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AFM imaging of functionalized carbon nanotubes on biological membranes

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Abstract

Multifunctional carbon nanotubes are promising for biomedical applications as their nano-size, together with their physical stability, gives access into the cell and various cellular compartments including the nucleus. However, the direct and label-free detection of carbon nanotube uptake into cells is a challenging task. The atomic force microscope (AFM) is capable of resolving details of cellular surfaces at the nanometer scale and thus allows following of the docking of carbon nanotubes to biological membranes. Here we present topographical AFM images of non-covalently functionalized single walled (SWNT) and double walled carbon nanotubes (DWNT) immobilized on different biological membranes, such as plasma membranes and nuclear envelopes, as well as on a monolayer of avidin molecules. We were able to visualize DWNT on the nuclear membrane while at the same time resolving individual nuclear pore complexes. Furthermore, we succeeded in localizing individual SWNT at the border of incubated cells and in identifying bundles of DWNT on cell surfaces by AFM imaging.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

Multifunctional carbon nanotubes (CNT) are promising for biomedical applications [1–3]. They can be used for biosensing [4] or act as nano-heaters [5, 6], temperature sensors [7] and drug-carrier systems for therapy and diagnosis at the cellular level [8, 9]. Functionalization of the outer surface of CNT with biomolecules such as nucleic acids, proteins, peptides and polymers leads to their specific internalization into the cell [10–14]. However, the exact uptake mechanism remains a controversial issue as it may well depend on cell type, bio-functionalization scheme, size of the nanotube and other factors [15–17]. In the cited publications, several uptake pathways have been discussed, such as needle-like penetration of the cell membrane, passive diffusion across the lipid bilayers, and energy-dependent endocytosis or phagocytosis. Experiments addressing the uptake of carbon nanotubes into cells usually comprise fluorescent labeling of carbon nanotubes and subsequent detection of the dyes by means of fluorescence microscopy. Label-free observation of carbon nanotube uptake has been achieved by transmission electron microscopy (TEM) [18]. The latter data support a passive diffusion mechanism, as well as the internalization via endocytotic and phagocytotic pathways. Another technique capable of resolving details of cellular surfaces at the nanometer scale is atomic force microscopy (AFM) which has the advantage of being able to operate in liquid [19], allowing for measurements under near physiological conditions [20]. In a previous publication we have presented AFM as a useful tool for a simple and
direct assessment of CNT surface bio-functionalization [21]. Here, we report high-resolution topographic AFM images of functionalized single walled carbon nanotubes (SWNT) and double walled carbon nanotubes (DWNT) immobilized on various relevant biological membranes, including nuclear membranes and cell surfaces.

2. Materials and methods

2.1. DWNT functionalization with bovine serum albumin (BSA) and RNA

Purified DWNT [22] and BSA (Roche) were mixed with deionized water at a concentration of 0.25 mg ml\(^{-1}\) and 1.0 mg ml\(^{-1}\), respectively. The mixture was sonicated for 45 min (Bandelin Sonoplus GM70 tip sonicator, MS 73 microtip, at 20\% power) under cooling in an ice bath. The DWNT BSA suspension was then filtered through centrifugal filters (nanosep, 300 kD cutoff, Pall) at around 5500 g (Eppendorf Centrifuge 5417C) and washed 5 times with deionized water to remove unbound excess BSA. After resuspension in deionized water, the concentration of DWNT was estimated to be 0.05–0.1 mg ml\(^{-1}\), due to losses of material at the sidewall of the filtration device and the filter membrane. The suspension was stored at 4\°C and used within 3 days. For functionalization of DWNT with RNA, purified DWNT and yeast RNA (Roche) were mixed in water with respective concentrations of 0.25 and 0.5 mg ml\(^{-1}\). Sonication and filtration were performed as stated above. The RNA–DWNT conjugates were resuspended in deionized water and stored at 4\°C until usage.

2.2. Functionalization of DWNT with biotinylated BSA

Biotin–BSA was prepared according to Kamruzzahan et al [23]. 5 mg (77 nmol) BSA (Roche Diagnostics) was dissolved in 0.5 ml buffer (100 mM NaCl, 35 mM boric acid, pH adjusted to 8.5 using NaOH) and 5 \(\mu\)l of 66 mM biotin–cap–NHS [24] in DMSO was slowly added with gentle vortexing. After 30 min reaction (with occasional gentle vortexing), the biotinylated BSA was purified by gel filtration. Functionalization of DWNT with biotinylated BSA was done with the same procedure as described for BSA–DWNT. The filtered and washed functionalized DWNT were resuspended in a 1/10 PBS solution.

2.3. Functionalization of SWNT with RNA

SWNT functionalized with RNA were prepared by adding an RNA solution to commercial SWNT (Sigma) resulting in a final concentration of 2.5 mg l\(^{-1}\) SWNT and 2.5 mg l\(^{-1}\) RNA. Tip sonication of the mixture was carried out six times at intervals of 10 s and followed by extensive sonication in a water bath for 1–2 h. The suspension was then centrifuged at 16,000 g (Eppendorf Centrifuge 5415R) for 50–90 min to remove impurities and large nanotube bundles. The supernatant was subsequently collected and filtered through centrifugal filters (nanosep 100 kD cutoff, Pall) to remove excess RNA molecules. The RNA–SWNT conjugates were resuspended in deionized water. The final concentration of the RNA–SWNT suspension was determined by UV–vis analysis [25].

2.4. Preparation of an avidin layer and incubation with biotin–BSA–DWNT conjugates

Freshly cleaved muscovite mica was mounted in a commercial AFM fluid cell prior to AFM imaging and immersed with an avidin solution [24, 26]. The avidin solution was prepared by diluting an avidin stock solution (1 mg ml\(^{-1}\) in PBS) with a 9:1 mixture of water and PBS (PBS\(^{+}\)) to a final avidin concentration of 0.1 mg ml\(^{-1}\). The avidin molecules were immobilized onto mica via electrostatic adsorption due to their positive net charge [24]. After 20 min, the mica substrate was thoroughly rinsed with PBS\(^{+}\). For incubation with biotin BSA–DWNT, the fluid cell was filled with 600 \(\mu\)l of a biotin–BSA–DWNT suspension in PBS\(^{+}\) at a concentration of 0.2 mg ml\(^{-1}\) for 20 min. Before AFM imaging, the CNT containing buffer suspension was replaced by PBS\(^{+}\).

2.5. Preparation of nuclear membrane samples and incubation with RNA–DWNT conjugates

Preparation of nuclear envelopes was performed as described previously [27]. Briefly, oocytes were obtained by surgery of Xenopus laevis females. Stage VI oocytes were used
for nucleus extraction. For this purpose, the oocytes were placed in nuclear isolation medium (NIM) (90 mM KCl, 26 mM NaCl, 5.6 mM MgCl₂, 1.1 mM EGTA, 1.5% w/v polyvinylpyrrolidone (PVP), 10 mM HEPES pH 7.4) and the nuclei were dissected with a pair of sharp tweezers. Nuclei were further purified from the remaining yolk by washing two times in NIM. A 200 μl drop of PVP-free NIM-buffer was placed on ethanol-cleaned glass slides. The nuclei were transferred into the drop, opened and prepared with the cytoplasmic side up using sharp needles. The samples were thoroughly washed with deionized water and air dried. For imaging of the DWNT on the surface of the nuclear envelope, 1 μl of the RNA–DWNT suspension in deionized water was placed on a dry nucleus sample and the liquid was allowed to evaporate.

2.6. Delivery of RNA–SWNT conjugates to mammalian cells

HeLa cells (human cervical carcinoma cells) were cultured in 35 mm plates containing poly-l-lysine-coated cover slips using Eagle’s minimal essential medium (MEM) containing 2 mM l-glutamine, 0.1 mM non-essential amino acids, 10% fetal bovine serum, and penicillin/streptomycin (100 units ml⁻¹ penicillin and 100 μg ml⁻¹ streptomycin in the final formulation). The medium is formulated for use with 5% CO₂ in air atmosphere. When the cells reached 80% confluence, they were incubated for 3 h with RNA-functionalized SWNT in deionized water placed on a dry nucleus sample and the liquid was allowed to evaporate.

2.7. Delivery of BSA–DWNT conjugates to mammalian cells

ECV 304 cells (human urinary bladder carcinoma cells) were cultured in RPMI 1640 medium (E15-848), containing 10% fetal calf serum, supplemented with 1% penicillin/streptomycin and 1% 1 M HEPES (all: PAA Laboratories, Vienna, Austria). Cells were passaged twice a week using a dilution of 1:2 and were maintained under air atmosphere with 5% CO₂ at 37 °C. For AFM experiments, cells were seeded on cleaned (with ethanol and sterile water) glass cover slips (22 × 22 mm²) with a dilution of 1:80 to obtain 50% confluence after 2 days. For imaging of DWNT on the surface of ECV cells, two different approaches were used. (a) Cells were fixed with 4% paraformaldehyde in PBS over a period of 30 min. After washing with PBS, the cells were exposed to RNA-functionalized DWNT diluted in HBSS at a concentration of ∼100 μg ml⁻¹ for several hours to allow for sedimentation of DWNT on the fixed cells. Prior to AFM imaging the DWNT solution was replaced by plain HBSS. (b) After having reached 60% confluence, the cells were incubated for 3 h with RNA–DWNT conjugates diluted in medium at a concentration of ∼100 μg ml⁻¹. Subsequently, the cells were washed several times with PBS buffer, fixed with 4% paraformaldehyde for 30 min, washed again, and kept in HBSS for AFM imaging.

2.8. MAC mode AFM imaging of avidin layer

Topographical imaging of biotin–BSA–DWNT incubated on an avidin layer was carried out in PBS buffer (phosphate buffered saline) at room temperature with a PicoPlus atomic force microscope (Agilent Technologies, Chandler, AZ) equipped with a conventional fluid cell. Images were acquired in magnetically driven dynamic mode (MACmode, Agilent Technologies, Chandler, AZ). AFM tips with a magnetic coating (Agilent Technologies, Chandler, AZ) and a nominal
spring constant of 0.1 N m\(^{-1}\) were used. The measurement frequency was set to 20% below the resonance frequency. Scanning was done with the feedback adjusted to 40% amplitude reduction at a lateral scan frequency of 1.0 Hz with 512 scan lines per image.

### 2.9. Contact mode imaging of nuclear membrane and cells

Topographical imaging of RNA–DWNT on a dried nuclear membrane sample was performed with a PicoPlus AFM (Agilent Technologies, Chandler, AZ) operated in contact mode at room temperature with a lateral scan rate of 1–1.5 Hz at 512 lines. AFM cantilevers with a nominal spring constant of 0.03 N m\(^{-1}\) (Veeco) were used. Contact mode imaging of fixed ECV cells was performed in physiological buffer (Hank’s buffered salt solution). Contact mode imaging of HeLa cells was done with a Digital Instruments dimension 3100 atomic force microscope (Veeco) under dry conditions with a lateral scan rate of 2 Hz at 265 sample lines using cantilevers with a nominal spring constant of 0.03 N m\(^{-1}\).

### 3. Results and discussion

The goal of this study was to achieve high-resolution imaging of CNT on the surfaces of cells and organelles. In preliminary test series, the soft cell surface was mimicked by an ultraflat support to which CNT were firmly attached. For this purpose, CNT were functionalized with biotin residues and then bound to a dense monolayer of avidin (a biotin-binding protein) which had been formed on freshly cleaved mica. Spontaneous formation of avidin monolayer on mica is due to the positive net charge of avidin and the negative charge of mica at neutral pH [23]. The intactness of the avidin layer was proven by scratching a small rectangular area (200 nm × 200 nm) with high force (10 nN) in order to expose bare mica (figure 1(a) inset). Cross-section analysis (figure 1(b) inset) revealed a layer height of ~1.5 nm and a surface roughness of less...
The nuclear envelope presents an interesting biological membrane which, in contrast to an artificial membrane, native biological membranes are less homogeneously structured and show significantly higher roughness. Here, it was possible to obtain very flat nuclear membrane patches. The average roughness of the underlying NPC area was 4–5 nm, while the height of individual DWNT ranged from 2 to 4 nm and the height of bundles from 6 to 15 nm. In this way the visualization of single DWNT on the nuclear membrane was achieved while, at the same time, resolving individual nuclear pore complexes.

For transport of CNT to the nucleic region of a cell and into the nucleus, the nanotubes first have to bind to the cell surface followed by internalization. Having a possibility to observe the binding of individual CNT to the cell directly, could provide further insight into the uptake pathway of bi-functional CNT for therapeutic applications. To visualize CNT immobilized on a cell surface, HeLa cells were imaged after incubation with RNA-coated SWNT under dry conditions. Figure 3 shows topographic images captured in contact mode. When gradually zooming-in on the border of an isolated cell, the images showed tangled single nanotubes in close proximity to the cell border. At the edge (figure 3(e)) of the cell, some of the SWNT are found to partially extend onto the cell surface. In order to improve the visibility of the SWNT, a polynomial background was subtracted in figure 3(f) in order to level the steep edge of the cell and the underlying glass slide. Furthermore, when imaging directly on the cell surface, no RNA–SWNT were observed. Scanning of several live and fixed HeLa cells showed that these cells generally grow in a rather compact form and exhibit a very soft surface. This softness obviously impairs the detection of individual CNT on top of the cell.

To meet the particular challenge of imaging small diameter CNT on cell surfaces, it is important to use flat cells which exhibit a low surface roughness and form widespread flat regions. We conducted systematic investigations on different cell lines (EAhy, DU-145 and MyEnd cells) via topographic AFM imaging of live and fixed mammalian cells to find a suitable cell line. ECV cells, a derivative of human urinary bladder carcinoma cells, grow in culture as an adherent monolayer. AFM imaging revealed the typical sizes of ECV cells in the range of 50–80 μm. The nucleus region has a height of ~3 μm while the surrounding cytoplasmic area is rather flat, with a surface roughness of ~50 nm. Figure 4 shows a glass slide with cells at about 40–50% confluence, which were incubated with BSA-coated CNT. The triangular AFM cantilevers are visible in the upper part of the picture. The black spots are BSA–CNT aggregates and lumps. Interestingly, most lumps are located on the cells, while the bare glass seems to be free of CNT aggregates. For successful imaging, the AFM cantilever was positioned on cells without CNT aggregates to allow for imaging of individual CNT.

Contact mode AFM images of ECV cells incubated with BSA–DWNT acquired under near physiological conditions in buffer solution are shown in figure 5. Larger bundles of CNT were typically observed on the cell surfaces. In figure 5(a), a long straight structure lies atop the cell surface, with a height of about 80 nm, probably representing a bundle of DWNT. Beneath the bundle, filamentous cellular substructures can be observed (medium-scale scan) which appear in the small scan-size image as an interlaced network. The filaments range in height between a few nanometers and 50 nm, causing an average roughness valley depth of 14.5 nm and an average roughness peak height of 14.4 nm. When comparing these values to the typical dimensions of a BSA-coated DWNT with roughly 5–12 nm in height, it appears difficult to discriminate...
subcellular structures as filaments from individual carbon nanotubes.

Further examples of DWNT bundles on cell surfaces are displayed in figures 5(b) and (c). In figure 5(b), a bundle of DWNT can be seen in the lower left part of the large-scale image. The bundle exhibits a height of 250–300 nm. The two bright round regions encircled in the upper left corner and in the right part of the large-scale image correspond to the nuclei of two ECV cells with a height of around 3 μm. In figure 5(c), the deflection images of two ECV cells are shown (nuclei and contact region of the cells are marked). In the lower part, the underlying cover slip is visible where several shorter rather straight features are observed. A zoom-in on the glass cover slip beside the cell border and flat parts of the cells between the nuclei reveals DWNT with a height of 30–80 nm. In contrast to the images in figures 5(a) and (b), here the CNT bundles appear to be sticking in the cell surface which suggests a possible internalization of BSA–DWNT bundles.

4. Conclusions

We presented a method for high-resolution topographical imaging of bio-functionalized DWNT in buffer solution. Stable immobilization of CNT on an ultraflat support was achieved by functionalization of CNT with biotinylated BSA and of mica with electrostatically bound avidin. Furthermore, we showed for the first time high-resolution topographical AFM images of DWNT on the nuclear membrane while resolving individual nuclear pore complexes. This demonstrates the potential for monitoring the transport of individual CNT through single nuclear pore complexes into the cell nucleus with the AFM, allowing for time-resolved studies with high lateral resolution. For such an experiment, an NPC transport protein, e.g. importin β, is needed as a bio-functional coating for CNT to study the binding to an individual NPC structure in comparison with a control sample of CNT coated with a non-binding protein [28].

Our experiments with biological cells showed great differences depending on the measurement conditions, cell line and the type of functionalized carbon nanotubes. On HeLa cells, SWNT could only be seen near the cell border and lack of detection of CNT in the center region was attributed to the surface topology. In contrast, DWNT were well detectable on ECV cells, yet here only large bundles of CNT could be identified with certainty because individual DWNT could not unequivocally be discriminated from filaments of the cytoskeleton which consist of microtubules (∼25 nm
diameter), intermediate filaments (roughly 10 nm) and actin filaments (~10 nm). For comparison, individual SWNT and DWNT have diameters ranging between 1–3 nm and 2–4 nm, respectively (without bio-functionalization). Accordingly, it was only possible to observe and identify larger bundles of CNT with diameters of 50 nm up to several hundreds of nanometers lying across cell surfaces. Although these preliminary experiments indicate certain limitations for the AFM to study binding of individual carbon nanotubes to cells and their uptake into the cell, complementary techniques may prove promising. The combination of fluorescence microscopy and AFM imaging [29] as well as simultaneous topographical and recognition imaging (TREC) [30, 31] with a ligand-functionalized AFM tip could facilitate the localization of functionalized carbon nanotubes on the cell surface and time-resolved uptake studies.

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