1	Combined use of bacteriophage K and a novel bacteriophage to reduce Staphylococcus
2	aureus biofilm
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# 23 ABSTRACT

24 Biofilms are major causes of impairment of wound healing and patient morbidity. One of the most common and aggressive wound pathogens is *Staphylococcus aureus*, displaying a 25 26 a large repertoire of virulence factors and commonly reduced susceptibility to antibiotics, 27 such as the spread of methicillin-resistant S. aureus (MRSA). Bacteriophages are obligate 28 parasites of bacteria. They multiply intracellularly and lyse their bacterial host releasing their progeny. We isolated a novel phage, DRA88 that has a broad host range amongst S. 29 30 *aureus*. Morphologically the phage belongs to the *Myoviridae* family and comprises a large 31 dsDNA genome of 141,907 bp. DRA88 was mixed with phage K to produce a high titre mixture that showed strong lytic activity against a wide range of S. aureus isolates 32 including representatives of the major international MRSA clones and coagulase-negative 33 34 Staphylococcus. Its efficacy was assessed both in planktonic cultures and when 35 treating established biofilms produced by three different biofilm producing S. aureus 36 isolates. A significant reduction of biofilm biomass over 48 hours of treatment was recorded in all cases. The phage mixture may form the basis of an effective treatment 37 38 for infectious caused by S. aureus biofilms.

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41 Key words: *Staphylococcus aureus*, bacteriophage therapy, biofilm, antibiotic resistant
42 bacteria, MRSA, sequence analysis.

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### 47 **INTRODUCTION**

Staphylococcus aureus is an opportunistic human bacterial pathogen that primarily 48 colonizes the anterior nares (1) but is frequently shed onto skin surfaces. When an 49 50 opportunity occurs that facilitates its penetration of the skin surface, it is able to cause a 51 broad spectrum of human diseases, from skin and soft-tissue infections to systemic 52 infections such as pneumonia, meningitis and osteomyelitis (1, 2). Host invasion and immune evasion is possible due to a myriad of virulence factors, including toxins, adhesins and 53 54 evasins (3). In addition, S. aureus infections often comprise strains with antibiotic resistance. Examples of these are penicillin and methicillin resistant S. aureus (MRSA) and isolates with 55 decreased susceptibility to vancomycin (4). 56

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S. aureus is one of the most common gram-positive causes of wound infectious (5, 6). The 58 59 wound environment is an ideal one for establishment of a bacterial infection as it contains 60 large aggregations of necrotic tissue and accumulations of protein exudate (7). It is also observed that wound infections are strongly associated with the formation of biofilm 61 62 communities (8). Once in a biofilm, bacterial cells experience greater protection against 63 antibiotics and against elements of the host immune system, when compared to cells growing in a planktonic state (9). For example, the exopolysaccharide matrix blocks 64 antibody penetration into biofilm (10) and phagocytes are unable to interact with bacterial 65 cells (11). Biofilm structures are believed to be present in many acute wounds, but are very 66 common in chronic wounds (12), and are a major factor delaying wound healing (12, 13). 67 Moreover, many biofilms colonize implanted medical devices (14) greatly increasing 68 69 patient morbidity and mortality and they are associated with larger health care costs (15). 70 Chronic wounds affect 6.5 million patients in the United States and more than US\$25 billion are spent every year on treatment (16). 71

Current antibiotic options to successfully treat *S. aureus* are becoming scarce despite the development of some novel drugs (17), and there is a growing need for effective agents to combat infections (18). Bacteriophage therapy is a viable alternative / adjunct to antibiotics in treating bacterial infection (for review see (19)). Bacteriophages (phages) are viruses able to infect highly specifically and kill the bacterial species targeted, but not eukaryotic cells. The phage-encoded lysis proteins – endolysin and holin - cause the breakdown of the bacterial membrane (20) resulting in cell death and release of phage particles.

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80 Several studies have shown the potential of using phages to treat S. aureus infections (21–23) 81 and it has been demonstrated that phages also have the ability to disrupt bacterial biofilms 82 (24). Phages are increasingly recognized as serious candidates in the fight against antibiotic 83 resistant bacteria in human therapeutics and as prophylaxis (25). Phages with strictly lytic life cycles, which result in a rapid killing of the target host and diminish the chances for 84 85 bacteria to evolve towards phage resistance, are preferred for bacteriophage therapy use (26). It is also of value for the phage (or phage mixture) used to have a polyvalent nature, 86 i.e. is able to infect a large set of strains within a species and may show improved 87 applicability in situations where the etiological agent of an infectious disease has not been 88 identified. 89

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Here we investigate the potential of using two lytic and polyvalent *S. aureus* phages: K, a well-documented staphylococcal phage (27–29) that attaches specifically to the cell wall teichoic acid (30) providing a wide host range; and a newly isolated phage, DRA88. Phage K was, previously, shown to be able to disrupt biofilms produced by *S. aureus* strains (28). Here, we characterized the antimicrobial activity of the phages alone and in combination in planktonic and biofilm systems.

### 97 MATERIALS AND METHODS

# 98 Bacteria and Growth Conditions

99 S. aureus strains (listed in Table 1) used in this study were from our collection of >5000100 clinical isolates and they were selected to be genetically diverse (by multilocus sequence 101 typing (31)) and also to contain members of the major MRSA and MSSA clones present 102 worldwide. Examples of eight coagulase-negative staphylococci: S. xylosus, S. sciuri subsp. 103 sciuri, S. chromogenes, S. hvicus, S. arlettae, S. vitulinus, S. simulans and S. epidermidis 104 were also included in this study. Bacteria from 5% (v/v) blood agar plates were grown at 37 105 °C with constant shaking (170 rpm) in Tryptic Soy Broth (TSB). Tryptic Soy Agar (TSA) 106 and TSB-soft-agar containing 0.65% of bacteriological agar were used for bacteriophage 107 propagation and plaque count assays. Note that media was supplemented with 1 mM CaCl<sub>2</sub> 108 and MgSO<sub>4</sub> to improve phage adsorption (32). Bacterial aliquots were stocked at  $-80^{\circ}$ C in 109 broth containing 15% glycerol (v/v).

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# 111 Bacteriophage Isolation

112 Bacteriophages were isolated from crude sewage (Thames Water PLC, Luton, UK). 113 Bacterial enrichments with S. aureus isolates were performed to increase phage numbers as follows: 5 ml of actively growing S. aureus cells (from overnight liquid culture in TSB) 114 and TSB supplemented with 1 mM MgSO<sub>4</sub> and 1 mM CaCl<sub>2</sub>. The enrichment was 115 incubated overnight at 37 °C. A 10 ml aliquot was taken from the overnight culture and 1 M 116 117 NaCl and 0.2% chloroform were added. The culture was then centrifuged (30 min, 3000 x g) to remove bacteria and the supernatant was filtered sterilized (0.22  $\mu$ m pore size, 118 119 Millipore filter). This lysate (supernatant) was used to check the presence of lytic phages 120 using the double layer method as described previously (33). Isolated single plaques were 121 picked into SM buffer (5 M NaCl, 1M MgSO4, 1 M Tris-HCl [pH 7.5], gelatine solution distilled water) and successive rounds of single plaque purification were carried out until
purified plaques were obtained, reflected by single plaque morphology. Purified phages
were stored in 50% glycerol (v/v) at -80 °C for long term use. Short term stock
preparations were maintained at 4 °C.

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### 127 Bacteriophage Propagation

128 Phage lysates were propagated on their respective bacterial hosts. Briefly, 100  $\mu$ l of phage 129 lysate and 100  $\mu$ l of host growing culture were mixed and left for 5 min at room temperature. 130 Afterwards 3 ml of soft-agar was added and poured onto TSA plates. The following day, 131 after an overnight incubation at 37 °C, plates displaying confluent lysis were selected and 3 132 ml of SM buffer and 2% chloroform were added before incubating at 37 °C for 4 h. High titre phage solution was removed from the plates, and centrifuged (8,000 x g, 10 min) to 133 134 remove cell debris, then filter sterilized (pore size, 0.22 µm) and stored at 4 °C. Both phages 135 were propagated in the prophage-free isolate S. aureus RN4220 (40) to avoid potential 136 contamination with mobilized phages (34).

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# 138 Sensitivity Assay

To determine phage sensitivity of bacterial isolates spot tests were performed. Briefly, 3 ml 139 140 of TSB-soft-agar was added to 100 µl of host growing culture and poured onto TSA. Plates were left to dry for 20 min at 37 °C. The different phage lysates were standardized to a 141 titre of  $10^9$  PFU/ml and 10 µl of each lysate were spotted onto the bacterial lawns. This 142 143 assay was performed in triplicate. The plates were allowed to dry before incubating 144 overnight at 37 °C. The following day the sensitivity profiles of each of the bacterial strains were determined: if the bacterial lawn was lysed; slightly disrupted; or not disrupted, the 145 146 bacterial isolate was labelled: sensitive; intermediate; or resistant to the phage infection.

### 147 **Bacteriophage adsorption**

The experiment was carried out at 37 °C under constant shaking (60 rpm) and a phage inoculum of MOI (Multiplicity Of Infection) = 0.01. The number of free phages was calculated from the PFU of chloroform-treated samples within 5 min after inoculation. The adsorption rate, assuming a first order kinetics was calculated in terms of the percentage of free phage loss by fitting phage decay curve (normalized as a percentage) to the rate equation:

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155 
$$\ln (\% \text{ phage})_t = \ln (\% \text{ phage})_o - k't$$
 (1)

156

157 Where k' is the pseudo 1st rate constant for free phage loss:

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159 
$$k' = k[bacteria]$$
 (2)

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161 From this, the percentage phage remaining at any time t can be easily calculated.

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# 163 **One-step growth curve**

164 Phage growth cycle parameters - the latent period (L), eclipse period (E) and burst size (B), were determined from the dynamical change of the number of free and total phages. 165 166 Hence, one-step growth curves were measured as described by Pajunen et al. (35) with 167 some modifications: 10 ml of a mid-exponential-phase culture were harvested by 168 centrifugation (7,000x g, 10 min, 4°C) and resuspended in 5 ml TSB to obtain OD<sub>600</sub> of 1.0. To this suspension, 5 ml of phage solution were added to obtain a MOI (multiplicity of 169 170 infection) of 0.001. Phages were allowed to adsorb for 5 min at room temperature. The 171 mixture was than centrifuged as described above and the pellet was resuspended in 10 ml of fresh TSB medium. Two samples were taken every 5 min over a period of 1 h at 37 °C under constant shaking. The first samples were plated immediately without any treatment and the second set of samples was plated after treatment with 1 % (v/v) chloroform to release intracellular phages.

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### 177 Measurement of phage zeta potential and size

The particle size and electrophoretic mobility of the phages was measured by Dynamic Light Scattering (DLS) using a Zetasizer Z/S (Malvern, UK) at 37 °C. Cuvettes filled with the sample were carefully inspected to avoid air bubbles. Phages were diluted in  $dH_2O$  to a final concentration of  $10^8$  PFU ml<sup>-1</sup>. Measurements were repeated at least three times.

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# **183** Electron Microscopy

Phage particles in water were deposited on carbon coated copper grids and negatively stained
with 1% uranyl acetate (pH 4). Visualization was performed using a transmission electron
microscope (TEM) (JEOL JEM1200EXII, Bath, UK) operated at 120 kV.

187

### **188 Phage DNA Extraction**

Concentrated phage preparations were obtained by a caesium chloride (CsCl) (Sigma, UK) gradient composed of three different solutions with densities of: 1.35, 1.50 and 1.7 g ml<sup>-1</sup>. They were prepared in a 36.5 ml ultracentrifuge tube (Beckman Coulter, Seton Scientific, UK). For the preparation of CsCl solution at a given density,  $\rho$  (g ml<sup>-1</sup>), the following equation was used to calculate the final CsCl concentration, c (g ml<sup>-1</sup>): c = $0.0478\rho^2 + 1.23\rho - 1.27$  as well as the protocol described previously by Boulanger (36). After ultracentrifugation for 3 hours (75,600 g at 4 °C) the phage band was collected and 196 dialysis was performed to remove CsCl residuals. Briefly, phage suspension was washed into dialysis cassettes (Slide-A-Lyser, Fisher, UK), which in turn were introduced in 500 197 volumes of dialysis buffer (10 mM sodium phosphate, pH 7) for 30 min. After performing 198 199 the process three times the concentrated and purified phage suspension was collected. 200 Phenol/chloroform extraction was performed according to (37). 1.8 ml of phage lysates were treated with 18µl DNase I (1 mg ml<sup>-1</sup>) (Sigma Aldrich) and 8 µl RNase A (10 mg ml<sup>-1</sup>) 201 (Sigma Aldrich) and incubated at 37 °C for 60 min. Subsequently, 18 µl proteinase K (10 202 mg ml<sup>-1</sup>) (Sigma Aldrich), 1% of sodium dodecyl sulphate (SDS) and 1 mM EDTA.Na<sub>2</sub> 203 204 were added to the samples and these were incubated at 65 °C for further 60 min. All 205 protein material was eliminated by using phenol:chloroform:isoamyl alcohol (25:24:1) 206 (Sigma Aldrich) and DNA extraction and precipitation was performed as described 207 previously elsewhere (38). Nucleic acid concentration and quality was assessed with a 208 Nanodrop spectrophotometer (Thermo Scientific, UK).

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# 210 DNA sequencing, analysis and assembly

DNA sequencing libraries were prepared using the Nextera® XT DNA kit (Illumina, San Diego, USA) according to the manufacturer's protocol. Individually tagged libraries were sequenced as a part of a flowcell as 2x250 base paired-end reads using the Illumina MiSeq platform (Illumina, San Diego, USA) at The Danish National High-Throughput DNA-Sequencing Centre. Reads were analysed, trimmed and assembled using 6.5.1 CLC Genomic Workbench as described before by Kot *et al.* (39). Genes were predicted and annotated using RAST server (40).

# 218 Planktonic cultures treated with phage mixture

Planktonic cultures were performed in 96-well microtitre plates and their optical densitywas measured. In summary, a 1:100 dilution was prepared in the wells of the microplate by

adding 2  $\mu$ l of an overnight culture to 198  $\mu$ l of TSB. After 2 hours of incubation at 37 °C, the single or mixed phage at a MOI of 0.1 was added and the microplates were incubated for a further 24 hours. The incubation was followed on a plate reader (FLUOstar Omega, BMG LabTech, UK) where the growth curves were established at an optical density at 590 nm (OD<sub>590</sub>). This approach allows observation of the phage-bacteria interaction over time and also allows for monitoring of the appearance of resistant mutants for each phage lysate.

228

# 229 **Biofilm formation**

230 Biofilm assay was performed similarly to previously described methods (41), but with 231 some modifications in order to optimize the system. Biofilm formation was performed 232 in 96-well polystyrene tissue culture microplates (Nunclon<sup>™</sup> Delta Surface, Nunc, UK) to 233 achieve an improved cell attachment. TSB supplemented with 1% D-(+)-glucose (TSBg) and 1% NaCl (TSBg+NaCl) was used to perform this assay as this helps to improve 234 biofilm formation (37, 38). An overnight culture was diluted to a titre of 10<sup>8</sup> cfu ml<sup>-1</sup>. 235 236 Briefly, in the microplate wells a 1:100 dilution was performed by adding 2  $\mu$ l of the bacterial suspension to 198 µl of TSBg+NaCl, making a starting inoculum of 10<sup>6</sup> cfu ml<sup>-1</sup>. 237 238  $200 \ \mu$ l of broth were added to a set of wells as a negative control. All wells were replicated three times. Afterwards, microplates were incubated at 37 °C for 48 hours with no shaking 239 240 for biofilm formation. During the incubation time (~24 hours after incubation), 50  $\mu$ l of 241 fresh TSBg + NaCl was added to all control and test wells. Following incubation, medium 242 was poured off and wells were carefully washed twice with sterile phosphate buffered 243 saline (PBS) solution (Sigma Aldrich, UK) to remove any planktonic cells. Microplates were 244 allowed to dry for 1 hour at 50 °C. To determine total biofilm biomass microplate wells were stained with 0.1% Crystal Violet (CV). After staining the wells were washed twice 245

with PBS solution and dried. Biofilm formation was determined by visual comparison of thestained wells and photographed.

248

# 249 **Biofilm treatment with phage mixture**

Biofilm formation was carried out as described above. Once biofilms were established and 250 251 washed once with PBS, 100 µl of phage mixture in PBS was added to a set of wells. Two 252 different MOIs were set up for the single or mixed phage: 1 and 10. 100 µl of PBS were 253 added both to the isolate positive and negative controls. All the experiments were performed three times. After static incubation at 37 °C microplates were washed and stained, as 254 255 described before, at pre-determined time-points. To perform optical density readings of the 256 staining intensity, 100  $\mu$ l of 95% ethanol (v/v) was added to each well and optical density at 257 590 nm (OD590) was taken using a plate reader.

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# 259 Data analysis

Comparisons between the different time points and the positive controls were made by performing a *Student's t-test* and a *p-value* < 0.05 was considered statistically significant for all cases. All tests were performed with a confidence level of 95%. Spread of data at the 95% confidence interval (CI) was estimated using the Winpepi freeware statistical analysis program (42).

265

# 266 Nucleotide sequence accession number

267 The genome sequence of phage DRA88 has been deposited in the GenBank database268 under accession number KJ888149.

269

# 271 **RESULTS**

# 272 Isolation and host range determination

273 Several lytic phages were isolated in crude sewage samples from a water treatment facility 274 and tested against the S. aureus bacterial collection listed in Table 1 in order to isolate phages with broad host ranges. One phage, DRA88, presented a host infectivity coverage of 60.0% 275 276 (95% CI: 50% - 69%; 57 isolates out of 95 were susceptible) and so was selected for further 277 studies. Phage K was also tested and showed host coverage of 64.2% (95% CI: 54% - 73%; 278 61 isolates out of 95 were susceptible). These two phages were mixed in a phage mixture giving a total coverage of 73.7% (95% CI: 64% - 82%) of the S. aureus isolates assigned 279 280 to 14 different MLST (Multilocus Sequence Typing) types. Infectivity of both phages was 281 also tested on a group of coagulase-negative isolates, where DRA88 did not kill efficiently 282 any of the isolates. Phage K showed also a weakly infectivity, however two of the isolates (S. 283 simulans and S. hyicus) were sensitive (Table. 2).

284

### 285 Phage Growth Characteristics

286 The life cycle and adsorption affinity of phage DRA88 and phage K were assessed when growing in RN4220 S. aureus host at 37 °C. Firstly, one-step growth studies were 287 288 performed to identify the different phases of a phage infection process. After infection, 289 phage growth cycle parameters (L - latent, E - eclipse, and B - burst size) were determined 290 (Fig. 1). In the system established, the eclipse and latent periods of DRA88 were, 15 291 min and 25 min, respectively. DRA88 yielded a burst size of 76PFU and phage K of 125 292 PFU per infected cell after 60 min. These phage life cycle values are in conformity with the 293 values normally observed for T7 group phages (43). The adsorption efficiency of phages to 294 the host was estimated with cells in the early logarithmic growth phase (Fig. 2). The adsorption rate (adsorption affinity) of the phages, when infecting actively growing S. 295

*aureus* RN4220, was measured. From equation (1), the rate constant for the adsorption (loss of free phage) for phage K and DRA88 were calculated (Fig. 2), k' = 0.352 min-1 and k' = 0.252 min-1, respectively. Hence, although similar, after 5 minutes, 80.3% of phage K was adsorbed to the bacteria and 71.6% of DRA88. After 10 min, values for free viral particles were below 5% for both phages.

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### **302** Morphology of phage particles

303 The isolated phage DRA88 was further characterized with regard to its morphology. Images of the phage DRA88 were developed using Transmission Electron Microscopy 304 305 (TEM). The results revealed that phage DRA88 has, along with an icosahedral head of  $\sim 78$ nm in diameter, a long contractile tail of ~179 nm with tail fibers (Fig. 3A). Therefore it 306 307 can be classified as belonging to the *Myoviridae* family (order *Caudoviridales*), 308 according to the classification system of Ackermann (44). It was observed for phage 309 DRA88 that besides single phage particles there were also various aggregates made through contact of their tail fibers (Fig. 3B). Phage K was also observed under TEM (Fig. 310 311 S1), revealing a viral particle also belonging to the *Myoviridae* family (27). The zeta 312 potential of phage DRA88 and phage K calculated from the electrophoretic mobility was 313 -17 mV and -26.3 mV, respectively. The size measured was 122 nm for phage K, but 314 according to the TEM micrographs and the literature this phage has an average size of 280 315 nm total (45). This discrepancy could be due to contraction of tails, which was observed 316 frequently, and can interfere with the measurement. Note that Dynamic Light Scattering 317 (DLS) may show lower accuracy when measuring irregular shaped particles, such as tailed 318 phages, where the size is not uniform in all dimensions. Regarding phage DRA88 we were 319 not able to obtain an accurate size measurement and this can be related to the phenomena of 320 aggregation observed under TEM.

### 321 Genomic characterization of DRA88 and comparison with phage K

322 To gain a more ample understanding of the phage DRA88, its DNA was extracted and 323 genome sequencing performed. Upon assembly and annotation it was found that phage 324 DRA88 has a large double stranded DNA genome with terminal redundancy, which 325 suggests that phage DRA88 has a headful packaging system (46). The genome comprises 326 141,907 bp and can be grouped into the class III of Staphylococcal phages (>125kbp) (47); 327 204 putative coding regions and four tRNA genes were identified (Fig. 5 and Table 1 in 328 Supplementary Data). The gene coding potential with 1.44 genes per kb, exhibits a high gene density. The majority of genes, 145 (71%), are found in the forward strand and 59 329 330 (29%) in the opposite strand. tRNA genes are all located in the reverse strand of the genome. Regarding the GC% content it shows 30.4%, a lower percentage than the one 331 332 found in the S. aureus host -32.9%. The amino acid sequence was found to share strong 333 similarities (>95%) with several other phages, such as JD007 and GH15 previously 334 sequenced (48, 49). A comparison between phage DRA88 and phage K was performed 335 using the BLAST algorithm (48). DRA88 genome seems to be organized into functional 336 modules - cell lysis, DNA replication and structural elements - highly similar to the 337 organization of phage K and other staphylococcal Myoviridae phages belonging to the 338 Twort-like viruses (47, 50, 51). Between these modules we can find several putative coding 339 regions that are not yet found in the NCBI database or have no attributed function (phage and hypothetical proteins). These unknown functions represent 84.81% of the 340 coding capacity. Three potential coding regions (orf178, orf192 and orf195) did not have 341 342 any identical match with phage genes in the NCBI database. DRA88 lysin and DNA 343 polymerase are not interrupted by introns (49), contrary to phage K, but similarly to phage 344 GH15. At the end of the genome (between orf164 and orf182) there is a large coding region with unidentified functions inserted into the DRA88 genome that is not 345

observed in phage K. Also, DRA88 genome analysis did not reveal similarity to known
virulent associated or toxin proteins.

348

### **Bacteriophage mixture inhibits planktonic bacterial growth**

350 The efficacy of single DRA88 and phage K and their combination was assessed when treating 351 bacterial cultures. Single phages and the phage mixture (MOI = 0.1) were introduced in a bacterial culture already growing for two hours under planktonic conditions and let to 352 353 incubate for further 22 hours (Fig. 5). Single phage treatment showed to be less 354 successful in general than the phage mixture and this was cleared observed for S. aureus 355 15981 isolate (Fig 5A). For all three S. aureus bacterial cultures - 15981, MRSA252 and H325 – no cell growth was observed when the phage mixture was present compared to 356 357 bacterial growth only. In fact, we observed efficient inhibition of the bacterial growth for 358 15981 and H325. The same was not observed for MRSA 252, where we observed 359 bacterial growth after 18 hours of treatment (time = 20 hours).

360

#### **Biofilm eradication**

362 Biofilms produced by the three S. aureus isolates were established in microtitre plates. 363 All isolates were strong biofilm producers, however MRSA 252 followed by H325 isolates produced lower biofilm biomass and more fragile structure (when performing the 364 365 mechanical washing steps the biofilm was more susceptible to disruption). The established 366 biofilms were treated with single and mixed phage at MOI 10 and the biofilm was assessed 367 (Figs 6 and 7). Measurement of biofilm density made using crystal violet / OD590 nm 368 over 48 hours showed a clear reduction following phage inoculation compared with nontreated controls (p = < 0.05). A decrease in biofilm biomass from the mixture compared 369 with the single phages after 48 hours of treatment was seen in all cases, although not 370

371	significant it can been observed by eye (along with CV stained wells). Established biofilms
372	were also treated with the phage mix at two different MOIs: 1 and 10, and the biofilm was
373	assessed at the time points: 0h, 2h, 4h, 24h and 48h (Fig 8). Unsurprisingly, phage mixtures
374	with higher MOIs (10 compared with 1) gave a more rapid reduction in biofilm density,
375	although both MOIs of 1 and 10 resulted in the same endpoint after 48 hours. For S.
376	aureus 15981 biofilms treated with MOI 10, 4 hours after treatment, there was already
377	more than 50% biofilm reduction ( <i>p</i> -value $< 0.05$ ) and after 48 hours of treatment the
378	biofilm biomass was almost completely disrupted (MOI 10, <i>p</i> -value = $4.82 \times 10^{-3}$ ; MOI 1,
379	<i>p-value</i> = $1.47 \times 10^{-5}$ ) (Fig. 8A). Figure 8B shows that biofilms produced by MRSA 252
380	were eliminated by more than 50% (MOI 10, <i>p-value</i> = 0.003; MOI 1, <i>p-value</i> = 0.012) after
381	48 hours of phage treatment. At last, biofilms produced by H325 were not initially as strong
382	as the other isolates. However we were able to observe a reduction of the biofilms as well
383	over 48 hours for MOI 10 ( <i>p-value</i> = $0.049$ ). For MOI 1 a reduction of the biofilm
384	structure was observed after 24 hours ( <i>p</i> -value = $0.034$ ). However such reduction was not
385	observed after 48 hours.
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#### 395 **DISCUSSION**

*S. aureus* biofilms in wounds and catheter sites present particular problems to patients , increasing morbidity, mortality and difficulty in delivering effective chemotherapy. For wound healing to occur, treatment of the biofilm infection is essential and often requires selection of the correct antibiotic. The choice of appropriate chemotherapy is however made more difficult due to the increasing prevalence of antibiotic resistance. Hence, solutions are required to avoid treatment delay or ineffectiveness.

402

The host range infectivity of DRA88 against a genetically diverse collection of S. aureus 403 404 isolates was broad, showing that this is a polyvalent phage. Based on TEM analysis DRA88 belongs to the Myoviridae tailed-phage family. DRA88 adsorption occurred 405 406 rapidly and at a similar rate to that observed for phage K, which is in accordance to several 407 S. aureus Myoviridae infecting phages (52, 53). An interesting observation for phage 408 DRA88 was the possible formation of phage aggregates. Phage aggregation is observed 409 occasionally in nature (54, 55) and is dependent on pH, ionic strength and the composition 410 of ions. Such phenomena could have been influenced by uranyl acetate (pH 4) used to stain the phages for TEM observation. However, DLS measurements (in d.H<sub>2</sub>O: ~pH 7) 411 412 were unsuccessful and possibly due to several sizes found in the sample (singles and 413 several aggregates), hence, we suspect that phage DRA88 is prone to form aggregates. When 414 interacting with bacterial cells, aggregation could impede phage access to the cells and 415 hence slow the rate of adsorption. Phage aggregation can be inhibited by optimization of 416 growth media composition or stabilised in nano-emulsions resulting in phages that more 417 efficiently attach to bacterial cells (56–58), consequently this could affect estimation of the 418 PFU which then may not directly correspond with the number of infective particles (MOI).

419

420 At the genome level, DRA88 was revealed to be a large dsDNA phage, usually a 421 common characteristic of *Myoviridae* virus (50), carrying a high gene density and low 422 GC%. It exhibits a high degree of relatedness to several other phages of the Twort-like 423 group, including phage K. No virulence factors in the genome were identified, according to 424 data available, suggesting that this phage could be safely used to treat S. aureus infections. 425 The majority of putative coding regions of DRA88 genome do not have yet any function 426 attributed yet, which is generally observed for the numerous phages being sequenced at the 427 current time (59). Consequently, there is a crucial need for a more comprehensive 428 investigation of phage genomes. Considering phage therapy as a therapeutic approach 429 option, it is extremely important that we expand our knowledge regarding phage genes and 430 proteins, and their respective functions and potentialities, as they can be involved in phage-431 host interaction and even code for novel virulence determinants (60).

432

433 The novel isolated phage was mixed into a phage mixture with phage K and as a result their lytic potential was increased (74 % of host coverage). The use of a phage mixture is 434 435 largely preferred over use of single phage as it results in a decreased rate of bacteria exhibiting 436 resistance (61, 62). Bacterial broth cultures growing with phage mixtures showed an 437 elimination of bacterial cells and suppression of resistant mutants. However, this was only observed for two strains; after 18 hours of growth, MRSA 252 was able to counteract the 438 439 infectivity of the phage particles present in the culture, possibly due to the presence of phage 440 receptor mutated clones present in the culture. This situation could be overcome or delayed by 441 including more phage types to the therapeutic mixture, a task far easier (more rapid and likely 442 to succeed) to undertake compared to the discovery of a new antibiotic. It has been 443 demonstrated that phage therapy could be delivered in synchronization with another antibacterial therapy, such as antibiotics (63). Reducing the initial bacterial load could be 444

- sufficient to bring the bacterial numbers under control so that the drug and also the action ofthe immune system can clear the infection (63–65) in an effective way.
- 447

More than 60% of all infections are related to the formation of biofilms (12). Hence, it is 448 449 important that biofilm model studies are investigated when testing a new antibacterial phage 450 mixture. Some studies already suggest the significant potential of phages to reduce and/or 451 eliminate biofilms. This is the case of phage K, where biofilms produced by S. aureus isolates 452 showed a remarkable decrease in their biofilm biomass after addition of the phage (28, 41). 453 Here, we showed that the phage mixture was able to reduce significantly the biofilm load on 454 the polystyrene surface a microtitre plate. For isolate 15981 the eradication effect started readily after addition of the killing mixture and after 48 hours of treatment the biofilm was 455 456 almost completely disrupted. Higher MOI showed a more rapid effect of biofilm reduction and a better prevention of biofilm regrowth was also observed for isolate H325. Comparing the 457 458 effect of phage mixture on broth cultures to biofilm we observed a more positive effect on 459 eliminating the bacterial load of broth cultures than that on biofilm, which is disrupted more 460 slowly. This scenario was hypothesized to be due the metabolic state of the cells. In biofilms, cells can be in a low metabolic activity stage and phages cannot proliferate as efficiently as in 461 462 active growing cells (41). To date, very few studies have been performed regarding phage 463 mixture treatments, especially to treat S. aureus biofilms and in particular MRSA-caused ones. Kelly et al (28) have already shown the efficacy of a phage K and derivatives mixture 464 465 on the eradication of S. aureus biofilms produced by non-human clinical isolates. Here we go 466 further by treating biofilms produced by prevalent human clinical isolates that also include 467 MRSA types. This study suggests that the utilisation of a mixture of bacteriophage i.e. phage 468 K and DRA88 in this case, could provide a practical alternative to antibiotic / antimicrobial treatments for combating some S. aureus infections and in particular the devastating effects of 469

470 MRSA infections and biofilms related, such as burn-wound or catheter infection. Even with 471 the contribution of this study on the effectiveness of bacteriophage therapy to fight established 472 bacterial infections, there is still a long way to go and several barriers to address. The safety 473 of phages and ethical and regulatory issues, for example, must be overcome in order to phages 474 become an available alternative therapeutic (see review (66)).

475

In summary, here we describe the isolation and characterization of a novel bacteriophage
against pathogenic *S. aureus* bacteria. The phage, in combination with phage K,
showed an improved range of infectivity of *S. aureus* isolates and a potent effect in
biofilm dispersion making it a good candidate for further therapeutic development.

480

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### 673 TABLE LEGENDS

Table 1: Sensitivity screening<sup>†</sup> of phage mixture against 95 *S. aureus* isolates.
Further information regarding the isolate sequence type (ST) and origin are supplied.
<sup>†</sup> Bacterial isolates were Susceptible – clear spot (S, light grey), Intermediate – turbid
spot (I, medium grey) or Resistant – no disturbance of bacterial lawn (R, dark grey) to
phage infection.

Table 2: Sensitivity screening of phage mixture against coagulase-negative staphylococci
isolates. <sup>†</sup> Bacterial isolates were Susceptible – clear spot (S, light grey), Intermediate –
turbid spot (I, medium grey) or Resistant – no disturbance of bacterial lawn (R, dark grey) to
phage infection.

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#### 684 FIGURE LEGENDS

**Figure 1:** Curve One-step growth of phage DRA88 (left) and phage K (right) in RN4220 *S. aureus* at 37 °C. Shown are the PFU per infected cell in untreated cultures (•) and in chloroform-treated cultures ( $\circ$ ) at several time points over 60 minutes. The phage growth parameters are indicated in the figure and correspond to: E - eclipse period; L - latent period and B - burst size. Each data point is the mean of three independent experiments and error bars indicate the mean  $\pm$  standard deviation.

**Figure 2:** Percentage of free (A) DRA88 and (B) Phage K phages after infection of actively growing RN4220 *S. aureus* at a MOI of 0.001 at several time points over 10 minutes. Rate constants for loss of phage are 0.352 min<sup>-1</sup> for Phage K and 0.252 min<sup>-1</sup> for DRA88. Each data point is the mean of three independent experiments and error bars indicate the mean  $\pm$  standard deviation.

697 Figure 3: Electron micrograph images of phage DRA88 negatively stained with

698 1% uranyl acetate, (A) showing the tail in a contracted position and (B)699 formation on phage particles aggregates. Scale bar is indicated.

**Figure 4:** Comparative genomic analysis of phage DRA88 and phage K. Nucleotide sequences were compared using the Artemis Comparison Tool (ACT). Predicted ORFs are denoted by arrows, tRNAs are indicated (vertical blue dashed line) and genes encoding proteins with at least 69% amino acid identity between the two genomes are indicated by shaded regions.

**Figure 5:** Dynamic of bacteria with single and phage mixture in liquid cultures over 24 hours incubation at 37 °C. Absorbance readings at 590nm were taken in a microtitre plate reader. *S. aureus* isolates: A – 15981; B – MRSA 252; C – H325 growing with only SM buffer ( $\circ$ ), with single DRA88 ( $\checkmark$ ), with single phage K ( $\triangle$ ) and with the phage mixture in SM buffer ( $\bullet$ ) and also a negative control only SM buffer ( $\bullet$ ) are shown in the figure. Assays were performed four times and OD590 was expressed as the mean ± standard deviation.

711 **Figure 6**: Normalized biofilm biomass treated with single phage K, DRA88 and the phage 712 mixture after 48 hours at MOI 10 (OD590 reading after CV staining). S. aureus isolates: 1 – 713 15981; 2 – MRSA 252; 3 – H325. Mean values for the three strains treated with: phage K =714 0.63 (SD±0.10), 0.30 (SD±0.16), 0.27 (SD±0.06); DRA88 = 0.11 (SD±0.04), 0.18  $(SD\pm0.08)$ , 0.29  $(SD\pm0.03)$  phage mixture = 0.06  $(SD\pm0.02)$ , 0.15  $(SD\pm0.02)$ , 0.23 715 716 (SD $\pm$ 0.02). Assays were performed three times and the mean  $\pm$  standard deviation is 717 indicated. Statistically significance of biofilm reduction was assessed by performing a Student's t-test. p-values are indicated (\* : <0.05). 718

Figure 7: Visualization of wells stained with 0.1% of crystal violet after 48 hours of phage treatment at MOI 10. Shown are the biofilm wells treated with PBS, phage K, DRA88 and phage mixture at 0h (A) and (B) 48 hours after. Experiments were performed in triplicate.

**Figure 8:** Normalized biofilm biomass treated with the phage mixture over 48 hours at two

723	different MOIs (OD590 reading after CV staining). S. aureus isolates: A - 15981; B -
724	MRSA 252; C - H325. Assays were performed three times and the mean $\pm$ standard
725	deviation is indicated. Statistically significance of biofilm reduction was assessed by
726	performing a Student's t-test. p-values are indicated (* : <0.05).
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748	Table 1:	Sensitivity	screening <sup>†</sup> of pl	hage mixture	against 95 S.	aureus isolates.	Further

749	information	regarding th	he isolate s	sequence type	(ST)	and origin	are supplied.
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Isolates	ST	Country	Phage K	DRA88	Phage mixture
WBG8343	1	Australia	S	I	S
MSSA H476	1	England	S	S	S
HT2001749	1	USA	S	I	S
H148	3	UK	S	S	S
CDC980193-USA800	5	USA	S	S	S
Mu3	5	Japan	S	R	S
963Small	5	USA	S	S	S
97.1948.S.	5	Scotland	R	R	R
C56	6	UK	S	S	S
C2	7	UK	Ι	Ι	I
CDC201114-USA300	8	USA	S	S	S
15981	8	Spain	I	S	S
HT20030203	8	USA	I	S	S
HT20030206	8	USA	S	I	S
C125	8	UK	Ι	Ι	Ι
Fra97392	8	France	Ι	R	Ι
EMRSA6	8	UK	Ι	Ι	Ι
99st22111	8	-	S	S	S
H169	9	UK	S	S	S
D316	11	UK	Ι	Ι	Ι
D329	12	England	S	S	S
H117	12	UK	S	S	S
H402	13	UK	S	S	S
C154	14	UK	S	S	S
C357	15	UK	S	S	S
H291	18	UK	S	Ι	S
H42	20	UK	S	S	S
HO50960412	22	UK	Ι	S	S
H182	22	England	S	S	S
C720	22	England	R	S	S
C13	22	Eire	S	S	S
C49	23	UK	S	S	S
D279	25	UK	Ι	I	I
Not116	27	UK	S	S	S
H118	28	UK	S	Ι	S
SwedenAO17934/97	30	Sweden	S	I	S
Cuba4030	30	Cuba	Ι	Ι	Ι

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C390	31	UK	S	S	S
Н399	33	UK	S	S	S
C160	34	UK	Ι	Ι	Ι
Btn766	36	UK	S	S	S
MRSA 252	36	England	S	S	S
H325	36	UK	S	S	S
EMRSA16	36	UK	S	S	S
H137	38	UK	Ι	Ι	Ι
H137MRSA	38	UK	S	S	S
C253	40	UK	Ι	I	Ι
C427	42	UK	S	S	S
Fin76167	45	Finland	S	S	S
C316	49	UK	S	S	S
H417	50	UK	S	S	S
C3	51	UK	S	S	S
D49	53	UK	S	S	S
D98	54	UK	Ι	Ι	Ι
D318	57	UK	S	S	S
D535	59	UK	I	I	Ι
HT20050306	59	Australia	I	I	Ι
H40	60	UK	S	S	S
D473	69	UK	Ι	I	I
CDC201078-USA700	72	USA	I	I	I
HT20040991	80	Algeria	S	R	S
SwedN8890/99	80	Sweden	Ι	Ι	Ι
BK1563	88	USA	Ι	S	S
HT20020635	93	Australia	S	I	S
HT2001634	93	Australia	Ι	S	S
HT20020635	93	USA	Ι	S	S
Cuba4005	94	Cuba	S	S	S
D302	97	UK	S	S	S
Not38	101	UK	S	S	S
D472	109	UK	S	S	S
H560	121	UK	S	S	S
D139	145	UK	Ι	Ι	Ι
D22	182	UK	Ι	I	I
Can6428-011	188	Canada	S	S	S
D470	207	UK	Ι	Ι	Ι
98/10618	217	UK	S	S	S
CDC12	225	USA	S	S	S
Germany131/98	228	Germany	S	S	S
CDC16	231	USA	S	S	S

99.3759V	235	Scotland	S	Ι	S
SwedenON408/99	246	Sweden	Ι	Ι	Ι
KD121618	250	Switzerland	S	Ι	S
KD12943	257	UK	Ι	Ι	Ι
Not271	264	UK	Ι	Ι	Ι
Not380	266	UK	Ι	S	S
Not98-53	280	UK	Ι	S	Ι
CAN6820-0616	289	Canada	Ι	Ι	Ι
Fin62305	296	Finland	S	Ι	S
Not266	301	UK	S	S	S
Btn2164	312	UK	S	S	S
Btn2299	322	UK	Ι	Ι	S
Btn2289	322	UK	S	S	S
515/09	398		S	S	S
Not161	517	UK	S	Ι	S
Not290	529	UK	S	S	S
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<sup>†</sup>Bacterial isolates were Susceptible – clear spot (S, light grey), Intermediate – turbid spot (I,

751 medium grey) or Resistant - no disturbance of bacterial lawn (R, dark grey) to phage

- 752 infection.

764 Table 2: Sensitivity screening of phage mixture against coagulase-negative staphylococci

# 765 isolates.

Species	Isolates	k	DRA88	Phage Cocktail
S. xylosus	ATCC 29971	Ι	Ι	Ι
S. sciuri subsp. sciuri	ATCC 29062	Ι	I	Ι
S. chromogenes	CCM 3387	Ι	Ι	Ι
S. hyicus	CCM 29368	S	Ι	S
S. arlettae	N910 254	Ι	Ι	Ι
S. vitulinus	ATCC 51145	Ι	Ι	Ι
S. simulans	N920 197	S	I	S
S. epidermidis	ATCC 14990	Ι	R	Ι

<sup>†</sup> Bacterial isolates were Susceptible – clear spot (S, light grey), Intermediate – turbid spot (I,

767 medium grey) or Resistant - no disturbance of bacterial lawn (R, dark grey) to phage

- 768 infection.



**Figure 1**: Curve One-step growth of phage DRA88 (left) and phage K (right) in RN4220 *S. aureus* at 37 °C. Shown are the PFU per infected cell in untreated cultures ( $\bullet$ ) and in chloroform-treated cultures ( $\circ$ ) at several time points over 60 minutes. The phage growth parameters are indicated in the figure and correspond to: E - eclipse period; L - latent period and B - burst size. Each data point is the mean of three independent experiments and error bars indicate the mean  $\pm$  standard deviation.

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**Figure 3:** Electron micrograph images of phage DRA88 negatively stained with 1% uranyl acetate, (A) showing the tail in a contracted position and (B) formation on phage particles aggregates. Scale bar (50 nm (A) and 100 nm (B) is indicated.



**Figure 4:** Comparative genomic analysis of phage DRA88 and phage K. Nucleotide sequences were compared using the Artemis Comparison Tool (ACT). Predicted ORFs are denoted by arrows, tRNAs are indicated (vertical blue dashed line) and genes encoding proteins with at least 69% amino acid identity between the two genomes are indicated by shaded regions.



**Figure 5:** Dynamic of bacteria with single and phage mixture in liquid cultures over 24 hours incubation at 37 °C. Absorbance readings at 590nm were taken in a microtitre plate reader. *S. aureus* isolates: A – 15981; B – MRSA 252; C – H325 growing with only SM buffer ( $\circ$ ), with single DRA88 ( $\checkmark$ ), with single phage K ( $\triangle$ ) and with the phage mixture in SM buffer ( $\bullet$ ) and also a negative control only SM buffer ( $\bullet$ ) are shown in the figure. Assays were performed four times and OD590 was expressed as the mean ± standard deviation.



Figure 6: Normalized biofilm biomass treated with single phage K, DRA88 and the phage mixture after 48 hours at MOI 10 (OD590 reading after CV staining). *S. aureus* isolates: 1 – 15981; 2 – MRSA 252; 3 – H325. Mean values for the three strains treated with: phage K = 0.63 (SD $\pm$ 0.10), 0.30 (SD $\pm$ 0.16), 0.27 (SD $\pm$ 0.06); DRA88 = 0.11 (SD $\pm$ 0.04), 0.18 (SD $\pm$ 0.08), 0.29 (SD $\pm$ 0.03) phage mixture = 0.06 (SD $\pm$ 0.02), 0.15 (SD $\pm$ 0.02), 0.23 (SD $\pm$ 0.02). Assays were performed three times and the mean  $\pm$  standard deviation is indicated. Statistically significance of biofilm reduction was assessed by performing a *Student's t-test. p-values* are indicated (\* : <0.05).

		15	981	MRS	A 252	H325		
	<b>Biofilm Treatment</b>	А	В	Α	В	Α	В	
	PBS	000						
	Phage K	000			$\bigcirc \bigcirc \bigcirc \bigcirc$	000	000	
	DRA88							
	Phage Combination							
	Figure 7: Vis	sualization of	f wells stained	d with 0.1%	of crystal vio	let after 48 h	ours of phage	
	treatment at N	4OI 10. Show	wn are the bi	ofilm wells t	reated with P	BS, phage K	, DRA88 and	
	phage mixture	e at 0h (A) ar	nd (B) 48 hou	ırs after. Expe	eriments were	performed in	triplicate.	
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