were gradually shifted to 96 and 48 kDa,
and finally converged to 48 kDa, strongly suggesting that nascent ZZ-
L proteins form a dimer by intramolecular disulfide bonds which then
form multimers by intermolecular disulfide bonds, as previously
proposed for recombinant mammalian cell-derived HBsAg L protein$^{20}$. Taken together, each protrusion on ZZ-BNCs may be a
ZZ-L dimer containing six membrane-spanning domains, which is
 corroborated by the fact that the diameter of the protrusions (6.1 $\pm$
3.6 nm) is consistent with that of seven membrane-spanning
rhodopsin ($\sim$4.3 nm)$^{21}$. 

Observation of mouse IgG3 in solution. We next observed mouse
IgG3 molecules in solution by HS-AFM (Fig. 2a). Conventional AFM
observation of IgG molecules on a mica surface in air have shown that
the hinge region of IgG is flexible, allowing two Fab subunits to rotate
about their axes to present the side to the mica surface, with
deposition of the Fc subunit on the opposite side in two forms
(extended and folded)$^{22}$. Assuming that the vertical length of Fab
subunits is consistent amongst IgG forms, each IgG was identified
as a Y-shaped structure including three subunits (two Fab and one
Fc) in extended or folded form. By measuring the size of the extended
form of IgG (n = 3), the distance between the two Fab subunits was
calculated at 26.8 $\pm$ 2.4 nm, and those between the Fc subunit and
each Fab subunit at 32.5 $\pm$ 5.7 nm and 31.0 $\pm$ 6.3 nm, respectively.
The width of the Fc subunit was 8.9 $\pm$ 1.6 nm. The greatest height
of mouse IgG3 was 3.7 $\pm$ 0.5 nm and the centre of mass was estimated
to be located at 19.1 $\pm$ 4.0 nm from the tip of the Fc subunit. These
sizes of mouse IgG3 molecules are consistent with those of various
IgG measurements obtained by conventional AFM observations$^{23}$. We
previously reported that one ZZ-BNC can capture a maximum of
60 IgG molecules$^6$. When the surface of ZZ-BNC (5,170 nm$^2$; radius,
20.3 nm) was filled with Fc subunits (base area, 32.8 $\pm$ 6.0 nm$^2$;
height, 19.1 $\pm$ 4.0 nm) in close-packed arrangement, ZZ-BNC
could be surrounded by 161 $\pm$ 29 molecules of Fc subunits. On the
other hand, ZZ-BNC fully covered with Fc subunits (surface area,
19,500 nm$^2$; radius, 39.4 nm) could be surrounded by 197 $\pm$ 18
molecules of F(ab')$_2$ subunits (base area, 99.3 $\pm$ 8.8 nm$^2$; height,
9.6 $\pm$ 2.0 nm), suggesting that Fc subunits on ZZ-BNC contact
one another intermolecularly, rather than F(ab')$_2$ subunits. Consequently, each IgG could be tethered onto the surface of ZZ-
BNC at appropriate intervals, thereby improving antigen recognition
by the Fv region.

Real-time movement of mouse IgG3 on ZZ-BNC in solution. For
developing the next generation of immunosensors, it is a prerequisite
to analyze and optimize the real-time movement of immobilized
IgGs in solution at the single molecular level. ZZ-BNCs adsorbed
onto gold surfaces were immersed into PBS (0 s), contacted with
mouse IgG3 (55 s, yellow arrows in Figs. 2b and d), and then subjected
to time-lapse imaging with HS-AFM (0.2 frames per second (fps), total 1,300 s, 260 frames (Fig. 2b and Supplementary
Movie 1). After 45 s from the addition of IgG (100 s from the start),
some of the ZZ-BNCs exhibited a rough surface. After 95 s from the
addition of IgG (150 s from the start), the height of three ZZ-BNCs
Figure 2 | HS-AFM analyses of mouse IgG3 and ZZ-BNC in solution.
(a) HS-AFM images of mouse IgG3 molecules. Bar, 20 nm. (b) Video image of the movement of mouse IgG3 on ZZ-BNC (0.2 fps). Yellow arrow, the addition of mouse IgG3. Red arrow, first appearance of attachment. Times (s) after the start of observation are indicated in the left margin. Bar, 20 nm. (c) HS-AFM images of ZZ-BNCs on a gold surface without (10 s, left panel)/with (150 s, right panel) mouse IgG3. ZZ-BNC used for (b) is indicated by a white box. The locations where mouse IgG3 appeared are indicated by red arrowheads. Bars, 20 nm. (d) Binarized video image of the movement of mouse IgG3 on ZZ-BNC (0.2 fps). Yellow arrow, the addition of mouse IgG3. Red arrow, first appearance of mouse IgG3. Times (s) after the start of observation are indicated in the left margin. Bar, 20 nm.

Analyses of the movement of mouse IgG3 on ZZ-BNC. The positions of the centre of mass of attachments were calculated from their projected areas using ImageJ software and were found to exhibit a two-dimensional Gaussian distribution (major axis, 25.6 nm; minor axis, 20.4 nm; Fig. 3a). Because the length between the centre of mass of mouse IgG3 and the tip of the Fc subunit was estimated to be 19.1 ± 4.0 nm (see above), the deflection angle of mouse IgG3 on ZZ-BNC was estimated to be 88.1 ± 20.5 degrees. These results demonstrate that mouse IgG3s were tightly bound onto ZZ-BNC through the Fc subunit with Fab subunits swinging outwardly (Fig. 3b). The average speed of movement of mouse IgG3 was estimated as 4.6 ± 3.0 nm (13.8 ± 9.0 degrees) per 5 s (Fig. 3c). The speed fluctuated from 0.24 to 15.6 nm (0.74 ± 0.14 to 49.7 ± 10.1 degrees) per 5 s (Fig. 3d). It has been reported that the diffusion coefficient of monomeric human IgG is $4 \times 10^{-7}$ cm$^2$ s$^{-1}$, implying that one IgG molecule can move spontaneously at 20 nm per 5 second$^{24}$. These results indicate that IgG molecules on ZZ-BNC undergo rotational Brownian motion. ZZ-BNC acts as a scaffold for oriented-immobilization of IgGs, being expected to enhance the sensitivity and specificity of a wide range of immunosensors and immunoassays.

Discussion
The surface structure of BNC (recombinant yeast-derived HBsAg L protein particle) has been analyzed by conventional transmission electron microscopy (TEM) and AFM in air$^{8,9,25}$. While rugged surface was suggested by TEM observation, no protrusion has so far been identified. Recently, the surface structure of native HBV virion has been analyzed by electron cryomicroscopy$^{17}$. The envelope of HBV virion was decorated with 160–200 surface protrusions that are spaced ~6 nm apart. In this study, HS-AFM observation revealed for the first time that ZZ-BNC (a derivative of BNC) in solution displays 54 surface protrusions that are spaced 8.2 ± 1.7 nm apart. The small differences in space may be attributed to either the conditions for observation (air versus solution) or the mass of displayed molecules (~14-kDa pre-S region for HBV virion versus ~20-kDa ZZ domains for ZZ-BNC). Furthermore, during the biosynthesis of HBsAg particle in recombinant mammalian cells, nascent HBsAgs were shown to translocate across endoplasmic reticulum (ER) membrane along with the formation of intramolecular disulfide bonds, dimerize by forming intermolecular disulfide bonds, and then spontaneously assemble into particle structure without forming multimers$^{25}$. But, while recombinant yeast-derived HBsAg L proteins (including HBsAg S and M proteins) form multimers by intermolecular disulfide bonds, it has not been determined if they form dimers or not prior to the formation of BNC. In this study, nascent ZZ-L proteins (a derivative of HBsAg L protein) translocated across yeast ER membrane were strongly suggested to dimerize firstly, assemble to particle structures, and then form multimers by intermolecular disulfide bonds. Therefore, it was considered that more oxidative conditions of protein biosynthesis in yeast cells rather than mammalian cells$^{27}$ might contribute to the formation of multimeric HBsAg proteins in yeast cells$^{8}$.

In the development of biosensors, it is indispensable to decipher the direction, shape, and movement of sensing molecules on solid phase for improving sensitivity, specificity, and analyte-binding capacity. Conventionally, these molecules have been analyzed by electron microscopy (EM) and AFM$^{22,24}$. While both methods allowed us to observe IgGs at single molecule level, the former could be operable only in air. However, they could neither achieve real-time observation of IgGs in solution nor distinguish Fc from Fab clearly. Exceptionally, Garcia et al. have succeeded in identifying the position of the Fc and Fab fragments in air by tapping-mode AFM$^{2,23}$. The technical defects of these methods have forced us to evaluate the sensing molecules on solid phase by indirect methods (i.e., functional analyses). For example, in case of immunosensors, the degree of
oriented immobilization of IgGs has been evaluated by the reactivity to antigens. These indirect methods are indeed suitable for the optimization of biosensors, but it is equivocal to judge if sensing molecules exert their ability as much as possible. In this study, we could directly observe real-time movement of mouse IgG3s tethered onto ZZ-BNCs in solution by HS-AFM. It was therefore confirmed that ZZ-BNCs are stable to heat, chemical, and mechanical stresses, ZZ-BNCs would be an ideal and practical scaffold for oriented immobilization of IgGs.

When observing the surface of biosensors by EM and AFM, it is essentially difficult to identify what molecules are adsorbed onto solid phase. For the molecular identification in EM, samples are essentially difficult to identify what molecules are adsorbed onto solid phase. For the molecular identification in EM, samples are usually pre-treated with labelling reagents (e.g., gold particle-labelled antibodies). Worse yet, since conventional AFM can detect the ruggedness of sample surface only, it is nearly impossible to identify the molecules on the surface of mixed samples. On the contrary, HS-AFM can identify the molecules on the surface of solid phase by their movements without any modification. Taken together, HS-AFM might be worthwhile for the refinement of other biosensing molecules (e.g., ligands, receptors, aptamers, sugar chains, lectins) in various biosensors at the single molecule level.

Methods

ZZ-BNCs. ZZ-BNCs were overexpressed in Saccharomyces cerevisiae AH22R cells carrying the ZZ-BNC-expression plasmid pGILD-ZZ50. Following a previously described purification protocol for BNC, ZZ-BNCs were extracted from yeast cells by disruption with glass beads and purified using an AKTAliquid chromatography system (GE Healthcare, Amersham, UK) by affinity chromatography on porcine IgG, followed by gel filtration. The ZZ-BNCs were analyzed by 12.5% SDS-PAGE, followed by silver staining (Wako Chemicals, Osaka, Japan).

High-speed AFM. Topology images of ZZ-BNCs and mouse IgG3s (Sigma-Aldrich, St Louis, MO, USA) were obtained on a high-speed AFM system (Nano Live Vision, RIBM, Tokyo, Japan) using a silicon nitride cantilever (BL-ACI0EGS, Olympus, Tokyo, Japan). To obtain static images, either ZZ-BNCs (50 μg ml⁻¹, 2 μl) or mouse IgG3s (1 μg ml⁻¹, 2 μl) were adsorbed on a mica surface (Ted Pella Inc., Redding, CA, USA), incubated for 10 min, washed with phosphate-buffered saline (PBS) three times, and then subjected to time-lapse imaging. To observe the real-time movement of IgGs on ZZ-BNC in solution, ZZ-BNCs (50 μg ml⁻¹, 2 μl) were adsorbed onto an atomically flat gold surface (Auro sheet (111) HS; Tanaka Kikinzoku Kogyo K.K., Tokyo, Japan), incubated for 10 min, washed with PBS three times, subjected to time-lapse imaging (0 s), and then mixed with mouse IgG3 (50 μg ml⁻¹, 4 μl) (53 s). Images (192 × 144 pixels) were obtained at a scan rate of 0.2 frames per second (fps) for 1,300 s. Images were analyzed by ImageJ 1.44o software (http://rsbweb.nih.gov/ij/) and PaintShop Pro software (COREL Corporation, Ontario, Canada). Semi-automated, single-particle reconstructions were performed using the EMAN software package, v1.9. Ten particles were selected from 48 frames of the HS-AFM movies (total 761 frames) using the boxer program in the EMAN package.

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