

Wilson-Nieuwenhuis, JST and Dempsey-Hibbert, N and Liauw, CM and Whitehead, KA (2017)*Surface modification of platelet concentrate bags to reduce biofilm formation and transfusion sepsis.* Colloids and Surfaces B: Biointerfaces, 160. pp. 126-135. ISSN 0927-7765

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Version: Accepted Version

Publisher: Elsevier

DOI: https://doi.org/10.1016/j.colsurfb.2017.09.019

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Surface modification of platelet concentrate bags to reduce biofilm formation and transfusion sepsis.

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Abstract

Bacterial contamination of blood products poses a major risk in transfusion medicine, including transfusions involving platelet products. Although testing systems are in place for routine screening of platelet units, the formation of bacterial biofilms in such units may decrease the likelihood that bacteria will be detected. This work determined the surface properties of p-PVC platelet concentrate bags and investigated how these characteristics influenced biofilm formation. *Serratia marcescens* and *Staphylococcus epidermidis*, two species commonly implicated in platelet contamination, were used to study biofilm growth. The platelet concentrate bags were physically flattened to determine if reducing the surface roughness altered biofilm formation. The results demonstrated that the flattening process of the platelet bags affected the chemistry of the surface and reduced the surface hydrophobicity. Flattening of the surfaces resulted in a reduction in biofilm formation for both species after 5 days, with *S. marcescens* demonstrating a greater reduction. However, there was no significant differences between any of the surfaces following 7 days' incubation for *S. marcescens* and no significant differences between any of the surfaces following 7 days' incubation for *S. epidermidis*. The results suggest that flattening the p-PVC surfaces may limit potential biofilm

formation for the current duration of platelet storage time of 5 days. It is hoped that this work will enhance the understanding of how surface properties influence the development of microbial biofilms in platelet concentrate bags in order to devise a solution to discourage biofilm formation.

Keywords: Biofilms; MATH; Physicochemistry; Chemistry; Topography; Surface.

Introduction

Bacterial contamination of blood products poses one of the biggest risks of transfusion medicine.¹ This contamination is most commonly associated with transfusion of platelet concentrates (PC). This occurs in part due to the storage conditions within the blood bank used to maintain the platelets, which inadvertently provide ideal conditions for bacterial growth and proliferation. It is reported that approximately 1 in 2000-5000 PCs are contaminated.²⁻⁴ Whilst the reported incidence of culture confirmed sepsis is relatively low, at around 1 in 100,000, a higher incidence rate of 1 in 25,000 has been estimated, with higher rates likely due to under reporting of transfusion-related sepsis.^{1,2} Given that PC's are frequently administered to immunocompromised patients such as those with major medical issues in oncology, haematology, paediatric and bone marrow transplant patients, bacterial contamination and resultant sepsis can prove fatal.⁵ The mandatory use of bacterial detection systems within the national blood banks, such as the BacT/Alert microbial detection system minimise the likelihood of transfusing a contaminated unit. Yet, whilst these systems are extremely valuable, occasionally false negatives can occur due to low initial bacterial concentration in the PCs or due to the formation of bacterial biofilms within the PCs during storage.^{6,7} Even with these systems in places, storage time is not always extended from the 5-day safer period to the longer 7-day storage period.^{1,6}

Whilst aseptic techniques are used during the processing of blood products, three main sources of contamination exist: the venepuncture site during phlebotomy, transient donor bacteraemia and contamination during PC processing.^{1,8} Of these, the most common contaminants are typically skin flora surrounding the venepuncture site, contaminating the blood during collection.⁸ Most isolated contaminants are Gram-positive commensal skin flora species, with the two most commonly identified bacteria being Staphylococcus epidermidis (often only identified as a coagulase-negative staphylococci) or *Propionibacterium acnes*.⁸⁻¹⁰ Whilst it is more common for contaminants to be Gram-positive, Gram-negative bacteria are responsible for 60 % - 80 % of transfusion septic fatalities.^{5,11} Gram-negative species such *Serratia spp*, Escherichia coli, Salmonella spp and Enterobacter spp have all been implicated in severe adverse transfusion reactions.¹¹ Of these, *Serratia spp* has been shown as being able to remain culture negative even after 9 days of storage, leading to several severe or fatal septic reactions.¹² Biofilms form when individual bacterial cells adhere to a surface and to other bacterial cells, becoming embedded in a matrix of extracellular polymeric substance (EPS) in a complex and multifactorial process.¹³⁻¹⁵ The EPS is usually comprised of mostly polysaccharides with some DNA and proteins.¹⁶ Biofilm formation enables the bacteria to alter their gene expression whilst also altering cell growth, making the bacteria much harder for screening systems to detect.^{13,17} Further, if the bacteria are adhered to the surface of the platelet bag in the form of biofilms then less cells may be in planktonic form within the unit and therefore are not collected into the aliquot for testing, leading to false negatives.¹⁸

Initially bacteria are weakly adhered to a surface via Lifshitz Van der Waals forces, after which stronger adhesion is formed due to bacterial cell adhesion structures.¹⁵ Lifshitz Van der Waals forces are properties of both the bacteria and the surfaces, and understanding their interactions can elucidate how the bacteria interact at the cell:surface interface. By altering the surface to

produce changes in the surface properties, it is possible to alter how bacteria interact with a surface and potentially reduce bacterial adherence. As such, whilst it may be possible to develop more sensitive detection systems, modifying the platelet bag surfaces in a way so that bacterial cells are either unable or take longer to adhere to the platelet bag surface could play a role in reducing false negatives in already established systems.

An approach to reduce initial surface contamination, which may lead to a subsequent reduction in biofilm formation, is to modify the substratum surface topography.¹⁹ Although a number of studies have been carried out on the effect of surface topography on microbial retention, findings are often conflicting especially when macro topography (> 10 μ m), micro topography (< 10 μ m – 0.5 μ m) and nano topographies (< 0.5 μ m) may be present, all of which may affect initial conditioning film and bacterial attachment.²⁰

It is proposed that surface modification of the p-PVC platelet bags could reduce bacterial attachment, which would have two major advantages; (1) bacteria would be planktonic and therefore more likely to be collected in the testing aliquot for BacT/Alert screening, (2) bacteria would be unable to increase their virulence properties and slow their growth rate as occurs upon biofilm formation, hence increasing the likelihood of bacterial detection.

The aim of this work was to determine if a reduction in surface roughness of p-PVC platelet transfusion bag surfaces reduced the formation of *S. marcescens* or *S. epidermidis* biofilms.

Methods

Coupon Preparation

Unmodified Coupons

Plasticized poly(vinyl chloride) (p-PVC) platelet storage bags were obtained from Terumo BCT (USA). Coupons of 11 mm in diameter were punched from p-PVC platelet bags using an 11 mm disc punch (Agar scientific, UK) and the coupons were then sterilised.

Flattened Coupons

Strips of p-PVC were cut from whole platelet bags, 150 mm x 25 mm, and were placed between two sheets of kapton. A template made of polyethylene terephthalate (PET) was used to ensure the p-PVC was not flattened beyond the depth of the PET for consistency. This was placed in between stainless steel sheets, which was subsequently placed between two plates of steel which had been pre-heated to 150 °C in the hot press. The p-PVC / PET / Kapton and stainless steel sheets were placed into the hot press and 10 tonnes pressure was applied at 150 °C for 10 min. The plates were transferred to a cold press and cooled with 10 tonnes of pressure for 10 min. Circular coupons (11 mm diameter) were punched out of the flattened strips; these were sterilised immediately.

Sterilisation of Coupons

Coupons were placed into a 30 mL sterile universal containing 70 % (v/v) ethanol and vortexed for one min. The coupons were then soaked for 15 min in the ethanol. The ethanol was removed and the coupons were aseptically transferred to a sterile Petri dish containing sterile distilled water (dH₂O) to remove the ethanol before being aseptically moved across to another sterile Petri dish containing sterile dH₂O. The washing sequence was performed in triplicate to ensure all the excess ethanol was removed before the coupons were air dried in a class II microbiological cabinet.

Scanning Electron Microscopy

An overnight culture was prepared and the washed bacterial inoculum was diluted to an optical density (OD) of 1.0 at 540 nm. Ten microliters of bacterial inoculum was pipetted onto a coupon and dried in a class II cabinet for 1 h before being processed by placing into 4 % glutaraldehyde (made up in phosphate buffered saline) overnight. The coupons were removed from the glutaraldehyde using tweezers and rinsed with sterile dH₂O to remove the remaining

solvent. The coupons were dried in a fume hood for 1 h before an ethanol gradient of 10 %, 30 %, 50 %, 70 %, 90 % and 100 % made up from absolute ethanol diluted with sterile dH₂O was performed. Each coupon was left in each ethanol concentration for 10 min moving from the lowest ethanol concentration to the greatest. Once removed from the 100 % ethanol solution, the coupons were placed into a desiccator and were later attached to SEM stubs with carbon tabs prior to being sputter coated with a gold and palladium coating (Model: SC7640, Polaron, Au/Pd target, deposition time: 1.5 min) before being stored in a desiccator until imaged.

Surface Roughness and Optical Surface Profiles

Surface roughness (S_a) values were calculated from surface profiles measured using a ZeGage 3D Optical Surface Profiler (Zygo, USA) for all p-PVC coupons. Measurements were taken using 50 x magnification with a scanning distance of 50 µm from the centre position (n = 3). *Attenuated Total Reflection-Fourier Transform Infrared Spectroscopy (ATR-FTIR)*

ATR-FTIR was used to determine the molecular structures and chemical bonds of the untreated and modified p-PVC (Nicolet 380 FTIR with a Smart iTR attachment, Thermo Scientific, UK). Background spectra was captured prior to each measurement and spectra were acquired at room temperature using Omnic 5.2 software with each run made up of 16 scans and a resolution of 4 cm⁻¹. Analysis of each sample was performed in triplicate and the average spectra were reported (n = 3).

Physicochemistry of surfaces

Contact angles (θ) using HPLC grade water (BDH, UK), ethylene glycol or dioodomethane (Alfa Aesar, USA) were measured with a MobileDrop goniometer (Krüss GMBH, Germany). Both advancing and receding angles were determined, with five measurements of each chemical on each sample taken (n = 10). Fresh coupons were used for each solvent to ensure there was no cross contamination of solvents on the surfaces. The method of Van Oss *et al.*²¹ was used for calculating the surface energy (γ_s^{SE}) of the films from these measurements, according to the following equation:

$$(1 + \gamma_l)\cos\theta = 2\left(\sqrt{\gamma_s^{LW}\gamma_l^{LW}} + \sqrt{\gamma_s^A\gamma_l^B} + \sqrt{\gamma_s^B\gamma_l^A}\right)$$
(1)

where the subscripts *s* and *l* denote the surface energy of the solid and liquid respectively. The superscript LW denotes the Lifshitz-van der Waals components of the surface energy, and the superscripts *A* and *B* denote the Lewis acid and Lewis base parameters of the surface energy. The acid and base terms can be combined into the Lewis acid base (superscript *AB*) component of the surface energy:

$$\gamma_i^{AB} = 2\sqrt{\gamma_i^A \gamma_i^B} \tag{2}$$

Subsequently the overall surface energy was calculated as the sum of the Lifshitz-van der Waals and Lewis acid base components:

$$\gamma_i = \gamma_i^{LW} + \gamma_i^{AB} \tag{3}$$

The components of the surface energy were then used to assess the hydrophobicity, or Gibbs free energy of attraction between the surface and liquid (surface energies are denoted by subscript w), and were calculated using the following²²:

$$\Delta G_{sw} = -2\left(\left(\sqrt{\gamma_s^{LW}} - \sqrt{\gamma_l^{LW}}\right)^2 + 2\left(\sqrt{\gamma_s^a \gamma_s^b} + \sqrt{\gamma_l^a \gamma_l^b} - \sqrt{\gamma_s^a \gamma_l^b} - \sqrt{\gamma_l^a \gamma_s^b}\right)\right)$$
(4)

Tensile Strength Determination

Tensile strength of the PVC was determined using a Hounsfield H10KS tensometer running Qmat 5.53. Samples were run with a 4 mm width and a 200 mm/min speed (n = 5).

Bacterial Growth Conditions

Strains of *Serratia marcescens* (NCTC 9741) and *Staphylococcus epidermidis* (DSM 28319) were obtained from frozen pure culture stock, plated onto tryptone soya agar (TSA) (Oxoid, UK) and incubated at 37 °C overnight. New plates were streaked from freezer stock every four weeks to ensure the continuity of the bacteria was maintained for all experiments, with streaked plates stored at 4 °C. A single colony was removed from the agar plate and inoculated into a volume (10 mL) of sterile tryptone soya broth (TSB) (Oxoid, UK) which was incubated overnight at 37 °C with shaking (150 *rpm*). The bacterial culture was centrifuged at 2210 *g* for 12 min, the supernatant removed and the cells washed with 10 mL of sterile dH₂O before repeating. The cells were re-suspended in sterile dH₂O and diluted to an OD of 1.0 (\pm 5 %) at 540 nm using a spectrophotometer (Jenway 6305, UK). A serial dilution with spread plates was performed in duplicate with three separate bacterial broth cultures to obtain 1.25 x 10⁹ colony forming units per mL (CFU mL⁻¹) for *S. marcescens* and 1.44 x 10⁸ CFU mL⁻¹ for *S. epidermidis* at an OD of 1.0 at 540 nm.

Biofilm formation

A 1.8 mL volume of sterile broth was added to each well of a sterile 12-well plate (Nunc Non-Treated Multidishes, Thermo Fisher Scientific, UK) along with a sterilised p-PVC coupon, 11 mm in diameter, test side of the surface face up. The reverse end of a sterile swab was used to push the coupon to the bottom of the well. The 12 well plates were left at room temperature overnight to check for any contaminant growth and to check that all coupons were fully sterilised. An overnight culture was prepared and diluted to obtain 1.0 x 10⁸ CFU mL⁻¹. Two hundred microliters of the bacterial suspension was added to the appropriate wells to give a working concentration of 1.0×10^7 CFU mL⁻¹. The 12 well plates were parafilmed to prevent the media from evaporating from the plates. The plates were incubated at 22 °C for either 5 or 7 days with gentle shaking (68 *rpm*). Following incubation, all the media in each well was removed by pipetting, leaving just the coupon behind and the wells containing the coupons were washed by adding 2 mL of sterile dH₂O to each well; carefully against the side of the well so as not to disturb the biofilm formed on each coupon. The liquid was gently swirled around the wells manually for 5 s before pipetting out the liquid to remove any unbound planktonic cells. The 12 well plates and coupons were air dried in a class II cabinet for 1 h.

Crystal Violet Assay

One millilitre of 0.03 % w/v crystal violet solution was added to each well following drying, ensuring full coupon coverage, and left for 30 min. After the allotted time, the stain was pipetted out of each well and 2 mL of sterile dH₂O was added and manually gently swirled around for 10 s to remove the excess stain. This was repeated once and the coupons were removed after the second time and transferred to a fresh 12 well plate. To each well in the new plate, 2 mL of 33 % (v/v) glacial acetic acid solution was added and left for 15 min with manual agitation every 5 min to elute the bound crystal violet from the biofilm. The absorbance of the eluted stain in the glacial acetic acid was measured at 590 nm to determine the optical density, thus quantifying biofilm growth. The absorbance value obtained was doubled to convert it to OD per mL. When the absorbance was too high to be read, the remaining 1 mL in the well was diluted down with another 1 mL of 33 % (v/v) glacial acetic acid solution in the dilution and measured, with values obtained multiplied by four to account for the dilution. (n = 6)

Microbial Adhesion to Hydrocarbons (MATH) Assay

Cultures were grown at 37 $^{\circ}$ C overnight in TSB. Bacterial cells were harvested via centrifugation at 2210 *g* for 12 min and then washed 3 times using PUM buffer pH 7.1 (PUM

buffer; K₂HPO₄.3H₂O 22.2; KH₂PO₄ 7.26; urea 1.8; MgSO₄.7H₂O 0.2 g/L) and re-suspended to an OD 1.0 at 400 nm. A 1.5 mL volume of washed cells suspended in PUM buffer was added to a round bottomed test tube 15 mm in diameter. A 250 μ L volume of one of the test liquids (Chloroform; Hexadecane; Ethyl Acetate or Decane (BDH, UK)) was added to the suspension. Suspensions were incubated at 37 °C for 10 min to equilibrate. The test suspension was vortexed for 2 min before re-incubation for 30 min at 37 °C to allow separation of the lower aqueous phase. The lower aqueous phase was removed from the test tube and the OD determined at 400 nm. This was used with a calculation developed by Rosenberg *et al.*²³ to determine cell surface hydrophobicity;

$$adhesion = \left(\frac{1-A}{A\phi}\right) x \ 100$$
[5]

Where $A\phi$ was the optical density of the microbial suspension when measured at 400 nm before adding test chemical and A was the optical density measured at 400 nm of the extracted lower aqueous phase after mixing with the hydrocarbon. (n = 3)

Statistics

All results were presented as mean \pm standard deviation (SD) for tables or \pm standard error (SE) for the figures. Mean values were compared using analysis of variance (ANOVA) to determine significant differences between mean values at the 95 % confidence level (p < 0.05).

Results

Surface analysis

Upon initial analysis of the bags, it was determined that they were made of two sheets of p-PVC melded together. The analysis of the PC bags was carried out on the inside faces of the bags against which the PCs come into contact. SEM images demonstrated that one inside face of the p-PVC had a diamond patterned surface, which showed roughened features (rough surface) (**Fig**. 1a.). The rough surface showed pits along the edges of the diamond imprints of varying sizes. In comparison to this, the other inward facing side of the p-PVC polymer demonstrated a smoother, more featureless surface (smooth surface) (**Fig.** 1c.). Due to the unexpected surface features, flattening of the surfaces was performed to remove any surface features and determine whether this was enough to reduce biofilm growth. Post flattening of the p-PVC, the surface was demonstrated to be smoother still and without any discernible surface features (flat surface) (**Fig.** 1e).

Following quantification of the surface roughness, optical surface profiling demonstrated that the rough surface inside of the platelet bag had the greatest levels of differing surface topography (**Fig**. 2a). The mean surface roughness (S_a) of the rough surface varied between the diamond bottom, ridges and the sides of the diamonds which were demonstrated as having pits. A statistically significant reduction in S_a value was observed from the rough surface (3549.8 nm) to the smooth surface (1021.2 nm) and a further reduction was observed on the flattened surface (108.1 nm).

The ATR-FTIR spectra of the surfaces were dominated by the major absorptions expected in p-PVC (**Fig.** 2b). The most dominant were C-H stretching (centred at ca 2900 cm⁻¹), ester carbonyl stretching from the plasticiser (1740 cm⁻¹) and C-O bending of the plasticiser ester (1170 cm⁻¹). The C-Cl stretching vibration from the PVC itself at ca. 690 cm⁻¹ was also present. Two very small absorptions appeared between the phthalate ester carbonyl stretch and the C-H bending vibration of the PVC / plasticiser at 1458 cm⁻¹ (see **Fig. 2** inset). The smooth unmodified surface showed two small absorption peaks at 1576 cm⁻¹ and 1541 cm⁻¹, however, on the rough unmodified surface only one peak at 1539 cm⁻¹ was evident. Interestingly, both peaks were absent on the flattened surface. The peak positions match the antisymmetric $v_{as}COO^{-}$ stretching vibration of carboxylate groups in unidendate (1577-1575 cm⁻¹) and bidendate (1544-1540 cm⁻¹) coordination with calcium.²⁵ The carboxylate carbonyl absorptions

for the zinc stearate were simpler, just a single peak at ca 1540 cm^{-1.26} A schematic representation of the possible adsorbed structure of calcium stearate on the unmodified p-PVC surfaces is demonstrated indicating the presence of calcium/zinc stearates on the surface were present for the smooth, and to a degree the rough surfaces (**Fig.** 3a). However, after the flattening process the two carboxylate carbonyl absorptions were no longer present, suggesting that during the flattening process the calcium stearate either diffused more deeply into the p-PVC or preferentially adsorbed onto the kapton separating sheets (**Fig.** 3b).

Physicochemistry measurements between the smooth and rough surfaces demonstrated that there were no significant differences in the results (**Table**. 1). An increase in the Gibbs Free (ΔG) energy demonstrated that flattening of the surfaces resulted in the surface becoming less hydrophobic, and an increase in the overall surface energy (γ_s). However, the flat modified surface maintained similar Lifschitz van der Waals force ($\gamma_s LW$) values as the unmodified (smooth and rough) surfaces. The higher base energy (γ_s -) on all the surfaces demonstrated that the surfaces were electron donating rather than electron accepting. Flattening the p-PVC resulted in an increase in both the acid (γ_s +) and base (γ_s -) energies demonstrating the surfaces' ability to donate and accept electrons had both increased.

Tensile strength measurements of both surfaces determined that there was no significant difference between the average tensile strength of the unmodified surface (15.679 Mpa) and the flattened surface (15.145 Mpa) apart from the higher variability apparent for the modified surface. Further, there was also no significant difference between the elongation percentage for the unmodified surface (329.760 %) and the flattened surface (358.720 %) (Table 2).

Microbiology

SEM of the surfaces with the bacteria demonstrated that the density of *S. marcescens* and *S. epidermidis* was similar on all the surfaces (**Fig**. 1). The bacteria were observed to be clumped

in and around the pits of the rough surface with fewer cells spread around the other surface features (**Fig.** 1a and b). The number of cells present on the smooth surface was significantly reduced, and most cells formed small groups (**Fig.** 1c and d). On the flat surface, both species demonstrated very low coverage with only a few cells visible (**Fig.** 1e and 1f).

The MATH assay for *S. marcescens* (**Fig.** 4a) demonstrated a low affinity for the apolar nalkanes decane (6.33 %) and hexadecane (5.24 %), whilst having a moderate affinity for the acidic (Lewis-acid) solvent chloroform (43.97 %) and the basic (Lewis-base) solvent ethyl acetate (50.12 %). Pairs were chosen as having one polar and one non-polar liquid, both with similar Lifschitz van der Waals forces.³¹ The extremely low affinity (< 10 %) towards both non-polar hydrocarbons decane and hexadecane determined that *S. marcescens* was highly hydrophilic, which was supported by the bacteria having a greater affinity to the polar solvents chloroform and ethyl acetate.²⁸⁻³⁰ The stronger affinity for both of the polar solvents (chloroform and ethyl acetate) when compared with their non-polar counterpart (hexadecane with chloroform and decane with ethyl acetate) further indicated that *S. marcescens* was both a moderate electron donor and a moderate electron acceptor.²⁹ However, when compared in their pairs (chloroform and hexadecane against ethyl acetate and decane), the value for ethyl acetate and decane was higher, which demonstrated that the bacteria were better electron donors than acceptors.²⁹ This was further supported by the higher affinity to chloroform than to hexadecane.³¹

In contrast, *S. epidermidis*' MATH assay results (**Fig.** 4b) were significantly different to that of *S. marcescens*, demonstrating a high affinity to the apolar n-alkanes decane (98.21 %) and hexadecane (97.76 %) whilst also having a high affinity to the acidic solvent chloroform (98.44 %). The high affinity (> 55.00 %) to both non-polar hydrocarbons decane and hexadecane determined that *S. epidermidis* was highly hydrophobic. This was supported by the higher

combined affinity to the non-polar hydrocarbons (decane and hexadecane) when compared against the polar hydrocarbons (chloroform and ethyl acetate). The significantly higher affinity of *S. epidermidis* to the acidic solvent chloroform compared to the basic solvent ethyl acetate (30.67 %) indicated that the bacteria were strong electron donors. *S. epidermidis* as an electron donor was further supported by a higher combined value of chloroform and hexadecane compared to ethyl acetate and decane. *S. epidermidis* also had a moderate affinity to the basic solvent ethyl acetate indicating that whilst it was a strong electron donor it also had moderate electron accepting properties. This determined that *S. epidermidis* was highly hydrophobic and was a strong electron donor, with lesser electron accepting properties.

Biofilm growth of both *S. marcescens* and *S. epidermidis* following incubation in platelet storage conditions (22 °C) on the p-PVC for 5 or 7 days was assessed using a crystal violet assay (**Fig.** 5). Following 5 days' incubation, *S. marcescens* demonstrated a significant decrease in the amount of biofilm formed on the flattened (1.75) surface when compared to the rough (3.26) or smooth (2.94) surfaces (**Fig.** 5a). Whilst a decrease in the amount of growth on the flattened (2.60) surface compared to with that on the rough (3.66) surface was observed at day 7, it was less apparent than after 5 days, and no significant decrease was observed between the smooth and flat surfaces after 7 days (**Fig.** 5a). Similarly, *S. epidermidis* demonstrated a reduced growth on the flat (1.06) surface compared to the rough (1.96) and smooth (1.42) surfaces after 5 days of incubation, despite demonstrating significantly less growth overall (**Fig.** 5b). However, after 7 days' incubation there was no significant difference between the rough (2.13), smooth (1.94) or flat (2.04) surfaces (**Fig.** 5b).

Discussion

Surface Analysis

The surface properties of PC bags were characterised in order to determine the effect on biofilm formation. The microbial pathogens S. marcescens and S. epidermidis were used in the study as both species are regularly identified in contaminated PCs. The platelet bag surfaces were also flattened to determine whether alterations to the surface could potentially reduce biofilm growth. Observations of the topography of the surfaces via SEM and of biofilm growth, as quantified via crystal violet assay, demonstrated that the rougher surface features of the p-PVC platelet bag typically accumulated the most biofilm mass. This demonstrated a bacterial preference for a rougher surface to grow on, which may be due to an increase in surface contact area between the surface and the bacteria, increasing the strength of the initial adherence. The preference of bacteria for surfaces with defined features has previously been shown, where differing surface features altered bacterial retention.^{19,32} The surface roughness did not appear to affect the surface energies, as the surface energy properties of both the rough and smooth surfaces were similar. However, the flattened surface, which demonstrated no surface features did present differences in its surface energies, indicating that the flattening procedure changed the surface energy. The changes observed in the surface energy of the flattened surface led to the surface becoming less hydrophobic as well as increasing its overall ability to accept and donate electrons, whilst maintaining a preference for accepting electrons. These changes may reduce or increase bacterial adherence to the flattened surface, depending on the surface energy components of the bacteria themselves.

Analysis of the surface chemistry via ATR-FTIR demonstrated that most of the spectra for all the surfaces were similar. Whilst the C-CL stretching vibration of PVC would be a strong band in non-plasticised PVC, the high level of plasticiser in the samples caused absorptions from the latter to dominate the spectrum. Aromatic C-H stretching and deformation absorptions (3060 - 3100 cm⁻¹ and 1600 - 1580 cm⁻¹, respectively) were absent from the spectrum confirming that

a phthalate plasticiser had not been used in this particular platelet bag formulation. Absence of phthalates was further supported by the specific wavenumber of the ester carbonyl stretching absorption; phthalate carbonyl groups absorb at ca. 1730 cm⁻¹, whereas adipate and sebecate carbonyl groups absorb at ca 1740 cm⁻¹.²⁴

The two carboxylate carbonyl peaks (1576 and 1541 cm⁻¹), indicating presence of calcium / zinc stearates on the surface and / or within ca. 1.8 µm of the surface, were present for the smooth, and to a degree the rough surfaces. However, after the flattening process the two carboxylate carbonyl absorptions were no longer present, suggesting that during the flattening process the calcium stearate either diffused more deeply (i.e., deeper than ca 1.8 μ m) into the p-PVC or was preferentially adsorbed on to the polyester separating sheets, either way, calcium / zinc stearate could no longer be resolved by ATR-FTIR. An identical observation has been made by Bodecchi et al.³³ in ATR-FTIR studies on plasticised PVC that had been sterilised by β -irradiation, where it was argued that the stearate diffused further into the p-PVC. Combinations of calcium and zinc stearates have been reported to form complexes during the stabilisation of PVC,³⁴ with the combined carboxylate carbonyl stretching vibrations of the complex resulting in an absorption at 1600 cm⁻¹, but no such absorption was evident. This indicated that the stearates had been removed or diffused deeper into the substrate, rather than becoming complexed / combined as a result of elevated temperatures experienced by the p-PVC during the flattening operation. The spectrum of the rough surface was noisier than the smooth and flat surface due to the rough surface having an impaired contact with the internal reflection element of the ATR accessory.

The contact angle measurements demonstrated that the unmodified surfaces had lower surface energies than the flattened surface, and this observation indicated that a fraction of the stearates had bloomed to the surface of the unmodified substrates; the increase in surface energy of the

flattened substrate is corroborated by the absence of stearates on this sample. As the p-PVC had a relatively polar surface it may be that the bloomed calcium / zinc stearates were adsorbed with the carboxylate groups facing the surface, leaving the stearyl ($C_{17}H_{35}$) tails pointing away from the surface. The latter may account for the unidentate form of calcium stearate, absorbing at ca. 1576 cm⁻¹. If the local surface concentration of calcium stearate was high enough, the stearyl tails of the bidentate form (absorbing at ca. 1541 cm⁻¹) in the form of a bi-layer with Ca²⁺ ions in between, could incorporate the unidentate forms in wax-like self-assembled surface structures.³⁵ Such structures could have contributed to the reduced surface energy of the unmodified surfaces. The marginally higher values of γ^- , γ^+ and γ_s^{AB} obtained for the smooth unmodified surface, relative to the rough unmodified surface, may have been due to incorporation of some unidentate carboxylate, with the carboxylate group pointing away from the p-PVC, within the self-assembled structures. The rough unmodified surface showed only one carboxylate absorption at 1539 cm⁻¹, indicating the presence of bidentate calcium stearate, possibly with a small amount of zinc stearate. However, a uniform monolayer of stearate composed of stearyl chains, pointing away from the substrate in a self-assembled array, will have a very low surface energy of 22-24 mJ/m^{2 36}, and the latter here is clearly not the case and the implication is that the surface coverage with calcium / zinc stearate was not uniform. It may be that the stearates exist as micron-scale patches on the surface of the p-PVC.

Microbiology

When observing biofilm growth of both bacterial species on the smooth and flattened surfaces, and comparing this with growth observed on the rough surface, it was demonstrated that there was significantly more growth on the rough surface. The only exception to this was after 7 day's incubation with *S. epidermidis* when there was no significant difference between the surfaces. Since the chemistry and physicochemistry of the rough and smooth surfaces were the

same, this can only be due to a topographical effect with respect to the smooth surface. The similar growth on all the surfaces after 7 day's incubation with *S. epidermidis* may have been caused by a build-up of conditioning film and cells after the 5 days which meant that any surface topography, chemistry or physicochemistry effects on biofilm growth were negated. The effect of flattening the surface resulted in the most pronounced biofilm reduction with *S. marcescens* after 5 days. It is suggested that for *S. marcescens*, which demonstrated moderate electron donor and accepting properties, as well as highly hydrophilic properties, the reduction in hydrophobicity of the flattened surface reduced initial adherence of the bacteria to the surface.²⁸⁻³¹ However, the reduced biofilm growth could also be due to the lack of features present on the flattened surface which reduced surface area contact between the bacteria and the surface. A combination of both factors is most likely to be the cause of the lowered retention of *S. marcescens* to the flattened surface. However, it is difficult to determine which surface parameter had the most influence.

With respect to *S. epidermidis*, this organism was found to be highly hydrophobic and have strong electron donor properties. Therefore, it would not be unreasonable to assume that on the flattened surface which has reduced hydrophobicity and electron accepting properties, there would be an increase in total biofilm mass.^{28-30,37} However, our results demonstrated initial decreased biofilm growth on the flattened surface when compared to the rough after 5 days, which may be attributed to the lack of surface features.

Although data exists to support the theory that bacteria, which are generally negatively charged, will preferentially adhere to negatively charged surfaces, there are conflicting results in this area due to assays being carried out in a number of different ways and under different environmental conditions.^{28,38,39} However, it is clear from the present study that each reduction in surface topography resulted in a reduction in biofilm formation on each surface for both

bacterial species after 5 days incubation. However there was no significant differences between any of the surfaces following 7 days' incubation for *S. epidermidis* or between the smooth and flat surfaces following 7 days' incubation for *S. marcescens*. Over time, it appeared that the reduction of biofilm formation between the unmodified surfaces and the flattened surface was lessened as the reduction in growth was much greater at the 5-day point than what was observed after 7 days. This may be due to a build-up of adhered cells to the surfaces, which would gradually mask the underlying surface properties and allow adherence of further cells.

There is conflicting evidence regarding how significant the surface properties such as the roughness and physicochemistry are in determining bacterial retention, and hence biofilm formation. It has previously been determined by Katsikogianni *et al.* that under the conditions of flow on p-PVC surfaces, bacterial adhesion was influenced by the surface roughness.⁴⁰ However, it has also been demonstrated that p-PVC surface colonisation by *Pseudomonas aeruginosa* and *S. epidermidis* was independent of surface roughness and wettability.⁴¹ Furthermore, previous work completed by our group has demonstrated that whilst hydrophobic bacteria tend to adhere more to a hydrophilic surface and vice versa, some bacterial species can adhere better to a surface that has the same wettability.³⁹

Conclusion

The results demonstrated that by reducing the surface topography of PC bag surfaces, a reduction in biofilm formation by both bacterial strains was observed after 5 days particularly for *S. marcescens*. The, polar energies of the cells and the surfaces appeared to have little influence on bacterial adhesion as both organisms adhered to the rough surface in greater numbers than the other surfaces, which was the least polar. Therefore, it is suggested that reducing the surface topographies of the PC bags may be an effective strategy to reduce biofilm formation during PC storage.

Acknowledgements

This work was undertaken by Manchester Metropolitan University and was funded by the Sir Halley Stewart Trust. The views expressed within this report are those of the authors and not necessarily those of the trust.

Conflict of Interest

There are no conflicts of interest in this work.

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Table. 1. Physicochemistry results of the rough, smooth and flattened surfaces. Gibbs free energy (ΔG), surface energy (γ_s), Lifschitz van der Waals forces ($\gamma_s LW$), acid base components ($\gamma_s AB$), acid component (γ_s +) and base component (γ_s -).

	ΔG mJ/m ²	γ_s mJ/m ²	$\gamma_s LW$ mJ/m ²	$\gamma_s AB$ mJ/m ²	$\gamma_s + mJ/m^2$	γ_{s} -mJ/m ²
Rough	-52.98	52.37	48.87	3.50	0.44	7.33
Smooth	-50.15	49.61	45.65	3.96	0.54	7.54
Flat	-40.21	53.30	45.16	8.14	1.79	9.67

Table.
 2. Tensile strength and elongation measurements for the unmodified and flattened

 surfaces with standard errors.

	Tensile Strength		Elongation		
	(Mpa)		(%)		
Unmodified	15.679	± 0.257	329.76	± 19.373	
PVC					
Flattened PVC	15.145	± 1.531	358.72	± 25.019	

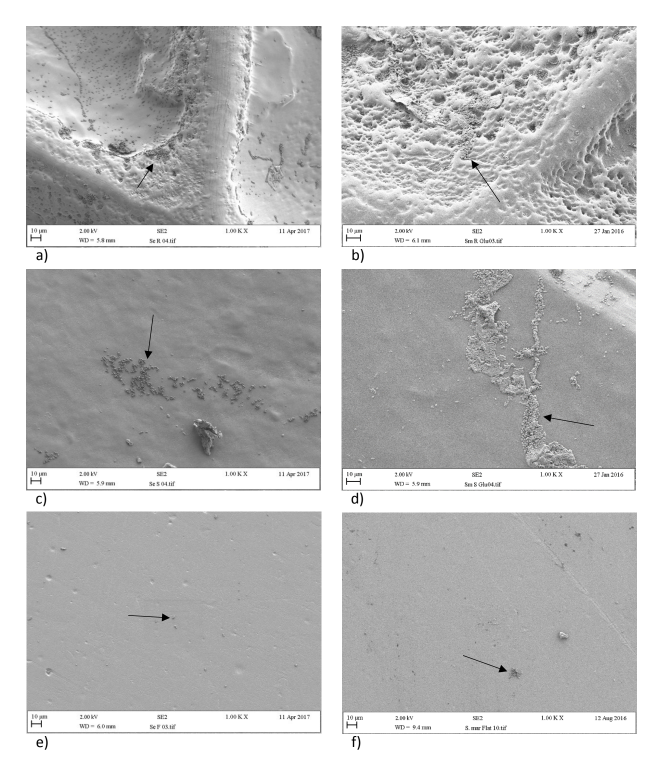


Fig. 1. SEM images of the inoculated p-PVC platelet bags a/b) rough (inside facing), c/d) smooth (inside facing) and e/f) flattened surfaces. Black arrows indicate bacterial cells with a, c and e representing *S. epidermidis* and b, d and f representing *S. marcescens*.

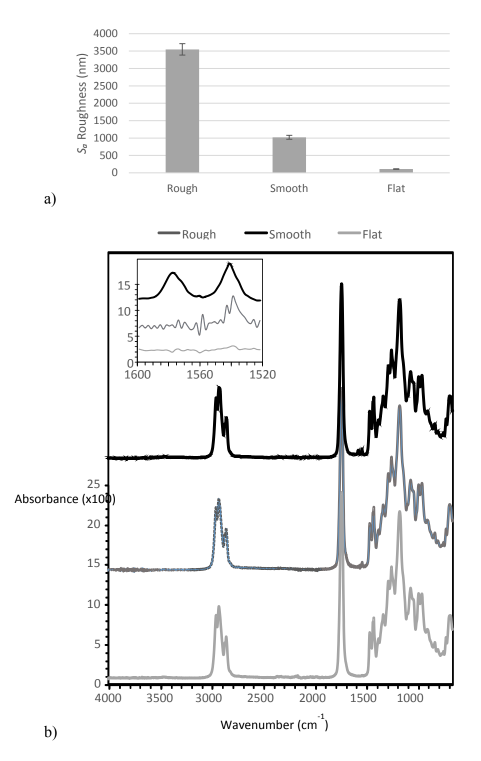


Fig. 2a. Surface roughness (S_a value) of rough, smooth and flat surfaces and b) ATR-FTIR spectra (average of n=3) of transfusion bag surfaces; unmodified rough side (dark grey), unmodified smooth side (black) and flattened surfaces (light grey). The carboxylate carbonyl antisymmetric stretching vibrations (at 1577 and 1544 cm⁻¹) are shown in the inset.

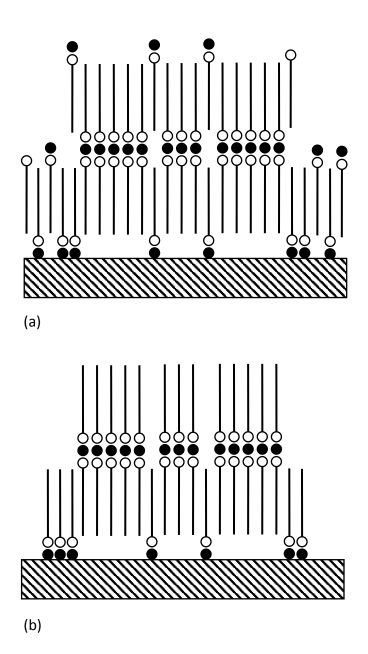


Fig. 3. Schematic representation of possible adsorbed structure of calcium stearate on the unmodified p-PVC surfaces. a) the presence of calcium/zinc stearates on the surface b) the carboxylate carbonyl absorptions are no longer present. The straight lines represent the $C_{17}H_{35}$ chains, the open circles represent the carboxylate group (COO⁻) and the closed (black) circles represent the calcium (Ca²⁺) ions (adapted from).³⁵

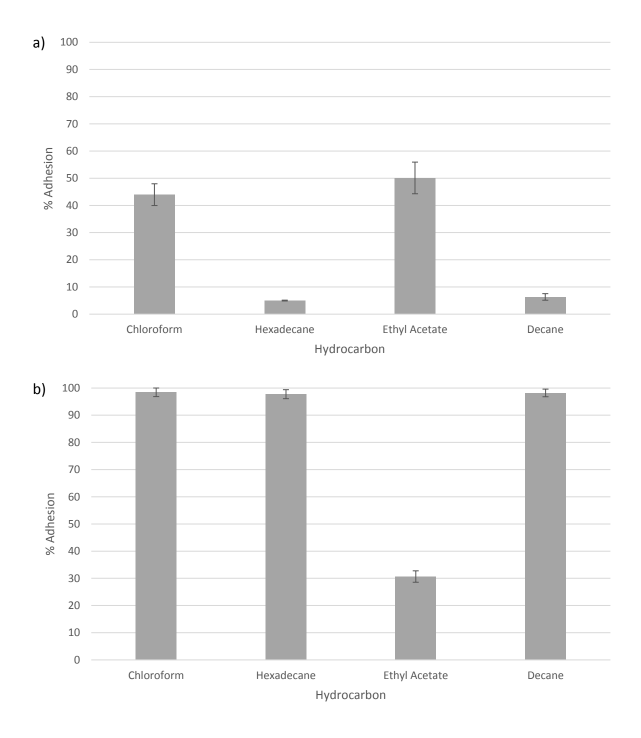


Fig. 4. MATH assays of a) *S. marcescens* and b) *S. epidermidis* demonstrating affinity to the apolar n-alkanes decane and hexadecane and to the solvents chloroform (acidic) and ethyl acetate (basic).

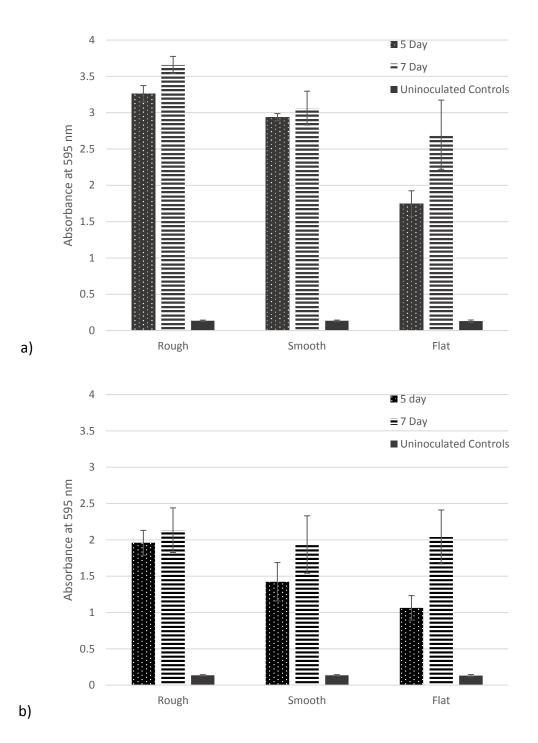


Fig. 5. Five and seven-day biofilm growth of a) *S. marcescens* and b) *S. epidermidis* on the unmodified (rough and smooth) and flattened p-PVC surfaces. The control surfaces followed the same procedure but were inoculated with sterile distilled water.