Endothelialization of Titanium Surfaces**

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In the extracellular matrix (ECM), chemical cues are present that control all aspects of cell biology.[1–3] These surface-bound and soluble factors provide the necessary adhesion and signaling for normal cellular activity, and without such matrix support, the cells will quickly apoptose. Implanted device surfaces lack the molecular features that provide guidance to the surrounding cells to afford optimal in vivo integration and function. One class of implanted materials are metal implants, which are commonly used in cardiovascular therapy (e.g., stents) and orthopedic procedures (e.g., hip replacement). While such materials are favored for their inertness and mechanical strength, negative consequences can arise from suboptimal tissue integration. For example, during a coronary angioplasty an occluded coronary artery is opened using a balloon catheter and then a metal stent is inserted to provide a permanent framework supporting vascular patency. If the artery re-occludes due to smooth muscle cell proliferation, in a process called restenosis, a second procedure is required to re-establish blood flow. Restenosis used to occur in about 25% of patients, but with the introduction of stents that elute a mitotic inhibitor, such as paclitaxel, this number has been reduced significantly.[4] However, there is new concern regarding the use of drug-eluting stents owing to an increased thrombolytic potential of two to three fold compared to bare metal stents.[5] An alternative approach to the use of drugs would be to develop materials that better integrate with the surrounding endothelial cells that line the artery to afford a prohealing response. Ideally, such materials would provide cues to appropriately direct cellular activity, for example, through integrin interactions that control an assortment of biological processes.[6,7] In order to impart suitable extracellular signals, we are developing interfacial coatings to study and ultimately control cell biology on a metal implant. As our first foray into this area, we have synthesized a procellular coating that adheres to a Ti surface under mild aqueous conditions and retains endothelial cells under flow.

In its simplest form this interfacial coating is a 20 to 30-mer peptide that possesses two domains, one that binds to Ti metal and a second domain that binds to endothelial cells (Fig. 1). For this set of experiments, the ubiquitous arginine–glycine–aspartic acid (RGD) integrin-binding motif was selected for the cell-binding domain. This tripeptide sequence, found on most ECM proteins, readily binds to a number of cell types, including endothelial cells. Specific endothelial cell-binding peptides have yet to be reported, although the identification of cell distinctive peptides is an active research area with some recent notable successes (e.g., neutrophils, hematopoietic stem cells, cancer cells).[8–11] Previous efforts to attach the RGD domain to Ti surfaces have typically required careful control over the multistep process required to coat the material.[12–18] Our next step was to identify peptide sequences that bind Ti metal. We chose to use a phage display combinatorial library that operates through affinity selection of phage-displayed peptides based on their ability to bind to Ti metal.

Figure 1. Schematic of an interfacial coating residing at the boundary between a material and a cell. Blue residue positions represent the material-binding domain, while orange represents those residues that bind the cell.

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encoded peptides to identify the binding motif(s). This iterative selection process has been used previously to isolate peptides that recognize a variety of diverse targets such as GaAs nanocrystals, iron oxide, gold, polystyrene, and a variety of other materials.

A filamentous M13 phage displaying a 19-mer peptide (SCX$_5$C, where the one letter abbreviations for the amino acids are used and X represents one of the twenty amino acids encoded by synthetic NNK (N: A, C, G, T; K: G, T) codons, $10^{10}$ total phage screened, $10^8$ complexity) attached to the amino terminus of the pIII coat protein was used to identify peptides that adhere to 6Al-4V Ti, a commonly used implant material. Our prior experiences with the commercial libraries have not yielded adequate binders to our previous targets, and thus we have designed new libraries such as the one used in these experiments. This library was designed to have one fixed cysteine residue in its sequence to favor the possibility of a cyclic disulfide motif being selected due to the reduced entropic penalty associated with binding. The phage that bound to the target Ti substrate were eluted and amplified by infection of a bacterial host. Once the target binding phage clones were identified, the base sequence of the DNA insert in the phage genome was located and then translated to yield the corresponding amino acid sequence displayed on the phage surface. After three rounds of screening, we identified 10 peptides with high Ti affinity, listed in Table 1. Sequence 4, SCSDCLKSVDFIPSSLASS, was selected for further study because its relative binding affinity on phage was two to four fold greater than the weakest binders. Previous experiments have suggested that this initial screen is an effective method to rapidly quantify the relative binding strengths of the sequences without having to fully synthesize them. Analysis of the 10 sequences shows that D occurs more frequently than expected (13 occurrences, 5.2 expected in 160 library-encoded residues, ratio of 2.5 times over expected), and that C and S are slightly enriched in the variable region (10 occurrences, 5.2 expected, ratio of 1.9 and 27 occurrences, 15.5 expected, ratio of 1.7 times over expected, respectively). This suggests a possible mode of coordination to the Ti oxide surface via hydrogen bonding for this peptide sequence. Finally, F is highly enriched (18 occurrences, 5.2 expected, ratio of 3.5 over expected), although the other aromatic residues are not (see Fig. S2 in the Supporting Information).

Next, peptide 4 was prepared using automated solid-phase peptide synthesis following standard N-9-fluorenylmethoxy-carbonyl (FMC) protocols containing either a C-terminal biotin, the prototypical RGD integrin-binding domain, or the closely related, but nonfunctional arginine–glycine–glutamic acid (RGE). The resulting peptides: SCSDCLKSVDFIPSSLASS-SSG-Biotin, 11, SCSDCLKSVDFIPSSLASS-SSG-RGDS, 12, and SCSDCLKSVDFIPSSLASS-SSG-RGESP, 13, contained a tripeptide spacer of SSG to separate the Ti binding domain from the biotin, RGD, or RGE domain. We coated Ti disks (height = 3.8 mm; diameter = 12.7 mm), which were lathe cut from a 6Al-4V Ti rod (root mean square = 150 nm), by dissolving 12 and 13 in aqueous buffer, and applying the solutions to the disk surfaces followed by washings to remove nonspecific binding. The disk surface was cleaned extensively prior to use and was characterized using atomic force microscopy (AFM), scanning electron microscopy (SEM), and X-ray photoelectron spectroscopy (XPS). The latter confirmed the presence of a TiO$_2$ coating (Ti2p/O1s = 0.48 ± 0.003) as well as some vanadium and aluminum as expected, with values similar to those reported in the literature. Prior to application the contact angle of the Ti surface was 53.4° ± 2.0°, which is in agreement with literature values for bulk Ti6Al4V materials where depending on the surface roughness the angle can vary from 35° to 75°C. After coating with 12 the contact angle decreased to 37.3° ± 1.4° ($n = 3$; $p < 5.10^{-5}$), with a similar decrease to 38.4° ± 1.5° recorded for 13. As a control, a coating of bovine serum albumin (BSA) did not change the contact angle of the Ti significantly (53.9° ± 3.8°; $p = 0.84$). We next determined the binding affinity of 11 to Ti by enzyme-linked immunosorbent assay (ELISA) using a streptavidin-horseradish peroxidase conjugate and a 2,2′-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (ABTS) readout. The affinity constant ($k_{\text{aff}}$) of the peptide material domain to the Ti disk was found to be 4.1 × 10$^{12}$ m$^{-1}$.

To assess the ability of the Ti-RGD macromolecule, 12, to affect endothelial biology on a Ti metal surface, we examined attachment morphology, proliferation, and cell binding. As a first step in the process, optical images of human umbilical vein endothelial cells (HUVECs) on a Ti surface were obtained by utilizing glass microscope coverslips coated with a 20 nm film of Ti metal (root mean square = 0.75 nm). Due to the difficulty in evaporating alloys because of the differing vapor pressures of the components, commercially pure Ti was substituted as a model for these studies. XPS data again reveals the presence of TiO$_2$ (Ti2p/O1s = (0.48 ± 0.01)), however as expected, this data verifies that the oxide layer on pure Ti is not identical to the oxide layer seen on the Ti6Al4V alloy, but is similar to values reported in the literature. Additionally, the XPS data shows complete Ti surface coverage as the Si2p peak is not significantly greater than that on the bulk Ti disks ($p = 0.23$). Next, we examined the binding affinity of the peptide to this new TiO$_2$ surface. Binding strength measure-

Table 1. Ti binding sequences isolated from the phage display combinatorial library and their functionalized derivatives. Italicized residues indicate nonvariable positions.

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<tr>
<td>1</td>
<td>SCFWLRSLSLFLFTCCS</td>
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<td>SCSENFMNMYCTGVCTES</td>
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<tr>
<td>2</td>
<td>SCESVDFCADSRMKSMS</td>
<td>7</td>
<td>SCSSFSEVMFCATAVIS</td>
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<tr>
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<td>8</td>
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</tr>
<tr>
<td>4</td>
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<tr>
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<td>10</td>
<td>12</td>
</tr>
<tr>
<td>6</td>
<td>SCMLFSSVFDCGMUSDSL</td>
<td>11</td>
<td>13</td>
</tr>
<tr>
<td>7</td>
<td>SCVDYVMHADSPGPDGLNS</td>
<td>12</td>
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ments determined using the same procedure as previously used showed that the affinity constant was slightly lower but within the same order of magnitude as with the Ti disks. This suggests that the Ti binding peptide, 12, has general micromolar affinity for TiO2 surfaces regardless of the underlying Ti composition. Contact-angle measurements on this substrate, when compared to the bulk Ti disks, showed a slightly more hydrophobic surface with an angle of 63.2°±3.7° similar but not identical to literature values, again, owing to the differing cleaning procedures and surface roughnesses. However, the effects created by a coating of peptides 12 and 13 were more pronounced than those witnessed earlier with angles of 8.7°±2.1° (n=5; p<5×10⁻⁸) and 9.7°±2.9°. Again, a slight increase in hydrophobicity was seen with the BSA coating to an angle of 69.7°±3.0°. In order to determine if the contact angle changes were due solely to leaching of the surface-bound molecules into the incident droplets, contact-angle experiments were performed where 30.7 μM concentrations of 12 (0.1 mg mL⁻¹) or arginine-glycine–aspartic acid–serine (RGDS) (0.013 mg mL⁻¹) were placed onto the surface of a cleaned Ti coated coverslip. Measurements taken show a small decrease in contact angle of 15.7°±5.6° for RGDS and 11.2°±2.5° for 12, but not the 50° change seen on the peptide-coated surfaces. The concentrations used in this experiment were orders-of-magnitude above the leachable amounts from the surface in coated conditions, and show that the changes in contact angles cannot be due to this effect alone but must also be caused by a change in surface character due to the bound peptide coating.

Next, the cell studies were performed. A 100 μL solution of 12, 13 (0.1 mg mL⁻¹ in phosphate buffered saline (PBS)), or plain PBS was added to the center of these slides and let stand for 2 h. The solutions were then removed and 100 μL of a 1% BSA solution was added and the slides were let stand for an additional 2 h. The BSA was removed and the slides were washed with PBS. Next, 100 μL of medium containing 2.0 × 10⁴ HUVEC was added and allowed to interact with the surfaces for 60 min. Microscope images were taken of the slides at 0, 15, 30, 45, and 60 min after addition of the cells. After 60 min, the slides were washed profusely with excess PBS and an image was taken of the cells that remained using an actin immunostain. The beginnings of HUVEC spreading can be observed after 15 minutes and is readily apparent after 60 min on the 12 coated (RGD) surface (Fig. 2A and D). In contrast, HUVECs on the 13 (RGE) and uncoated surfaces did not spread as well over the time course of the experiment with only slight pseudopodia observed in most cases (Fig. 2B, C, E, and F). Staining the actin filaments of the cytoskeleton with phalloidin-tetramethyl rhodamine iso-thiocyanate (TRITC) further demonstrates cell spreading on the coated Ti with marked membrane ruffling, and corroborates the differences witnessed with the cells observed in the phase-contrast images (Fig. 2G–I). Surface areas were calculated for 180+ cells in each image revealing that the cells on the uncoated or 13 coated surfaces had approximately identical mean areas at all five time points (p>0.01) and doubled in area during the 60 min on the 12 coated and uncoated Ti disk surfaces and the relative proliferation rate was calculated as follows. Thirty Ti disks were treated with either 1 mg mL⁻¹ solution of 12 (n=15) or buffer (n=15) for 1 h at room temperature (RT), and then washed with PBS. BSA was not utilized in this experiment so that attachment rate to the surfaces and spreading rates did not become a factor. Next, 20 000 HUVECs in medium were seeded onto ten disks in each group, the other five in each group were exposed to plain media for a background reading. Five of the coated and uncoated disks from each group were immediately developed using an 3(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS) proliferation assay at 1 h. The remaining two groups of five disks were

![Figure 2. Phase contrast image of HUVEC on a Ti-coated glass slide treated with 12 (top), 13 (middle), or PBS (bottom) for 1 h. Images were taken at 15 min (A,B,C), and 60 min (D,E,F) post cell addition, at which point a washing step was performed to remove any unbound cells and actin immunostain was applied (G,H,I) (scale bars in all photos represent 100 μm).](image-url)
incubated for 24 h prior to assaying. The proliferation doubling times were determined using an MTS assay and subtracting the background readings from the resultant data points. The proliferation doubling times were found to be 

\(25.0 \pm 2.4\) and 

\(22.0 \pm 2.7\) h for the coated and uncoated Ti surfaces, respectively. The coating did not alter the proliferation of the cells on the surface as the difference between the timings is negligible 

\(p = 0.23\), and both are within the expected population doubling times reported for HUVECs.\(^{[42,43]}\)

The above experiments were conducted under static conditions, but many biological processes occur under dynamic conditions, such as flow. Given our interest in endothelialization of metal surfaces and the importance this process has in stenting and ultimate clinical efficacy, we investigated whether this coating affected HUVEC retention under flow. Eight Ti disks were placed in a custom-built flow chamber and treated with either 250 \(\mu\)L of \(12\) (0.1 mg mL\(^{-1}\)) or 250 \(\mu\)L of PBS for 2 h (Fig. 3A). A 1 % BSA solution in PBS was added for an additional 2 h because albumin, a major component of blood, would be present during the attachment process, and is known to block nonspecific binding. The solutions were removed and the disks washed before adding a 250 \(\mu\)L media solution (2 % fetal bovine serum (FBS)) containing 5 \(\times\) 10\(^4\) HUVECs. The cells were allowed to incubate for 15 min before the chamber was filled with medium and the flow started (shear ca. 5 dynes cm\(^{-2}\)). The flow was maintained for 1 h at which point the disks were removed and placed into a 24-well plate. An MTS assay was then used to quantify the number of cells still adhering. Using appropriate controls with known cell counts, we determined that (1.5 \(\pm\) 0.3) \(\times\) 10\(^4\) cells remained on the uncoated disks, while (4.6 \(\pm\) 0.2) \(\times\) 10\(^4\) cells remained on the coated surfaces, which is over a 3-fold increase in retention \((p < 5 \times 10^{-5})\). The same 3-fold differential was again witnessed when a 10 % serum medium solution was used in place of the 2 % serum used earlier \((p < 5 \times 10^{-5} \text{; Fig. 3B})\). Even with this relatively short preflooding incubation time, the coating increased the cellular binding to the point that almost no cells were lost once the flow began. Additionally, the coating was not affected by the adsorbed proteins such as albumin. The flow rate chosen is at the upper end of venous flow,\(^{[44]}\) and we are currently designing a system that can simulate arterial flow rates. Importantly, these initial experiments show the benefits of an endothelializing coating to improve cellular attachment under physiological conditions.

In order to confirm integrin-RGD binding was responsible for the attachment difference observed, we performed experiments using soluble tetrapeptide sequences to preblock integrins on the HUVECs. Eight Ti disks were treated with \(12\) for 2 h followed by 2 h of a BSA block. In the meantime, trypsinized cells were placed into a medium solution supplemented with either 2 mm soluble RGDS or arginine–glycine–glutamic acid–serine (RGES) for 1 h. These treated cells were then placed onto the \(12\)/BSA-coated Ti disks for 15 min and the flow was started and maintained for 1 h. As before, an MTS assay was used to quantify the number of cells still present on the discs. As shown in Figure 3B, treatment of the HUVECs with RGDS \((n = 4)\) prior to placement on the coated Ti disks resulted in a significant decrease in the number of cells adhered, when compared to HUVECs treated with RGES \((n = 4)\) \((p < 5 \times 10^{-4})\). Only the soluble RGDS binds to the cell integrin in this modified competitive binding assay. The results of the experiment verify the RGD motif as the primary method through which coating \(12\) adheres HUVECs. These results are encouraging because integrins are one of the primary methods through which cells receive biological cues from their environment and these specific integrin interactions regulate many aspects of cellular biology. Although, we have initially opted to use the ubiquitous but well-studied RGD domain, more specific endothelial cell binders may be substituted in the future to create a coating with precise phenotype binding control. These studies are ongoing.

In summary, we have designed, synthesized, and characterized a peptide-based implant coating that mediates endothelialization of a synthetic surface. As a prototypical example we examined Ti metal, which lacks a natural ability for directing cellular activity, necessitating surface modification to ensure proper in vivo integration with the surrounding biological matrix. The peptide coating is readily applied to the Ti surface under mild aqueous conditions and shows functionality under dynamic conditions, thereby lending itself to the idea of a prohealing approach to device modification. The materials and methods to coat metal surfaces described herein
should be applicable to other medically important metallic surfaces (e.g., stainless steel) and biologies including proteins (e.g., growth factors) and cells (e.g., osteoblasts) because the coating process is mild, can be performed in aqueous solution at neutral pH, and the interfacial coating system is modular with readily interchangeable domains. An interfacial biomaterial coating for a metallic surface (IFBM) that recruits, attaches, and organizes endothelial cells mimics several functions of the ECM, and thus can potentially guide endothelialization on the coated surface. Continued research in this area will foster the development of next-generation biologically inspired tissue-integrated devices for fields ranging from cardiology to implant-based drug delivery.

Experimental

Phage Display Metallic Substrates: Ti 6Al-4V beads of ca. 300 μm in diameter were obtained from Dynomet (Washington, PA). All beads were exhaustively washed in detergent, then rinsed alternately with deionized water and ethanol prior to use. Beads were characterized by using SEM and XPS. SEM showed a smooth bead surface and XPS revealed the presence of a TiO₂ layer on the surface (Ti2p/O1s = 0.313).

Ti Disk Manufacture and Cleaning: Ti disks were lathe cut from a Ti 6Al-4V rod (12.7 mm diameter; 3.8 mm high). Disks were cleaned by first rubbing with 400 grit sandpaper. The disks were then transferred into a beaker containing acetone and shaken for 10 min. This was followed by 10 min of shaking in 95 % ethanol. The disks were then placed into deionized water and cooled for 20 min (20 % amplitude, 1 s on/1 s off) using an ultrasonic probe (Sonics & Materials, Vebra Cell, Newton, CT). The disks were then placed into a 40 % nitric acid solution for 30 min. Finally, the disks were treated with the ultrasonic tip using the same settings, but for only 10 min, and transferred into a 70 % ethanol holding solution for use in future experiments. Characterization with AFM, SEM, and XPS of the cleaned disks was performed. AFM and SEM analysis showed a surface with roughness of rms = 150 nm, most likely caused by the cutting and subsequent sanding (Fig. S1). XPS analysis showed the presence of a TiO₂ layer on the surface (Ti2p/O1s = 0.27 ± 0.003) as well as the presence of some surface impurities (C1s/Ti2p = 2.91 ± 0.01).

Ti Coated Class Substrates: Glass microscope slide covers were coated with a 20 nm layer of Ti (cp Ti from Kurt J. Lesker Co., Clair-an oxidized layer (Ti2p/O1s = 0.48 ± 0.07) was performed. AFM and SEM analysis showed a surface with roughness of rms = 150 nm, most likely caused by the cutting and subsequent sanding (Fig. S1). XPS analysis showed the presence of a TiO₂ layer on the surface (Ti2p/O1s = 0.27 ± 0.003) as well as the presence of some surface impurities (C1s/Ti2p = 2.91 ± 0.07).

Phase Separation Phage Panning of Ti: In the first round of screening, 25 Ti beads were placed in a 1.4 mL eppendorf tube. 10 μL of the SCA6 library and 200 μL 1 % BSA were added to the beads and incubated for 1.5 h with gentle shaking. Next, 400 μL of a 15:1 solution of dibutyl phthalate: cyclohexanone (Sigma Aldrich; St. Louis, MO) was added to the phage-bead solution. The tube was inverted several times then rested on the bench to allow phase separation. Both the organic and aqueous phases were removed from the tube, leaving only the Ti beads and strongly bound Ti phage. The phage-bead complexes were transferred to a fresh eppendorf tube and the phage eluted from the beads. Next, 400 μL of concentrated exponential phase TG-1 e-coli cells were added to the solution of phage for the first step in infection. This suspension was incubated at 37 °C for 30 min, then for 1 h at 37 °C with shaking. Next, the cell suspension was decanted from the eppendorf into a 13 mL conical tube and incubated for 4 h at 37 °C with shaking to amplify the phage population. This amplified population was then used as the input for the next round of screening, carried out in an identical fashion to that of the first round. Three such rounds were completed.

On Phage ELISA: Freshly amplified phage particles were added to each vials containing the Ti beads. After incubation for 1 h at RT with gentle shaking, HRP-anti-M13 Ab conjugate (Amer sham-Pharmacia, Piscataway, NJ), 1:5000 diluted in PBS-T, was then added for 1 h. The vials were washed five times with PBS-T buffer, followed by addition of 1 % BSA. One hundred microliters of ABTS was then added and incubated at RT for 15 min. The absorbance was read using a plate reader at 405 nm. A plot of absorbance versus log peptide concentration yielded a sigmoidal plot, the inverse of the half point was used as the affinity constant.

Contact-Angle Determination: The experimental substrate was placed onto the stage of a Kruss DSA 100 contact Angle tester. A bead of 2.5 μL of nanopure water was added and the resulting image was captured and analyzed using drop shape analysis software (Kru ss). Reported angles were calculated using a tangential method and averaging the angles from both the left and right sides of the droplet.

Binding-Affinity Calculations: Affinity constants were calculated by performing a modified ELISA assay. First, successive dilutions of the biotin-termininated synthesized peptide (1 mg mL⁻¹ stock in 10 % DMSO/PBS) were made in PBS-T (0.05 % PBS-Tween 20). 100 μL of each concentration was then spotted onto the appropriate metal substrate (Ti disks or slides) and incubated at RT for 1 h. After five washes in PBS-T, 100 μL of 1:2500 streptavidin-HP (Promega) was added to each well and incubated for 1 h at RT. Unbound streptavidin was removed during five PBS-T washes, then the chromogenic substrate ABTS was added and incubated on the bench for 10 min. Supernatants were transferred to a 96-well ELISA plate (CoStar) and the color intensity indicative of bound peptide was measured using a plate reader at 405 nm. A plot of absorbance versus log peptide concentration yielded a sigmoidal plot, the inverse of the half point was used as the affinity constant.
Metal-Binding Phage Isolation and Peptide Translation: Phage from the final rounds of pandering were plaque-purified and assayed by ELISA to confirm their binding to the Ti substrate. The DNA of these positive ELISA clones (so labeled due to an absorbance reading at or above 0.50 units) were then isolated using a mini-prep kit (Qiagen) and sequenced following the conventional chain-terminator method.

Peptide Residue Analysis: Expected expression levels for the random domain residues were calculated using the reduced genetic code produced by the synthetic NNK codons. Observed frequencies for the 16 variable positions on the 10 Ti binding sequences were tabulated and an expression ratio was calculated (Obs/Exp) for all 20 amino acids (Fig. S2 in Supporting Information).

Peptide Synthetic Chemistry: Peptides were commercially synthesized by solid-phase peptide synthesis techniques. The resultant peptides were purified to at least 95 % purity and included high-performance liquid chromatography and mass spectrometry analysis. The biotinylated peptide for affinity constant calculation was synthesized using a C-terminal biotin attached through the epsilon amide of a lysine residue.

Cell Attachment Immunostaining and Phase-Contrast Microscopy: One Ti-coated glass slide was treated by placing a 100 L droplet of DPBS in its center, and a second and third slide had a 100 L droplet of EBM-2 medium coating 12 or 13 (0.1 mg mL\(^{-1}\)). After all slides were blocked for 10 L at RT for 2 h. The slides were blocked with a 0.1 % BSA solution for an additional 2 h at RT. The BSA solution was removed and the slides were washed with DPBS prior to the addition of 20000 HUVECs in 100 L EGM-2 medium (Cambrex, system contains 2 % serum and a variety of growth factors) supplemented with 2 mm of RGDS (American Peptide Company; 0.867 mg mL\(^{-1}\)) or RGES (American Peptide Company 0.895 mg mL\(^{-1}\)). Cells and solutions were kept at 37°C for 1 h with occasional brief mixing and then added on top of the coated Ti disks.

Cell Proliferation Quantification: HUVECs were cultured using the conditions listed above. Cells were grown to passage three before use. Thirty cleaned Ti disks were treated with either a concentrated 1 mg mL\(^{-1}\) solution of 12 (n = 15), or PBS (n = 15) for 1 h at RT. The solutions were removed and 10 disks (5 in each treatment) had 200 mL of the plain medium listed above added. The other 20 disks had 20000 cells in 200 mL of the above medium added. The acellular groups, and five in each treatment group of the cellular disks had 40 mL of the MTS assay solution (Promega CellTiter 96 AQueous One Solution Cell Proliferation Assay) added and were incubated for 1 h (37°C and 5 % CO2). 200 mL aliquots were taken from the disk surface and were put into a polystyrene plate for absorbance reading at 492 nm. The remaining 10 disks were incubated for 24 h before 40 mL of MTS was added, incubated for an additional hour, and aliquots were read on the platerader. The results from three disks, each from a separate group, could not be determined as the media leaked off of the disks before the aliquots could be taken, leaving a total of 27 usable data points. By subtracting the background readings (acellular disks) from the resultant data points (20000 and unknown cell amounts) and determining the time needed to exponentially double the absorbance reading, and therefore proportionally the cell count, doubling times for the IFBM treated and untreated Ti surfaces were calculated.

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