Inhibition of Staphylococcus epidermidis Biofilms Using Polymerizable Vancomycin Derivatives

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Abstract

Background Biofilm formation on indwelling medical devices is a ubiquitous problem causing considerable patient morbidity and mortality. In orthopaedic surgery, this problem is exacerbated by the large number and variety of material types that are implanted. Metallic hardware in conjunction with polymethylmethacrylate (PMMA) bone cement is commonly used.

Questions/purposes We asked whether polymerizable derivatives of vancomycin might be useful to (1) surface modify Ti-6Al-4V alloy and to surface/bulk modify PMMA bone cement to prevent Staphylococcus epidermidis biofilm formation and (2) whether the process altered the compressive modulus, yield strength, resilience, and/or fracture strength of cement copolymers.

Methods A Ti-6Al-4V alloy was silanized with methacryloxypropyltrimethoxysilane in preparation for subsequent polymer attachment. Surfaces were then coated with polymers formed from PEG(375)-acrylate or a vancomycin-PEG(3400)-PEG(375)-acrylate copolymer. PMMA was loaded with various species, including vancomycin and several polymerizable vancomycin derivatives. To assess antibiofilm properties of these materials, initial bacterial adherence to coated Ti-6Al-4V was determined by scanning electron microscopy (SEM). Biofilm dry mass was determined on PMMA coupons; the compressive mechanical properties were also determined.

Results SEM showed the vancomycin-PEG(3400)-acrylate-type surface reduced adherent bacteria numbers by approximately fourfold when compared with PEG(375)-acrylate alone. Vancomycin-loading reduced all mechanical properties tested; in contrast, loading a vancomycin-acrylamide derivative restored these deficits but demonstrated no antibiofilm properties. A polymerizable, PEGylated vancomycin derivative reduced biofilm attachment but resulted in inferior cement mechanical properties.

Clinical Relevance The approaches presented here may offer new strategies for developing biofilm-resistant orthopaedic materials. Specifically, polymerizable derivatives of traditional antibiotics may allow for direct polymerization into existing materials such as PMMA bone cement while minimizing mechanical property compromise. Questions remain regarding ideal monomer structure(s) that confer biologic and mechanical benefits.

Introduction

Biofilms have been referred to as the “hallmark characteristic of periprosthetic joint infection” [8]. One report
suggests many cases of presumed aseptic prosthetic loosening may actually be the result of undetected biofilms [45]. The antibiotic concentration required for bactericidal activity against sessile organisms can be several orders of magnitude higher than for planktonic bacteria [20], the rule of thumb being 1000 to 1500 times higher [14]. This places a premium on extensive wound irrigation and débridement at the time of surgery followed by lengthy antibiotic therapy.

A classic strategy used in orthopaedic infections, including those of biofilm origin, relies on loading poly(methylmethacrylate) (PMMA) bone cement with antibiotics that elute in vivo [4, 12, 13, 17, 18, 24, 26, 27, 29, 35, 37, 41, 42, 49, 50, 66]. Antibiotic-loaded bone cements tend to release entrapped pharmaceuticals in a burst-type fashion and, hence, are effective at delivering high local levels of an antibacterial agent for a short postimplantation period, usually a few days [4, 12, 37, 41, 66]. When loaded at high dose, effective antibiotic levels may be maintained in surrounding tissues for several months [1, 30, 41]. However, high-dose loading compromises mechanical behavior of bone cements [12, 26, 35] likely through direct physical changes in material architecture related to the addition of nonpolymerizable species. Antibiotic-loaded PMMA may continue to elute therapeutic agents for weeks or months at suboptimal levels. These problems may be partially overcome by adding water-soluble fillers that increase cement porosity and facilitate antibiotic release, and vacuum processing techniques can lead to more homogenous cement compositions [9, 43, 46]. Unfortunately, bone cement has a surface quite suitable for colonization, and although antibiotic loading can reduce biofilm formation, organisms are still able to grow, for example, on gentamicin-loaded bone cements [33, 61].

Additionally, there is much literature dedicated to polymeric surface modification of metal oxides, including those of titanium, with the goal of blocking protein adhesion and bacterial attachment [15, 22, 28, 40, 54]. Derivatives of poly(ethylene glycol) (PEG) are common in these antibiofouling and antibiofilm strategies [11, 15, 19, 22, 28, 40, 54, 62, 65]. PEG blocks the nonspecific adsorption of proteins, cells, and cellular debris through mechanisms generally attributed to its hydrophilic character and high surface mobility (steric repulsion) [2, 36, 52]. It is particularly attractive in biomaterial systems because PEG is nontoxic [2] and well tolerated at high doses [64]. Polymerizable derivatives of traditional therapeutics potentially offer a convenient advantage over other compounds in that they can be polymerized to implant materials such as Ti-6Al-4V alloy [38] or copolymerized with polymeric biomaterials to provide surface-active antibiotic moieties [39].

We asked whether (1) vancomycin-PEG(3400)-acrylate [VPA(3400)] polymerized to Ti-6Al-4V alloy reduces bacterial attachment with respect to PEG controls; (2) whether the antibiofilm effects of VPA(3400) and two vancomycin-acrylamide derivatives when individually copolymerized with PMMA bone cement would be statistically superior to vancomycin loading in terms of reduced adherent biofilm mass; and (3) whether polymerizable antibiotics when incorporated into PMMA would yield cement constructs with compressive mechanical properties similar to PMMA homopolymer and statistically superior to vancomycin-loaded PMMA.

Materials and Methods

This study involved the synthesis of three polymerizable vancomycin monomers followed by experimentation along three lines: (1) coating of Ti alloy with a subset of these monomers followed by assessment of bacterial surface attachment; (2) incorporation of monomers into PMMA bone cement followed by assessment of biofilm proliferation on composite surfaces; and (3) incorporation of monomers into PMMA bone cement followed by compressive mechanical testing (Fig. 1).

All monomer synthesis, purification, and characterization have been described previously [38, 39]. In a slight modification of those procedures, VPA(3400) was purified by dialysis only. This modification was necessary because of the large quantities of material required for PMMA bone cement biofilm experiments. We obtained three polymerizable vancomycin derivatives. These compounds are illustrated (Fig. 2).

The procedure for functionalizing surgical-grade Ti-6Al-4V alloy bar stock for subsequent polymer attachment has also been described in detail elsewhere [38, 44] but is illustrated briefly (Fig. 3). To form a comonomer solution for polymerization to silanized Ti-6Al-4V, 40 mg VPA(3400) (estimated to be approximately 50% product with residual PEG[3400]-acrylate and some high-molecular-weight byproducts) was dissolved in 320 μL Ar-purged dimethylsulfoxide containing the photoinitiator 2,2-dimethoxy-2-phenyl acetophenone at a concentration of 1 mg/mL along with 80 μL PEG(375)-acrylate. This formulation results in a calculated PEG(375)-acrylate:VPA(3400) ratio of approximately 100:1. When nonantibiotic controls were desired, we omitted the VPA(3400). Surfaces were modified by photoinitiated polymerization of 15 μL of a comonomer solution under 45-mW/cm² intensity, collimated, broad-range, ultraviolet light (Hg arc-lamp centered at 365 nm) for 900 seconds using a Hybralign Series 200 mask alignment system (Oriel Instruments, Stratford, CT) by techniques detailed elsewhere [31, 38, 56–58]. A 3-day
wash protocol [38] was used to elute any detectable residual monomer, unfunctionalized vancomycin or loosely entangled polymer chains.

To verify the surfaces were appropriately silanized, x-ray photoelectron spectroscopy (XPS) was used with a 45° incident monochromatic x-ray beam to characterize

**Fig. 1** The flow chart illustrates a basic research design having three separate arms of inquiry following chemical synthesis of polymerizable vancomycin derivatives. The number of samples of each material type examined are shown by “n”.

**Fig. 2** Vancomycin was modified at the primary amines illustrated. Both PEG-acrylate and acrylamide polymerizable derivatives of vancomycin were synthesized. The PEG-acrylate derivative had a 3400 g/mol PEG spacer. For VA-1 and VA-2, the numeral refers to the site of modification, primary or secondary amine, respectively. R₁ is classically referred to as the vancomycin V₃ position, and R₂ is termed the X₁ position.
oxidized and silanized surfaces; SEM was also used as a qualitative adjunct to explore the surface features of oxidized surfaces (Fig. 4).

Vancomycin was loaded into “C-ment 1” PMMA bone cement (Jorgensen Laboratories, Inc, Loveland, CO) at a concentration of 1 g vancomycin to 40 g bone cement powder (2.5 wt%, a standard, prophylactic loading dose) or at a concentration of 4 g to 40 g bone cement powder (10 wt%, a high, therapeutic loading dose). VPA(3400), VA-1, VA-2, and PEG(3000)-acrylate were loaded at 10 wt%. Cylinders were constructed by first dissolving the desired additive in water and mixing with bone cement powder to create a slurry. The slurry was then flash-frozen using liquid nitrogen and lyophilized to recover a homogeneous powder. Bone cement cylinders were polymerized by addition of monomer liquid according to the manufacturer’s directions. Cylinders of dimensions 10 mm x 5 mm (height x diameter) were formed using a Teflon mold.

Substrates (polymer-coated Ti alloy) were placed in 50 mL brain heart infusion (BHI) medium, and the medium was inoculated with Staphylococcus epidermidis ATCC 35984 (clinical, biofilm-forming isolate). BHI medium was used in previous studies of polymerizable vancomycin derivatives [38, 39] and, for consistency, was used here. Samples were incubated on a rotary shaker at 100 rpm at 37°C. Polymer-coated Ti samples were allowed to incubate for 24 or 72 hours. Twenty-four-hour samples were removed, rinsed with copious amounts of water to wash away unbound organisms, and prepared for SEM. Seventy-two-hour specimens were inspected visually. The SEM preparatory procedure consisted of 1-hour fixation in 10% (v/v) phosphate-buffered formalin (Fisher Scientific, Kalamaoo, MI) followed by an ethanol dehydration series: 15 minutes in 50:50 ethanol:H2O, 15 minutes in 75:25 ethanol:H2O, 15 minutes in 95:5 ethanol:H2O, and 30 minutes in 100% ethanol. The samples were then placed in hexamethyldisilazane (Electron Microscopy Sciences, Hatfield, PA) for 1 hour 45 minutes, removed, frozen, lyophilized, and gold-coated. SEM images were obtained at 750x magnification using a JSM-6480LV system (JOEL Ltd, Tokyo, Japan). A minimum of 10 random images was taken of each sample coupon by panning the microscope to non-predicted regions about the test coupon and then quantifying adherent organisms. Organisms were not visible during the panning procedure, only when the microscope came to a complete stop. Adherent Staphylococcal organisms were counted, and an average surface density was calculated for each coupon using area measurements from ImageJ Version 1.33u software (National Institutes of Health, Bethesda, MD).
Institutes of Health, Bethesda, MD). This approach was repeated for each Ti sample of varying surface type.

PMMA cylinders were incubated analogously except they were transferred to fresh media every 22 hours. Twenty-two hours was chosen as a time point for media collection as this marked the onset of bacterial suspension turbidity, and there was concern for inappropriate bacterial death with incubation under turbid conditions. At 22, 44, 66, and 88 hours, cylinders were collected, freeze-dried before adherent biofilm mass was determined gravimetrically (biofilm dry weight) to the closest 1/10 milligram using a standard analytical balance and a modification of methods described elsewhere [59]. We used 4-day growth data for comparing surface types. Vancomycin-loaded cylinders released active antibiotic for the first 22 hours at levels sufficient to inhibit bacterial growth in the incubation vessel as these vancomycin-containing vessels did not go on to reach turbidity if allowed to incubate further. However, if fresh media were supplied, bacterial growth commenced over subsequent time intervals. Accordingly, for vancomycin-loaded samples, reported time points were adjusted by subtracting 22 hours and represent hours after observed burst elution.

PMMA and PMMA-composite compressive mechanical properties were measured using an MTS 858 MiniBionix II system (MTS Systems Corp, Eden Prairie, MN) with a platen speed of 1 mm/min. Stress strain data were analyzed external to the system software. Compressive modulus was calculated as the slope of the elastic deformation region using a standard linear least squares fit. Yield strength (stress at the transition between elastic and plastic deformation) was calculated using a 0.002 offset method. Resilience was calculated as the area under the stress-strain curve up to yield strength. Fracture strength, when applicable, was taken as the stress at catastrophic failure.

We determined by SEM differences in the average number of adherent S. epidermidis organisms between Ti alloy surfaces coated with PEG(375)-acrl (n = 6) and...
VPA(3400)-PEG(375)-acrl (n = 7) using the Student’s t-test. We determined differences in adherent biofilm dry mass between PMMA cylinders (n = 11), PMMA cylinders incorporating vancomycin at 2.5 wt% (n = 3), vancomycin at 10 wt% (n = 10), VPA(3400) at 10 wt% (n = 10), VA-1 at 10 wt% (n = 5), VA-2 at 10 wt% (n = 9), or PEG(3000)-acrylate at 10 wt% (n = 9) using the one-way analysis of variance (ANOVA) in combination with the Tukey Test for pairwise comparisons. We determined differences in compressive modulus, yield strength, resilience, fracture strength, and strain at fracture between PMMA coupons (n = 5), PMMA coupons incorporating vancomycin 10 wt% (n = 5), VPA(3400) 10 wt% (n = 5), PEG(3000)-Acrl 5 wt% (n = 5), PEG(3000)-Acrl 10 wt% (n = 5), and VA-2 10 wt% (n = 3) using the one-way ANOVA in combination with the Tukey test for pairwise comparisons. Levine’s test was used to check the assumption of equal sample variances, and the Kolmogorov-Smirnov test along with visual examination of normal probability plots was used to verify data normality for all ANOVA analyses.

Results

In the 24-hour growth experiment to assess the initial attachment of organisms to the polymer-coated Ti alloy surfaces the SEM observations followed by quantification of adherent organisms suggested the VPA(3400) copolymer reduced (p = 0.007) the number of organisms by approximately fourfold relative to the PEG(375)-acrylate surface (Fig. 5). The bare oxide surface was covered with a confluent layer of organisms, making enumeration impractical. In a separate 72-hour growth experiment with oxidized Ti-6Al-4V alloy, PEG(375)-acrylate coated alloy, and VPA(3400)-PEG(375)-acrylate coated alloy, the bare

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**Fig. 5A–E** (A) Three Ti alloy discs exposed to biofilm-forming Staphylococcus epidermidis culture for 66 hrs are shown. The composition of the surface coatings (top surface only) are illustrated (arrows). (B) The same three discs are shown from a second perspective. Bare oxide surface is completely covered with biofilm. The coated portion of the PEG(375)-acrylate disc is visually clean. The coated portion of the VPA(3400)-PEG(375)-acrylate disc is visually clean as well. These figures clearly demonstrate the antibiofilm properties of PEG-type coatings. (C) A scanning electron microscopy (SEM) image of S. epidermidis biofilm at ×5000 magnification is shown (66 hours in culture). Cocci are apparent as is glycocalyx. (D) An SEM image at ×750 magnification shows attachment of S. epidermidis on PEG(375)-acrylate-coated Ti alloy (24 hours in culture). Images of this sort (without obscuring glycocalyx) were used to quantify attached organisms on both PEG and VPA(3400) surfaces. The white spots are bacteria. (E) Both surfaces reduced the number of attached organisms. However, data suggest that having VPA(3400) in the copolymer reduces bacterial attachment more effectively than PEG(375)-acrylate alone.
oxide surface was completely covered with biofilm organisms and glycocalyx (Fig. 5A–B). Both polymer-coated surfaces were visually clear of bacteria.

In separate experiments with PMMA-antibiotic composites, after 4 days under growth conditions, the mass of adherent organisms and glycocalyx was reduced on 10 wt% VPA(3400)-PMMA and 10 wt% PEG(3000)-acrylate-PMMA relative to PMMA controls (p < 0.01 for both comparisons). Ten wt% VA-1 and VA-2 composites were associated with a similar mass of adherent organisms and glycocalyx on PMMA. Bacteria colonized and grew on high-dose vancomycin-loaded PMMA (10 wt% vancomycin), and biofilm mass showed an exponential increase with time (Fig. 6A). Low-dose or high-dose loading of vancomycin did not reduce biofilm mass. However, both 10 wt% VPA(3400)-PMMA and 10 wt% PEG(3000)-acrylate-PMMA had less biofilm than high-dose vancomycin cement (p = 0.08 and p = 0.09, respectively) (Fig. 6B).

We observed adverse mechanical effects of adding vancomycin to PMMA bone cement and also the potential mechanical property advantages of adding a polymerizable antibiotic derivative to PMMA (Table 1; Fig. 7A). Vancomycin loaded at 10 wt% in PMMA bone cement decreased the compressive modulus by 20% (p = 0.01), yield strength by 16% (p < 0.0001), resilience by 17% (p = 0.02), fracture strength by 41% (p < 0.0001), and strain at fracture by 35% (p = 0.02). Conversely, VA-2 did not decrease any mechanical property and rather increased (p = 0.1) the compressive modulus 13% with respect to PMMA. Relative to PMMA, VPA(3400) decreased the compressive modulus by 36% (p < 0.001), yield strength by 38% (p < 0.001), and resilience by 43% (p < 0.001). Relative to PMMA, PEG(3000)-acrylate incorporated at 10 wt% decreased the same mechanical properties to a similar degree (Table 1).

Discussion

Biofilm-forming organisms continue to be a central concern in orthopaedic procedures which involve the placement of metallic hardware in conjunction with PMMA bone cement. We present initial work to evaluate the usefulness of vancomycin-PEG-acrylate and vancomycin-acrylamide species for the inhibition of S. epidermidis biofilm formation on both Ti-6Al-4V orthopaedic alloy and on PMMA bone cement. We evaluated: (1) whether VPA(3400) polymerized to Ti-6Al-4V orthopaedic alloy reduces bacterial attachment; (2) the antibiofilm effects (reduced biofilm mass) of VPA(3400) and two vancomycin-acrylamide derivatives when individually copolymerized with PMMA bone cement; and (3) the compressive mechanical effects of adding polymerizable antibiotics to PMMA bone cement.

We acknowledge limitations to our experiment. First is that the biofilm growth conditions presented may not reflect those encountered in vivo. A biofilm bioreactor might offer a more sophisticated approach to simulating growth conditions [3, 10, 34]. However, it is inherently difficult to define the complex milieu of an infection site, and the assays used here were sufficient to define differences between material types. Second, the traditional method of quantifying adherent biofilm is by measurement of viability by prior desorption (ultrasonication) and subsequent agar plating [25], though there is often considerable variability in the absolute bacterial count obtained by different methods [48]. Quantification by SEM is often limited [25] secondary to problems of bacterial...
Table 1. Compressive mechanical properties for PMMA bone cement with various additives are shown. VA-2 composite properties are very similar to those of PMMA. Species with PEG functionalities lead to mechanical property compromise

| Material          | Compressive modulus (MPa) | Yield strength (MPa) | Resilience (J/cm³) | Fracture strength (MPa) | Strain at fracture (%) |
|-------------------|---------------------------|---------------------|--------------------|-------------------------|------------------------|
| PMMA              | 25 ± 1 (100%)             | 85 ± 3 (100%)       | 3.0 ± 0.2 (100%)   | 96 ± 13 (100%)          | 31 ± 7 (100%)          |
| Vancomycin 10wt%  | 20 ± 1 (80%)              | 71 ± 2 (84%)        | 2.5 ± 0.3 (83%)    | 57 ± 5 (59%)            | 20 ± 3 (65%)           |
| VA-2 10wt%        | 28.2 ± 0.4 (113%)         | 87 ± 1 (102%)       | 2.8 ± 0.3 (93%)    | 90 ± 2 (94%)            | 31 ± 3 (100%)          |
| VPA(3400) 10wt%   | 16 ± 3 (64%)              | 53 ± 2 (62%)        | 1.7 ± 0.2 (57%)    | 76 ± 8 (79%)*           | 34 ± 2 (110%)*         |
| PEG(3000)-Acrl 5wt% | 20.7 ± 0.5 (83%)       | 59 ± 2 (69%)        | 1.9 ± 0.1 (63%)    | 72 ± 6 (75%)*           | 33 ± 3 (106%)*         |
| PEG(3000)-Acrl 10wt% | 15 ± 1 (60%)            | 44 ± 4 (52%)        | 1.4 ± 0.2 (47%)    | 50 ± 7 (52%)*           | 27 ± 6 (87%)*          |

Asterisks indicate materials with potential microcracking.

Enumeration arising from bacterial agglomerates and matrix-embedded bacteria [25, 55, 60] and from the possibility that some of the biofilm structure will be lost during preparation [23, 63]. For this reason, our SEM quantification methods were limited to measurement of initial bacterial attachment where extensive glycocalyx was not present and where relative bacterial surface density was low. We expect the reported bacterial surface densities (Fig. 5) to be technique dependent as with other methods [48]. Dry mass has also been used to quantify biofilm proliferation [51, 59]; however, the gravimetric assay used here to evaluate adherent biofilm mass on PMMA surfaces does not provide adequate precision for discriminating between the effects of VPA(3400) and PEG(3000)-acrylate. Third, although we demonstrated various statistically significant effects attributable to polymerizable antibiotic species, whether these effects will have clinical importance is a question for future work.

In experiments with polymer-coated Ti alloy, VPA(3400) appears to add an additional level of protection to PEG-type coatings (Fig. 5). Surface-contact killing may contribute to the decreased number of adherent bacteria during the initial stages of biomaterial colonization. It is likely this period is critical in true physiological infections in which the bacterial load in the surrounding fluids is generally expected to be low. In comparison, a recently described antimicrobial technique involves the covalent attachment of an antibiotic (presumably a monolayer) to titanium [5–7, 21, 32, 47]. Building on surface modification methods described by Nanci et al. [44], the authors were able to attach vancomycin to titanium via two aminoethoxyethoxyacetate linkers. One study suggested Staphylococcus aureus attachment may be blocked using this surface-modification platform [7]. However, because aminoethoxyethoxyacetate linkers (sold commercially as “Mini-PEG”) effectively put a thin PEG layer on a surface (four ethylene glycol units), some of the observed antibiofilm properties may be related to the linker chemistry. Another study reported as few as four ethylene glycol repeats can provide antibiofouling properties for 3 weeks or longer with some cell types [22].

The data show incorporation of PEG moieties into the PMMA architecture retards but ultimately does not prevent biofilm adherence. This effect may be related to blockage of initial bacterial attachment, as suggested in other studies with PMMA [16, 53], but may be overcome as the surface is saturated with bacterial glycocalyx and cellular debris.
Neither of the vancomycin-acrylamide derivatives showed an antibiofilm effect. This behavior is consistent with previous observations [39] suggesting the PEG linker is critical for antibacterial activity once vancomycin derivatives are polymerized from solid substrates. In an alternative approach, a quaternary amine dimethacrylate (QADMA) species was copolymerized with PMMA bone cement, and antibiofilm properties against Escherichia coli were examined [16, 53], but no quantification of adherent organisms was undertaken. Data suggested QADMA blocked the attachment of Escherichia coli; whether this was related to the killing of microorganisms is less clear.

Mechanical testing of antibiotic-loaded bone cements (Table 1) confirms that adding vancomycin at 10 wt% decreases compressive modulus, yield strength, resilience, and fracture strength, which is consistent with previous reports [12, 26, 35]; for example, Klekamp et al. [35] showed that, in fatigue tests, loading vancomycin at 7.5 wt% decreased the number of cycles to failure by 50%. Substituting the acrylamide-modified vancomycin derivative VA-2 at 10 wt% restores all measured mechanical deficits and may actually increase the compressive modulus. These improvements likely occur because the acrylamide functionality copolymerizes with methacrylate groups during bone cement curing, whereas vancomycin does not. Conversely, vancomycin may leave gross structural cavities in the cement or act as a plasticizer. Unfortunately, VA-2 and VA-1 were not effective at blocking biofilm formation in the assay presented here, suggesting it was the PEG functionality that conferred antibiofilm properties and also mechanically compromised the cements. PEG is known to confer resistance to cellular or protein attachment [19, 22, 65], and the PEG tether is reportedly important for increasing surface-based activity of VPA-type species [39]. There may be some optimal number of ethylene glycol units that confers adequate antibacterial activity without sacrificing bone cement mechanical integrity.

Data with coated Ti alloy surfaces suggest copolymerizing a PEGylated vancomycin species, VPA(3400), with PEG(375)-acrylate is more effective than PEG alone at blocking S. epidermidis biofilms. The PEG spacer itself likely contributes to the antibiofilm effect, but SEM data indicate the pendant vancomycin molecule improves the antimicrobial effect under some growth conditions. Loading PMMA bone cement with certain polymerizable vancomycin derivatives may eventually be useful for retarding biofilm adherence without compromising mechanical properties, although the current formulations did one or the other, but not both. A vancomycin-acrylamide additive resulted in compressive mechanical properties almost identical to PMMA controls but did not inhibit biofilm growth. VPA(3400) reduced biofilm growth but compromised mechanical properties. Experiments described here should facilitate the development of new antibiofilm biomaterial surfaces.

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