A genetically engineered protein domain binding to bacterial murein, archaeal pseudomurein, and fungal chitin cell wall material

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Abstract The major murein and pseudomurein cell wall-binding domains, i.e., the Lysin Motif (LysM) (Pfam PF01476) and pseudomurein cell wall-binding (PMB) (Pfam PF09373) motif, respectively, were genetically fused. The fusion protein is capable of binding to both murein- and pseudomurein-containing cell walls. In addition, it also binds to chitin, the major polymer of fungal cell walls. Binding is influenced by pH and occurs at a pH close to the pI of the binding protein. Functional studies on truncated versions of the fusion protein revealed that murein and chitin binding is provided by the LysM domain, while binding to pseudomurein is achieved through the PMB domain.

Keywords Murein · Pseudomurein · Chitin · Domain

Introduction

Cells of most Gram-positive bacteria and methanogenic archaea of the orders Methanobacteriales and Methanopyrales are surrounded by a protective cell wall made of murein or pseudomurein, respectively (Kandler and König 1978, 1998; Steenbakkers et al. 2006; Visweswaran et al. 2010). Although murein and pseudomurein perform a similar function in these organisms, some fundamental differences exist in their composition and architecture. Murein is made of polymers of β(1-4) linked N-acetylmuramic acid and N-acetyl-D-glucosamine (NAG), whereas pseudomurein is composed of chains of β(1,3)-linked NAG and N-acetyltalosaminuronic acid residues. Murein and pseudomurein are substrates for a variety of cell wall hydrolases involved in various physiological processes like cell separation and autolysis. To facilitate in these biological processes, most cell wall hydrolases are equipped with one or more cell wall-binding domains that non-covalently attach the enzymes to their polymeric substrates.

In this study, a fusion protein was generated by the union of two cell wall-binding domains, i.e., the Lysin Motif (LysM) domain, which binds to bacterial cell walls, and the pseudomurein cell wall-binding (PMB) domain, which binds to pseudomurein-containing archaeal cell walls. The LysM domain is a ubiquitous, murein-binding domain present in proteins of both prokaryotes and eukaryotes (Audouy et al. 2006, 2007; Bosma et al. 2006; Audouy et al. 2007). For instance, LysM domains are currently employed in the development of pneumococcal and influenza vaccines (Audouy et al. 2006, 2007; Bosma et al. 2006; Saluja et al. 2010). They are also used for the display of various heterologous proteins on the bacterial cell surface (Steen et al. 2003; Bosma et al. 2006; Okano et al. 2008; Tarahomjoo et al. 2008; Hu et al. 2010). In bacteria, LysM domains are present in murein hydrolases helping in the non-covalent attachment of the enzymes to the murein layer, thus facilitating substrate hydrolysis. Functional studies suggest that LysM domains
recognize the NAG moiety of murein (Ohnuma et al. 2008; Petutschnig et al. 2010). In plants, LysM domains play a vital role in the symbiotic relationship between plants and some of their bacterial hosts (Mulder et al. 2006; Radutoiu et al. 2007; Wan et al. 2008). Plant LysM domains may also bind to chitin, the cell wall polymer of fungi, as part of the plants’ defense mechanism against pathogenic fungi (Ohnuma et al. 2008; Petutschnig et al. 2010).

On the other hand, the PMB domain is narrowly distributed and present only in a few methanogenic archaeal proteins and in two archaea-specific viral hydrolases (PeiP and PeiW) (Kiener et al. 1987; Stax et al. 1992; Pfister et al. 1998; Luo et al. 2001, 2002; Steenbakkers et al. 2006; Visweswaran et al. 2010, 2011a, b). Until now, there are no experimental reports demonstrating the occurrence of LysM domains in archaea (Buist et al. 2008) or PMB domains in bacteria. The function of the PMB domain in archaeal hydrolases is analogous to that of the LysM domains in bacterial hydrolases, i.e., binding of the enzyme to its polymeric substrate, in this case the pseudomurein layer in methanogenic archaeal cell walls (Visweswaran et al. 2010, 2011a, b). Deletion of the PMB domain in one of the pseudomurein endoisopeptidases, PeiW, led to loss of binding of the enzyme to the host cell wall (Steenbakkers et al. 2006). In analogy to the use of the LysM domain, the PMB domain might be employed for the surface display of heterologous proteins on pseudomurein-containing cell walls.

To obtain the combined properties of both the LysM and PMB domains and also to increase the number of binding partners, we developed a fusion protein by genetic fusion of the three C-terminal LysM motifs of a major autolysin, AcmA from Lactococcus lactis, and three C-terminal PMB motifs of the surface S-layer protein MTH719 from Methanothermobacter thermautotrophicus (see Fig. 1). The genetically engineered fusion protein was investigated with respect to binding to bacterial and methanogenic archaeal cell walls and the effect of pH thereon. Interestingly, the fusion protein was shown to bind to chitin flakes and to fungal cell walls through its LysM domain.

Materials and methods

Strains and plasmids

The LysM_{AcmA}, PMB_{MTH719}, and the M–P fusion proteins used in this study were constructed in pBADcLIC-GFP plasmid, bearing an arabinose-inducible promoter, using the ligation-independent cloning method described previously (Geertsma and Poolman 2007). Oligos used for the amplification of specific DNA fragments are shown in Table 1. Primers LR and PF contain homologous nucleotide sequences to the PMB_{MTH719} domain and the LysM_{AcmA} domain, respectively, to facilitate overlap PCR. Plasmids annealed with DNA fragments of individual constructs of LysM_{AcmA}, PMB_{MTH719} domains, and the M–P fusion protein were used for transformation of Escherichia coli Rosetta gami 2 (Novagen, Darmstadt, Germany) using the heat shock method (van Die et al. 1983). Transformed cells were plated on selective TY 1.5% (w/v) agar plates and transformants were checked by colony PCR and plasmid DNA sequencing (ServiceXS, Leiden, the Netherlands) for the correct sequence and right orientation of the desired gene products. For overexpression of specific constructs, E. coli cells were grown with shaking at 37°C in TY broth (Difco, Sparks, MD, USA), containing ampicillin (50 μg/ml) or chloroamphenicol (50 μg/ml), as required. The murein-lacking planctomycetes cells (Rhodopirellula baltica) were grown on M607 agar plates; Sulfolobus acidocaldarius cells were grown on Brocks medium, supplemented with 0.2% (w/v) tryptone.

Protein expression, isolation, and purification

Fresh cultures of E. coli Rosetta gami 2 bearing the constructs of C-terminally GFP- and H10-tagged LysM_{AcmA}

![Fig. 1](image-url) Molecular architecture and binding activity of the LysM_{AcmA}, PMB_{MTH719}, and the M–P–GFP–H10 fusion proteins. The three LysM motifs of AcmA and three PMB motifs of MTH719 are abbreviated as L1, L2, and L3 and P1, P2, and P3, respectively. GFP and H10 indicate the green fluorescent protein and the His_{10} tag located at the C-terminus, respectively. The binding activity of the different domain constructs to the cell wall materials is indicated with + (bound) or – (not bound).
Table 1  Primers for the amplification of the M–P–GFP–H₁₀ fusion protein construct

| Oligo | Primer sequence; 5’>3’ |
|-------|------------------------|
| LF    | ATGGGTGGTGGATTTGCTGGAAATCTAATTCTGGTGCC |
| LR    | CTAATTTAACAATCGCTTTATTCGTAAGATACCTGCAAC |
| PF    | TCAGTACCTCAGAATAAAAAGCGAGTGTTAAGATTAG |
| PR    | TGGAAGATATAAATTCTGAGCTAGCAAGTCTTTGCC |

LF and PR are forward and reverse primers of the LysM<sub>Acma</sub> and PMB<sub>MTH719</sub> domains, respectively. Bases denoted in italics were added before the gene sequence to each forward and reverse primer as described previously (Geertsma and Poolman 2007) and the bases in bold are the starting and reverse complementary bases of the respective genes. LR is the reverse primer for the LysM<sub>Acma</sub> domain with a short domain, PMB<sub>MTH719</sub> domain, and the M domain, PMB<sub>MTH719</sub> domain, and the M–P fusion protein were started from the mother cultures. Cells were grown at 37°C in selective TY medium until an OD at 600 nm of 0.6–0.8 was reached and were induced with 0.2% (w/v) arabinose (Merck KGaA, Darmstadt, Germany) followed by incubation for another 2 h at 37°C on a rotating shaker at 250 rpm (Innova 4000, New Brunswick Scientific, Edison, NJ, USA). The overexpressed proteins were isolated and purified separately according to the manufacturer’s protocol (Ni-NTA superflow, Qiagen GmbH, Hilden, Germany). GFP (control) was purified by hydrophobic interaction chromatography (HIC) according to the Bio-Rad experimental protocol.

In-gel fluorescence and western hybridization

The LysM<sub>Acma</sub>, PMB<sub>MTH719</sub>, and M–P fusion protein bearing the C-terminal GFP and H₁₀ tag were subjected to SDS 12.5%-PAGE and in-gel GFP fluorescence was visualized using a Gel Documentation System (Bio-Rad Laboratories Inc, Hercules, CA, USA). The same gel was blotted-transferred onto polyvinylidene fluoride transfer membrane (GE Healthcare UK Limited, Little Chalfont, Buckinghamshire, UK). Anti-His-tag polyclonal rabbit IgG (Santa Cruz Biotechnology, Santa Cruz, CA, USA) and anti-rabbit IgG peroxidase (GE Healthcare UK Limited) were used as primary and secondary antibodies, respectively. Washing and incubation steps were performed using a SNAP-id system (Millipore). Enhanced chemiluminescence was used to detect the signals, according to the manufacturer’s protocol (GE Healthcare UK Limited).

Binding studies

Preparation of <i>L. lactis</i> Gram-positive Enhancer Matrix (GEM) particles containing purified murein was previously explained (Audouy et al. 2006, 2007; Bosma et al. 2006). Ni-NTA-purified LysM<sub>Acma</sub>, PMB<sub>MTH719</sub>, and M–P fusion protein and HIC-purified control GFP protein were incubated separately with the <i>L. lactis</i> GEM particles, pseudomurein of <i>Methanobacterium</i> sp. cells (Sigma, Zwijndrecht, the Netherlands) chitin from shrimp shells (Sigma) and fungal cells (Psilocybe cubensis taxid: 181762) in 50 mM NaHCO₃, pH 9.2 and 10.0 buffers. Additionally, the M–P–GFP–H₁₀ fusion protein, purified GFP (control), and the substrates were incubated in 50 mM NaH₂PO₄, pH 6.0 and 8.0 buffers. As negative controls, cells of <i>R. baltica</i>, a murein-lacking bacterium, and <i>S. acidocaldarius</i>, an archaean deficient in pseudomurein, were used in all the binding experiments. The incubation was performed at room temperature for 30 min on a rotatory shaker. The mixture was spun down at 14,000 rpm on a desktop centrifuge. The pellet was washed three times with the same buffer supplemented with 150 mM NaCl followed by centrifugation after every wash. After the incubation and washing steps, the material was resuspended in the same buffers. The material was placed on a microscope slide and viewed under a phase-contrast microscope (Zeiss Axioshot, Thornwood, USA) fitted with a digital camera and a green filter to view fluorescence. All photographs were taken at 1,250-fold magnification.

Results

Nucleotide sequences encoding the LysM domain from the autolysin Acma of <i>L. lactis</i> subsp. cremoris MG1363 (plasmid-free derivative of strain NCDO712 of the National Collection of Dairy Organisms) (Buist et al. 1995) and the PMB domain from the S-layer protein MTH719 of <i>M. thermotogatus</i> str. str. (strain DSMZ1053 of the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH) (Visweswaran et al. 2011a) were amplified separately by PCR using the primer pair LF/LR and PF/PR, respectively (see Table 1). The individual LysM and PMB amplification products were used as templates for the generation of the fusion gene by overlap PCR with the primer pair LF/PR. The three amplified DNA fragments were each cloned into pBADcLIC-GFP (Geertsma and Poolman 2007) using <i>E. coli</i> Rosetta gami 2. In this way, each protein domain was fused to a GFP and H₁₀ tag. The molecular architecture of the individual LysM, PMB, and LysM–PMB fusion protein (from now onwards denoted as M–P) domains is shown in Fig. 1. The expression of the Acma LysM (LysM<sub>Acma</sub>), MTH719 PMB (PMB<sub>MTH719</sub>), and M–P fusion domains and their proper fusion to the C-terminal GFP–H₁₀ tag, was confirmed by in-gel fluorescence detection and western hybridization using anti-His-tag antibodies (Fig. 2). The extra band in lane 5 (Fig. 2a) is most likely
caused by protein degradation, liberating GFP, as the protein in this band also fluoresces (Fig. 2b). A non-fluorescent breakdown product can be seen in lane 3 (Fig. 2a, b).

**Homologies of LysM and PMB motifs**

LysM motifs contain 40–65 amino acids, whereas PMB motifs are 30–35 residues in length (Fig. 3) (Buist et al. 2008; Visweswaran et al. 2010, 2011a, b). No significant homology exists between the LysM and the PMB motifs. The three C-terminal LysM motifs of AcmA share homology with one another (Buist et al. 1995). A multiple sequence alignment using the ClustalW2 program (EMBL-EBI, Chenna et al. 2003) reveals 16 amino acid residues that are identical in all three motifs (Fig. 3a). Six of these amino acid residues tend to be highly conserved in most of the known LysM sequences including those of the LysMs of which 3D structures are known (MltD, PDB: 1E0G, and YkuD, PDB: 1Y7M) (Fig. 3a).

In contrast, the three PMB motifs in PMB_{MTH719} are poorly conserved. Only two amino acid residues (R26, P28) are commonly found in most other PMB motif sequences (Fig. 3b and Pfam, http://pfam.sanger.ac.uk/family/pmb#tabview=tab4). However, motifs 1 and 3 and motifs 2 and 3 possess 11 and nine identical amino acid residues between them, respectively, while motifs 1 and 2 have only four residues in common (Fig. 3b). The significance of the three PMB motifs of the MTH719 protein is that they contribute an overall positive charge to MTH719. The pI of PMB_{MTH719} is 10.6 while the pI of the MTH719 protein without the PMB domain is only 4.5 (ExPASy). Because of this high pI value of the PMB domain, the MTH719 protein as a whole possesses a relatively high pI (8.7) (ExPASy). In contrast, the pI of the AcmA protein is not significantly affected by the presence of the LysM domain.

**Fig. 2** Expression of M–P, LysM_{AcmA}, and PMB_{MTH719} GFP–H10 fusion protein constructs. a Western blot using anti-His antibodies and b in-gel fluorescence. Lane 1, molecular mass marker; lanes 2, 4, and 6 and 3, 5, and 7, uninduced and 0.2% L-arabinose-induced cell free extracts of M–P–GFP–H10 fusion protein; LysM_{AcmA}–GFP–H10 and PMB_{MTH719}–GFP–H10, respectively. Arrows (↓) indicate specific protein band.

**Fig. 3** Multiple sequence alignments of the AcmA LysM and MTH719 PMB motifs using ClustalW2 (EMBL-EBI, Chenna et al. 2003). a The three C-terminal LysM motifs of the AcmA protein. b The three C-terminal PMB motifs of the MTH719 protein. Bold residues indicate highly conserved amino acids in consensus sequences (Bateman and Bycroft 2000) and asterisks (*) indicate identical residues in the three motifs. Amino acid residues common between MTH719 PMB repeats 1 and 3, 2 and 3, and 1 and 2 are indicated in blue, red, and green, respectively.

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The LysM and PMB domains specifically bind to their respective substrates

To demonstrate the functionality of the individual domains in the LysMAcmA–GFP–H₁₀ and PMBMTH719–GFP–H₁₀ proteins, both were purified using Ni-NTA column chromatography. LysMAcmA–GFP–H₁₀ was incubated with L. lactis GEM particles (Audouy et al. 2006, 2007; Bosma et al. 2006), shrimp shell chitin flakes, pseudomurein of Methanobacterium sp. cells, and cells of P. cubensis at pH 10.0. LysMAcmA–GFP–H₁₀ specifically bound to the murein-containing L. lactis GEM particles (Figs. 1 and 4a). Interestingly, it also bound to shrimp shell chitin flakes and to the chitin-containing fungal (P. cubensis) cell walls (Figs. 1 and 4b, c). No binding was observed to the pseudomurein of Methanobacterium sp. cells (Figs. 1 and 4a). HIC-purified GFP was used as a negative control in the binding experiments and did not give any fluorescence signal with all tested substrates (data not shown). Additionally, cells of R. baltica and S. acidocaldarius were used as murein and pseudomurein-deficient controls, respectively. No protein binding was observed (data not shown). This indicates that the LysMAcmA domain is responsible for binding to the three substrates (L. lactis GEM particles, shrimp shell chitin flakes, and cells of P. cubensis) and shows, for the first time, that a LysM domain from a Gram-positive bacterium can also bind chitin, as can their plant counterparts (Ohnuma et al. 2008; Petutschnig et al. 2010).

When purified PMBMTH719–GFP–H₁₀ was incubated with the same substrates at pH 10.0, binding was only observed with pseudomurein-containing Methanobacterium sp. cells (Figs. 1 and 4d); no GFP fluorescence was seen with the other substrates (Fig. 4d and data not shown) indicating the specificity of the PMBMTH719 domain towards pseudomurein-containing Methanobacterium sp. cells.

The M–P fusion protein is capable of binding to murein, pseudomurein, and chitin

Ni-NTA-purified M–P–GFP–H₁₀ fusion protein was incubated at pH 10.0 with the same substrates tested earlier for the individual LysMAcmA and PMBMTH719 domains. The fusion protein bound to both whole cells of L. lactis, to GEM particles, and to pseudomurein of Methanobacterium sp. cells at room temperature (Figs. 1 and 5a). The M–P–GFP–H₁₀ fusion protein also bound to shrimp shell chitin flakes and to P. cubensis fungal cell walls (Figs. 1 and 5b, c). In contrast, the M–P–GFP–H₁₀ fusion protein did not bind to the negative controls (cells of R. baltica and S. acidocaldarius) (Fig. 5a). As HIC-purified GFP protein (control) did not give a fluorescence signal with the substrates (data not shown), the binding of the M–P–GFP–H₁₀ fusion protein was specific.

Binding is pH dependent

Different pH conditions were tested to study the effect of pH on binding of the M–P–GFP–H₁₀ fusion protein. To rule out the influence of pH on the GFP signal, HIC-purified GFP was tested at different pH prior to the binding experiments. Under all pH conditions tested, the GFP signal remained the same. The Ni-NTA-purified M–P–GFP–H₁₀ fusion protein (pH 10.3) was mixed with L. lactis GEM particles, pseudomurein of Methanobacterium sp. cells, chitin flakes, and cells of the fungus P. cubensis in buffers of pH 6.0, 8.0, 9.6, or 10.0. No GFP signal was seen at pH 6.0 and 8.0, indicating that no binding occurred (data not shown). A very low GFP signal was detected at pH 9.6 (data not shown). At pH 10.0, a strong GFP signal was observed in all tested samples (Fig. 5), indicating that the M–P–GFP–H₁₀ protein had bound to the four substrates.

We conclude that M–P fusion protein is capable of binding to multiple cell wall substrates due to the combined properties of the two major murein and pseudomurein cell wall-binding domains, i.e., LysMAcmA and PMBMTH719. We also showed that the LysMAcmA domain from bacterial origin can bind chitin.

Discussion

AcmA, the widely characterized autolysin of L. lactis, possesses an N-terminal catalytic glucosaminidase domain followed by a C-terminal LysM domain consisting of three motifs for binding to murein. Using LysMAcmA–GFP fusion studies, we have determined that the LysMAcmA domain binds to substrates other than murein. LysMAcmA did not only bind to murein-containing L. lactis GEM particles but also to chitin and to fungal cell walls. To date, only eukaryotic LysM domains were shown to bind to chitin (Ohnuma et al. 2008; Petutschnig et al. 2010). Here, we show for the first time the prokaryotic LysM domain’s ability to bind eukaryotic cell walls, i.e., chitin and to the cell wall of the fungus P. cubensis. The prokaryotic and eukaryotic LysM domains vary quite considerably in their secondary structures and mode of bonding. Unlike eukaryotic LysM domains, the LysM domains of prokaryotes do not possess disulfide bonds, rather the latter domains are supported by extensive secondary structure and hydrogen bonding (Buist et al. 2008). Furthermore, the amino acid sequences of LysM domains of chitinases and bacterial cell wall hydrolases are strikingly different (Steen et al. 2003).

NAG is the only common moiety between murein and chitin, and moreover, the mode of bonding is also the same, i.e., β(1,4). Previously, it was shown that LysM domains from plant origin bind to NAG derivatives (Ohnuma et al. 2008; Petutschnig et al. 2010). We presume that, like its
Fig. 4 Binding specificity of the LysM<sub>Acma</sub>-GFP-H<sub>10</sub> and PMB<sub>MTH719</sub>-GFP-H<sub>10</sub> proteins. The figures (a–c) show fluorescence microscopy views of the LysM<sub>Acma</sub>-GFP-H<sub>10</sub> binding selectively and specifically to <i>L. lactis</i> GEM particles (cocci), chitin flakes, and <i>P. cubensis</i> cells, respectively, at pH 10.0. The figure (d) shows binding specificity of PMB<sub>MTH719</sub>-GFP-H<sub>10</sub> protein to pseudomurein-containing <i>Methanobacterium</i> sp. cells (long rods), at pH 10.0.
eukaryotic counterparts, LysMAcmA might also recognize NAG in both murein and chitin. For a better understanding of LysM functionality and to investigate the nature of the substrate and the mode of binding, a ligand-bound structure of the LysM domain would be essential and an interesting target for future research.

Unlike the LysMAcmA domain, the PMBMTH719 domain specifically binds to pseudomurein-containing *Methanobacterium* sp. cells (Fig. 4d). This result is supported by studies on the PMB domain of PeiW, a pseudomurein endoisopectidase containing four N-terminal PMB motifs, which showed that the protein specifically bound to pseudomurein-containing archaeal cells (Steenbakkers et al. 2006). Although the PMB domains of MTH719 and PeiW are functionally similar in specifically binding to pseudomurein, both domains have less than 20% amino acid sequence homology (data not shown). PMB_MTH719 does not bind to chitin nor to intact murein, but it was recently shown that the PMB domains of both PeiW and MTH719 bind to bacterial spheroplasts (Visweswaran et al. 2011a). As these were made by lysozyme treatment of whole lactococcal cells, it was suggested that NAG is the binding ligand. The only common moiety in all three cell wall polymers (murein, pseudomurein and chitin) is NAG. Whether PMB_MTH719 recognizes NAG or β(1,3)-linked NAG might be resolved by a 3D structure of the domain with its ligand.

**Fig. 5** Multiple substrate-binding ability of M–P–GFP–H10 fusion protein. The figure shows fluorescence microscopy views of M–P–GFP–H10 fusion protein binding at pH 10.0 to a *L. lactis* GEM particles (cocc) and pseudomurein-containing *Methanobacterium* sp. cells (long rods). Cells of *R. baltica* (round phase—dark, arrows) and *S. acidocaldarius* (round phase—gray, arrow heads) show no binding to b chitin flakes and c *P. cubensis* cells.
The M–P–GFP–H₁₀ fusion protein obtained by the union of the LysMₐcmₐ domain and the PMBₘ₆₇₁₉ domain was shown to be functionally active even after the genetic fusion of these two domains from rather distinct origins (bacteria and archaea) and despite the presence of the GFP–H₁₀ tag on the C-terminus of the triple fusion. This indicates that the individual domains (LysMₐcmₐ and PMBₘ₆₇₁₉) in the M–P fusion protein are correctly folded. Due to their combined properties, M–P–GFP–H₁₀ could bind to all the substrates tested (murin, pseudomurin, and chitin). Based on these results and together with previously obtained results from the individual domains, we conclude that binding of M–P–GFP–H₁₀ to GEM particles, chitin flakes, and to *P. cubensis* fungal cell walls was mediated by the LysMₐcmₐ domain, and binding to pseudomurin-containing *Methanobacterium* sp. cells is effectuated by the PMBₘ₆₇₁₉ domain. The binding of M–P–GFP–H₁₀ is greatly influenced by the pH of the binding environment and depends on the individual pH values of the LysMₐcmₐ and PMBₘ₆₇₁₉ domains. Previous work from our group on the individual LysMₐcmₐ (pH 10.0) and PMBₘ₆₇₁₉ (pH 10.6) domains has shown that both bind to their substrates at a pH close to their respective pH values (Buist et al. 1995; Steen et al. 2003, 2005; Visweswaran et al. 2011a). The M–P–GFP–H₁₀ fusion protein (pH 10.3) bound to substrates at a pH close to its pI, i.e., pH 10.0, indicating pH dependency. The LysM domain of AcmD, an N-acetylglucosaminidase from *L. lactis* and a homologue of AcmA, binds to *L. lactis* cells at pH 4.0, which is close to the pI value of LysMₐcmₐ (pI 4.3) (Visweswaran et al., unpublished results). An M–P–GFP–H₁₀ fusion protein (pI 9.6) made of the LysMₐcmₐ (pI 4.3) and PMBₘ₆₇₁₉ (pI 10.6) bound to *L. lactis* GEM particles at pH 4.0 and to pseudomurin of *Methanobacterium* sp. cells at pH 10.0, but not to both substrates at an intermediate pH (pH 7.0), due to the quite different pH values of the two domains (data not shown).

In vivo, the prokaryotic LysM domains present in cell wall hydrolases aid in non-covalent binding of the enzyme to their substrates allowing some of the physiological functions like cell separation and cellular autolysis to work efficiently (Steen et al. 2005; Buist et al. 2008). Similarly, the PMB domains of archaeal hydrolases (PeiW and PeiP) allow binding of the enzymes to pseudomurin, facilitating hydrolysis by the catalytic domains (Luo et al. 2002; Steenbakkers et al. 2006; Visweswaran et al. 2010). In recent years, the LysM domains have been widely applied for various biotechnological purposes such as in the preparation of oral influenza vaccine and for surface display of heterologous proteins on bacterial cell surfaces (Bosma et al. 2006; Okano et al. 2008; Shao et al. 2009; Hu et al. 2010; Sahuja et al. 2010; Xu et al. 2011). Similarly, the PMB domain could be employed for the display of target proteins on pseudomurin-containing archaeal cell surfaces; it could also be used as a marker protein to identify pseudomurin-containing methanogens from their counterparts. The fusion of the PMB domain to the LysM domain adds additional functionality to the latter, widening its application potential.

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