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# New insights into the role of plasmids from probiotic *Lactobacillus pentosus* MP-10 in Aloreña table olive brine fermentation

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*In silico* analysis of *Lactobacillus pentosus* MP-10 plasmids (pLPE-1 to pLPE-5) suggests that plasmidborne genes mediate the persistence of lactobacilli during olive fermentation and enhance their probiotic properties and their competitiveness in several ecological niches. The role of plasmids in the probiotic activities of *L. pentosus* MP-10 was investigated by plasmid-curing process which showed that plasmids contribute in increased metal tolerance and the biosequestration of several metals such as iron, aluminium, cobalt, copper, zinc, cadmium and mercury. Statistically significant differences in mucin adhesion were detected between the uncured and the cured *L. pentosus* MP-10, which possibly relied on a serine-rich adhesin (*sraP*) gene detected on the pLPE-2 plasmid. However, plasmid curing did not affect their tolerance to gastro-intestinal conditions, neither their growth ability under predetermined conditions, nor auto-aggregation and pathogen co-aggregation were changed among the cured and uncured *L. pentosus* MP-10. These findings suggest that *L. pentosus* MP-10 plasmids play an important role in gastro-intestinal protection due to their attachment to mucin and, thus, preventing several diseases. Furthermore, *L. pentosus* MP-10 could be used as a bioquencher of metals in the gut, reducing the amount of these potentially toxic elements in humans and animals, food matrices, and environmental bioremediation.

Table olive fermentation is the oldest practice by our ancestors to preserve vegetables and to also produce different flavours and textures. Additionally, fermented table olives remain an important economy for many production countries and a component of the Mediterranean diet (and recommended as part of the Healthy Eating Pyramid published in 2010, https://dietamediterranea.com/). The high nutritional value of fermented table olives (e.g., their content of carbohydrates, fiber, minerals, vitamins, fatty acids, and amino acids) and their role as potential source of probiotic lactobacilli of vegetable origin<sup>1-5</sup> make them very attractive from an economic and social point of view. Lactobacillus genus is the most representative and heterogeneous member of lactic acid bacteria (LAB) group currently consisting of 237 species (as of December 2018 in www.bacterio.net) since they harbour in their genome a plethora of genes involved with a wide array of functional properties<sup>6,7</sup>. Lactobacillus spp. are principal bacteria in olive fermentation processes, possessing many biochemical and physiological traits to ferment several carbohydrates and tolerate stress<sup>8</sup>. These phenotypes are important as the brine environment represent harsh conditions for bacterial growth with low nutrient availability, saltiness, low pH and the presence of antimicrobials (e.g., phenolic compunds and oleuropein); thus, highly robust L. plantarum and L. pentosus are frequently isolated from the end of olive fermentation<sup>1,8,9</sup>. Furthermore, Perpetuini, et al.<sup>10</sup> demonstrated by transposon mutagenesis that the high capacity of L. plantarum and L. pentosus to survive in the hostile, brine environments was due to critical genes encoding proteins involved in carbohydrate metabolism, membrane structure and function, and

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					Similarity to plasmids (BlastN)	
Plasmid	Size (bp)	G+C(%)	CDs	Hypothetical proteins	Identity in <i>Lactobacillus</i> (Isolation source)	Coverage (%)
pLPE-1	29,077	40.77	35	20	92% in <i>L. plantarum</i> subsp. <i>plantarum</i> TS12 plasmid pLP12-4 ("Stinky tofu")	59
pLPE-2	34,764	39.93	36	13	99% in <i>L. pentosus</i> IG1, annotated genomic scaffold00003 (Spanish-style green-olive fermentations)	47
pLPE-3	38,717	42.50	42	10	91% in <i>L. plantarum</i> strain BLS41 plasmid pLPBLS41_3 (Kimchi)	28
pLPE-4	43,946	40.09	53	32	91% in <i>L. casei</i> str. Zhang plasmid plca36 (Koumiss)	75
pLPE-5	46,498	39.52	58	32	99% in <i>L. plantarum</i> WCFS1 1.25 plasmid pWCFS103 (Human saliva)	51

Table 1. General features of circular plasmids from L. pentosus MP-10.

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gene-expression regulation. They further suggested that the *obaD* gene, which encodes a putative membrane protein strictly specific to *L. pentosus/L. plantarum* species, may be one of the key elements involved in their efficient adaptation to several conditions in many fermented food processes and natural ecosystems<sup>10</sup>.

Aloreña green table olive fermentation is a spontaneous process relying on *L. pentosus* strains and yeasts<sup>1,9</sup>. Resistance, persistence and predominance of *Lactobacillus* spp. in green table olive fermentation is due to their genetic variation and plasticity related to their chromosome and plasmids. In fact, *L. pentosus* species isolated from olive fermentation harbours the largest genome recognized to date and several plasmids (range: n = 5 to 7)<sup>11–13</sup>. However, *L. plantarum* contains the largest plasmids among the genus *Lactobacillus* plasmids are cryptic; however, they possess important properties such as antibiotic resistance, exopolysaccharide production, antimicrobial activity, bacteriocin synthesis, bacteriophage resistance, carbohydrate metabolism, host colonization and probiotic activity<sup>17–22</sup>. On the other hand, megaplasmids were also detected in *Lactobacillus* sp., up to 490 kb<sup>23</sup>. In this study, we analyzed *in silico* five plasmids harboured by *L. pentosus* MP-10 isolated from naturally fermented Aloreña green table olives<sup>2,9,12</sup>. Moreover, we aimed to better understand the underlying functional and probiotic properties of these plasmids using curing plasmid experiments; in particular, we examined their physiological traits in metal tolerance and biosorption, antimicrobial activity and adaptation to gastro-intestinal conditions to determine possible probiotic applications of this bacterium.

#### Results

**General features of** *L. pentosus* **MP-10 plasmids.** We have already reported the sequencing of *L. pentosus* MP-10 genome<sup>12</sup>, which consisted of a single circular chromosome of 3,698 kbp and five plasmids ranging 29–46 kbp (accession numbers FLYG01000001 to FLYG01000006). Sequence annotation was done using the Prokka annotation pipeline, version  $1.11^{24}$  as previously reported by Abriouel, *et al.*<sup>12</sup>. The general features of the circular five plasmids<sup>2</sup> are reported in Table 1. The average GC content of *L. pentosus* MP-10 plasmids ranged 39.52–42.50%, slightly lower than the host chromosome (with GC value of 46.32%). Furthermore, the GC contents of *L. pentosus* MP-10 plasmids were among the highest of known *L. pentosus* plasmids. All open reading frames in *L. pentosus* MP-10 plasmids are greater than 34 amino acids (Tables 2–6). Blast search for homology revealed lower identity with other plasmids in the database; however depending on coverage percentage, some regions harboring several genes in *L. pentosus* MP-10 plasmids were highly related with plasmids of *Lactobacillus* species isolated from foods like fermented olives, kimchi, koumiss, tofu or raw sausages, and also from human saliva (Table 1).

Figure 1 shows the frequency of KEGG functional annotations by BlastKOALA (KEGG tool; last updated March 4, 2016), which assigned plasmid genes to KEGG annotations corresponding to environmental information processing (pLPE-3, pLPE-4 and pLPE-5), genetic information processing (pLPE-2, pLPE-3, pLPE-4 and pLPE-5), carbohydrate metabolism (pLPE-3 and pLPE-5), amino acid metabolism (pLPE-3 and pLPE-5), cellular processes (pLPE-1, pLPE-2, pLPE-4 and pLPE-5), nucleotide metabolism (pLPE-2), metabolism of cofactors and vitamins (pLPE-3), and enzyme families (pLPE-3).

*In silico* analysis of plasmid properties in *L. pentosus* MP-10. Analysis of the annotated CDs of each *L. pentosus* MP-10 plasmid revealed the presence of five genes involved in mobilization (*mobA* gene) distributed in all plasmids except the pLPE-2 plasmid (Tables 2–6). These genes are likely required for plasmid relaxation and mobilization by conjugative plasmids. Also, conjugation-related genes were found, e.g., *traG* in pLPE-4 (*traG\_1* and *traG\_2*) and pLPE-5 (*traG\_3*) plasmids (Tables 5 and 6). A gene encoding for a bacteriophage peptidoglycan hydrolase that may have been involved in growth was found in pLPE-4 (*XX999\_00013* and *XX999\_00049*) and pLPE-5 (*XX999\_03566*) plasmids (Tables 5 and 6).

The presence of mobile genetic elements in *L. pentosus* MP-10 plasmids (pLPE-2, pLPE-3, pLPE-4 and pLPE-5) was already reported by Abriouel *et al.*<sup>2</sup> such as four putative transposon Tn552 DNA-invertase bin3 (four different genes of the same family), transposase DDE domain proteins (4 genes in pLPE 2 and pLPE5 plasmids), transposases of the mutator family (3 genes in pLPE2, pLPE3 and pLPE5 plasmids) and transposases (2 genes in pLPE-2 and pLPE-3 plasmids). Concerning integrases, one phage integrase family protein (pLPE-1 plasmid) and 9 integrase core domain proteins were detected in pLPE-2, pLPE-3 and pLPE-5 plasmids (Tables 3, 4 and 6). A gene *pinR* coding for DNA invertase from prophage was detected in pLPE-5 plasmid (Table 5).

Chloride- (*clcA\_2*) and sodium- (*nhaS3\_4*) transport genes harboured by pLPE-2 plasmid (Table 3) indicated that this plasmid was involved in salt-tolerance in brine solutions (plasmid curing experiments). Furthermore, a

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03518	XX999_03518	804-950	-	147	Hypothetical protein	_	98% identity in <i>L. paracasei</i> subsp. <i>paracasei</i> Lpp70
XX999_03519	XX999_03519	963-1271	-	309	Phage integrase family protein	_	87% identity in <i>Lactobacillus</i>
XX999_03520	XX999_03520	1238-1651	_	414	Hypothetical protein	_	99% identity in <i>L. plantarum</i> IPLA88
XX999_03521	XX999_03521	1871–2215	+	345	Toxin MazF	DNA binding (MF); RNA binding (MF); endoribonuclease activity (MF); endoribonuclease activity, producing 5'-phosphomonoesters (MF); negative regulation of cell growth (BP); regulation of mRNA stability (BP); RNA phosphodiester bond hydrolysis, endonucleolytic (BP)	100% identity in <i>L. pentosus</i>
XX999_03522	XX999_03522	2675-3739	-	1065	Hypothetical protein	—	99% identity in L. xiangfangensis
XX999_03523	XX999_03523	3901-4380	-	480	Hypothetical protein	-	100% identity in L. pentosus
XX999_03524	XX999_03524	4989-5576	-	588	Initiator Replication protein	_	98% identity in <i>L. plantarum</i>
XX999_03525	XX999_03525	6296-6490	-	195	Hypothetical protein	_	100% identity in L. pentosus IG1
XX999_03526	mobA_4	7058-8221	+	1164	Mobilization protein A	Conjugation (BP); DNA binding (MF); DNA-directed RNA polymerase activity (MF); DNA topoisomerase type I activity (MF); cytoplasm (CC); metal ion binding (MF)	100% identity in <i>L. pentosus</i>
XX999_03527	XX999_03527	8218-8910	+	693	Hypothetical protein	_	100% identity in L. pentosus
XX999_03528	XX999_03528	9111-9866	_	756	Initiator Replication protein	_	100% identity in <i>L. plantarum</i> IPLA88
XX999_03529	XX999_03529	10508-10957	+	450	Hypothetical protein	—	100% identity in L. pentosus
XX999_03530	XX999_03530	10954-11157	+	204	Hypothetical protein	-	100% identity in L. pentosus
XX999_03531	XX999_03531	11306-11668	-	363	Hypothetical protein	—	100% identity in L. pentosus
XX999_03532	XX999_03532	11912-12271	-	360	Hypothetical protein	_	99% identity in L. brevis
XX999_03533	XX999_03533	12284-12871	-	588	Site-specific tyrosine recombinase XerC	_	99% identity in <i>L. plantarum</i> 2025
XX999_03534	XX999_03534	12949-13212	+	264	Putative regulator PrlF	Regulation of cell growth (BP); DNA binding (MF); sequence-specific DNA binding transcription factor activity (MF); cytoplasm (CC); transcription, DNA-templated (BP); enzyme binding (MF); negative regulation of transcription, DNA-templated (BP)	100% identity in <i>L. plantarum</i>
XX999_03535	ndoA_2	13212–13559	+	348	mRNA interferase EndoA	DNA binding (MF); RNA binding (MF); endoribonuclease activity (MF); endoribonuclease activity, producing 5'-phosphomonoesters (MF); negative regulation of cell growth (BP); regulation of mRNA stability (BP); RNA phosphodiester bond hydrolysis, endonucleolytic (BP)	98% identity in <i>Lactobacillus</i>
XX999_03536	XX999_03536	14021-15085	-	1065	Hypothetical protein	-	99% identity in <i>L. xiangfangensis</i>
XX999_03537	XX999_03537	15164-15751	-	588	Hypothetical protein	—	100% identity in <i>L. pentosus</i>
XX999_03538	XX999_03538	15993-16928	-	936	Initiator Replication protein	-	99% identity in <i>L. plantarum</i> subsp. <i>plantarum</i>
XX999_03539	XX999_03539	17648-17842	-	195	Hypothetical protein	-	100% identity in <i>L. pentosus</i> IG1
XX999_03540	mobA_5	18410-19573	+	1164	Mobilization protein A	Conjugation (BP); DNA binding (MF); DNA-directed RNA polymerase activity (MF); DNA topoisomerase type I activity (MF); cytoplasm (CC); metal ion binding (MF)	95% identity in <i>L. plantarum</i>
XX999_03541	XX999_03541	19570-20262	+	693	Hypothetical protein	_	98% identity in <i>L. plantarum</i> 2025
XX999_03542	XX999_03542	20463-21218	-	756	Initiator Replication protein	_	100% identity in <i>L. plantarum</i> IPLA88
XX999_03543	XX999_03543	21860-22309	+	450	Hypothetical protein		100% identity in L. pentosus
XX999_03544	XX999_03544	22306-22509	+	204	Hypothetical protein	-	100% identity in L. pentosus
XX999_03545	XX999_03545	22658-23020	-	363	Hypothetical protein		100% identity in L. pentosus
XX999_03546	XX999_03546	23264-23623		360	Hypothetical protein	-	100% identity in L. pentosus
XX999_03547	XX999_03547	23636-24223	-	588	Site-specific tyrosine recombinase XerC	_	99% identity in <i>L. plantarum</i> 2025
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Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03548	XX999_03548	24300-24563	+	264	Putative regulator PrlF	Regulation of cell growth (BP); DNA binding (MF); sequence-specific DNA binding transcription factor activity (MF); cytoplasm (CC); transcription, DNA-templated (BP); enzyme binding (MF); negative regulation of transcription, DNA-templated (BP)	100% identity in <i>L. plantarum</i> 2025
XX999_03549	ndoA_3	24563-24910	+	348	mRNA interferase EndoA	DNA binding (MF); RNA binding (MF); endoribonuclease activity (MF); endoribonuclease activity, producing 5'-phosphomonoesters (MF); negative regulation of cell growth (BP); regulation of mRNA stability (BP); RNA phosphodiester bond hydrolysis, endonucleolytic (BP)	98% identity in <i>Lactobacillus</i>
XX999_03550	XX999_03550	25372-26436	_	1065	Hypothetical protein	-	99% identity in L. xiangfangensis
XX999_03551	XX999_03551	26515-27102	-	588	Hypothetical protein	—	100% identity in L. pentosus
XX999_03552	XX999_03552	27344-28279	-	936	Initiator Replication protein	_	99% identity in <i>L. plantarum</i> subsp. <i>plantarum</i>

**Table 2.** Genes determined in pLPE-1 plasmid of *Lactobacillus pentosus* MP-10 isolated from naturallyfermented Aloreña table olives. BP, biological process; CC, celular component; MF, molecular function.

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copy of the same genes *clcA\_1*, *nhaS3\_1*, *nhaS3\_2* and *nhaS3\_3* were also found in *L. pentosus* MP-10 chromosome with the aim to potentiate chloride and sodium tolerance in brines.

Genes related to carbohydrate metabolism were found on plasmids (besides on the chromosome) such as L-Lactate dehydrogenase in pLPE-5 plasmid (*ldh\_7* and *ldh\_8* genes) (Table 6), genes involved in glucose uptake and metabolism such as *glcU\_1* and *gdhIV\_1* genes in pLPE-3 plasmid (Table 4), and a gene involved in xylan catabolic process (*axeA1\_3*) in pLPE-5 (Table 5). However, another gene involved in xylan catabolic process (*XX999\_00089*) was only detected in pLPE-3 plasmid, but not on the chromosome (Table 4).

Toxins reported in *L. pentosus* MP-10 plasmids include mazF-toxin encoding gene (*XX999\_03521*) detected in pLPE-1 plasmid, genes coding for Zeta toxins in pLPE-3 (*XX999\_00053*) and pLPE-4 (*XX999\_00024*) plasmids, and also for antitoxins such as RelB antitoxin (*XX999\_00026*) in pLPE-4 plasmid and the bifunctional antitoxin/ transcriptional repressor RelB in pLPE-5 plasmid (*XX999\_03554*) (Tables 2, 4–6). MazF toxin is a desirable property in probiotic bacteria, and is only detected in plasmid DNA of *L. pentosus* MP-10, not in the chromosome. However, *L. pentosus* MP-10 has to protect itself from the MazF toxin without any MazE antitoxin. On the other hand, RelB antitoxins were found both on plasmids and on the chromosome; however, no RelB toxins were detected both on the chromosome (one gene) and also on plasmid DNA (two genes); however, no antitoxin was detected.

Other coding genes for several functions, such as a serine-rich adhesin for platelets precursor (*sraP* gene), were detected in pLPE-2 plasmid but not on the chromosome (Table 3); genes coding for vitamin biosynthesis such as *panE\_1* and *panE\_2* genes coding for 2-dehydropantoate 2-reductase (biosynthesis of vitamin B5), a gene XX999\_00068 coding for prephenate dehydratase (biosynthesis of phenylalanine, tyrosine and tryptophan), were detected on the pLPE-3 plasmid (Table 4) and also on the chromosome.

Regarding their responses to stress, *in-silico* analysis of plasmid sequences revealed the presence of *yhdN\_1* gene coding for a general stress protein 69 (in pLPE-3, Table 4) and several genes coding for metal tolerances, such as cadmium [cadmium resistance transporter (*XX999\_03594*) and a putative positive regulator of cadmium resistance (*cadC*)] and two operons of arsenic resistance (in pLPE-5, Table 6). One *ars* operon consists of *arsR\_3* (arsenical resistance operon repressor ArsR) and *arsB* [arsenical pump membrane protein (ArsB)], but lacks *arsC* gene (arsenate reductase ArsC); the other *ars* operon contains *arsA* [arsenical pump-driving ATPase (ArsA)] and *arsD* gene [arsenical resistance operon trans-acting represor (ArsD)] in pLPE-5 (Table 6). The synteny of arsenic-resistance genes was examined by comparing the annotated sequences of pLPE-5 and pWCFS103 plasmids (aligned by MAUVE algorithm) from *L. pentosus* MP-10 and *L. plantarum* WCFS1, respectively. Comparison revealed that the synteny of genes was similar (Fig. 2), being arsenic operons in pLPE-5 of *L. pentosus* MP-10 composing of two copies each gene: *arsB* [coding for trivalent As(III) efflux permease ArsB], *arsA* [coding for trivalent As(III)-stimulated ATPase ArsA], *arsD* [coding for trivalent As(III) metallochaperone ArsD] and *arsR\_3* gene [a trivalent As(III)-responsive repressor (ArsR)]. On the other hand, *arsC* gene (*arsC2* coding for reductase ArsC), as a part of *ars* operon with *arsB* and *arsR* genes, was found in *L. pentosus* MP-10 chromosome, as well as two *arsR* gene copies (*arsR\_1* and *arsR\_2*).

*In vitro* detection of functional properties in *L. pentosus* MP-10 plasmids. *Effect of plasmid curing on growth of L. pentosus MP-10.* The MIC of acridine orange (AO) was of 0.15 mg/ml; as such, we used 0.1 mg/ml as the sub-MIC for plasmid curing in this strain. After confirming *L. pentosus* MP-10 being cured of plasmids (data not shown), we compared the growth kinetics of uncured and cured *L. pentosus* MP-10C. The presence of plasmids did not affect the growth in MRS broth at 37 °C in any experimental conditions: presence/

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03611	clcA_2	189-1568	+	1380	H(+)/Cl(-) exchange transporter ClcA	Voltage-gated chloride channel activity (MF); integral component of plasma membrane (CC); antiporter activity (MF)	99% identity in <i>L. pentosus</i> SLC13 plasmid pSLC13
XX999_03612	XX999_03612	2377-3297	+	921	Integrase core domain protein	_	99% identity in L. pentosus
XX999_03613	XX999_03613	4093-4353	-	261	Phd_YefM	DNA binding (MF); transcription, DNA- templated (BP); regulation of transcription, DNA-templated (BP)	100% identity in <i>L.</i> <i>plantarum</i> CMPG5300
XX999_03614	XX999_03614	4535-5902	-	1368	Transposase DDE domain protein	_	82% identity in L. backii
XX999_03615	sraP	6332-8032	+	1701	Serine-rich adhesin for platelets precursor	Calcium ion binding (MF); extracellular region (CC); cell wall (CC); pathogenesis (BP); membrane (CC)	60% identity in <i>L.</i> <i>plantarum</i> O2T60C
XX999_03616	yusO	8090-8527	+	438	Putative HTH-type transcriptional regulator YusO	DNA binding (MF); sequence-specific DNA binding transcription factor activity (MF); intracellular (CC); transcription initiation from RNA polymerase II promoter (BP)	99% identity in <i>L. pentosus</i> IG1
XX999_03617	XX999_03617	8891-9229	+	339	Hypothetical protein	_	100% identity in <i>L.</i> <i>plantarum</i> plasmid pLP9000_06
XX999_03618	XX999_03618	9187-9690	+	504	Transposase DDE domain protein	-	100% identity in <i>L.</i> <i>plantarum</i> UCMA 3037
XX999_03619	XX999_03619	9650–9883	-	234	Hypothetical protein	_	62% identity in <i>L.</i> <i>plantarum</i> subsp. <i>plantarum</i>
XX999_03620	XX999_03620	10191-11729	-	1539	Hypothetical protein	_	-
XX999_03621	soj_3	12508-13425	+	918	Sporulation initiation inhibitor protein Soj	ATP binding (MF); oxidoreductase activity (MF); hydrolase activity (MF); sporulation resulting in formation of a cellular spore (BP); negative regulation of sporulation resulting in formation of a cellular spore (BP)	99% identity in <i>L. pentosus</i> DSM 20314
XX999_03622	XX999_03622	13409-13696	+	288	Hypothetical protein	_	100% identity in <i>L.</i> plantarum
XX999_03623	XX999_03623	13862-15037	+	1176	Transposase, Mutator family	-	100% identity in <i>L. pentosus</i>
XX999_03624	XX999_03624	15547-15732	-	186	Hypothetical protein	_	53% identity in <i>L. plantarum</i>
XX999_03625	XX999_03625	16181-16477	-	297	Hypothetical protein	_	100% identity in <i>L.</i> pentosus DSM 20314
XX999_03626	XX999_03626	16698-17015	+	318	Hypothetical protein	_	100% identity in <i>L. plantarum</i>
XX999_03627	XX999_03627	17186-17482	+	297	Transposase DDE domain protein	_	100% identity in <i>L. plantarum</i>
XX999_03628	XX999_03628	17582-17986	+	405	D-alanine/D-serine/glycine permease	_	99% identity in <i>L.</i> plantarum
XX999_03629	XX999_03629	18547-18774	-	228	Hypothetical protein	_	100% identity in <i>L.</i> <i>pentosus</i> DSM 20314
XX999_03630	soj_4	19677-20477	+	801	Chromosome-partitioning ATPase Soj	ATP binding (MF)	100% identity in <i>L. pentosus</i> IG1
XX999_03631	XX999_03631	20479-20826	+	348	Hypothetical protein	_	100% identity in <i>L. plantarum</i> CMPG5300
XX999_03632	XX999_03632	21412-22314	-	903	Hypothetical protein	_	100% identity in <i>L. pentosus</i> IG1
XX999_03633	bin3_4	22401-23033	_	633	Putative transposon Tn552 DNA-invertase bin3	Recombinase activity (MF); DNA binding (MF); DNA integration (BP); transposition (BP)	99% identity in <i>L.</i> <i>plantarum</i> 16
XX999_03634	XX999_03634	23329-23958	+	630	Integrase core domain protein	-	100% identity in <i>L. pentosus</i> IG1
XX999_03635	nrdF2_2	24087-25037	_	951	Ribonucleoside-diphosphate reductase subunit beta nrdF2	Ribonucleoside-diphosphate reductase activity, thioredoxin disulfide as acceptor (MF); ribonucleoside-diphosphate reductase complex (CC); DNA replication (BP); deoxyribonucleoside diphosphate metabolic process (BP); deoxyribonucleotide biosynthetic process (BP); metal ion binding (MF)	100% identity in <i>L. pentosus</i> IG1
XX999_03636	nrdF	25052-25978	-	927	Ribonucleoside-diphosphate reductase 2 subunit beta	Ribonucleoside-diphosphate reductase activity, thioredoxin disulfide as acceptor (MF); ribonucleoside-diphosphate reductase complex (CC); DNA replication (BP); deoxyribonucleoside diphosphate metabolic process (BP); deoxyribonucleotide biosynthetic process (BP); metal ion binding (MF)	100% identity in <i>L.</i> pentosus IG1
XX999_03637	nrdE	26085-28253	_	2169	Ribonucleoside-diphosphate reductase 2 subunit alpha	Ribonucleoside-diphosphate reductase activity, thioredoxin disulfide as acceptor (MF); ATP binding (MF); DNA replication (BP)	100% identity in <i>L.</i> <i>pentosus</i> DSM 20314
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Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03638	XX999_03638	28260-28697	-	438	Putative NrdI-like protein	-	100% identity in <i>L. plantarum</i> AY01
XX999_03639	XX999_03639	29395-29496	-	102	Hypothetical protein	_	100% identity in <i>L. plantarum</i> 2165
XX999_03640	XX999_03640	29486-29845	-	360	Putative hydrolase	-	99% identity in <i>L. plantarum</i> 2165
XX999_03641	XX999_03641	30683-30943	-	261	Hypothetical protein	Recombinase activity (MF); DNA binding (MF); DNA integration (BP)	100% identity in <i>L.</i> <i>plantarum</i> AY01
XX999_03642	XX999_03642	30999-31250	+	252	Transposase	-	100% identity in <i>L. pentosus</i>
XX999_03643	XX999_03643	31304-32146	+	843	Integrase core domain protein	_	99% identity in <i>L.</i> plantarum
XX999_03644	XX999_03644	32416-32805	-	390	Integrase core domain protein	-	99% identity in <i>L. plantarum</i>
XX999_03645	XX999_03645	32896-33381	-	486	Hypothetical protein	-	100% identity in <i>L. plantarum</i>
XX999_03646	nha\$3_4	33487-34641	+	1155	High-affinity Na(+)/H(+) antiporter NhaS3	Plasma membrane (CC); sodium ion transmembrane transporter activity (MF); antiporter activity (MF); solute:proton antiporter activity (MF); integral component of membrane (CC); sodium ion transmembrane transport (BP)	100% identity in <i>L. pentosus</i> IG1

**Tablze 3.** Genes determined in pLPE-2 plasmid of *Lactobacillus pentosus* MP-10 isolated from naturally fermented Aloreña table olives. BP, biological process; CC, celular component; MF, molecular function.

absence of 6.5% NaCl, different pH ranges (1.5 to 7.0), nor the presence of bile salts (1.8 or 3.6%) -no differences in 600 nm absorbances were detected over 24 h of incubation- (Figs S1–A,B, S2). In a similar manner, pH monitoring during their incubation also did not exhibit any significant differences between cured and uncured strains in regards to their acidification capacity (Fig. S1–C). Furthermore, no differences in the growth were detected between the cured and uncured *L. pentosus* MP-10 strains in the presence of xylan as the only carbohydrate source (Fig. S1–D). However, at high salt concentration of 8% usually found in brine, significant differences were detected between the cured and uncured *L. pentosus* MP-10 strains, with the uncured strain being the most tolerant (Fig. S1–E).

Table 7 shows that curing had no significant effect on the growth of uncured and cured *L. pentosus* MP-10 in the presence of phenolic compounds naturally present in the brines; both the cured and uncured strains tolerated more than 200 mg/ml of olive-leaf extract.

*Effect of plasmid curing on metal tolerance.* Plasmid annotations predicted gene clusters involved in arsenateand/or arsenite-, and cadmium resistance. First, we precisely determined metal concentrations that inhibit the visible growth of the wildtype *L. pentosus* MP-10; results showed that this strain tolerated high concentrations of metals depending on the metal with  $1 < MIC < 4096 \mu g/ml$ , and tolerances were observed to be in order Fe > [Al/ Cu/Co] > Zn > Cd > Hg (Table 8). When we compared the uncured and the cured *L. pentosus* MP-10, we found that mercury and cadmium exibited different MICs among strains by 2–8 fold increase (Table 8) in those uncured; as such, plasmids have a key role in mercury and cadmium tolerances.

The removal of different metals was shown in Table 8, which demonstrated that *L. pentosus* MP-10 was able to remove different metals, thus exhibiting high removal capacity of mercury ( $81.74\% \pm 2.04$ ), cadmium ( $67.10\% \pm 0.88$ ) and aluminium ( $57.14\% \pm 0.99$ ). However, the cured *L. pentosus* MP-10C demonstrated statistically significant reduced performance. Metal removal differences between the uncured and the cured *L. pentosus* MP-10 highlight the role of plasmids to remove iron, cadmium, aluminium, cobalt, copper, zinc and mercury (Table 8).

To understand how *L. pentosus* MP-10 interact with selected metals, SEM analysis was performed and showed the biosorption potential of the uncured *L. pentosus* MP-10 (Fig. 3). The micrographs and EDX spectra obtained before and after the biosorption process showed clearly that the cell morphology of the uncured *L. pentosus* MP-10 changed and exhibited the presence of bright particles on the surface of the bacteria exposed to some metals. Regarding cadmium, mercury and zinc, we couldn't detect these metals by EDX analysis. Furthermore, in the presence of either aluminium, cobalt, copper, mercury or zinc, higher potential for biofilm formation was observed. These results, confirmed by EDX analyses, support that these metals remained adsorbed entirely on the cell surface.

*Effect of plasmid curing on antimicrobial resistance and probiotic features.* We determined the MIC of different antibiotics and biocides between uncured and cured strains, and the results did not show any significant differences in response between both strains except for clindamycin, which exibited 20 fold increase in the MIC in the uncured *L. pentosus* MP-10. Thus, plasmids have no role in the suceptibility to the antibiotics and biocides tested, except clindamycin (Table 7).

Regarding the probiotic features, the uncured and the cured *L. pentosus* MP-10 had performed similarly in auto-aggregation and co-aggregation with all pathogens tested (Table 7), which suggest that plasmids had neither

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms
XX999_00053	XX999_00053	146-412	-	267	Zeta toxin	—
XX999_00054	XX999_00054	586-783	_	198	Hypothetical protein	_
XX999_00055	XX999_00055	1002-1931	+	930	Integrase core domain protein	_
XX999_00056	XX999_00056	1934-2152	+	219	Hypothetical protein	_
XX999_00057	soj_1	3395-4204	+	810	Chromosome-partitioning ATPase Soj	DNA binding (MF); ATP binding (MF); chromosome segregation (BP); hydrolase activity (MF)
XX999_00058	XX999_00058	4197-4532	+	336	Hypothetical protein	_
XX999 00059	XX999 00059	4598-4771	+	174	Hypothetical protein	_
 XX999_00060	 XX999_00060	5611-6453	_	843	Integrase core domain protein	_
XX999_00061	 XX999_00061	6507-6758	_	252	Transposase	_
XX999_00062	XX999_00062	6826-7092	_	267	Divergent AAA domain protein	_
XX999_00063	ilvE_1	7372-8394	_	1023	Putative branched-chain-amino- acid aminotransferase	Isoleucine biosynthetic process (BP); leucine biosynthetic process (BP); valine biosynthetic process (BP); L-leucine transaminase activity (MF); L-valine transaminase activity (MF); L-isoleucine transaminase activity (MF)
XX999_00064	panE_1	8444-9463	-	1020	2-dehydropantoate 2-reductase	Cytoplasm (CC); 2-dehydropantoate 2-reductase activity (MF); pantothenate biosynthetic process from valine (BP); NADP binding (MF)
XX999_00065	yvdD_1	9990-10559	-	570	LOG family protein YvdD	-
XX999_00066	XX999_00066	10970-11968	+	999	Integrase core domain protein	-
XX999_00067	panE_2	12688-13698	+	1011	2-dehydropantoate 2-reductase	Cytoplasm (CC); 2-dehydropantoate 2-reductase activity (MF); pantothenate biosynthetic process from valine (BP); NADP binding (MF)
XX999_00068	XX999_00068	13686-14087	-	402	Prephenate dehydratase	_
XX999_00069	XX999_00069	14032-14613	-	582	Transposase, Mutator family	—
XX999_00070	asnB_1	14954-16543	_	1590	Asparagine synthetase B [glutamine- hydrolyzing]	Asparagine synthase (glutamine-hydrolyzing) activity (MF); aspartate-ammonia ligase activity (MF); ATP binding (MF); cytoplasm (CC); asparagine biosynthetic process (BP); glutamine metabolic process (BP); cellular amino acid biosynthetic process (BP); cellular amino acid catabolic process (BP); amino acid binding (MF); identical protein binding (MF); L-asparagine biosynthetic process (BP)
XX999_00071	bin3_2	17298-17972	_	675	Putative transposon Tn552 DNA- invertase bin3	Recombinase activity (MF); DNA binding (MF); DNA integration (BP); transposition (BP)
XX999_00072	ltrA_1	18520-19686	+	1167	Group II intron-encoded protein LtrA	RNA-directed DNA polymerase activity (MF); endonuclease activity (MF); intron homing (BP); mRNA processing (BP)
XX999_00073	hosA_1	20060-20479	+	420	Transcriptional regulator HosA	DNA binding (MF); sequence-specific DNA binding transcription factor activity (MF); intracellular (CC); transcription, DNA-templated (BP); pathogenesis (BP)
XX999_00074	XX999_00074	20536-20991	+	456	hypothetical protein	_
XX999_00075	XX999_00075	20988-21206	+	219	hypothetical protein	_
XX999_00076	XX999_00076	21421-21912	+	492	hypothetical protein	-
XX999_00077	XX999_00077	22017-22805	+	789	flavodoxin	-
XX999_00078	XX999_00078	22823-23476	+	654	NmrA-like family protein	-
XX999_00079	XX999_00079	23512-24384	+	873	Alpha/beta hydrolase family protein	—
XX999_00080	hsrA_2	24631-24924	+	294	putative transport protein HsrA	Plasma membrane (CC); integral component of membrane (CC); transmembrane transport (BP)
XX999_00081	efpA	24921-25958	+	1038	putative MFS-type transporter EfpA	Plasma membrane (CC); integral component of membrane (CC); transmembrane transport (BP)
XX999_00082	XX999_00082	26043-26618	+	576	flavodoxin	-
XX999_00083	glcU_1	26631-27491	+	861	Glucose uptake protein GlcU	Plasma membrane (CC); rhamnose transmembrane transporter activity (MF); integral component of membrane (CC); sporulation resulting in formation of a cellular spore (BP)
XX999_00084	yvgN_1	27580-28431	+	852	Glyoxal reductase	Methylglyoxal reductase (NADPH-dependent) activity (MF)
XX999_00085	gdhIV_1	28460-29245	+	786	Glucose 1-dehydrogenase 4	Identical protein binding (MF); glucose 1-dehydrogenase [NAD(P)] activity (MF)
XX999_00086	adhR_1	29308-29703	+	396	HTH-type transcriptional regulator AdhR	DNA binding (MF); transcription, DNA-templated (BP); regulation of transcription, DNA-templated (BP)
XX999_00087	XX999_00087	29700-30434	+	735	putative oxidoreductase	Oxidoreductase activity (MF)
XX999_00088	yhdN_1	30459-31436	+	978	General stress protein 69	Oxidoreductase activity (MF)
XX999_00089	XX999_00089	31514-32101	+	588	Polysaccharide deacetylase	Hydrolase activity, acting on carbon-nitrogen (but not peptide) bonds (MF); polysaccharide binding (MF); endo- 1,4-beta-xylanase activity (MF); xylan catabolic process (BP)
XX999_00090	XX999_00090	32681-33805	-	1125	hypothetical protein	-
XX999_00091	XX999_00091	33809-34024	-	216	hypothetical protein	-
Continued						

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms
XX999_00092	topB_4	34147-35697	-	1551	DNA topoisomerase 3	Magnesium ion binding (MF); DNA binding (MF); DNA topoisomerase type I activity (MF); DNA topological change (BP); DNA recombination (BP); chromosome separation (BP)
XX999_00093	mobA_2	35779-37842	_	2064	Mobilization protein A	Conjugation (BP); DNA binding (MF); DNA-directed RNA polymerase activity (MF); DNA topoisomerase type I activity (MF); cytoplasm (CC); metal ion binding (MF)
XX999_00094	XX999_00094	38344-38622	+	279	hypothetical protein	-

**Table 4.** Genes determined in pLPE-3 plasmid of *Lactobacillus pentosus* MP-10 isolated from naturallyfermented Aloreña table olives. BP, biological process; CC, celular component; MF, molecular function.

any role in auto-aggregation nor co-aggregation processes. Regarding acid and bile tolerance, no differences were detected between the uncured and the cured *L. pentosus* MP-10 (Table 7).

Adhesion to mucin was measured in both the uncured and the cured *L. pentosus* MP-10, and the results showed a statistically significant increase in adhesion capacity to mucin in the uncured *L. pentosus* MP-10 (Table 7).

#### Discussion

Olive brine represents a stressful environment for the growth and survival of many bacteria due to the harsh conditions (i.e., high salt concentration, presence of phenolic compounds and low-nutrient availability), which provide selective pressures for the maintenance of LAB. As such, *L. plantarum* and *L. pentosus* have the genetic tools to survive and grow in the hostile olive-brine conditions<sup>10</sup>, and these genetic traits are widely distributed on both the chromosome and the plasmids, with several genes having multiple copies to enhance their adaptability and fitness in different ecological niches.

In this study, *L. pentosus* MP-10, isolated from Aloreña green table olives, harboured five plasmids with an average GC content (39.52–42.50%) slightly lower than the host chromosome (46.32%), this difference was less than 10% as reported by Nishida<sup>25</sup> for the majority of plasmids. pLPE-5 had remarkably the lowest average GC content (39.52%) than the other four plasmids (pLPE-1, pLPE-2, pLPE-3 and pLPE-4), suggesting it is possibly a recent acquisition from another bacterium. *In-silico* analysis of plasmid sequences revealed the presence of genes involved in mobilization (*mobA*) and conjugation (*traG*) distributed in several plasmids, which suggest their role in gene mobilization and secretion using a type-IV secretion mechanism<sup>26</sup>. Furthermore, mobile genetic elements (e.g., transposon, transposase, integrase and invertase) were also found in several plasmids<sup>2</sup> suggesting a frequent genetic diversification among the *L. pentosus* MP-10. Furthermore, bacteriophage peptidoglycan hydrolases were found in pLPE-4 and pLPE-5 plasmids; these lysozyme-like proteins may play a key role in *L. pentosus* MP-10 growth, its cell-wall structure, and immunomodulatory properties as reported by Rolain, *et al.*<sup>27</sup>.

Metabolic profile within L. pentosus MP-10 plasmids include carbohydrate enzymes such as L-lactate dehydrogenase, glucose uptake and metabolism and xylan catabolic enzymes. L-lactate dehydrogenase was codified by two genes (*ldh\_7* and *ldh\_8*) located on pLPE-5 plasmid; however, six L-lactate dehydrogenase (*ldh\_1*, *ldh\_2*, ldh\_3, ldh\_4, ldh\_5 and ldh\_6) and four D-lactate dehydrogenase (XX999\_00315, XX999\_00955, XX999\_02047 and XX999\_02719) coding genes were also present on the chromosome. Both enantiomers (L-lactate and D-lactate) are produced by L. pentosus MP-10 being D-and L-lactate dehydrogenases involved in the reversible metabolism of D- and L-lactate, respectively. This finding is of great interest suggesting that the use of L. pentosus MP-10 as a probiotic may help human to metabolise D-lactate obtained from exogenous sources (e.g., diet and the carbohydrate-fermenting bacteria normally present in the gastrointestinal tract) since mammalian cells lack sufficient D-lactate dehydrogenase required to utilise D-lactic acid—leading to chronic fatigue syndrome and D-lactic acidosis or D-lactate encephalopathy associated with short bowel syndrome<sup>28-30</sup>. Further, L-lactate dehydrogenase genes present on the plasmids may enhance their metabolic activity during the fermentation process to produce more L-lactate and energy. However, the presence of L-lactate dehydrogenase (*ldh\_7* and *ldh\_8*) coding genes on pLPE-5 plasmid did not enhance the acidification capacity, as results were similar after 8 and 24 h incubation in both cured and uncured L. pentosus MP-10, suggesting that these genes either have a minor role in lactate production or they are regulated. Further experiments, based on differential relative expression of *ldh* gene in both the cured and uncured *L. pentosus* MP-10 strains, revealed low expression level in the cured strain (Fig. S3), thus the low activity of lactate dehydrogenase gene in the cured strain is enough to give rise to a substantial lactate accumulation in the fermentation broth in a manner similar as the uncured strain. Regarding glucose uptake and metabolism, glcU\_and gdhIV genes were over-expressed in the uncured L. pentosus MP-10 indicating the role of plasmid in this process (Fig. S3).

Among defense mechanisms found on plasmids, gene encoding the mazF toxin (pLPE-1), Zeta toxins (pLPE-3 and pLPE-4), and also antitoxins such as RelB antitoxin (pLPE-4) and the bifunctional antitoxin/transcriptional repressor RelB (pLPE-5) were detected in *L. pentosus* MP-10 plasmids. RelBE and MazEF are known as sequence-specific endo-ribonucleases that inhibit the global translations of cellular mRNAs<sup>31</sup>. MazF toxin is a desirable trait for probiotic bacteria, as its antimicrobial property inhibits several pathogens in foods and the gastrointestinal tract<sup>32</sup>. However, *L. pentosus* MP-10 must protect itself from the mazF toxin, as no MazE antitoxin was detected. Either their protection relies on other mechanisms because *mazF* is functional being only expressed in the uncured strain (Fig. S3). On the other hand, genes for RelB antitoxins were found both on plasmids and on the chromosome; however, no RelB-toxin genes were detected. So this antitoxin may contribute a greater defense against other bacteria possessing RelB toxins, possibly increasing its competitiveness and survival in



**Figure 1.** BlastKOALA results of functional gene-categories predicted in *Lactobacillus pentosus* MP-10 plasmids and their frequencies.

several ecological niches including gastrointestinal tract. This feature was mainly linked to plasmid being *relB* antitoxin gene over-exopressed in the uncured strain (Fig. S3). Zeta toxins, which are kinases that kill bacteria through global inhibition of peptidoglycan synthesis<sup>33</sup>, are detected both on the chromosome and also on plasmid DNA of *L. pentosus* MP-10, however no antitoxin was detected. Overall, *L. pentosus* MP-10 harbored in their plasmids incomplete toxin-antitoxin systems unlike what occur naturally in bacterial genomes, since several toxins or antitoxins were detected without self protection.

Data obtained by *in-silico* analysis suggests that plasmid-borne genes mediate the persistence of lactobacilli under olive fermentation conditions and enhance their probiotic properties; however, this hypothesis requires further studies for confirmation. As such, plasmid curing experiments carried out with L. pentosus MP-10 showed several differences between the uncured and the cured strains regarding metal tolerances, removal and mucin adhesion. However, plasmid curing did not affect their tolerance to gastro-intestinal conditions (e.g., acids and bile salts); neither their ability to grow under determined conditions (i.e., different pH intervals, bile salts or sodium chloride of 6.5%) nor their colony morphology were changed after plasmid curing (data not shown). However, at high concentration of chloride of 8% (commonly added to brines), L. pentosus pLPE-2 plasmid plays a key role in salt tolerance. In this sense, the results suggest that the plasmids did not govern the fermentation of carbohydrates under these conditions, however different results were obtained by Adeyemo and Onilude<sup>34</sup> which showed that plasmid curing had a significant negative effect on growth, physiological characteristics and colony morphology of L. plantarum isolated from fermented cereals. In this study, plasmids in L. pentosus MP-10 may confer a selective advantage, providing other physiological properties in certain environments such as gut and brines and thus allowing metal tolerance and removal, salt tolerance and adherence to mucin and thus their persistence in competitive ecological niches. Mucin adhesion declined in the cured L. pentosus MP-10 since a serine-rich adhesin for platelets precursor gene (sraP, detected in pLPE-2 plasmid) may be involved in mucin adhesion mechanisms similarly as reported by Hevia, et al.<sup>34</sup> for an extracellular serine/threonine-rich protein as a novel aggregation-promoting factor with affinity to mucin in Lactobacillus plantarum NCIMB 8826. The role of L. pentosus MP-10 plasmids in mucin adhesion was confirmed by relative expression gene analysis as reported by Pérez Montoro et al.<sup>35</sup>, since recA and pgm genes considered as potential biomarkers of mucin adhesion were over-expressed in the uncured strain (Fig. S3). However, auto-aggregation and co-aggregation with some pathogens were not changed after plasmid curing of L. pentosus MP-10.

With respect to metals, which are considered non-biodegradable and non-thermodegradable and are of high concern in both developing and developed countries because of their impact on the environment and health (water and food), the wild strain *L. pentosus* MP-10 showed greater tolerance to their increased concentrations (MICs higher than 1 mg/ml, except for cadmium and mercury) of iron, cobalt, copper, aluminium and zinc. This suggests that high contamination of metals in the environment from natural and anthropogenic sources<sup>36</sup> may be tolerable by the bacteria. The self-protective mechanisms displayed by *L. pentosus* MP-10 as a response to metals is promoted by their architecture (cell wall and membrane) and also by their resistance determinants located on the chromosome and the plasmids. Moreover, several chromosomally encoded cation transporters (e.g., encoded by *czcD* gene) have a predicted substrate range, including cadmium, cobalt and zinc; although the increased

				Gen length			Similarity to proteins in
Gene ID	Gene	Position	Strand	(bp)	Protein description	GO terms	Lactobacillus
XX999_00001	XX999_00001	99–314	-	216	hypothetical protein	_	100% identity in <i>L. plantarum</i> 90sk
XX999_00002	XX999_00002	435-803	_	369	DNA topoisomerase III	_	100% identity in <i>L</i> . paraplantarum DSM 10667
XX999_00003	topB_1	808-1116	-	309	DNA topoisomerase 3	Magnesium ion binding (MF); DNA binding (MF); DNA topoisomerase type I activity (MF); DNA topological change (biological_process); DNA recombination (BP); chromosome separation (BP)	100 identity in <i>L.</i> paraplantarum DSM 10667
XX999_00004	topB_2	1194–2567	-	1374	DNA topoisomerase 3	Magnesium ion binding (MF); DNA binding (MF); DNA topoisomerase type I activity (MF); DNA topological change (BP); DNA recombination (BP); chromosome separation (BP)	98% identity in <i>L. pentosus</i> IG1
XX999_00005	XX999_00005	2574-2984	-	411	hypothetical protein	_	100% identity in <i>L. plantarum</i> Lp1610
XX999_00006	XX999_00006	3000-3857	-	858	hypothetical protein	_	100% identity in <i>L. sakei</i> WiKim0063
XX999_00007	XX999_00007	3863-4237	-	375	hypothetical protein	-	100% identity in L. pentosus
XX999_00008	traG_1	4252-5796	_	1545	Conjugal transfer protein TraG	Conjugation (BP); DNA binding (MF); plasma membrane (CC); integral component of membrane (CC)	99% identity in <i>L.</i> <i>kefiranofaciens</i> subsp. <i>kefiranofaciens</i> DSM 5016
XX999_00009	XX999_00009	5840-6010	-	171	hypothetical protein	_	86% identity in <i>L. fermentum</i> MTCC 8711
XX999_00010	XX999_00010	6026-6496	-	471	hypothetical protein	—	97% identity in <i>L. paraplantarum</i>
XX999_00011	XX999_00011	6499–6867	-	369	hypothetical protein	_	91% identity in L. plantarum
XX999_00012	XX999_00012	6854-7471	-	618	hypothetical protein	_	99% identity in <i>L. brevis</i> DmCS_003
XX999_00013	XX999_00013	7486-8640	-	1155	Bacteriophage peptidoglycan hydrolase	_	99% identity in <i>L. brevis</i> KB290
XX999_00014	XX999_00014	8641-10059	-	1419	hypothetical protein	-	98% identity in <i>L. plantarum</i> Nizo2259
XX999_00015	XX999_00015	10052-12070	-	2019	AAA-like domain protein	_	99% identity in <i>L. parabuchneri</i> DSM 15352
XX999_00016	XX999_00016	12082-12741	-	660	hypothetical protein	_	100% identity in <i>L. plantarum</i> 2025
XX999_00017	XX999_00017	12710-13072	_	363	hypothetical protein	_	100% identity in <i>L. plantarum</i> CMPG5300
XX999_00018	XX999_00018	13093-13431	-	339	hypothetical protein	_	100% identity in <i>L. plantarum</i> Nizo2259
XX999_00019	XX999_00019	13433-14047	-	615	hypothetical protein	_	98% identity in <i>L. paracollinoides</i> DSM 15502
XX999_00020	XX999_00020	14061-14390	_	330	hypothetical protein	_	100% identity in <i>L. parakefiri</i> JCM 8573
XX999_00021	mobA_1	14473-16533	_	2061	Mobilization protein A	Conjugation (BP); DNA binding (MF); DNA- directed RNA polymerase activity (MF); DNA topoisomerase type I activity (MF); cytoplasm (CC); metal ion binding (MF)	100% identity in <i>L. pentosus</i>
XX999_00022	XX999_00022	16804-17013	+	210	hypothetical protein	_	100% identity in L. pentosus
XX999_00023	XX999_00023	17036-17314	+	279	hypothetical protein	-	100% identity in L.
XX999_00024	XX999_00024	17304-17984	-	681	Zeta toxin	-	100% identity in L.
XX999_00025	XX999_00025	17981-18172	-	192	hypothetical protein	-	100% identity in L.
XX999_00026	XX999_00026	18213-18494	-	282	RelB antitoxin	-	100% identity in <i>L</i> .
XX999_00027	XX999_00027	18899-19993	-	1095	hypothetical protein	-	100% identity in <i>L</i> .
XX999_00028	XX999_00028	20530-20805	-	276	hypothetical protein	-	100% identity in <i>L</i> .
XX999_00029	XX9999_00029	20808-21857	-	1050	StbA protein	-	100% identity in L.
XX999_00030	XX999_00030	22519-23808	-	1290	hypothetical protein	-	100% identity in <i>L</i> .
XX999_00031	dpnM	23823-24686	-	864	Modification methylase DpnIIA	Nucleic acid binding (MF); site-specific DNA-methyltransferase (adenine-specific) activity (MF); DNA restriction-modification system (BP)	100% identity in <i>L</i> .
XX999_00032	bin3_1	24835–25416	-	582	Putative transposon Tn552 DNA- invertase bin3	Recombinase activity (MF); DNA binding (MF); DNA integration (BP); transposition (BP)	
XX999_00033	XX999_00033	25526-26605	-	1080	FRG domain protein	-	
XX999_00034	hsrA_1	27276-28667	+	1392	putative transport protein HsrA	Plasma membrane (CC); integral component of membrane (CC); transmembrane transport (BP)	
XX999_00035	XX999_00035	28667-29329	+	663	putative hydrolase		
XX999_00036	XX999_00036	29319-29720	+	402	hypothetical protein	—	
Continued							

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_00037	XX999_00037	29820-30473	+	654	S-adenosyl-L-homocysteine hydrolase	Adenosylhomocysteinase activity (MF); cytoplasm (CC); one-carbon metabolic process (BP)	
XX999_00038	XX999_00038	31017-32141	-	1125	hypothetical protein	-	
XX999_00039	XX999_00039	32145-32360	-	216	hypothetical protein	-	
XX999_00040	topB_3	32482-34617	_	2136	DNA topoisomerase 3	Magnesium ion binding (MF); DNA binding (MF); DNA topological change (BP); DNA recombination (BP); chromosome separation (BP)	
XX999_00041	XX999_00041	34624-35034	-	411	hypothetical protein	-	
XX999_00042	XX999_00042	35050-35907	-	858	hypothetical protein	-	
XX999_00043	XX999_00043	35913-36287	-	375	hypothetical protein	-	
XX999_00044	traG_2	36302-37846	-	1545	Conjugal transfer protein TraG	Conjugation (BP); DNA binding (MF); plasma membrane (CC); integral component of membrane (CC)	
XX999_00045	XX999_00045	37890-38060	-	171	hypothetical protein	-	
XX999_00046	XX999_00046	38076-38546	-	471	hypothetical protein	-	
XX999_00047	XX999_00047	38549-38917	-	369	hypothetical protein	-	
XX999_00048	XX999_00048	38904-39521	-	618	hypothetical protein	-	
XX999_00049	XX999_00049	39536-40690	-	1155	Bacteriophage peptidoglycan hydrolase	-	
XX999_00050	XX999_00050	40691-42004	-	1314	hypothetical protein	—	
XX999_00051	XX999_00051	42001-42108	-	108	hypothetical protein	-	
XX999_00052	XX999_00052	42101-43795	-	1695	AAA-like domain protein	Conjugation (BP); plasma membrane (CC)	

**Table 5.** Genes determined in pLPE-4 plasmid of *Lactobacillus pentosus* MP-10 isolated from naturally fermented Aloreña table olives. BP, biological process; CC, celular component; MF, molecular function.

resistance towards different metals are displayed by plasmids (especially the pLPE-5 plasmid). Similar results were obtained by van Kranenburg *et al.*<sup>22</sup>, which reported that the plasmid-borne (pWCFS103) *cadC* gene coding for a transcription regulator of the cadmium operon was responsible of the increased resistance to cadmium in *L. plantarum* WCFS1. Furthermore, the synteny of *ars* genes in both *L. pentosus* MP-10 and *L. plantarum* WCFS1<sup>22</sup> was similar suggesting their evolutionary relatedness. Arsenic and cadmium are among the most toxic elements widely ocurring in the environment, often a threat to food and water supply. Arsenic is known as a group A "known" carcinogen according to the United States Environmental Protection Agency (USEPA) and contributes to a range of other illnesses such as cardiovascular and peripheral vascular diseases, neurological disorders, diabetes mellitus and chronic kidney disease<sup>37–39</sup>. Detoxification of this metal was earlier established by bacteria. Thus, tolerance of *L. pentosus* MP-10 is necessary to prevent damage to their cells.

The ability of *L. pentosus* MP-10 to bind different metals was demonstrated by SEM and EDX analysis. This is of great importance with regards to their application as an adjunct to improve food safety and quality by bioquenching metals and probiotically reduce metal toxicity among human intestinal microbiota and thus protecting the host<sup>40</sup>. Also, we demonstrated that *L. pentosus* MP-10 contributed to metal removal, especially mercury and cadmium (81 and 67%, respectively).

Metal- and antibiotic-resistance genes often co-exist on the same plasmid, however in this case, we did not find any genes coding for clindamycin resistance on plasmids, which was the only antibiotic with different susceptibility after plasmid curing. Thus, clindamycin resistance in *L. pentosus* MP-10 may rely on other plasmid-associated genes that we could not deciphered yet.

#### Conclusions

*In-silico* analysis of *L. pentosus* MP-10 plasmids suggests that plasmid-borne genes mediate the persistence of lactobacilli under olive-fermentation conditions and enhance their probiotic properties with genes encoding for carbohydrate metabolism, defense mechanisms, metal tolerance and mobilization increasing subsequently its competitiveness and survival in several ecological niches. Plasmid curing demonstrated the role of plasmids in the increased metal tolerance, and bioremoval of several metals (e.g., iron, aluminium, cobalt, copper, zinc, cadmium and mercury). This probiotic property by *L. pentosus* MP-10 should be exploited to detoxify metals in intestines; basically they could bioquench the metals in the gut thus reducing their toxic exposure to humans and animals, in the food matix and in environmental bioremediation.

#### **Materials and Methods**

**Bacteria and growth conditions.** Lactobacillus pentosus MP-10 isolated from naturally-fermented Aloreña green table olives<sup>1</sup> were cultured in de Man Rogosa and Sharpe (MRS) broth (Fluka, Madrid, Spain) at 37 °C for 24 h. Pathogenic bacteria used in this study included *Listeria innocua* CECT 910, *Staphylococcus aureus* CECT 4468, *Escherichia coli* CCUG 47553, and *Salmonella* Enteritidis UJ3449, which were cultured in Tryptone

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03553	XX999_03553	763-1230	+	468	Hypothetical protein	_	95% identity in <i>L. plantarum</i> Nizo2814
XX999_03554	XX999_03554	1634-1915	+	282	Bifunctional antitoxin/ transcriptional repressor RelB	DNA binding (MF); transcription, DNA-templated (BP); regulation of transcription, DNA-templated (BP)	99% identity in <i>L. plantarum</i> 16
XX999_03555	XX999_03555	1956–2147	+	192	Hypothetical protein	_	98% identity in <i>L. plantarum</i> Nizo2814
XX999_03556	XX999_03556	2224-2502	-	279	Hypothetical protein	_	100% identity in <i>L. farraginis</i> DSM 18382
XX999_03557	XX999_03557	2525-2734	-	210	Hypothetical protein	_	100% identity in <i>L. diolivorans</i> DSM 14421
XX999_03558	mobA_6	3004–5061	+	2058	Mobilization protein A	Conjugation (BP); DNA binding (MF); DNA-directed RNA polymerase activity (MF); DNA topoisomerase type I activity (MF); cytoplasm (CC); metal ion binding (MF)	99% identity in <i>L. plantarum</i> 2025
XX999_03559	XX999_03559	5168-5461	+	294	Hypothetical protein	-	100% identity in <i>L. plantarum</i> 2025
XX999_03560	XX999_03560	5501-6115	+	615	Hypothetical protein	—	100% identity in L. plantarum
XX999_03561	XX999_03561	6117-6455	+	339	Hypothetical protein	—	100% identity in L. plantarum 2025
XX999_03562	XX999_03562	6476-6838	+	363	Hypothetical protein	_	100% identity in <i>L. plantarum</i> IPLA88
XX999_03563	XX999_03563	6807-7466	+	660	Hypothetical protein	_	100% identity in <i>L. plantarum</i> CMPG5300
XX999_03564	XX999_03564	7478-9496	+	2019	AAA-like domain protein	Conjugation (BP); plasma membrane (CC)	99% identity in <i>L. plantarum</i>
XX999_03565	XX999_03565	9489-10904	+	1416	Hypothetical protein	_	99% identity in <i>L. plantarum</i> TMW 1.25 pL125–4 plasmid
XX999_03566	XX999_03566	10906-12075	+	1170	Bacteriophage peptidoglycan hydrolase	_	99% identity in <i>L. paraplantarum</i>
XX999_03567	XX999_03567	12090-12707	+	618	Hypothetical protein	_	100% identity in L. plantarum 2025
XX999_03568	XX999_03568	12685-13059	+	375	Hypothetical protein	_	99% identity in <i>L. plantarum</i> Nizo2814
XX999_03569	XX999_03569	13060-13518	+	459	Hypothetical protein	_	100% identity in L. plantarum 2025
XX999_03570	traG_3	13515-15044	+	1530	Conjugal transfer protein TraG	Conjugation (BP); DNA binding (MF); plasma membrane (CC); integral component of membrane (CC)	99% identity in <i>L. plantarum</i>
XX999_03571	XX999_03571	15057-15470	+	414	Hypothetical protein	_	99% identity in <i>L. plantarum</i> Nizo2814
XX999_03572	XX999_03572	15483-16352	+	870	Hypothetical protein	_	99% identity in <i>L. L. plantarum</i> Nizo2814
XX999_03573	topB_5	16369-18507	+	2139	DNA topoisomerase 3	Magnesium ion binding (MF); DNA binding (MF); DNA topoisomerase type I activity (MF); DNA topological change (BP); DNA recombination (BP); chromosome separation (BP)	98% identity in <i>L. plantarum</i> SRCM101060
XX999_03574	XX999_03574	18629–18844	+	216	Hypothetical protein	_	100% identity in <i>L. plantarum</i> Nizo1838
XX999_03575	XX999_03575	18848-19039	+	192	Hypothetical protein	_	83% identity in <i>L. collinoides</i> 237
XX999_03576	XX999_03576	18993-19250	+	258	Hypothetical protein	_	99% identity in <i>L. plantarum</i> Nizo2029
XX999_03577	axeA1_3	19530-20243	+	714	Acetylxylan esterase precursor	Xylan catabolic process (BP); acetylxylan esterase activity (MF)	100% identity in <i>L. plantarum</i> Nizo2029
XX999_03578	XX999_03578	20326-20994	_	669	Integrase core domain protein	_	100% identity in <i>L. tucceti</i> DSM 20183
XX999_03579	XX999_03579	20957-21163	-	207	Hypothetical protein		100% identity in <i>L. brevis</i> 47 f
XX999_03580	ldh_7	21343-22305	+	963	L-lactate dehydrogenase	L-lactate dehydrogenase activity (MF); cytoplasm (CC); glycolytic process (BP); cellular carbohydrate metabolic process (BP)	100% identity in <i>L. plantarum</i> Nizo2029
XX999_03581	XX999_03581	22735-22869	+	135	Hypothetical protein	_	95% identity in <i>L. backii</i> TMW 1.1991
XX999_03582	XX999_03582	23089-23295	-	207	Hypothetical protein	-	100% identity in <i>L. brevis</i> 47 f
XX999_03583	ldh_8	23475-24437	+	963	L-lactate dehydrogenase	L-lactate dehydrogenase activity (MF); cytoplasm (CC); glycolytic process (BP); cellular carbohydrate metabolic process (BP)	100% identity in <i>L. plantarum</i> Nizo2029
XX999_03584	XX999_03584	24867-25001	+	135	Hypothetical protein	_	95% identity in <i>L. backii</i> TMW 1.1991
XX999_03585	XX999_03585	24998-25501	-	504	Transposase DDE domain protein	_	100% identity in <i>L. plantarum</i> subsp. <i>plantarum</i> P-8
Continued							

Gene ID	Gene	Position	Strand	Gen length (bp)	Protein description	GO terms	Similarity to proteins in Lactobacillus
XX999_03586	XX999_03586	25459-25797	-	339	Hypothetical protein	_	98% identity in <i>L. plantarum</i> IPLA88
XX999_03587	XX999_03587	26046-26384	-	339	Hypothetical protein	_	100% identity in <i>L. plantarum</i> IPLA88
XX999_03588	XX999_03588	26499-27041	+	543	Hypothetical protein	-	100% identity in L. pentosus
XX999_03589	XX999_03589	27059-27859	+	801	Adenylate and Guanylate cyclase catalytic domain protein	_	100% identity in <i>L. pentosus</i>
XX999_03590	ubiE_3	27925-28629	+	705	Demethylmenaquinone methyltransferase	Methyltransferase activity (MF); menaquinone biosynthetic process (BP)	98% identity in <i>L. parakefiri</i> DSM 10551
XX999_03591	XX999_03591	28680-28781	+	102	Hypothetical protein	_	100% identity in <i>L. parakefiri</i> DSM 10551
XX999_03592	XX999_03592	28794-29021	+	228	ASCH domain protein	_	100% identity in <i>L. plantarum</i> Nizo2029
XX999_03593	XX999_03593	29337-30368	+	1032	Integrase core domain protein	_	99% identity in <i>L. plantarum</i> WCFS1
XX999_03594	XX999_03594	30453-31067	-	615	Cadmium resistance transporter	_	100% identity in <i>L. plantarum</i> SF2A35B
XX999_03595	cadC	31069-31437	_	369	putative positive regulator of cadmium resistance	Sequence-specific DNA binding transcription factor activity (MF); regulation of transcription, DNA- templated (BP)	100% identity in <i>L. plantarum</i> WCFS1
XX999_03596	npr_2	31787-33187	_	1401	NADH peroxidase	NADH peroxidase activity (MF); cell redox homeostasis (BP); flavin adenine dinucleotide binding (MF)	100% identity in <i>L. plantarum</i> Nizo1839
XX999_03597	XX999_03597	33361-33615	+	255	Hypothetical protein	—	_
XX999_03598	XX999_03598	33533-33862	-	330	Hypothetical protein	_	100% identity in <i>L. plantarum</i> WCFS1
XX999_03599	arsB	33879-35174	_	1296	Arsenical pump membrane protein	Plasma membrane (CC); arsenite transmembrane transporter activity (MF); arsenite transport (BP); integral component of membrane (CC); response to arsenic-containing substance (BP)	99% identity in <i>L. plantarum</i> SF2A35B
XX999_03600	arsA	35233-36963	_	1731	Arsenical pump-driving ATPase	ATP binding (MF); arsenite- transmembrane transporting ATPase activity (MF); detoxification of arsenic-containing substance (BP)	100% identity in <i>L. plantarum</i> WCFS1
XX999_03601	arsD	37047-37409	_	363	Arsenical resistance operon trans- acting repressor ArsD	DNA binding (MF); transcription, DNA-templated (BP); negative regulation of transcription, DNA- templated (BP); response to arsenic- containing substance (BP)	100% identity in <i>L. plantarum</i> WCFS1
XX999_03602	arsR_3	37396–37755	_	360	Arsenical resistance operon repressor	DNA binding (MF); sequence- specific DNA binding transcription factor activity (MF); intracellular (CC); transcription, DNA-templated (BP); response to arsenic-containing substance (BP)	100% identity in <i>L. plantarum</i> WCFS1
XX999_03603	pinR	39098-39679	+	582	Putative DNA-invertase from lambdoid prophage Rac	Recombinase activity (MF); DNA binding (MF); DNA integration (BP)	100% identity in <i>L. backii</i> TMW 1.1992
XX999_03604	bin3_3	40077-40709	+	633	Putative transposon Tn552 DNA- invertase bin3	Recombinase activity (MF); DNA binding (MF); DNA integration (BP); transposition (BP)	100% identity in <i>L. backii</i> TMW 1.1992
XX999_03605	XX999_03605	40806-41168	+	363	Hypothetical protein	_	98% identity in <i>L. backii</i> TMW 1.1992
XX999_03606	XX999_03606	41577-41990	-	414	Hypothetical protein	_	100% identity in <i>L. backii</i> TMW 1.1992
XX999_03607	parA	41987-42871	-	885	Chromosome partitioning protein ParA	ATP binding (MF); chromosome segregation (BP)	100% identity in <i>L. hokkaidonensis</i> JCM 18461
XX999_03608	XX999_03608	43459-44988	+	1530	Hypothetical protein	-	100% identity in <i>L. backii</i> TMW 1.1992
XX999_03609	XX999_03609	45128-45235	-	108	Hypothetical protein	-	-
XX999_03610	XX999_03610	45885-46475	-	591	Transposase, Mutator family	_	99% identity in <i>L. brevis</i> TMW 1.2113

**Table 6.** Genes determined in pLPE-5 plasmid of *Lactobacillus pentosus* MP-10 isolated from naturallyfermented Aloreña table olives. BP, biological process; CC, celular component; MF, molecular function.

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Soya Broth (TSB; Fluka, Madrid, Spain) at 37 °C for 24 h. Cultures were maintained in 20% glycerol at -20 °C and -80 °C for short- and long-term storage, respectively.

*In silico* analysis of *L. pentosus* MP-10 plasmid sequences. The genome sequence of *L. pentosus* MP-10 consisted of a single circular chromosome of 3,698,214 bp, with an estimated mol% G + C content of

pLPE-5 plasmid of L. pentosus MP-10



#### pWCFS103 plasmid of L. plantarum WCFS1

**Figure 2.** MAUVE visualization of the alignment of the pLPE-5 plasmid from *L. pentosus* MP-10 with the pWCFS103 plasmid from *L. plantarum* WCFS1. Arsenic- and cadmium-resistance genes are indicated.

		MIC (µg/ml)	
		L. pentosus MP-10 (uncured)	L. pentosus MP- 10C (cured)
	Amoxicillin	0.2	0.2
	Ampicillin	2	2
	Chloramphenicol	8	8
	Ciprofloxacin	16	16
	Clindamycin	2	0.1
	Gentamicin	0.1	0.1
Antibiotic	Kanamycin	4	4
	Streptomycin	4	4
	Teicoplanin	256	256
	Tetracycline	16	16
	Trimethoprim	0.125	0.125
	Trimethoprim/sulfometoxazole	0.125/2.38	0.125/2.38
	Vancomycin	2048	2048
Dianida	Benzalconium Chloride	2	2
biocide	Triclosan	32	32
Phenolic componds		$>2 \times 10^5$	$>2 \times 10^{5}$
	Auto-aggregation (%)	$20.58 \pm 2.54^{a}$	$13.49 \pm 0.54^a$
	Co-aggregation + L. innocua CECT 910 (%)	$32.87 \pm 2.14^a$	36.13±2.33 <sup>a</sup>
	Co-aggregation + S. aureus CECT 4468 (%)	$28.61 \pm 0.99^{a}$	$28.69 \pm 0.72^{a}$
	Co-aggregation + E. coli CCUG 47553 (%)	$16.14 \pm 2.09^{a}$	$14.15 \pm 3.24^{a}$
	Co-aggregation + S. Enteritidis UJ 3449 (%)	$12.27 \pm 1.50^{a}$	$13.17 \pm 2.87^a$
	Acid tolerance pH 2.0 (%)	$100 \pm 0.04^{a}$	$100 \pm 0.01^{a}$
Probiotic properties	Acid tolerance pH 2.5 (%)	$100 \pm 0.03^{a}$	$100 \pm 0.02^{a}$
	Acid tolerance pH 3.0 (%)	$100 \pm 0.01^{a}$	$100 \pm 0.02^{a}$
	Bile tolerance at 1%	+	+
	Bile tolerance at 2%	+	+
	Bile tolerance at 3%	+	+
	Bile tolerance at 4%	+	+
	Mucin adhesion (%)	55.93±0.34ª	51.92±1.06 <sup>b</sup>

**Table 7.** Antibiotic and biodice susceptibility, and probiotic properties of cured and uncured *L. pentosus* MP-10 isolated from Aloreña Green table olives.  $\pm$ SD, standard deviations of three independent experiments. \*Different lowercase letters represent significant differences according to 2-sided Tukey's HSD between strains (*p* < 0.05). +, Presence of growth in MRS-agar with different concentrations of bile salts.

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	MIC (µg/ml)	
Metal	<i>L. pentosus</i> MP-10 (uncured)	L. pentosus MP-10C (cured)
Mercury (Hg)	2	1
Cobalt (Co)	2048	2048
Copper (Cu)	2048	2048
Zinc (Zn)	1024	1024
Aluminium (Al)	2048	2048
Iron (Fe)	4096	4096
Cadmium (Cd)	8	1
	% heavy metal removed	
Mercury (Hg)	81.74±2.04 <sup>a</sup>	$63.68 \pm 1.09^{b}$
Cobalt (Co)	$10.65 \pm 1.03^{a}$	$10.18 \pm 0.67^{b}$
Copper (Cu)	$11.92 \pm 0.45^{a}$	$7.41\pm0.89^{\mathrm{b}}$
Zinc (Zn)	$37.03 \pm 1.02^{a}$	34.73±2.0 <sup>b</sup>
Aluminium (Al)	57.14±0.99ª	49.92±0.72 <sup>b</sup>
Iron (Fe)	$21.04 \pm 1.50^{a}$	$14.36 \pm 0.78^{b}$
Cadmium (Cd)	67.10±0.88 <sup>a</sup>	$55.40 \pm 0.67^{b}$

**Table 8.** Tolerance of cured and uncured *L. pentosus* MP-10 isolated from Aloreña Green table olives to heavy metals.  $\pm$ SD, standard deviations of three independent experiments. \*Different lowercase letters represent significant differences according to 2-sided Tukey's HSD between strains (p < 0.05).

46.32% and 5 plasmids ranging 29–46 kb (accession numbers FLYG01000001 to FLYG01000006) were annotated using the Prokka annotation pipeline, version 1.11 (Seemann, 2014) as previously reported by Abriouel, *et al.*<sup>12</sup>. The predicted CDSs of plasmids<sup>2,12</sup> were annotated by using BLAST (Basic Local Alignment Search Tool) and the associated GO (Gene Ontology) terms were obtained by using Swiss-Prot database.

The general metabolic pathways of *L. pentosus* MP-10 plasmids were reconstructed using BlastKOALA (last updated March 4, 2016) as part of the KEGG (Kyoto Encyclopedia of Genes and Genome) tool in the pathway database (http://www.genome.jp/kegg/pathway.html) for annotating genomes; here, we used the annotated genes predicted in each *L. pentosus* MP-10 plasmid as the input query.

To evaluate the alignment and the synteny of genes between the *L. pentosus* MP-10 and *L. plantarum* WCFS1 plasmid data sets, comparison was done by using Mauve algorithm in Lasergene's MegAlign Pro software (Lasergene 14).

*In vitro* analysis of *L. pentosus* MP-10 plasmid properties. *Plasmid curing.* First, we determined the minimum inhibitory concentrations (MIC) of acridine orange (AO) to *L. pentosus* MP-10 using the broth micro-dilution method. Overnight cultures, grown in MRS broth at 37 °C for 24 h, were diluted 1/10 (v/v) in fresh MRS broth and 20 µl were added to each well of 96-well microtiter plates. 180 µl of MRS broth supplemented with AO at different concentrations (12.5–400 µg/ml) were then added to the wells and incubated at 37 °C under aerobic conditions for 24 h. Bacterial growth was evaluated by the presence of turbidity. MIC was defined as the lowest concentration of AO that inhibited visible growth. Each experiment was done in triplicate.

Plasmid curing (eliminating the plasmid from cells) of *L. pentosus* MP-10 was done as described by Adeyemo and Onilude<sup>41</sup> with some modifications. Briefly, MRS broth (4 ml) supplemented with the sub-MIC of AO, as determined in this study, was inoculated with a selected colony of *L. pentosus* MP-10 grown onto MRS agar; then the cultures were incubated at 37 °C for 72 h. Serial dilutions of bacterial cultures in NaCl (0.85%) were plated onto MRS agar, and the resulting colonies, obtained after incubation for 48 h at 37 °C, were inoculated into MRS broth to obtain a pure culture. Cultures were maintained in 20% glycerol at -20 °C and -80 °C for short- and long-term storage, respectively.

To confirm that the resulting colonies were cured of plasmids, bacterial cultures (uncured and cured) were subjected to plasmid isolation as described by Abriouel, *et al.*<sup>42</sup> and then visualized on 0.8% agarose gel electro-phoresis (iNtRON Biotechnology) in 1xTBE (Tris-Boric acid-EDTA) buffer.

For additional confirmation, total genomic DNA (uncured and cured strains) was extracted using DNA Extraction Kit (Xtrem Biotech SL, Spain) according to the manufacturer's instructions and tested for plasmid-borne genes. DNA quantification and quality assessment were carried out using a NanoDrop 2000 spectrophotometer (Thermo Scientific). DNAs were frozen at -20 °C until required and then subjected to PCR amplification of genes harboured by pLPE5, the biggest plasmid detected in *L. pentosus* MP-10. The PCR primers were designed in this study: Ars-pl5-F (5'-ATTATTTTGATCTCATTGATTTT-3') and Ars-pl5-R (5'-TGAATAAACGAAACGGGAATGT-3'), yielding an amplicon of 570 bp. The 50µl PCR mixture contained 20 ng of DNA,  $0.5 \mu$ m of each primer (Ars-pl5-F and Ars-pl5-R), 200 µm of each deoxyribonucleoside triphosphate (Bioline), and 1 U of *Taq* DNA polymerase in 1X buffer according to the manufacturer's instructions (Bioline). PCR was performed under the following conditions: one cycle at 95 °C for 3 min, 35 cycles at 95 °C for 30 s, 58 °C for 30 s, and 72 °C for 1 min and the final hold for 3 min at 72 °C. Analysis of PCR products was done by electrophoresis through a 1% agarose gel electrophoresis in 1xTBE (Tris-Boric acid-EDTA).



**Figure 3.** SEM (**A**,**C**,**E**,**G**,**I**,**K**,**M**,**O**) and EDX (**B**,**D**,**F**,**H**,**J**,**L**,**N**,**P**) analysis of uncured *L*. *pentosus* MP-10 without metal (**A**,**B**) and with Al (**C**,**D**), Cd (**E**,**F**), Co (**G**,**H**), Cu (**I**,**J**), Fe (**K**,**L**), Hg (**M**,**N**) and Zn (**O**,**P**).

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**Effect of plasmid curing on growth, safety and functional properties of** *L. pentosus* **MP-10.** *Growth properties.* To test whether there is any differences in growth between the uncured and the cured *L. pentosus* **MP-10** strains, MRS broth was inoculated (1% v/v) with overnight cultures of each strain and then incubated at 37 °C for 24 h. Growth rates (OD<sub>600nm</sub>) were measured each hour using Microtiter plate reader (iMark Microplate Absorbance Reader, Bio-Rad instrument). Additionally, we measured pH at different time intervals (following 0, 8 and 24 h of incubation at 37 °C).

To determine the effect of pH on the growth of both strains, MRS broth was adjusted to different pH ranges (1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, 5.0, 5.5, 6.0, 6.5 and 7.0) with phosphate buffer, and they were inoculated (1% v/v) overnight cultures of both strains and then incubated at 37 °C for 24 h, as described above.

To test whether brine conditions had an effect on the growth of the plasmid-cured versus uncured *L. pentosus* MP-10 strains in MRS broths under the following experimental conditions: unsupplemented vs. those supplemented with either 6.5% (or high concentration of 8%) NaCl or phenolic compounds, or modified MRS broth (without glucose) added with xylan (5 g/l) were inoculated with both strains as described above. Phenolic compounds were obtained from freshly pulverized olive leaves using RETSCH laboratory ball mills (Retsh MM 400). The leaf extracts were resuspended in LSM broth, centrifuged and the resulting supernatant was filtered ( $0.45 \mu m$ ) and added at different concentrations (0.780 to 200 mg/ml) to MRS broth. The cultures were incubated at 37 °C for 24 h and the OD<sub>600nm</sub> was measured as described above.

In all cases, experiments were done in triplicate.

*Evaluation of metal tolerance.* The sensitivity of both *L. pentosus* strains (MP-10 and MP-10C (cured)) towards metals: cadmium (CdSO<sub>4</sub>·8/3H<sub>2</sub>O), cobalt (CoCl<sub>2</sub>), copper (CuCl<sub>2</sub>·2H<sub>2</sub>O), iron (FeSO<sub>4</sub>·7H<sub>2</sub>O), mercury (HgCl<sub>2</sub>), aluminium (Al<sub>2</sub>O<sub>3</sub>), or zinc (ZnCl<sub>2</sub>) was tested in LSM broth supplemented with 0 to 10 mg/ml of each metal and then inoculated with 2% (v/v) of an overnight culture of each strain. After 24 h of incubation at 37 °C, the MIC from each metal exposure was determined as described above, which corresponded to the lowest concentration that completely inhibited visible growth.

To analyse the removal of metals by cured and uncured *L. pentosus* MP-10, MRS broth supplemented with  $\frac{1}{2}$ MIC of each metal was inoculated with 2% (v/v) of an overnight culture of each strain and then incubated 24 h at 37 °C. After incubation, the bacterial cells were removed by centrifugation and kept for the subsequent examination of metal sorption. The resulting supernatants were filter sterilized using a 0.22 µm filter (Millipore, Spain) and then used to check metal removal. MRS broth added either with different metals (with  $\frac{1}{2}$ MIC) or not were used as positive and negative controls, respectively. The positive controls (MRS broth with individual metal added: Fe at 2 mg/ml; Al, Co and Cu at 1 mg/ml; Zn at 0.5 mg/ml; Cd at 4 µg/ml and 0.5 µg/ml; and Hg at 1 µg/ml and 0.5 µg/ml) were considered "100%" baselines to calculate relative metal removal rates (as a percentage).

Metal concentrations were measured using 7900 ICP-Mass Spectrometer (Agilent, USA) with graphite tube atomizer and autosampler, a superior matrix tolerance and advanced collision/reaction cell (CRC) technology to remove the polyatomic interferences that can affect some of the trace elements. The spectrometer software was Agilent ICP-MS MassHunter Work Station, which provides simple autotuning functions, and a Method Wizard automates the method setup process.

Biosorption of metals by *L. pentosus* MP-10 was further examined using scanning electron microscope (SEM) coupled with energy dispersive X-ray spectroscopy before and after metal uptake. For this, a drop of the bacterial pellet, which had been previously exposed to a metals (as previously described), were disposed into microporous capsules (ANAME, Spain), dried and then dehydrated in a series of 20, 40, 60, 80, and 100% ethanol solutions (15 min each) before suspension in acetone for 1 h. After this, the capsules were subjected to critical-point drying before examination by SEM (FESEM, MERLIN de Carl Zeiss, Oxford).

*Safety and probiotic properties.* To determine differences in antimicrobial (antibiotic and biocide) susceptibility of *L. pentosus* MP-10C versus wild strain, we determined the MIC of several antimicrobials following the method previously described by Casado Muñoz, *et al.*<sup>42,43</sup> using LSM broth (Oxoid).

To determine if plasmids further play a role in several probiotic peroperties, we analyzed acid- and bile- tolerances, auto-aggregation, co-aggregation with pathogens (*L. innocua* CECT 910, *S. aureus* CECT 4468, *E. coli* CCUG 47553, and *S.* Enteritidis UJ3449) and mucin adhesion in both *L. pentosus* strains (MP-10 and MP-10C) according to the methods reported by Pérez Montoro *et al.*<sup>35</sup>.

*Gene expression analysis.* To analyse the role of plasmid in several metabolic and probiotic properties, both the uncured and cured L. pentosus strains were subjected to RNA extraction using Direct-zol<sup>TM</sup> RNA Miniprep (Zymo Research, California, USA) according to the manufacturer's instructions. RNA quantification and quality assessment were carried out by using a NanoDrop 2000 spectrophotometer (Thermo Scientific). RNAs were adjusted to a concentration of 500 ng/ml and frozen at -80 °C until required for analysis.

The expression of selected genes (Table S1) was determined by quantitative, real-time PCR (qRT-PCR) using SensiFASTTM SYBR & Fluorescein One-Step Kit (BIOLINE) as reported in Pérez Montoro *et al.*<sup>35</sup>.

**Statistical analysis.** All analyses were performed in triplicate. Statistical descriptors were calculated using Excel 2007 (Microsoft Corporation, Redmond, Washington, US), e.g., determining averages and standard deviations. Statistical comparison of growth and probiotic properties assays were conducted by analysis of variance (*ANOVA*) using Statgraphics Centurion XVI software (Statpoint Technologie, Warrenton, Virginia, US). The same software was used to perform Shapiro–Wilk and the Levene tests to check data normality and to perform 2-sided Tukey's multiple contrast to determine the pair-wise differences between strains. Level of significance was set at P < 0.05.

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#### Author Contributions

H.A. and N.B. designed the experiments. H.A., N.B. and C.K. wrote the main manuscript text. H.A., B.P.M., J.F.O. and L.L.L. did the experiments and prepared figures and tables. All authors reviewed the manuscript.

#### Additional Information

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