Secreted biofilm factors adversely affect cellular wound healing responses \textit{in vitro}

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Although most chronic wounds possess an underlying pathology, infectious agents also contribute. In many instances, pathogens exist as biofilms forming clusters surrounded by a secreted extracellular substance. We hypothesized that compounds secreted by biofilm bacteria may inhibit normal wound healing events including cell proliferation and migration. Conditioned media from two common bacterial species associated with chronic skin wounds and chronic tympanic membrane perforations, \textit{Staphylococcus aureus} and \textit{Pseudomonas aeruginosa}, were evaluated for their capacity to affect keratinocyte proliferation and migration. Additionally, proteomic analysis was performed to identify proteins within the biofilm conditioned media that may contribute to these observed effects. Biofilm conditioned media from both species inhibited proliferation in human tympanic membrane derived keratinocytes, whereas only biofilm conditioned media from \textit{S. aureus} inhibited migration. Human epidermal keratinocytes were found to be more sensitive to the effects of the conditioned media resulting in high levels of cell death. Heat treatment and microfiltration suggested that \textit{S. aureus} activity was due to a protein, while \textit{P. aeruginosa} activity was more likely due to a small molecule. Proteomic analysis identified several proteins with putative links to delayed wound healing. These include alpha hemolysin, alcohol dehydrogenase, fructose-bisphosphate aldolase, lactate dehydrogenase and epidermal cell differentiation inhibitor.

The primary function of the skin is to serve as a barrier to the external environment. Wound healing of the skin is a complex process involving cellular proliferation, migration and tissue remodelling leading to reestablishment of its primary function. Failure of a wound to heal in a timely manner may lead to further complications, including septicemia, chronic pain, prolonged hospitalization, amputations, and mortality\textsuperscript{1}. Chronic wounds in humans include wounds of the lower limb, due to an underlying pathology such as diabetes, pressure or venous hypertension, and chronic tympanic membrane perforations typically caused by chronic supplicative otitis media. Bacteria are present in chronic wounds and are proposed to play a role in delayed tissue repair\textsuperscript{2,3}.

It is considered that the bacteria present in chronic wounds not only exist in a planktonic form, but are also characterized by polymicrobial colonies of bacteria existing in communities known as biofilms\textsuperscript{4}. Biofilms consist of a cluster of bacteria adhering to a surface surrounded by an extracellular polymeric matrix\textsuperscript{5}. The function of the biofilm is varied and includes protection from the environment, nutrient storage and aiding in cell-cell communication. Consequently, gene expression and phenotype of bacteria in biofilm mode is different to that when in free living or planktonic mode\textsuperscript{6,7}. The presence of a biofilm in chronic wounds gives rise to several common features; they render the bacteria virtually impervious to antimicrobial agents and can sequester cargo such as DNA and antibiotics. Biofilms can remain in situ for many years and can reactivate when conditions become more permissive or when the body’s defenses are compromised. This biological resistance to treatment has significantly hindered patient recovery.

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to antibiotic treatments and affected tissues remain in a persistent inflammatory state. Additionally, it has been shown that the presence of a biofilm can delay wound healing. However, the mechanisms of delayed wound healing due to biofilms are poorly understood.

We hypothesize that bioactive compounds secreted by bacteria whilst in biofilm mode inhibit key wound healing mechanisms including cell proliferation and migration. Therefore, the aims of this research were to grow isolates of bacteria in biofilm mode and planktonic mode and produce conditioned media. We have selected perhaps the two most common bacterial species related to chronic wound colonization or infection, *Pseudomonas aeruginosa* and *Staphylococcus aureus*. In vitro cell culture was then used to compare the effects of secreted biofilm and planktonic compounds on key wound healing events, cell proliferation and migration. For this purpose we utilized human derived primary keratinocytes from two distinct areas, the tympanic membrane and the normal epidermis of the upper arm. Subsequently, proteomic analysis was undertaken to identify potential protein products responsible for any observed effects.

**Results**

**Untreated Samples.** *Proliferation.* A 10% dilution of biofilm conditioned media (BCM) and planktonic conditioned media (PCM) from *S. aureus* and *P. aeruginosa* were examined for their effect on cell proliferation *in vitro*. Included controls were a 10% dilution of tryptic soy broth (TSB) plus a positive control of cell culture media (DMEM +) only (Fig. 1). Compared to the TSB media control, PCM from *P. aeruginosa* significantly reduced proliferation (*p* = 0.0025) in hEKs, but PCM from *P. aeruginosa* did not (*p* = 0.7). However, the BCM of both bacterial species significantly reduced proliferation (indicated by *) in both cell types. Error bars are standard deviation of the mean with *n* = 4. Control is cell culture media only.

![Figure 1. Effect of conditioned media on cell proliferation.](image-url)

Figure 1. Effect of conditioned media on cell proliferation. BCM and PCM were tested for their effect on cell proliferation using hTMks (a) and hEKs (b). For hEKs the BCM was diluted further to 5% due to high levels of cell death. Additionally, compared to the TSB media control, PCM from *S. aureus* significantly reduced proliferation (*p* = 0.0025) in hEKs, but PCM from *P. aeruginosa* did not (*p* = 0.7). However, the BCM of both bacterial species significantly reduced proliferation (indicated by *) in both cell types. Error bars are standard deviation of the mean with *n* = 4. Control is cell culture media only.
Migration. The effect of diluted PCM and BCM on cell migration was examined (Fig. 2). In hTMks, PCM from S. aureus and P. aeruginosa plus BCM from P. aeruginosa had no significant effect on migration compared to the TSB media control (p = 0.27, 0.29 and 0.72 respectively) (Fig. 2a). However, the BCM of S. aureus significantly reduced migration to 9.8 ± 7.2% of the media-only control and approximately 13% of the TSB media control (p < 0.01) in this cell type.

In hEKs (Fig. 2b), the TSB media and PCM from P. aeruginosa seemed to accelerate migration to over 100% of the media-only control. However, the BCM from P. aeruginosa appeared to be highly cytotoxic.

Under microscopy, control hEKs stained with crystal violet exhibited the characteristic cobblestone appearance of keratinocytes (Fig. 3a). When treated in 5% P. aeruginosa BCM for the duration of the migration experiment (15 hours) a large proportion of the cells were killed and appeared contracted (Fig. 3b). The PCM of S. aureus (Fig. 3c) resulted in a reduced number of cells with altered morphology, which appeared to be losing close contact with their neighbours. Treatment with BCM of S. aureus resulted in highly reduced cell numbers (Fig. 3d). In addition, the presence of apoptotic cells appeared to be evident (Fig. 3d, arrow heads). It was therefore not possible to interpret these results in terms of altered migration. However, the observations indicate that hEKs are more sensitive to conditioned media than hTMks.

Effect of Heat Treatment. Only hTM-derived keratinocytes were used for analyses.

Proliferation. As the PCM has no observable effect on hTMks proliferation, it was excluded from this experiment. Heat treatment completely abrogated all anti-proliferative activity of S. aureus BCM, which showed no significant difference to the TSB media control (p = 0.97). However, heat treatment had no effect on the anti-proliferative activity of P. aeruginosa BCM (Fig. 4a).

Migration. Similar to proliferation, heat treatment of S. aureus BCM removed the ability to inhibit migration (Fig. 4b), which showed no significant difference to the TSB control (p = 0.23). Heat treated BCM from P. aeruginosa also demonstrated no significant difference compared to TSB control (p = 0.19), similar to its untreated form.

Effect of Filtration. Proliferation. Fractionating the samples using a 3 KDa filter showed all the anti-proliferative activity of S. aureus was found in the filtrate (p < 0.01), with no significant (p = 0.45) activity found in the flow through fraction (Fig. 5a). By contrast, P. aeruginosa activity was primarily in the flow through fraction with significant activity (p = 0.002) also detected in the filtrate. However, this activity was significantly less than that of the flow through (p < 0.01).

Migration. Only BCM from S. aureus was tested (Fig. 5b) due to lack of activity in P. aeruginosa conditioned media. Similar to the effects on proliferation, the inhibitory effect of S. aureus on cellular migration was found only in the filtrate, which was significantly reduced to 40 ± 26.1% of the TSB control (p = 0.019). No significant activity (p = 0.255) was recorded for the flow through fraction.

PAGE and Proteomics. PAGE analysis was performed on the BCM and PCM for both bacterial species. No protein bands were detectable in the TSB media control or PCM (gel not shown). In the BCM gel (Fig. 6), raw BCM from P. aeruginosa and S. aureus (Lanes 2 and 5 respectively) produced several discernable bands ranging in size from approximately 12 KDa to 80 KDa. Samples of flow through following fractionation were also included (lane 3 and 6) along with samples of the filtrate (Lane 4 and 7) for
P. aeruginosa and S. aureus respectively. All proteins were retained in the filtrate. Six identifiable bands from S. aureus (arrows) were removed from the gel for peptide sequencing and identification. These are summarized in Table 1. Identification of proteins from P. aeruginosa BCM was not performed due to the inability of heat treatment to abrogate its effect and lack of activity in the protein fraction. These results suggest that activity was not due to a protein.

**Discussion**

In this study, conditioned media derived from P. aeruginosa and S. aureus cultured as biofilms were examined for their ability to disrupt the normal wound healing processes of cell proliferation and migration. Keratinocytes from the tympanic membrane (TM) and the normal skin epidermis from the upper arm were chosen for the *in vitro* analysis due to their different healing patterns. When the dermis heals in response to injury, it undergoes sequential phases of haemostasis, inflammation, cellular proliferation, and finally cellular migration. In TM wound healing, the first two stages remain, but migration precedes proliferation.

For hTMKs, at the 10% dilution tested, BCM from S. aureus was capable of a modest reduction in proliferation, while it completely inhibited cellular migration. Conversely, the BCM of P. aeruginosa appeared to only affect proliferation with no significant effect on migration. However, the PCM from both species had no significant effect on either proliferation or migration. The hEKs appeared to be far more sensitive to the conditioned media. At 10% dilution, BCM from both species was highly toxic, effectively killing the hEKs in both migration and proliferation tests. However, when reduced to 5% dilutions a clear inhibitory effect on proliferation was seen. The BCM of both species significantly reduced the level of proliferation compared to the TSB control, with S. aureus BCM appearing to have a larger effect than that of P. aeruginosa. However, a comparison between S. aureus and P. aeruginosa BCM was found to be not significant (p = 0.083). In addition, the PCM of S. aureus also caused a significant decrease in proliferation, but was much less potent than the BCM. On the other hand, migration was more difficult to interpret. Both the PCM and BCM of S. aureus had a toxic effect on hEKs in addition...
Figure 4. Effect of heat treated BCM on cell migration and proliferation. Heat treatment completely removed the anti-proliferative effect of *S. aureus* but had no effect on *P. aeruginosa* BCM, which significantly reduced proliferation (a). For migration (b) heat treatment also removed the inhibitory activity of *S. aureus* BCM. *P* < 0.05 * and error bars are the standard deviation of the mean with *n* = 4. Control is cell culture media only.

Figure 5. Effect of micro-filtered BCM on cell migration and proliferation. For proliferation (a) the effect of *S. aureus* were confined to the filtrate. However, the majority of activity for *P. aeruginosa* was seen in the flow through (FT) fraction. The migration assay was performed only on *S. aureus* BCM (b) and not *P. aeruginosa* due to the lack of activity seen in that species. All the inhibitory activity was seen in the filtrate fraction. Error bars are standard deviation of the mean with *n* = 4. Control is cell culture media only.
to the BCM of *P. aeruginosa* (Fig. 3a–d). Therefore, meaningful migration results could not be firmly established. However, it should be noted that the PCM of *P. aeruginosa* had no significant effect compared to the TSB control, although both seemed to increase the rate of migration compared to the cell culture media only control.

Previous studies using cultured cells have examined the effects of PCM and BCM derived from *S. aureus*. Using wound scratch assays, both were found to significantly reduce gap closure in keratinocytes and fibroblast cells\(^\text{11,12}\). However, a difference in cell morphology was noted between PCM and BCM treated cells indicating potentially different mechanisms of action. Similarly, both PCM and BCM

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**Figure 6. Page separation of proteins.** Raw samples of BCM plus the microfiltered fractions of BCM were run on a 4–12% gradient polyacrylamide gel and stained with coomassie blue. Lane 1 = molecular size marker, 2 = *P. aeruginosa* raw BCM, 3 = *P. aeruginosa* flow through fraction, 4 = *P. aeruginosa* filtrate fraction, 5 = *S. aureus* raw BCM, 6 = *S. aureus* flow through fraction, 7 = *S. aureus* filtrate fraction. The arrows indicate bands that were extracted for peptide/protein identification.

| Band Number | Estimated Mass (KDa) | Sequence Matches | Mass (KDa) |
|-------------|----------------------|------------------|------------|
| 1           | 80                   | Lipase 1         | 76714      |
|             |                      | Bifunctional Autolysin | 92280    |
| 2           | 75                   | Transketolase    | 72256      |
| 3           | 60                   | No sequence match|            |
| 4           | 35                   | Alpha-hemolysin  | 35953      |
|             |                      | Alcohol dehydrogenase | 36304    |
|             |                      | Fructose-biphosphate aldolase class 1 | 33007    |
|             |                      | L-lactate dehydrogenase 1 | 34551    |
|             |                      | Glycerophosphoryl diester phosphodiesterase | 35289    |
|             |                      | Chaperone hchA    | 32141      |
| 5           | 28                   | 2,3-bisphosphoglycerate-dependent phosphoglycerate mutase | 26663    |
|             |                      | Triosephosphate isomerase | 27303    |
|             |                      | Epidermal cell differentiation inhibitor (EDIN) | 27663    |
|             |                      | Diacetyl reductase [(S)-acetoin forming] | 27227    |
|             |                      | Enterotoxin A     | 29656      |
| 6           | 25                   | Purine nucleoside phosphorylase | 25892    |
|             |                      | Glucosamine-6-phosphate deaminase | 22330    |
|             |                      | 3-hexulose-6-phosphate synthase | 22436    |

Table 1. Identification of *S. aureus* proteins. Bands excised from the PAGE gel were submitted for identification.
resulted in reduced viability in both cell types with the BCM being significantly more potent than PCM, which was also reflected in our studies using hEKS, but this did not seem the case in hTMks. For apoptosis, BCM and PCM possessed equal potency in fibroblasts. However, in keratinocytes, BCM treatment caused significantly higher levels of apoptosis than PCM, which was also suggested in our experiments. In the case of P. aeruginosa, much exists in the literature regarding the presence of biofilm in wounds and their effects. However, no similar in vitro studies into effects on proliferation and migration have been performed.

In an effort to determine which components of the BCM were exerting the effects seen, further analyses were performed on hTMks. Heat treatment and microfiltration on the BCM revealed the most likely cause for the effects seen for S. aureus was a secreted protein. This was based on heat treatment completely abrogating all S. aureus BCM effects, presumably due to denaturation. Similarly, after filtering the BCM with a 3 KDa cut-off, the majority of activity was preserved in the filtrate and not the flow through, which is suggestive of a retained, and relatively large molecular weight protein and not a small molecule. On the other hand, the effects caused by P. aeruginosa BCM appeared to be caused by a small molecule due to the ineffectiveness of heat treatment and the anti-proliferative effect being seen within the flow through fraction following microfiltration.

To further establish possible bioactive components of the conditioned media, SDS PAGE was performed. This revealed a complex pattern of secreted proteins for both P. aeruginosa and S. aureus BCM. As the data suggested only S. aureus activity was due to a protein(s), six of the more prominent S. aureus protein bands were excised from the gel and identified (Table 1). A number of identified proteins including transketolase, triosephosphate isomerase, purine nucleoside phosphorylase and glucosamine-6-phosphate deaminase are typically involved with essential metabolic processes with no known link to wound healing. However, of the proteins identified, several have been previously implicated in delayed wound healing and/or cell death and may be contributors to the effects seen in this study. These proteins include alpha hemolysin, a highly studied S. aureus cytotoxin that is particularly potent in lysing rabbit erythrocytes. In addition, it is also known to be dermonecrotic and neurotoxic. Alcohol dehydrogenase (AD) may also require further study. Currently, no explicit reports on the direct effects of this enzyme with respect to wound healing have been reported. However, there exists anecdotal evidence that suggests the presence of AD may have a negative impact on wound healing. A transcriptome microarray study on healing tympanic membranes in rats revealed AD to be the most potently downregulated gene in the first 12 hours following an induced perforation. Other studies have shown that ethanol exposure prior to injury may inhibit normal wound healing processes and have suggested the presence of ethanol itself plus its metabolites to be the likely cause. However, the effects of elevated levels of AD were overlooked on these occasions. Therefore, combined with current data it is possible to consider that the presence of AD may also cause a negative impact on early wound healing processes. Fructose-bisphosphate aldolase (ALDOA) is a glycolytic enzyme and has been shown to directly bind to actin filaments of the cytoskeleton and modulate polymerization. In keratinocytes, endogenous ALDOA is involved in forming actin based lamellipodia, which are structures important for cell migration. Therefore, its plausible that exogenous ALDOA from S. aureus biofilm may alter the actin polymerisation process leading to a downstream effect in cell migration. Lactate dehydrogenase is an enzyme that reversibly converts lactate into pyruvate. In a healing environment, lactate accumulation stimulates the production of growth factors such as VEGF and TGF-β and is capable of enhancing cellular migration. Similar to ALDOA, exogenous lactate dehydrogenase would effectively remove lactate from the wound environment and potentially reduce the pro-healing effects. Another interesting candidate identified was epidermal cell differentiation inhibitor (EDIN) protein. EDINs are able to ADP-ribosylate members of the Rho GTPase family causing their inactivation. Inhibition of the Rho GTPase Rac1 results in disturbed migration of keratinocytes both in vitro and in vivo. Additionally, EDIN produces a loss of barrier function in endothelial cells through actin cable disruption leading to the formation of microapertures. This provides direct access of the endothelium basement membrane to S. aureus bacterium. Furthermore, EDIN genes from S. aureus strains have also been found to correlate to higher grade diabetic foot ulcers. However, in this instance the presence of biofilm was not established.

In terms of wound healing there is the potential that biofilm secretions from different species may act in synergy. For example, there is a correlation between the in vitro biofilm - producing capacity of P. aeruginosa and S. aureus and unfavourable evolution after endoscopic sinus surgery. In addition, using a porcine wound model, it was shown that while individual infections with strains of S. aureus and P. aeruginosa were sufficient to delay wound healing, only a co-infection resulted in decreased expression of KGF1. Normal wound healing circumstances KGF1 acts through its receptor to increase migration and proliferation.

There were several limitations inherent in this study, primarily in controlling the many variables. For example, quantifying the amount of total protein present in each sample may have enhanced our results. However, this process was complicated by the presence of the trypsinised soy protein present in the bacterial culture media. Therefore, normalising for protein content was not possible. Similarly, controlling growth conditions of the planktonic and biofilm cultures were difficult to perform equally. This is due to the different growth rates and potential nutrient requirements for the different phases. Also, the process of freeze drying may result in the loss of compounds such as volatile organic chemicals. Additionally, it has also been noted that protein structure may be adversely affected.
In this study we have shown that BCM from two of the most common bacterial strains associated with infections have a profound effect on the wound healing processes of migration and proliferation. Additionally, we have shown a differential effect in keratinocytes of different anatomical sites. Furthermore, the data suggests that effects caused by *P. aeruginosa* BCM were due to a small cytotoxic molecule while the effects from *S. aureus* BCM were protein based. Proteomics on the PAGE separated *S. aureus* BCM was able to reveal a potential cocktail of proteins that may act either individually or in synergy to produce the effects seen. Our data would suggest that in addition to identifying which species of pathogens are present in a chronic wound or infection, a more detailed study of biofilm components is also necessary to elucidate how the presence of biofilm may delay wound healing.

**Methods**

**Bacterial Isolates.** *P. aeruginosa* isolates used in this study were obtained from a clinical case of canine otitis externa from Australia. Swabs were cultured onto 5% Columbia sheep blood agar (SBA) (Thermo Fisher Scientific, Scoresby, Vic, Australia) and incubated at 37°C and 42°C; overnight. Following incubation, cultures were stained by the Gram-method for determination of purity and morphology and an oxidase test was performed. *S. aureus* ATCC 29213 was routinely streaked out onto SBA and incubated at 37°C overnight. All bacterial isolates were stored at −80°C in brain heart infusion broth (BD Biosciences, Sydney, NSW, Australia) with 20% glycerol (Ajax Finechem (Thermo Fisher)).

**Conditioned Media.** *Planktonic and biofilm culture conditions.* Inoculated tryptic soy broth (TSB) (25 ml culture concentration of 1.0 × 10⁷ cfu/ml) was incubated in 50 ml tubes at 35 ± 2°C for 24 hours on a shaking incubator. Biofilms were formed using the MBEC™ high-throughput assay (MBEC™ Innovotech, Canada) according to the manufacturer’s instructions²⁸ with incubation temperature at 35 ± 2°C and times extended to 24 hours. In brief, this device consists of a 96-peg-lid (top plate) and a ridged trough (bottom plate). The top plate fits over the trough containing 10 ml of TSB. The inoculated device is placed on a rocking platform (Select BioProducts, Model SBS300-2, New Jersey, United States), which allows a shear force to be created forming equivalent biofilms on all the pegs. The inoculum is incubated at 35 ± 2°C for 24 hours in the MBEC device before collection.

**Supernatant preparation.** Planktonic and/or biofilm cultures were centrifuged at 4500 rpm for 20 minutes at 4°C (MPW-351R centrifuge, Warsaw, Poland), the pellet discarded, the supernatant filtered (Millex, PES 0.22 μm, Darmstadt, Germany) and frozen at −80°C. Samples were freeze-dried (Dynavac Freeze Dryer, Model FD12, Australia) and stored at −20°C. The freeze-dried conditioned media was then resuspended in an equivalent volume of DMEM+ (DMEM 4,500 g/l D-glucose, supplemented with 100 U/ml penicillin, 100 g/ml streptomycin (Invitrogen (Thermo Fisher)), and 10% FBS (Invitrogen)) before further dilutions.

**Cell Culture.** *Ethical Statement.* Isolation of human tympanic membrane keratinocytes was performed with approval from the St. John of God Healthcare Ethics Committee (Subiaco, Western Australia). Human normal skin keratinocytes were isolated from upper arm skin with approval from the University of Western Australian Human Research Ethics Committee (Approval # RA/4/1/5489). Patients signed a written, informed consent before intervention and specimens were treated anonymously. All collections and experiments were conducted according to the principles and ethical standards expressed by the Declaration of Helsinki.

**Human Tympanic Membrane Keratinocytes (hTMk).** Culture of primary human tympanic keratinocytes was performed as previously described⁴⁰. Briefly, fresh intact hTM tissue was obtained from consenting patients who underwent otological procedures requiring the permanent surgical removal of the tympanic membrane. All hTM tissue received was disease-free and considered morphologically normal. Small pieces of hTM tissue, approximately 0.25 mm² pieces. The pieces of skin were placed in 5 ml of EpigrowTM human epidermal keratinocyte complete media with 20 μl of fungizone (Gibco®, Invitrogen™, Carlsbad, California, USA) and 100 μl of kanamycin (Gibco®). Dispase (Life technologies™, Carlsbad, California, USA) was added at 0.0127 g per 5 ml (2.4 U/ml). This solution was incubated for 24 hours at 40°C. This method is a modified version of a protocol published by the CELLNTEC company (CellNTEC, 2011).

After 24 hours, the epidermis was separated from the dermis using sterile fine tip forceps and the pieces of epidermis transferred to a sterile petri dish with 500 μl of TrypLE™ Select (Gibco®). The epidermal tissue pieces were spread out flat on the surface of TrypLE™ Select and covered with a lid to prevent evaporation. After incubating for 20 to 30 minutes at room temperature, the petri dish was
tilted at a 30° angle and 2 ml of human epidermal keratinocyte complete media was added. Using sterile
forceps, epidermal tissue pieces were rubbed gently against the base of the petri dish in order to gen-
erate a single cell suspension. The resulting cell suspension was collected in a 15 ml centrifuge tube and
the previous step repeated to increase the yield of keratinocytes. The second preparation of a single cell
suspension was then added to the same 15 ml centrifuge tube. The combined cell preparation was then
centrifuged at 200 g for 3 minutes. The cell pellet was resuspended in 5 ml of human epidermal ker-
atinocyte complete media. Cells were cultured in a T25 tissue culture flask (Sarstedt Australia Pty Ltd,
Belmore, NSW, Australia)

**Filtration.** Samples of conditioned media were filtered through Amicon® Ultra-4 (Ultracel® - 3k)
Centrifugal filters (Merk Millipore Ltd, Tullagreen, Cork, Ireland) with a 3 KDa cutoff. Both flow through
and the filtrate were retained for analysis.

**Heat Treatment.** Undiluted samples of conditioned media were subjected to heat treatment of 75°C
for 20 minutes. The samples were allowed to cool to room temperature prior to use in downstream assays.

**Proliferation Assay.** Cell numbers following proliferation were calculated using a modified cell
attachment assay31. This method uses a crystal violet solution to stain cells attached to the culture plate.
After solubilizing the stained cells, the intensity of the solution is proportional to the number of cells
present.
Cells were seeded into the wells of a 96 well plate (Greiner Bio-One GmbH, Frickenhausen, Germany)
at a density of 3000 cells/well in 200 μl of DMEM+. The plates were incubated as described overnight
(approx. 16 hours) to equilibrate. The media was then replaced with conditioned media diluted to 10% in
DMEM+. Cells were incubated for 48 hours. The media was then removed and the cells fixed using
100 μl of 10% neutral buffered formalin (NBF). The fixed cells were washed twice in 100 μl PBS with
gentle shaking. The PBS was aspirated and the cells stained with 100 μl of crystal violet solution (0.1% w/v
crystal violet in 200 μm MES buffer, pH 6.0 filtered through a 0.22 μm filter) for 15 minutes. The
crystal violet solution was removed and the cells washed 3 times in 100 μl ddH2O with gentle shaking.
All the ddH2O was aspirated and the cells solubilized in 100 μl of 10% acetic acid for 10 minutes with
gentle shaking. Absorbance was read at 570 nm on an EPOCH microplate reader (BioTek Instruments Inc.,
Winooski, VT, USA)

**Migration Assay.** Using IBIDI cell migration culture inserts (Ibidi GmbH, Martinsried, Germany),
5 × 104 cells in a volume of 70 μl of DMEM+ were seeded into adjoining Ibidi insert chambers that had
been placed into the wells of a 48-well plate (Greiner). The cells were then incubated overnight at 37°C
in a humidified atmosphere with 5% CO2. The inserts were removed after 24 hours to reveal a well defined
gap approximately 500 μm wide. The wells were washed once with PBS containing fresh DMEM+ and
containing diluted conditioned media. Cells were photographed at 0 and 6 hours for hTKMs and at 0 and
15 hours for hEKs using a Nikon Eclipse TE2000-5 microscope. Multiple photographs were taken to ensure the entire “gap” was captured. The gap was then measured using Imagel software (NIH, USA) at 20 different points along the gap and averaged to provide the mean gap
distance. The relative distance of cell migration is calculated as:

% Gap closure = (Test (G-t₀ hr – G-t₆/₁₅ hr)/Control (G-t₀ hr – G-t₆/₁₅ hr)) X 100%

Where G-t₀ hr is the mean gap distance prior to cellular migration immediately following the removal of
the Ibidi culture insert (zero hour) and G-t₆/₁₅ hr is the mean gap distance at the 6 or 15 hour endpoint.
The results are expressed as a percentage of the control treatment, which contains no conditioned media
or diluted TSB media.

**Protein Analysis.** PAGE. BCM and PCM samples (300 μl) were vacuum desiccated and resuspended
in 20 μl of standard 1 × Laemmili loading buffer. Samples were heated to 95°C for 10 minutes and allowed
to cool to room temperature for 10 minutes. The tubes were centrifuged to collect the contents. Premade
4–12% gradient polyacrylamide gels were used for protein separation (Bio-Rad Laboratories, Hercules,
CA, USA). The gel cassette was loaded into a Protean II gel apparatus (BioRad) as per the manufactures
instructions with standard 1 × Tris/Glycine running buffer. The protein samples were loaded in the
wells along with a sample of Kaleidoscope size marker (Promega Corporation, Sydney, Australia) and
the gel run at 130 V for 30 minutes. After separation, the gel was removed from the cassette into a small
dish and stained with 100 ml of coomassie blue stain (BioRad) for 2 hours with gentle rocking. The stain
was decanted and the gel rinsed for several seconds in ddH2O to remove excess stain. 100 ml of destain
solution (50% ddH2O, 40% methanol, 10% acetic acid) was then added to the gel along with 2–3 small
pieces (2 cm³) of sponge and gently rocked overnight. The destained gel was then photographed.

**Sequencing and Identification.** Prominent bands in the polyacrylamide gel were excised and submitted
to Proteomics International (Perth, Western Australia, Australia) for identification. Protein samples
were trypsin digested and peptides extracted according to standard techniques32. Peptides were analysed
by electrospray ionisation mass spectrometry using the Agilent 1260 Infinity HPLC system (Agilent Technologies, Santa Clara, CA, USA) coupled to an Agilent 6540 mass spectrometer (Agilent). Tryptic peptides were loaded onto a C18 column 300 SB, 5 μm (Agilent) and separated with a linear gradient of water/acetonitrile/0.1% formic acid (v/v). Spectra were analysed to identify proteins of interest using Mascot sequence matching software (Matrix Science Inc., Boston, MA, USA) with Ludwig NR database. A match was considered positive if two or more peptides map to the same protein and were within the predicted size parameters as estimated by the standard on the acrylamide gel.

Statistics. ANOVA with a post hoc Dunnett’s test was used to analyse migration and proliferation results. Four replicates (n = 4) were used for each test. All comparisons were made against the TSB (vehicle) control. Significance was set at p < 0.05.

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**Author Contributions**
R.M. designed and executed experiments, analysed data and prepared the manuscript. H.J.W. and M.W. helped design experiments and prepare the manuscript. D.W. performed laboratory experiments and assisted in interpretation of results. HSW and R.O.H. aided experimental design, established the bacterial cultures, produced the conditioned media, interpreted results and aided in manuscript preparation.

**Additional Information**
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