Inhibition of bacterial attachment and biofilm formation by a novel intravenous catheter material using an in vitro percutaneous catheter insertion model

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Introduction: Despite sterile barrier precautions and vigorous skin antisepsis, percutaneous insertion of intravenous catheters has been shown to result in attachment to the catheter surface of bacteria residing in the deep structures of the skin. Such attachment poses the risk of biofilm formation and eventual catheter-related bloodstream infection (CRBSI). This study was undertaken to assess whether the non-coated surface treatment of a unique catheter material (ChronoFlex C® with BioGUARD™) could inhibit bacterial attachment and biofilm formation.

Methods: A novel in vitro model and fluorescence microscopy were used to compare two intravascular catheter materials with respect to bacterial attachment and biofilm formation. The control material was a commonly used polyurethane. The study material was a unique copolymer, treated so as to remove surface additives, alter hydrophobicity and create surface micro-patterning. Outcomes were assessed using both a membrane potential indicator and a cell death reporter with appropriate fluorescent channels. Thus, bacterial cells attached to the catheter surface (living and dead) were imaged without mechanical disruption.

Results: Both bacterial attachment and biofilm formation are significantly inhibited by the study catheter material. In fact, over 5 times more bacteria were able to attach and grow on the control polyurethane material than on the study material (P = 0.0020). Moreover, those few bacteria that were able to attach to the study material had a 1.5 times greater likelihood of dying.

Conclusion: Using a novel in vitro percutaneous catheter insertion model, ChronoFlex C with BioGUARD is proven to significantly inhibit bacterial attachment and biofilm formation as compared with a commonly used polyurethane catheter material.

Keywords: biofilm, CRBSI, CLABSI, catheter-related infections, catheter-associated infections, bacterial resistance

Introduction

In 1998, Livesley et al demonstrated that percutaneous insertion of catheters—despite rigorous aseptic technique, maximum barrier protection and chlorhexidine skin antisepsis—resulted in staphylococcal “impaction” (i.e., attachment) on the external surface of catheters in 16.7% of cases.1 Once attached, of course, numerous factors including material characteristics, type of bacteria and hematologic conditions mediate the progression from attachment to cell-to-cell adhesion, biofilm formation, maturation and eventually planktonic dispersion into the bloodstream.2–4 While disquieting, bacterial contamination upon insertion may well explain, at least in part, the observation by Crnich and Maki5 and Trimsit et al6 as to the similarity between central venous catheter (CVC) and arterial line bloodstream infection rates: namely, despite their obvious
differences, both are placed percutaneously and have similar average dwell times. It may further explain rising bloodstream infections rates (predominantly staphylococcal) when peripheral IV catheters are allowed to dwell an average of 6 days as opposed to shorter time periods.5,7,8

Livesley further demonstrated that catheter insertion through an introducer sheath can prevent skin contamination of the catheter’s external surface during insertion.1 However, introducer sheaths entail their own risks. If they do not peel away, the external surfaces of the sheaths may be expected to acquire bacteria during percutaneous insertion. On the other hand, if they do peel away, the to-and-fro movement of the sheath and catheter during sheath removal may well damage the vessel intima, provoking thrombophlebitis; the larger hole left by the sheath, once removed, may result in leakage and back bleeding, especially during computerized tomography power injection of the contrast media; also, when used for CVC placement, an open, unvalved introducer sheath certainly enhances the risks of bleeding and air embolism.10

Coating the catheters with antimicrobials, in “pre-CVC bundle” studies, has been shown to reduce associated bloodstream infections roughly to half.11 However, it is not without risk. Anaphylaxis to chlorhexidine-containing coatings has been reported, and with chlorhexidine usage on the rise, so goes chlorhexidine sensitivity and allergy.12

What would be ideal is a percutaneous catheter somehow resistant to bacterial attachment and biofilm formation, but devoid of special coatings or embedded substances. Unfortunately, polyurethane – the polymer out of which most intravenous catheters are now made – is the exact opposite of this ideal.13 Nouman et al have recently demonstrated that during natural aging, additives in commercially used polyurethanes bloom to the surface, and that these additives then “facilitate bacterial attachment”.14

The present study was undertaken to determine whether a novel catheter material (ChronoFlex C; AdvanSource Biomaterials, Wilmington, MA, USA), which undergoes proprietary processing intended to discourage bacterial attachment and biofilm formation, could outperform a commonly used polyurethane catheter with respect to bacterial attachment and biofilm formation, using an in vitro percutaneous catheter insertion model. The study catheter is neither coated nor impregnated; its base material is a unique copolymer consisting of both polyurethane and polycarbonate. According to the catheter manufacturer (Access Scientific, LLC., San Diego, CA, USA), it is treated in such a way as to remove surface additives, alter hydrophobicity and create surface micro-patterning – measures which have all been shown to discourage microbial attachment.14–17

Materials and methods

Bacillus subtilis NCIB 3610 trpC2 pheA1 espA::Pspac-hyepsA-0 (Bs) was selected, based on its evolutionarily shared attachment and biofilm formation mechanisms with numerous biofilm-forming bacteria.18–20 In other words, with respect to bacterial attachment, adhesion and biofilm formation, Bs is a nonpathogenic model organism for Gram-positive bacteria, including Staphylococcus aureus.21 B. subtilis is well characterized, allowing one to determine using fluorescent microscopy techniques whether failed biofilm formation is the result of impaired attachment, growth or matrix production.22

Control catheters were made of a polyurethane material: 18 guage (PowerGlide™; C.R. Bard, Inc., Salt Lake City, UT, USA). Study catheters were made of the copolymer ChronoFlex C (polycarbonate, polyurethane): 17 guage (note: no 18 guage is manufactured; POWERWAND™ Access Scientific, LLC.).

The percutaneous catheter insertion model consisted of a central Bs biofilm culture, an underlying pH-adjusted (pH=7.0) agar/nutrient substrate (~3–4 mm thick) and an orifice through the bottom of the Petri dish covered with a Parafilm® (Bemis NA; Neenah, WI, USA) (Figure 1). This model was designed so as to simulate skin, such that a catheter passing through it – from the bottom to top – must first penetrate the epidermal layer (the Parafilm), then the dermis (the agar) and, finally, enter the hypodermal layer (Bs colony) wherein pockets of bacteria are known to reside within the various deeper structures (eg, sweat glands or follicles) despite chlorhexidine antisepsis.23 All catheters were placed in this manner in accordance with the manufacturers’ directions for use.

Upon withdrawal of the needle (and dilator, in the case of study catheter), the internal lumen of the catheter was evacuated using pressurized, filtered air. (This was done in order to study the biofilm formed on the exterior of the catheter only, since bloodstream infections associated with short-term percutaneously placed catheters – as opposed to the catheters placed through introducer sheaths – are associated more commonly with external surface biofilm).24 Catheters were then cut into segments and incubated at 30°C for 48 hours. Thereafter, the catheter segments were exposed to Sytox™ (Thermo Fisher Scientific; Waltham, MA, USA); a commercially available dye that reports directly on cell death and to Thioflavin-T (ThT; Abcam, Cambridge, MA, USA); a Nernstian membrane potential indicator dye that reports on cell viability, allowed to incubate for 1 hour and imaged using a phase contrast microscope (Olympus IX81 or IX83): the yellow fluorescent protein (YFP) channel was used for Sytox and the cyan fluorescent
Protein (CFP) fluorescent channel for ThT. Thus, bacterial cells attached to the catheter surface (living and dead) were imaged without mechanical disruption, as often happens with “semi-quantitative” methods (eg, the roll plate technique).

Total fluorescence was then meticulously recorded over the entire external catheter surface of the distal tip of each catheter, for each indicator, over nine separate trial runs.

P-values were calculated using paired t-test for the means, signed-rank test for the medians and t-test for ratios for the geometric means (ie, folds). The test for normality was based on the Shapiro–Wilk test.

**Results**

**Bacterial attachment and biofilm formation**

In Table 1, Comparison 1 summarizes total CFP channel fluorescence – reflecting live bacterial attachment and biofilm formation – over the distal tip (2 cm) of both control and study catheters for all nine runs. The study catheter exhibited on average significant inhibition of bacterial attachment and biofilm formation as compared with the control catheter \( P = 0.0133 \) based on a paired t-test of the differences, 95%CL 4.578E9-2.915E10, mean difference=1.687E10.

On average, the amount of attached, biofilm-producing cells on the polyurethane control catheter was 5.407 times greater than on the study catheter (\( P = 0.0020 \)). Figure 2 (ThT uptake using CFP channel) illustrates typical catheter segments as seen with the CFP channel using phase-contrast fluorescent microscopy. Exuberant biofilm growth can be seen extending broadly over a large area of the control catheter’s external surface. In contrast, the study catheter can be seen to host a relative paucity of bacteria.

**Bacterial attachment and cell death**

In Table 1, Comparison 2 compares total YFP channel fluorescence – reflecting attachment and subsequent cell death – of both control and study catheters. Cell death was on average greater on the study catheter than on the control polyurethane catheter; however, this trend did not achieve statistical significance (\( P = 0.1533 \)). Nevertheless, on average, ~1.5-fold increase in bacterial cell death was observed on the study catheter as compared with the control catheter.

In sum, all catheters encountered actively growing \( B_{s} \) colonies upon “percutaneous” insertion. Yet, despite incubation in conditions favorable for attachment and growth, the study catheter exhibited significantly less bacterial attachment and cell death.

### Table 1 Total fluorescence

<table>
<thead>
<tr>
<th></th>
<th>Polyurethane</th>
<th>ChronoFlex C®</th>
<th>Difference</th>
<th>Lower 95% CL</th>
<th>Upper 95% CL</th>
<th>P-valuea</th>
<th>Normality testb</th>
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<tbody>
<tr>
<td><strong>Comparison 1: CFP channel (membrane potential indicator)</strong></td>
<td></td>
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<tr>
<td>Mean</td>
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<td>1.657E10</td>
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<td>SD</td>
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<td>1.159E10</td>
<td>0.103E10</td>
<td>3.063E10</td>
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<tr>
<td>Median</td>
<td>1.97E10</td>
<td>2.61E9</td>
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<td></td>
<td></td>
<td>0.0117</td>
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<tr>
<td>Geometric mean (fold)</td>
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<td>2.277</td>
<td>12.837</td>
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<tr>
<td><strong>Comparison 2: YFP channel (cell death reporter)</strong></td>
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<tr>
<td>Mean</td>
<td>2.405E11</td>
<td>1.733E11</td>
<td>0.672E10</td>
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<td>2.556</td>
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<td></td>
<td>0.1276</td>
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</table>

Notes: P-values reported on the mean row come from paired t-test. P-values reported on the median row come from signed-rank test. P-values reported on the geometric mean (fold) row come from the t-test of ratio (Poly/Chrono). *Test for normality is based on Shapiro–Wilk test.

Abbreviations: CEP, cyan fluorescent protein; YEP, yellow fluorescent protein; E, exponent base 10.
biofilm formation than the control catheter. Moreover, when bacteria were able to attach to the study catheter, they were 1.5 times more likely to die as compared with attached bacteria on the polyurethane control catheter.

**Discussion**

This study utilizes a novel in vitro model and fluorescence microscopy to compare two intravascular catheter materials with respect to bacterial attachment and biofilm formation. The control material is a commonly used polyurethane. The study material is a unique copolymer consisting of polycarbonate and polyurethane, treated so as to remove surface additives, alter hydrophobicity and create surface micro-patterning – measures intended to discourage bacterial colonization and bloodstream infection. In clinical use, catheters made of the study material have consistently yielded zero bloodstream infections despite extensive use in a variety of challenging clinical settings. However, none of these reports feature catheter-related bloodstream infection as a primary endpoint; also, all but one of these studies lack the rigor of a randomized controlled trial (RCT). Therefore, it seemed prudent to begin with a controlled in vitro trial comparing these two materials prior to embarking on an RCT.

The results show that both bacterial attachment and biofilm formation are significantly inhibited by the study catheter material. In fact, over 5 times more bacteria were able to attach and grow on the control polyurethane material as compared with the study material ($P=0.0020$). Moreover, those bacteria that were able to attach to the study catheter material had 1.5 times greater likelihood of dying – possibly due to their inability to congregate with other cells and form a viable community.

If confirmed clinically in an RCT, these results may have far-reaching implications for vascular access. Presently, despite enormous strides in infection prevention, catheter-related/catheter-associated bloodstream infection remains a major scourge in hospitals worldwide. For example, bloodstream infection rates for CVCs range from 1% to 3%; similar rates exist for arterial catheters, and percutaneously placed polyurethane midlines have recently reported bloodstream infection rates as high as 3.2%. Even peripheral IV catheters, which were once removed in 72–96 hours (thereby limiting the time for biofilm to reach maturity), have recently been reported to have an associated bloodstream infection rate of 1.64 per 1,000 hospital admissions. The vast majority of these catheters are made of polyurethane, the very material recently shown to bloom additives with normal aging – additives proven to “facilitate bacterial attachment”.

To the best of our knowledge, this study demonstrates for the first time how removal of the aforementioned additives, coupled with both micro-patterning and altered surface hydrophobicity (ie, the BioGUARD™ [Access Scientific, San Diego, CA, USA] secondary processes, performed by the catheter manufacturer), can inhibit bacterial viability on the surface of an extruded tube – specifically, an intravascular catheter. It has been known for some time that flat surfaces, when subjected to micro-patterning and altered hydrophobicity, can “create energetic conditions unfavorable for bacterial attachment and induce repulsive surface–bacterial interaction forces that impairs attachment and subsequent biofilm formation”. It has also been shown, again on flat surfaces, that increased hydrophobicity and decreased availability of contact area can contribute to a reduction in bacterial adhesion. The question has been how to translate these findings into real-world applications. Figure 3 shows a scanning electron micrograph of the study catheter’s surface micro-patterning. This surface appears to offer little purchase for one or two microbes, much less a microbial community. If our results are confirmed clini-
mentally, one can easily imagine how this technology might be put to use across the entire range of infection-prone intravascular devices, including all central venous access devices, midlines, extended-dwell catheters and so on.

**Limitations**

While the present study models percutaneous catheter contamination, it should be noted that other routes of contamination exist. For example, intravascular catheters can also become contaminated from hematogenous spread of microorganisms and from accidents of mishandling. Certainly, no direct inference should be made from the present study as to the other potential mechanisms of catheter infection.

Additionally, care should be exercised when extrapolating the in vitro results to in vivo clinical circumstances. Proteins and formed elements present in blood, for example, are not present in the experimental model. Further, the bacteria used in this experiment, *B. subtilis*, are not the actual pathogens encountered in usual device-related bloodstream infections. Nevertheless, a compelling literature attests to the shared characteristics of the study bacteria and more pathogenic species, such as *Staphylococcus aureus*, especially with respect to attachment and biofilm formation.18–21

A similar study for in vivo catheters following careful removal could confirm our findings. Also, a study looking at bacterial attachment and biofilm formation on the intraluminal surface of both control and study catheters would provide additional insights and information. Finally, a study comparing the study material to available central lines and arterial lines with regard to biofilm formation both in vivo and in vitro would also be very informative.

**Conclusion**

Using a novel in vitro percutaneous catheter insertion model, Chronoflex C with BioGUARD is proven to significantly inhibit bacterial attachment and biofilm formation as compared with a commonly used polyurethane catheter material. These findings may offer insight into the low bloodstream infection rates previously reported with catheters made of this material. A large-scale, randomized controlled clinical trial comparing bloodstream infections associated with extended-dwell catheters made of the study catheter material vs a polyurethane catheter material is now indicated.

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**Disclosure**

Steve Bierman MD is the Chief Medical Officer of Access Scientific, LLC. Pieter d’Arnaud is an employee of Instat Consulting, Inc. The authors report no other conflicts of interest in this work.

**References**


