Enhanced endothelial cell functions on rosette nanotube-coated titanium vascular stents

Eli Fine¹
Lijie Zhang¹
Hicham Fenniri²
Thomas J Webster¹
¹Department of Engineering, Brown University, Providence, RI, USA; ²National Institute for Nanotechnology and Department of Chemistry, University of Alberta, Edmonton, AB, Canada

Abstract: One of the main problems with current vascular stents is a lack of endothelial cell interactions, which if sufficient, would create a uniform healthy endothelium masking the underlying foreign metal from inflammatory cell interference. Moreover, if endothelial cells from the arterial wall do not adhere to the stent, the stent can become loose and dislodge. Therefore, the objective of this in vitro study was to design a novel biomimetic nanostructured coating (that does not contain drugs) on conventional vascular stent materials (specifically, titanium) for improving vascular stent applications. Rosette nanotubes (RNTs) are a new class of biomimetic nanotubes that self-assemble from DNA base analogs and have been shown in previous studies to sufficiently coat titanium and enhance osteoblast cell functions. RNTs have many desirable properties for use as vascular stent coatings including spontaneous self-assembly in body fluids, tailorable surface chemistry for specific implant applications, and nanoscale dimensions similar to those of the natural vascular extracellular matrix. Importantly, the results of this study provided the first evidence that RNTs functionalized with lysine (RNT–K), even at low concentrations, significantly increase endothelial cell density over uncoated titanium. Specifically, 0.01 mg/mL RNT–K coated titanium increased endothelial cell density by 37% and 52% compared to uncoated titanium after 4 h and three days, respectively. The excellent cytocompatibility properties of RNTs (as demonstrated here for the first time for endothelial cells) suggest the need for the further exploration of these novel nanostructured materials for vascular stent applications.

Keywords: stents, nanotechnology, self-assembly, rosette nanotubes, endothelial cells, titanium

Introduction
Vascular stents are predominantly metal structures inserted into partially clogged arteries to promote normal blood flow and prevent myocardial infarctions. There is an enormous market for vascular stents with $918 million in US sales in 2005 alone.¹ The constant contact of vascular stents with blood has been the source of several unresolved efficacy issues.²,³ For example, conventional bare metal stents often trigger an inflammatory response leading to scar tissue formation, known as restenosis, which constricts blood flow defeating the original purpose of the stent.³ Drug-eluting stents counteract such events by releasing immunosuppressants, however, these immunosuppressants can also suppress desirable endothelial cell growth and sometimes lead to late-stage thrombosis.³ For the above reasons, this study focused on the design, synthesis, and investigation of a new non-drug containing material called rosette nanotubes (RNTs) to coat titanium stent surfaces in order to enhance endothelial cell adhesion and proliferation.

RNTs are novel materials composed of DNA base analogs featuring the hydrogen bonding arrays of both guanine and cytosine, the G>C motif. This module self-assembles spontaneously in aqueous solutions first into a rosette supermacrocycle...
maintained by 18 hydrogen bonds (Figures 1A–B). Then the rosettes undergo self-organization via noncovalent stacking interactions to form helical nanotubes with an inner diameter of 11 Å and lengths ranging from a few nanometers to several micrometers (Figure 1C). Such nanometer dimensions of RNTs mimic the dimensions of the natural components of tissues, such as collagen fibrils. Furthermore, the outside of the tubes can be functionalized to possess various peptides and amino acid side chains (eg, K, RGD [arginine–glycine–aspartic acid]) to interact with cells and stimulate specific tissue growth.

For orthopedic applications, previous studies have shown that various functionalized RNTs have attractive properties including remarkably improved osteoblast (bone-forming cell) functions when coated on traditional implant materials (such as titanium) or embedded in hydrogels.4–8 Promoted osteoblast functions have been attributed to the nanoscale biologically-inspired features and the rich lysine or peptide moieties of RNTs. In addition, RNTs have been investigated as novel injectable materials which solidify when heated to body temperatures.4–7 Of course as with any new nanotechnology-derived material, toxicity could be a concern. However, in a recent in vivo study, Journeay and colleagues reported that RNTs did not trigger severe inflammatory responses and had low pulmonary toxicity in vivo at mg/mL concentrations (toxic concentrations above which RNTs have been shown to promote cell functions).9

For all of the above reasons, especially due to the promise of coating titanium with RNTs for orthopedic applications, this report focuses on endothelial cell functions on RNT-coated titanium with the aim of improving vascular stent performance.

**Materials and methods**

**Substrate preparation and characterization**

Titanium squares (Ti, 10 × 10 × 0.5 mm, Alfa Aesar titanium foil, 99% metal basis) and glass coverslips (Fisher Scientific, Pittsburgh, PA) were cleaned in acetone for 15 min, sonicated for 15 min, and rinsed three times with Millipore distilled/deionized water (ddH2O; Millipore, Billerica, MA). They were then soaked, sonicated, and rinsed in 70% ethanol then in ddH2O. The glass was etched (1 M NaOH, 1 h) and thoroughly rinsed in ddH2O. The substrates were then placed in a drying oven overnight and autoclaved before the cell adhesion experiments.

The RNT–K (RNTs functionalized with lysine) were synthesized as previously reported,10 and the corresponding solutions were sterilized by filtration via a 0.22 µm syringe filter.4 To confirm the formation of the RNTs, a sample (0.01 mg/mL) was imaged on a Philips, EM410 PW6008 transmission electron microscope (Philips, Amsterdam, The Netherlands). One day prior to seeding cells, the titanium substrate samples were placed in solutions of either 0.01 mg/mL or 0.001 mg/mL RNT–K for 45 min, then removed and left to dry overnight.

**Endothelial cell culture**

Rat aortic endothelial cells (RAEC; VEC Technologies, Rensselaer, NY) were cultured in MCDB–131 Complete medium (VEC Technologies) under standard cell culture conditions (37 °C, humidified, 5% CO2/95% air). Medium was replaced every other day.

**Endothelial cell adhesion experiment**

For the 4 h cell adhesion studies, a 12-well plate was prepared with three samples consisting of 0.01 mg/mL...
Rosette nanotubes as vascular stent coatings

RNT-coated titanium, 0.001 mg/mL RNT-coated titanium, uncoated titanium, and a glass reference. Cells were seeded onto the substrates at a density of 3500 cells/cm² in 2 mL of MCDB-131 Complete Medium (Sigma-Aldrich, St. Louis, MO) and were incubated for 4 h. The medium was then removed and the substrates were rinsed with phosphate-buffered saline (PBS; 3×) to remove non-adherent cells. The adherent cells were then fixed with 10% formalin (Fisher Scientific) for 10 min and then rinsed with PBS (3×). The cells were then fixed with 0.1% Triton X–100 (Sigma-Aldrich) for 5 min. After sequential PBS rinsing, endothelial cells were stained with rhodamine–phallolidin (Molecular Probes, Carlsbad, CA) for 20 min, rinsed with PBS (2×), stained with DAPI (Invitrogen, Carlsbad, CA) for 5 min and rinsed with PBS. 4',6-Diamidino-2-phenylindole (DAPI) was chosen as a fluorescent dye since it stains the cell nucleus and can be used to count the number of cells present on opaque substrates. Rhodamine phallloidin was also chosen because it stains the F-actin filaments of endothelial cells and, thus, highlights cell morphology. After staining, endothelial cells were observed using a Zeiss Axiovert 200 M fluorescence microscope (Carl Zeiss Microimaging Inc., Thornwood, NY) and five different areas of each sample were imaged. The cell density was determined by counting cells using Image Pro Analyzer (Media Cybernetics, Inc., Bethesda, MD).

Endothelial cell proliferation experiment

For the proliferation studies, three sets of substrates were seeded with endothelial cells at a density of 1000 cells/cm² and were placed in an incubator for 1, 3, and 5 days. The medium was changed every other day. After each time period, cells were stained and counted as described above.

Statistical analysis

All cellular experiments were run in triplicate and repeated at least three different times for each substrate. Numerical data were analyzed with a Student’s t-test to make pairwise comparisons. Statistical significance was considered at p < 0.1.

Results and discussion

RNT–K morphology

Transmission electron microscopy images showed that the RNT–K used in the present study formed a dense network of long nanotubes each ∼3.5 nm in diameter and several hundred nanometers in length (Figure 2), consistent with earlier reports.4

Enhanced endothelial cell adhesion

The 4 h adhesion study provided the first evidence that endothelial cells adhered at a higher density to RNT-coated titanium compared to uncoated titanium substrates. Specifically, the 0.01 mg/mL and 0.001 mg/mL RNT–K-coated titanium increased endothelial cell density by 37% and 22%, respectively, over uncoated titanium (Figures 3 and 4).

One hypothesis concerning why increased endothelial cell adhesion was observed on RNT-coated compared to uncoated titanium is that the biomimetic features of the nanostructured RNTs favored endothelial adhesion whereas the uncoated titanium did not possess such biologically-inspired surface roughness. In addition, another study reported that poly-L-lysine-coated vascular stents significantly improved endothelial cell adhesion compared to uncoated controls perhaps due to the electrostatic or optimal initial protein interactions between poly-L-lysine and endothelial cells.11 Considering the rich lysine side chains and well controlled nanometer spatial distributions of such lysine side chains on RNTs, the surface chemistry of titanium was altered through RNT coatings to possibly promote greater endothelial cell attachment and spreading as shown in Figure 5.

Endothelial cell proliferation

Endothelial cell density increased on all substrates after 5 days of culture (Figure 6). More importantly, results of this
Figure 3 Enhanced endothelial cell adhesion on 0.01 and 0.001 mg/mL RNT–K coated on titanium after 4 h compared to uncoated titanium. Notes: Data are mean values ± SEM; N = 3. *p < 0.05; **p < 0.1 when compared to uncoated titanium; and #p < 0.005 when compared to all other substrates. Abbreviations: RNT–K, rosette nanotube–lysine side chain; SEM, standard error of mean.

The study provided the first evidence that RNT–K-coated titanium greatly improved endothelial cell functions compared to uncoated titanium (Figure 6). Specifically, the 0.01 mg/mL and 0.001 mg/mL RNT-coated titanium increased endothelial cell density 52% and 21%, respectively, compared to uncoated titanium after three days of culture. In addition, endothelial cell densities were significantly higher on 0.01 mg/mL RNT–K coated titanium than on 0.001 mg/mL RNT–K-coated titanium after three days of culture. Such data supports the positive role that higher concentrations of RNTs can play in promoting endothelial cell coverage of vascular stents by (a) presenting more lysine side chains per unit of surface area, and/or by (b) more closely resembling the natural nanostructure of vascular tissue when coated on titanium.

By day 5, the RNT-coated titanium still had a higher average endothelial cell density, but the differences were no longer statistically significant (Figure 6). However, when analyzing endothelial cell numbers from day 1 to 5, it was clear that endothelial cells reached confluency faster on the RNT-coated titanium than on the uncoated titanium. Furthermore, Figures 7 and 8 illustrate the difference in endothelial cell confluence on titanium when coated with RNTs.

In short, this study demonstrated for the first time improved endothelial cell density on RNT-coated titanium

Figure 4 Fluorescence microscopy images of increased endothelial cell densities after 4 h on A) 0.01 mg/mL RNT–K coated on titanium and B) 0.001 mg/mL RNT–K coated on titanium compared to C) uncoated titanium. Dots in the images represent DAPI-stained endothelial cell nuclei. Scale bars = 50 μm (left), 40 μm (middle), and 50 μm (right). Abbreviations: DAPI, 4′,6-diamidino-2-phenylindole; RNT–K, rosette nanotube–lysine side chain.
compared to uncoated titanium controls. Importantly, this was achieved without embedding or coating drugs on the titanium stent surface, rather, such positive data was obtained after simply soaking titanium in RNT-containing solutions for 45 min. This study, thus, suggests the potential of using biomimetic RNTs as a quickly applied, efficient coating on titanium to improve cytocompatibility properties for vascular stent applications.

Conclusions
In summary, this initial investigation concerning endothelial cell functions on RNT-coated titanium stents yielded
Figure 7 Fluorescence microscopy images of increased endothelial cell densities (DAPI stain) on 0.01 mg/mL and 0.001 mg/mL RNT–K coated on titanium compared to uncoated titanium after 1, 3, and 5 days of culture.

Abbreviations: DAPI, 4′,6-diamidino-2-phenylindole; RNT–K, rosette nanotube-lysine side chain.

Figure 8 Fluorescence microscopy images of double-stained endothelial cells on 0.01 mg/mL and 0.001 mg/mL RNT–K coated on titanium and uncoated titanium after 1, 3, and 5 days of culture. F-actin filaments were stained by rhodamine phallolidin and cell nuclei were stained by DAPI.

Abbreviations: DAPI, 4′,6-diamidino-2-phenylindole; RNT–K, rosette nanotube-lysine side chain.
Rosette nanotubes as vascular stent coatings

very positive results. Titanium coated with RNTs showed a significant concentration effect of increased endothelial cell density compared to uncoated titanium after 4 h and after three days. While further investigation is clearly needed, such as using dynamic flow rather than the static tests conducted here, the results of this study provided promise for the use of RNTs to improve vascular stent applications.

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References
