

The mef(A)/msr(D)-carrying streptococcal prophage **Φ**1207.3 encodes an SOS-like system, induced by UV-C light, responsible for increased survival and increased mutation rate

This is the peer reviewed version of the following article:				
Original:				
Fox, V., Santoro, F., Apicella, C., Diaz-Diaz, S., Rodriguez-Martínez, J.M., Iannelli, F., et al. (2023). The mef(A)/msr(D)-carrying streptococcal prophage Φ1207.3 encodes an SOS-like system, induced by UV-C light, responsible for increased survival and increased mutation rate. JOURNAL OF BACTERIOLOGY, 205(9), 1-13 [10.1128/jb.00191-23].				
Availability:				
This version is availablehttp://hdl.handle.net/11365/1246694 since 2023-10-04T08:37:18Z				
Published:				
DOI:10.1128/jb.00191-23				
Terms of use:				
Open Access				
The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. Works made available under a Creative Commons license can be used according to the terms and conditions of said license. For all terms of use and more information see the publisher's website.				

(Article begins on next page)

1	The <i>mef</i> (A)/ <i>msr</i> (D)-carrying streptococcal prophage Φ1207.3 encodes
2	an SOS-like system, induced by UV-C light, responsible for increased
3	survival and increased mutation rate
4	
5	Valeria Fox ¹ , Francesco Santoro ^{1,*} , Carmen Apicella ¹ , Sara Diaz-Diaz ² , Josè Manuel Rodriguez-
6	Martinez ³ , Francesco Iannelli ^{1,*} , Gianni Pozzi ¹
7	
8	¹ Laboratory of Molecular Microbiology and Biotechnology, Department of Medical
9	Biotechnologies, University of Siena, 53100 Siena, Italy
10	² Unidad Clínica de Enfermedades Infecciosas, Microbiología y Medicina Preventiva, Hospital
11	Universitario Virgen Macarena, Sevilla, Spain
12	³ Departamento de Microbiología, Facultad de Medicina, Universidad de Sevilla, Sevilla, Spain
13	
14	*Corresponding author.
15	Mailing address: LAMMB, Biotecnologie Mediche/Università di Siena, Policlinico Le Scotte, V
16	Lotto I Piano, Viale Bracci, 53100 Siena, Italy.
17	Phone: +39-0577-233156; +39-0577-233156. Fax: +39-0577-233334.
18	E-mail: francesco.iannelli@unisi.it; santorof@unisi.it
19 20	
20	
21	Keywords: SOS response, S. pneumoniae, S. pyogenes, prophage, Φ1207.3, mef(A)/msr(D),
22	mutation rate, bacterial survival, UV-C LED light, fluctuation test, DNA repair
23	
24	

25 Abstract

26 Bacterial SOS response is an inducible system of DNA repair and mutagenesis. Streptococci lack a canonical SOS response, but an SOS-like response was reported in some species. The mef(A)-27 28 msr(D)-carrying prophage Φ 1207.3 of Streptococcus pyogenes contains a region, spanning orf6 to 29 orf11, showing homology to characterized streptococcal SOS-like cassettes. Genome-wide 30 homology search showed the presence of the whole Φ 1207.3 SOS-like cassette in three S. pyogenes 31 prophages, while parts of it were found in other bacterial species. To investigate whether this cassette confers an SOS-mutagenesis phenotype, we constructed Streptococcus pneumoniae R6 32 33 isogenic derivative strains: (i) FR172, streptomycin resistant, (ii) FR173, carrying Φ 1207.3, and (iii) FR174, carrying a recombinant Φ 1207.3, where the SOS-like cassette was deleted. These strains 34 were used in survival and mutation rate assays using a UV-C LED instrument for which we 35 36 designed and 3D-printed a customized equipment, constituted of an instrument support and swappable-autoclavable mini-plates and lids. Upon exposure to UV fluences ranging from 0 to 6400 37 J/m^2 at four different wavelengths, 255, 265, 275 and 285 nm, we found that the presence of 38 Φ1207.3 SOS-like cassette increases bacterial survival up to 34-fold. Mutation rate was determined 39 by measuring rifampicin resistance acquisition upon exposure to UV fluence of 50 J/m^2 at the four 40 41 wavelengths by fluctuation test. The presence of Φ 1207.3 SOS-like cassette resulted in a significant 42 increase in the mutation rate (up to 18-fold) at every wavelength. In conclusion, we demonstrated that Φ 1207.3 carries a functional SOS-like cassette responsible for an increased survival and 43 44 increased mutation rate in S. pneumoniae.

45 **Importance**

Bacterial mutation rate is generally low, but stress conditions and DNA damage can induce stress response systems, which allow for improved survival and continuous replication at the cost of elevated mutagenesis. The SOS response is a DNA repair mechanism activated by some bacteria in response to stressful conditions, which leads to a temporary hypermutable phenotype and is usually absent in streptococcal genomes. Here, using a reproducible and controlled UV irradiation system, we demonstrated that the SOS-like gene cassette of prophage Φ 1207.3 is functional, responsible for a temporary hypermutable phenotype, and enhances bacterial survival to UV irradiation. Prophage Φ 1207.3 also carries erythromycin resistance genes and can lysogenize different pathogenic bacteria, constituting an example of a mobile genetic element which can confer multiple phenotypes to its host.

56 Introduction

Bacteria evolve through chromosomal rearrangements, acquisition of DNA by horizontal gene 57 58 transfer and mutation. Mutations arise by a process referred to as mutagenesis which is measured 59 determining the mutation frequency or the mutation rate. Mutation frequency represents the proportion of mutant cells in a specific culture, while the mutation rate is the probability of a 60 mutation occurring per cell division and thus is also an estimation of the probability of a mutation 61 62 occurring during the life of a bacterial cell (1-3). Mutation rates differ among bacterial species and among strains of the same species, and are influenced by the fidelity of DNA replication, DNA 63 64 damage and repair pathways, and exposure to chemical or physical agents. In Escherichia coli DNA replication is carried out by high fidelity polymerases which misinsert nucleotides at a rate of 10⁻⁴-65 10^{-6} per base pair (bp), but possess a 3'-exo proofreading activity that reduces mutation load by 66 67 about 100-fold (4). E. coli has also evolved the SOS response, which is inducible by DNA damage 68 and allows for improved survival and continuous replication at the cost of elevated mutagenesis (5, 6). The main proteins involved in SOS response are LexA, a homodimeric transcriptional repressor 69 of the SOS regulon, and RecA, which, upon DNA damage, binds to ssDNA, becomes activated, and 70 71 promotes the autocleavage of LexA and subsequent transcription of the SOS genes (7). The SOS 72 response induces the expression of genes involved in DNA-damage tolerance and repair, including the Translesion Synthesis (TLS) system (8). TLS is involved in the tolerance to DNA damage, and 73 can be both error-free or error-prone, when the polymerase V is recruited. Polymerase V is encoded 74 by the *umuDC* operon, where *umuC* encodes a translession DNA polymerase, of the Y family, while 75

umuD encodes a regulatory protein. The low fidelity polymerase V has a high rate of nucleotide misincorporation $(10^{-3}-10^{-4} \text{ per bp})$ (9) and also lacks proofreading activity (10).

Streptococci lack a canonical SOS response (11), but SOS-like gene cassettes are present in the 78 genomes, either on the chromosome or on Mobile Genetic Elements, of Streptococcus uberis, 79 80 Streptococcus agalactiae, Streptococcus pyogenes, Streptococcus sanguinis, Streptococcus mitis, and Streptococcus thermophilus (12). In S. uberis, the SOS-like cassette contains 4 genes arranged 81 82 in an operon including (i) hdiR, coding for a protein homologous to the HdiR regulator of Lactococcus lactis, induced by heat shock and DNA-damage (13), (ii) umuC encoding an UmuC-83 like protein, (iii) 2 genes of unknown function. The HdiR regulator of S. uberis undergoes 84 spontaneous self-cleavage at alkaline pH, similar to the RecA-mediated self-cleavage of LexA-like 85 86 proteins. HdiR specifically binds to an inverted repeat sequence downstream the *hdiR* putative promoter, suggesting a role as transcriptional repressor of the SOS-like cassette. The pneumococcal 87 Integrative and Conjugative Element (ICE) Tn5253 confers UV resistance to the cell, inducing a 88 89 mutagenic SOS response (14). Tn5253 contains an SOS-like cassette similar to that described in S. 90 uberis, comprising: (i) orf69, which codes for an LexA-like repressor, (ii) orf70, which codes for an 91 UmuC-like protein, (iii) orf71 and orf72 of unknown function (15–17). In S. pyogenes, a gene cassette homologous to the already known streptococcal SOS-like cassettes is associated with the 92 mef(A)-msr(D)-carrying prophages Φ 1207.3 (18–21), Φ 10394.4, Φ m46.1 and its variant 93 VP 00501.1 (22–25). Φ 1207.3 is a functional bacteriophage capable to lysogenize different strains 94 95 of the human pathogens S. pyogenes and S. pneumoniae, which colonize the upper respiratory tract (26). Since both of these species can cause invasive infections, the transmission of antibiotic 96 97 resistance genes by lysogenic transfer of Φ 1207.3 assumes clinical relevance. In this work, we transferred Φ 1207.3 prophage from the original S. pyogenes host to the S. pneumoniae standard 98 laboratory strain R6 to investigate if the SOS-like cassette carried by the Φ 1207.3 prophage is able 99 to activate an SOS-like response, which could increase the acquisition of mutations associated to 100 101 antimicrobial resistance.

102 Materials and methods

103 *Bioinformatic analysis*

104 Homology searches were conducted in public nucleotide and protein databases, namely Microbial 105 Genomes Database (https://www.ncbi.nlm.nih.gov/genome/microbes/), Conserved Domain 106 Database (CDD, https://www.ncbi.nlm.nih.gov/cdd/) and Pfam protein family database (https://pfam.xfam.org/), using default parameters, and only alignment with E-values <0.001 were 107 considered. Protein sequence analysis was conducted with the tools available at Softberry website 108 (http://www.softberry.com/berry.phtml) the 109 and with software Phyre2 (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index) (27). Nucleotide sequence of the orf6-110 111 orf11 region was used as a probe for Microbial Genomes Database searches, while ORF predicted 112 amino acid sequences were used for protein databases searches.

113 Bacterial strains and growth

Streptococcal strains used in this work and their relevant properties are reported in Table 1. Strains were grown in Tryptic Soy Broth (TSB, BD) at 37°C. Starter cultures were taken at an optical density at 590 nm (OD_{590}) ranging from 0.2 to 0.3 and were frozen in 10% glycerol at -70°C. Solid media were obtained by supplementing TSB with 1.5% agar (BD Difco) and 3% defibrinated horse blood (Liofilchem). When required, both liquid and solid media were supplemented with antibiotics at the following concentrations: 500 µg/ml kanamycin, 500 µg/ml streptomycin, 0.5 µg/ml erythromycin, 5 µg/ml chloramphenicol and 20 µg/ml rifampicin.

121 Lysogenization assays

122 Transfer of bacteriophage Φ 1207.3 was obtained through a plate mating experiment, as previously 123 described (28). Briefly, donor cells, carrying the phage, and recipient cells were grown separately in 124 TSB in the presence of the appropriate antibiotics. Upon reaching the end of the exponential phase, 125 cells were mixed at a donor-recipient 1:10 ratio, centrifuged, and the pellet was plated on TSA 126 plates supplemented with 5% horse blood. Plates were incubated at 37°C in the presence of 5% CO₂ 127 for 4h, cells were then recovered with a cotton swab and resuspended in TSB/10% glycerol. To select for lysogens, cell suspension was plated following a multilayer plating procedure in presence of erythromycin and streptomycin which select for lysogens and counterselect the Φ 1207.3 donor cells. To confirm the lysogens phenotype, a genetic analysis was performed as reported (16,28), while PCR and direct Sanger sequencing of PCR template were used for genotype analysis according to the protocol already described (32).

133 PCR Gene SOEing mutagenesis

To generate a knock-out mutant of orf6-orf11 region, PCR Gene Splicing by Overlap Extension 134 135 (SOEing) was used to produce a mutagenic construct containing the kanamycin resistance cassette 136 (876 bp) flanked by the DNA fragments located upstream of orf6 (555 bp) and downstream of orf11 137 (829 bp) as previously described (29). The primer pair IF149/IF210 was used to amplify the 876-bp 138 ami/aphIII kanamycin resistance cassette (30), while IF242/IF239 and IF240/IF241 amplified the 139 flanking fragments. Assembly of the final construct (2,260 bp) was obtained with the IF242/IF241 primer pair. PCR was carried out as previously described (31). The PCR products were separated in 140 141 a 0.8% agarose gel at 200 V for 30 minutes, stained for 15 minutes in a 0.1% Ethidium Bromide 142 solution and visualized with UV light. PCR products after gel purification (NucleoSpin Gel and 143 PCR Clean-up kit, Macherey-Nagel) were quantified using the Qubit 2.0 Fluorometer (Thermo 144 Fisher Scientific). Oligonucleotide primers are listed in Table S1. The PCR mutagenic construct 145 was directly used as donor DNA in transformation experiments. Since integration of Φ 1207.3 into 146 S. pneumoniae chromosome leads the disruption of the celB competence gene with a consequent 147 impairment of genetic transformation, we constructed transformable S. pneumoniae Rx1 derivative 148 strains carrying Φ 1207.3 integrated elsewhere in pneumococcal chromosome to be used as 149 tranformation recipients (Table 1) (28). Pneumococcal competent cells were prepared as previously 150 described (29). Competent cells were thawed, supplemented with 25 ng/ml Competence Stimulating 151 Peptide (CSP) and 1 µg/ml of the purified mutagenic construct, and the transformation mixture was 152 incubated at 37°C for 45 minutes. The resulting recombinant strains were selected for acquisition of 153 kanamycin resistance by multilayer plating (28). The deletion of the orf6-11 region (nucleotides

154 6,189 to 9,297, GenBank accession no. <u>AY657002</u>) was confirmed by PCR and direct sequencing

155 of the PCR products (32).

156 UV fluence measurement

157 UV-C light experiments were carried out using a PearlLab Micro UV-C LED instrument 158 (AquiSense Technologies, Erlanger, Kentucky, USA). The instrument is a compact device equipped 159 with 4 interchangeable USB UV-C LEDs emitting at different wavelengths (255, 265, 275 and 285 160 nm) and a dimmer switch that allows irradiation with 10 different intensities. For our purposes, the lower portion of the instrument was un-screwed, and the lamp was used as a general light source. A 161 162 customized equipment, constituted of an instrument support and swappable and autoclavable miniplates and lids, was 3D-designed and printed in Nylon-12 (Figure 163 S1, https://doi.org/10.5281/zenodo.8208699). The irradiance was measured 3 times with a ILT1400 164 165 radiometer and SED240/W detector (International Light Technologies, Peabody, MA). Mean and 166 standard deviation for each intensity of the different wavelengths used are reported in Table S2. The UV fluence (J/m^2) is calculated as "irradiance (W/m^2) x exposure time (sec)", thus the same value 167 of the UV fluence can be obtained varying the values of the 2 parameters. Since the PearlLab Micro 168 169 UV-C LED instrument emits light at 10 different intensities, allowing for 10 different values of irradiance for each of the 4 wavelengths, different combinations of intensity and time were possible. 170 171 The choice of the optimal parameters to be used in UV-C light survival and fluctuation assays was based on a preliminary survival experiment where an UV fluence of 50 J/m^2 was obtained varying 172 the irradiance and the time. Since the combination of high intensities and low exposure times, or 173 174 low intensities and high exposure times, resulted in a wide variability of survival among the 175 technical replicates, we chose, when possible, to maintain similar exposure times across all wavelengths and to use intermediate intensities for the 50 J/m^2 UV fluence for fluctuation assays 176 (Table S3). 177

178 Survival assay

179 Bacterial survival upon UV-C light irradiation at different wavelengths (255 to 285 nm) was assessed by plating the cells after exposure to different UV fluences from 0 to 6400 J/m^2 (0, 25, 50, 180 100, 200, 400, 800, 1600, 3200, 6400) and by calculating the cell viability for each strain. Cells 181 were grown at 37°C in 10 ml of TSB until reaching an OD₅₉₀ of 0.6 (about 1 x 10^8 CFU/ml), when 182 183 they were centrifuged at 5,000 x g for 15 minutes, washed with sterile phosphate buffered saline (PBS, Sigma Aldrich) and resuspended in 10 ml of PBS. UV-C survival experiments were carried 184 185 out by placing 1 ml of cells in the sterile Nylon-12 mini-plate and exposing them to different fluences, under continuous stirring. UV-C irradiated and untreated control bacterial cells were 186 plated by multilayer plating. Survival was calculated as the ratio between UV-C irradiated and not 187 188 irradiated control cells.

189 Luria-Delbrück fluctuation test

190 Mutation rate was determined by fluctuation analysis in absence or presence of UV-C light at 191 different wavelengths ranging from 255 to 285 nm. Strains were grown in TSB at 37°C until an 192 OD₅₉₀ of 0.6 and the number of total and rifampicin-resistant CFUs was determined prior to freezing at -70°C in 1 ml-aliquots containing 10% glycerol. Rifampicin was used at a final 193 194 concentration of 20 μ g/ml in the third layer of TSA, when appropriate (28). For mutation rate determination, a set of 10 independent cultures was obtained by inoculating 5 ml of TSB with 1 x 195 10⁵ CFU of the same 1 ml frozen aliquot. For mutation rate determination following UV-C light 196 treatment, the 1 ml frozen aliquot was irradiated at an UV fluence of 50 J/m² before inoculum. 197 Then, to prepare 10 independent cultures, an inoculum corresponding to 1×10^5 CFU was calculated 198 199 by inferring bacterial viability after irradiation from the data obtained with the UV-C light survival 200 assay. Cultures were grown at 37°C until they reached the mid-log phase, corresponding to about 12 generations, the number of total and rifampicin-resistant CFUs was determined by plating 0.1 ml of 201 culture and incubating the plates at 37°C for 48 h. For each strain and condition, 4 replicates of the 202 203 fluctuation test were carried out and the rifampicin-resistant CFUs numbers of the 40 cultures were 204 pooled into a single data set (33). The mutation rate determination and the statistical analysis were

carried out using the R package rSalvador (34). The expected number of mutations per culture (m) was first obtained with the use of the function *newton.LD.plating*, which computes the maximum likelihood estimates (MLE) of m, adjusting to the plating efficiency, i.e., when only a portion of the culture is plated. Once the estimate of m was obtained, the mutation rate (μ) was determined dividing m by the number of cell divisions that had taken place. Since the final number of cells in a culture, N_t, arises from N_t-1 divisions, the mutation rate is:

$$\mu = \frac{m}{Nt - 1} \approx \frac{m}{Nt}$$

where N_t -1 is equal to the number of generations and can be approximated to N_t (3, 34). 211 Differences in mutation rates were compared using the rSalvador likelihood ratio test (LRT) (34-212 213 36), which takes into consideration both the plating efficiency and the final cell number N_t , which 214 differs among different strains or experimental conditions, by applying the *compare.LD.plating* 215 function pairwise among all conditions of each strain and among the different strains. The *p.adjust* 216 R function was then used to correct the overall false positive rate of the obtained p values for 217 multiple comparisons. Differences in the mutation rate were considered significant only when the 218 adjusted p-values were less than 0.05 (33). The comparison was also observed graphically by 219 checking the overlapping of the 84% confidence intervals (C.I.s) (36).

220 **Results**

221 Φ 1207.3 carries an SOS-like cassette

The Φ 1207.3 DNA region located between the *mef*(A)-*msr*(D) erythromycin resistance cassette (20) and the type II restriction-modification cassette (37), contains 6 open reading frames (namely *orf6* to *orf11*) likely arranged in an operon (Figure 1). Homology search revealed that *orf11* predicted gene product is homologous to the HdiR LexA-like transcriptional repressor of *Lactococcus lactis*, *orf8* is homologous to the *Streptococcus pneumoniae* Tn5253 *umuC/orf70*, while *orf7* gene product is homologous to the *Bacillus subtilis* YolD-like protein, which is predicted to be a functional equivalent of UmuD (38) (Table 2). The deduced amino acid sequences of *orf6* to *orf11* share homology (40-73%) with the predicted gene products of the *S. uberis* and of the *S. pneumoniae* Tn5253 SOS-like cassettes, respectively (12, 14), with the exception of *orf9* and *orf10*, which appear to be unique for Φ 1207.3. For these reasons, the *orf6* to *orf11* region of Φ 1207.3 was referred to as an SOS-like cassette.

233 Presence of the Φ 1207.3 SOS-like cassette in bacterial genomes

A genome-wide homology search of the Φ 1207.3 SOS-like cassette in the 67,343 complete 234 235 bacterial genomes available in the GenBank Microbial Genomes Database (accessed in March 236 2022) revealed the presence of the cassette, or parts of it, in 104 complete bacterial genomes (Figure 237 S2, Table S4). In 3 S. pyogenes genomes, a cassette 100% homologous to that of Φ 1207.3 was 238 found associated to mef(A)-msr(D)-carrying prophages, including Φ 10394.4. A copy of the SOS-239 like cassette, containing a short deletion at the 5' end of orf11, was found in 12 additional 240 streptococcal genomes, including Streptococcus gallolyticus, Streptococcus dysgalactiae, and 241 Streptococcus suis. In the remaining 89 genomes, fragments of the cassette were found, including a 242 fragment spanning orf6 to orf9 in 6 S. suis genomes and a fragment spanning orf6 to the partial 3' 243 end of orf8 in 43 genomes, including 16 pneumococcal genomes. The fragment belongs to the 244 mef(A)-msr(D)-carrying element Tn1207.1 (39, 40) in the S. pneumoniae INV200 genome and to 245 the mef(E)-msr(D)-carrying mega element (41, 42) in the remaining 15 genomes.

246 Construction of isogenic S. pneumoniae recombinant strains

To investigate if the presence of this cassette confers an SOS-mutagenesis phenotype we constructed three isogenic derivative strains of the *S. pneumoniae* standard laboratory strain R6: (i) FR172, streptomycin resistant derivative of R6, (ii) FR173, derivative of FR172 carrying Φ 1207.3, and (iii) FR174, derivative of FR172 carrying a recombinant Φ 1207.3, where a 3109-bp DNA fragment, corresponding to the *orf6* to *orf11* region, was deleted by substitution with a kanamycin resistance cassette (Table 1). Isogenic strains were constructed in R6 since it harbours a functional Mismatch Repair System (*hex*+) which results in a low basal mutation rate (43–46).

254 The presence of the Φ 1207.3 SOS-like cassette increases bacterial survival upon UV-C light 255 irradiation

To assess whether the SOS-like cassette of Φ 1207.3 is involved in increased survival upon UV-C 256 257 light treatment, the bacterial viability of the three isogenic strains was measured upon exposure to UV fluences ranging from 0 to 6400 J/m² at four different wavelengths, 255, 265, 275 and 285 nm 258 (Figure 2). No significant difference in bacterial survival was observed between strain FR172, not 259 260 carrying $\Phi_{1207.3}$, and FR174, carrying the recombinant $\Phi_{1207.3\Delta orf6-orf11}$ devoid of the SOSlike cassette. The presence of the Φ 1207.3 in FR173 increases bacterial survival upon UV-C 261 irradiation compared to FR172 and FR174, regardless of UV-C light fluences and wavelengths. 262 Noticeably, at a 6400 J/m^2 fluence we did not observe any increase in survival due to the SOS-like 263 cassette presence. It is likely that, at high UV fluences, the accumulation of DNA damages with the 264 265 associated high rate of mortality hinders the activation of the SOS-like response. FR173, carrying 266 Φ 1207.3, resulted in an increased bacterial survival compared to FR172, not carrying Φ 1207.3, up 267 to (i) 8.46-fold at 255 nm, (ii) 29.11-fold at 265 nm, (iii) 16.10-fold at 275 nm, (iv) 7.26-fold at 285 268 nm. When compared to FR174, carrying the recombinant Φ 1207.3 Δ SOS-like cassette, FR173 269 resulted in an increased bacterial survival up to (i) 10.93-fold at 255 nm, (ii) 34.21-fold at 265 nm, 270 (iii) 10.69-fold at 275 nm, (iv) 8.23-fold at 285 nm (Table S5).

271 The Φ 1207.3 SOS-like cassette increases mutation rate upon UV-C light irradiation

272 Activation of the Φ 1207.3 SOS-like response upon UV-C exposure was assessed determining the 273 mutation rate by measuring the acquisition of rifampicin resistance, associated to point mutations in 274 the *rpoB* gene coding for the β subunit of RNA polymerase (47, 48). A significant increase of 275 mutation rate was observed at every wavelength tested in FR173, carrying Φ 1207.3, whereas no significant increase was observed in the FR172, not carrying Φ 1207.3, and FR174, carrying the 276 277 recombinant Φ 1207.3 Δ SOS-like cassette (Figure 3). In particular, in FR173 the mutation rate increase of irradiated cells compared to not irradiated cells was: (i) 6.64-fold at 255 nm (p-value = 278 279 0), (ii) 9.50-fold at 265 nm (p-value = 0), (iii) 8.63-fold at 275 nm (p-value = 0), and (iv) 18.38-fold

at 285 nm (p-value = 0), whereas in FR172 and FR174 no significant increase in mutation rate was observed (Table S6). In absence of UV-C light irradiation, the three isogenic strains showed similar mutation rate values, regardless of the presence of the Φ 1207.3 prophage or of the SOS-like cassette.

284 Discussion

285 Mutation rate is generally low in bacteria, but hypermutable phenotype can arise as a consequence 286 of the loss of DNA proofreading during replication Sundin and Weigand 2007; Gressel and Levy 287 2010; Jolivet-Gougeon et al. 2011). Stress conditions and DNA damage can also induce stress 288 response systems, which allow for improved survival and continuous replication at the cost of 289 elevated mutagenesis (6, 53). The SOS response is a DNA repair mechanism activated by some 290 bacteria in response to stressful conditions, which leads to a temporary hypermutable phenotype (5, 291 6). In the present work, we demonstrated that the S. pyogenes prophage Φ 1207.3 carries a 292 functional gene cassette responsible for the activation of an SOS-like response in S. pneumoniae in 293 response to UV-C light irradiation. We studied this cassette after its introduction in the S. 294 pneumoniae R6 strain, which contains a Mismatch Repair System (hex) conferring a low mutation 295 rate (43). We observed that the activation of this SOS-like cassette, upon UV-C exposure, makes the R6 strain temporarily hypermutable, resulting in a higher ability of surviving and evolving. 296 297 Hypermutable strains have been found in pathogenic bacteria (49) and represent a huge problem in 298 clinical settings (54-57). Clinical strains are frequently exposed to variable and changing stressful 299 conditions, such as antibiotic treatments and host defenses, which can lead to evolution of resistance 300 and virulence phenotypes, with consequent therapeutic failures (33, 51, 58-62). Hypermutable 301 strains were also characterized in S. pyogenes, where the DNA Mismatch Repair (MMR) gene mutL 302 can be interrupted by the insertion of a genomic island. This genomic island is capable of growth-303 phase-dependent excision and functions as a switch for the hypermutator phenotype (63). Prophage Φ 1207.3, originally found in a clinical isolate of S. pyogenes, carries the erythromycin resistance 304 305 genes mef(A)-msr(D) (18-21) and is able to transfer among S. pyogenes strains with different

306 genetic background (unpublished data) and other streptococcal species such as S. gordonii and S. 307 pneumoniae (18). Furthermore, genomic analysis showed that DNA segments with homology to the 308 Φ 1207.3 SOS cassette are present in different streptococci and are carried on mobile genetic 309 elements, which have the potential to spread the cassette. A recent work showed that the 310 Streptococcal Mobilizable family of phages are widely distributed among different streptococcal 311 species, carry antibiotic resistance and virulence genes associated with the SOS cassette and are 312 mobilizable by a mechanism resembling conjugative transfer, possibly facilitated by the 313 concomitant presence of ICEs ((64)). Here we demonstrated that Φ 1207.3 also carries a functional 314 SOS-like cassette conferring a transient hypermutable phenotype in response to stresses, which, 315 under the selective pressure of antibacterial therapy, could result in the accumulation of mutations 316 conferring additional antibiotic resistance phenotypes. The presence of the Φ 1207.3 SOS-like 317 cassette in S. pneumoniae confers a higher survival rate compared to the isogenic strains without 318 prophage or carrying $\Phi 1207.3 \Delta or f6$ -or f11 devoid of the SOS-like cassette. On the one hand, the 319 cells harbouring Φ 1207.3 are capable of surviving when exposed to stresses, on the other hand 320 Φ 1207.3 ensures its survival within the lysogenic cell population. Further experiments will be 321 needed to assess the effect of UV light exposure on prophage gene expression and production of phage particles. This would clarify whether phage replication or lysogenic conversion is favoured. 322 323 The presence of erythromycin resistance genes, the ability to transfer among different bacterial 324 species and the ability to confer a transient hypermutable phenotype in response to stresses 325 highlight the clinical relevance of Φ 1207.3.

326 Acknowledgments

This work was supported in part from the Italian Ministry of Education, University and Research (MIUR-Italy) under grant number 20177J5Y3P (call "Progetti di Ricerca di Rilevante Interesse Nazionale – Bando 2017"), in part from the Italian Ministry of University and Research (MUR-Italy) under grant number 202089LLEH ("Transition from asymptomatic colonization to disease by human respiratory-tract bacteria as a target for vaccines and antimicrobial therapy: The CoDiCo

332	(colonisation to disease concepts)" project call "Progetti di Ricerca di Rilevante Interesse Nazionale
333	- Bando 2020"), and in part by EU funding within the MUR PNRR Extended Partnership Initiative
334	on Emerging Infectious Diseases (Project no. PE00000007, INF-ACT, "One Health Basic and
335	Translational Research Actions addressing Unmet Needs on Emerging Infectious Diseases").

336 FIGURE LEGENDS

337 Figure 1. S. pyogenes Φ 1207.3 SOS-like cassette. The prophage Φ 1207.3 contains an SOS-like 338 cassette located between the mef(A)-msr(D) gene pair conferring erythromycin resistance and the 339 type II restriction-modification cassette constituted by orf12, orf13 and orf14. The cassette contains 340 6 open reading frames (namely orf6 to orf11) likely to be arranged in an operon. The orf11 gene 341 product is homologous to L. lactis HdiR LexA-like transcriptional repressor, Orf8 to the S. 342 pneumoniae Tn5253 UmuC/Orf70, Orf7 to the B. subtilis YolD-like protein, a putative functional 343 homolog of UmuD. The putative promotor sequence of the SOS-like operon is indicated, ORFs and 344 their direction of transcription are reported as arrows (in red *mef*(A)-*msr*(D), in green the SOS-like 345 cassette, in blue the restriction-modification cassette).

346 Figure 2. Bacterial survival upon UV-C light irradiation. Effect of the Φ 1207.3 SOS-like 347 cassette on the S. pneumoniae bacterial survival upon UV-C light irradiation was investigated in absence of Φ 1207.3 prophage (strain FR172, black circle), in presence of Φ 1207.3 (FR173, green 348 349 square) and in the presence of the recombinant $\Phi 1207.3 \Delta or f6$ -or f11, lacking the SOS-like cassette 350 (FR174, red triangle). Irradiation was carried out with UV-C light at wavelengths: A) 255 nm; B) 351 265 nm; C) 275 nm; D) 285 nm. No significant difference was observed between strain FR172, not 352 carrying Φ 1207.3, and FR174, carrying the recombinant Φ 1207.3 Δ orf6-orf11 devoid of the SOS-353 like cassette. The presence of the Φ 1207.3 in FR173 resulted in an increased bacterial survival 354 compared to FR172, not carrying Φ 1207.3, up to: i) 8.46-fold at 255 nm, ii) 29.11-fold at 265 nm, 355 iii) 16.10-fold at 275 nm, iv) 7.26-fold at 285 nm, while compared to FR174, carrying the 356 recombinant Φ 1207.3 Δ SOS-like cassette, up to i) 10.93-fold at 255 nm, ii) 34.21-fold at 265 nm, 357 iii) 10.69-fold at 275 nm, iv) 8.23-fold at 285 nm. Results are reported as means of at least three 358 independent experiments with standard deviations (error bars). In each experiment, 100% survival 359 was based on the cell viability measured in plates not receiving UV-C light irradiation (0 J/m^2) .

360 Figure 3. Mutation rates upon UV-C light irradiation.

361 Effect of the Φ 1207.3 SOS-like cassette on the mutation rate in S. pneumoniae upon UV-C light 362 irradiation was investigated in absence of Φ 1207.3 prophage, in presence of Φ 1207.3 and in the presence of the recombinant \$\Psi_1207.3 \Delta SOS-like cassette. Irradiation was carried out with UV-C 363 light at wavelengths 255 nm, 265 nm, 275 nm, and 285 nm at a fluence of 50 J/m². Strain FR173, 364 365 carrying Φ 1207.3, showed a significant increase in the mutation rate (up to 18.38-fold at 285 nm) 366 upon UV-C light treatment at every wavelength, whereas no significant increase was observed for 367 strain FR172, not carrying Φ 1207.3, and strain FR174, carrying the recombinant Φ 1207.3 Δ SOS-368 like cassette. The mutation rate was assessed by measuring the acquisition of rifampicin resistance. Results are reported as means of four independent experiments with 84% confidence intervals. *** 369

370

adjusted *p*-value < 0.001.

|--|

372	1.	Pope CF, O'Sullivan DM, McHugh TD, Gillespie SH. 2008. A Practical Guide to Measuring Mutation Pates in Antibiotic Pasistence. AAC 52:1200–1214
373		Rates in Antibiotic Resistance. AAC 52:1209–1214.
374	2.	Rosche WA, Foster PL. 2000. Determining Mutation Rates in Bacterial Populations. Methods 20:4–17.
375	3.	Foster PL. 2006. Methods for Determining Spontaneous Mutation Rates, p. 195–213. In Methods in
376		Enzymology. Elsevier.
377	4.	Goodman MF. 2016. Better living with hyper-mutation: Hypermutation. Environ Mol Mutagen
378		57:421–434.
379	5.	Radman M. 1975. SOS Repair Hypothesis: Phenomenology of an Inducible DNA Repair Which is
380		Accompanied by Mutagenesis, p. 355-367. In Hanawalt, PC, Setlow, RB (eds.), Molecular
381		Mechanisms for Repair of DNA. Springer US, Boston, MA.
382	6.	Maslowska KH, Makiela-Dzbenska K, Fijalkowska IJ. 2019. The SOS system: A complex and tightly
383		regulated response to DNA damage. Environ Mol Mutagen 60:368-384.
384	7.	Baharoglu Z, Mazel D. 2014. SOS, the formidable strategy of bacteria against aggressions. FEMS
385		Microbiol Rev 38:1126–1145.
386	8.	Fijalkowska IJ, Schaaper RM, Jonczyk P. 2012. DNA replication fidelity in Escherichia coli: a multi-
387		DNA polymerase affair. FEMS Microbiology Reviews 36:1105–1121.
388	9.	Tang M, Pham P, Shen X, Taylor J-S, O'Donnell M, Woodgate R, Goodman MF. 2000. Roles of E.
389		coli DNA polymerases IV and V in lesion-targeted and untargeted SOS mutagenesis. Nature
390		404:1014–1018.
391	10.	Friedberg EC, Walker GC, Siede W, Wood RD, Schultz RA, Ellenberger T. 2006. DNA Repair and
392		Mutagenesis. ASM press.

393	11.	Gasc AM, Sicard N, Claverys JP, Sicard AM. 1980. Lack of SOS repair in Streptococcus pneumoniae.
394		Mutation Research/Fundamental and Molecular Mechanisms of Mutagenesis 70:157–165.
395	12.	Varhimo E, Savijoki K, Jalava J, Kuipers OP, Varmanen P. 2007. Identification of a Novel
396		Streptococcal Gene Cassette Mediating SOS Mutagenesis in Streptococcus uberis. JB 189:5210-5222.
397	13.	Savijoki K, Ingmer H, Frees D, Vogensen FK, Palva A, Varmanen P. 2003. Heat and DNA damage
398		induction of the LexA-like regulator HdiR from <i>Lactococcus lactis</i> is mediated by RecA and ClpP:
399		RecA and CIpP modulate the expression of HdiR. Molecular Microbiology 50:609–621.
400	14.	Munoz-Najar U, Vijayakumar MN. 1999. An Operon That Confers UV Resistance by Evoking the
401		SOS Mutagenic Response in Streptococcal Conjugative Transposon Tn5252. J Bacteriol 181:2782-
402		2788.
403	15.	Iannelli F, Santoro F, Oggioni MR, Pozzi G. 2014. Nucleotide sequence analysis of integrative
404		conjugative element Tn5253 of Streptococcus pneumoniae. Antimicrob Agents Chemother, 58:1235-
405		1239.
406	16.	Santoro F, Romeo A, Pozzi G, Iannelli F. 2018. Excision and Circularization of Integrative
407		Conjugative Element Tn5253 of Streptococcus pneumoniae. Front Microbiol 9:1779.
408	17.	Santoro F, Fox V, Romeo A, Lazzeri E, Pozzi G, Iannelli F. 2021. Chromosomal integration of Tn5253
409		occurs downstream of a conserved 11-bp sequence of the rbgA gene in Streptococcus pneumoniae and
410		in all the other known hosts of this integrative conjugative element (ICE). Mobile DNA 12:25.
411	18.	Santagati M, Iannelli F, Cascone C, Campanile F, Oggioni MR, Stefani S, Pozzi G. 2003. The Novel
412		Conjugative Transposon Tn1207.3 Carries the Macrolide Efflux Gene mef(A) in Streptococcus
413		pyogenes. Microbial Drug Resistance 9:243–247.
414	19.	Iannelli F, Santagati M, Santoro F, Oggioni MR, Stefani S, Pozzi G. 2014. Nucleotide sequence of
415		conjugative prophage Φ 1207.3 (formerly Tn <i>1207.3</i>) carrying the <i>mef</i> (A)/ <i>msr</i> (D) genes for efflux
-		

416 resistance to macrolides in Streptococcus pyogenes. Front Microbiol 5:687-687.

417	20.	Iannelli F, Santoro F, Santagati M, Docquier J-D, Lazzeri E, Pastore G, Cassone M, Oggioni MR,
418		Rossolini GM, Stefani S, Pozzi G. 2018. Type M Resistance to Macrolides Is Due to a Two-Gene
419		Efflux Transport System of the ATP-Binding Cassette (ABC) Superfamily. Front Microbiol 9:1670.
420	21.	Fox V, Santoro F, Pozzi G, Iannelli F. 2021. Predicted transmembrane proteins with homology to
421		Mef(A) are not responsible for complementing <i>mef</i> (A) deletion in the <i>mef</i> (A)- <i>msr</i> (D) macrolide efflux
422		system in Streptococcus pneumoniae. BMC Res Notes 14:432.
423	22.	Banks DJ, Porcella SF, Barbian KD, Martin JM, Musser JM. 2003. Structure and Distribution of an
424		Unusual Chimeric Genetic Element Encoding Macrolide Resistance in Phylogenetically Diverse
425		Clones of Group A Streptococcus. J Infect Dis 188:1898–1908.
426	23.	Brenciani A, Bacciaglia A, Vignaroli C, Pugnaloni A, Varaldo PE, Giovanetti E. 2010. Φm46.1, the
427		Main Streptococcus pyogenes Element Carrying mef (A) and tet (O) Genes. Antimicrob Agents
428		Chemother 54:221–229.
429	24.	Di Luca MC, D'Ercole S, Petrelli D, Prenna M, Ripa S, Vitali LA. 2010. Lysogenic Transfer of mef(A)
430		and tet(O) Genes Carried by 4m46.1 among Group A Streptococci. Antimicrob Agents Chemother
431		54:4464-4466.
432	25.	Vitali LA, Di Luca MC, Prenna M, Petrelli D. 2016. Correlation between genetic features of the
433		mef(A)- msr(D) locus and erythromycin resistance in Streptococcus pyogenes. Diagnostic
434		Microbiology and Infectious Disease 84:57–62.
435	26.	Santoro F, Pastore G, Fox V, Petit M-A, Iannelli F, Pozzi G. 2023. Streptococcus pyogenes Φ1207.3 Is
436		a Temperate Bacteriophage Carrying the Macrolide Resistance Gene Pair <i>mef</i> (A)- <i>msr</i> (D) and Capable
437		of Lysogenizing Different Streptococci. Microbiol Spectr e04211-22.
438	27.	Kelley LA, Mezulis S, Yates CM, Wass MN, Sternberg MJE. 2015. The Phyre2 web portal for protein
439		modeling, prediction and analysis. Nat Protoc 10:845–858.

- 440 28. Iannelli F, Santoro F, Fox V, Pozzi G. 2021. A Mating Procedure for Genetic Transfer of Integrative
 441 and Conjugative Elements (ICEs) of Streptococci and Enterococci. MPs 4:59.
- 442 29. Iannelli F, Pozzi G. 2004. Method for Introducing Specific and Unmarked Mutations Into the
 443 Chromosome of *Streptococcus pneumoniae*. MB 26:81–86.
- 444 30. Pearce BJ, Iannelli F, Pozzi G. 2002. Construction of new unencapsulated (rough) strains of
- 445 *Streptococcus pneumoniae*. Research in Microbiology 5.
- 446 31. Santoro F, Oggioni MR, Pozzi G, Iannelli F. 2010. Nucleotide sequence and functional analysis of the
- 447 *tet*(M)-carrying conjugative transposon Tn5251 of *Streptococcus pneumoniae*. FEMS Microbiol Lett.
- 448 32. Iannelli F, Giunti L, Pozzi G. 1998. Direct sequencing of long polymerase chain reaction fragments.
 449 Mol Biotechnol 10:183–185.
- 33. Silayeva O, Engelstädter J, Barnes AC. 2020. Evolutionary epidemiology of *Streptococcus iniae*:
 Linking mutation rate dynamics with adaptation to novel immunological landscapes. Infect Genet Evol
 85:104435.
- 453 34. Zheng Q. 2017. rSalvador: An R Package for the Fluctuation Experiment. G3 (Bethesda) 7:3849–3856.
- 454 35. Zheng Q. 2015. Methods for comparing mutation rates using fluctuation assay data. Mutation

455 Research/Fundamental and Molecular Mechanisms of Mutagenesis 777:20–22.

456 36. Zheng Q. 2016. Comparing mutation rates under the Luria–Delbrück protocol. Genetica 144:351–359.

- 457 37. Euler CW, Ryan PA, Martin JM, Fischetti VA. 2007. M.SpyI, a DNA methyltransferase encoded on a
 458 *mef*A chimeric element, modifies the genome of *Streptococcus pyogenes*. J Bacteriol, 2006/11/03 ed.
 459 189:1044–1054.
- 460 38. Permina EA, Mironov AA, Gelfand MS. 2002. Damage-repair error-prone polymerases of eubacteria:
 461 association with mobile genome elements. Gene 293:133–140.

462	39.	Santagati M, Iannelli F, Oggioni MR, Stefani S, Pozzi G. 2000. Characterization of a Genetic Element
463		Carrying the Macrolide Efflux Gene mef(A) in Streptococcus pneumoniae. 9. Antimicrob Agents
464		Chemother 44:2585–2587.

- 465 40. Pozzi G, Iannelli F, Oggioni M, Santagati M, Stefani S. 2004. Genetic Elements Carrying Macrolide
 466 Efflux Genes in Streptococci. 3. CDTID 4:203–206.
- 467 41. Del Grosso M, Scotto d'Abusco A, Iannelli F, Pozzi G, Pantosti A. 2004. Tn2009, a Tn916-Like

Element Containing *mef*(E) in *Streptococcus pneumoniae*. Antimicrob Agents Chemother 48:2037–
2042.

470 42. Del Grosso M, Camilli R, Iannelli F, Pozzi G, Pantosti A. 2006. The mef(E)-Carrying Genetic Element

471 (mega) of *Streptococcus pneumoniae*: Insertion Sites and Association with Other Genetic Elements.

472 Antimicrob Agents Chemother 50:3361–3366.

- 473 43. Shoemaker NB, Guild WR. 1974. Destruction of low efficiency markers is a slow process occurring at
 474 a heteroduplex stage of transformation. Molec Gen Genet 128:283–290.
- 475 44. Guild WR, Shoemaker NB. 1976. Mismatch correction in pneumococcal transformation: donor length
 476 and hex-dependent marker efficiency. J Bacteriol 125:125–135.
- 477 45. Claverys J-P, Prats H, Vasseghi H, Gherardi M. 1984. Identification of *Streptococcus pneumoniae*478 mismatch repair genes by an additive transformation approach. Mol Gen Genet 196:91–96.
- 479 46. Hoskins J, Alborn WE, Arnold J, Blaszczak LC, Burgett S, DeHoff BS, Estrem ST, Fritz L, Fu D-J,
- 480 Fuller W, Geringer C, Gilmour R, Glass JS, Khoja H, Kraft AR, Lagace RE, LeBlanc DJ, Lee LN,
- 481 Lefkowitz EJ, Lu J, Matsushima P, McAhren SM, McHenney M, McLeaster K, Mundy CW, Nicas TI,
- 482 Norris FH, O'Gara M, Peery RB, Robertson GT, Rockey P, Sun P-M, Winkler ME, Yang Y, Young-
- 483 Bellido M, Zhao G, Zook CA, Baltz RH, Jaskunas SR, Rosteck PR, Skatrud PL, Glass JI. 2001.
- 484 Genome of the Bacterium *Streptococcus pneumoniae* Strain R6. J Bacteriol 183:5709–5717.

485 47. Wehrli W. 1983. Rifampin: Mechanisms of Action and Resistance. Clinical Infectious Diseases
486 5:S407–S411.

487 48. Goldstein BP. 2014. Resistance to rifampicin: a review. J Antibiot 67:625–630.

- 488 49. Sundin GW, Weigand MR. 2007. The microbiology of mutability. FEMS Microbiology Letters
 489 277:11–20.
- 490 50. Gressel J, Levy AA. 2010. Stress, Mutators, Mutations and Stress Resistance, p. 471–483. In Pareek,
- 491 A, Sopory, SK, Bohnert, HJ (eds.), Abiotic Stress Adaptation in Plants: Physiological, Molecular and
 492 Genomic Foundation. Springer Netherlands, Dordrecht.
- 493 51. Jolivet-Gougeon A, Kovacs B, Le Gall-David S, Le Bars H, Bousarghin L, Bonnaure-Mallet M, Lobel
- B, Guillé F, Soussy C-J, Tenke P. 2011. Bacterial hypermutation: clinical implications. Journal of
 Medical Microbiology 60:563–573.
- 496 52. Tanaka MM, Bergstrom CT, Levin BR. 2003. The Evolution of Mutator Genes in Bacterial
 497 Populations: The Roles of Environmental Change and Timing. Genetics 164:843–854.

498 53. Marles-Wright J, Lewis RJ. 2007. Stress responses of bacteria. Current Opinion in Structural Biology

- **499** 17:755–760.
- 500 54. Negri M-C, Morosini M-I, Baquero M-R, Campo R del, Blázquez J, Baquero F. 2002. Very Low

501 Cefotaxime Concentrations Select for Hypermutable *Streptococcus pneumoniae* Populations. AAC
502 46:528–530.

- 503 55. Gould CV, Sniegowski PD, Shchepetov M, Metlay JP, Weiser JN. 2007. Identifying Mutator
- 504 Phenotypes among Fluoroquinolone-Resistant Strains of *Streptococcus pneumoniae* Using Fluctuation
 505 Analysis. Antimicrob Agents Chemother 51:3225–3229.
- 506 56. Cortes PR, Piñas GE, Albarracin Orio AG, Echenique JR. 2008. Subinhibitory concentrations of
- 507 penicillin increase the mutation rate to optochin resistance in *Streptococcus pneumoniae*. Journal of
- 508 Antimicrobial Chemotherapy 62:973–977.

- 509 57. Henriques-Normark B, Blomberg C, Dagerhamn J, Bättig P, Normark S. 2008. The rise and fall of
 510 bacterial clones: *Streptococcus pneumoniae*. Nat Rev Microbiol 6:827–837.
- 511 58. Giraud A, Matic I, Radman M, Fons M, Taddei F. 2002. Mutator Bacteria as a Risk Factor in

512 Treatment of Infectious Diseases. Antimicrob Agents Chemother 46:863–865.

- 513 59. Labat F, Pradillon O, Garry L, Peuchmaur M, Fantin B, Denamur E. 2005. Mutator phenotype confers
- advantage in *Escherichia coli* chronic urinary tract infection pathogenesis. FEMS Immunology &
 Medical Microbiology 44:317–321.
- 516 60. Denamur E, Matic I. 2006. Evolution of mutation rates in bacteria. Mol Microbiol 60:820–827.
- 517 61. Oliver A, Mena A. 2010. Bacterial hypermutation in cystic fibrosis, not only for antibiotic resistance.
- 518 Clinical Microbiology and Infection 16:798–808.
- 62. Gifford DR, Berríos-Caro E, Joerres C, Galla T, Knight CG. 2019. Mutators drive evolution of multiresistance to antibiotics. bioRxiv 643585.
- 521 63. Nguyen SV, McShan WM. 2014. Chromosomal islands of *Streptococcus pyogenes* and related
- 522 streptococci: molecular switches for survival and virulence. Frontiers in Cellular and Infection523 Microbiology 4.
- 64. Huang J, Dai X, Wu Z, Hu X, Sun J, Tang Y, Zhang W, Han P, Zhao J, Liu G, Wang X, Mao S, Wang
 Y, Call DR, Liu J, Wang L. 2023. Conjugative transfer of streptococcal prophages harboring antibiotic
 resistance and virulence genes. ISME J 1–15.
- 527 65. Claverys JP, Lacks SA. 1986. Heteroduplex Deoxyribonucleic Acid Base Mismatch Repair in Bacteria.
 528 Microbiol Rev 50:33.
- 529 66. Salles C, Créancier L, Claverys JP, Méjean V. 1992. The high level streptomycin resistance gene from
 530 *Streptococcus pneumoniae* is a homologue of the ribosomal protein S12 gene from *Escherichia coli*.
- 531
 22. Nucleic Acids Res 20:6103–6103.

- 532 67. Muñoz R, Bustamante M, de la Campa AG. 1995. Ser-127-to-Leu substitution in the DNA gyrase B
 533 subunit of *Streptococcus pneumoniae* is implicated in novobiocin resistance. Journal of Bacteriology
- 534 177:4166–4170.

Strain	Relevant properties ^a	Origin (Reference)
R6	Unencapsulated transformation recipient, Hex ⁺	(Hoskins et al. 2001)
Rx1	Unencapsulated transformation recipient, Hex-	(Cuppone et al. 2021)
FP10	Rx1 competence deficient derivate, $\Delta comC$; str-41, Cm ^R , Sm ^R	(Santoro et al 2010)
FP11	Rx1 competence deficient derivate, $\Delta comC$; nov-1, Cm ^R , Nov ^R	(Santoro et al 2010)
FR125	FP10 derivative carrying Φ 1207.3; $\Delta comC$, str-41, $\Delta celB$; Cm ^R , Sm ^R , Spc ^R , Em ^R	(Iannelli unpublished)
FR3	FP10 derivative carrying Φ 1207.3 (by transformation with FR125 chromosomal DNA), Cm ^R , Sm ^R , Em ^R	(Iannelli unpublished)
FR169	FP11 derivative carrying Φ 1207.3 (by conjugation with FR3), Cm ^R , Nov ^R , Em ^R	(Iannelli unpublished)
FR170	FR3 derivative carrying Φ 1207.3 Δ orf6-orf11 (by transformation with the orf6-orf11 PCR mutagenic construct), Cm ^R , Sm ^R , Em ^R , Km ^R	(Iannelli unpublished)
FR171	FP11 derivative carrying Φ 1207.3 Δ <i>orf6-orf11</i> (by conjugation with FR170), Cm ^R , Nov ^R , Em ^R , Km ^R Em ^R	(Iannelli unpublished)
FR172	R6 derivative, <i>str-41</i> (by transformation with a PCR fragment obtained from FP10), Sm^{R}	This study
FR173	FR172 derivative carrying Φ 1207.3 (by conjugation with FR169), Sm ^R , Em ^R	This study
FR174	FR172 derivative carrying Φ 1207.3 Δ orf6-orf11 (by conjugation with FR171), Sm ^R , Em ^R , Km ^R	This study

^aHex is the DNA Mismatch Repair System (65), *str-41* and *nov-1* indicate point mutations conferring resistance to streptomycin and novobiocin, respectively (66, 67). Cm, chloramphenicol;

- 542 Sm, streptomycin; Nov, novobiocin; Spc, spectinomycin; Em, erythromycin; Km, kanamycin. The
- *orf6-orf11* region encodes the SOS-like cassette of the S. pyogenes Φ 1207.3 prophage.

Table 2. Φ 1207.3 SOS-like cassette predicted structural proteins.

ORF (aa) ^a	Predicted protein	Pfam domain(s) (aa) ^b [E value]	Conserved Domain Database (aa) ^c [E value]	Homologous protein ID / Origin (identity %) [E value] ^d	Phyre2 (confidence) ^e [identity %]
orf6 (99)		DUF5960 (11-92) [2.0e-32]			Chain B lambda repressor, Escherichia virus Lambda (100%) [21%]
orf7 (122)	YolD-like protein	YolD-like protein (28–111) [3.2e-12]		WP_052006222.1/ Bacillus subtilis plasmid pLS20 (38) [2e-5]	Ribosome maturation factor rimp, <i>Escherichia coli</i> (88.9%) [16%]
orf8 (471)	SOS response UmuC- like protein	IMS (109-207) [2.3e-11] IMS_HHH (223-255) [4.5e-06] IMS_C (303-406) [4.9e-14]	PolY_Pol_V_umuC (15-400) [2.37e- 121]	EU351020 orf70/Tn5253 Streptococcus pneumoniae (73) [0.0]	Y-family DNA Polymerase, Saccharolobus solfataricus (100%) [25%]
orf9 (46)				,	Bacterial cell-division inhibitor MinC, <i>Thermotoga maritima</i> (30.4%) [36%]
orf10 (75)					Putative DNA mismatch repair protein, <i>Bacteroides</i> <i>thetaiotaomicron</i> (31.6%) [30%]
orf11 (229)	LexA-like repressor	HTH_3 (7-61) [1.1e-12] Peptidase_S24 (111-220)[4.3e-22]	HTH_XRE (6-61) [1.64e-14] Peptidase_S24 (111-220) [1.62e-29]	CAD89881.1/ Lactococcus lactis subsp. cremoris (31) [9e-32]	

5487e number of the amino acid (aa) of the predicted protein is reported in parenthesis
5487he number in parenthesis indicates the region of the protein homologous to the Pfam domain or to the domain presnt in the Conserved Domain Database
5490termined by compositional matrix adjustment
5500nfidence indicates the reliability of the alignment





