

1 Cloning and functional expression of a food-grade circular bacteriocin, plantacyclin B21AG,
2 in probiotic *Lactobacillus plantarum* WCFS1

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24 **Abstract**

25 There is an increasing consumer demand for minimally processed, preservative free and
26 microbiologically safe food. These factors, combined with risks of antibiotic resistance, have
27 led to interest in bacteriocins produced by lactic acid bacteria (LAB) as natural food
28 preservatives and as protein therapeutics. We previously reported the discovery of
29 plantacyclin B21AG, a novel circular bacteriocin produced by *Lactobacillus plantarum* B21.
30 Here, we describe the cloning and functional expression of the bacteriocin gene cluster in the
31 probiotic *Lactobacillus plantarum* WCFS1. Genome sequencing demonstrated that the
32 bacteriocin is encoded on a 20 kb native plasmid, designated as pB21AG01. Seven open
33 reading frames (ORFs) putatively involved in bacteriocin production, secretion and immunity
34 were cloned into an *E. coli/Lactobacillus* shuttle vector, pTRKH2. The resulting plasmid,
35 pCycB21, was transformed into *L. plantarum* WCFS1. The cell free supernatants (CFS) of
36 both B21 and WCFS1 (pCycB21) showed an antimicrobial activity of 800 AU/mL when
37 tested against the WCFS1 (pTRKH2) indicator strain, indicating functional expression of
38 plantacyclin B21AG. Real-time PCR analysis revealed that the relative copy number of
39 pB21AG01 was 7.60 ± 0.79 in *L. plantarum* B21 whilst pCycB21 and pTRKH2 was $0.51 \pm$
40 0.05 and 25.19 ± 2.68 copies, respectively in WCFS1. This indicates that the bacteriocin gene
41 cluster is located on a highly stable, low copy number plasmid pB21AG01 in *L. plantarum*
42 B21. Inclusion of the native promoter for the bacteriocin operon from pB21AG01 may result
43 in similar inhibitory zones observed in both wild type and recombinant hosts despite the low
44 copy number of pCycB21.

45

46 **Introduction**

47 Bacteriocins are ribosomally synthesised, extracellularly released peptides or peptide
48 complexes that possess antibacterial activity against species usually closely related to the
49 producer strains or a wider range of microorganisms [1,2]. Interestingly, the bacteriocins
50 produced by gram-positive bacteria seem to exhibit broader spectrum activity compared to
51 the gram-negative bacteria [3]. Among the gram-positive bacteria, bacteriocins produced by
52 the food-grade lactic acid bacteria (LAB) have attracted considerable interest because they
53 are generally regarded as safe (GRAS). Being proteins they can be easily degraded by
54 proteases in the mammalian gastrointestinal tract, making them safe for human consumption
55 and minimizing the risk of developing resistant bacteria [4,5]. They have been widely used as
56 natural food preservatives for controlling food-borne and food-spoilage bacteria without
57 affecting sensory qualities. They also have huge potential in veterinary applications and as
58 next-generation antibiotics against multi-drug resistant (MDR) pathogens [5-7]. One of the
59 advantages of bacteriocins over conventional antibiotics is that they are directly gene
60 encoded, making bioengineering feasible to enhance their productivity or specificity towards
61 target pathogens [5,8].

62

63 The classification of bacteriocins produced by gram-positive bacteria has been constantly
64 revised due to the extensive research performed over the last two decade [9-11]. Here we use
65 the classification proposed by Acedo et al. [12]. Class I contains modified peptides including
66 lantibiotics, lipolanthines, linear azol(in)e-containing bacteriocins, thiopeptides,
67 bottromycins, sactibiotics, lasso peptides, glycocins and circular bacteriocins. Class II are
68 unmodified peptides such as YGNG-motif containing bacteriocins, two-peptide bacteriocins,
69 leaderless bacteriocins and other linear bacteriocins. Class III are large heat labile
70 bacteriocins such as bacteriolysins, non-lytic large bacteriocins and tailocins. Of these, the
71 circular bacteriocins have gained considerable attention as they generally exhibit broad

72 antimicrobial activity. They are synthesised as linear pre-peptides where the leader peptides
73 of variable sizes (3 – 35 amino acids) are cleaved off during maturation, forming 58 – 70
74 amino acid peptides which are covalently linked by a largely unknown cyclisation
75 mechanism [13,14]. The circular structures appear to enhance their pH and thermal stability
76 as well as protease resistance. These properties make them a preferred candidate for potential
77 industrial applications compared to the other classes of bacteriocins [13,15].

78

79 Among the LAB, bacteriocins produced by *Lactobacillus*, in particular *Lactobacillus*
80 *plantarum* have been widely studied for several reasons. *L. plantarum* is a versatile species
81 that is widely found in a variety of sources, including meat, dairy, fish, fruit and vegetables
82 [16]. It is also one of the natural inhabitants of the human gastrointestinal tract (GIT) where
83 its ability to survive passage through the GIT makes it an attractive vector for vaccine
84 delivery [17,18]. The availability of the complete genome sequence of *L. plantarum* WCFS1
85 and genome mining tools have facilitated the characterisation of the genetic organisation of
86 the plantacyclin (*pln*) loci from this species [19]. Hitherto, several other class II linear two-
87 peptide bacteriocins produced by *L. plantarum* strains have been described. For example,
88 plantaricin C-19 produced by *L. plantarum* C-19, isolated from fermented cucumber, and
89 plantaricin NA produced by *L. plantarum*, isolated from ‘ugba’, an African fermented oil-
90 bean seed showed strong antimicrobial activity against the food-borne pathogen, *Listeria*
91 *monocytogenes* [20,21]. Bacteriocin AMA-K produced by *L. plantarum* AMA-K, isolated
92 from fermented milk exhibited strong adsorption to cells of *L. monocytogenes*, *L. ivanovii*
93 subsp. *ivanovii* and *L. innocua* [22]. Plantaricin ST8KF produced by *L. plantarum* ST8KF,
94 isolated from kefir, demonstrated antimicrobial activity against *L. casei*, *L. salivarius*, *L.*
95 *curvatus*, *Enterococcus mundtii* and *L. innocua* [23,24]. In contrast, only one circular
96 bacteriocin, plantaricyclin A produced by *L. plantarum* NI326 has been reported to date.

97 Similarly, this circular bacteriocin is active against beverage-spoilage bacterium
98 *Alicyclobacillus acidoterrestris* [25]. These antimicrobial peptides appear to have great
99 potential in food preservation, particularly in controlling food-borne pathogens. Discovery of
100 more circular bacteriocins is highly favourable over linear peptides due to their superior
101 stability against various stresses [26].

102

103 In recent years, research on bacteriocins has progressed from producing the inhibitory
104 compounds in native systems to heterologous production in diverse producer organisms
105 which have the potential to be employed as starters, protectors and/or probiotics [27]. Several
106 strategies for heterologous expression of bacteriocins have been investigated either for
107 overproduction of the bacteriocin or structure-function studies [7,27-29]. We previously
108 reported the discovery of plantacyclin B21AG, a food-grade circular bacteriocin produced by
109 *Lactobacillus plantarum* B21 [30,31]. It is shown to be active against food-borne pathogens
110 including *Clostridium perfringens* and *Listeria monocytogenes*; food spoilage bacteria such
111 as *L. arabinosus*; as well as other LAB including *L. plantarum*, *L. brevis* and *Lactococcus*
112 *lactis* [30-32]. This study aimed to transfer the production of the broad antimicrobial
113 spectrum of plantacyclin B21AG to a probiotic strain, *L. plantarum* WCFS1 [33]. We
114 demonstrated that the bacteriocin gene cluster can be recombinantly expressed in *L.*
115 *plantarum* WCFS1 at a level comparable to the native producer *L. plantarum* B21. The
116 mobilization of the plantacyclin B21AG operon into the probiotic, *L. plantarum* WCFS1,
117 enhances the antimicrobial activity spectrum of the strain, potentially making it more useful
118 for use in the food industry and for clinical applications.

119

120 **Materials and methods**

121 **Bacterial strains, plasmids and culture conditions**

122 Bacterial strains and plasmids used in this study are listed in Table 1. All *Lactobacillus*
123 strains were cultured statically in deMan, Rogosa and Sharpe (MRS) broth (Becton,
124 Dickinson and Company, USA) at 37 °C under aerobic conditions. *Escherichia coli* strains
125 were grown in Luria Bertani (LB) medium (Becton, Dickinson and Company, USA) at 37 °C
126 with continuous agitation at 250 rpm. For selection, medium were supplemented with 100
127 µg/mL of ampicillin and/or 150 µg/mL of erythromycin for *E. coli* and 15 µg/mL of
128 erythromycin for *Lactobacillus*.

129

Table 1. Bacterial strains, plasmids and primers.

Strain/plasmid	Characteristics ^a	Reference/Source
Strains		
<i>Lactobacillus plantarum</i>		
B21	Wild-type strain; plantacyclin B21AG producer	[34]
WCFS1	Wild-type strain obtained from NIZO ^b	[19]
WCFS1 (pCycB21)	<i>L. plantarum</i> type strain transformed with pCycB21; Em ^r	This study
WCFS1 (pTRKH2)	Indicator strain; Bac ⁻ ; sensitive to plantacyclin B21AG	[35,36]
<i>Escherichia coli</i>		
JM110 (pTRKH2)	<i>E. coli</i> strain containing pTRKH2; Amp ^r Em ^r	[37] NEB
DH5α	Host strain; <i>fhuA2 Δ(argF-lacZ)U169 phoA glnV44 Φ80 Δ(lacZ)M15 gyrA96 recA1 relA1 endA1 thi-1 hsdR17</i>	
Plasmids		
pB21AG01	20.4 kb; native plasmid in <i>L. plantarum</i> B21; contains seven genes necessary for the production, immunity, export and processing of plantacyclin B21AG	This study
pTRKH2	6.7 kb; <i>E. coli/L. plantarum</i> shuttle vector; Amp ^r Em ^r	[37] This study
pCycB21	10.1 kb; contains the full plantacyclin operon cloned into the <i>Bam</i> HI and <i>Sac</i> I sites of pTRKH2; Em ^r	
Primers		
B21AG_F	CTGCAGGGATCCGTTCAACCTCCTTTCTGAC	
B21AG_R	GGCCGGGAGCTCACAGGTTCTTAGAATACTG	

130

131 ^a Em^r and Amp^r, erythromycin and ampicillin resistant, respectively; bac⁻, bacteriocin non-producing132 ^b NIZO food research, The Netherlands

133 **Sequence determination and genetic analysis of pB21AG01**

134 The full genome sequence of *L. plantarum* B21, including the 20.4 kb plasmid, pB21AG01,
135 was sequenced at the Beijing Genomics Institute (BGI) using the Illumina HiSeq 2000
136 platform (Illumina, USA) and assembled with SOAPdenovo software [38]. The plasmid was
137 annotated using RAST [39]. The resulting open reading frames (ORFs) were confirmed using
138 blastp against the NCBI non-redundant protein database [40]. The obtained putative protein
139 sequences were searched for conserved domains using the NCBI Conserved Domain
140 Database (CDD) site [41] and also examined for transmembrane domains using the TMHMM
141 transmembrane prediction algorithm at <http://www.cbs.dtu.dk> [42].

142

143 **DNA manipulations, plasmid constructions and *E. coli* transformations**

144 Total gDNA from *E. coli* and *Lactobacillus* was isolated using the GeneElute™ Bacterial
145 Genomic DNA Kit (Sigma-Aldrich, USA) as described in the user manual. Plasmids from *E.*
146 *coli* were extracted using the ISOLATE II Plasmid Mini kit (Bioline, Australia) according to
147 manufacturer's instruction. Plasmids from *Lactobacillus* were prepared using QIAGEN®
148 Plasmid Midi Kit (Qiagen, Germany) following supplier's direction with a few modifications
149 to cell wall lysis. One hundred millilitre of overnight culture was harvested by centrifugation
150 and washed in 20 mL STE buffer (6.7 % sucrose; 50 mM Tris-HCl, pH 8.0; 1 mM EDTA)
151 [43] to remove and neutralise acids produced during cell growth. The bacterial pellet was
152 then resuspended in 4 mL STE buffer containing 10 mg/mL lysozyme and incubated at 37 °C
153 for 1 hour.

154

155 For the construction of pCycB21, the full plantacyclin B21AG operon with its native
156 promoter was amplified from *L. plantarum* B21 plasmids using OneTaq® 2X Master Mix
157 with Standard Buffer (NEB, USA) in a T100™ Thermal Cycler (Bio-Rad, USA). PCR, using

158 the primers indicated in Table 1, was performed as follows: initial denaturation for 30 s at 94
159 °C; followed by 30 cycles of denaturation for 30 s at 94 °C, annealing for 30 s at 55 °C and
160 extension for 4 min at 68 °C; and a final extension for 10 min at 68 °C. The amplified
161 product was purified using the ISOLATE II PCR and Gel Kit (Bioline, Australia) and cloned
162 into the *Bam*HI and *Sac*I sites of the *E. coli*/*Lactobacillus* shuttle vector, pTRKH2. The
163 construct was transformed into *E. coli* DH5 α (NEB, USA) according to manufacturer's
164 protocol in order to obtain sufficient amount of plasmid DNA for subsequent transformation
165 into LAB. The recombinant plasmid is confirmed by PCR, double restriction enzyme
166 digestion and DNA sequencing.

167

168 **Electroporation of LAB**

169 Electroporation of LAB was performed as described by Mason et al. [44] with a few
170 modifications. Briefly, 8 mL of overnight LAB cultures were diluted into 40 mL of fresh pre-
171 warmed MRS broth containing 2 % glycine. The diluted culture was incubated for 1.5 hr at
172 37 °C. The cells were pelleted by centrifugation at 4000 x g for 2 min at 4 °C and washed
173 with 40 mL of ice-cold Milli-Q water. Cells were then resuspended in 40 mL of ice-cold 50
174 mM EDTA and incubated on ice for 5 min. Centrifugation was repeated followed by washing
175 the cells in 40 mL of ice-cold 0.3 M sucrose. The cells were resuspended in 200 μ L of 0.6 M
176 sucrose. Finally, 3 μ g of DNA in 50 μ L of sterile Milli-Q water was added to 50 μ L of
177 freshly prepared competent cells and transferred into a pre-chilled electrocuvette with a 0.2-
178 cm electrode gap (Cell Projects, UK). The cell suspension containing plasmid DNA was
179 electroporated using a Gene Pulser electroporator (Bio-Rad, USA) with the following
180 parameters: 1.5 kV, 200 Ω parallel resistance and 25 μ F capacitance. The cells were
181 transferred immediately after electroporation into 1.3 mL of pre-warmed MRS broth and
182 incubated for 3 hrs at 37 °C. Two hundred microliters of the cells were plated onto MRS agar

183 containing erythromycin and incubated for 2 days at 37 °C. Recombinant plasmids were
184 confirmed by PCR and double restriction enzyme digestion.

185

186 **Antimicrobial activity assay**

187 The antimicrobial activity of the bacteriocin produced by LAB was evaluated using the well
188 diffusion agar (WDA) method [45]. Briefly, cell free supernatants (CFS) of *L. plantarum* B21
189 and WCFS1 (pCycB21) were harvested from 15 mL overnight LAB culture by centrifugation
190 at 4,000 x g for 20 min at 4 °C. The CFS was then concentrated 15-fold using an Amicon®
191 Ultra-15 Centrifugal Filter Devices (Merck Millipore, Germany) and stored at 4 °C until
192 used. To evaluate the antimicrobial activity of plantacyclin B21AG, MRS agar plates
193 supplemented with 10 µg/mL of erythromycin were seeded with 10⁶ cfu/mL of *L. plantarum*
194 WCFS1 (pTRKH2), used as the indicator strain. Wells were made in the agar using a sterile
195 8-mm cork borer. One hundred microliters of the 2-fold serial diluted CFS was then loaded
196 into the wells and the plates were incubated at 30 °C for 16 – 18 h. Antimicrobial activity was
197 expressed as arbitrary unit (AU/mL) using the following equation, $a^b \times 100$, where “a” is the
198 dilution factor, “b” is the last dilution showing an inhibition zone of at least 2 mm in diameter
199 [46].

200

201 **Extraction of Plantacyclin B21AG with 1-butanol**

202 Plantacyclin B21AG was purified from *L. plantarum* B21 and *L. plantarum* WCFS1
203 (pCycB21) using 1-butanol as described by Abo-Amer [47] with the following modifications:
204 the concentrated CFS was mixed with ½ volume of water-saturated butanol for 20 s. The
205 mixture was incubated at room temperature for 10 min to allow phase separation before
206 centrifugation at 10,000 x g for 10 min. The butanol phase was transferred to a clean 1.5 mL
207 tube whilst the aqueous phase was subjected an additional butanol extraction. The two

208 butanol fractions containing plantacyclin B21AG were combined and the solvent was
209 removed using a freeze dryer (FDU-8612, Operon Co. Ltd, Korea). The lyophilised protein
210 was dissolved in 20 mM sodium phosphate buffer (pH 6.0).

211

212 **Mass spectrometry analysis**

213 The protein was subjected to matrix-assisted laser desorption/ionization time-of-flight mass
214 spectrometry (MALDI-TOF-MS) analyses as described by Vater et al. [48]. The MALDI-
215 TOF mass spectra were recorded using an Autoflex Speed MALDI-TOF instrument (Bruker,
216 Germany) containing a 355 nm Smartbeam II laser for desorption and ionization. 10 mg of α -
217 cyano-4-hydroxycinnamic acid dissolved in 70 % acetonitrile (ACN) containing 0.1 % (v/v)
218 trifluoroacetic acid (TFA) was used as matrix solution. Five microliters of bacteriocin
219 samples were mixed with equal volume of matrix solution and 1 μ L of the mixture was
220 spotted onto the target, air dried and measured.

221

222 **Plasmid copy number determination by real-time PCR**

223 The copy number of the native (pB21AG01) and recombinant (pCycB21) plasmids were
224 determined using real-time PCR according to Škulj et al. [49]. A 5-fold serial dilution of total
225 DNA extracted from *L. plantarum* B21 was used for the standard curves (final 1 ng/ μ L to
226 0.0016 ng/ μ L). Real-time PCR reactions were performed in 12 μ L mixtures containing 1 x
227 SensiFAST SYBR No-ROX mix (Bioline, Australia), 400 nM of each forward and reverse
228 primer (Table 3) and 1 μ L of DNA. The alanine racemase gene (*alr*), a single copy,
229 chromosomal gene from *L. plantarum* WCFS1, was selected as the reference gene
230 (GeneBank Accession No. AL935253) whilst the bacteriocin structural (*orf19*) was chosen as
231 the target for detection of the recombinant plasmid pCycB21. The replication (*rep*) gene was
232 used as the target to detect pTRKH2 in WCFS1. Separate reactions were prepared for the

233 detection of chromosomal and plasmid amplicons. All reactions were performed in duplicate
234 using the Rotor-Gene™ Q (Qiagen, Germany). Thermocycling conditions were: initial
235 denaturation for 3 min at 95 °C, followed by 40 cycles of 5 s at 95 °C, 10 s at 55 °C and 20 s
236 at 72 °C. Fluorescence signal was acquired at the end of each 72 °C step.

237

238 **Table 2. Primers used for plasmid copy number (PCN) detection with real-time PCR.**

Target gene	Location	Name	Sequence 5' → 3'	Amplicon size
<i>alr</i>	Chromosome	<i>alrF</i>	TGGGACGAATCGGGTTTCAG	208 bp
		<i>alrR</i>	GACACGTGGACATAGCGTGG	
<i>orf19</i>	pB21AG01	<i>orf19F</i>	CTGCAGCCTGCAGGGTTCAACCTCCTTTCTGAC	277 bp
		<i>orf19R</i>	GGTGGTCCTGCAGGCCTAACCTGCTACGATATGC	
<i>rep</i>	pTRKH2	<i>repF</i>	CGCTCAATCACTACCAAG	102 bp
		<i>repR</i>	CTCGGAAGTCAGAACAAC	

240

241 The slope of the relative standard curve with a condition that $r^2 > 0.99$ was used to calculate
242 the amplification efficiency (E) using equation 1.

243
$$E = 10^{(-1/\text{slope})}$$

244
$$E(\%) = (10^{(-1/\text{slope})} - 1) \times 100 \quad (1)$$

245

246 The plasmid copy number (PCN) was calculated based on equation 2 using efficiency (E) and
247 Ct values for both chromosomal (c) and plasmid (p) amplicons.

248
$$\text{PCN} = (\text{Ec})^{\text{Ctc}} / (\text{Ep})^{\text{Ctp}} \quad (2)$$

249

250 Results

251 Sequence analysis of pB21AG01

252 Sequence analysis revealed that *L. plantarum* B21 (GenBank Accession No. CP010528)
253 harboured at least two cryptic plasmids, designated as pB21AG01 (GenBank Accession No.
254 CP025732) and pB21AG02 (GenBank Accession No. CP025733). For the purpose of this
255 study, we focused our analysis on pB21AG01 as it was found to encode the genes responsible
256 for the production of a circular bacteriocin. Plasmid profile analysis revealed that pB21AG01
257 is a 20,429 bp circular DNA molecule with a GC content of 37.3 %. A total of 26 open
258 reading frames (ORF) were identified (Table 3). 14 ORFs of the pB21AG01 were
259 homologous to proteins with known or predicted functions whilst the remaining 12 ORFs
260 were either homologous to hypothetical proteins lacking functional predicts or had no
261 significant homology with any protein sequences in the GenBank databases.

262

263

264

Table 3. Putative genes and their proposed function deduced from the amino acid sequences of pB21AG.

Gene	Codon		No. of amino acids	Best homolog, GenBank Accession No. [organism]	% Identity (No. of amino acids overlapping)	Proposed function of gene product
	Start	Stop ^a				
<i>orf1</i>	2150	156C	664	Hypothetical protein, WP_057741928.1 [<i>Lactobacillus capillatus</i>]	97 (643)	Hypothetical protein
<i>orf2</i>	2808	2143C	221	DNA replication and relaxation protein, WP_057741930.1 [<i>Lactobacillus capillatus</i>]	93 (206)	Plasmid replication and relaxation
<i>orf3</i>	3590	3477C	37	No significant similarity		Hypothetical protein
<i>orf4</i>	4239	4030C	69	Hypothetical protein, WP_053266991.1 [<i>Lactobacillus plantarum</i>]	92 (55)	Hypothetical protein
<i>orf5</i>	5585	4599C	328	Hypothetical protein LVISKB P8-0006, BAN08211.1 [<i>Lactobacillus brevis</i> KB290]	92 (301)	Hypothetical protein
<i>orf6</i>	5748	5885	45	No significant similarity		Hypothetical protein
<i>orf7</i>	6592	5909C	227	Helix-Turn-Helix DNA binding domain of transcription regulators from the MerR superfamily, WP_057741904.1 [<i>Lactobacillus capillatus</i>]	85 (193)	Transcriptional regulator
<i>orf8</i>	7581	6667C	304	Hypothetical protein, WP_057741908.1 [<i>Lactobacillus capillatus</i>]	79 (246)	Hypothetical protein
<i>orf9</i>	8205	7597C	202	Hypothetical protein, WP_057741914.1 [<i>Lactobacillus capillatus</i>]	94 (189)	Hypothetical protein
<i>orf10</i>	9312	8230C	360	Cell wall hydrolase, WP_057741916.1 [<i>Lactobacillus</i>]	98 (352)	Hydrolysis of beta-1,4-

				<i>capillatus</i>]			linked polysaccharides
<i>orf11</i>	11112	9313C	599	Domain of unknown function DUF20, WP_057741917.1 [<i>Lactobacillus capillatus</i>]	90 (558)		Hypothetical protein
<i>orf12</i>	11484	11125C	119	Hypothetical protein, WP_053266985.1 [<i>Lactobacillus plantarum</i>]	93 (111)		Hypothetical protein
<i>orf13</i>	11657	11481C	58	Hypothetical protein FC81_GL002105, KRL03443.1 [<i>Lactobacillus capillatus</i>]	90 (52)		Hypothetical protein
<i>orf14</i>	13984	11657C	775	AAA-like domain containing a P-loop motif, KRL03444.1 [<i>Lactobacillus capillatus</i> DSM_19910]	99 (768)		Conjugative transfer
<i>orf15</i>	14588	14031C	185	TcpE family, WP_003688369.1 [<i>Lactobacillus mali</i>]	93 (172)		Conjugative transfer
<i>orf16</i>	14933	14601C	110	Hypothetical protein, WP_053266999.1 [<i>Lactobacillus</i>]	95 (104)		Hypothetical protein
<i>orf17</i>	15908	14949C	319	Conjugative transposon protein TcpC, WP_003688364.1 [<i>Lactobacillus mali</i>]	91 (290)		Conjugative transfer
<i>orf18</i>	16272	15979C	97	Hypothetical protein, WP_053266998.1 [<i>Lactobacillus</i>]	95 (92)		Hypothetical protein
<i>orf19</i>	16521	16796	91	Plantaricyclin A precursor, PlcA, PCL98053.1 [<i>Lactobacillus plantarum</i>]	88 (67)		Bacteriocin production
<i>orf20</i>	16888	17361	157	Plantaricyclin A immunity protein, PlcD, PCL98052.1 [<i>Lactobacillus plantarum</i>]	94 (147)		Involved in immunity to Plantaricyclin A
<i>orf21</i>	17364	17528	54	Plantaricyclin A immunity protein, PlcI, PCL98051.1 [<i>Lactobacillus plantarum</i>]	89 (48)		Involved in immunity to Plantaricyclin A

<i>orf22</i>	17548	18231	227	ABC transporter ATP-binding protein, PlcT, PCL98050.1 [<i>Lactobacillus plantarum</i>]	95 (215)	Transport
<i>orf23</i>	18234	18878	214	ABC-2 transporter permease, PlcE, PCL98049.1 [<i>Lactobacillus plantarum</i>]	94 (202)	Transport
<i>orf24</i>	18881	19402	173	Plantaricyclin A related protein, PlcB, PCL98048.1 [<i>Lactobacillus plantarum</i>]	90 (155)	Involved in Plantaricyclin A production
<i>orf25</i>	19567	19397C	56	Plantaricyclin A related protein, PlcC, PCL98047.1 [<i>Lactobacillus plantarum</i>]	95 (53)	Involved in Plantaricyclin A production
<i>orf26</i>	20229	19696C	177	Transposase DDE domain, CCB82565.1 [<i>Lactobacillus pentosus</i> MP-10]	100 (177)	DNA transposition

266 ^a C: complementary sequence

267 Seven ORFs were predicted to encode genes putatively responsible for the production,
268 immunity and transport of plantacyclin B21AG (*orf19* – *orf25*) [30]. *Orf19* showed 88 %
269 identity to Plantaricyclin A precursor (PlcA), a circular bacteriocin produced by *L. plantarum*
270 (PCL98053.1), presumably the structural gene responsible for the production of plantacyclin
271 B21AG. It encodes 91 amino acids consisting of a 33 amino acid leader peptide and a 58
272 amino acid bacteriocin mature peptide, which is 86 % identical to *plcA*, the structural gene of
273 Plantaricyclin A [25]. *Orf20* and *orf21* are 94 % and 89 % identical to *plcD* (PCL98052.1)
274 and *plcI* (PCL98051.1), putatively involved in immunity to plantaricyclin A. *Orf22* and *orf23*
275 are possible transporters of the plantacyclin B21AG. *Orf22* showed 95 % identity to *plcT*, a
276 gene encoded for ABC transporter ATP-binding protein in *L. plantarum* (PCL98050.1) whilst
277 *orf23* is 94 % identical to *plcE*, an ABC-2 transporter permease encoding gene in *L.*
278 *plantarum* (PCL98049.1), respectively. *Orf24* and *orf25* are predicted as plantaricyclin A-
279 related proteins, putatively involved in the production of the circular bacteriocin. Amino acid
280 sequence comparison with gassericin A revealed that *orf20*, *orf24* and *orf25* may be
281 membrane associated proteins, but their roles in bacteriocin production/immunity remain
282 unknown [50]. Transmembrane analysis using TMHMM revealed the presence of 2
283 transmembrane domains in *orf19*, *orf21* and *orf25*. *Orf20* and *orf23* and *orf24* contain 4, 6
284 and 5 transmembrane domains, respectively. This result is consistent with the properties of
285 the proteins involved in gassericin A production. No transmembrane domain was predicted
286 for *orf22*.

287

288 Although no *rep* gene or direct repeats were found in pB21AG01, *orf2* showed 93 % identity
289 to a protein in *L. capillatus* (WP_057741930.1) which is essential for plasmid replication and
290 relaxation. In addition, *orf7* is predicted to be a Helix-Turn-Helix DNA binding domain of
291 transcription regulators from the MerR superfamily, accession number WP_057741904.1

292 [*Lactobacillus capillatus*] [41]. They have been shown to mediate responses to environmental
293 stress including exposure to heavy metals, oxygen radicals and antibiotics or drugs in a wide
294 range of bacterial genera [51]. *Orf10* is predicted as cell wall hydrolase, accession number
295 WP_057741916.1 [*Lactobacillus capillatus*]. It is putatively involved in the hydrolysis of
296 beta-1,4-linked polysaccharides [41]. *Orf26* is 100 % identical to the pfam01609 transposase
297 DDE domain of *Lactobacillus pentosus* MP-10 (CCB82565.1) which is essential for efficient
298 DNA transposition. Although we did not find any *tra* genes responsible for bacterial
299 conjugation in pB21AG01, three ORFs were annotated as genes related to plasmid
300 mobilisation. *Orf15* and *orf17* showed 93 % and 91 % identity to genes encoded for the
301 conjugative transposon proteins TcpE (WP_003688369.1) and TcpC (WP_003688364.1) of
302 *L. mali*, respectively. *Orf14* is 99 % identical to the pfam 12846 AAA-like domain containing
303 a P-loop motif from *L. capillatus* DSM_19910 (KRL03444.1), putatively involved in
304 conjugative transfer [41].

305

306 **Cloning of the B21AG gene cluster and LAB transformation**

307 The 3,424 bp sequence corresponding to seven genes putatively involved in plantacyclin
308 B21AG production, secretion and immunity was cloned into the pTRKH2 shuttle vector. The
309 resulting plasmid, pCycB21, transformed into *L. plantarum* WCFS1 is detailed in Fig 1.

310

311 **Fig. 1 Map of pTRKH2, pB21AG and pCycB21.** A: Map of pTRKH2 consisting an origin
312 of replication from plasmid P15A and a replication gene (green arrows) for replication in
313 both *E. coli* and gram positive bacteria; an erythromycin resistance gene (ErmR, yellow
314 arrow) for selection in *E. coli* and LAB; and a multiple cloning site (MCS). B: Map of
315 pB21AG containing 26 open reading frames, with seven ORFs corresponding to bacteriocin
316 production, immunity and transportation (red arrows); a plasmid replication and relaxation

317 gene (purple arrow); a transcription regulator (orange arrow), three conjugation transfer genes
318 (blue arrows) and 14 other ORFs (grey arrows). C: Map of pCycB21 harbouring seven
319 bacteriocin-associated genes cloned into *SacI* and *BamHI* restriction sites.

320

321 The pTRKH2 and pCycB21 plasmids were electrotransformed into *L. plantarum* WCFS1 with
322 efficiencies of 2.4×10^2 and 3.4×10^2 transformants per μg DNA, respectively. Addition of
323 glycine in the growth medium inhibits formation of cross-linkages in the cell wall where L-
324 alanine is replaced by glycine, thereby weakening the cell wall and facilitating DNA uptake
325 by the cells [52,53].

326

327 **Assay of plantacyclin B21AG expression**

328 Antimicrobial activities of the wild type *L. plantarum* B21 and recombinant host *L.*
329 *plantarum* WCFS1 were assayed using the well diffusion agar (WDA) method. Since
330 pTRKH2 conferred erythromycin (Em) resistance on WCFS1, culture supernatants
331 containing Em were assayed using indicator WCFS1 (pTRKH2) without removing the
332 antibiotics. To eliminate the effect of acid production, the pH of cell free supernatants were
333 neutralised to pH6.5. Both wild type and recombinant hosts were found to produce inhibition
334 zones against the indicator strain up to 1:8 dilution (Fig 2). The antimicrobial activity was
335 calculated as approximately 800 AU/mL for both strains. This indicates that the inhibitory
336 activity is not due to acid production but to an antimicrobial substance secreted into the broth
337 [54]. The CFS of control cultures containing WCFS1 (pTRKH2) did not show any inhibitory
338 activity, confirming that the recombinant plasmid pCycB21 was responsible for the
339 antimicrobial activity.

340

341 **Fig. 2 Antimicrobial activity of a two-fold serial dilution of Plantacyclin B21AG**
342 **secreted by (a) *Lactobacillus plantarum* B21 and (b) WCFS1 harbouring pCycB21. *L.***
343 ***plantarum* WCFS1 (pTRKH2) was used as the indicator strain. Numbers above the wells**
344 **correspond to the CFS dilution in each well. 1, Undiluted CFS; 2, 1:2 dilution of CFS; 3, 1:4**
345 **dilution of CFS; 4, 1:8 dilution of CFS; 5, 1:16 dilution of CFS; 6, 1:32 dilution of CFS;**
346 **7,1:64 dilution of CFS; 8, 1: 128 dilution of CFS. Well 9 (indicator strain) and well 10 (MRS**
347 **broth) are negative control.**

348

349 **Mass spectrometry analysis**

350 The plantacyclin B21AG produced by the wild type B21 and recombinant host WCFS1
351 (pCycB21) was purified by extraction into butanol. MALDI-TOF-MS analysis revealed a
352 major peptide of molecular mass of 5663.9 Da, essentially identical to the plantacyclin B21AG
353 produced by the wild type *L. plantarum* B21 (5664.7 Da) (Fig 3). No major peaks were observed
354 for WCFS1 transformed with the shuttle vector pTRKH2, corroborating the results from the
355 functional expression assay described previously (Fig 2, well 9).

356

357 **Fig. 3 MALDI-TOF-MS spectrum of plantacyclin B21AG, showing a single peak at**
358 **molecular mass of 5664.69 for B21 and 5663.92 for WCFS1 (pCycB21). No major peaks**
359 **were observed for WCFS1 (pTRKH2).**

360

361 **Copy number of pB21AG01 and pCycB21**

362 The relative copy number of pB21AG01 and pCycB21 was determined by real-time PCR
363 using the single copy alanine racemase (*alr*) as reference gene. In our experiment, the
364 standard curves obtained for *alr*, *orf19* and *rep* were linear ($R^2 > 0.99$) over the range tested;
365 whilst the amplification efficiency for all experiments ranged between 90 – 102 %, which is

366 within the acceptable range (90-110%) [55]. Analysis of the results revealed that
367 approximately 7.6 ± 0.79 copies per cell of pB21AG01 was detected in *L. plantarum* B21
368 whilst pCycB21 in *L. plantarum* WCFS1 was present at a noticeably lower level of just $0.5 \pm$
369 0.05 copies, indicating that only half the cells carry the recombinant plasmid. In contrast, the
370 copy number of the shuttle vector pTRKH2 in *L. plantarum* WCFS1 was high at 25.19 ± 2.68
371 copies per cell (average for two clones \pm standard deviation) per chromosome equivalent.

372

373 **Discussion**

374 Several lactic acid bacteria species have been recognised as probiotics that possess important
375 traits such as the production of bacteriocins and organic acids, adhesion to host cells, and
376 resistance to antibiotics and heavy metals [56,57]. A number of native plasmids that encode
377 these probiotic traits have been sequenced from *L. plantarum* [56,58], *L. salivarius* [59] and
378 *L. fermentum* [57]. In this study, seven bacteriocin-associated genes were found to be located
379 on a 20 kb native plasmid, pB21AG01 in *L. plantarum* B21. No replication protein (repB) or
380 initiator replication family protein (repA) was found in pB21AG01. However, a DNA
381 replication and relaxation conserved domain was detected in *orf2*. We could not detect any
382 clear repeats in the region upstream of *orf2*, suggesting that the plasmid may replicate
383 through a mechanism yet to be determined [60].

384

385 In addition, we identified two *tcp* loci, TcpE (*orf15*) and TcpC (*orf17*), which are involved in
386 the transfer of conjugative plasmid, pCW3 from *Clostridium perfringens*. TcpE was shown
387 to play a role in the formation of Tcp transfer apparatus in the gram positive *Clostridium*
388 *perfringens* [61]. TcpC was identified as a bitopic membrane protein, where membrane
389 localisation is important for its function, oligomerisation and interaction with other
390 conjugation proteins [62]. Bantwal et al. [63] proposed that TcpC may initiate accumulation

391 of peptidoglycan hydrolase at the cell wall of *Clostridium*, resulting in degradation of
392 peptidoglycan, thus facilitate the formation of transfer apparatus. Interestingly, a cell wall
393 hydrolase (*orf10*) was also found in pB21AG01, suggesting that TcpC may play a role in
394 promoting the hydrolysis of *Lactobacillus* cell wall and subsequently the transfer of
395 pB21AG01. However, we did not transfer the plasmid by mating because we could not
396 determine a full set of genes responsible for the mobilisation of the plasmid. pCW3 has a
397 novel conjugation region consisting of 11 genes encoding the Tcp proteins (TcpA, TcpB,
398 TcpC, TcpD, TcpE, TcpF, TcpG, TcpH, TcpI, TcpJ and TcpM) [61,64]. Several studies have
399 demonstrated that TcpA, TcpD, TcpE, TcpF and TcpH are essential to form the conjugation
400 complex. Moreover, we could not detect any antibiotic resistance and/or heavy metal
401 resistance genes in pB21AG01, which could be used as natural selection markers if we were
402 to transfer the plasmid by mating. Due to the lack of *tcp*-encoded proteins and selection
403 markers, we decided to transfer the plasmid by electroporation.

404

405 Electroporation seems to be an efficient method to transfer plasmid DNA into LAB to
406 enhance their probiotic functionality, or to secrete therapeutic proteins into the culture
407 medium for human and animal health [65]. However, the success rate of LAB transformation
408 is extremely low compared to *E. coli* due to various restriction modification (RM) system
409 encoded by the host. RM systems are required to protect bacteria from foreign DNA such as
410 genetically transferred plasmid DNA or the bacteriophage DNA [65,66]. Since DNA
411 manipulation is easier in *E. coli* than in *Lactobacillus*, we have built the recombinant plasmid
412 in the shuttle vector pTRKH2, followed by propagating the plasmids in *E. coli* to obtain
413 sufficient amount of plasmid DNA for LAB transformation. Numerous attempts at
414 electrotransformation were performed according to various protocols described in the
415 literature but without any success. The parameters for electrotransformation that we have

416 tried included varying the percentage of glycine added to LAB culture prior to pelleting the
417 cells; addition of PEG₁₅₀₀ and EDTA in the washing steps; various concentrations of plasmid
418 DNA; different electroporation buffers and diverse combinations of voltage (V), capacitance
419 (μ F) and resistance (ohm). The final modified version of the method described [44], that we
420 eventually had success with is described in the materials and methods section.

421

422 A few attempts have been made to heterologously expressed bacteriocins in different LAB
423 species because they promise a food grade background, where the expression of bacteriocins
424 would enhance their probiotic functionality [28,67]. However, all of these peptides that have
425 been successfully expressed heterologously belong to the class II bacteriocins. To date, only
426 one circular bacteriocin, plantaricyclin A from *L. plantarum* NI326, has been successfully
427 cloned into a nisin-inducible plasmid and expressed in *L. lactis* pNZPlc. Both the
428 recombinant host and the wild type producer exhibited similar level of antimicrobial activity
429 [25], indicating that circular bacteriocin can be heterologously expressed in other LAB
430 species. Our results are in accordance to Borrero et al. [25], where both native producer and
431 the recombinant host expressed similar antimicrobial activity up to 800 AU/mL. This result is
432 also confirmed by a single peak observed in our mass spectrometry analyses. In contrast, *L.*
433 *plantarum* WCFS1 transformed with the empty vector pTRKH2 produced no bacteriocin
434 activity. These results indicate that the bacteriocin activities observed are due to the cloned
435 genes (*orf19 – orf25*). We demonstrated that a 3.4 kb plasmid region of *L. plantarum* B21 is
436 sufficient for functional expression of plantacyclin B21AG. However, our attempt to
437 transform pCycB21 into *Lactobacillus agilis* La3, a type of LAB found to be colonising
438 chicken gastrointestinal tract (GIT) [68], did not result in heterologous expression of the
439 bacteriocin. This result suggests that the expression of recombinant protein in LAB is
440 species-specific. One possible explanation is that the native promoter cloned is specific to *L.*

441 *plantarum*, and a promoter from *L. agilis* is probably required for heterologous expression.

442 Although bacteria promoters share similar features, promoter strength is strain- and context-

443 specific, and can vary significantly within LAB [67,69,70].

444

445 The isolation and purification of bacteriocins from their LAB producers is often very time-

446 consuming and labour intensive [71]. Many studies have been performed to heterologously

447 express and overexpress the class II bacteriocins in *E. coli* to facilitate the production of these

448 antimicrobial peptides. For instance, sakacin P, pediocin PA-1, divercin V41 and plantaricin

449 NC8 have been successfully expressed in *E. coli* [71-74]. However, no circular bacteriocins

450 have been successfully expressed in *E. coli*. Kawai et al. [75] tried to express a circular

451 bacteriocin, gassericin A, in *E. coli* JM109 as a biotinylated fusion protein. However, a

452 positive clone which accumulated the bacteriocin did not show any antimicrobial activity.

453 Further treatment with factor Xa protease released the N-terminal leader peptide, resulting in

454 an active unclosed gassericin A. The results indicate that expression of circular bacteriocins is

455 host-specific, where a yet-to-be identified host-encoded peptidase is required to cleave the

456 leader peptide, allowing the ligation of N- and C- terminal to happen [13,76].

457

458 Plasmid copy number (PCN) analysis showed that the native pB21AG01 is a highly stable,

459 low copy number plasmid in B21. pTRKH2 was selected as a shuttle cloning vector because

460 it has been shown that it is structurally stable in *Lactobacillus vaginalis* Lv5, a common

461 feature of the theta-replicating mechanism [68]. It also has good structural stability in *E. coli*,

462 possibly due to the lack of a resolvase-encoding gene [37]. PCN analysis revealed that

463 pTRKH2 is more stable than pCycB21 in *L. plantarum* WCFS1, indicating that the

464 bacteriocin gene cluster may cause instability of the vector pTRKH2. Erythromycin selection

465 is required to maintain the pCycB21 in *L. plantarum* WCFS1. One possible reason which

466 may contribute to the instability of pCycB21 is plasmid incompatibility, where two plasmids
467 containing the same origin of replication cannot co-exist stably in the cell. Plasmids that have
468 growth advantages, such as faster replication and less toxicity will rapidly outgrow the other
469 plasmids [77]. The host used in this study, *L. plantarum* WCFS1 is known to harbour three
470 native plasmids size 1.9 kb, 2.4 kb and 36 kb [56]. Thus, the introduced bacteriocin gene
471 cluster could be a plausible reason for pCycB21 instability in *L. plantarum* WCFS1.
472 Similarly, the copy number of pCycB21 is extremely low compared to the native plasmid
473 pB21AG01. This suggests that other fitness factors present on the native plasmid pB21AG01
474 play a role in positive plasmid selection. For instance, apart from the immunity genes which
475 are known to protect bacteriocin-producing strains against its own toxins, the gene encoding
476 for ABC transporter also plays an important role. It has been shown to translocate the
477 bacteriocin across the cytoplasmic membrane, thereby avoiding toxin accumulation in the
478 host cells [78,79]. In our case, the presence of ABC transporter could potentially stabilise
479 pB21AG01 in *L. plantarum* B21. Despite the PCN variation between pB21AG01 and
480 pCycB21, plantacyclin B21AG was expressed at a similar level. The production of
481 plantacyclin B21AG would depend on plasmid stability and copy number differences
482 between pB21AG01 and pCycB21, but more likely, might be caused by the promoters used
483 to drive gene expression [7]. We have cloned the native promoter from pB21AG01 into
484 pCycB21AG, presumably resulting in similar levels of plantacyclin B21AG production. In
485 the future, inducible or controlled promoters may be tested to optimise heterologous
486 production of plantacyclin B21AG [67].

487

488 In summary, circular bacteriocins are thought to have more potential to form the next
489 generation of biopreservatives as a consequence of their stability and activity [8]. The ability
490 to transfer vectors harbouring the pB21AG01 gene cluster into an industry standard probiotic

491 *L. plantarum* WCFS1 highlights its biotechnological interest for the overproduction of the
492 antimicrobial peptide with high antimicrobial activity against food-borne pathogens.

493

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497

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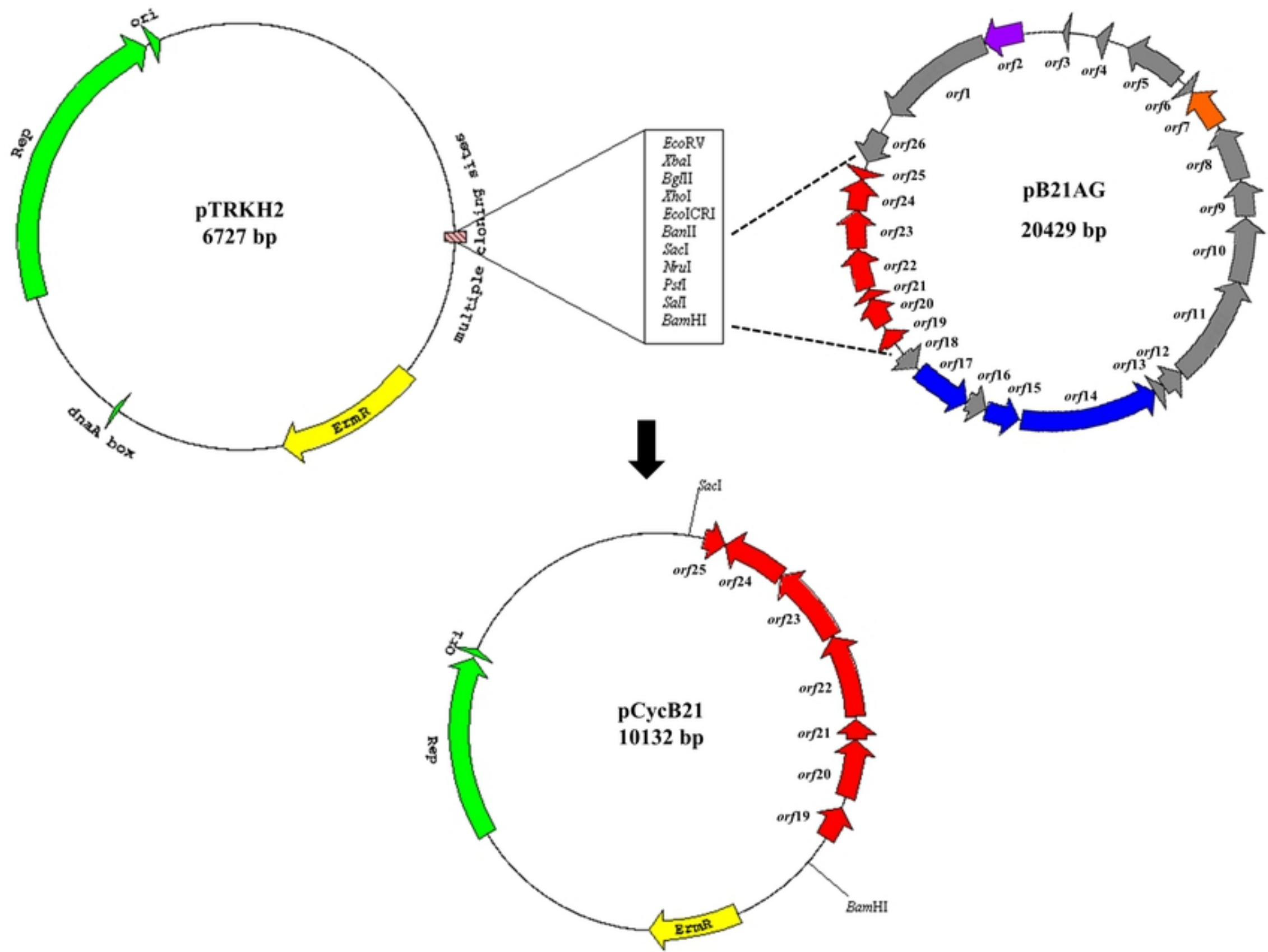


Figure 1

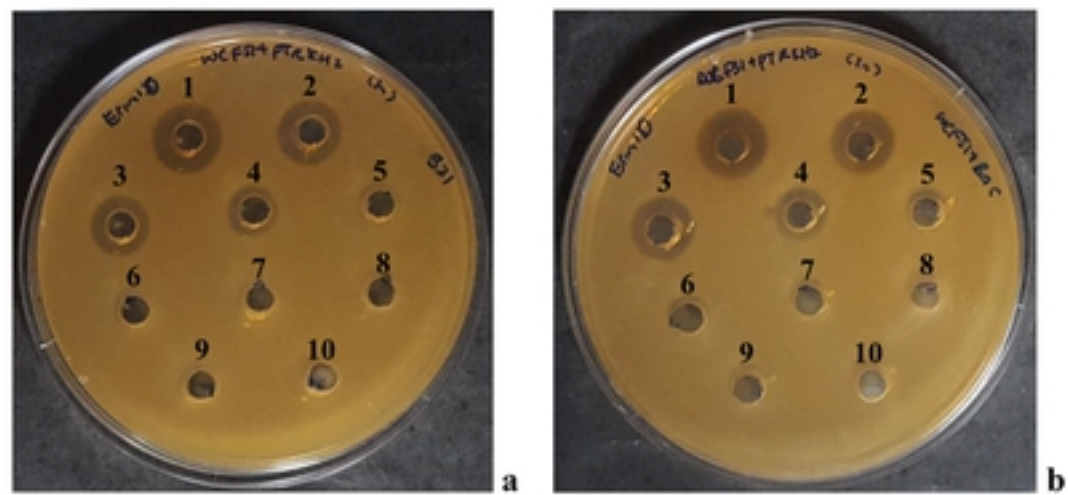


Figure 2

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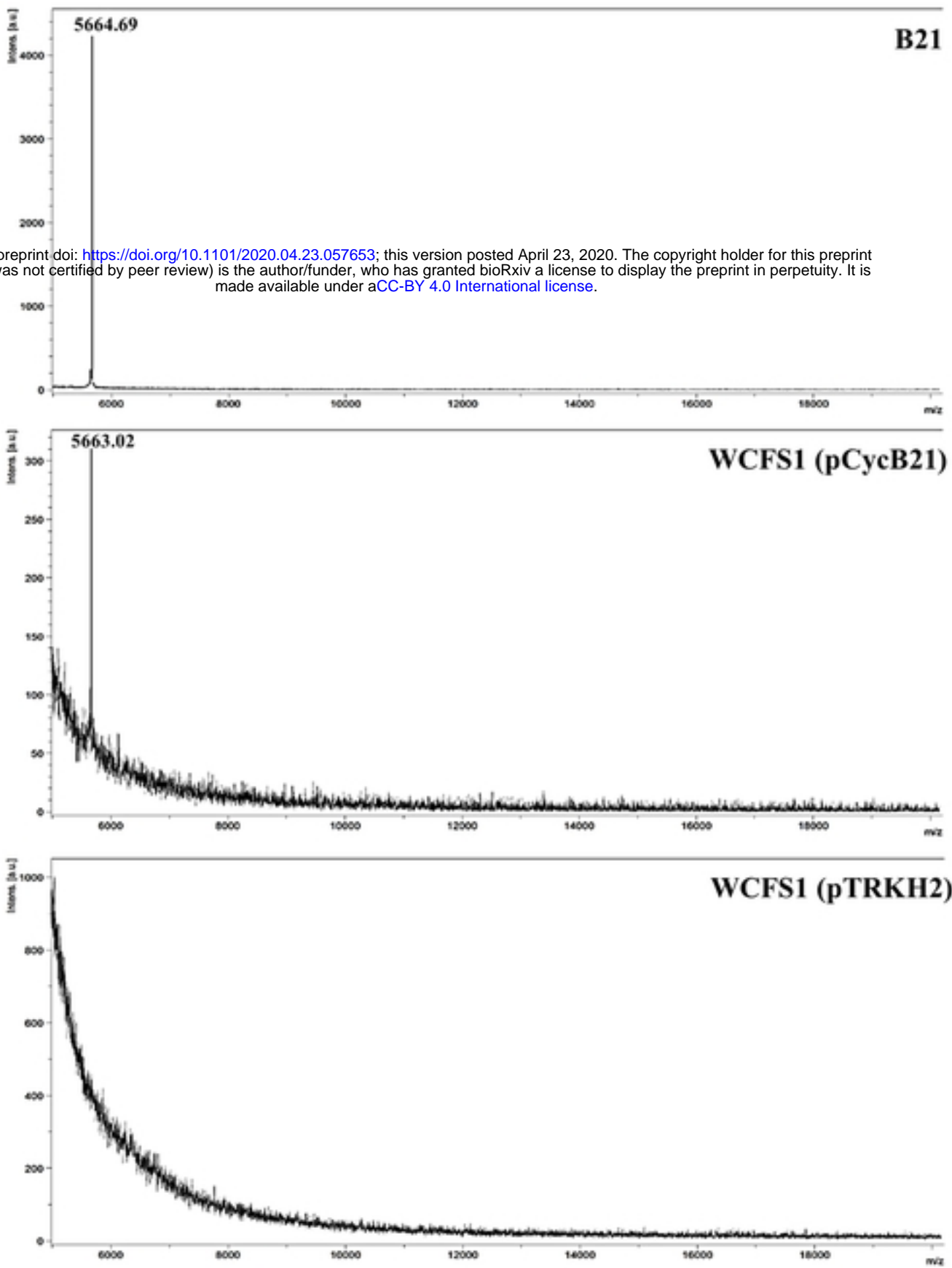


Figure 3