# **Analyst**



View Article Online **PAPER** 



Cite this: Analyst, 2023, 148, 618



Wagar Ahmed, Da Emmanuelle Bardin, b,c Michael D. Davis,d 

Early detection of lung infection is critical to clinical diagnosis, treatment, and monitoring. Measuring volatile organic compounds (VOCs) in exhaled breath has shown promise as a rapid and accurate method of evaluating disease metabolism and phenotype. However, further investigations of the role and function of VOCs in bacterial-host-stress response is required and this can only be realised through representative in vitro models. In this study we sampled VOCs from the headspace of A549 cells at an air-liquid interface (ALI). We hypothesised VOC sampling from ALI cultures could be used to profile potential biomarkers of S. aureus lung infection. VOCs were collected using thin film microextraction (TFME) and were analysed by thermal desorption-gas chromatography-mass spectrometry. After optimising ALI cultures, we observed seven VOCs changed between A549 and media control samples. After infecting cells with S. aureus, supervised principal component-discriminant function analysis revealed 22 VOCs were found to be significantly changed in infected cells compared to uninfected cells (p < 0.05), five of which were also found in parallel axenic S. aureus cultures. We have demonstrated VOCs that could be used to identify S. aureus in ALI cultures, supporting further investigation of VOC analysis as a highly sensitive and specific test for S. aureus lung infection.

Received 22nd July 2022, Accepted 12th December 2022 DOI: 10.1039/d2an01205g

rsc.li/analyst

## Introduction

As a commensal microorganism Staphylococcus aureus does not normally cause harm to a healthy individual. Complications arise however with tissue damage<sup>1</sup> and a weakened immune system unable to fight off an opportunistic infection, for example in critically ill patients.<sup>2,3</sup> Rapid and early detection of S. aureus is critical to diagnose infection,

Measuring exhaled volatile organic compounds (VOCs) offers potential for rapid non-invasive detection of lung infection.<sup>5,6</sup> To this end, several studies have explored VOC analysis with S. aureus infection from respiratory samples.7-9 However, VOCs from complex samples may contain several endogenous (e.g., host tissues and metabolism) and exogenous (e.g., sampling apparatus, environment, and other pathogens) sources, and these may likely confound VOCs directly associated with microbes. Investigating how these sources impact changes in the volatilome may provide novel insights into their metabolic pathway activation and role in infection pathogenesis. For example, studies have explored how changing the culture media environment can cause variations in the volatilome of bacterial cultures. 10,11 Confounders such as host response metabolic response to an infection may be responsible for poor reproducibility between in vitro and in vivo studies.12,13

Infection models of microbes and mammalian cells are more representative of the human lung environment than standard microbial cultures. 14,15 Air-liquid interface (ALI) culture models are widely used to emulate the cellular structure of lung disease in controlled environments and in

10.1039/d2an01205g

prescribe the right antibiotic at the right time, and reduce the risks associated with drug-resistant strains.4

<sup>&</sup>lt;sup>a</sup>Division of Immunology, Immunity to infection & Respiratory Medicine, Faculty of Biology, Medicine, and Health, University of Manchester, Manchester, UK <sup>b</sup>Institut Necker-Enfants Malades, Paris, France

<sup>&</sup>lt;sup>c</sup>Université Paris-Saclay, UVSQ, INSERM, Infection et inflammation, Montigny le Bretonneux, France

 $<sup>^</sup>d$ Herman B Wells Center for Pediatric Research, Pediatric Pulmonology, Allergy, and Sleep Medicine, Indiana University School of Medicine, Indianapolis, USA <sup>e</sup>Service de Pneumo-Pédiatrie. Université René Descartes. Hôpital Necker-Enfants Malades, Paris, France

<sup>&</sup>lt;sup>f</sup>Hôpital Foch, Exhalomics, Département des maladies des voies respiratoires,

<sup>&</sup>lt;sup>g</sup>NIHR Manchester Biomedical Research Centre, Manchester University Hospitals NHS Foundation Trust, Education and Research Centre, Wythenshawe Hospital, Manchester, M23 9LT, UK. E-mail: stephen.fowler@manchester.ac.uk † Electronic supplementary information (ESI) available. See DOI: https://doi.org/

**Analyst** Paper

some cases reduce the dependence on animal models for experimentation.<sup>16</sup> Studies have also profiled the headspace from air-liquid interface (ALI) models, to investigate viral infection and the metabolic impact of oxidative stress from smoking.<sup>17–19</sup> These studies used either used stir-bar sorptive extraction or solid phase microextraction to sample VOCs from modified cell culture apparatus. All three studies used gas chromatography-mass spectrometry (GC-MS) for analysis of VOCs.

The aim of this study was to model epithelial damage caused by S. aureus and analyse the volatilome for potentially diagnostic biomarkers for S. aureus. The hypothesis was that VOC sampling could be used to profile potential biomarkers of S. aureus lung infection. We developed a method to sample VOCs using thin-film microextraction (TFME) from the apical compartment of an alveolar epithelial ALI culture, analysed offline by thermal desorption-gas chromatography-mass spectrometry (TD-GC-MS). Univariate and multivariate statistical comparisons will be presented between mammalian cells with and without S. aureus inoculation, using cell culture media and S. aureus culture as control samples.

#### 2. Methods

#### 2.1. Strains and cell culture

The immortalised A549 alveolar epithelial cell line was used in this study, kindly donated from the Blaikley lab at the University of Manchester (passage no. 117). Cells were cultured in Dulbecco's Modified Eagle's medium (DMEM) supplemented with 4500 mg L<sup>-1</sup> glucose (Sigma Aldrich, UK), 10% fetal bovine serum (PAA, UK), 2% 200 mM L-glutamine (Sigma Aldrich, UK), 1% 100 mM sodium pyruvate solution (Sigma Aldrich, UK), 1% penicillin-streptomycin (1000 µg mL<sup>-1</sup>-10 mg mL<sup>-1</sup> in solution with 0.9% NaCl, Sigma Aldrich, UK), and 0.1% 250  $\mu g \text{ mL}^{-1}$  Amphotericin B (Gibco, UK).

Cells were incubated at 37 °C in 5% CO<sub>2</sub>, passaged when at least 80% cell confluence was reached using trypsin-EDTA (0.25% solution, Sigma Aldrich, UK), centrifuged at 100 RCF for 5 min, and the pellet resuspended in fresh media to a concentration of  $1 \times 10^6$  cells per mL. Cell concentration was determined using a trypan blue stain (0.4%, Sigma Aldrich, UK) and cells counted using a Fuchs-Rosenthal haemocytometer (C-chip DHC-F01, NanoEntek, South Korea).

To generate an ALI culture, approximately  $1 \times 10^5$  cells were seeded onto sterile 12-well plates with well inserts (ThinCert™, PET transparent membrane, pore size 3.0µm, pore density 0.6 × 10<sup>6</sup> cm<sup>2</sup>, insert surface area 113.1mm<sup>2</sup>, Gibco, UK). Spent media in the apical compartment was removed after 24 h to generate a cell layer exposed to air on the apical side of the insert membrane. Basal compartment media was replenished every other day. Cell growth was visually inspected compared with control wells. In addition, transepithelial electrical resistance (TEER) measurements were carried out using an epithelial voltohmmeter according to manufacturer protocols (EVOM<sup>2</sup>, World Precision Instruments, USA). Cell monolayer

TEER was calculated using the following equation: TEER ( $\Omega$ cm<sup>2</sup>) =  $R_{\text{Tissue}} \times S$ , where R is resistance ( $\Omega$ ),  $R_{\text{Tissue}} = R_{\text{blank}}$  –  $R_{\text{culture}}$ , and S is the insert surface area (cm<sup>2</sup>). Both apical and basal compartments were washed with phosphate buffered saline (PBS) before measurement.

#### 2.2. Infection with S. aureus and VOC sample collection

A reference strain of S. aureus (ATCC 29213) was initially subcultured in tryptic soy media. Apical compartments were inoculated with 100 µL of S. aureus suspended in PBS and incubated for 24 h at 37 °C. Prior to this, basal media was replaced with DMEM without antibiotic supplementation and incubated at 37 °C for a further 24 h to confirm absence of contamination.

TFME was carried out using PDMS strips (20 mm × 15 mm × 0.45 mm, Goodfellow Cambridge Ltd, Huntingdon, UK), as previously described.20 Strips were initially washed with water then methanol and conditioned in a stream of N2 at 350 °C for 1 h. Conditioned strips were stored in capped stainless-steel tubes until sample collection and analysis and were used within 24 h of conditioning. Strips were reused after reconditioning with the same parameters. VOCs were sampled by inserting a PDMS strip into the apical compartment of each well insert. Cultures were then incubated for a further 24 h before removing PDMS strips for VOC analysis. S. aureus was aspirated from the apical side with a 100 µL PBS wash and growth observed on tryptic soy agar plates using the drop plate method (10 µL of the PBS wash dropped on a segment of the agar plate and incubated at 37 °C for 18 h). Basal media was also collected to observe S. aureus migration through the cell layer and insert membrane. The PBS wash and basal media were diluted (1:10) up to a maximum of four dilutions.

Absorbance change of Phenol Red, a pH indicator and a constituent of DMEM, was measured as an orthogonal method to detect cell culture acidification by S. aureus infection. A standard curve of DMEM at different pH was created by adding citric acid, measured at 415 and 560 nm (Biochrom Asys UVM340, Cambridge, UK). The standard curve was calculated based on the 415/560 nm ratio for Phenol Red and used to interpolate pH levels of basal media ( $R^2$  0.96). This method was used as opposed to direct measurement using a pH probe due to the low volume of spent media available per well and risk of cross contamination between infected and uninfected samples.

# 2.3. Thermal desorption-gas chromatography-mass spectrometry (TD-GC-MS)

PDMS strips were dry purged with N<sub>2</sub> at 50 mL min<sup>-1</sup> for 4 min to remove excess water on the sorbent before analysis by TD-GC-MS. Prior to thermal desorption, a gas standard (1 ppmV p-bromofluorobenzene in N2; Thames Restek, Saunderton, UK) was automatically spiked onto each tube. Tubes were then desorbed at 280 °C for 5 min (TD100, Markes International, Llantrisant, UK) then transferred with splitless injection to a cold trap (kept at 0 °C) for cryofocussing which was subsequently flash heated to 280 °C for 2 min. Desorbed

**Paper** Analyst

VOCs were transferred to the GC column (DB-5 ms, length 30 m  $\times$  internal diameter 0.25 mm  $\times$  film thickness 0.25  $\mu$ m, Agilent Technologies, Cheadle, UK) using a constant helium flow of 1 mL min<sup>-1</sup>. The GC column oven (7890B, Agilent Technologies, Cheadle, UK) was set to a linear temperature programme starting at 30 °C and increasing to 250 °C at 7.5 °C min<sup>-1</sup> (total GC cycle time of 29.33 min). After GC separation, VOCs were transferred to the MS (7010, Agilent Technologies, Cheadle, UK) where they were ionised (EI 70 eV) in the source kept at 150 °C. Data was acquired in full scan mode with an acquisition rate of 5 scans s<sup>-1</sup> between m/z 40-300. In addition, validation of cell culture VOCs was performed on a different GC-MS method, as described previously. 21

#### 2.4. Data analysis

Masshunter Quantitative Analysis software (version B.07.00, Agilent, Cheadle, UK) was used to automatically integrate peaks (Agile2) and identify VOCs in each sample to MSI level 2<sup>22</sup> using the NIST mass spectral library (version 2014, match factor > 70) and retention indices calculated using an alkane ladder. An aligned  $n \times p$  matrix was created of peak intensities. Missing values were imputed using Random Forest imputation (R package missForest). All samples were then normalised by the internal standard (p-bromofluorobenzene), and features were log transformed and mean centred (i.e. z-normalised). Statistical analyses were performed in Graphpad Prism (version 9.1) and R (v 4.1.1) and consisted of univariate tests including repeated measures ANOVA, non-parametric Mann-Whitney-U and multivariate analysis in the form of unsupervised principal component analysis (PCA) and supervised classification via principal component-discriminant function analysis (PC-DFA). PC-DFA was cross validated (80:20 training: test set) using stratified random sampling with 1000 repeats. Comparisons were made between culture media and

mammalian cells, and infected and uninfected cells. The Benjamini-Hochberg one way ANOVA was applied (q < 0.05)where appropriate to control the false discovery rate (FDR).

#### 3. Results

#### Blank PDMS strip reproducibility

Reproducibility of PDMS strip adsorption was assessed by calculating RSD for the internal standard purged onto blank PDMS strips. Repeated desorption of a single blank PDMS strip (n =10) resulted in an RSD of 1.35% whereas ten repeat desorption runs of a tube without sorbent material resulted in an RSD of 1.10%. Desorption of several individual PDMS strips (n = 10)resulted in an RSD 7.27%. The most consistently produced compounds associated with the sorbent background were hexamethylcyclotrisiloxane and octamethylcyclotetrasiloxane.

#### 3.2. Optimal ALI culture time and CFU inoculum for S. aureus

To assess the formation of a A549 cell layer on the well insert, transepithelial integrity was measured by TEER (n = 11) on day 2, 5, 7, 9, and 12 after seeding (Fig. 1). A repeated measures ANOVA showed TEER increased on all days compared to the initial measurement on day 2 (p = < 0.001), whereas no difference was found between days 5, 7, 9, and 12. Day 9 was selected as the optimal time to infect cells based on the representative TEER (55.2  $\pm$  6.5) and low RSD (11.7%), in comparison to other time points (day  $5 = 52.3 \pm 14.2$  RSD 27.2%, day  $7 = 56.1 \pm 16.8 \text{ RSD } 30.0\%$ , day  $12 = 39.7 \pm 3.6 \text{ RSD } 9.1\%$ ). TEER measurements were also used to evaluate the impact of S. aureus on membrane integrity.

To select an infection CFU inoculum concentration for infection, A549 cells were cultured at ALI and the apical side

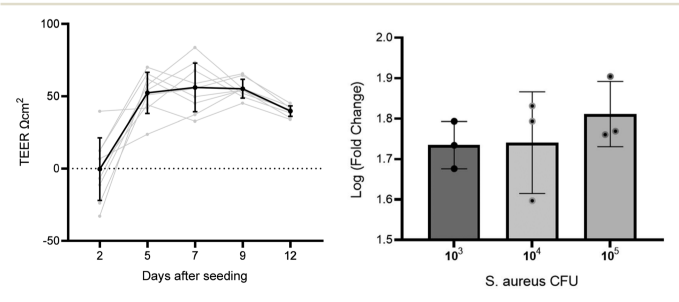


Fig. 1 Left: line graph with TEER measurements across several days on the same well inserts. Right: graph comparing three S. aureus CFU concentration of TEER fold change before and after inoculation.

was inoculated with *S. aureus* at three CFU concentrations –  $1 \times 10^3$ ,  $1 \times 10^4$ , and  $1 \times 10^5$  CFU (n=3 per concentration). All *S. aureus* CFU concentrations decreased cell culture TEER after inoculation and incubation for 24 h (paired t test p < 0.001). Mean log fold changes for each CFU concentration were  $1 \times 10^3$  CFU mL<sup>-1</sup> = 1.7,  $1 \times 10^4$  CFU mL<sup>-1</sup> = 1.8, and  $1 \times 10^5$  CFU mL<sup>-1</sup> = 1.8. No significant differences were found between TEER measured before or after inoculation between CFU concentrations (p > 0.05). The lowest concentration tested ( $10^3$  CFU) was taken forward for infecting cells in the head-space sampling experiments. A low initial CFU was selected to reduce detection of saturated high-abundance VOCs from *S. aureus*, reducing the risk of sample carry over.

#### 3.3. Volatiles from A549 cells without infection

Headspace volatiles were sampled from A549 cells and mediaonly controls after 9 days in ALI culture, under standard growth conditions without infection. In addition, a validation experiment was carried out using the same culture methods but with a different analytical method as described in<sup>21</sup> (cells n = 6, media n = 3). All samples were confirmed free of microbial contamination after plate culture and visual inspection of the culture colour remaining red (media) or pink (cells) and absence of turbidity. After controlling for FDR, 107 features were significantly different between media and cells. For the validation experiment, 111 features were found significantly different between the two groups. Separation across PC1 was observed between sample groups for both original (Fig. 2A) and validation experiments (Fig. 2B).

Seven VOCs extracted from PCA loadings were significantly different between media and cells across both experiments as shown in Fig. 2. These included benzene 2-nitroethyl-(log 2 fold change [FC] 2.7, q < 0.001, validation experiment FC 5.2, q < 0.001), 1-hexanol 2-ethyl-(FC 2.0, q = 0.003 validation FC 2.3, q = 0.023) and a methylated furanone (Fig. 2C). The latter compound was identified as 2(3H)-furanone 5-methyl-(FC 2.5, q = 0.026) in the original experiment and as 2(3H)-furanone dihydro-4-methyl-(FC 2.9, q = 0.004) in the validation experiment. VOCs with a decreased abundance in cells (Fig. 2D) were benzeneacetaldehyde (FC -2.5, q = 0.003, validation FC -3.5, q = 0.001), benzaldehyde (FC -2.0, q = 0.003, validation FC -1.6 q = 0.007), 4-quinolinecarboxaldehyde (FC -8.1, q = 0.010, validation FC -4.1 q = 0.007) and benzaldehyde 4-methyl-(FC -1.2, q = 0.032, validation FC -2.7 q < 0.001).

#### 3.4. Infection of A549 cells with S. aureus

After 9 days of maintaining an ALI, cells were infected with  $10^3$  CFU of *S. aureus* (n=6) on the apical side alongside cells without infection inoculated with  $100\mu$ L PBS without *S. aureus* (n=6) and control samples (media and *S. aureus* in media). After 24 h, leakage of media from the basal to the apical side was observed for A549 cultures with *S. aureus* infection, whereas no such leakage was seen for samples uninfected with *S. aureus*. TEER measurements decreased in *S. aureus* infected cells after 24 h (paired t test p=0.0003), as shown in Fig. 3A. Basal media pH suggested increased acidity in infected cell

cultures compared to uninfected cells (Fig. 3B) which is consistent with bacterial contamination of mammalian cell culture. No visible *S. aureus* growth was observed for A549 cultures alone compared to infected A549 for both apical and basal samples (Fig. 3C). Saturated microbial growth was observed across all inoculated cultures, including the basal compartment cultures (>10<sup>7</sup>CFU) suggesting exponential growth of *S. aureus* in the presence of A549 cells.

#### 3.5. Volatile profile of A549 cells infected with S. aureus

Data were initially assessed using PCA (Fig. S1†) and inspecting total ion current chromatograms, where no clear separation was found between groups. In addition, two samples failed analysis due to water retention. Therefore, to extract VOC features which differentiate infected and uninfected cells, a cross-validated PC-DFA (with eight PCs) was performed with A549 cells infected with S aureus (n = 4) in comparison to uninfected A549 cells (n = 6), media (n = 8), and S. aureus (n = 8)samples alone (Fig. 4). The PC-DFA model showed separation across the first discriminant function (DF1, S3) between samples with A549 cells (infected and uninfected) and those without (media and S. aureus alone). The second discriminant function (DF2, S4) showed separation of infected and uninfected sample groups. After applying an arbitrary noise cut off to the loadings (0.05), features that contributed to separation were extracted for DF1 (n = 70) and DF2 (n = 50).

## Discussion

In this study, we found several VOCs associated with uninfected and infected alveolar epithelial cells in ALI cultures. VOCs were associated with the infection response alone and included VOCs also identified in axenic cultures of *S. aureus*. Therefore, these results demonstrate the diverse sources of VOCs in the infection process (*i.e.* bacterial infection and cell response) and are potentially good candidates as diagnostic biomarkers of the multiple contributors to infection pathogenesis.

An assessment of PDMS material adsorption showed good reproducibility without degradation after 10 sequential desorption cycles of the same PDMS strip. Desorption between different PDMS strips was less reproducible (higher RSD). This may be due to minor inconsistencies between strip length and width. To minimise the number of water-logged samples, the dry purge time would require optimisation in future studies. When sampling from cell headspace, only seven compounds were observed to be consistent between two experiments con-

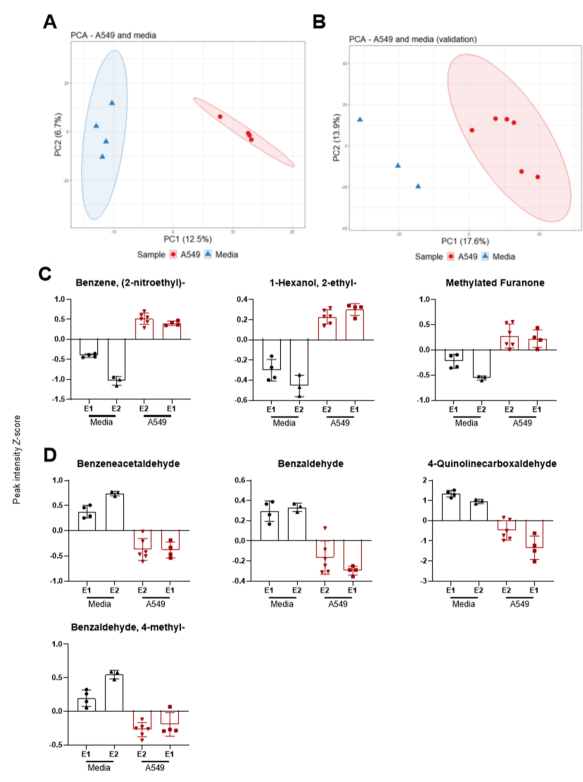


Fig. 2 PCA scores plot showing PC1 and PC2 of headspace sampling experiments where VOCs were sampled after 24 h of incubation at 37 °C comparing (A) A549 and media, and (B) validation experiment comparing A549 and media using a different analysis method (see methods). Z-score plots of (C) of significantly increased (p < 0.05) VOCs in A549 and (D) significantly increased VOCs in media, across both original (E1) and validation (E2) experiments.

ducted at different times and with different GC-MS methods. This shows that different analytical methods can impact on the total number of VOCs observed. Furthermore, improving

sampling and analytical parameters, such as introducing a secondary polar GC column, will increase the abundance and range of detectable VOCs. For sampling headspace, research-

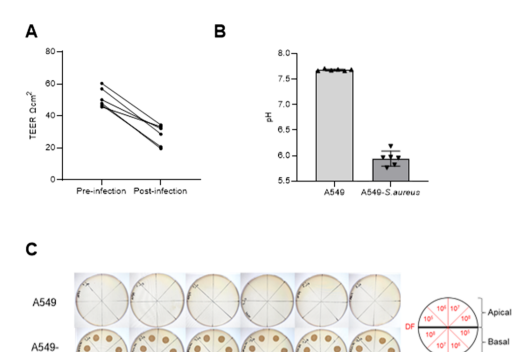
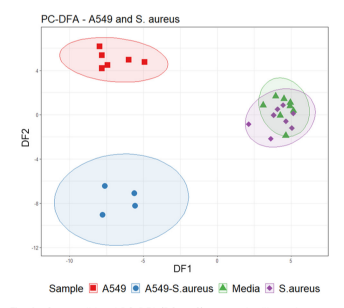


Fig. 3 (A) TEER measurements per well before and after infection of A549 cells with *S. aureus*. (B) Basal media pH after 24 h of infection (A549-S. *aureus*) compared to uninfected (A549) cells. (C) Tryptic soy agar plates with serial dilutions (DF = estimated CFU mL $^{-1}$ ) of apical PBS wash and basal media from both sample groups.



S. aureus

Fig. 4 Cross-validated PC-DFA (PCs = 8) scores plot illustrating separation between sample groups on DF1 and DF2.

ers have developed multi-bed TFME films which may also increase the range of VOCs sampled from cultures.  $^{23}$ 

Cells were successfully infected with *S. aureus* as demonstrated by increased CFU post-infection, reduced pH in co-culture (*i.e.*, acidification due to aerobic bacterial contamination), and reduced TEER suggesting membrane damage.

Epithelial damage by S. aureus and subsequent membrane rupture resulting in leakage in ALI culture is consistent with previous studies using a similar airway infection model.<sup>24</sup> Alcohol and fatty acid compounds were increased in the VOC profile of infected cells. Phenylethyl alcohol has previously been identified in S. aureus headspace.<sup>25</sup> The VOC 1-hexanol 2-ethyl in ALI culture was also shown by previous studies investigating A549 cells.<sup>26</sup> Fatty acids have been associated with the cell membrane phospholipid structure and their increased detection in infected cells may potentially be caused by membrane damage by S. aureus. Fatty acids are also found to control cell membrane integrity and biofilm dispersion in S. aureus. 27-30 Aromatic volatiles were decreased under normal cell growth suggesting uptake from the media and have been reported in previous studies investigating A549 cells. 26,31 Several methylated pyridine compounds were increased in infected cells compared to uninfected cells. Pyridine 3-methyl-2-phenyl was also detected in axenic S. aureus cultures. Pyridine is a potential marker for smoking and structurally associated with nicotine.32

Although it was expected that VOCs which are very volatile will not be captured by the PDMS sorbent, not all VOCs within the detectable range were consistent with previous literature. For example, dimethyl disulfide (DMDS), a VOC known to be emitted by *S. aureus*, was comparatively low in infected cells, which suggests altered metabolism during infection. Furthermore, DMDS is a ubiquitous VOC produced by several microbial and plant species and therefore has little value as a

 Table 1
 List of VOCs which differentiate infected and uninfected A549

Main		Quantifier	NIST match	Retention	NIST library	Calculated	Detected in infection	Detected in S. aureus	Normalis intensity	Normalised peak intensity		,	d	95% CI	95% CI
Mainten	VOC name (NIST ID)	z/w	factor	time (min)	.KI	kovats RI	model	cultures	Mean		Median		value	lower	upper
cis acid         22         3.81         NA         721         1         0.056         1.015         0.095         1.015         0.095         1.015         0.095         1.015         0.095         1.015         0.095         1.015         0.095         1.015         0.015         2.32         0.010         2.32         0.010         2.32         0.010         2.32         0.010         2.32         0.010         0.32         0.010         0.317         0.319         0.010         2.32         0.010         0.32         0.010         0.32         0.010         0.32         0.010         0.32         0.010         0.32         0.02         0.02         0.03         0.010         0.32 <t< td=""><td>Triethylamine</td><td>101</td><td>87.0</td><td>3.44</td><td>711</td><td>693</td><td></td><td></td><td>-0.101</td><td>0.978</td><td>-0.545</td><td></td><td>0.171</td><td>-0.099</td><td>2.235</td></t<>	Triethylamine	101	87.0	3.44	711	693			-0.101	0.978	-0.545		0.171	-0.099	2.235
tine cacid at the	2-Propenoic acid	72	92.7	3.81	NA	721	<b>←</b>		0.584		869.0		0.010	-2.550	-0.475
liting 2-billone-N/N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-	Propanoic acid	74	6.68	3.82	NA	722	- ←		0.566		0.00		0.010	-2.379	-1.397
cicald, amethyly         94         90.6         419         722         740         1         -0.190         1319         1316         0.00         2.01         0.11         131         140         0.00         2.03         1.03         0.01	2,2′-Bioxirane	57	81.9	3.94	NA	729			-0.330		0.370		0.610	-3.350	0.913
sing 2-chilorowA, M.         107         87.0         4.29         75.0         75.3         †         0.349         0.68         0.629         0.68         0.629         1.66         0.034         0.06         0.04<	Disulfide, dimethyl <sup>a</sup>	94	9.06	4.10	722	740	$\rightarrow$		-0.190		0.436		0.010	2.031	2.991
oct acid, 2-methyl         86         71.1         4.63         71.1         776         1         0.012         0.377         0.013         0.89         0.347           cell pell acid, 2-methyl         85         3.4         5.54         816         88         1         0.022         0.858         0.220         0.858         0.220         0.858         0.220         0.858         0.11         0.230         0.00         0.220         0.868         0.11         0.230         0.00         0.220         0.668         0.11         0.12         0.13	Ethanamine, 2-chloro-N,N-	107	87.0	4.29	750	753	- ←		0.349		0.629		0.038	-2.602	-0.085
circle and between byte and control and con	dimethyl														
Second	2-Propenoic acid, 2-methyl	98	71.1	4.63	711	922	$\rightarrow$		-0.012	0.717	-0.015		0.010	0.347	1.935
sed of selection of s	Acetyl valeryl	85	93.4	5.56	816	830	<b>←</b>		0.622		0.658	1.352	0.010	-2.296	-0.696
septimely         74         82.9         6.42         86.8         6.42         86.8         1.25         0.611         0.711         -2.83           septimely         7         9.4         6.42         86.7         87.4         1.25         0.11         0.71         0.11         0.71         0.13         0.13         0.13         0.13         0.13         0.13         0.13         0.13         0.13         0.13         0.13         0.03	Formic acid hydrazide	09	72.4	5.94	NA	849	- ←		0.687		0.530		0.010	-1.918	-0.538
14.3mmthyl, acetate 96 843 845 874 874 1 0.000 0.050 0.068 1.884 1.887 0.349 1.341 1.348 0.349 0	Butanoic acid, 2-methyl	74	82.9	6.31	894	898			0.638		0.111		0.171	-3.833	0.281
1.   1.   1.   1.   1.   1.   1.   1.	1-Butanol, 3-methyl-, acetate	70	92.4	6.42	867	874	←		0.700		0.668		0.010	-2.187	-0.875
object         3.5-dimethyl         87         44.1         8.15         NAA         958         †         —0.003         1.15         0.389         1.03         0.239         1.03         0.239         1.03         0.207         0.007 </td <td>Cyclopent-4-ene-1,3-dione</td> <td>96</td> <td>84.3</td> <td>6.47</td> <td>880</td> <td>876</td> <td></td> <td></td> <td>0.584</td> <td></td> <td>0.131</td> <td></td> <td>0.352</td> <td>-0.928</td> <td>1.815</td>	Cyclopent-4-ene-1,3-dione	96	84.3	6.47	880	876			0.584		0.131		0.352	-0.928	1.815
hylacohol*         128         84.5         10.11         1097         1052         0.391         0.391         1.33         0.389         1.205         0.391         1.203         0.389         1.205         0.134         1.134         1113         1         4         0.239         0.802         0.201         0.754         0.124         0.001         2.114         0.113         1<	4-Heptanol, 3,5-dimethyl	87	74.1	8.15	NA	928	←		-0.093	1.415	-0.303		0.010	-3.423	-1.378
thy alcohol/- the chone 2         91         98.2         11.36         1186         1112         †         0.097         0.990         0.591         1.754         100         0.214         1.756         0.064         0.051         1.764         1.00         0.024         0.020         0.021         0.020         0.054         1.175         0.064         0.064         0.054         0.071         0.071         0.034         1.175         0.064         0.053         0.072         0.064         0.064         0.064         0.064         0.064         0.064         0.064         0.064         0.064         0.064         0.073         0.076         0.064         0.064         0.073         0.076         0.064         0.073         0.076         0.064         0.073         0.076         0.066         0.073         0.076         0.066         0.073         0.076         0.066         0.073         0.076         0.066         0.073         0.076         0.068         0.113         0.073         0.078         0.073         0.076         0.069         0.073         0.078         0.073         0.078         0.073         0.078         0.073         0.078         0.073         0.078         0.073         0.078         0.073	Furaneol	128	84.5	10.11	1097	1052			0.391		0.389		0.171	-0.201	2.563
14   14   15   1134   1143   1144   1143   1144   1143   1144	Phenylethyl alcohol $^a$	91	98.2	11.36	1086	1112	←	←	0.097		-0.541		0.010	-2.114	-1.675
Propose monthy    12, 22, 1169   1155   1158   1158   1158   1158   1158   1159   1158   11	4H-Pyran-4-one, 2,3-dihydro-	144	79.8	11.97	1134	1143			0.239		0.201		0.762	-0.664	1.982
Part	3,5-dihydroxy-6-methyl														
1.   1.   1.   1.   1.   1.   1.   1.	Dehydromevalonic lactone	54	9.06	12.22	1169	1155			0.154	1.144	-0.104		0.114	-3.150	0.365
rowsethyl)-2-         100         73.1         12.30         NA         1159         0.190         1.123         -0.29         1.008         0.171         -0.725           none         82         89.1         13.22         12.00         1205         -0.086         0.752         0.078         0.753         1.009         0.071         -0.083           proparentitile         91         95.3         13.82         1249         1270         1         4         6         0.752         0.078         0.078         0.070         -0.063           ged, 2-bydroxy-, ethyl         120         71.2         14.45         1249         1270         1         7         0.482         1.08         0.752         0.078         0.079         0.070         2.033           ged, 2-bydroxy-, ethyl         120         14.45         1249         1270         1         4         4         6         6         6         7         1         0.010         -2.631           ged, 4-chloro         139         91.1         137.0         1446         1453         1         4         6         6         6         9         1         1         1         1         1         1         <	2-Isobutyl-4-methylpyridine	107	90.3	12.27	1154	1158	←		0.469		0.440		0.038	-1.346	-0.038
Public   P	3-(2-Hydroxyethyl)-2-	100	73.1	12.30	NA	1159			0.190	1.123	-0.229		0.171	-0.725	2.948
82         89.1         13.22         1200         1205         120         1205         0.05         0.078         0.75         0.078         0.75         0.078         0.75         0.078         0.079         0.070         -0.063           acid, 2-hydroxy, ethyl         120         71.2         1244         1249         1270         †         †         0.363         1.198         -0.185         1.281         0.010         -2.861           acid, 2-hydroxy, ethyl         120         95.6         14.48         1292         1293         †         †         0.492         0.73         -0.492         0.73         0.010         -2.861           acid, 4-chloro         139         91.3         15.69         NA         1337         0.418         1.265         -0.492         0.73         0.010         -1.739           neceramide         92         85.5         16.12         1387         1361         †         4.46         1.25         0.743         0.743         0.743         0.010         -1.739           neceramide         92         84.4         17.71         1446         1453         †         4.41         0.743         0.743         0.743         0.741         0.010 <td>oxazolidinone</td> <td></td>	oxazolidinone														
proparenitrile 91 95.3 13.82 1243 1237 † † 0.343 1.068 0.725 1.851 0.010 -2.631 acid, 2-hydroxy, ethyl 120 71.2 14.45 1249 1270 † † † 0.343 1.068 0.725 1.851 0.010 -2.631 acid, 2-hydroxy, ethyl 120 71.2 14.45 1249 1270 1293 † † † 0.343 1.08 0.753 1.98 0.753 0.010 0.2.93 acid, 4-chloro 139 0.01 1.292 1.293 † † † 0.492 0.735 0.753 0.898 0.510 0.735 0.753 0.898 0.510 0.293 0.895	Decanal	82	89.1	13.22	1200	1205			-0.086		0.078	0.598	0.067	-0.063	2.038
acid, 2-hydroxy, ethyl 120 71.2 14.45 1292 1293 ↑ ↑ ↑ 0.363 1.198 -0.187 2.122 <b>0.010</b> -2.861 1.39 acid, 4-chloro 139 91.3 15.69 NA 1337	Benzenepropanenitrile	91	95.3	13.82	1243	1237	←		0.343		0.725		0.010	-2.631	-0.634
scid, 4-chloro         39         95.6         14.88         1292         1293         †         —0.492         0.738         —0.533         1.295         —0.503         1.295         —0.492         0.783         —0.533         1.295         —0.492         0.783         —0.593         1.598         0.010         —1.739           sic acid         73         85.5         16.12         1387         1361         1389         †         —0.493         0.785         0.763         0.783         0.171         —0.945           sectamide         92         87.6         16.80         1412         1399         †         —0.093         0.879         0.747         0.010         —2.938           sectamide         92         87.4         17.71         1446         1453         †         —0.093         0.879         0.747         0.010         —2.938           nac, 1-cthenyl-1-         85         91.1         1492         1562         †         0.433         1.054         0.410         1.574         0.010         —2.747           hydro         129         94.5         19.49         1462         1562         †         0.433         1.054         0.010         1.574         0.010 <td>Benzoic acid, 2-hydroxy-, ethyl</td> <td>120</td> <td>71.2</td> <td>14.45</td> <td>1249</td> <td>1270</td> <td>←</td> <td>←</td> <td>0.363</td> <td></td> <td>-0.187</td> <td></td> <td>0.010</td> <td>-2.861</td> <td>-1.857</td>	Benzoic acid, 2-hydroxy-, ethyl	120	71.2	14.45	1249	1270	←	←	0.363		-0.187		0.010	-2.861	-1.857
scid, 4-chloro         139         95.6         14.88         1292         1293         †         —0.492         0.738         —0.503         1.298         0.010         —1.731           sic acid, 4-chloro         139         91.3         15.69         NA         1337         1         —0.492         0.738         —0.503         1.258         —0.078         1.651         0.171         —0.345           sic acid, 4-chloro         139         11.2         1387         1361         7         6.263         0.748         1.651         0.171         —0.045           sectamide         92         87.5         16.80         1412         1399         †         6.39         0.747         0.010         —2.738         0.601         —0.045           resence         93         84.4         17.71         1446         1453         †         †         0.030         0.154         0.101         —2.038         0.610         —0.045           resence         85         91.1         1452         1465         †         †         0.030         0.154         0.010         —2.734           resence         85         91.2         1465         †         †         0.030	ester														
139         91.3         15.69         NA         1337         0.418         1.265         -0.078         1.651         0.171         -0.312           73         85.5         16.12         1387         1361         6.529         0.805         0.763         0.783         0.810         -0.945           92         87.6         16.80         1412         1399         †         4.037         1.102         0.689         0.763         0.879         -0.473         1.629         0.010         -0.945           93         84.4         17.79         1463         1465         †         4.093         0.879         -0.473         1.629         0.010         -2.017           85         91.1         17.92         1463         1465         †         †         0.433         1.096         0.090         1.564         0.010         -2.047           129         94.5         19.49         1492         1562         †         †         0.433         1.096         0.090         1.572         0.010         -2.047           168         86.8         20.09         1513         †         †         0.433         1.096         0.050         1.112         0.010	$Indole^a$	06	92.6	14.88	1292	1293	←		-0.492		-0.503		0.010	-1.739	-0.395
73         85.5         16.12         1387         1361         0.529         0.529         0.805         0.763         0.888         0.610         -0.945           92         87.6         16.80         1412         1399         †         0.376         1.102         0.639         0.747         0.010         -2.938           93         84.4         17.71         1446         1453         †         -0.093         0.879         -0.473         1.629         0.010         -2.938           85         91.1         17.92         1463         1465         †         0.800         1.154         0.410         1.964         0.010         -2.747           93         76.0         19.49         1492         1562         †         0.433         1.096         0.090         1.572         0.010         -2.674           129         94.5         19.53         1562         †         †         0.433         1.096         0.090         1.572         0.010         -2.794           93         72.9         24.79         NA         1915         †         †         0.745         0.919         0.456         1.124         0.019         -2.784	Benzoic acid, 4-chloro	139	91.3	15.69	NA	1337			0.418		-0.078		0.171	-0.312	3.358
92         87.6         16.80         1412         1399         †         0.376         1.102         0.639         0.747         0.010         -2.938           93         84.4         17.71         1446         1453         †         -0.093         0.879         -0.473         1.629         0.010         -2.017           85         91.1         17.92         1463         1465         †         0.800         1.154         0.410         1.964         0.010         -2.047           93         76.0         19.49         1492         1562         †         0.433         1.096         0.090         1.572         0.010         -2.074           168         86.8         20.09         1513         1598         †         0.202         1.163         -0.151         1.713         0.010         -2.794           93         72.9         24.79         NA         1915         †         0.745         0.919         0.450         1.124         0.019         -2.794           93         72.9         24.78         1908         1921         †         0.745         0.919         0.450         1.124         0.019         -2.794           94 <td< td=""><td><i>n</i>-Decanoic acid</td><td>73</td><td>85.5</td><td>16.12</td><td>1387</td><td>1361</td><td></td><td></td><td>0.529</td><td></td><td>0.763</td><td></td><td>0.610</td><td>-0.945</td><td>2.131</td></td<>	<i>n</i> -Decanoic acid	73	85.5	16.12	1387	1361			0.529		0.763		0.610	-0.945	2.131
93       84.4       17.71       1446       1453       †       †       -0.093       0.879       -0.473       1.629       0.010       -2.017         85       91.1       17.92       1463       1465       †       0.800       1.154       0.410       1.964       0.010       -2.047         93       76.0       19.53       1562       †       †       0.433       1.096       0.090       1.572       0.010       -2.674         129       94.5       19.53       1562       †       †       0.202       1.163       -0.208       1.176       0.010       -2.674         168       86.8       20.09       1513       1598       †       †       0.191       1.129       -0.151       1.713       0.010       -2.794         93       72.9       24.79       NA       1915       †       †       0.745       0.919       0.450       1.124       0.019       -2.280         154       73.0       24.88       1908       1921       †       0.051       1.413       0.107       0.738       0.038       -3.777         284       92.5       27.69       2170       2111       0.051       1.054	Benzeneacetamide	95	87.6	16.80	1412	1399	←		0.376		0.639		0.010	-2.938	-0.280
85       91.1       17.92       1463       1465       †       0.800       1.154       0.410       1.964       0.010       -2.747         93       76.0       19.49       1492       1562       †       †       0.433       1.096       0.090       1.572       0.010       -2.574         129       94.5       19.53       1562       1564       †       †       0.202       1.163       -0.208       1.176       0.010       -2.574         168       86.8       20.09       1513       1598       †       †       0.191       1.129       -0.151       1.713       0.010       -2.794         93       72.9       24.79       NA       1915       †       †       0.745       0.919       0.450       1.124       0.019       -2.794         93       72.9       24.78       1908       1921       †       †       0.375       1.413       0.107       0.738       0.038       -3.777         284       92.5       27.69       2170       2111       0.051       1.054       0.322       0.268       0.476       -1.252	<i>cis</i> -β-Farnesene	93	84.4	17.71	1446	1453	←	<b>←</b>	-0.093		-0.473		0.010	-2.017	-1.190
93         76.0         19.49         1492         1562         †         †         0.433         1.096         0.090         1.572 <b>0.010</b> -2.674           129         94.5         19.53         1562         1564         †         0.202         1.163         -0.208         1.176 <b>0.010</b> -3.492           168         86.8         20.09         1513         1598         †         0.091         1.129         -0.151         1.713 <b>0.010</b> -2.794           93         72.9         24.79         NA         1915         †         0.745         0.919         0.450         1.124 <b>0.019</b> -2.794           93         72.9         24.79         NA         1921         †         0.745         0.919         0.450         1.124 <b>0.019</b> -2.794           154         73.0         24.88         1921         †         0.379         1.413         0.107         0.738 <b>0.038</b> -3.777           284         92.5         27.69         2170         2111         0.051         1.054         0.322         0.268         0.476         -1.252	2(3H)-Furanone, 5-hexyldihydro	85	91.1	17.92	1463	1465	←		0.800		0.410	1.964	0.010	-2.747	-1.414
129 94.5 19.53 1562 1564 ↑ 0.202 1.163 -0.208 1.176 <b>0.010</b> -3.492 168 86.8 20.09 1513 1598 ↑ ↑ 0.191 1.129 -0.151 1.713 <b>0.010</b> -2.794 93 72.9 24.79 NA 1915 ↑ 0.379 1.413 0.107 0.738 <b>0.038</b> -3.777 284 92.5 27.69 2170 2111 0.001 2111 0.001 1.054 0.322 0.268 0.476 -1.252	Cyclohexane 1-ethenyl-1-	93	0.92	19.49	1492	1562	<b>←</b>	<b>←</b>	0.433		0.090		0.010	-2.674	666.0-
henyl 168 86.8 20.09 1513 1598 † † 0.191 1.129 -0.151 1.713 <b>0.010</b> -3.492 henyl 168 86.8 20.09 1513 1598 † † † 0.191 1.129 -0.151 1.713 <b>0.010</b> -2.794 93 72.9 24.79 NA 1915 † 0.379 1.413 0.107 0.379 1.413 0.107 0.738 <b>0.038</b> -3.777 284 92.5 27.69 2170 2111 2111 0.015 1.051 1.054 0.322 0.268 0.476 1.125 2.280	methyl-2-(1-methylethenyl)-4- (1-methylethylidene)						-	_							
henyl 168 86.8 20.09 1513 1598 † † 0.191 1.129 -0.151 1.713 <b>0.010</b> -2.794 henyl 168 72.9 24.79 NA 1915 † 0.745 0.919 0.450 1.124 <b>0.019</b> -2.280 1.44 154 73.0 24.88 1908 1921 † 0.379 1.413 0.107 0.738 <b>0.038</b> -3.777 284 92.5 27.69 2170 2111 0.019 2111 0.051 1.054 0.322 0.268 0.476 1.252	Dodecanoic acid	129	94.5	19.53	1562	1564	<b>←</b>		0.202	1.163	-0.208		0.010	-3.492	-0.455
93 72.9 24.79 NA 1915 † 0.745 0.919 0.450 1.124 <b>0.019</b> -2.280 1.4- 154 73.0 24.88 1908 1921 † 0.379 1.413 0.107 0.738 <b>0.038</b> -3.777 284 92.5 27.69 2170 2111 0.051 1.054 0.322 0.268 0.476 -1.252	Pvridine, 3-methyl-2-phenyl	168	86.8	20.09	1513	1598	- ←	<b>←</b>	0.191		-0.151		0.010	-2.794	-1.217
1,4-     154     73.0     24.88     1908     1921     †     0.379     1.413     0.107     0.738     0.038     -3.777       284     92.5     27.69     2170     2111     0.051     1.054     0.322     0.268     0.476     -1.252	Pyridine, 2-hexyl	93	72.9	24.79	NA	1915	- <b>←</b>	-	0.745		0.450		0.019	-2.280	-0.126
284 92.5 27.69 2170 2111 0.051 1.054 0.322 0.268 0.476 -1.252	Dyrrolo[1 2-albyrozine-1 4-	7 T	73.0	24.88	1008	1021	- +		0 370		0.107		0.020	-3 777	-0.113
284 92.5 27.69 2170 2111 0.051 1.054 0.322 0.268 0.476 -1.252	dione, hexahydro-3-(2-	+CT	73.0	7.00	1300	1971	_		6/6.0		0.107		0.030	777.6	C11.0_
25.3 27.0 0.470 -1.232	metrypropys,	707	л <b>с</b> о	07 60	0170	0111			0.051		0 200		9470	.1 050	0 544
	סרומחרימווטוי מיות	F07	5.76	60.14	0/17	7117			1000		445.0		0 / # • 0	4.4.4	44.5

Values highlighted in Bold indicate statistically significant changes in abundance ( $\alpha < 0.05$ ). <sup>a</sup> Identified to metabolomics standards initiative level 1. <sup>b</sup> NIST library retention indices based on non-polar capillary column stationary phase (references in Table S1†).

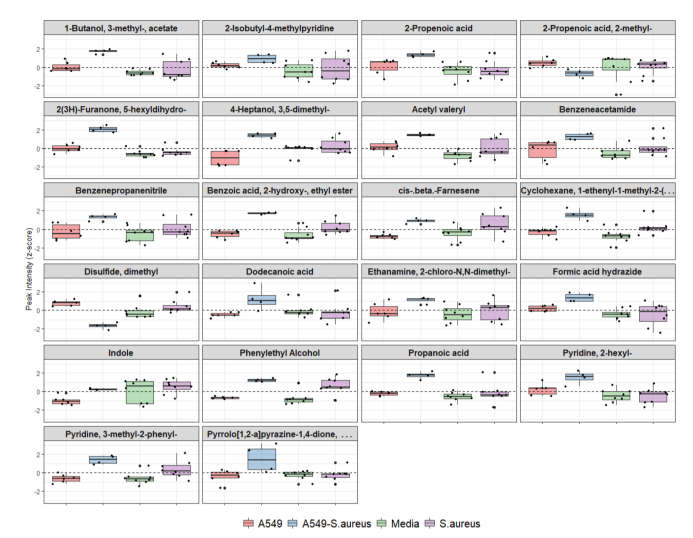


Fig. 5 Boxplots of VOCs which are significantly different between infected and uninfected cells, showing all sample groups for comparison. Dashed line represents peak intensity mean after z-normalisation.

clinically useful diagnostic biomarker. However, further optimisation of the method (*e.g.* use of labelled sulfur-containing amino acids to track DMDS) is required to evaluate metabolic changes during infection. Culture conditions could be further improved by using primary cell culture, other cell types, by targeting specific bacterial and mammalian growth phases, and by collecting VOCs through dynamic or active sampling which has the advantage of measuring VOC metabolism in relation to VOC release time and rate.<sup>33,34</sup> Finally, we recognise the use of immortalised mammalian cells lines such as A549 may not represent normal mammalian cell metabolism and response to infection and a more representative model would include patient-derived primary cells.

We demonstrated for the first time a direct VOC sampling method for ALI cultures of bacterial infection and using standard 2D culture well inserts. TFME is a highly versatile method<sup>31,35</sup> and was applied to ALI cultures with minimal intervention, thus preserving routine mammalian lung cell propagation techniques. ALI culture is physiologically rele-

vant and resembles the environment of the respiratory epithelium more closely than standard sedimented liquid cultures. Furthermore, ALI is an important tool to study the effects of nebulised drugs, exogenous pollutants and particulate matter on airway cells. Cultures can be further enhanced to develop a true lung infection model to include bronchial epithelial cells for cilia, goblet cells for mucus production, and include inflammatory cells for host response to xenobiotics.<sup>36</sup>

# 5. Conclusions

In this study we have detected VOCs characteristic of *S. aureus* infection in lung epithelial cells, using a novel approach which harnesses the adaptability of TFME to sample from a two-dimensional air–liquid interface culture model. We identified VOCs that are unique to *S. aureus*, and to infected and uninfected mammalian cells. These results demonstrate the poten-

Paper Analyst

tial for investigating VOCs as potential biomarkers of infection using complex two-dimensional culture models.

# Conflicts of interest

There are no conflicts to declare.

# **Acknowledgements**

We acknowledge the experimental support provided by Kai Wang and Professor Clare Mills at the Manchester Institute of Biotechnology and Dr John Blaikley and Dr Conal Hayton for donation of the immortalised cell line. E. B. also acknowledges the fellowship support of the European Cystic Fibrosis Society. W. A. and S. J. F. are supported by the NIHR-Manchester Biomedical Research Centre.

# References

- 1 A. J. Fischer, S. B. Singh, M. M. LaMarche, L. J. Maakestad, Z. E. Kienenberger, T. A. Peña, et al., Sustained Coinfections with Staphylococcus aureus Pseudomonas aeruginosa in Cystic Fibrosis, Am. I. Respir. Crit. Care Med., 2021, 203(3), 328-338.
- 2 J. Vincent, J. Rello, J. Marshall, E. Silva, A. Anzueto, C. Martin, et al., International Study of the Prevalence and Outcomes of Infection in Intensive Care Units, J. Am. Med. Assoc., 2009, 302(21), 2323-2329.
- 3 D. Koulenti, E. Tsigou and J. Rello, Nosocomial pneumonia in 27 ICUs in Europe: perspectives from the EU-VAP/CAP study, Eur. J. Clin. Microbiol. Infect. Dis., 2017, 36(11), 1999-
- 4 D. G. Wootton, S. J. Aston and T. W. Felton, The challenge of antimicrobial prescribing for hospital-acquired pneumonia, J. Hosp. Infect., 2020, 104(2), 198-199.
- 5 W. M. Ahmed, O. Lawal, T. M. Nijsen, R. Goodacre and S. J. Fowler, Exhaled Volatile Organic Compounds of Infection: A Systematic Review, ACS Infect. Dis., 2017, 3(10), 695-710.
- 6 L. D. J. Bos, P. J. Sterk and M. J. Schultz, Volatile Metabolites of Pathogens: A Systematic Review, PLoS Pathog., 2013, 9(5), e1003311.
- 7 A. H. Neerincx, B. P. Geurts, M. F. J. Habets, J. A. Booij, J. van Loon, J. J. Jansen, et al., Identification of Pseudomonas aeruginosa and Aspergillus fumigatus mono- and co-cultures based on volatile biomarker combinations, J. Breath Res., 2016, 10(1), 016002.
- 8 M. Nasir, H. D. Bean, A. Smolinska, C. A. Rees, E. T. Zemanick and J. E. Hill, Volatile molecules from bronchoalveolar lavage fluid can 'rule-in' Pseudomonas aeruginosa and 'rule-out' Staphylococcus aureus infections in cystic fibrosis patients, Sci. Rep., 2018, 8(1), 1-11.
- 9 W. Filipiak, R. Beer, A. Sponring, A. Filipiak, C. Ager, A. Schiefecker, et al., Breath analysis for in vivo detection of

- pathogens related to ventilator-associated pneumonia in intensive care patients: a prospective pilot study, J. Breath Res., 2015, 9(1), 016004.
- 10 C. L. Jenkins and H. D. Bean, Influence of media on the differentiation of Staphylococcus spp. By volatile compounds, I. Breath Res., 2020, 14(1), 016007.
- 11 O. Lawal, H. Muhamadali, W. M. Ahmed, I. R. White, T. M. E. Nijsen, R. Goodacre, et al., Headspace volatile organic compounds from bacteria implicated in ventilatorassociated pneumonia analysed by TD-GC/MS, J. Breath Res., 2018, 12(2), 026002.
- 12 J. Zhu, H. D. Bean, M. J. Wargo, L. W. Leclair and J. E. Hill, Detecting bacterial lung infections: in vivo evaluation of in vitro volatile fingerprints, J. Breath Res., 2013, 7(1),
- 13 H. D. Bean, J. Jimenez-Diaz, J. Zhu and J. E. Hill, Breathprints of model murine bacterial lung infections are linked with immune response, Eur. Respir. J., 2015, 45(1), 181-190.
- 14 S. Traxler, G. Barkowsky, R. Saß, A. C. Klemenz, N. Patenge, B. Kreikemeyer, et al., Volatile scents of influenza A and S. pyogenes (co-)infected cells, Sci. Rep., 2019, 9(1), 1-12.
- 15 O. Lawal, H. Knobel, H. Weda, L. D. Bos, T. M. E. Nijsen, R. Goodacre, et al., Volatile organic compound signature from co- culture of lung epithelial cell line with Pseudomonas aeruginosa, Analyst, 2018, 3148-3155.
- 16 D. Baldassi, B. Gabold and O. M. Merkel, Air-Liquid Interface Cultures of the Healthy and Diseased Human Respiratory Tract: Promises, Challenges, and Future Directions, Adv. Biomed. Res., 2021, 1(6), 2000111.
- 17 M. Schivo, A. A. Aksenov, A. L. M. M. McCartney, J. Simmons, R. W. Harper, et al., Volatile emanations from in vitro airway cells infected with human rhinovirus, J. Breath Res., 2014, 8(3), 037110.
- 18 M. S. Yamaguchi, M. M. McCartney, A. L. Linderholm, S. E. Ebeler, M. Schivo and C. E. Davis, Headspace sorptive extraction-gas chromatography-mass spectrometry method to measure volatile emissions from human airway cell cultures, J. Chromatogr. B: Anal. Technol. Biomed. Life Sci., 2018, 1090(January), 36-42.
- 19 M. S. Yamaguchi, M. M. McCartney, A. K. Falcon, A. L. Linderholm, S. E. Ebeler, N. J. Kenyon, et al., Modeling cellular metabolomic effects of oxidative stress impacts from hydrogen peroxide and cigarette smoke on human lung epithelial cells, J. Breath Res., 2019, 13(3), 036014.
- 20 M. Ashrafi, Y. Xu, H. Muhamadali, I. White, M. Wilkinson, K. Hollywood, et al., A microbiome and metabolomic signature of phases of cutaneous healing identified by profiling sequential acute wounds of human skin: An exploratory study, PLoS One, 2020, 15(2), e0229545.
- 21 W. M. Ahmed, P. Geranios, I. R. White, M. Bromley, O. Lawal, T. Nijsen, et al., Development of an adaptable headspace sampling method for metabolic profiling of the fungal volatome, Analyst, 2018, 4155-4162.

- 22 L. W. Sumner, A. Amberg, D. Barrett, M. H. Beale, R. Beger, C. A. Daykin, *et al.*, Proposed minimum reporting standards for chemical analysis, *Metabolomics*, 2007, 3(3), 211– 221.
- 23 J. J. Grandy, E. Boyaci and J. Pawliszyn, Development of a Carbon Mesh Supported Thin Film Microextraction Membrane As a Means to Lower the Detection Limits of Benchtop and Portable GC/MS Instrumentation, *Anal. Chem.*, 2016, 88(3), 1760–1767.
- 24 M. R. Kiedrowski, A. E. Paharik, L. W. Ackermann, A. U. Shelton, S. B. Singh, T. D. Starner, *et al.*, Development of an in vitro colonization model to investigate Staphylococcus aureus interactions with airway epithelia, *Cell. Microbiol.*, 2016, **18**(5), 720–732.
- 25 G. Preti, E. Thaler, C. W. Hanson, M. Troy, J. Eades and A. Gelperin, Volatile compounds characteristic of sinusrelated bacteria and infected sinus mucus: Analysis by solid-phase microextraction and gas chromatography-mass spectrometry, J. Chromatogr. B: Anal. Technol. Biomed. Life Sci., 2009, 877(22), 2011–2018.
- 26 R. Thriumani, A. Zakaria, Y. Z. H. Y. Hashim, A. I. Jeffree, K. M. Helmy, L. M. Kamarudin, *et al.*, A study on volatile organic compounds emitted by *in vitro* lung cancer cultured cells using gas sensor array and SPME-GCMS, *BMC Cancer*, 2018, 18(1), 1–17.
- 27 C. N. H. Marques, D. G. Davies and K. Sauer, Control of biofilms with the fatty acid signaling molecule cis-2-Decenoic acid, *Pharmaceuticals*, 2015, 8(4), 816–835.
- 28 D. G. Davies and C. N. H. Marques, A fatty acid messenger is responsible for inducing dispersion in microbial biofilms, *J. Bacteriol.*, 2009, **191**(5), 1393–1403.
- 29 M. W. Frank, S. G. Whaley and C. O. Rock, Branched-chain amino acid metabolism controls membrane phospholipid

- structure in Staphylococcus aureus, *J. Biol. Chem.*, 2021, **297**(5), 1–14.
- 30 X. Chen, W. Ping Teoh, M. R. Stock, Z. J. Resko and F. Alonzo, Branched chain fatty acid synthesis drives tissue-specific innate immune response and infection dynamics of Staphylococcus aureus, *PLoS Pathog.*, 2021, 17(9), 1–22.
- 31 W. Filipiak, K. Jaroch, P. Szeliska, K. Żuchowska and B. Bojko, Application of thin-film microextraction to analyze volatile metabolites in a549 cancer cells, *Metabolites*, 2021, **11**(10), 704.
- 32 V. Kapishon, G. K. Koyanagi, V. Blagojevic and D. K. Bohme, Atmospheric pressure chemical ionization mass spectrometry of pyridine and isoprene: potential breath exposure and disease biomarkers, *J. Breath Res.*, 2013, 7(2), 026005.
- 33 A. Baranska, A. Smolinska, A. W. Boots, J. W. Dallinga and F. J. van Schooten, Dynamic collection and analysis of volatile organic compounds from the headspace of cell cultures, *J. Breath Res.*, 2015, 9(4), 047102.
- 34 W. M. Ahmed, P. Geranios, I. R. White, M. Bromley, O. Lawal, T. Nijsen, et al., Development of an adaptable headspace sampling method for metabolic profiling of the fungal volatome, Analyst, 2018, 143, 4155–4162.
- 35 T. Koehler, I. Ackermann, D. Brecht, F. Uteschil, J. Wingender, U. Telgheder, *et al.*, Analysis of volatile metabolites from in vitro biofilms of Pseudomonas aeruginosa with thin-film microextraction by thermal desorption gas chromatography-mass spectrometry, *Anal. Bioanal. Chem.*, 2020, **412**(12), 2881–2892.
- 36 V. V. Thacker, N. Dhar, K. Sharma, R. Barrile, K. Karalis and J. D. McKinney, A lung-on-chip infection model reveals protective and permissive roles of alveolar epithelial cells in tuberculosis, *bioRxiv*, 2020.