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8	Production of multiple bacteriocins from a single locus by gastrointestinal strains
9	of Lactobacillus salivarius
10	
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## 30 Abstract

31 Bacteriocins produced by *Lactobacillus salivarius* isolates derived from

32 gastrointestinal origin have previously demonstrated efficacy for in vivo protection 33 against Listeria monocytogenes infection. In this study, comparative genomic analysis 34 was employed to investigate the intraspecies diversity of seven L. salivarius isolates 35 of human and porcine intestinal origin, based on the genome of the well characterised 36 bacteriocin-producing strain L. salivarius UCC118. This revealed a highly conserved 37 megaplasmid-encoded gene cluster in these strains involved in the regulation and 38 secretion of two-component class IIb bacteriocins. However, considerable 39 intraspecific variation was observed in the structural genes encoding the bacteriocin 40 peptides. These ranged from close relatives of abp118 such as salivaricin P, which 41 differs by 2 amino acids, to completely novel bacteriocins such as salivaricin T, which 42 is characterized in this study. Salivaricin T inhibits closely related lactobacilli and 43 bears little homology to previously characterized salivaricins. Interestingly, the two 44 peptides responsible for salivaricin T activity, SalT $\alpha$  and SalT $\beta$ , share considerable 45 identity with the component peptides of thermophilin 13, a bacteriocin produced by 46 Streptococcus thermophilus. Furthermore, the salivaricin locus of strain DPC6488 47 also encodes an additional novel one-component class IId anti-listerial bacteriocin, 48 salivaricin L. These findings suggest a high level of redundancy in the bacteriocins 49 that can be produced by intestinal *L. salivarius* isolates using the same enzymatic 50 production and export machinery. Such diversity may contribute to their ability to 51 dominate and compete within the complex microbiota of the mammalian gut.

## 52 Introduction

53 There is increasing evidence to suggest that bacteriocin production is a 54 desirable probiotic trait which enables the establishment and persistence of the 55 producing strains within the gastrointestinal tract (GIT) (2, 9, 15, 35). These 56 antimicrobials may have a narrow or broad spectrum of inhibition (16). Although 57 broad spectrum bacteriocins, such as nisin, have been useful with respect to the 58 control of spoilage and pathogenic organisms in food preservation (8), there has been 59 increasing interest in narrow spectrum bacteriocins, such as thuricin CD, due to their 60 narrow specificity and minimal impact on non-target beneficial GIT microbes (30). 61 As such, these bacteriocins offer potential alternatives to traditional antibiotics with 62 respect to controlling pathogens within the gut (29). In situ production of such narrow 63 spectrum bacteriocins by probiotics would overcome complications such as the 64 proteolytic degradation of orally delivered antimicrobial peptides during gastric 65 transit. Lactobacillus salivarius is a promising probiotic candidate frequently isolated 66 from human, porcine and avian GITs, many of which are producers of unmodified 67 bacteriocins of class IIa (pediocin-like bacteriocins), class IIb (two-component 68 bacteriocins) and class IId (linear non-pediocin-like bacteriocins) (4, 13, 31). 69 Significantly, an *in vivo* demonstration of the anti-infective properties of the abp118-70 producing strain L. salivarius UCC118 has established the in vivo functionality of 71 such bacteriocins (6). In addition, purified OR7, a class IIa bacteriocin produced by 72 the chicken intestinal isolate L. salivarius NRRL B-30514, has also been successfully 73 employed to reduce *Campylobacter jejuni* colonization in poultry (31). 74 Closely related variants of abp118 (a two-component class IIb bacteriocin) 75 frequently occur in intestinally-derived L. salivarius from different hosts (2, 17, 26, 76 27), suggesting that this feature may be important for the successful establishment of

77	L. salivarius within the GIT. Further evidence of the ecological advantage that this
78	trait bestows upon the producing strains was provided by the salivaricin P-producing
79	L. salivarius DPC6005, which prevailed over four components of a probiotic
80	formulation within the porcine ileum (34). The production of analogous bacteriocins
81	by genetically-distinct strains and species is frequently ascribed to extensive
82	horizontal gene transfer events occurring within the GIT (1, 2, 15, 22, 33). It is thus
83	notable that abp118 is encoded on a megaplasmid in L. salivarius UCC118. Although
84	the transfer-associated genes within the conjugative megaplasmid pMP118 appear
85	non-functional in L. salivarius UCC118 (10), corresponding RepA-type megaplasmids
86	are universally present in L. salivarius (17). Interestingly, Wescombe et al., (2006)
87	demonstrated the in vivo conjugative transmission of bacteriocin-rich megaplasmids
88	in Streptococcus salivarius species, which was associated with the numerical
89	prominence of the species within the oral cavity (35).
90	The variable nature of <i>L. salivarius</i> with respect to their ability to produce
91	bacteriocins, and the nature of the bacteriocins which they produce, is becoming
92	increasingly apparent. It has been noted that many L. salivarius strains which harbour
93	homologues of the abp118 structural genes do not display anti-Listeria activity (17).
94	A recent comprehensive genomic analysis revealed that the three-component
95	regulatory system responsible for the transcriptional regulation of abp118 production
96	was not well conserved and was likely responsible for the bacteriocin negative
97	phenotype of these strains (28). Other strains produce closely related, yet distinct,
98	bacteriocins such as salivaricin P which differs from abp118 with respect to two
99	amino acids in their respective $\beta$ peptides (2). Interestingly, the nine associated
100	polymorphisms (three in the $\alpha$ gene, six in the $\beta$ gene) appear to be specific features
101	associated with L. salivarius isolates derived from porcine intestinal origin (2). It

102	would thus seem that gene acquisition, mutation and/or gene decay, all of which are
103	potential consequences of species adaptation to a specific ecological niche (3, 24),
104	have impacted on the adaptation of this widespread bacteriocin locus of L. salivarius,
105	driving genome-wide specialisation in response to a particular niche.
106	We conducted a genome-wide comparison of seven genetically distinct
107	bacteriocin-producing intestinal L. salivarius strains isolated in our laboratory which
108	revealed that the salivaricin/abp118-associated bacteriocin locus is a site of high
109	variability. In the case of the neonatal isolate L. salivarius DPC6488, two novel
110	narrow spectrum bacteriocins, salivaricin T, which displays considerable homology to
111	a two-component class IIb bacteriocin associated with Streptococcus thermophilus,
112	and salivaricin L, a one component class IId bacteriocin exhibiting anti-Listeria

113 activity, have been characterised in this study.

## 114 Materials and Methods

115 1	Bacterial strains and culture conditions. L. salivarius strains of both human and
116 p	porcine intestinal origin, previously isolated and characterised in our laboratory, were
117 i	ncluded in this study (Table 1). Lactobacilli were routinely cultured under anaerobic
118 c	conditions at 37°C in MRS medium (Difco Laboratories, Detriot, MI). Anaerobic
119 c	conditions were maintained with the use of anaerobic jars and Anaerocult A gas packs
120 (	Merck, Darmstadt, Germany). Listeria innocua DPC3572 and Listeria
121 n	nonocytogenes NCTC 11994 were grown aerobically at 37°C in BHI (Merck).
122	
123 <b>H</b>	Pulsed-field gel electrophoresis analysis. Differentiation of the L. salivarius isolates
124 v	was confirmed by PFGE as previously described (23). The megaplasmid content of
125 t	he strains was also determined by PFGE following S1 nuclease treatment of high-
126 n	nolecular-weight DNA, also previously described (17).
127	
128 <b>I</b>	<b>DNA amplification, sequencing and analysis.</b> Template DNA was extracted from <i>L</i> .
129 s	salivarius DPC6005 and L. salivarius DPC6488 to amplify the genetic loci
130 r	responsible for salivaricin production in the corresponding strains using primers
131 d	designed specific to the locus of abp118 (accession number AF408405). A series of
132 a	abp118-specific primers were used to amplify the salivaricin P locus of strain
133 I	DPC6005 by routine PCR generating a contiguous sequence of approximately 13 kb.
134 7	The primer pair 5' CCGCCGATATACTATTCGTGG 3' and 5'
135 C	GAGAGTTAGACCTGATGAAG 3' was also used to amplify the 13 kb salivaricin P
136 g	gene cluster in its entirety using Extensor Hi-Fidelity PCR mastermix (Abgene,

138 A PCR amplicon previously generated using DPC6488 template DNA and 139 primers specific for the amplification of the salivaricin P structural genes (23) 140 generated approximately 900 bp sequence data in the present study. Following 141 sequence analysis, the abp118-specific primer pair above was also employed to 142 amplify the 12 kb salivaricin gene cluster of strain DPC6488. Oligonucleotide primers 143 were synthesised by Sigma-Genosys (Poole, Dorset, UK) and purified amplicons were 144 sequenced by Beckman Coulter Genomics (Essex, UK). Alignments and analyses of 145 sequence data were performed using LASERGENE 6 software (DNAStar Inc., 146 Madison, WI). Database searches were performed using the basic local alignment 147 search tool (BLAST) on the National Centre for Biotechnology information (NCBI) 148 server (http://www.ncbi.nlm.nih.gov). Open reading frames (ORFs) were identified 149 using LASERGENE 6, ORF finder and glimmer on the NCBI server and Genemark, 150 gene prediction software (http://exon.biology.gatech.edu). 151

152 Microarray-based Comparative Genomic Hybridisation. Comparative genomic 153 hybridizations were performed using a highly replicated custom microarray (Agilent 154 Technologies, CA, USA) designed based on the genome of L. salivarius UCC118 as 155 previously described (5, 11, 28). The experimental procedures for genomic DNA 156 extraction, fragmentation and fluorescent labelling and co-hybridisation experiments 157 of fluorescently-labelled gDNA of the test and reference strains employed in this 158 study were recently described (28). Two-dye-swap replicate hybridisations were 159 performed for each strain tested and self-hybridisation of the reference strain was 160 carried out as a control experiment. Analysis of the microarray data were also as 161 described by Raftis et al. (28). The sequencing data obtained for our strains was in 162 agreement with the empirically determined cutoff intervals used to discriminate

163 between the presence, divergence and absence of genes previously selected on the 164 basis of a comparison of the output data of a BLASTn comparison of the UCC118-165 specific oligonucleotide probe set with the sequence of the draft genome of the type 166 strain Lactobacillus salivarius DSM20555 (accession number NZ ACGT00000000) with the log<sub>2</sub>-transformed signal ratios determined for the hybridisation reaction of the 167 168 gDNA of the corresponding strain (28). Therefore, these cuttoff intervals were also applied to the present dataset. Cutoff intervals of  $(\geq -1.5)$ ,  $(< -1.5, \geq -2.4)$ ,  $(< -2.4, \geq -2.4)$ 169 170 4.5),  $(< -4.5, \geq -5.8)$  and (< -5.8) corresponded to highly conserved, conserved, 171 divergent, highly divergent, and absent features in the test strain relative to L. 172 salivarius UCC118, respectively. Hierarchical clustering of strains was performed 173 using the complete linkage clustering and results were visualised using Genesis 174 software (32).

175

## 176 Purification of the antimicrobial peptides produced by *L. salivarius* DPC6488.

177 The antimicrobial peptides were purified from a 2 L overnight culture of L. salivarius 178 DPC6488 grown in MRS media. The cells were harvested by centrifugation at 8,000 179  $\times$  g for 15 min, resuspended in 250 mL 70% (vol/vol) propan-2-ol containing 0.1% 180 (vol/vol) trifluoric acid (TFA) and stirred at room temperature for 3 h. Cells were 181 removed by centrifugation, and propan-2-ol by rotary evaporation. The resultant 182 preparation was applied to a 5.0 g (20 mL volume) Strata C18 solid phase extraction 183 (SPE) column (Phenomenex, Cheshire, UK) pre-equilibrated with methanol and 184 water. The column was washed with 35% (vol/vol) ethanol and the antimicrobial 185 peptides were subsequently eluted with 70% (vol/vol) propan-2-ol containing 0.1% 186 (vol/vol) TFA. Following removal of the propan-2-ol from the preparation, 4 ml 187 aliquots were applied to a Jupiter proteo C12 reversed phase-high pressure liquid

chromatography (RP-HPLC) column (250.0  $\times$  10.0 mm, 4  $\mu$ m particle size, 90 Å pore 188 189 size; Phenomenex) pre-equilibrated with 25% (vol/vol) acetonitrile 0.1% (vol/vol) TFA. The column was developed in a gradient of 25% to 55% (vol/vol) acetonitrile 190 191 containing 0.1% (vol/vol) TFA from 5 to 50 min at a flow rate of 2.5 mL min<sup>-1</sup>. Individual fractions were assayed by well diffusion using the sensitive indicator strain 192 193 L. bulgaricus LMG 6901, and matrix-assisted laser desorption ionisation-time of 194 flight (MALDI-TOF) mass spectrometry (MS) was performed to determine fractions 195 containing the peptides of interest. Fraction 27, containing a mass of 4433 Da 196 corresponding to the salivaricin B peptide, was applied to a Luna analytical SCX 197 cation exchange HPLC column ( $250.0 \times 4.6$  mm, 5 µm particle size, 100 Å pore size; 198 Phenomenex), for further purification, following removal of acetonitrile by rotary 199 evaporation. Using buffer A (20 mM potassium phosphate containing 25% (vol/vol) 200 acetonitrile, pH 2.5), and buffer B (20 mM potassium phosphate, 25% (vol/vol) 201 acetonitrile, 1 M potassium chloride, pH 2.5) the column was pre-equilibrated with 202 10% buffer B and subsequently developed in a gradient of 10% to 65% buffer B, from 5 to 45 min at a flow rate of 1.0 ml min<sup>-1</sup>. Individual fractions of interest were applied 203 204 to a 200 mg (3 ml volume) Strata C18 SPE column (Phenomenex) pre-equilibrated 205 with methanol and water. The column was washed with 30% (vol/vol) ethanol and 206 40% (vol/vol) propan-2-ol, followed by the elution of the antimicrobial component 207 with 70% (vol/vol) propan-2-ol containing 0.1% (vol/vol) TFA. Similarly, fractions 208 38-40, containing masses of 5655 Da and 5267 Da corresponding to the respective 209 SalT $\alpha$  and SalT $\beta$  peptides, were pooled, concentrated and applied to the Luna 210 analytical SCX cation exchange HPLC column for separation and purification as 211 described above, as were fractions 44-46, in which a mass of 4117 Da was detected 212 corresponding to the mature salivaricin L peptide. Bacteriocin activity was monitored

throughout purification by well diffusion assay using the sensitive indicator strain *L*. *bulgaricus* LMG 6901. MALDI-TOF MS was performed as previously described (7).

216 Peptide synthesis. Synthetic analogues of mature salivaricin B and salivaricin L were 217 synthesised according to the deduced amino acid sequence of *slnT3* and ORF13 (*slnL*) 218 using microwave-assisted solid phase peptide synthesis (MW-SPPS) performed on a CEM Liberty<sup>TM</sup> microwave peptide synthesiser using a H-Ser-HMPB-ChemMatrix® 219 220 and a H-Cys(TRT)-HMPB-ChemMatrix® resin (PCAS Biomatrix Inc. Quebec, 221 Canada), respectively. The molecular masses of the synthetic analogues were 222 confirmed using MALDI TOF MS and the antimicrobial activity of the crude peptides 223 was assayed by well diffusion. Synthetic salivaricin L was purified by RP-HPLC using a Jupiter C5 column (250.0  $\times$  10.0 mm, 10  $\mu$ m particle size, 300 Å pore size; 224 225 Phenomenex) developed in a gradient from 25% to 55% (vol/vol) acetonitrile containing 0.1% TFA from 10 to 50 min at a flow rate of 2.5 mL min<sup>-1</sup>. Fractions 226 227 containing the desired molecular mass, identified by MALDI TOF MS, were pooled 228 and lyophilised using a Genevac HT 4X lyophiliser (Genevac Ltd. Ipswitch, UK). The 229 peptide was dissolved in 50% (vol/vol) acetonitrile at a concentration of 5 mg/ml and 230 stored at -20°C under nitrogen. Appropriate dilutions of the peptides in 50mM 231 potassium phosphate buffer pH 6.8 were used for bacteriocin assays. 232

Specific activity analysis of salivaricin L. A microtiter plate assay system was used to determine the minimum concentration of the synthetic salivaricin L analogue required to inhibit growth of the indicators, *L. innocua* DPC3572 and *L. monocytogenes* NCTC 11994 by 50% (MIC<sub>50</sub>). The microtitre plate was first treated with bovine serum albumin (BSA) to prevent adherence of the peptide to the sides of

238 the wells, as described previously (12). Each plate included triplicate assays at each 239 concentration examined. Each well contained a total volume of 200 µl, comprised of 240 purified synthetic salivaricin L, 150 µl of a 1-in-10 dilution of the indicator culture 241  $(A_{590} \text{ of } 0.5)$  in BHI broth. Control wells contained media only (blanks), or untreated 242 indicator culture. The microtiter plate cultures were then incubated at 37°C and the 243 optical density at 590 nm (OD<sub>590</sub>) recorded at hourly intervals for 6 h (GENios plus; 244 TECAN, Switzerland). Triplicate readings were averaged and blanks were subtracted 245 from these readings. The amount of bacteriocin that inhibited the indicator strain by 246 50% was defined as 50% of the final  $OD_{590} \pm 0.05$  of the untreated control culture.

247

260

## 248 **Results**

249 The seven L. salivarius test strains included in this study were of either human or 250 porcine intestinal origin (Table 1), and were specifically selected by their ability to 251 produce bacteriocins which we propose constitutes an important probiotic trait (6, 31, 252 34). For comparative reasons, one isolate of human origin with a bacteriocin negative 253 phenotype, DPC6196, was also included in the genotypic analysis as it harbours 254 homologues of the abp118 bacteriocin structural genes (2). Due to the localisation of 255 the abp118 bacteriocin gene cluster on the megaplasmid of strain UCC118 (17), the 256 megaplasmid content of the test strains was first compared with that of UCC118. 257 258 Analysis of the megaplasmid content of bacteriocin-producing L. salivarius 259 isolates. Pulsed field gel electrophoresis (PFGE) of total genomic DNA confirmed

that the seven test strains were genetically distinct (data not shown), and the

- 261 megaplasmid content of each, as determined by S1 PFGE, is outlined in Table 2. The
- human derived isolates DPC6196 and DPC6488 were found to harbour megaplasmids

263 of 180 and 195 kb, respectively. Smaller plasmids were also identified in these strains, 264 two in DPC6488 of approximately 20 kb and 30 kb, and one in DPC6196 of 265 approximately 48 kb. The porcine derived isolates, DPC6005, DPC6502 and 7.3, each 266 contained megaplasmids of approximately 242 kb, similar to pMP118 (5), while 267 DPC6027 and DPC6189 contained larger plasmids of 320 kb and 360 kb, 268 respectively. Multiple megaplasmids were observed in L. salivarius DPC6189 and L. 269 salivarius 7.3 which both appear to harbour an additional linear megaplasmid of 270 approximately 195 kb. The linear nature of these megaplasmids was determined 271 when, under different switching conditions, PFGE analyses revealed that the 272 corresponding bands migrated to the same location on the gel regardless of whether 273 DNA from these isolates was digested with S1 nuclease or not (data not shown). 274 275 Genomic diversity of L. salivarius strains. Array comparative genomic hybridisation 276 (aCGH) analyses revealed considerable genomic diversity between strain UCC118, 277 the five Bac<sup>+</sup> (bacteriocin-producing) porcine isolates and the human-derived Bac<sup>+</sup> 278 strain DPC6488 and Bac<sup>-</sup> (bacteriocin negative, despite harbouring homologues of the 279 abp118 structural genes) strain DPC6196, consistent with results reported by Raftis et 280 al., (2011) (28). We particularly noted a high level of plasticity within the UCC118 281 megaplasmid-associated bacteriocin abp118-encoding locus. 282 283 Genetic diversity of the salivaricin bacteriocin locus in L. salivarius. The genetic 284 determinants responsible for the production of abp118 are comprised of the abp118  $\alpha$ 285 and  $\beta$  structural genes, accompanied by the genes encoding a cognate immunity 286 protein, a three-component regulatory system, an ABC-transporter and transport 287 accessory protein which are responsible for cleavage and secretion of the mature

288 active bacteriocin (13) (Fig. 1). Six additional ORFs are also present on the 10.7 kb 289 abp118 locus, three of which encode putative bacteriocin-like precursor peptides, two 290 of which have no homologues while one was identified as a pre-salivaricin B 291 homologue (4, 13). Each of the L. salivarius test isolates included in this study are 292 bacteriocinogenic, with the exception of DPC6196, and sequence analysis revealed 293 that the abp118 gene cluster shares greater than 90% sequence similarity with that of 294 salivaricin P (data not shown (13)). Despite this, CGH revealed considerable genetic 295 diversity with respect to the bacteriocin locus across the test strains (Fig. 2). The 296 porcine intestinal strain DPC6502 was particularly notable by virtue of lacking 297 abp118-related homologues, despite having an antimicrobial-producing phenotype 298 (Fig. 2, (23)). This observation eliminates the possibility that this strain produces 299 abp118, salivaricin P and/or salivaricin B. Although CGH data indicated that the 300 bacteriocin structural genes, the genes encoding the response regulator and the 301 transport system of abp118 were conserved in each of the remaining porcine derived 302 isolates, DPC6005, DPC6027, DPC6189 and 7.3, the genes encoding the immunity, 303 induction peptides and the histidine kinase component of the salivaricin regulatory 304 system of these isolates were divergent from their respective abp118 counterparts, as 305 illustrated in Fig. 2. Indeed, sequence analysis of the salivaricin P locus of the 306 prototype producing strain DPC6005 (data not shown) confirmed this. 307 Overall, the abp118 locus was most highly conserved in the human isolate 308 DPC6196 (Fig. 2). Although displaying a Bac<sup>-</sup> phenotype, this isolate harbours 309 homologues of  $abp118\alpha$  and  $abp118\beta$ , differing by just 1 and 2 nucleotides, 310 respectively (2). CGH analysis revealed divergence with respect to the *abpT* 311 (LSL\_1910) and *abpD* (LSL\_1909) homologues, required for secretion of the mature 312 active bacteriocin, in DPC6196 which is likely responsible for the Bac<sup>-</sup> phenotype of

313 this strain. Previously, PCR analysis using primers specific for the salivaricin P 314 structural genes indicated the presence of related genes in L. salivarius DPC6488 315 (23). However, the unique genetic variability pattern of this neonatal isolate instead 316 indicated that the genes corresponding to  $abp118\alpha$  (LSL 1917) and  $abp118\beta$ 317 (LSL\_1916) were divergent and absent, respectively (Fig. 2). In contrast, the genes 318 encoding the abp118 regulatory and transport systems were perfectly conserved in this 319 strain, relative to UCC118, as was the gene encoding pre-salivaricin B (LSL\_1921). 320 As this strain is of human origin, an important consideration in the selection of 321 commercial probiotics, the antimicrobial activity and corresponding genetic 322 determinants of this isolate were further investigated. 323 324 Analysis of the gene products encoded by the salivaricin locus of L. salivarius 325 **DPC6488.** A single primer pair successfully amplified the individual salivaricin loci 326 of L. salivarius UCC118, DPC6005 and DPC6488 in their entirety. Sequencing of the 327 corresponding amplicon of L. salivarius DPC6488 allowed for elucidation of the most 328 divergent region of the bacteriocin locus of this strain, which on the basis of CGH 329 data corresponds to the sequence between LSL\_1913 and LSL\_1924 of strain 330 UCC118 (Fig. 2), and confirmed the conservation of the regulatory and transport 331 systems of abp118 in this strain. In silico analysis of the sequence data resulted in the 332 identification of 20 ORFs (Fig. 1; Table 3), five of which, ORF3-5, 7 and 13, encoded 333 bacteriocin-like prepeptides. The putative products of ORF3 and ORF4 334 (correspondingly designated *slnT*3 and *slnT*4) were 98% and 97% identical to the 335 salivaricin B precursor peptide (LSL\_1921) and a bacteriocin-like prepeptide 336 (LSL\_1918) of UCC118, respectively. The deduced products of ORF5 and ORF7 also 337 resembled bacteriocin precursor peptides with double-glycine leader sequences, and

338 were designated *slnTa* and *slnTβ* (Fig. 3A). The first 57 nucleotides of *slnTa* closely 339 resemble those encoding the leader peptide of Abp118 $\alpha$  and Sln1, differing by two 340 nucleotides (and, in turn, one aa) (Fig. 3B). However, no significant homology was 341 observed between the corresponding propeptide (61 aa) and Abp118 $\alpha$  (45 aa). Indeed 342 a BLAST search revealed that this peptide instead shared 47% identity with ThmA, 343 one component of the two-peptide bacteriocin thermophilin13 produced by 344 Streptococcus thermophilus (20) (Fig. 3B). The bacteriocin-like precursor peptide 345 encoded by  $slnT\beta$  displayed similarity to acidocin LF221 A, one peptide of a putative 346 two-component bacteriocin produced by Lactobacillus gasseri (18) (Fig. 3B). Only 347 partial sequence of the complementary peptide of LF221 A (37 C-terminal aa) was 348 available for comparison with the propeptide encoded by  $slnT\alpha$ , with which it shared 349 30% similarity (Fig. 3B). While the leader sequence of SalT $\beta$  also shares 42% 350 similarity with that of Abp118α, no significant similarity was observed between the 351 leader sequence of SalT $\beta$  and that of Abp118 $\beta$  (Fig. 3B). Indeed, the SalT $\beta$  leader 352 sequence was most similar (59%) to that of the LafX peptide of lacticin F, produced 353 by Lactobacillus johnsonii (Fig. 3B, (14)), while, perhaps most notably, the 354 propeptide encoded by  $slnT\beta$  shares 43% similarity with ThmB, the complementary 355 peptide of ThmA (20). 356 Further *in silico* analysis of this cluster identified two ORFs encoding putative 357 immunity proteins downstream of the potential bacteriocin structural genes  $slnT\alpha$  and

358  $slnT\beta$  (Fig. 3A). The deduced protein sequences corresponding to slnT IM1 (ORF6)

and *slnT IM2* (ORF8) displayed 81% and 76% identity with Abp118 IM. Despite

being present in two copies, these LSL\_1915-homologues were not detected by CGH

361 due to the divergence in the nucleotide sequences of the genes in DPC6488 (72% and

362 69% identity, respectively, with *abpIM*). Three ORFs downstream of the putative

363 bacteriocin immunity gene, ORFs 9, 10 and 11, encode a putative induction peptide, a 364 sensory transduction histidine kinase and a response regulator, which shared 100%, 365 99% and 94% identity with their respective abp118 counterparts, thus confirming 366 conservation of the regulatory system of abp118 in strain DPC6488. The deduced 367 products of ORFs 12 and 14 did not share homology with any previously known 368 proteins. The 59 aa putative bacteriocin prepeptide encoded by ORF 13 (slnL) consists 369 of a double-glycine leader sequence of 18 aa and a mature peptide of 41 aa designated 370 salivaricin L, and shared greatest homology (59% identity) with a putative bacteriocin 371 precursor of the Streptococcus sp. C150 genome (accession no. EFX55741). This 372 putative bacteriocin precursor also shared very weak homology (38% and 37%) with 373 the precursor peptides of cerein 7B produced by Bacillus cereus Bc7 (25) and sakacin 374 Q produced by Lactobacillus sakei (21), respectively. Interestingly, both cerein 7B 375 and sakacin Q are single peptide class IId bacteriocins which are simultaneously 376 produced with cerein 7A and sakacin P by their respective producing strains (21, 25). 377 The deduced protein products of ORFs 15 and 16 share 99% and 97% identity with 378 those involved in the transport of abp118, AbpT and AbpD, respectively, thus 379 confirming conservation of the abp118 transport system in strain DPC6488. Three 380 ORFs downstream of the putative bacteriocin transport genes encoded proteins 381 homologous to the IS1223 family transposases. The protein products of overlapping 382 ORFs 17 and 18 shared 97% and 94% identity with the 61 C-terminal aa and 246 N-383 terminal aa of transposase ISLasa2b encoded by LSL\_1957, approximately 30 kb 384 downstream of the abp118 bacteriocin locus on pMP118. In addition, the deduced 385 product of ORF 19 shares 98% identity with transposase ISLasa1b encoded by 386 LSL\_1958 of *L. salivarius* UCC118, indicative of a possible recombination event. 387 The final ORF identified on the salivaricin locus of DPC6488 encoded a hypothetical

protein of 274 aa, the C- and N-terminus of which shared 97% and 87% identity with
the smaller hypotetical proteins encoded by LSL\_1907 (174 aa) and LSL\_1908 (76
aa) of the abp118 locus, respectively.

391

#### 392 Isolation and characterisation of the bacteriocin-like peptides produced by L.

393 salivarius DPC6488. The spectrum of activity of L. salivarius DPC6488, as

determined by well diffusion assay of the neutralised cell free supernatant (CFS), was

395 previously found to be limited to 12 closely related species of lactic acid bacteria

396 (including *Enterococcus*, *Lactobacillus* and *Streptococcus* sp.) of a total of 62

397 indicator strains investigated. This activity was lost upon protease treatment with

398 proteinase K,  $\alpha$ -chymotrypsin, trypsin or pepsin (23). Cross sensitivity assays using

the CFS of DPC6488 and of the abp118 and salivaricin P producers (UCC118 and

400 DPC6005, respectively) revealed that all 3 strains are immune to both abp118 and the

401 DPC6488-associated antimicrobials. Interestingly, however, both human isolates were

402 sensitive to the CFS of the salivaricin P producer DPC6005, despite the fact that their

403 respective immunity peptides share greater than 76% sequence identity.

404 Further investigations were performed to establish the nature of the 405 antimicrobials produced by DPC6488. Mass spectral analysis revealed that peptides of 406 mass corresponding to those of the mature products of slnT3,  $slnT\alpha$ ,  $slnT\beta$ , and slnL407 were present in the culture supernatant of DPC6488. To determine if these peptides 408 were responsible for the antimicrobial activity of the strain, the peptides were 409 separated and purified using cation exchange chromatography. MS data confirmed the 410 predicted molecular masses of the mature peptides encoded by  $slnT\alpha$ ,  $slnT\beta$ , and slnL, 411 5,655 Da, 5,269 Da, and 4,117 Da, respectively, in the individually eluted active

412 fractions. The individual SalT $\alpha$  and SalT $\beta$  peptides exhibited antimicrobial activity

413 against the sensitive indicator strain *L. delbrueckii ssp. bulgaricus* LMG 6901, which

414 was further enhanced when they were combined (Fig. 4). Thus  $slnT\alpha$  and  $slnT\beta$ 

415 appear to encode a two-peptide bacteriocin that most closely resembles thermophilin

416 13, which we designated salivaricin T.

MS analysis also established that the mature salivaricin B analogue encoded
by *salT3* is secreted by strain DPC6488 and identified the fraction containing a

419 peptide of corresponding mass (4,433 Da). However, this peptide did not inhibit the *L*.

420 *delbreueckii* ssp. *bulgaricus* indicator strain, nor did it display the anti-*Listeria* 

421 activity previously attributed to salivaricin B (4) (data not shown). As noted above, a

422 gene encoding a homologue of the salivaricin B precursor peptide is also located on

423 the abp118 gene cluster (LSL\_1921) but was designated non-functional in *L*.

424 salivarius UCC118 (5, 13). Indeed, CGH demonstrates that this gene is conserved in

425 six of the seven test strains employed in this study and DNA sequence and MS data

426 confirmed that the peptide encoded by *slnT3* of DPC6488 shares 100% identity with

427 salivaricin B. To investigate further, a synthetic analogue of mature salivaricin B was

428 generated based on the deduced aa sequence of *slnT3*. However, we were again

429 unable to detect antimicrobial activity from the synthetic peptide (data not shown).

430 Furthermore when either the synthetic analogue or the purified salivaricin B-

431 containing fraction were combined with those containing SalT $\alpha$  and SalT $\beta$ , or the

432 individual purified component peptides of salivaricin P, Sln1 and Sln2, salivaricin B

433 failed to enhance their antimicrobial activity against *L. delbrueckii ssp. bulgaricus* 

434 LMG 6901 or *Listeria innocua* (data not shown). These findings thus question the

435 previously reported anti-*Listeria* potency of this bacteriocin (4).

Interestingly, production of the fourth bacteriocin-like prepeptide encoded onthe salivaricin gene cluster of strain DPC6488, with similarity to one-peptide class IId

438 bacteriocins, was also confirmed by MALDI-TOF MS. While this peptide inhibited 439 three of the 12 indicator strains sensitive to salivaricin T, L. delbrueckii ssp. 440 bulgaricus, L. delbrueckii ssp. lactis and L. ruminis, synergistic activity was not 441 observed when combined with the salivaricin T component peptides SalT $\alpha$  and SalT $\beta$ . 442 Moreover, the purified salivaricin L peptide also exhibited anti-Listeria activity. A 443 synthetic analogue of the mature 41 aa peptide displaying similar activity to the 444 natural peptide was employed to determine the specific activity of the bacteriocin, 445 revealing an MIC<sub>50</sub> of 20 µM for both L. innocua DPC3572 and L. monocytogenes 446 NCTC 11994 (Fig. 5). 447 Given the inactivity of salivaricin B, and the antimicrobial activity of purified

449 *bulgaricus* LMG 6901 strain by *L. salivarius* DPC 6488 can be attributed to two novel

SalT $\alpha$ , SalT $\beta$  and salivaricin L, it is apparent that the inhibition of L. delbrueckii ssp.

450 bacteriocins encoded within a single gene cluster, which we designate salivaricin T,

451 corresponding to the two component peptide products encoded by  $slnT\alpha$  and  $slnT\beta$ ,

452 and salivaricin L, corresponding to the one-peptide class IId bacteriocin encoded by

453 salL. Furthermore, the regulatory and export systems of corresponding salivaricin

454 locus are analogous to those of abp118 and salivaricin P, thereby highlighting the

455 ability of these systems to recognise and produce various antimicrobial peptides.

456

448

#### 457 Discussion

458 CGH analysis revealed a high level of intraspecies diversity within the intestinal *L*.
459 *salivarius* isolates employed when compared to the genome of *L. salivarius* UCC118,
460 largely consistent with a recent survey of the genomic diversity of 33 *L. salivarius*

461 strains (28).

462 Bacteriocin production is a megaplasmid encoded feature of importance for 463 the probiotic functionality of L. salivarius UCC118 (5, 6) and CGH confirmed the 464 presence of *repA*-type megaplasmids in each of the test strains investigated in this 465 study. The megaplasmids of DPC6488 and DPC6196 were considerably smaller in 466 size than pMP118, notably, a 67 kb non-functional conjugation transfer locus present 467 in UCC118 (10) is absent in these strains. Indeed, this locus is also absent in 468 DPC6502 despite the fact that it harbours a megaplasmid similar in size to pMP118. 469 This indicates a considerable number of additional megaplasmid-encoded features in 470 DPC6502, which also is the case for DPC6027 and DPC6189 which harbour 471 megaplasmids considerably larger than pMP118. CGH analysis of the bacteriocin 472 locus in the porcine salivaricin P-producing test strain DPC6005 produced results 473 which were consistent with subsequent sequence analysis. This revealed that the 474 salivaricin P gene cluster was organised in a similar arrangement to the genetic locus 475 of abp118 (13, 27), with the most notable difference between the genes encoding the 476 induction peptides (60% identity). It was not surprising that the N-terminal region of 477 the corresponding sensory histidine kinase genes also varied slightly (93% identity). 478 Similar variations were previously observed between the regulatory operons of the *pln* 479 loci of Lactobacillus plantarum C11 and L. plantarum NC8 which are also otherwise 480 highly homologous (19). Indeed, similar levels of variability were observed with 481 respect to the bacteriocin locus of all of the salivaricin P-producing porcine isolates, 482 relative to UCC118, indicating that they may all be derived from a single ancestor. CGH also revealed that the bacteriocin-negative phenotype of the human isolate 483 484 DPC6196 is likely due to a genetic defect in the bacteriocin transport system in this 485 isolate.

486 Sequence analysis of the salivaricin locus of strain DPC6488 further validated 487 the CGH data. Although the genes involved in regulation and transport of abp118 488 were highly conserved in DPC6488, this strain harbours structural genes for the 489 production of two novel antimicrobials, a two-component class IIb bacteriocin 490 salivaricin T, and a one-peptide class IId bacteriocin salivaricin L. It is understandable 491 that strain DPC6488 was previously mistakenly characterised as a salivaricin P 492 producer as the primers designed to amplify the bacteriocin structural genes were 493 complementary to the sequence encoding the double-glycine leader of Sln1 and the 494 immunity genes of salivaricin P, which share considerable identity with the leader 495 sequence of SalTa and the respective immunity genes of the corresponding salivaricin 496 locus. Notably, the porcine isolate DPC6502, which was also previously characterised 497 as a salivaricin-P producer, lacked all abp118-related homologues and so the Bac<sup>+</sup> 498 phenotype of this strain will be the basis of further investigation. 499 The leader sequence of the SalT $\alpha$  precursor is almost identical to that of the 500 Abp118a and Sln1 prepeptides (95% identity, with just one conservative aa difference [I/V] in 19), which may be a requirement for the efficient processing and export of the 501 502 mature active peptide by the conserved ABC transporter. Indeed, while the mature 503 salivaricin T peptides most closely resembled the component peptides of thermophilin 504 13 (20), the proteins involved in providing immunity to, regulating production of and 505 transporting these peptides are analogous to those of abp118. However, it should be

506 noted that, at present, the thermophilin 13 gene cluster has not been reported. The 507 presence of two putative immunity genes downstream of each of the salivaricin T 508 structural genes is interesting and this fact, combined with the individual activity of 509 each of the component peptides, may indicate that salivaricin T evolved from two 510 synergistically acting one-peptide bacteriocins. Interestingly, the one-peptide class IId

511 bacteriocins cerein 7B and sakacin Q with which salivaricin L shares weak homology 512 are also simultaneously produced with cerein 7A and sakacin P by their respective 513 producing strains B. cereus and L. sakei (21, 25). As observed with salivaricin T and 514 salivaricin L, neither of these pairs of simultaneously produced bacteriocins exhibit 515 synergistic activity. Moreover, salivaricin L exhibited antagonistic activity to the 516 gastrointestinal pathogen L. monocytogenes, with an MIC<sub>50</sub> of 20  $\mu$ M. 517 L. salivarius DPC6488 was selectively isolated from the neonatal faecal 518 population as a consequence of its antimicrobial phenotype. Therefore, it is tempting 519 to suggest that salivaricin T and salivaricin L production may be an important 520 mechanism for host colonisation and prevalence of the producing strain allowing it to 521 outcompete closely-related populations within the intestine. Notwithstanding this 522 hypothesis, the apparent ability of L. salivarius to produce very differing bacteriocins 523 using the same cellular machinery suggests a hitherto unknown versatility with 524 respect to production of this dominant probiotic trait.

525

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663		

### 664 **Figure legends**

**Figure 1.** Comparative representation of the abp118 and salivaricin T gene clusters.

666 The bacteriocin structural and predicted immunity genes are indicated by the colours

black and grey with a black outline, respectively. The homologues of the putative

protein products encoded by the salivaricin T gene cluster are outlined in Table 3. The

669 most notable regions of similarity between the respective structural and immunity

670 genes are highlighted by the black dashed lines.

671



673 within *L. salivarius* test strains as determined by aCGH. Black, blue and yellow

674 regions represent absence, conservation or overrepresentation of CDS, respectively,

675 corresponding to the colour legend.

676

677 Figure 3. Nucleotide sequence and deduced peptide sequence of the salivaticin T 678 structural genes and the immunity genes of salivaricin locus of L. salivarius 679 DPC6488. (A) The 1087-bp DNA sequence shown encodes the component peptides 680 of salivaricin T produced by L. salivarius 6488 (SalTa and SalTB) together with their 681 leader sequences (underlined). The GG-processing sites are indicated by bold 682 triangles ( $\blacktriangle$ ). Two putative immunity genes are present downstream of *slnTa* and 683  $slnT\beta$ . Putative ribosome binding sites (RBS), start and stop codons are indicated in 684 bold, start codons are also underlined. An inverted repeat sequence (indicated by 685 arrows) typical of a Rho-independent terminator sequence was identified downstream 686 of  $slnT\beta$  also indicated in bold. (B) Alignment of salivaricin T component precursor 687 peptides with those of abp118, acidocin LF221 A, lactacin F, and thermophilin 13. 688 The leader sequences of the Abp118 $\alpha$  and SalT $\alpha$  prepeptides and SalT $\beta$  and LafX

689	prepeptides share 95%	and 59% identity,	respectively. The	e SalTα propeptide	displays
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- 690 greatest identity (46%) with ThmA and their respective complementary peptides,
- 691 SalT $\beta$  and ThmB share 33% identity. The SalT $\beta$  propertide shares greatest identity
- 692 (41%) with acidocin LF221 A, the partial available sequence of the LF221 A
- 693 complementary peptide shares 24% identity with the propeptide of SalTα.
- 694
- **Figure 4.** MALDI-TOF MS data for purified SalT $\alpha$  and SlaT $\beta$  and individual and
- 696 combined antimicrobial activity of SalTα and SlaTβ against *L. delbrueckii* subsp.
- 697 *bulgaricus* LMG6901.
- 698

**Figure 5.** Inhibitory effect of synthetic salivaricin L on the growth of indicator strains *Listeria innocua* DPC3572 (A), and *Listeria monocytogenes* NCTC 11994 (B) at concentrations of  $0 \ \mu M$  ( $\blacklozenge$ ),  $1.0 \ \mu M$  ( $\square$ ),  $5.0 \ \mu M$  ( $\blacktriangle$ ),  $10.0 \ \mu M$  ( $\circ$ ),  $15.0 \ \mu M$  ( $\times$ ), 20.0  $\mu M$  ( $\triangle$ ) and 50.0  $\mu M$  ( $\diamondsuit$ ). Error bars represent standard deviations based on triplicate data.

Source	Strain	Relevant features	Reference	
	Lactobacillus delbrueckii subsp. bulgaricus LMG 6901	Indicator strain	(2)	
	Listeria innocua DPC3572	Indicator strain	(2)	
Human	Lactobacillus salivarius UCC118	Abp118 producer	(13)	
	Lactobacillus salivarius DPC6196	Harbours bacteriocin structural genes	(2)	
	Lactobacillus salivarius DPC6488	Produces salivaricin T and salivaricin L	(23)	
Porcine	Lactobacillus salivarius DPC6502	Bacteriocin producer	(23)	
	Lactobacillus salivarius 7.3	Salivaricin P producer	(2)	
	Lactobacillus salivarius DPC6189	Salivaricin P producer	(2)	
	Lactobacillus salivarius DPC6027	Salivaricin P producer	(2)	
	Lactobacillus salivarius DPC6005	Salivaricin P producer	(2)	

**Table 1.** Bacterial strains used in this study

Strain	UCC118	DPC6196	DPC6488	DPC6502	DPC6027	DPC6189	7.3	DPC6005
Approximate length of circular megaplasmids (kb)	242	180	195	242	320	360	195	242
Additional plasmid content								
Approximate length of linear megaplasmids (kb)	-	-	-	-	-	195	195	-
Approximate length of smaller plasmids (kb)	44, 20	48	30, 20	-	-	-	-	-

**Table 2.** Megaplasmid content of intestinal *L. salivarius* strains

708	Table 3. Homologues of the deduced proteins encoded by the salivaricin T bacteriocin locus of L. salivarius DPC6488

ORF (gene)	Size (aa)	Function	Homologue	Identity (%) <sup>a</sup>	Reference	
ORF 1 (slnT1)	65	Conserved hypothetical protein	Conserved hypothetical protein of L. salivarius DSM20555	95 [62/65]	EEJ73426 <sup>b</sup>	
ORF 2 (slnT2)	87	Conserved hypothetical protein	Conserved hypothetical protein of L. salivarius DSM20555	100 [87/87]	EEJ73427 <sup>b</sup>	
ORF 3 (slnT3)	57	Bacteriocin-like prepeptide	Salivaricin B prepeptide	98 [56/57]	(4)	
ORF 4 (slnT4)	89	Bacteriocin-like prepeptide	LSL_1918 of L. salivarius UCC118	98 [83/85]	(5)	
ORF 5 ( $slnT\alpha$ )	80	Salivaricin T prepeptide SalT alpha	ThmA, amphipathic pore-forming peptide precursor of S. thermophilus	45 [35/77]	(20)	
ORF 6 (slnT IM1)	59	Putative bacteriocin immunity protein	Abp118 IM (LSL_1915) of L. salivarius UCC118	82 [45/55]	(13)	
ORF 7 $(slnT\beta)$	75	Salivaricin T prepeptide SalT beta	Acidocin LF221A produced by L. gasseri	46 [31/67]	(18)	
ORF 8 (slnT IM2)	54	Putative bacteriocin immunity protein	Abp118 IM (LSL_1915) of L. salivarius UCC118	76 [38/50]	(13)	
ORF 9 (slnT IP)	38	Putative induction peptide	Abp118 IP (LSL_1914) of L. salivarius UCC118	100 [38/38]	(13)	
ORF 10 $(slnT K)$	429	Sensory transduction histidine kinase	AbpK of L. salivarius UCC118	99 [424/429]	(13)	
ORF 11 ( <i>slnT R</i> )	265	Response regulator	Salivaricin response regulator of strain CECT5713/ (AbpR 94%)	95 [249/263]	ADJ79880 <sup>b</sup>	
ORF12	41	Hypothetical protein	no homologues			
ORF 13 (Sln L)	59	Bacteriocin-like prepeptide	Putative bacteriocin of Streptococcus sp. C105	60 [35/60]	EFX55741 <sup>b</sup>	
ORF 14	74	Hypothetical protein	no homologues			
ORF 15 $(slnTT)$	719	Salivaricin T ABC-transporter protein	AbpT (LSL_1910) of L. salivarius UCC118	99 [709/719]	(13)	
ORF 16 ( <i>slnT D</i> )	384	Salivaricin T export accessory protein	AbpD (LSL_1909) of L. salivarius UCC118	97 [370/384]	(13)	
ORF 17	63	Truncated transposase	Transposase ISLasa2b, IS1223 family (LSL_1957)	97 [59/61]	(5)	
ORF 18	246	Truncated transposase	Transposase ISLasa2b, IS1223 family (LSL_1957)	94 [231/246]	(5)	
ORF 19	172	Transposase	IS1223 family transposase (LSL_1958, LSL_0049) of UCC118	98 [169/172]	(5)	
ORF 20	273	Hypothetical protein	Conserved hypothetical protein of L. salivarius CECT5713	95 [257/270]	ADJ79877 <sup>b</sup>	

<sup>a</sup>Percentage identity was determined using BLAST

709 <sup>b</sup>Accession number of sequence directly submitted to EMBL Database