Mini-Review	Making sense of quorum sensing in lactobacilli: a special focus on <i>Lactobacillus plantarum</i> WCFS1							
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	<i>In silico</i> identification criteria were defined to predict if genes encoding histidine protein kinases (HPKs) and response regulators (RRs) could be part of peptide-based quorum sensing (QS) two-component regulatory systems (QS-TCSs) in Firmicutes. These criteria were used to screen HPKs and RRs annotated on the completed genome sequences of <i>Lactobacillus</i> species, and several (putative) QS-TCSs were identified in this way. The five peptide-based QS-TCSs that were predicted on the <i>Lactobacillus plantarum</i> WCFS1 genome were further analysed to test their (QS) functionality. Four of these systems contained an upstream gene encoding a putative autoinducing peptide (AIP), of which two were preceded by a double-glycine-type leader peptide. One of these was identical to the <i>plnABCD</i> regulatory system of <i>L. plantarum</i> C11 and was shown to regulate plantaricin production in <i>L. plantarum</i> WCFS1. The third TCS was designated <i>lamBDCA</i> for <i>Lactobacillus agr</i> -like module, where the <i>lamD</i> gene was shown to encode a cyclic thiolactone peptide. The fourth TCS was paralogous to the <i>lam</i> system and contained a putative AIP-encoding gene but lacked the <i>lamB</i> gene. Finally, a genetically separated orphan HPK and RR that showed clear peptide-based QS characteristics could form a fifth peptide-based QS-TCS. The predicted presence of multiple (peptide-based) QS-TCSs in some lactobacilli and in particular in <i>L. plantarum</i> might be a reflection of the ability of these species to persist in a diverse range of ecological niches.							

Two-component regulatory systems in lactic acid bacteria

The lactic acid bacteria (LAB) comprise a diverse group of Gram-positive bacteria that are applied in the production of fermented food products such as dairy, meat and vegetable products (Caplice & Fitzgerald, 1999). Several LAB are also found in the gastrointestinal tracts of humans and other animals (Vaughan *et al.*, 2002). Among the LAB the genus *Lactobacillus* forms a large and diverse group, and several *Lactobacillus* strains are considered to exert health-promoting effects in man and animals (Ouwehand *et al.*, 2002). The genome sequences of a number of *Lactobacillus* species from different ecological niches are currently available (Makarova

et al., 2006; Makarova & Koonin, 2007; Siezen et al., 2004). This enables a comparative genomics analysis of lactobacilli that are restricted to a specific niche and have limited physiological abilities, such as Lactobacillus johnsonii in the human gastrointestinal tract (Pridmore et al., 2004) and Lactobacillus delbrueckii subsp. bulgaricus in dairy products (Van de Guchte et al., 2006), with more adaptable species such as Lactobacillus plantarum (Kleerebezem et al., 2003), which is found in fermented food products, on plant material (Caplice & Fitzgerald, 1999) and as a natural inhabitant of the human gastrointestinal tract (Ahrne et al., 1998). To allow for efficient colonization and persistence or effective adaptation to changing environmental conditions, lactobacilli require sensory systems to detect (specific) environmental signals. In bacteria, this function is commonly mediated by two-component regulatory systems (TCSs), which consist of a membrane-located histidine protein kinase (HPK) that monitors one or more environmental factors, and a cytoplasmic response regulator (RR), which modulates expression of specific genes. The HPK and RR

Abbreviations: AIP, autoinducing peptide; HPK, histidine protein kinase; QS, quorum sensing; RR, response regulator; TCS, two-component system; TMS, transmembrane segment.

Three supplementary figures are available with the online version of this paper.

function as a phospho-relay signal-transduction system (Fig. 1), and the genes encoding the cognate HPK and RR are generally organized in an operon structure (Hoch & Silhavy, 1995). TCSs monitor and respond to changes in environmental conditions such as osmolarity, nutrient availability (C, N, P) or temperature (Hoch & Silhavy, 1995). However, in Gram-positive species, TCSs are also known to respond to specific secreted signalling molecules involved in quorum sensing (QS) (Sturme *et al.*, 2002).

QS-TCSs in Gram-positive bacteria regulate the expression of genes involved in diverse functions such as virulence, competence or bacteriocin production (Kleerebezem *et al.*, 1997). This modulation is done in a coordinated and celldensity-dependent manner, using specific autoinducing signalling peptides (AIPs) that are often post-translationally modified and exported by dedicated transport systems (Ansaldi & Dubnau, 2004; Håvarstein *et al.*, 1995; Zhang & Ji, 2004), and sensed by responsive cells via dedicated HPKs. Bacteria may contain multiple QS-TCSs, underlining the importance of intercellular communication.

This mini-review describes *in silico* criteria for the identification of (putative) peptide-based QS-TCSs in lactobacilli and other Firmicutes. In addition, it describes the application of those criteria in the identification of QS-TCSs in the model organism *L. plantarum* WCFS1 (Kleerebezem *et al.*, 2003). Detailed analysis and experimental confirmation of the functionality of predicted peptide-based QS-TCSs in *L. plantarum* WCFS1 is also discussed.



Fig. 1. General signal transduction mechanism of two-component regulatory systems. The cellular localization and sequence motifs characteristic of two-component histidine protein kinases of the HPK₁₀ subfamily (membrane-bound) and response regulators (cytoplasmic) are depicted (see also Table 1). TMS, transmembrane segments I–VII. Homology boxes of the HPK₁₀ subfamily as described in the text are: H, X, N and G. Conserved residues in the N-terminal CheY-like receiver domain of response regulators as described in the text are DD, D, and K.

Protein architecture of peptide-based QS-TCSs

A distinctive functional feature of TCSs is the ability to transfer a phosphoryl group from ATP to a receptor protein. As a result, the two 'components' of the system contain characteristic functional domains. The HPKs have a conserved C-terminal ATP-binding domain in which the phosphoryl-accepting histidine residue is located, and related to that they also have highly conserved clusters of residues called homology boxes (Grebe & Stock, 1999; Parkinson & Kofoid, 1992). Based on the presence and structure of the various homology boxes, a comprehensive classification of HPKs was made by Grebe & Stock (1999). In this classification the vast majority of peptide-based QS-TCSs comprise specific HPKs belonging to the subfamily called HPK₁₀. The only exception known to date is ComP of Bacillus subtilis, which is a QS-HPK that belongs to a different HPK subfamily (HPK₇). The HPK₁₀ homology boxes have the following characteristics: the H-box (histidine phosphorylation site) contains a tyrosine two residues downstream from the conserved histidine (characteristic motif F+HDYxN) and lacks an otherwise conserved proline residue at position 5 downstream; the X-box (a hydrophobicity pattern conserved in several subfamilies) is present; the N-box has only one conserved asparagine residue (characteristic motif DNAIE); and the G-box, which plays a critical role in phosphoryl transfer, has a characteristic FSTKGxGxGLGL motif (Fig. 1). The D-box that in most subfamilies is part of the nucleotide-binding domain is absent in this subfamily. Furthermore, HPK₁₀ subfamily members commonly possess five to seven N-terminal transmembrane segments (TMSs) (Fig. 1).

In general, RRs contain a C-terminal DNA-binding domain and a N-terminal CheY-like receiver domain (REC: SM00448) that includes a conserved phosphorylaccepting aspartate residue (Volz, 1993) (Fig. 1). Analogous to the HPKs, the RRs have been classified based on the receiver and the DNA-binding domains (Grebe & Stock, 1999). In both classifications the RRs related to HPKs of the HPK₁₀ subfamily comprise a separate subfamily: the RD and ComE subfamily, respectively. Most established QS-RRs are encompassed within this RD/ ComE subfamily of RRs. However, in accordance with the unusual sequence of its cognate HPK, the competenceregulating RR (ComA) of B. subtilis belongs to the RE rather than the RD subfamily of RRs and contains a HTH-LuxR DNA-binding domain (SM00421) (Fuqua et al., 1994). In a more recent analysis most RRs of the RD/ComE subfamily were classified in the LytTR family of response regulators (PF04397), based on a conserved motif in the Cterminal helix-turn-helix (HTH) DNA-binding domain (Nikolskaya & Galperin, 2002). For RRs the presence of a HTH-LytTR DNA-binding domain does not in all cases classify a RR as a peptide-based QS-RR (Nikolskaya & Galperin, 2002). However, the presence of an adjacent HPK with HPK₁₀-subfamily characteristics can be used to classify the RR as such.

Genomic context: linkage of peptide-based QS-TCSs with AIPs and other QS-related functionalities

In Gram-positive bacteria, genes encoding peptide-based QS-TCSs are in general preceded by genes encoding the cognate autoinducing peptide (AIP). Many AIPs have a structure similar to class I bacteriocins (lantibiotics) (McAuliffe et al., 2001) or to class II bacteriocins (nonlantibiotics) (Ennahar et al., 2000). These AIPs contain recognizable leader peptides with conserved residues, such as the double-glycine leader peptides of most class II and some class I bacteriocins (the consensus for residues -12to -1 of GG-leader is LSxxELxxIxGG) (Nes & Eijsink, 1999). The web-based bacteriocin genome-mining tool BAGEL (http://bioinformatics.biol.rug.nl/websoftware/bagel/ bagel_start.php) might be used to identify such bacteriocin-like AIPs that contain leader peptides (de Jong et al., 2006). In addition, some QS-TCS genes are genetically linked to genes encoding AIP transport and/or modification proteins, bacteriocins and bacteriocin-immunity proteins (Kleerebezem et al., 1997). For double-glycinetype AIPs the cognate transporters are ABC transporters that contain a characteristic N-terminal peptidase C39 domain (COG2274) (Håvarstein et al., 1995), while in the case of agr-like systems AgrB-type cysteine proteases are involved in transport and modification of AIPs (Nakayama et al., 2006; Qiu et al., 2005; Zhang & Ji, 2004).

In silico identification of candidate QS-TCSs in lactobacilli

To identify new putative QS-TCSs, protein sequences of experimentally verified QS-TCSs in *Staphylococcus aureus* (AgrCA) (Ji *et al.*, 1997) and *L. plantarum* C11 (PlnBCD) (Diep *et al.*, 1996) were collected from public databases (http://www.ncbi.nlm.nih.gov/). Potential system homologues were collected from the genomes of lactobacilli and other Firmicutes via iterative BLASTP searches for HPKs or RRs, using default settings (PSI-BLAST, *E*-value threshold 1×10^{-5}) (Altschul *et al.*, 1990).

Then, various peptide-based OS-specific protein characteristics were used to reduce the list of homologues. These characteristics included: (i) for HPKs the presence of HPK₁₀-subfamily domains (Grebe & Stock, 1999) and five to seven N-terminal transmembrane segments (TMSs), and (ii) for RRs the presence of RD/ComE subfamily domains (Grebe & Stock, 1999) or HTH_LytTR DNA-binding domains (Nikolskaya & Galperin, 2002). In addition, (iii) the presence of adjacent AIP-like genes and additional peptide-based QS-related genes was investigated for these putative OS-TCSs. Protein domains were predicted using the HMMs of SMART, including outlier homologues and PFAM domains (Schultz et al., 1998), and membrane topology was predicted using TMHMM 2.0 (Krogh et al., 2001). The remaining HPK and RR sequences were aligned, and bootstrapped neighbour-joining trees were constructed with CLUSTAL X (Thompson et al., 1997). The The in silico analysis confirmed previously described QS-TCSs and identified novel putative QS-TCSs and AIPs in lactobacilli. The analysis predicted the presence of five QS-TCSs in L. plantarum WCFS1, two QS-TCSs in both the intestinal species Lactobacillus acidophilus NCFM and L. johnsonii NCC533, one QS-TCS in the intestinal species L. salivarius subsp. salivarius UCC118 and the food species L. delbrueckii subsp. bulgaricus ATCC BAA-365, and no QS-TCS in the intestinal species L. gasseri ATCC 33323. In most cases these putative QS-TCSs had adjacent genes encoding class II bacteriocin-like or lantibiotic-like peptides, which might serve as AIPs, as well as ABC transporters with a peptidase C39 domain, which supports a role in QS-regulated bacteriocin production (see Fig. S2). The QS-functionality of abpIPKR in L. salivarius and LBA1798-LBA1800 in L. acidophilus was previously shown (Flynn et al., 2002; Dobson et al., 2007), and the functional analysis of the putative QS-TCSs in L. plantarum is discussed in the next section.

In addition, in each of the food species Lactobacillus brevis ATCC 367, L. casei ATCC 334 and L. sakei subsp. sakei 23K one putative QS-TCS was identified. However, for those systems the functionality of the predicted QS-TCS is doubtful, since their HPK genes seemed to be (i) interrupted (internal deletions) or (ii) incomplete (Nterminus absent), or (iii) the cognate HPK was apparently absent (i.e. not adjacent to the RR). A good example of this is the sppIPKR QS system, which was detected and functional in several L. sakei strains, but not functional in L. sakei subsp. sakei 23K (Møretrø et al., 2005). This is caused by a 4 bp internal deletion in the HPK gene (*sppK*) of this strain, resulting in two truncated HPK fragments (*sppkN* and *sppKC*) and thereby a non-functional QS-TCS. In L. casei ATCC 334 there are two genes present encoding RRs with RD and LytTR domains. One appears to be an orphan gene (LSEI_2389) with a putative AIP downstream (LSEI_2390). The other RR (LSEI_2599) is part of a putative QS-TCS, where the HPK encoded by LSEI_2600 contains the HPK₁₀-subfamily motifs but lacks most of the N-terminus with TMS. Finally, for L. brevis ATCC 367 there are two HPKs present (typical is only one HPK) downstream of a RR with RD and LytTR domains (LVIS_0163), but they lack either a clear H-box (LVIS_0164) or N-box (LVIS_0165), which are typical of the HPK₁₀ subfamily.

Functional analysis of candidate QS-TCSs in *L. plantarum* WCFS1

Previous annotation of the 3.3 Mb genome sequence of *L. plantarum* WCFS1 revealed the presence of 13 genetically linked TCSs, and one orphan HPK and RR (Kleerebezem

Table 1. General features of HPKs and RRs of candidate QS-TCSs in *L. plantarum* WCFS1 and related lactobacilli (see also Figs S2 and S3)

Accession numbers for Lactobacillus genomes: L. plantarum WCFS1 (AL935263); L. acidophilus NCFM (CP000033); L. johnsonii NCC533 (AE017198); L. salivarius subsp. salivarius UCC118 (CP000233); L. sakei subsp. sakei 23K (CR936503); L. brevis ATCC 367 (CP000416); L. delbrueckii subsp. bulgaricus ATCC BAA-365 (CP000412); L. casei ATCC 334 (CP000423).

Gene*/ accession no.	Locus†	Size (aa)	TMS‡	HPK10 subfamily§	Gene*/ accession no.	Locus†	Size (aa)	LytTR domains	Gene*/ accession no.	Locus†
НРК					RR				AIP	
L. plantarum WCFS1 (3.34 Mb)										
plnB	lp_0416	442	711	+	plnC/plnD	lp_0417/ lp_0418	247	+	plnA	lp_0415
pltK	lp_1355	420	6	+	pltR	lp_1356	255	+	pltA	lp_1354a
hpk4	lp_1488	343	_	- (HPK ₇)	rrp4	lp_1487	217	- (LuxR)	_	
hpk6	lp_1943	367	5	- (HPK ₇)	rrp6	lp_1942	201	- (LuxR)		
hpk9	lp_3063	422	6	+	rrp8	lp_2665	249	+		
hpk10	lp_3088	416	6	+	rrp10	lp_3087	248	+	—	lp_3089
lamC	lp_3581	419	5	+	lamA	lp_3580	247	+	lamD	lp_3581a
L. acidothilus N	ICFM (2.0 Mb)									
YP 193512	LBA0602	426	6	+	YP 193513	LBA0603	265	+		
YP 194634	LBA1799	440	7	+	VP 194633	LBA1798	200	+	VP 194635	I BA 1800
		110	,		11_171055	LDITITY	271	I	11_171055	LDIII000
L. johnsonii NCC533 (2.0 Mb)										
NP_964473	LJ0448	419	6	+	NP_964474	LJ0449	255	+		
NP_964617	LJ0764	435	6	+	NP_964619	LJ0766	265	+	NP_964616	LJ0763b
Leading site and an endingering UCC118 (2.12 Mb)										
aboK	1 SI 1013	A20	7	-	ahoP	ISI 1012	264	_L	aboID	ISI 1017
иорк	L3L_1915	429	/	Ŧ	иорк	L3L_1912	204	Ŧ	uopir	L3L_1914
Lactobacillus sakei subsp. sakei 23K (1.9 Mb)										
sppKN+sppKC	LSA0561	69 ^{N,Δ}	0	+	sppR	LSA0563	248	+	sppIP	LSA0560_b
	+ LSA0562	$+183^{N,\Delta}$								
L. brevis ATCC										
YP_794364	LVIS_0164	444	6	\pm	YP_794363	LVIS_0163	251	+		
_	_			(no H-box)	_	_				
YP 794365	LVIS 0165	443	7	+						
_				(no N-box)						
I delhruechii oubon hulgaricus ATCC RAA 365 (1.0 Mb)										
VP 812154	IBUI 0022	134	6	- ×	VP 812153	IBUL 0021	260	<u>т</u>		
11_012134	LDOL_0022	1,71	0	1	11_012155	LDCL_0021	200	I		
L. casei ATCC 334 (2.93 Mb)										
		N			YP_807570	LSEI_2389	268	+	YP_807571	LSEI_2390
YP_807763	LSEI_2600	303 ^{IN}	2	+	YP_807762	LSEI_2599	234	+		

*, †Gene name and locus number from http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi.

‡TMS, transmembrane segments as predicted from TMHMM 2.0 (http://www.cbs.dtu.dk/services/TMHMM/).

\$HPK classification after Grebe & Stock (1999).

IExperimentally confirmed data for PlnB in L. plantarum C11 (Johnsborg et al., 2003).

^N, Incomplete gene: N-terminal domain (partially) missing.

^{Δ}, 4 bp deletion in *sppK* gene (Møretrø *et al.*, 2005).

et al., 2003). Out of these, five TCSs were identified as candidate QS-TCSs based on the *in silico* approach described above. They were predicted to comprise five HPKs that showed characteristics of the HPK₁₀ subfamily and six RRs that contained a HTH-DNA-binding domain of the LytTR family. Of the remaining TCSs, which did not fit the general *in silico* criteria for peptide-based QS-TCSs,

we found on closer examination that there were two TCSs where the HPKs and RRs showed similarity to another type of QS-TCS, as found for the ComPA system of *B. subtilis.* These HPKs showed HPK₇-subfamily characteristics (Grebe & Stock, 1999) and the RRs contained a HTH-DNA-binding domain of the LuxR family (Fuqua *et al.*, 1994) (Table 1).

Four adjacent HPK- and RR-encoding genes constituted complete TCSs that were classified as candidate peptidebased QS-TCSs (pln, plt, TCS10 and lam). In addition, the orphan HPK and RR could constitute a fifth peptide-based QS-TCS. The relevant features of these HPKs and RRs are summarized in Table 1 and their genetic organization is shown in Supplementary Figs S2 and S3. For all of the complete QS-TCSs a putative AIP was encoded upstream (Fig. 2 and Fig. S3), two of which (pln and plt) contained a putative double-glycine-type leader peptide, while the third and fourth were of a different type. The gene connected to the lam QS-TCS (lamD) encodes a cyclic thiolactone AIP (Sturme et al., 2005), whereas the one connected to TCS10 encodes a putative AIP that shows little similarity to known AIPs. Details of the characterization and functionality of these putative OS-TCSs are discussed below.

Plantaricin TCS pln (lp_0415 to lp_0418)

The plantaricin TCS pln (lp_0415 to lp_0418) was identical to the previously described *plnABCD* system of L. plantarum C11, which regulates the production of class II antimicrobial peptides (AMPs) (Diep et al., 1996). This system contains a gene (plnB: lp 0416) encoding a typical HPK of subfamily HPK₁₀ that was shown to have seven TMSs (confirmed in strain C11 by Johnsborg et al., 2003). Downstream of plnB, two RRs are encoded (plnC and plnD) that both contain the RD receiver domains and LytTR HTH-DNA-binding domains. Upstream of plnB, plnA encodes a 48 amino acid double-glycine-type AIP precursor (Nes & Eijsink, 1999). Genes encoding class II bacteriocins (PlnE-PlnF, PlnJ-PlnK and PlnN) were localized near the *pln* regulatory module (Supplementary Fig. S3). Cleavage of the double-glycine-type leader peptide from the PlnA precursor peptide results in a linear AIP of 26 amino acids without modifications and an amphipathic character (Fig. 2). The *pln* system was found to be functional in *L. plantarum* WCFS1, as was shown by a bacteriocin agar-well diffusion assay with *L. plantarum* 965 as an indicator strain (Fig. 3). The *L. plantarum* WCFS1 native state was bacteriocin-negative (Bac⁻), but bacteriocin production could be induced with either bacteriocin-positive supernatant of strain C11 (Bac⁺) or purified PlnA peptide.

The *pln* system of WCFS1 is expected to play a role in competition with other bacteria, as in strain C11 it regulates the production of the bacteriocins PlnE-PlnF, PlnJ-PlnK and PlnN. These C11 plantaricins showed activity against closely related species (Anderssen *et al.*, 1998), which can be found in the same ecological niches as *L. plantarum* strains WCFS1 and C11 (Vaughan *et al.*, 2002). However, some strain-specific differences in plantaricin activity and target specificity might be expected for various *L. plantarum* strains, considering the differences in *pln* gene composition and sequence variations identified in the *pln* operons of strains WCFS1, C11 and NC8 (Maldonado *et al.*, 2004; Molenaar *et al.*, 2005).

TCS plt (lp_1354a, lp_1355 and lp_1356)

The *plt* locus (lp_1354a, lp_1355 and lp_1356) encodes a typical HPK (*pltK*: lp_1355) of the HPK₁₀ subfamily with five or six predicted TMSs, and a RR (*pltR*: lp_1356) that contains a RD-type receiver domain and a LytTR HTH-DNA-binding domain. Upstream of *pltK*, a 58 amino acid double-glycine-type AIP precursor appears to be encoded (*pltA*: lp_1354a). The predicted mature PltA peptide is a 28 amino acid candidate AIP that is expected to be unmodified (Fig. 2). Northern blot analysis showed that the *pltAKR* operon is transcribed as a single polycistronic, 2.4 kb transcript in a cell-density-dependent manner (Fig. 4). The predicted mature PltA peptide was chemically



Ip_3089 MVQWAKRFSETKEPVVLISHNQNRCAGKIVVLMMSRLELWGS

Fig. 2. AIPs encoded on the *L. plantarum* WCFS1 genome. Triangles indicate the cleavage site between the double-glycine leader peptide and the (predicted) mature peptide in PlnA and PltA (shown in bold). The underlined bold residues in the LamD precursor peptide are processed to the mature thiolactone peptide shown on the right. For the putative AIP encoded by lp_3089 the mature peptide sequence and structure are unknown.



Fig. 3. Agar well diffusion assay for plantaricin production in L. plantarum WCFS1 and the related strain C11, using indicator strain L. plantarum 965. To induce plantaricin production in L. plantarum WCFS1, this strain was grown in MRS medium containing 1 µg ml⁻¹ of purified PlnA peptide or in MRS medium containing 2% of spent culture supernatant of bacteriocinproducing L. plantarum C11. To obtain a bacteriocin-negative derivative of L. plantarum C11, bacteriocin-producing cells of this strain were inoculated in MRS at a density below 10⁴ c.f.u. ml⁻¹ (previously reported to result in loss of plantaricin production). A, C11, Bac+; B, WCFS1, Bac-, native state of WCFS1; C, Pln production in WCFS1, induced with purified PInA peptide; D, C11, Bac⁻, obtained by <10⁴ c.f.u. ml⁻¹ dilution; E, Pln production in C11 ($<10^4$ c.f.u. ml⁻¹ dilution), induced with purified PlnA; F, Pln production in WCFS1, induced with C11, Bac⁺ supernatant; G, Pln production in C11, induced with WCFS1, Bac+ supernatant. Strain C11, indicator strain L. plantarum 965 and purified PInA were generous gifts from Dzung Diep and Ingolf Nes, Norwegian University of Life Sciences, Ås, Norway.

synthesized, but was water-insoluble and acted as a gel above 35–40 % purity. Therefore it could not be used in induction experiments. The water insolubility could result from the high hydrophobicity and lack of α -helical amphipathic characteristics of this peptide, which is typical in class II bacteriocin-like AIPs (Anderssen *et al.*, 1998). Alternatively, the *pltA*-encoded peptide could be subject to unpredicted post-translational modifications that affect its solubility and render a functional secreted AIP. In conclusion, the role of the proposed (modified) mature PltA peptide in the cell-density-dependent expression of the *pltAKR* locus and possible secondary target loci remain to be established. Nevertheless, its canonical genetic organization suggests a role of the *pltAKR* operon in QS, which is supported by its observed cell-density-dependent



Fig. 4. Northern blot analysis of temporal gene expression of the *lamBDCA* and *pltAKR* operons of *L. plantarum* WCFS1, when grown in MRS at 30 °C, without agitation. Growth phases are indicated above the samples: E, M and L indicate early-, mid- and late-exponential growth phases and S the stationary growth phase. Equal amounts of RNA (10 μ g) were loaded. Expression of *lamBDCA* was monitored using a *lamC* internal probe, and expression of *pltAKR* using a *pltK* internal probe. Transcript sizes are shown on the right.

expression pattern. The *plt* locus does not appear to be genomically directly linked to any ABC transporter or bacteriocin-encoding gene (Supplementary Figs S2 and S3). However, the PltA pre-peptide could be transported by another ABC transporter with a peptidase C39 protease-family domain (Håvarstein *et al.*, 1995), such as the ABC transporter encoded within the *pln* locus (encoded by *plnG* and *plnH*: Fig. S3).

TCS lam (lp_3580 to lp_3582)

The TCS encoded by lp_3580 to lp_3582 was designated lam for Lactobacillus agr-like module, as it showed a similar gene organization to the agrBDCA QS system of S. aureus (Ji et al., 1995). The L. plantarum lamBDCA locus encodes a HPK (lamC: lp_3581) of the HPK₁₀ subfamily with five predicted TMSs, and a RR (lamA: lp_3580) that contains a RD-type receiver domain and a LytTR HTH-DNA-binding domain. The locus is organized as an operon and is transcribed as a single transcript in a cell-densitydependent manner (see Fig. 4 and Sturme et al., 2005). Analogous with the staphylococcal agr system (Ji et al., 1995), the L. plantarum lamBDCA locus is involved in production of a cyclic thiolactone peptide (CVGIW) which is predicted to derive from the LamD precursor (lp_3581a) that is processed by LamB (lp_3582) (Fig. 2). Analysis of a lamA mutant revealed a role for the lam operon in L. plantarum biofilm-forming capacity. Transcriptome profiling of wild-type and lamA mutant strains in early-, midand late-exponential phase uncovered only a small set of clustered genes (2% of all genes) that were significantly modulated by the lamA mutation. These data confirmed the autoregulation of lamBDCA, and showed the regulation of a surface polysaccharide biosynthesis gene-cluster, and several cell envelope and sugar utilization functions (Sturme et al., 2005). The same study suggested that a direct repeat sequence within the lamB promoter region (5'-TCTTTAAAT - 12 bp - TCTTAAAA-3') that displays similarity with previously established cognate cis elements of LamA homologues (Morfeldt et al., 1996; Qin et al., 2001; Quadri et al., 1997; Risøen et al., 2000) acts as the LamA DNA-binding site.

TCS10 (lp_3087 and lp_3088)

The bioinformatic analysis suggests that the HPK and RR of TCS10 (lp_3087 and lp_3088) are inparalogues of the lamCA-encoded proteins, with the cognate HPKs and RRs showing 55 % and 70 % identity, respectively, at the amino acid level. This module lacks a lamB homologue and originally seemed to lack a lamD homologue. However, a region somewhat upstream of the hpk10 gene is remarkably similar to that of the region upstream of the lam system. This (promoter) region contains a direct repeat (5'-TCTTGAAAT - 12 bp - TCTTAAAG-3'), displaying very high similarity with the proposed regulatory element of the lamB promoter (see above). Interspersed between this regulatory region of TCS10 and the *hpk10* gene is a region of about 130 nucleotides that on closer inspection was shown to encode a small peptide (lp 3089) that initially was not annotated (Fig. 2 and Supplementary Fig. S3). Although the peptide shows some conservation with respect to the lamD gene product, it is not clear whether it is a genuine AIP. Taken together these findings support an inparalogous relationship between the RRs and HPKs of the TCS10 and lam systems, and suggests that there could be cross-talk between these two regulatory systems.

Orphan genes rrp8 (lp_2665) and hpk9 (lp_3063)

The two orphan *hpk* and *rrp* genes show typical characteristics of peptide-based QS systems. The orphan gene hpk9 (lp 3063) was predicted to encode a HPK₁₀subfamily protein containing six TMSs. Interestingly, immediately downstream (7 bp) of hpk9 a small ORF is located (lp_3062) that shows 46% homology to the Cterminal domain of the TfoX protein of Haemophilus influenzae (PF04994). The TfoX protein is proposed to play a key role in cell-density-dependent regulation of genetic competence in H. influenzae, and the C-terminal domain is suggested to function autonomously (Zulty & Barcak, 1995). The lp_3062 gene product might therefore function in co-operation with hpk9. However, no clearly identifiable cognate RR appears to be encoded in the vicinity of *hpk9*. The downstream-located response regulator lp_3060 probably does not fulfil this function, as it was not transcriptionally linked and lacks a RD-type receiver domain and contains a typical HTH-AraC DNA-binding domain (SM00342) rather than the canonical LytTR or HTH-LuxR DNA-binding domains (see Supplementary Fig. S3). It is possible that the hpk9-encoded HPK in combination with the rrp8-encoded orphan RR forms a functional TCS, which could be involved in peptide-based QS. This possibility is supported by the finding that rrp8 (lp_2665) encodes a RR with a typical LytTR HTH-DNAbinding domain. Moreover, the neighbour-joining trees based on protein sequence alignments of HPKs and RRs belonging to QS-TCS showed that the hpk9- and rrp8encoded proteins (LPL03063 and LPL02665 in Supplementary Fig. S1) cluster with several HPKs and RRs of other Firmicutes that are adjacent on their respective genomes, such as LEUM_0009/LEUM_0008 from *Leuconostoc mesenteroides* or EF1820/EF1822 from *Enterococcus faecalis*. Interestingly, the gene downstream of *rrp8* (lp_2664) contains a HDc-superfamily-type phosphohydrolase-domain (SM00471) that might have a role in dephosphorylation of the *rrp8* gene product. Overall, the role of *hpk9* in QS-mediated regulation and the identity of its eventual partnering transcriptional regulator remain to be established.

Additional candidate QS-TCSs and genes involved in non-peptide-based QS

Besides the typical peptide-based QS-TCSs, TCS4 (lp_1487 and lp_1488) and TCS6 (lp_1942 and lp_1943) both encode HPK₇-subfamily-type HPKs (Grebe & Stock, 1999) adjacent to a RR containing a HTH-LuxR DNA-binding domain (see Table 1 and Supplementary Fig. S3). This resembles the protein architecture of the ComPA QS-TCS of B. subtilis that is involved in OS regulation of competence (Weinrauch et al., 1989, 1990). However, both the HPK₇ subfamily and the HTH-LuxR DNA-binding domain are not exclusively associated with QS-TCSs (Gray & Garey, 2001; Grebe & Stock, 1999), indicating that on the basis of in silico analysis the involvement of TCS4 and TCS6 in QS can only be tentatively suggested. Interestingly, next to the putative QS-TCS an isolated homologue of the luxS gene was identified (lp_0774), encoding the autoinducer-2 (AI-2) synthase (Schauder et al., 2001). AI-2 is thought to play a role in QS, although there is still some debate on its actual physiological role (Sun et al., 2004). Associated functions like lsr, which is involved in AI-2 uptake in Salmonella typhimurium, or luxPQ involved in AI-2 sensing and signal transduction in Vibrio harveyi, were not identified in the L. plantarum genome (Sun et al., 2004). AI-2 could be involved in interspecies communication of L. plantarum with other bacteria within the same niche, as the *luxS* gene was detected in both Gram-negative and Gram-positive bacteria (Xavier & Bassler, 2003). Interestingly, AI-2 activity was detected in rumen samples (Mitsumori et al., 2003), suggesting a natural role of the AI-2 QS system in gastrointestinal ecosystems.

Concluding remarks

L. plantarum WCFS1 contains a relatively high number of (putative) peptide-based QS-TCSs (five, of which at least four can be directly coupled to a putative AIP), as well as other putative QS genes. This could reflect the ecological flexibility of this species, which can be found in plants, fermented foods and the gastrointestinal tract. In comparison, the genomes of related lactobacilli that are more restricted to specific environments seem to encode significantly fewer peptide-based QS-TCSs, despite the differences in genome size (see Table 1). Depending on

niche-specific conditions, different signalling systems might be triggered, resulting in a gene-regulatory network that controls a variety of phenotypic traits in response to environmental conditions in combination with cell density. The potential for cross-talk between the *lamBDCA* and TCS10 systems might be exemplary for the complexity of a regulatory network that controls surface adherence of *L. plantarum* under specific conditions (Sturme *et al.*, 2005). Moreover, the presence of competing micro-organisms could activate specific QS-TCSs involved in competition, as has been shown for the plantaricin system in *L. plantarum* NC8 (Maldonado *et al.*, 2004). Interestingly, the involvement of *agr*-like TCSs in host–microbe interactions of commensal bacteria has recently been suggested for *Roseburia inulinivorans* (Scott *et al.*, 2006).

Further experimental studies on the regulatory mechanisms of the different QS systems of *L. plantarum* and other *Lactobacillus* species and their effects on (global) gene regulation will be necessary, to provide more insight into the role of these systems in the survival of these organisms in their natural environments.

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