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Highlights

▶ We construct a plasmid vector for regulated gene expression in pneumococcus. ▶ Expression of target genes is induced by maltose. ▶ We test plasmid vector by cloning and expressing the *gfp* reporter gene. ▶ The expression system is stable under repression and induction conditions.

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Construction of a plasmid vector based on the pMV158 replicon 2 for cloning and inducible gene expression in Streptococcus pneumoniae 3

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ABSTRACT

We report the construction of a plasmid vector designed for regulated gene expression in Streptococcus pneumoniae. The new vector, pLS1ROM, is based on the replicon of the streptococcal promiscuous rolling circle replication (RCR) plasmid pMV158. We inserted the controllable promoter P_M of the S. pneumoniae malMP operon, followed by a multi-cloning site sequence aimed to facilitate the insertion of target genes. The expression from P_M is negatively regulated by the transcriptional repressor MalR, which is released from the DNA operator sequence by growing the cells in maltose-containing media. To get a highly regulated expression of the target gene, MalR was provided in cis by inserting the malR gene under control of the constitutive Ptet promoter, which in pMV158 directs expression of the *tetL* gene. To test the functionality of the system, we cloned the reporter gene gfp 33 from Aequorea victoria, encoding the green fluorescent protein (GFP). Pneumococcal cells 34 35 harboring the recombinant plasmid rendered GFP fluorescence in a maltose-dependent 36 mode with undetectable background levels in the absence of the inducer. The new vector, 37 pLS1ROM, exhibits full structural and segregational stability and constitutes a valuable tool for genetic manipulation and regulated gene expression in S. pneumoniae.

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1. Introduction 42

43 Streptococcus pneumoniae is a Gram-positive human 44 pathogen that causes invasive infections such as pneumo-45 nia, meningitis or sepsis. In addition, pneumococcus is the leading bacterial cause of childhood mortality due to acute 46 47 respiratory infections and nonepidemic meningitis in 48 developing countries (WHO, 2009). Increasing antibiotic resistance of pathogenic S. pneumoniae is still an important 49 50 problem despite the use of the conjugated pneumococcal vaccine and improvements in antimicrobial therapy 51 52 (Perez-Trallero et al., 2010; Williams et al., 2002). Even though S. pneumoniae is a naturally transformable 53

bacterium that is able to accept and transfer plasmids originally isolated from other species (Barany and Tomasz, 1980; Stassi et al., 1981), the occurrence of indigenous plasmids in pneumococcus seems to be very low. Most of the pneumococcal plasmids characterized so far are related to pDP1 (Smith and Guild, 1979). An exception is plasmid pSpnP1, which was identified in a multidrug-resistant strain of S. pneumoniae (Romero et al., 2007). Sequence analysis showed that they are rolling circle plasmids of the pC194/pUB110 family.

The availability of plasmid-based genetic tools to study the expression of native and heterologous genes in different microorganisms has a special interest for both basic research and practical applications. To date, the number of non-integrative plasmid vectors available to manipulate S. pneumoniae is still very poor. In fact, most of the constructs are based on the broad-host-range RCR plasmid

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71 pMV158, naturally isolated from *Streptococcus* agalactiae 72 (Burdett, 1980), and on its derivative pLS1 (Stassi et al., 73 1981). In addition, the pneumococcal cryptic plasmid 74 pRMG1 (related to pDP1) has been used to construct a 75 shuttle vector between pneumococcus and Escherichia coli 76 (Muñoz et al., 1999). Vectors based on the promiscuous 77 pMV158 replicon have been used to clone pneumococcal 78 genes and to analyze its heterologous expression (Espinosa 79 et al., 1984; Lacks et al., 1986). Recently, a plasmid 80 vector based on pLS1 has been presented as an useful ge-81 netic tool to study promoter and terminator sequences in S. pneumoniae and Enterococcus faecalis (Ruiz-Cruz et al., 82 2010). Plasmids pLS1RGFP (Nieto et al., 2000), pLS1GFP 83 84 (Fernández de Palencia et al., 2000), pMV158GFP (Nieto and Espinosa, 2003), and pLS70GFP (Acebo et al., 2000) 85 constitute another set of pMV158 replicon-based tools 86 for the regulated expression of the gfp reporter gene. In 87 88 these constructs, expression of gfp was controlled by the 89 regulatory mechanism of maltosaccharides utilization in 90 S. pneumoniae (Nieto et al., 1997). This system has been validated in pneumococcus (Nieto et al., 2000) and used 91 92 to detect pneumococcal cells by fluorescence microscopy 93 in culture medium (Nieto and Espinosa, 2003) or to con-94 firm its intracellular location in granulocytes and microglia 95 (Letiembre et al., 2005; Ribes et al., 2010).

In the present work we have constructed pLS1ROM, a 96 97 new pLS1-based vector designed for cloning and regulated gene expression in S. pneumoniae. The presence of a 98 99 multi-cloning site that helps the insertion of the target gene immediately after the maltose-inducible promoter 100 P_M , the tight transcriptional control of P_M arising from 101 the presence in cis of the MalR repressor (Nieto et al., 102 2000), and the minimization of the plasmid size by 103 104 removal of dispensable regions are the more remarkable features of the pLS1ROM pneumococcal vector. The suit-105 ability of pLS1ROM as a regulated expression vector has 106 107 been tested by cloning the *gfp* reporter gene and analyzing 108 the fluorescence of pneumococcal cells containing the resulting recombinant plasmid pLS1ROM-GFP upon malt-109 ose induction. In contrast to the previously constructed 110 pLS1RGFP plasmid, pLS1ROM-GFP exhibits structural and 111 segregational stability under induction conditions, which 112 113 represents a significant improvement over the prior maltose-inducible pneumococcal system. 114

115 **2. Materials and methods**

116 2.1. Bacterial strains and plasmids

S. pneumoniae cells were grown in AGCH medium 117 (Lacks, 1968) supplemented with 0.3% sucrose and 0.2% 118 119 yeast extract, with selection for resistance to erythromycin (Ery, $1 \mu g/ml$) when required. All cultures were grown at 120 37 °C. The constructs were done in S. pneumoniae 708 121 122 (end-1 exo-1 trt-1 hex-4 malM594; (Lacks and Greenberg, 123 1977). S. pneumoniae R6 (wild type; Lacks, 1968) was used for the expression of gfp and detection of GFP fluorescence. 124 125 Other plasmids used were pLS1R (initially named as 126 pAPM22; (Puyet et al., 1993), pCL1 (Nieto et al., 2000) 127 and pGreenTIR (Miller and Lindow, 1997).

2.2. Plasmid DNA preparation and DNA manipulations

Pneumococcal cells were grown and transformed 129 with plasmid DNA as previously reported (Lacks et al., 130 1986; Stassi et al., 1981). Plasmid DNAs purified from 131 S. pneumoniae were prepared as described (del Solar 132 et al., 1987). Restriction endonucleases, the Klenow 133 fragment of DNA polymerase I (PolIK), and T4 DNA ligase 134 were purchased from New England Biolabs or Roche, and 135 were used according to the manufacturer's instructions. 136 Restriction- or PCR-derived DNA fragments employed for 137 cloning experiments were purified from agarose gels by 138 phenol extraction as described (Sambrook et al., 1989). 139 The duplex oligonucleotide MCS having 5'-protuding ends 140 compatible with BamHI-generated ends was obtained by 141 annealing complementary oligonucleotides MCSUP and 142 MCSDOWN (MCSUP: <u>5'-GATCAAGCTTCCGCGGGCCCGGGC</u> 143 ATGCGGCC- GCACTAG (); MCSDOWN: 5'-GATCCACTAG TGCGGCCG—CATGCCCGGGGCCCGCGGAAG 144 145 ing ends are shown underlined in both ongonucleotides), 146 which were previously purified from gel as described 147 (Maniatis et al., 1982). Reduction of the plasmid vector size 148 was performed by inverse PCR (Ochman et al., 1988). Two 149 different sets of divergent primers were employed to 150 eliminate the Px promoter (DelX1: 5'-TCGATTCTGAGG 151 CCTCTGAG-3'; DelX2: 5'-TACCTCCCTGGTTCTAATCC-3') 152 and the tetL gene (Deltet1: 5'-GTCATTAGTTGGCTGGTTAC-153 3'; Deltet2: 5'-GGTTAATGATACGCTTCC-3'), by using 154 pLS1ROXM-MCS and pLS1ROMtelL-MCS as template, 155 respectively. After amplification with the Phusion poly-156 merase (Finnzymes), the resultant DNA fragments were 157 gel-purified and incubated with T4 DNA ligase to generate 158 circular molecules that were used to transform S. pneumo-159 niae cells. To allow ligation-sealing of the cyclized DNA 160 molecules, the primers were phosphorylated at their 5' 161 end previously to being used in the amplification reaction. 162 All new plasmid constructs were confirmed by automated 163 sequencing. 164

2.3. Determination of structural and segregational stability

Plasmid structural and segregational stability was 166 determined in exponentially growing cultures of S. pneu-167 moniae R6 containing either pLS1ROM or pLS1ROM-GFP. 168 Pneumococcal cells were grown under different conditions 169 (0.3% sucrose or maltose as carbon source; presence or ab-170 sence of selective pressure (Ery $1 \mu g/ml$) for the resident 171 plasmid). Samples were taken after the indicated number 172 of generations in order to analyze the integrity of the plas-173 mid DNA (by electrophoresis of total DNA preparations in 174 1% agarose gels; del Solar and Espinosa, 1992; Lacks 175 et al., 1986), as well as the percentage of plasmid-contain-176 ing (and hence Ery resistant) cells. 177

2.4. Measurement of GFP fluorescence

To measure the GFP fluorescence inside the pneumococ-179cal cells we followed the procedure described in (Nieto et al.,1802000). Pneumococcal cells harboring pLS1ROM-GFP were181grown in sucrose-containing media, with Ery resistance182(Ery^R) selection, to an OD₆₅₀ of 0.5 (about 4×10^8 c.f.u./ml)183

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and then diluted 1:1000 into medium containing maltose, 184 185 with or without selective pressure, and allowed to again 186 reach an OD₆₅₀ of 0.5. Cells (1 ml from each culture) were 187 sedimented by centrifugation and suspended in the same 188 volume of PBS buffer (10 mM Na₂HPO₄, 1 mM KH₂PO₄, 189 140 mM NaCl, 3 mM KCl), pH 7.2. Aliquots (200 µl) were 190 used to measure the fluorescence intensity on a Varioskan 191 Flash spectral scanning multimode reader (Thermo Scien-192 tific) by excitation at a wavelength of 488 nm and detection 193 of emission at 510 nm. As a control to determine the background fluorescence, cells harboring the plasmid vector 194 195 pLS1ROM (lacking the gfp gene) were used.

196 3. Results and discussion

197 3.1. Construction of the regulated expression vector pLS1ROM

To construct the pLS1ROM vector we started with plas-198 mid pLS1R, which has the pMV158 replicon and contains 199 the *malR* gene cloned under the control of the P_{tet} promoter 200 201 that directs constitutive expression of the tetracycline 202 resistance tetL gene of pMV158 (Puyet et al., 1993) 203 (Fig. 1). DNA from pLS1R was digested with HindIII and EcoRI, and treated with PolIK before purifying the larger 204 fragment. To obtain a DNA region containing the opera-205 tors/promoters P_X and P_M we used plasmid pCL1 (Nieto 206 et al., 2000). DNA from this plasmid was digested with Sall, 207 208 treated with PolIK, and then digested with SspI. The resultant \sim 1 kb fragment was purified and ligated to the pLS1R 209 210 larger fragment. The ligation mixture was used to transform S. pneumoniae, and transformants containing the 211 resultant construction pLS1ROXM (Fig. 1) were selected 212 213 for Ery^R. Transcription initiated from P_M is oriented in

opposite direction to the other protein-encoding genes of 214 pLS1ROXM (Fig. 1). Next step in the construction of the 215 cloning vector was to introduce a multi-cloning site se-216 quence (MCS) immediately after promoter P_M. To this 217 end, the DNA from pLS1ROXM was linearized with BamHI 218 and ligated to the 41-bp duplex oligonucleotide MCS, 219 whose sequence had been designed to reconstitute a 220 BamHI site at only one of its ends upon ligation with 221 BamHI-generated fragments. Hence, the reconstituted 222 BamHI site, which is to remain unique in the new construct 223 pLS1ROXM-MCS, constitutes an additional single restric-224 tion site within the MCS. Moreover, the Xbal single site 225 located between the P_M promoter and the MCS increases 226 the number of restriction sites that can be used for cloning 227 target genes (Fig. 1). Plasmid pLS1ROXM-MCS was 228 obtained after transforming the ligation mixture into 229 S. pneumoniae and selecting for Ery^R. 230

In order to improve the plasmid pLS1ROXM-MCS as an 231 expression vector several modifications were carried out. 232 First, we eliminated the P_X/O_X , region, which is unnecessary 233 for the MalR-mediated repr<mark>ssion</mark> of P_M. In fact, the tran-234 scription rate from P_M has been reported to increase when 235 the region encompassing P_X is deleted (Nieto et al., 2001). 236 Region P_X/O_X , was removed by inverse PCR, using the diver-237 gent phosp or lated primers DelX1 and DelX2, and DNA 238 from pLS1R MCS as template. The amplification reac-239 tion yielded a linear DNA fragment corresponding to almost 240 the entire vector except the P_x/O_x region. The amplified frag-241 ment was gel-purified and subjected to auto-ligation to ren-242 der circular plasmids molecules that were used to transform 243 S. pneumoniae. The resultant plasmid was termed pLS1ROM-244 tetL-MCS. Second, a ~1.4 kb DNA fragment containing the 245 tetL gene of pMV158 was eliminated. This gene was ex-246 pressed neither in pLS1 nor in pLS1ROMtetL-MCS, as it was 247



Fig. 1. Regulated expression vector pLS1ROM. (A) Construction of pLS1ROM. The various steps in the construction of pLS1ROM are indicated. PMV158-born genes (black) and the *ermAM* gene (gray) are indicated by arrows. Chromosomal-born *malR* gene and the region containing the promoters P_M and P_X are indicated by open arrows and open boxes respectively. The promoter sequences are indicated by triangles pointing in the direction of transcription. (B) Sequence of the plasmid vector region encompassing the inducible promoter P_M , the operator O_M and the MCS.

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248 uncoupled from its promoter. To delete this sequence we 249 followed the same strategy of inverse PCR but using the 250 divergent phosphorylated primers Deltet1 and Deltet2, 251 and DNA from pLS1ROMtetL-MCS as template. This deletion 252 did not include the transcriptional terminator of the *tetL* 253 gene, which was left to attenuate possible convergent 254 transcription from the ermAM gene. The final plasmid vector 255 pLS1ROM is 6.8 kb in size, confers Ery^R due to the presence of 256 the *ermAM* gene, and facilitates the insertion of the target 257 gene immediately after the regulated promoter P_M (Gen-Bank accession JN381945). 258

3.2. Functionality of pLS1ROM: cloning and expression of the
 gfp gene in the regulated expression vector, and stability of the
 vector and recombinant plasmids

262 To assess the functionality of the constructed expression system, the gfp reporter gene from Aequorea victoria, 263 encoding GFP, was fused to the promoter \overline{P}_{M} , and the fluo-264 rescence emission was measured. Specifically, we have 265 used the gfp cassette from plasmid pGreenTIR, which was 266 designed for prokaryotic transcriptional fusions and con-267 tains a mutated gfp gene (harboring the S65T "red shift" 268 and F64L "protein solubility" amino acid changes), a trans-269 270 lational enhancer and a consensus ribosome binding site, so that synthesis of GFP is enhanced. This optimized gfp 271 cassette has been efficiently expressed in S. pneumoniae 272 273 (Nieto et al., 2000) and in other Gram-positive hosts like 274 Bacillus subtilis (Serrano-Heras et al., 2005), Lactococcus lactis (Fernández de Palencia et al., 2000), and E. faecalis 275 276 (Lorenzo-Díaz and Espinosa, 2009; Ruiz-Cruz et al., 2010). 277 Recombinant plasmid pLS1ROM-GFP was constructed by inserting the gfp cassette from pGreenTIR under the control 278 279 of P_M. To this end, DNA from pLS1ROM was linearized with 280 BamHI and the resultant fragment was purified. The gfp cassette was extracted by digesting DNA from pGreenTIR 281

with BamHI. The resultant 810 bp DNA fragment was purified and ligated to the pLS1ROM fragment.

The stability of pLS1ROM and pLS1ROM-GFP was tested in cultures of *S. pneumoniae* R6 growing exponentially in media containing sucrose (repression conditions) and in the absence of Ery. The results showed that both plasmids were stably inherited, with no detectable appearance of Ery-sensitive plasmid-free cells over at least 100 generations (Fig. 2). Moreover, analysis of the total DNA content confirmed that the plasmids analyzed were stably maintained with no modification or rearrangement observed after 100 generations (Fig. 2).

Expression vectors and their recombinant derivatives harboring the gene to be expressed should ideally exhibit structural and segregational stability. It has been reported that RCR vectors for use in Gram-positive bacteria frequently become structurally and/or segregationally unstable upon insertion of even relatively small foreign DNA fragments (Grkovic et al., 2003; Leer et al., 1992). This is shown here not to be the case with <u>pLS1ROM-GFP</u>, which is stably maintained in pneumococcal cells, hence proving the suitability of pLS1ROM as a cloning vector for this bacterium.

We also analyzed the stability of the expression from the regulated promoter P_M by following the fluorescence emitted by the GFP protein in pLS1ROM-GFP-carrying pneumococcal cells that had been grown for several generations in the absence of both Ery and maltose. To induce GFP fluorescence, pneumococcal cultures grown under the above conditions for the indicated generations (Fig. 3A) were diluted 1:1000 into medium containing maltose and allowed to reach an OD₆₅₀ of 0.5. The results showed that the fluorescence emission is maintained at similar levels in the interval of generations analyzed (Fig. 3A), indicating that the integrity and functionality of the transcriptional fusion between P_M and the *gfp* gene is



Fig. 2. Structural and segregational stability of pLS1ROM and pLS1ROM-GFP. Total DNA extracted from cultures of pLS1ROM- or pLS1ROM-GFP-harboring pneumococcal cells that had been grown for the indicated number of generations in medium lacking Ery was analyzed by 1% agarose gel electrophoresis (left panel). To facilitate the identification of the main plasmid forms, pLS1ROM and pLS1ROM-GFP DNA samples were obtained by the alkaline lysis method and loaded to the left of the corresponding total DNA extracts. OC, open circular DNA; SC, supercoiled monomeric plasmid DNA. The percentage of Ery-resistant colonies is given in the right panel.

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Fig. 3. GFP fluorescence in pneumococcal cells harboring pLS1ROM-GFP. (A) Pneumococcal cells carrying pLS1ROM or pLS1ROM-GFP were grown in medium lacking both Ery and maltose for the indicated number of generations. Production of GFP was subsequently induced by growing these cells for 10 generations in the presence of maltose. Fluorescence background levels were obtained from cells grown in the absence of maltose for 10 more generations. (B) The fluorescence emitted by pneumococcal cells carrying pLS1ROM-GFP under induction (maltose) and repression (sucrose) conditions was compared with the fluorescence due to the previously reported construction pLS1RGFP (Nieto et al., 2000) under the same conditions. The fluorescence emission data of pneumococcal cells carrying the S1R and pLS1ROM were included as a control.



Fig. 4. Structural and segregational stability of pLS1RGFP, pLS1ROM and pLS1ROM-GFP under induction conditions. Pneumococcal cultures harboring the indicated plasmids were grown to an OD₆₅₀ of 0.5 in medium containing sucrose with Ery^R selection (generation 0), and then diluted 1:2000 into medium containing maltose, with or without Ery. After growing the cultures for the indicated number of generations total DNA preparations were analyzed by 1% agarose gel electrophoresis. As a control the same pneumococcal cultures were diluted into medium containing sucrose, with or without Ery, and subjected to the same analysis. The main plasmid forms of pLS1ROM and pLS1ROM-GFP were identified in DNA samples prepared by alkaline lysis (lanes C). OC, open circular DNA; SC, supercoiled monomeric plasmid DNA; Del, deleted plasmid DNA.

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318 preserved. These results are consistent with the new 319 expression system being structurally and segregationally 320 stable even in the absence of selective pressure, as demon-321 strated by the results shown in Fig. 2.

322 Since transcription from the regulated promoter P_M converges on and might collide with that of the ermAM 323 324 gene in both pLS1ROM (Fig. 1) and pLS1ROM-GFP (the 325 cloned gfp gene lacked an intrinsic terminator), it was 326 interesting to know whether, under induction conditions, expression of *ermAM* was affected so that selectable levels 327 of Ery^R could not be achieved. To test this, we monitored 328 the growth rate of pneumococcal cells carrying pLS1ROM 329 or pLS1ROM-GFP under induction conditions (maltose) 330 with or without Ery^R selection. As a control, the same cells 331 were grown in medium with sucrose (repression condi-332 tions), with or without Ery. The results (not shown) dem-333 onstrated that the duplication time of the pneumococcal 334 335 cells grown in the presence of Ery was not significantly affected by the presence or absence of maltose, indicating 336 337 that induction of transcription from P_M did not severely af-338 fect expression of the ermAM gene.

339 When analyzing the expression of gfp from the regulated 340 promoter P_M, we observed that the fluorescence emitted in 341 induced cultures of pneumococcal cells carrying pLS1ROM-342 GFP was remarkably higher than that of pneumococci harboring the previously-constructed plasmid pLS1RGFP 343 (Fig. 3B). Furthermore, the cells carrying pLS1RGFP took 344 approximately four times longer to duplicate in maltose-345 346 containing medium than in sucrose-containing medium, irrespective of the presence or absence of the antibiotic 347 (not shown). Inhibition of the bacterial cell growth under 348 induction conditions was accompanied by progressive plas-349 mid loss, close to 100% after 11 generations, in the absence 350 351 of Ery, or by deletions reducing significantly the plasmid size in the presence of the antibiotic (Fig. 4). In contrast, 352 353 the cells harboring pLS1ROM-GFP did not show any increase in their duplication time and the plasmid was 354 355 stably maintained, at least during the generations analyzed, 356 when grown in maltose-containing medium (Fig. 4). The 357 instability of pLS1RGFP in induced pneumococcal cultures 358 can explain the lower level of fluorescence observed in cells carrying this plasmid compared with those harboring 359 360 pLS1ROM-GFP. The stability difference between these two pMV158 replicon-based plasmids may arise from promoter 361 P_M directing transcription toward the replication origin in 362 pLS1RGFP (Nieto et al., 2000), which could destabilize the 363 plasmid under induction conditions. In contrast, 364 365 pLS1ROM-GFP has P_M orientated in opposite direction relative to the plasmid replicon, and lacks the divergent 366 367 P_X promoter.

4. Conclusions 368

In this manuscript we describe the construction of a 369 370 new regulated expression vector, pLS1ROM, which is based 371 on the streptococcal pMV158 replicon and has been 372 specially designed to clone and express both native and 373 heterologous genes in S. pneumoniae. We have included a 374 MCS sequence to facilitate the insertion of the target gene, 375 improved the expression from the regulated promoter P_M

and reduced the plasmid size to increase the stability of 376 377 the vector. We have proved the suitability of this vector by cloning and willfully expressing the gfp gene. In conclu-378 sion, pLS1ROM is a stable and functional expression vector 379 that increases the spectrum of genetic tools available to manipulate S. pneumoniae.

Acknowledgments

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