

Covalent anchoring of proteins onto gold-directed NHS-terminated self-assembled monolayers in aqueous buffers: SFM images of clathrin cages and triskelia

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Abstract *N*-Hydroxysuccinimide-terminated self-assembled monolayers with linear (CH₂)₁₀ chains were prepared on ultraflat Au(111) surfaces from dithiobis(succinimidylundecanoate). These monolayers, which are covalently chemisorbed to gold via thiolate bonds, form a highly reactive amino-group specific carpet at the liquid–solid interface. Proteins bind to it covalently in aqueous buffers under mild conditions; this provides a (general) procedure for protein immobilization for scanning probe microscopy. Using this technique, we have obtained what we believe are the first scanning force microscopy images of clathrin cages and of their in situ disassembly, yielding typical triskelia under non-denaturing conditions.

Key words: Atomic force microscopy; Scanning force microscopy; Gold; Self-assembled monolayer; *N*-Hydroxysuccinimide; Clathrin; Triskelion

1. Introduction

Scanning probe microscopy (SPM) potentially allows direct imaging of individual macromolecules under 'physiological' conditions, i.e. undenatured, in aqueous buffers, and also in the presence of appropriate ligands or effectors; for reviews see [1–5]. Even if this has to be paid for with a lower resolution, as compared to that of high-resolution electron microscopy (and of course also to that of SPM of many inorganic and organic materials), SPM should in principle allow structure–function studies to be carried out on native biological objects, particularly when major structural changes are involved.

One of the conditions to be fulfilled for SPM imaging of macromolecules in aqueous solutions is that they should be firmly anchored on appropriate substrates and thus not be displaced by the tip while scanning. Such substrates should be flat, i.e. featureless over large areas, be chemically inert, yet allow fixation of the objects. Mere electrostatic interactions, for example, would severely limit the conditions under which biological objects can be eventually investigated (e.g. only at low ionic strength or within a limited pH range). A variety of crystalline or amorphous substrates have been used to date, having different chemi- or physisorption properties. The substrates mostly used are mica, glass, silicon wafers, highly oriented pyrolytic graphite (HOPG), and gold.

Mica has excellent flatness. It binds biomolecules via electrostatic interactions (thus with the limitations indicated above), also after appropriate derivatization [6–10]. Glass can be derivatized to make it bind proteins covalently via silanization

[11,12]; it has the additional advantage of allowing imaging of the very same sample by both SPM and other techniques requiring a transparent support. Also HOPG (if used underivatized, but see [13]), binds macromolecules by way of weak electrostatic or 'adsorption' forces (and, moreover, it has fallen into, perhaps excessive, disrepute when it was shown to yield artifactual images mimicking DNA [14,15]). Gold is, we think, an even more promising substrate: it is inert against O₂, yet it can form very stable covalent Au–thiolate bonds [16–19]. Additional advantages of gold are related to its conductivity: it allows STM imaging and also potentiostatic deposition of charged macromolecules. We have recently worked out a simple and reproducible procedure to prepare ultraflat Au(111) surfaces (template-stripped gold, TSG) with a mean roughness as small as 2–5 Å over more than 25 μm² [20,21], thus overcoming the serious drawbacks of the irregular topography (on μm scale) of epitaxially grown or of other gold surfaces, as prepared by established procedures (for a comparative review until 1992, see [22]).

Some biomolecules may bind to these gold surfaces without additional treatment (e.g. via pre-existing thiol groups, or physisorption), but the two main all-purpose routes are: (i) introducing (extra) thiol groups into the biomolecule (e.g. DNA [23], see also [24]), proteins (via Traut's reagent: [25] and unpublished data from our group), phospholipids [26]), or (ii) formation of derivatised gold-directed self-assembled monolayers (SAM) onto these ultraflat Au(111) surfaces with subsequent non-covalent [27] or covalent anchoring of biological macromolecules (see below).

Here we deal with the latter approach. Hydrophobic thiols (e.g. dodecanethiol) easily form regular monolayers on epitaxially grown gold surfaces (e.g. [16–19]) with a commensurate ($\sqrt{3} \times \sqrt{3}$) R30° overlayer structure. Au(111) directs the thiolates to form two-dimensional arrays, which are further stabilized by their lateral hydrophobic interactions. The ultraflat Au(111) surfaces mentioned above [20] are exquisitely flat templates onto which to build very regular, extended monolayers, for

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Abbreviations: SPM, scanning probe microscopy; SFM, scanning force microscopy; STM, scanning tunneling microscopy; SAM, self-assembled monolayer; NHS, *N*-Hydroxysuccinimide; TSG, template-stripped gold; DSU, dithiobis(succinimidylundecanoate); HOPG, highly oriented pyrolytic graphite.

example of palmitoyl-*N*-cysteamine [21], or of ω -functionalized thiols. If the ω -substituent is a highly reactive group, it can act as the anchoring site, docking proteins onto the monolayer carpet. We have used here dithiobis(succinimidylundecanoate) (DSU). It readily forms regular and extended monolayers; the *N*-hydroxysuccinimidyl groups, which are exposed at the monolayer-water interface, react with amino groups under very mild conditions (pH ~6.5–7.5).

In this paper we focus on clathrin and its association states. This protein forms the regular polyhedral surface lattice of clathrin-coated vesicles, which are involved in intracellular protein transport. This function involves the reversible association of three-legged clathrin protomers (triskelia), composed of three heavy and three light chains, into regular coat structures. In vitro purified clathrin triskelia associate in the absence of membranes into empty cages which are very similar to the surface lattice of coated vesicles. Both triskelia and cages have been extensively characterized by electron microscopy [28,29].

This system was chosen because (i) cages and triskelia differ considerably in size and shape, and would thus be easily distinguished by SFM, (ii) clathrin is readily purified and the two association states are interconvertible by simple and established manipulations [30], thus enabling in situ studies of dynamic events in the nanometer range, (iii) clathrin contains enough lysine residues for immobilization, and (iv) our understanding of the dynamics of clathrin assembly and disassembly is likely to improve by using SFM.

2. Materials and methods

2.1. Materials and instrumentation

Clathrin coated vesicles were prepared from bovine brains according to [31], and clathrin (cages) were purified according to [30]. All chemicals and solvents were commercial grades of highest purity. Gold, mica, Si-wafer and epoxy-glue Epo-tek No. 377 were purchased as reported previously [20]. Synthesis of dithiobis(succinimidylundecanoate) was carried out via oxidation of the Bunte salt of 11-bromoundecanoic acid (Wagner et al., in preparation).

Scanning force microscopy was carried out on a NanoScope III from Digital Instruments Inc. (Santa Barbara, CA, equipped with an E-scanner with a $10 \times 10 \mu\text{m}$ scan range), and scanning tunneling microscopy on a home-built system. For SFM we used microfabricated monocrystalline silicon tips (with force constants ranging from 0.06 to 0.17 N/m, purchased from LOT, Darmstadt, Germany), and for STM mechanically cut Pt/Ir tips.

2.2. Preparation of template-stripped gold surfaces (TSG)

According to our procedure described in [20], gold was deposited 200 nm thick onto freshly cleaved, preheated (300°C) ruby muscovite mica sheets. After glueing onto a Si-wafer piece using the epoxy glue, the mica was removed by immersing the [Si-epoxy glue-gold-mica] multilayer into tetrahydrofuran. This resulted in exposure of the ultraflat Au(111) surfaces (TSG).

2.3. Formation of the NHS-SAM

The monolayer was prepared on these ultraflat Au surfaces by immersing the TSG in a 1 mM solution of DSU in acetone for 2 h at room temperature. After rinsing with acetone, the NHS-terminated monolayer was dried under a stream of nitrogen and immediately used for the immobilization step.

2.4. Protein immobilization and SFM imaging

The general procedure for immobilizing amino groups containing biomolecules was as follows: 50 μl of the protein solution (amine free buffers are mandatory) at a concentration of 1 $\mu\text{g}/\text{ml}$ were placed on a piece of parafilm; a TSG platelet was carefully put upside down onto the drop. After 1 h the platelet, now carrying the covalently immobi-

lized biomolecules, was rinsed with 2 ml buffer solution and placed into the fluid cell of the NanoScope. The buffer used here in the immobilization step and in the first imaging (see Fig. 3A below) was 100 mM MES (pH 6.5), 1 mM EGTA, 0.5 mM MgCl_2 and 0.05% NaN_3 . Subsequent injection of 200 μl of 0.5 M Tris-HCl (pH 7) into the fluid cell nearly completely exchanged the buffer, which triggered the in situ disassembly of the immobilized cages.

Image acquisition in the constant-force imaging mode was carried out in the buffers indicated. The proteins were never left to dry at any stage. Forces were in the range of approximately 1 nN and scan speeds were lower than 1 $\mu\text{m}/\text{s}$. The piezo scanners were calibrated in the *x* and *y* dimensions, by using a Si calibration standard with a periodicity of 200 nm, and in the *z*-dimension from the known monoatomic step heights of Au(111). All images shown in the following are based on unfiltered data.

3. Results and discussion

Ultraflat gold substrates were prepared and characterized as reported previously in detail [20,21]. NHS-SAM spontaneously formed onto them from DSU by self-assembly in a rather straight-forward manner (see section 2). In Fig. 1 the resulting complete multilayer system is shown, consisting of NHS-SAM–Au(111)–epoxy glue–Si(100). The NHS-SAM covered the

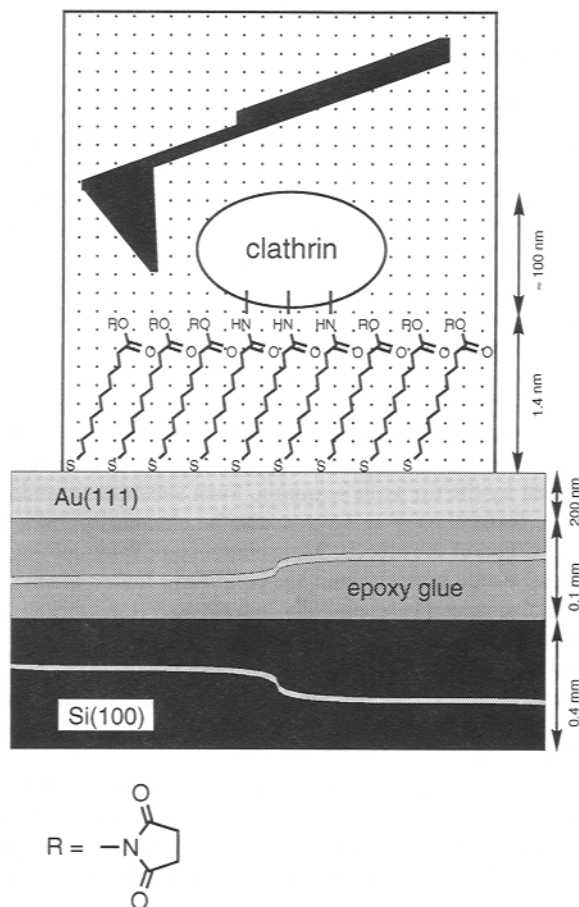


Fig. 1. A schematic view of the experimental set-up developed in the present paper for SPM imaging of a protein (not to scale). (Some of the amino groups of clathrin have reacted with the NHS-activated carboxylic acid groups at the water–solid interface of a self-assembled monolayer. The monolayer is covalently bound via S–Au bonds to ultraflat template-stripped gold (TSG) surfaces (roughness, 2–5 Å over $25 \mu\text{m}^2$) [20]. The Au(111) film is supported by a Si(100) wafer, to which it is glued via a special epoxy resin (Epo-tek).

