

Antibodies to a synthetic 1–9-N-terminal amino acid fragment of mature pediocin PA-1: sensitivity and specificity for pediocin PA-1 and cross-reactivity against Class IIa bacteriocins

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Polyclonal antibodies specific for pediocin PA-1 (Peda1) were generated by immunization of rabbits with a chemically synthesized 1–9-N-terminal amino acid fragment of this bacteriocin (PH1) conjugated to the carrier protein keyhole limpet haemocyanin (KLH). The PH1 fragment holds a highly conserved amino acid sequence with closely related Class IIa bacteriocins. The sensitivity and specificity of the PH1–KLH-generated rabbit polyclonal antibodies were evaluated by the development of various ELISAs, such as a non-competitive indirect ELISA (NCI-ELISA), a competitive indirect ELISA (CI-ELISA), a competitive direct ELISA (CD-ELISA) and a sandwich ELISA (S-ELISA), and by protein slot-blotting and Western blotting. NCI- and CI-ELISA were valuable for detecting the existence of Peda1-specific antibodies in the sera of immunized rabbits. The limit of detection of Peda1 in MRS medium was found to be 0.5 µg ml⁻¹ in NCI-ELISA, while CI-ELISA on plates coated with purified Peda1 increased the affinity of the PH1–KLH-generated antibodies for Peda1; the limit of detection of Peda1 was less than 0.01 µg ml⁻¹ and 50% binding inhibition was achieved with 0.1 µg Peda1 ml⁻¹. Similarly, the limits of detection of Peda1 in MRS medium were found to be 5 µg ml⁻¹ by protein slot-blotting and 0.01 µg ml⁻¹ by Western blotting. Most importantly, PH1–KLH-generated polyclonal antibodies detected the presence of Peda1 in the supernatants of the producing strains of *Pediococcus acidilactici* 347, Z102, A172, X13 and P20, with no reactivity or negligible immunoreactivity with the supernatants of other lactic acid bacteria producing or not producing closely related or different bacteriocins. The approaches taken for the selection of the bacteriocin peptide fragment, the generation of antibodies and the development of immunoassays could prove useful for the generation and evaluation of antibodies of adequate specificity for other bacteriocins of interest in the food industry.

Keywords: pediocin PA-1, bacteriocin, immunodetection, lactic acid bacteria

INTRODUCTION

Many lactic acid bacteria (LAB) are known to secrete

Abbreviations: ADT, agar diffusion test; CB, coating buffer; CD-, CI-, NCI- and S-ELISA, competitive direct, competitive indirect, non-competitive indirect and sandwich ELISA; KLH, keyhole limpet haemocyanin; LAB, lactic acid bacteria; MPA, microtitre plate assay; OA, ovalbumin; Peda1, pediocin PA-1; TMB, 3,3',5,5'-tetramethylbenzidine.

small, ribosomally synthesized antimicrobial peptides called bacteriocins (Klaenhammer, 1993; Jack *et al.*, 1995; Nes *et al.*, 1996). Some of these peptides have received considerable attention in recent years due to their potential application in the food industry as natural food preservatives. The bacteriocins produced by LAB are often cationic, amphiphilic, membrane-permeabilizing peptides classified into two main groups: Class I consisting of the modified bacteriocins, the

lantibiotics, and Class II consisting of the unmodified peptide bacteriocins, such as the pediocin-like bacteriocins, the non-pediocin-like bacteriocins and the two-peptide bacteriocins (Nissen-Meyer & Nes, 1997). A major subgroup of Class II bacteriocins (IIa) has been given the generic name of the pediocin-like family (Nes *et al.*, 1996) after its most extensively studied member, pediocin PA-1 (PedA1). The pediocin-like family is the only grouping that has been classified by sequence similarities, and bacteriocins of this family share structural homologies and are composed of a conserved N-terminal hydrophilic domain and a variable C-terminal domain (Aymerich *et al.*, 1996; Casaus *et al.*, 1997; Cintas *et al.*, 1997). PedA1 has been characterized at the biochemical (Henderson *et al.*, 1992; Nieto Lozano *et al.*, 1992) and genetic (Marugg *et al.*, 1992; Venema *et al.*, 1995) levels. The mature PedA1 molecule is a 44 aa peptide with a molecular mass of 4629 Da that contains four cysteine residues that participate in the formation of two disulfide bridges. The peptide is predicted to exist largely as a random coil, with only a small hydrophobic region in residues 21–25 with a propensity to form a β sheet (Klaenhammer, 1993).

Because of the potential use of bacteriocins as food preservatives and since most industrial strains do not produce such antagonistic peptides, interest in the heterologous expression or co-expression of Class II bacteriocins is growing rapidly (van Belkum *et al.*, 1997; Biet *et al.*, 1998; Horn *et al.*, 1998). However, an adequate identification, detection and quantification of the bacteriocin(s) produced by the heterologous hosts is demanded. The use of a bioassay-based method that assesses the inhibitory effect of bacteriocins in a test or indicator micro-organism is the most commonly used tool for detection and quantification of bacteriocins. The importance of the bioassay is undeniable, but it also has some drawbacks, such as lack of specificity and low sensitivity. However, the antimicrobial efficiency of different methods of food preservation can be improved through the application of the hurdle concept (Leistner & Gorris, 1995; Abriouel *et al.*, 1998). Bacteriocins either alone or in combination with other antimicrobial barriers may be useful tools to substantially reduce the load of foodborne pathogens and food spoilage bacteria. However, bacteriocin preparations with adequate purity must be provided. Accordingly, the development of efficient detection, quantification and purification procedures for PedA1 and other bacteriocins could greatly facilitate their use as food preservatives. The generation of antibodies against bacteriocins may provide sensitive and specific methods for the identification and detection of producing strains and for the quantification of bacteriocins in different substrates by the use of immunochemical assays (Martínez *et al.*, 1998). Antibodies also offer potential alternative methods for the purification of bacteriocins to homogeneity by the use of immunoaffinity chromatography strategies (Suárez *et al.*, 1997).

The scarcity of appropriate immunochemical methods for use in the bacteriocin research field is most probably due to the difficulties encountered in raising antibodies

and in the development of sensitive immunoassays. Reports on the generation of antibodies against bacteriocins have been scarce and have been based on the use of whole bacteriocin molecules, either alone or conjugated to carriers, as the immunogen (Bhunia *et al.*, 1990; Falahae *et al.*, 1990; Bhunia, 1994; Stringer *et al.*, 1995; Suárez *et al.*, 1996a, b; Bouksaim *et al.*, 1998). Recently, the use of a chemically synthesized fragment deduced from the C-terminal amino acid sequence and unique to PedA1 has facilitated the development of antibodies of predetermined specificity against PedA1 (Martínez *et al.*, 1998). However, the specificity of antibodies generated against bacteriocin sequences sharing strong consensus similarities has not been evaluated yet. We report in this communication the sensitivity and specificity of antibodies generated against a synthetic 1–9-N-terminal amino acid fragment of PedA1, a short peptide with a strong amino acid sequence homology with Class IIa bacteriocins and the development of sensitive immunoassays for PedA1 analysis.

METHODS

Materials. The amino acid sequence of the N-terminal fragment of PedA1 (peptide PH1) used in this work was NH₂-KYYGNGVTC-COOH. Peptide PH1 (residues 1–9, 1104 Da, 30 mg) was synthesized by Fmoc chemistry with an Applied Biosystems 431A automated solid-phase peptide synthesizer in the Protein Chemistry Facility at the Centro de Biología Molecular Severo Ochoa, Madrid, Spain. Purity of the peptide was monitored by RP-HPLC, being higher than 95%, and peptide identity was confirmed by MS. A chemically synthesized fragment corresponding to the C-terminal region of PedA1 (peptide PH2, residues 34–44, 1009 Da, 10 mg) was used as a control. The amino acid sequence of peptide PH2 was NH₂-ATGGHQGNHKC-COOH. Ovalbumin (OA) (grade III and fraction VII), horseradish peroxidase (HRP) (fraction VI), Tween 20, glutaraldehyde and Freund's adjuvants were obtained from Sigma. The Imject Activated Immunogen Conjugation Kit containing maleimide-activated keyhole limpet haemocyanin (KLH), maleimide-activated OA, conjugation buffer and gel filtration columns was obtained from Pierce. Goat anti-rabbit immunoglobulin G conjugated to horseradish peroxidase was obtained from Cappel. Pure nisin A (30 000 U mg⁻¹) was purchased from NBS Biologicals. Rabbits (New Zealand white females) were purchased from a local supplier (Navarra, Spain).

Preparation of immunoconjugates and immunization. PH1 was conjugated to maleimide-activated KLH (PH1–KLH, 1:2, w/w) using the components of the Imject Activated Immunogen Conjugation Kit, for use as the immunogen. The chemically synthesized PH1 fragment was also conjugated to maleimide-activated OA (PH1–OAM, 12.5:1, mol/mol) and to OA (PH1–OAG, 12:1, mol/mol) by the glutaraldehyde method (Avrameas & Ternynck, 1969; Briand *et al.*, 1985) for use as a solid-phase antigen. Peptide PH2 was also conjugated to OA by the glutaraldehyde method (PH2–OAG, 12.5:1, mol/mol) for use as a solid-phase antigen. PH1 and purified PedA1 were also conjugated to horseradish peroxidase (PH1–HRP, 1:5, w/w; PedA1–HRP, 1:5, w/w) by the periodate method (Nakane & Kawoi, 1974) for use in competitive direct ELISAs. Rabbits were immunized with PH1–KLH according

Table 1. Reactivities of serum polyclonal antibodies against culture supernatants of LAB as determined by a NCI-ELISA and CI-ELISA

Strain (bacteriocin produced)	Source*	Cross-reactivity (%)	
		NCI-ELISA †	CI-ELISA ‡
<i>P. acidilactici</i> 347 (PedA1)	Our collection	76.1	99.3
<i>P. acidilactici</i> Z102 (PedA1)	Our collection	100	99.4
<i>P. acidilactici</i> A172 (PedA1)	Our collection	83.7	99.2
<i>P. acidilactici</i> X13 (PedA1)	Our collection	90.7	100
<i>P. acidilactici</i> P20 (PedA1)	Our collection	4.5	8
<i>P. acidilactici</i> 347 Ped ⁻ (non-PedA1 producer)	Our collection	NR	NR
<i>P. pentosaceus</i> FBB61 (pediocin A)	TNO	NR	NR
<i>E. faecium</i> T136 (enterocin A and B)	Our collection	2.9	4.2
<i>E. faecium</i> P13 (enterocin P)	Our collection	NR	NR
<i>Lb. sakei</i> LTH673 (sakacin P)	NHL	NR	NR
<i>Lb. sakei</i> 706 (sakacin A)	NHL	NR	NR
<i>E. faecium</i> L50 (enterocin L50A and L50B)	Our collection	NR	0.6
<i>E. faecalis</i> INIA4 (enterocin AS-48)	INIA	2.1	NR
<i>Lb. sakei</i> 148 (lactocin S)	Our collection	NR	NR
<i>Lc. lactis</i> BB24 (nisin A)	Our collection	3.9	NR
<i>Lc. lactis</i> MG1614 (non-bacteriocin producer)	IFR	NR	1.2

NR, No reactivity.

* IFR, Institute of Food Research, Norwich Laboratory, Norwich, UK; INIA, Instituto Nacional de Investigación y Tecnología Agraria y Alimentaria, Madrid, Spain; TNO, Nutrition and Food Research, Zeist, The Netherlands; NHL, Laboratory of Microbial Gene Technology, Agricultural University, Ås, Norway.

† Cross-reactivity defined as [(absorbance reading produced by a culture supernatant above the absorbance reading produced by MRS/absorbance reading produced by supernatant of *P. acidilactici* Z102 above the absorbance reading produced by MRS) × 100].

‡ Cross-reactivity defined as [(antibody binding inhibition produced by a culture supernatant above the antibody binding inhibition produced by supernatant of *P. acidilactici* Ped⁻/antibody binding inhibition produced by supernatant of *P. acidilactici* X13 above the antibody binding inhibition produced by supernatant of *P. acidilactici* Ped⁻) × 100].

to a previously described scheme (Martínez *et al.*, 1998). Rabbits were bled via marginal ear veins on days 28 and 63 and a final bleed was performed on day 72 by cardiac puncture.

ELISAs. Most of the procedures were performed as previously described (Martínez *et al.*, 1998). Briefly, for antisera titration flat-bottom polystyrene microtitre plates (Maxisorp) were coated overnight (4 °C) with 100 µl PH1–OAG (5 µg ml⁻¹) in 0.1 M sodium carbonate/bicarbonate buffer, pH 9.6 (coating buffer, CB). Plates were washed three times with 300 µl washing solution (0.05% Tween 20 in PBS). Wells were blocked for 30 min at 37 °C with 300 µl 1% (w/v) OA (grade III) in PBS (OA-PBS) and then washed six times. Next, 50 µl serially diluted serum was added to each well and incubated for 1 h at 37 °C. Unbound antibody was removed by washing four times and 100 µl goat anti-rabbit IgG peroxidase conjugate (diluted 1:500 in OA-PBS) was added to each well. Plates were incubated for 30 min at 37 °C, washed eight times and bound peroxidase was determined with ABTS [2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid)] as substrate by measuring A₄₀₅. The titre of each serum was arbitrarily set as the maximum dilution that yielded at least twice the absorbance of the same dilution of non-immune control serum.

For antiserum specificity and sensitivity to PedA1, four types of ELISA were designed. In non-competitive indirect ELISA

(NCI-ELISA) wells of microtitre plates were coated with 100 µl of different concentrations of the analytes. The plates were maintained for 3 h at 40 °C, then blocked and washed as described for the antiserum titration procedure. Next, 50 µl antiserum, diluted 1:200 in PBS, was added and the plates incubated for 1 h at 37 °C. After the washing step and addition of the goat anti-rabbit IgG peroxidase conjugate (diluted 1:500 in OA-PBS), the bound peroxidase was determined with ABTS substrate as described above. In competitive indirect ELISA (CI-ELISA), microtitre plates were coated with 100 µl of either PH1–OAG or PedA1, both at 0.75 µg ml⁻¹ in CB, and blocked and washed as described for the antiserum titration procedure. Next, 50 µl of different concentrations of the analytes was simultaneously incubated with 50 µl antiserum (diluted 1:250 in PBS) for 1 h at 37 °C. After the washing step and addition of the goat anti-rabbit IgG peroxidase conjugate (diluted 1:500 in OA-PBS), the bound peroxidase was determined with the 3,3',5,5'-tetramethylbenzidine (TMB) liquid substrate system (Sigma). A₄₅₀ was measured after acidification of the samples with 100 µl of a 1 M H₂SO₄ stopping solution. Relative antibody affinity was arbitrarily designated as the bacteriocin concentration required to inhibit antibody binding by 50%.

A competitive direct ELISA (CD-ELISA) was also developed. In this assay, the plates were coated overnight by air drying at

40 °C with 125 µl PH1–KLH-generated antibodies (diluted 1:100 in CB). After washing and blocking, 50 µl of the analytes and 50 µl of either PH1–HRP (diluted 1:100) or PedA1–HRP (diluted 1:100) in OA-PBS, was added to each well consecutively. After 1 h incubation at 37 °C, the plates were washed and the amount of bound peroxidase was determined by addition of the TMB substrate. For sandwich ELISA (S-ELISA), the plates were coated overnight by air drying at 40 °C with 125 µl of the goat anti-rabbit IgG Fc fragment (diluted 1:540 in CB). After washing and blocking, 50 µl PH1–KLH-generated antibodies diluted 1:150 in PBS were added. After 30 min incubation at 37 °C the plates were washed and 50 µl standards, control samples or samples were added per well. After 45 min incubation at 37 °C, the plates were washed and 50 µl mouse PH2–KLH-generated antibodies (J. M. Martínez, M. I. Martínez, C. Herranz, L. M. Cintas, J. M. Rodríguez & P. E. Hernández, unpublished results) diluted 1:100 in PBS was added. After another 45 min incubation at 37 °C, the washing step and the addition of goat anti-mouse IgG peroxidase conjugate (diluted 1:500 in OA-PBS), the plates were washed and the amount of bound peroxidase was determined by addition of TMB substrate.

Protein slot-blot assay. Eighty microlitres of different concentrations of PH1–OAG, PH2–OAG, OA, PedA1 or pure nisin A dissolved in MRS, and the same volume of neutralized and filter-sterilized supernatants from 16 h cultures of various LAB, were deposited onto a nitrocellulose membrane (pore size, 0.2 µm; Bio-Rad) in a Bio-dot SF microfiltration apparatus (Bio-Rad) and the membrane was processed essentially as described previously (Martínez *et al.*, 1998). The membrane was incubated with 30 ml PH1–KLH antiserum (diluted 1:200 in PBS) and further incubated with 30 ml goat anti-rabbit IgG peroxidase conjugate (diluted 1:5000 in blocking solution). Specific antigens for PH1–KLH-generated antibodies were visualized by chemiluminescence with the ECL detection kit (Amersham). The light emission was detected by a short exposure of the membrane to a blue-light-sensitive autoradiography film (Hyperfilm ECL, Amersham).

Protein electrophoresis, Western hybridization and overlay assay. Fifteen microlitres of pure PedA1 (0.02, 0.001, 0.005 and 0.002 µg ml⁻¹) and the supernatants from 16 h cultures of *Pediococcus acidilactici* 347 (Ped⁺), *P. acidilactici* 347, *Enterococcus faecium* T136, *E. faecium* P13, *Lactobacillus sakei* 706, *Lb. sakei* LTH673, *Lactococcus lactis* BB24, *Lc. lactis* FI9181 (*Lc. lactis* IL1403 derivative, producer of PedA1; N. Horn, M. I. Martínez, J. M. Martínez, P. E. Hernández, M. J. Gasson, J. M. Rodríguez & H. Dodd, unpublished results) and *Lc. lactis* IL1403 were subjected to Tricine-SDS-PAGE (Shägger & Von Jagow, 1987). After protein electrophoresis, one of the gels was blotted onto a PVDF membrane (pore size, 0.2 µm; Bio-Rad) by the application of an electrical potential of 80 mV for 50 min. Further blocking and washing of the PVDF membrane, treatment with the PH1–KLH antiserum and goat anti-rabbit IgG peroxidase conjugate and visualization of the expected antigen–antibody interaction by chemiluminescence with an ECL detection kit were performed as described above for the protein slot-blot assay. To determine the antimicrobial activity of pure PedA1 and the supernatants of the tested strains, an overlay assay was performed (Bhumia *et al.*, 1987). After the gel was fixed, washed and drained it was overlaid with the indicator strain *Lb. sakei* ATCC 15521 (1 × 10⁵ c.f.u. ml⁻¹ in soft agar) and incubated overnight at 30 °C.

Micro-organisms, media and bacteriocin assays. The LAB tested for PedA1 production or antibody cross-reactivity are listed in Table 1. All micro-organisms were propagated in

MRS broth (Oxoid) at 32 °C and the supernatants were obtained by centrifugation at 12000 g for 10 min at 4 °C, adjusted to pH 6.2 with 1 M NaOH, filtering through 0.2 µm-pore filters (Whatman) and stored at –20 °C until use. The antimicrobial activity of the supernatants was evaluated by an agar diffusion test (ADT) and, when stated, by a microtitre plate assay (MPA). The ADT and the MPA assays were performed as described previously (Martínez *et al.*, 1998).

Purification of PedA1. The antimicrobial activity of *P. acidilactici* 347, used as the source of PedA1, was purified to homogeneity as described previously (Cintas *et al.*, 1997; Horn *et al.*, 1998; Martínez *et al.*, 1998). The final concentration of pure bacteriocin was estimated by using the extinction coefficient of PedA1 (A_{280} of 3.1 corresponds to 1 mg ml⁻¹).

RESULTS

Sensitivity of rabbit anti-peptide antibodies for PedA1

The PH1 fragment conjugated to KLH was used in the immunization of rabbits. On day 72 of the immunization process and after six doses of the immunogen had been administered, the animals had apparent titres in the serum of 1:25 600, 1:51 200, and 1:102 400. The serum with the highest titre for fragment PH1 was used throughout. The sensitivity of the anti-PH1–KLH antibodies for PedA1 was initially determined by NCI-ELISA. As seen in Fig. 1, the polyclonal antibodies showed a higher recognition of the peptide fragment

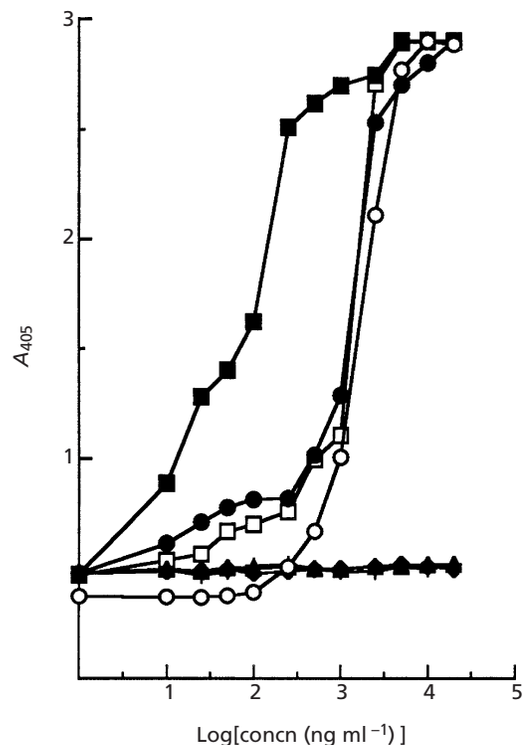


Fig. 1. Results of NCI-ELISA for the detection of PH1–OAM (■), PH1–OAG (●), PH2–OAG (▲), purified PedA1 in CB (□), purified PedA1 in MRS (○), pure nisin A in CB (◆) and pure OA in CB (+).

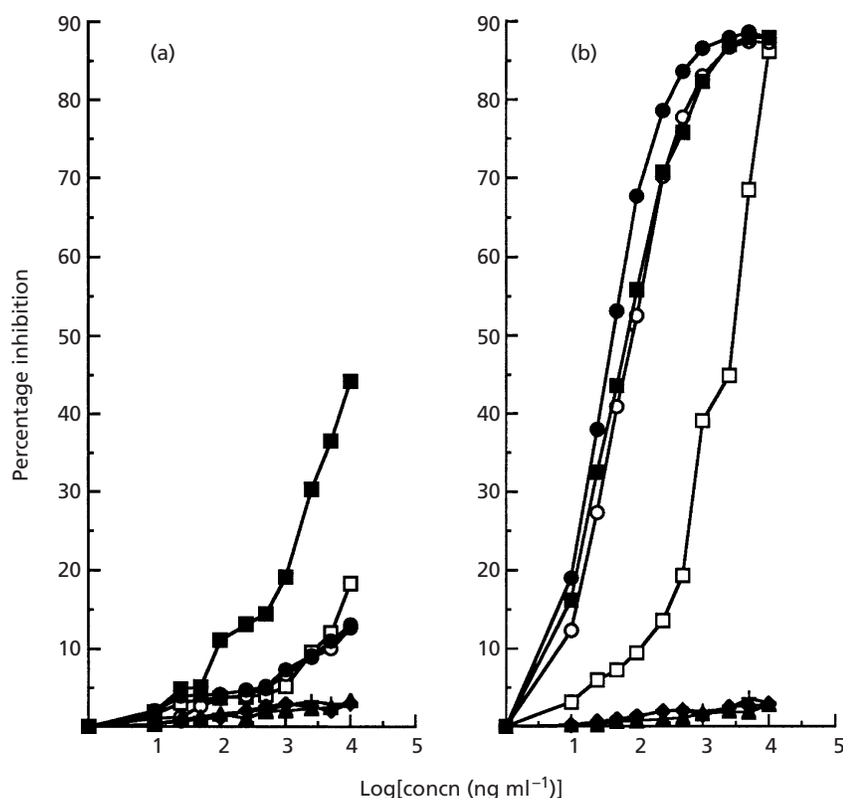


Fig. 2. Results of CI-ELISA for recognition of the PH1 fragment in PBS (■) and MRS (●), and purified PedA1 in PBS (□) and MRS (○), as well as for recognition of the PH2 fragment (▲), pure OA (+) and pure nisin A (◆) in PBS, using microtitre ELISA plates coated with PH1-OAG (a) or purified PedA1 (b).

conjugated to OA through the maleimide method (PH1-OAM) than to the fragment conjugated to OA by the glutaraldehyde method (PH1-OAG), suggesting that a large number of antibodies recognized the same chemical bridge between the peptide fragment and KLH. More importantly, the antibodies recognized PedA1 present in the wells of the microtitre plates, although recognition was higher for PedA1 in CB ($0.05 \mu\text{g ml}^{-1}$) than in MRS broth ($0.5 \mu\text{g ml}^{-1}$). Such antibodies could not detect the presence of equivalent concentrations of OA, PH2-OAG or pure nisin A in the wells of the microtitre plates.

The specificity for PedA1 of the PH1-KLH-generated antibodies was also investigated by a CI-ELISA. The mean detection limit for fragment PH1 was much higher on plates coated with PedA1 ($0.01 \mu\text{g ml}^{-1}$; Fig. 2b) than with PH1-OAG ($0.1\text{--}1 \mu\text{g ml}^{-1}$; Fig. 2a), while for PedA1 the detection limits were $1\text{--}2.5 \mu\text{g ml}^{-1}$ in PBS and $1 \mu\text{g ml}^{-1}$ in MRS broth on plates coated with PH1-OAG, and $0.01\text{--}0.025 \mu\text{g ml}^{-1}$ in PBS and $0.01 \mu\text{g ml}^{-1}$ in MRS broth on plates coated with PedA1. The amount of free PH1 required for 50% binding inhibition was not measurable on plates with PH1-OAG: $0.1 \mu\text{g ml}^{-1}$ in PBS and $0.05 \mu\text{g ml}^{-1}$ in MRS broth on plates coated with PedA1. Similarly, the amount of free PedA1 for 50% binding inhibition was not measurable on plates coated with PH1-OAG: $2.5 \mu\text{g ml}^{-1}$ in PBS and of $0.01 \mu\text{g ml}^{-1}$ in MRS broth on plates coated with PedA1. Thus, the performance of this assay is tremendously improved when PedA1 was used as the solid-phase antigen, thus increasing the relative antibody-binding

affinity and decreasing the free PedA1 concentration required to inhibit antibody binding.

A CD-ELISA was also developed to determine the specificity of the PH1-KLH-generated antibodies for PedA1. In this assay, the plates were coated with an appropriate dilution of the serum of immunized animals and with HRP conjugated to either PH1 or PedA1. However, PedA1 did not effectively compete with either PH1-HRP or PedA1-HRP for binding to the antibody-coated microtitre wells (results not shown). Similarly, the development of S-ELISA for evaluation of the specificity of the anti-peptide antibodies for PedA1 was also unsuccessful (results not shown), since it was not possible to observe an increase in the absorbance of the assay with increasing concentrations of PedA1 (until $10 \mu\text{g ml}^{-1}$).

Immunoreactivity of the rabbit anti-peptide antibodies to different bacteriocins

The specificities of the serum polyclonal antibodies in neutralized and filter-sterilized supernatants of 16-h-old cultures of representative LAB strains were evaluated by NCI- and CI-ELISA (Table 1). The antibodies in both immunoassays reacted with the supernatants of the *P. acidilactici* strains 347, Z102, A172, X13 and P20, all potential PedA1 producers since they have been reported to harbour the *pedA* gene by the use of rapid molecular biology techniques (Rodríguez *et al.*, 1997), but did not react with the supernatant of a derivative of *P. acidilactici* 347 (Ped⁻), a non-PedA1 strain (Martínez *et*

Table 2. Multiple sequence alignment of mature PedA1 with other Class IIa bacteriocins

The bar refers to the sequence of fragment PH1 of PedA1 against which the polyclonal antibodies were generated. Sequences were obtained from the following: piscicocin V1a, Bhugaloo-Vial *et al.* (1996); divercin V41, Metivier *et al.* (1998); mundticin, Bennik *et al.* (1998); all others, Martínez *et al.* (1998).

Bacteriocin	Sequence	
	1 5 10 15 20 25 30 35 40 45	
Pediocin PA-1	----KYYGNVTCGKHSCSVDWGKATTCIINNGAMAWATGGHQGNHKC----	44
Enterocin A	TTHSGKYYGNVYCTKNKCTVDWAKATTCIAGMSIGGFLGGAI-----	43
Enterocin P	---ATRSYNGVYCNNSKCWVNWGEAKENIAGIVISGWASGLAGMGH-----	44
Sakacin A	----ARSYNGVYCNKKCWVNRGEATQSIIGGMISGWASGLAGM-----	41
Sakacin P	----KYYGNVHCGKHSCVDWGTAINIGNNAAANWATGGNAGWNK-----	43
Carnobacteriocin BM1	----AISYNGVYCNKEKCWVNKAENKQAITGIVIGGWASGLAGMGH-----	43
Leucocin A-UAL187	----KYYGNVHCTKSGCSVNWG----EAFSAGVHRLANGGNGFW-----	37
Mesentericin Y105	----KYYGNVHCTKSGCSVNWG----EAASAGIHLRANGGNGFW-----	37
Carnobacteriocin B2	----VNYNGVSCSKTKCSVNWGQAFQERYTAGINSFVSGVASGAGSIGRRP	48
Bacteriocin 31	----ATYYNGLYCNKQKCWVDWNKASREIGKIIVNGWVQHGPWAPR-----	43
Piscicocin V1a	----KYYGNVSCNKNKCTVDWSKAIGIIGNNAAANLTTGGAAGWNKG----	44
Mundticin	----KYYGNVSCNKKGCSVDWGKAIGIIGNNSAANLATGGAAGWSK-----	43
Divercin V41	----TKYYGNVYCNKCKWVDWGQASGCIQTVVGGWLGAIPG--KC----	43

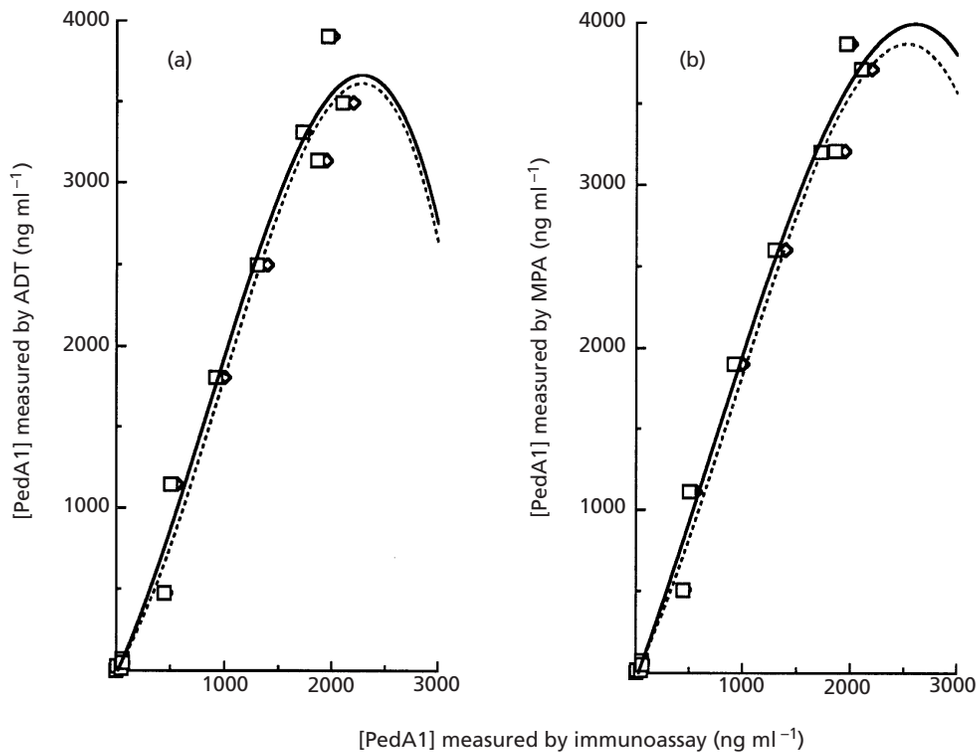


Fig. 3. Relationship between PedA1 concentrations by ADT (a) and MPA (b), and by NCI- (□) and CI-ELISA (◇). The resulting regression equations are: (a) $Y_{\text{NCI-ELISA}} = -19.211 + 1.4305x + 8.7312e - 4x^2 - 3.4726e - 7x^3$ ($r^2 = 0.986$), $Y_{\text{CI-ELISA}} = -9.7686 + 1.0798x + 1.1410e - 3x^2 - 4.0248e - 7x^3$ ($r^2 = 0.985$); and (b) $Y_{\text{NCI-ELISA}} = -37.931 + 1.7645x + 4.3088e - 4x^2 - 1.9740e - 7x^3$ ($r^2 = 0.990$), $Y_{\text{CI-ELISA}} = -28.544 + 1.4030x + 7.1950e - 4x^2 - 2.6280e - 7x^3$ ($r^2 = 0.991$).

al., 1998). However, the reactivity for the *P. acidilactici* P20 strain was much lower than for the other *P. acidilactici* strains. A much lower reactivity, negligible

reactivity or no reactivity was observed with the supernatants of *Pediococcus pentosaceus* FBB61, a pediocin A producer (Piva & Headon, 1994), *E. faecium*

Table 3. PedA1 concentration in culture supernatants of PedA1 producers as determined by CI-ELISA

<i>P. acidilactici</i> strain	PedA1 concentration (ng ml ⁻¹)*
347	1972
Z102	2005
A172	1946
X13	2200
P20	12
347 Ped ⁻ (non-PedA1 producer)	NR

* Mean value from four independent determinations. NR, No reactivity.

T136, an enterocin A and B producer (Casaus *et al.*, 1997), *E. faecium* P13, an enterocin P producer (Cintas *et al.*, 1998), *Lb. sakei* LTH673, a sakacin P producer (Tichaczek *et al.*, 1994), *Lb. sakei* 706, a sakacin A producer (Holck *et al.*, 1992), *E. faecium* L50, an enterocin L50A and L50B producer (Cintas *et al.*, 1998), *Lb. sakei* 148, a lactocin S producer (Rodríguez *et al.*, 1995a), *Lc. lactis* BB24, a nisin A producer (Rodríguez *et al.*, 1995b) and *Lc. lactis* MG1614, a non-bacteriocin producer (Gasson, 1983). Table 2 shows the amino acid sequence alignment of mature PedA1 with other Class IIa bacteriocins. It is important to note that enterocin A, sakacin P, leucocin A, mesentericin Y105, pisciocin V1a, mundticin and divercin V41 share the longer N-terminal consensus amino acid motif (KYYGNGVxC) of the pediocin family of bacteriocins, while enterocin P, sakacin A, carnobacteriocin BM1, carnobacteriocin B2 and bacteriocin 31 share the shorter (YGNGVxC) motif.

The concentration of PedA1 in the supernatant of a 16-h-old culture of *P. acidilactici* 347 grown in MRS broth was evaluated by NCI- and CI-ELISA, and by the bioassay-based antimicrobial tests ADT and MPA. The concentrations of PedA1 were determined to be 3198 ng ml⁻¹ by the MPA and 3310 ng ml⁻¹ by ADT, while the PedA1 concentrations were lower but similar in NCI-ELISA (1724 ng ml⁻¹) and CI-ELISA (1972 ng ml⁻¹). With the use of the previously described PH2-KLH-generated antibodies (Martínez *et al.*, 1998), the concentrations of PedA1 in the same supernatant were determined to be 1670 ng ml⁻¹ by NCI-ELISA, 1689 ng ml⁻¹ by CI-ELISA and 1701 ng ml⁻¹ by CD-ELISA. The differences in PedA1 concentrations as determined by the immunoassays and the bioassays may be attributed to the fact that the purified PedA1 used as the standard in both assays may have a lower activity than the native PedA1 present in the supernatants. For this reason, the haloes generated by the native PedA1 may be larger than those generated by the purified PedA1, magnifying the true concentration of PedA1 in the supernatants. Since the immunoassays detect and quantify PedA1 independently of their activity, the results obtained with the immunoassays are probably more suitable for evaluating the real concentrations of PedA1 in the supernatants of

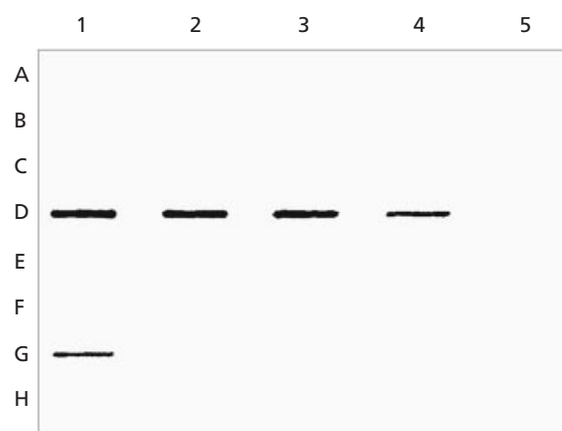


Fig. 4. Slot-blot of various standards and supernatants from LAB. Row A: supernatants from *P. acidilactici* 347 (lane 1), *P. acidilactici* Z102 (2), *P. acidilactici* X13 (3), *P. acidilactici* A172 (4) and *P. acidilactici* P20 (5). Row B: supernatants from *P. acidilactici* 347 (Ped⁻) (1), *E. faecium* L50 (2), *P. pentosaceus* FBB61 (3), *E. faecium* P13 (4) and *E. faecium* T136 (5). Row C: supernatants from *E. faecalis* INIA4 (1), *Lb. sakei* 148 (2), *Lc. lactis* BB24 (3), *Lb. sakei* 706 (4) and *Lb. sakei* LTH673 (5). Row D: PH1-OAG at 5 (1), 2.5 (2), 1 (3), 0.5 (4) and 0 µg ml⁻¹ (5). Rows E, PH2-OAG; F, pure OA; G, purified PedA1; H, pure nisin A (concentrations in lanes 1–5 in rows E–H as for row D, all in MRS).

the producer strains. The relationship between ADT (Fig. 3a) and MPA (Fig. 3b), and NCI-ELISA and CI-ELISA, using PH1-KLH-generated antibodies was also evaluated. Significant ($P < 0.001$) regression equations could be established for PedA1 concentrations detected by ADT and the immunoassays (r^2 , 0.986–0.985) and by MPA and the immunoassays (r^2 , 0.991–0.990). CI-ELISA was also used to determine the concentrations of PedA1 in the supernatants of the five independently isolated *P. acidilactici* strains (Table 3). While most of the strains displayed a similar production of PedA1, *P. acidilactici* P20 produced a significantly lower concentration of this bacteriocin, resolving the usefulness of this assay to detect and quantify low concentrations of PedA1. The PH1-KLH-generated antibodies and CI-ELISA also demonstrated their usefulness to quantify the heterologous production of PedA1 (from 30 to 1908 ng ml⁻¹) by a number of genetically modified *Lc. lactis* IL1403 derivatives (N. Horn, M. I. Martínez, J. M. Martínez, P. E. Hernández, M. J. Gasson, J. M. Rodríguez & H. Dodd, unpublished results).

The immunoreactivity of the PH1-KLH-generated antibodies to different conjugates, standards and bacteriocins was also evaluated by protein slot-blotting. Results of a slot-blot peptide assay of standards and supernatants of 16-h-old cultures of various LAB strains probed with the cited antibodies (Fig. 4), indicated that the antibodies recognized fragment PH1, but not fragment PH2, OA or pure nisin A. Similarly, the antibodies recognized pure PedA1 at a detection limit of 5 µg ml⁻¹, with a non-detectable response signal in the supernatants of *P. acidilactici* 347, X13, Z102, A172 and P20,

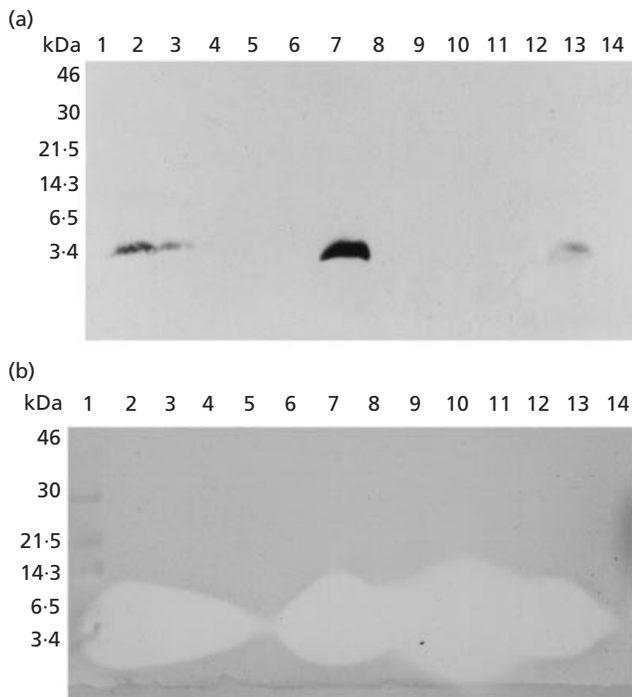


Fig. 5. Tricine-SDS-PAA gel: (a) Western blot and (b) overlaid with the indicator strain *Lb. sakei* ATCC 15521. Lane 1, molecular mass marker. Lanes 2 to 5 contain pure PedA1 at 0.02 (2), 0.01 (3), 0.005 (4) and 0.002 $\mu\text{g ml}^{-1}$ (5). Lanes 6 to 14 contain supernatants from *P. acidilactici* 347 Ped⁻ (6), *P. acidilactici* 347 (7), *E. faecium* T136 (8), *E. faecium* P13 (9), *Lb. sakei* 706 (10), *Lb. sakei* LTH673 (11), *Lc. lactis* BB24 (12), *Lc. lactis* FI9181 (13) and *Lc. lactis* IL1403 (14).

as well as in the supernatants of different bacteriocin producer or non-producer LAB strains. Finally, a Western hybridization assay was also evaluated to determine the immunoreactivity of rabbit anti-peptide antibodies to PedA1. The results obtained with purified PedA1 and supernatants from 16-h-old cultures of various LAB strains probed with PH1-KLH-generated antibodies (Fig. 5a) clearly indicated that the antibodies recognized PedA1 at a detection limit of 0.01 $\mu\text{g ml}^{-1}$, being able to detect the presence of this bacteriocin in the supernatants of *P. acidilactici* 347 and *Lc. lactis* FI9181, with no reactivity from supernatants of Class IIa bacteriocin producer or non-producer LAB strains. The presence of active bacteriocins in the supernatants was demonstrated by the results of the overlay assay as shown in Fig. 5b.

DISCUSSION

Contrary to the situation for other analytes, the identification and detection of bacteriocinogenic LAB and bacteriocins has relied mostly on the use of bioassay-based tests in which non-specificity is a major drawback. Moreover, although highly specific immunochemistry-based methods have been developed and routinely used as analytical tools in many areas of research, surprisingly the impact of these techniques in the bacteriocin research

field has been marginal (Suárez *et al.*, 1996a). Although other reports have described the generation of antibodies against nisin A and nisin Z (Falahaee *et al.*, 1990; Stringer *et al.*, 1995; Suárez *et al.*, 1996a, b; Bouksaim *et al.*, 1997), pediocin AcH (Bhunia *et al.*, 1990), and against pediocin RS2 (Bhunia, 1994), antibodies were generated against the whole bacteriocin molecule either alone or conjugated to a carrier protein. However, the lack of purified bacteriocins means the availability of these molecules for use as immunogens is poor and the low molecular masses (< 5000 Da) of these compounds make them poorly immunogenic or non-immunogenic. In addition, peculiar characteristics of bacteriocin molecules, such as their hydrophobicity, the formation of intrachain disulfur rings and, in the case of lantibiotics, the presence of modified amino acids, might also interfere with the development of immunoassays. Antibodies generated against a short chemically synthesized fragment of a bacteriocin of interest may be useful to generate antibodies of predetermined specificity (Martínez *et al.*, 1998).

Since PedA1 belongs to the pediocin family of bacteriocins, it exhibits strong amino acid sequence homology with other bacteriocins at the N terminus (Table 2). Moreover, since the PH1 fragment was evaluated to be a highly potential immunogenic fragment, according to its hydrophilicity and antigenic index determined by the use of the sequence analysis software package (Devereux *et al.*, 1984), antibodies generated against such a fragment could behave as specific for PedA1 or could display a significant cross-reactivity against other Class IIa bacteriocins, making them valuable for detection, quantification and/or immunopurification of a large number of bacteriocins. Accordingly, the sensitivity and specificity of antibodies generated against the 1–9-N-terminal amino acid fragment of mature PedA1 were evaluated against closely related bacteriocins through the development of sensitive immunoassays.

The specificity of PH1-KLH-generated polyclonal antibodies for PedA1 was evaluated by NCI-, CI-, CD- and S-ELISA, protein slot-blotting and Western blotting. The limit of detection of PedA1 in NCI-ELISA (Fig. 1) was lower in CB than in MRS broth, reflecting the masking effect of the latter in the detection of PedA1 in this assay. Results obtained with CI-ELISA clearly indicated that coating of the plates with PedA1 instead of PH1-OAG enhanced the detection of free PedA1 (Fig. 2). Moreover, the limit of detection of the immunoassay was improved to 0.01 $\mu\text{g PedA1 ml}^{-1}$ in MRS broth and 0.01–0.025 $\mu\text{g ml}^{-1}$ in PBS, and the 50% binding inhibition was achieved with 0.1 $\mu\text{g PedA1 ml}^{-1}$ in MRS broth and 2.5 $\mu\text{g ml}^{-1}$ in PBS. As shown also in Fig. 2, the competition curves for PedA1 in PBS and MRS broth differed notably, showing higher binding inhibition values in MRS broth. This effect was also observed with antibodies against the C-terminal fragment of PedA1 (Martínez *et al.*, 1998). The limit of detection and sensitivity of NCI- and CI-ELISA developed for PedA1 were in the range of the values reported for nisin A

(Falahaee *et al.*, 1990; Suárez *et al.*, 1996a, b) but were more effective than those obtained for pediocin RS2 (Bhunja, 1994).

However, contrary to what has been observed with antibodies against nisin A (Suárez *et al.*, 1996a, b) and against the C-terminal fragment of PedA1 (Martínez *et al.*, 1998), PedA1 did not effectively compete with either PH1-HRP or PedA1-HRP for binding to the antibody-coated microtitre wells in CD-ELISA. These results heighten the importance of the development of proper immunoassay formats for detection of each bacteriocin and confirm our previous observation that mice serum and ascites antibodies against the PH1-KLH conjugate did not recognize the whole PedA1 molecule by the use of CD-ELISA (Martínez *et al.*, 1997). Similarly, the development of S-ELISA for evaluation of the specificity of PH1-KLH-generated antibodies for PedA1 was also unsuccessful. It is possible that because of the short length of PedA1, the capture antibodies against its N-terminal end mask the recognition of the molecule by the detection antibodies generated against its C-terminal end.

PH1-KLH-generated antibodies showed a high affinity for PedA1 in the supernatants of *P. acidilactici* 347, Z102, A172 and X13 grown in MRS broth, and a small cross-reactivity to the supernatant of *P. acidilactici* P20 (Table 1), previously reported to harbour the *pedA* gene for production of PedA1 (Rodríguez *et al.*, 1997). The antibodies did not show a significant cross-reactivity with cell culture supernatants from enterocin A, enterocin P, sakacin P and sakacin A producer strains, bacteriocins which share the longer (KYYGNGxG) or shorter (YGNGVxG) consensus amino acid motifs with PedA1 (Table 2). This absence of cross-reactivity is not surprising, since it has been reported that closely related proteins have been distinguished by the use of antisera as probe for a specific substrate within the protein molecule (Groome, 1994) and that changes in a single amino acid residue drastically affects protein recognition (Rolland *et al.*, 1995). PedA1 produced by *P. acidilactici* 347, Z102, A172, X13 and P20 grown in MRS broth was quantified by CI-ELISA (Table 3). The lower level of production of PedA1 by *P. acidilactici* P20 may be reasonably explained by genetic defects perhaps affecting expression, processing or secretion of this bacteriocin. The use of PH1-KLH-generated antibodies as probe for the identification and quantification of PedA1 in the supernatants of bacteriocin-producing strains, may be valuable as a tool to avoid the use of complex biochemical techniques involved in the purification to homogeneity and determination of the amino acid sequence of unknown antimicrobial activities.

When the immunoreactivity of the PH1-KLH-generated antibodies was evaluated by protein slot-blotting, the antibodies recognized PedA1 at a detection limit of 5 µg ml⁻¹ (Fig. 4). The low level of detection of PedA1 in this assay compared to the other immunoassays may reflect differences in PedA1 solubility, conformation of the bacteriocin, preferential attachment of the bacteriocin to the membrane by its N-terminal end, aggregation of

the bacteriocin molecules with components of MRS broth and oxidation of amino acid residues. However, using the same assay and conditions, the limit of detection of PedA1 in MRS broth was found to be 2.5 µg ml⁻¹ when antibodies against the C-terminal end of PedA1 were used (Martínez *et al.*, 1998). This allows us to hypothesize that perhaps the preferential attachment of PedA1 to the nitrocellulose membrane through its N-terminal end is masking in part the recognition of the molecule by PH1-KLH-generated antibodies. When the immunoreactivity of the rabbit polyclonal antibodies to PedA1 was ascertained by Western blotting, the limit of detection of PedA1 was determined to be 0.01 µg ml⁻¹ (Fig. 5a). The absence of cross-reactivity of the antibodies against the supernatants of various pediocin-like bacteriocin producer or non-producer LAB strains heightens the importance and significance of the Western blotting technique for the rapid detection and identification of PedA1 in the supernatants of producer strains.

The strategy of using a synthetic peptide for generating antibodies against a bacteriocin epitope(s) sharing a highly conserved amino acid sequence with closely related bacteriocins has been shown to be both conceptually simple and practically convenient. Furthermore, the sensitivity and specificity of PH1-KLH-generated rabbit polyclonal antibodies for PedA1 and the absence of cross-reactivity against Class IIa bacteriocins or other bacteriocins, either lantibiotic or non-lantibiotic, suggest that all the techniques described here for selection of the peptide fragment, carrier molecule, conjugation methods and immunoassay development can be used as models for the generation of antibodies against other bacteriocins of interest in the food industry. Potential specific applications of these antibodies include the rapid identification and isolation of PedA1 producer strains from many sources, an application based on reports on the production of PedA1 by a vegetable-associated *Pediococcus parvulus* strain (Bennik *et al.*, 1997) and by *Lactobacillus plantarum* WHE 92 isolated from cheese (Ennahar *et al.*, 1996). The PH1-KLH-generated antibodies may also serve as a tool for studies on the regulation of PedA1 production, processing-secretion, identification of target cell specificity-determining regions, evaluation of structure-function relationships and analysis by ELISA of PedA1 in foods. Of great interest is the availability of specific antibodies to well characterized bacteriocins for studies on the expression of different bacteriocins in heterologous hosts (Horn *et al.*, 1998; N. Horn, M. I. Martínez, J. M. Martínez, P. E. Hernández, M. J. Gasson, J. M. Rodríguez & H. Dodd, unpublished results). The antibodies described in this work can also be used for the purification of PedA1 to homogeneity in a single step by immunoaffinity chromatography.

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