Substrate Specificity of Low-Molecular Mass Bacterial DD-Peptidases

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ABSTRACT: The bacterial DD-peptidases or penicillin-binding proteins (PBPs) catalyze the formation and regulation of cross-links in peptidoglycan biosynthesis. They are classified into two groups, the high-molecular mass (HMM) and low-molecular mass (LMM) enzymes. The latter group, which is subdivided into classes A–C (LMM, -B, and -C, respectively), is believed to catalyze DD-carboxypeptidase and endopeptidase reactions in vivo. To date, the specificity of their reactions with particular elements of peptidoglycan structure has not, in general, been defined. This paper describes the steady-state kinetics of hydrolysis of a series of specific peptidoglycan-mimetic peptides, representing various elements of stem peptide structure, catalyzed by a range of LMM PBPs (the LMM enzymes, Escherichia coli PBP5, Neisseria gonorrhoeae PBP4, and Streptococcus pneumoniae PBP3, and the LMMC enzymes, the Actinomadura R39 DD-peptidase, Bacillus subtilis PBP4a, and N. gonorrhoeae PBP3). The R39 enzyme (LMMC), like the previously studied Streptomyces R61 DD-peptidase (LMMB), specifically and rapidly hydrolyzes stem peptide fragments with a free N-terminus. In accord with this result, the crystal structures of the R61 and R39 enzymes display a binding site specific to the stem peptide N-terminus. These are water-soluble enzymes, however, with no known specific function in vivo. On the other hand, soluble versions of the remaining enzymes of those noted above, all of which are likely to be membrane-bound and/or associated in vivo and have been assigned particular roles in cell wall biosynthesis and maintenance, show little or no specificity for peptides containing elements of peptidoglycan structure. Peptidoglycan-mimetic boronate transition-state analogues do inhibit these enzymes but display notable specificity only for the LMMC enzymes, where, unlike peptide substrates, they may be able to effectively induce a specific active site structure. The manner in which LMM (and HMM) DD-peptidases achieve substrate specificity, both in vitro and in vivo, remains unknown.

The bacterial DD-peptidases catalyze the final steps of peptidoglycan (cell wall) biosynthesis and are efficiently inhibited by b-lactam antibiotics. Resistance to these antibiotics continues to emerge, particularly from evolution of new b-lactamases, enzymes that catalyze hydrolytic destruction of b-lactams, but also by the evolution and dispersal of b-lactam-resistant DD-peptidases. Important examples of pathogenic bacteria in which b-lactam-resistant DD-peptidases play a significant role in resistance are Staphylococcus aureus (MRSA), Streptococcus pneumoniae, Enterococcus faecium, and Neisseria gonorrhoeae. Many attempts have been made to develop new b-lactams effective against mutant DD-peptidases, but, to date, these have not been completely successful. The ideal of a broad-spectrum b-lactam not susceptible to loss of effectiveness through DD-peptidase mutation has been difficult to achieve. To a considerable extent, this is likely due to the largely trial and error approach to new b-lactams. The alternative approach of targeting the reaction center and the common and essential structural features of the DD-peptidase active site has not been pursued as vigorously. Classically, one would expect substrate analogue inhibitors, mechanism-based and transition-state analogues, to be effective. Few such examples are known, although the reasons for this are not clear. The first issue to be taken into account in the design of such inhibitors is the structure of the substrate itself and the affinity of the enzyme for it. They can first be divided into two groups, high-molecular mass (HMM) and low-molecular mass (LMM) enzymes, where the dividing line is approximately 50 kDa. Each of these groups can be subdivided, the HMM group into HMM-A and HMM-B classes and the LMM group into LMM, -B and -C classes. The HMM DD-peptidases are bifunctional enzymes that catalyze both the transglycosylation and transpeptidation reactions of incorporation of stem peptide monomers into peptidoglycan, whereas members of the HMMB group are thought to catalyze only the transpeptidation reactions.

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reaction. Both of these classes are essential to bacterial survival and are therefore the primary targets of β-lactam antibiotics. The LMM groups are also inhibited by β-lactams, but because they are not essential to bacterial survival, at least in the short term, they are not considered viable antibiotic targets. The LMM enzymes are thought to be DD-carboxypeptidases and/or endopeptidases, which regulate the extent of peptide cross-linking in peptidoglycan. Alternatively, the terminal D-alanine may be hydrolytically cleaved in a DD-carboxypeptidase reaction. The sites of these reactions are also shown in 1. Transpeptidation and carboxypeptidation reactions also may occur in previously cross-linked peptidoglycan. Structure 2 is a dimer in which two polysaccharide strands are cross-linked and the sites of the transpeptidase and carboxypeptidase reactions are indicated. Finally, dimers and more highly cross-linked peptidoglycan are also subject to endopeptidase hydrolysis, as shown in 2. Together, the carboxypeptidase and endopeptidase reactions regulate the degree of cross-linking of the peptidoglycan. All of these reactions are thought to proceed through a covalent acyl–enzyme intermediate derived from reaction of a nucleophlic serine residue at the DD-peptidase active site (Scheme 1).

Scheme 1

One would expect, therefore, that the substrate(s) of DD-peptidases would include elements of 1 and 2 and that these enzymes would show specificity toward peptides containing these elements. To a large extent, however, such specificity has not yet been systematically investigated, even in vitro. The situation in vivo is even more uncertain for most of these enzymes.

A clear exception to this statement is provided by one particular DD-peptidase, the LMMC enzyme from Streptomyces R61. This enzyme was studied for many years by the Liege group of Ghuysen and Frère. They showed that the R61 DD-peptidase catalyzed the hydrolysis of certain small peptides such as N,N-diacetyl-l-lysyl-d-alanyl-D-alanine, 3, and similar (thio)depsipeptides. More recently, it has been shown that peptide 4 is a very specific substrate for this enzyme \( k_{cat}/K_m = 8.7 \times 10^{6} \text{ s}^{-1} \text{ M}^{-1} \), much more reactive than generic peptide 3 \( k_{cat}/K_m = 3.5 \times 10^{3} \text{ s}^{-1} \text{ M}^{-1} \). The specificity of 4 as a substrate is certainly understandable in view of the Streptomyces R61 stem peptide structure that includes the N-glycyl-l-α-aminopimelyl moiety as the N-terminus.

X-ray crystal structures of complexes of 4 with the enzyme show what seems to be a very tight and specific binding site for the N-glycyl-l-α-aminopimelyl side chain of 4. The R61 DD-peptidase appears to have only weak endopeptidase activity in vitro, and although it does catalyze transpeptidation reactions between 4 and small acceptors, extended acceptors, such as one would expect in a natural substrate, are ineffective. In summary, these data support the proposition that the R61 DD-peptidase is a DD-carboxypeptidase, at least in vitro, whose preferred substrate is the stem peptide monomer, 1. Data of this detail pertaining to substrate specificity are not available for virtually any other DD-peptidase, although there is considerable information concerning the Actinomadura R39 DD-peptidase, including a recent systematic study from this laboratory. In this case, a peptidoglycan-mimetic boronate was also shown to be a very potent inhibitor, most likely a transition-state analogue.

In this paper, we explore the specificity of several LMA and LMMC DD-peptidases, employing a series of peptidoglycan-mimetic peptide substrates and boronate inhibitors. The aim was to determine whether peptidoglycan mimetics are generally useful leads to DD-peptidase inhibitors and thus, perhaps, to new antibiotics. In particular, the issue of whether more extended peptides, related to dimer and oligomer peptidoglycan structures, might lead to stronger interactions with these enzymes was evaluated.

### MATERIALS AND METHODS

The purified enzymes were prepared as described: the *Actinomadura* R39 DD-peptidase, *Escherichia coli* PBP5 (EcPBP5), *N. gonorrhoeae* PBP3 (NgPBP3) and PBP4 (NgPBP4), *Bacillus subtilis* PBP4a (BsPBP4a), and *S. pneumoniae* PBP3 (SpPBP3). The following peptides were purchased commercially and used as received: 10 (NeoMPS), 3 and 13 (Sigma-Aldrich), and 14–19 (New England Peptide). Peptides 7–9 were prepared by standard peptide methodology; the details are given in the Supporting Information. The syntheses of 5, 22, and 23 have been previously described.
and that of 6 will be described elsewhere. Synthesis of boronate 11 has been described previously; and that of 12, 20, and 21 were conducted similarly and are described in the Supporting Information.

A general 1H NMR experiment was first used to determine the products of reactions of the R39 DD-peptidase and BsPBP4a toward peptides 7−9. 1H NMR spectra of solutions of these peptides (3−5 mM) in deuterated phosphate buffer (20 mM) were recorded before and after the addition of enzyme (0.1−0.3 μM). Further spectra were recorded at various time intervals until the reaction was complete. Similar experiments were performed with SpPBP3 and peptides 14−18. The lack of reaction of 16 and 17 with BsPBP3 was checked by high-performance liquid chromatography (HPLC) where the absence of α-N-acetyllysinine was confirmed.

Kinetic Methods. Steady-state kinetics experiments were conducted under conditions previously established to generate high, stable activity: R39 DD-peptidase and BsPBP4a, 20 mM phosphate buffer (pH 7.5) at 25 °C; EcPBP5, NgPBP4, and NgPBP4, 0.1 M sodium pyrophosphate and 10% glycerol (pH 8.5) at 37 °C; SpPBP3, 20 mM phosphate buffer (pH 7.0) at 37 °C. Measurements of enzymatic activity with peptides were determined spectrophotometrically, where the loss of a product or substrate at 375 nm. Initial rates were fitted to the Henri−Michaelis−Menten equation by a nonlinear least-squares procedure to yield values of $k_{cat}$ and $K_m$. The range of peptide substrate concentrations employed was 0−1 or 2 mM, and enzyme concentrations were, typically, 0.1−0.4 μM. In cases where substrate concentrations approaching $K_m$ were not achieved, values of $k_{cat}/K_m$ were obtained by nonlinear least-squares exponential fits to the total progress curve. In the case of the Actinomadura R39 DD-peptidase enzyme, the $K_m$ of peptide substrate 9 was determined separately as the $K_m$ value from a competition experiment in which thiolactate 24−21 was the substrate. Equilibrium constants for inhibition of the enzymes by 11, 12, 20 and 21 were obtained from steady-state competition experiments in which N-phenylacetylglutaryl-D-thiolactate35 was employed as a chromophoric (245 nm, $\Delta e = 2500$ cm$^{-1}$ M$^{-1}$) substrate (0.5 mM). The reaction conditions were as follows: R39, boronate 11 and 12 concentrations of 0.1−1.0 μM and 0.1−1.0 mM, respectively, enzyme concentration of 94.5 nM, and substrate $K_m$ of 38.4 μM (S. A. Adediran and R. F. Pratt, unpublished observations); PBP4a, boronate 11 and 12 concentrations of 0.1−1.0 μM and 0.1−1.0 mM, respectively, enzyme concentration of 167 nM, and substrate $K_m$ of >1 mM; NgPBP3, boronate 11 and 12 concentrations of 10−250 μM and 0.1−1.0 mM, respectively, enzyme concentration of 10.4 nM, and substrate $K_m$ of >1 mM; NgPBP4, boronate 11 and 12 concentrations of 1.0−10 μM and 0.1−1.0 mM, respectively, enzyme concentration of 150 nM, and substrate $K_m$ of 1.6 μM; EcPBP5, boronate 11 and 12 concentrations of 0.1−1.0 mM, enzyme concentration of 137 μM, and substrate $K_m$ of 220 μM; SpPBP3, boronate 20 and 21 concentrations of 0.1−10 μM and 1.0−50 μM, respectively, enzyme concentration of 104 nM, and substrate $K_m$ of 350 μM. Competitive inhibition was assumed in fitting the data to a steady-state scheme. In a few cases, BsPBP4a with 11 and SpPBP3 with 20 and 21, boronate binding was slow (minutes) on a manual mixing time scale. Details of this will be published elsewhere. The $K_i$ values listed in Table 3 represent the steady-state values.

## RESULTS AND DISCUSSION

The LMM DD-peptidases examined in this work can be divided into two groups, one derived from *E. coli*, *N. gonorrhoeae*, *Actinomadura* R39, and *B. subtilis*, where the stem peptide has structure 22 with meso-diaminopimelic acid at position 3, and the other from *S. pneumoniae*, where the stem peptide has structure 23 with L-lysine at position 3 (Chart 1). Specifically, the LMMA enzymes studied were *E. coli* PBP5, *N. gonorrhoeae* PBP4, and *S. pneumoniae* PBP3, and the LMMC enzymes were the *Actinomadura* R39 DD-peptidase, *N. gonorrhoeae* PBP3, and *B. subtilis* PBP4a.

The panel of substrates and inhibitors 5−21 (Chart 2) was assembled for the examination of two major issues. The first of these is focused on the specificity of typical LMM DD-peptidases for extended acyl groups (acyl donor specificity), in particular to examine whether the in vivo substrate is likely to have a free amine at the N-terminus of the acyl group as in 1 or whether its amine is acylated as in dimer or oligomer formation. The second point to be addressed is whether, for any of these enzymes, there is specificity for an extended (beyond D-Ala) leaving group, e.g., for 9 versus 5 or 16 versus 13. A narrow specificity for a D-Ala leaving group would presumably signal purely carboxypeptidase activity, whereas preference for an extended leaving group might rather indicate transpeptidase or endopeptidase activity; furthermore, one might expect that the preference for the extended leaving group by an endopeptidase would be more pronounced than for a transpeptidase where there must also be affinity for D-alanine.
The boronates were prepared to investigate whether the affinity of these enzymes for elements of peptidoglycan structure could be detected through the stronger binding of transition-state analogues compared to that of substrates. For example, these inhibitors might be more effective than substrates at inducing reactive conformations from inactive forms that may exist in solution.

First, the response of the enzyme to N-terminal acylation, such as would occur on expansion of monomer 1 to dimer 2 and to oligomers, was examined. The reactivity of the enzymes from bacteria with the D-α-aminopimelyl stem peptide N-terminus against peptides 7, 8, and 10 (vs 5) and boronate 12 (vs 11) was assessed. Second, to evaluate the preference of the various enzymes for extension of the leaving group beyond D-alanine, as would occur in transpeptidase (actually, the preference for the nucleophile in the forward reaction, as depicted in 1 and 2) and endopeptidase reactions, peptide 9 (vs 5) was examined. The enzyme of *S. pneumoniae*, which has an ε-lysyl N-terminus of the stem peptide, was evaluated analogously by reference to peptides 3, 13−15, 18, and 19 (vs 13) and boronate 21 (vs 20). Preference for transpeptidase and endopeptidase (vs carboxypeptidase) activity was assessed by considering the results from peptides 16 and 17 (vs 13).

The products of hydrolysis of the peptides, as catalyzed by the enzymes named above, were identified by 3H NMR experiments. Both the R39 DD-peptidase and BsPBP4a...
Hydrolysis of 5 by these enzymes is known to yield D-alanine from the C-terminus. \( ^{34} \) Similarly, SpPBP3 was shown to catalyze a dd-carboxypeptidase reaction of 14, 15, 18, and 19 with release of D-Ala, under the same conditions, \( ^{1} \)H NMR and HPLC experiments showed no reaction of 16 and 17. Previous experiments have established that the R39 DD-peptidase, EcPBP5, NgPBP3, and NgPBP4 catalyze DD-carboxypeptidase reactions of 5–10, 3, and 13. \( ^{3,16,30,31} \)

Steady-state kinetic studies of the hydrolysis of the peptides described above by the enzyme panel were conducted as described in Materials and Methods. The steady-state parameters obtained are listed in Tables 1 and 2. The steady-state boronate inhibition constants were also determined as described and are listed in Table 3.

### LMMC dd-Peptidases

Comming with the LMMC enzymes, we see that the R39 enzyme, as suggested by previously available data, \( ^{26,27,34,35,38} \) has a clear preference for a nonacylated peptide N-terminus, as seen in the \( k_{cat}/K_m \) values of 5 and 9 versus 7, 8, and 10 and the much more potent inhibition of this enzyme by boronate 11 than by 12. The results, however, appear slightly ambiguous with respect to the transpeptidase/endopeptidase question. Peptide 9 is comparable to 5 in its reactivity with the enzyme, but not more so. On the other hand, it has been shown that extended acceptors such as N-\( \epsilon \)-acetyl-D-lysine are considerably better nucleophiles than D-alanine against the acyl–enzyme intermediate derived from 24, a thia analogue of 5. \( ^{26} \) The analogous and arguably more peptidoglycan-mimetic peptide 25 was an even better nucleophile. These results were interpreted to indicate that the dominant role in activity of the R39 dd-peptidase is as an endopeptidase. \( ^{26} \) Because the same transition state should be traversed in acylation of the enzyme by a peptide and aminolysis of the cognate acyl–enzyme intermediate by an amino acid, these results with 5 and 9 appear to some extent inconsistent with the earlier ones. The difference may come about because of the greater absolute reactivity of 5 and 9 (\( k_{cat}/K_m = 5.7 \times 10^6 \text{ and } 2.0 \times 10^6 \text{ s}^{-1} \text{ M}^{-1} \)), respectively) compared to that of 25 (\( k_{cat}/K_m = 7.3 \times 10^4 \text{ s}^{-1} \text{ M}^{-1} \)), the difference probably deriving from the fact that 5 is more hydrophobic than 25. \( ^{30} \) It is possible that a physical step has taken over as the rate-determining step for the acylation by 5 (and 9), whereas for 25, the chemical step of acyl transfer is rate-determining. The simplest interpretation of the evidence mentioned above is that the optimal substrate of the R39 DD-peptidase would be a peptidoglycan fragment with a free N-terminus but with an extended C-terminus, i.e., a dimer such as 2 or an oligomer. The preferred reaction catalyzed by the enzyme would be an endopeptidase reaction at the D-alanyl peptide bond closest to the free N-terminus of a dimer, such as that represented by 2, or of an oligomer.

![Image](http://dx.doi.org/10.1021/bi201326a)
N-terminus of a peptidoglycan fragment.\textsuperscript{27,39} The crystal structures show a well-defined binding site for a D-\textalpha-aminopimelyl terminus. The important elements of the binding site are Tyr147 and Met414, which abut the hydrophobic tetramethylene chain, Asp142, which interacts strongly with the terminal ammonium ion, and Arg351 and Ser415, which interact with the ligand carboxylate. Finally, Trp139 appears to form a π-cation complex with the terminal ammonium ion. Thus, the kinetics and structural data appear to be in accord regarding the interaction of a peptidoglycan N-terminus with the R39 DD-peptidase. The situation strongly resembles that found for the Streptomyces R61 DD-peptidase, described in the introductory section.\textsuperscript{18–21}

An example in which the kinetics and structural data are apparently less in accord is the LMMC enzyme BsPBP4a. This enzyme is generally less reactive ($k_{\text{cat}}/K_m$) with the relevant peptides than is the R39 DD-peptidase (Table 1). Its $k_{\text{cat}}/K_m$ value for 5 is more than 100-fold lower than that of the R39 enzyme, which has a strikingly low $K_m$ for this substrate; the $K_m$ of 5 in the reaction with BsPBP4a is greater than 1 mM. Most directly, this would suggest much weaker interactions with and less specificity for the N-terminus than in the R39 case. Despite this, as for the R39 DD-peptidase, $k_{\text{cat}}/K_m$ values for 7, 8, and 10 are considerably smaller than that of 5, suggesting that the enzyme has little interest in an extended N-terminus. Similar to that of the R39 enzyme, the activity of BsPBP4a against 9 is slightly lower than against 5 but considerably higher than against 7 and 8, suggesting a reasonable affinity for an extended C-terminus and thus possible transpeptidase/endopeptidase activity.

More emphatically supporting the affinity of BsPBP4a for a free N-terminus is the dramatically stronger inhibition demonstrated by boronate 11 than by 12 (Table 3). The apparent difference between the affinity of the enzyme for 11 and for the analogous substrate 5 is rather astonishing, even given that 11 is probably a transition-state analogue; for the R39 enzyme, the difference is much smaller because of the much stronger apparent binding ($K_m$) of 5.

It thus may be that BsPBP4a is, like the R39 DD-peptidase, an endopeptidase,\textsuperscript{41} preferring a free N-terminus adjacent to the cleavage site. In solution, however, it may exist in a largely inactive conformation (hence the high $K_m$ values for 5 and 9) from which an active conformation can be induced only poorly by a specific substrate such as 5 or 9 or, more effectively, by a boronate transition-state analogue such as 11 (e.g., Scheme 2). It is interesting to note, in this regard, that the crystal structure of a complex of BsPBP4a with 5 shows the latter covalently attached to the active site serine as a D-\textalpha-aminopimelyl-D-alanyl acyl–enzyme intermediate.\textsuperscript{32} This certainly suggests the trapping of the substrate by an inactive conformation. An acyl–enzyme intermediate has also been observed in a crystal structure of S. pneumoniae PBP1b, although in this case, the substrate was nonspecific.\textsuperscript{43} It might also be noted that the association of boronate 11 with BsPBP4a is an at least two-phase process with a slower phase lasting for minutes. The $K_i$ quoted in Table 3 is the steady-state value achieved after completion of the transient phase. Details of this reaction will be published elsewhere, but it is possible that this observation also derives from enzyme isomerization. The binding of 11 to the R39 DD-peptidase shows only a single phase on the same manual mixing time scale.

\begin{table}[h]
\centering
\caption{Inhibition of DD-Peptidases by Boronates}
\begin{tabular}{|c|c|c|c|c|}
\hline
\textbf{enzyme} & \textbf{11} & \textbf{12} & \textbf{20} & \textbf{21} \\
\hline
R39 (LMMC) & 0.032 ± 0.006 & 70 ± 15 & 0.16 ± 0.04 & " \\
BsPBP4a (LMMC) & 0.0090 ± 0.0002 & 29.7 ± 0.5 & " & " \\
NgPBP3 (LMMC) & 13 ± 3 & >1000 & " & " \\
EcPBP5 (LMMMA) & 180 ± 50 & 250 ± 30 & " & " \\
NgPBP4 (LMMMA) & 2.2 ± 0.4 & 0.95 ± 0.07 & " & " \\
SpPBP3 (LMMMA) & 8.8 ± 1.7 & " & 0.22 ± 0.02 & 0.91 ± 0.04 \\
\hline
\end{tabular}
\textsuperscript{"Not determined.}
\end{table}

Another complicating factor with respect to BsPBP4a is that some 99% of peptidoglycan in vegetative cells of B. subtilis have the free carboxyl group of the diaminopimelyl residue amidated; i.e., the stem peptide has the structure of 26.\textsuperscript{34,45}

It was possible, therefore, that the poor affinity of 5 for BsPBP4a reflected the specificity of the binding site for an amide CONH$_2$ group, rather than a carboxylate. We therefore prepared peptide 6, the amidated analogue of 5, and examined its reactivity with the relevant enzymes (Table 1). Stratepeptide 6 was ~1 order of magnitude less reactive ($k_{\text{cat}}/K_m$) with the R39 DD-peptidase, mainly because of a larger $K_m$ value, and also with NgPBP3. This result may reflect the ability of the amide group to adapt to the polar residues of the carboxylate binding site and the absence of any need for these enzymes to strongly discriminate between carboxylate and amide because they, presumably, would be unlikely to interact with the latter in vivo.

\begin{scheme}
\text{Scheme 2}
\end{scheme}
The kinetics of the reaction between BsPBP4a and 6 are also interesting. The \(K_{cat}/K_m\) value for 6 is essentially the same as for 5, with no sign of enhanced (or weakened) binding. It is likely that BsPBP4a does encounter free pimelyl carboxylates as well as amides, and thus, a versatile binding site may be advantageous. Although the role of BsPBP4a in vivo is not known, it is apparently expressed in late vegetative stages\(^46\) when spore formation may be beginning; it has also been reported that the aminopimelyl carboxyl of \(B.\ subtilis\) spore peptidoglycan is not amidaded.\(^47\)

The picture presented by NgPBP3, another LMMC enzyme, in Table 2 is rather different from that of the enzymes described above. This enzyme exhibits no particularly notable specificity toward any of the structural motifs presented by the relevant peptides. Apparent binding (\(K_m\)) to the enzyme is also uniformly weak. It seems that this enzyme may be able to nonspecifically catalyze all three reactions, carboxypeptidase, transpeptidase, and peptidase, with comparable facility and on any substrate. These data, therefore, taken alone, could be interpreted in terms of a versatile, nonspecific enzyme that can undertake any task presented to it, but the presence of such an enzyme in the carefully regulated process of bacterial cell wall metabolism\(^1,2,48,49\) seems unlikely, unless it could be selectively regulated through its location and interaction with other ligands.

Guthel and co-workers\(^30,50\) have shown that this enzyme is much more effective against \(L-\alpha\)-lysyl-\(D\)-alanine peptides where the lysine amine groups are acylated by quite large hydrophobic moieties, e.g., tert-butyloxycarbonyl and benzzyloxycarbonyl. It is not evident how these substituents directly relate to peptidoglycan structure, but a possible interpretation is that these substituents serve to reduce the mobility of the enzyme—substrate complex, leading, by entropic advantage, to more effective catalysis. This rationale would suggest that in vivo, NgPBP3 may function in a space more congested than that of the R39 and BsPBP4a enzymes. Such congestion could be caused by its restriction to the membrane and its proximity to other proteins, peptidoglycan, or both. NgPBP3 and BsPBP4a may be membrane-associated\(^30,46\) even though they do not have a terminal peptide for membrane attachment as do the typical LMMAs and HMM enzymes. In this regard, however, it is worth noting that NgPBP3 derives from a Gram-negative bacterium while the R39 DD-peptidase and BsPBP4a derive from Gram-positive organisms. The site of activity of these enzymes may differ in character, depending on this division. Under this scenario, one would expect that in vivo, the absolute reactivity of the peptide elements of peptidoglycan would be enhanced relative to the results in Table 2, again through entropic advantages. With respect to substrate specificity, it may be that, as for BsPBP4a, the much stronger preference of NgPBP3 for the N-terminal-free boronate 11 than for the N-acylated 12 is the most relevant result, again pointing to a class C specificity for a free N-terminus.

Prior to concluding this discussion of the LMMC enzymes, we find it is useful to note that a quite similar strength does exist between the putative (established for the R39 DD-peptidase and BsPBP4a) peptide N-terminus binding sites (Table 4, where the relevant data for four LMMC enzymes are given). As well as for the three enzymes discussed above, data for EcPBP4 are also given. This is a well-established LMMC PBP where a crystal structure is available,\(^31\) although not with the enzyme bound to a peptidoglycan-mimetic peptide. It seems that the \(D-\alpha\)-aminopimelyl-binding site is well-conserved in all of these enzymes. Asp142 of the R39 DD-peptidase is fully conserved; Arg351, or the alternative, potentially cationic, His352 of BsPBP4a, is conserved, and hydrophobic residues corresponding to Tyr147 and Met414 of the R39 DD-peptidase are retained. Trp139, the putative \(\pi\)-donor of R39 to the cationic ammonium terminus, is not strictly conserved and is replaced with the less electron-rich Phe and Pro in the other enzymes. Finally, Ser415 is replaced with Glu or Asn in two of the enzymes; the backbone carbonyl of this residue, which appears to hydrogen bond to the ligand in both the R39 and BsPBP4a crystal structures,\(^27,42\) is, of course, retained, and polar interactions of the ligand with the terminal amide groups of Glu and Asn are certainly possible. The crystal structure of the BsPBP4a−5 complex\(^25\) shows the N-terminus of the substrate bound to the enzyme in much the same way as in the R39 DD-peptidase, but with His352 directly replacing Arg351. Therefore, despite some counterindications from the peptide hydrolysis kinetics, in accord with the specific binding of boronate 11, it does seem likely that the R39 DD-peptidase, BsPBP4a, NgPBP3, and, quite likely, on the basis of the conservation of the substrate binding motif illustrated by Table 4, all LMMC enzymes have a specific binding site for the free N-terminus of the cognate stem peptide. A FASTA search showed that the 500 proteins of the UniProtKP database most similar to the R39 DD-peptidase retain homologues of Asp142 and Arg/His351. This high degree of active site similarity may be partly obscured in peptide hydrolysis kinetics in vitro by the presence of unreactive enzyme conformations in solution.

**LMMA DD-Peptidases.** The class A enzymes examined responded less positively, in general, to variations in substrate structure than the class C enzymes. EcPBP5 reacted very slowly with the generic N-terminally acylated peptide 3 and with 10 but not detectably with 7−9. Boronates 11 and 12 were also relatively ineffective as inhibitors, and no real sign of specificity for either of them was observed.

This result is not entirely unexpected. As previously noted, solubilized *E. coli* PBP5 rests in an active site-distorted conformation in the crystalline solid state, and, most likely, also in membrane,\(^52\) and in solution.\(^35\) It is pertinent to note at this point that a nonspecific boronate has been shown to induce organization of the reaction center,\(^53\) but this boronate did not have a peptidoglycan-mimetic side chain; thus, no direct evidence about how a specific side chain might bind was obtained. These results indicate that none of the structural motifs in the substrates tested was able to induce a catalytically effective conformation. As discussed above for NgPBP3, however, we presume that this induction does occur in vivo. There is evidence that very specific attachment of EcPBP5 to the cell membrane is required for activity in vivo.\(^54\) Recent experiments suggest that not only membrane attachment but also “ongoing peptidoglycan synthesis” may be needed for strong EcPBP5 substrate binding and catalysis.\(^55\) This is all certainly suggestive of the presence of a specific environment in vivo very different from that in vitro.

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**Table 4. N-Terminal Binding Site in LMMC PBPs**

<table>
<thead>
<tr>
<th>R39</th>
<th>BsPBP4a</th>
<th>NgPBP3</th>
<th>EcPBP4</th>
<th>interaction with $S$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trp139</td>
<td>Pro142</td>
<td>Pro152</td>
<td>Phe147</td>
<td>NH$_3^+$</td>
</tr>
<tr>
<td>Asp142</td>
<td>Asp145</td>
<td>Asp155</td>
<td>Asp150</td>
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<tr>
<td>Tyr147</td>
<td>Tyr150</td>
<td>Phe160</td>
<td>Phe155</td>
<td>(CH$_3$)$_2$</td>
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<tr>
<td>Arg351</td>
<td>His352</td>
<td>Arg361</td>
<td>Arg352</td>
<td>CO$_2^-$</td>
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<tr>
<td>Met414</td>
<td>Leu415</td>
<td>Leu421</td>
<td>Leu408</td>
<td>(CH$_3$)$_2$</td>
</tr>
<tr>
<td>Ser415</td>
<td>Ser416</td>
<td>Gln422</td>
<td>Asn409</td>
<td>CO$_2^-$</td>
</tr>
</tbody>
</table>
NgPBP4 is a likely orthologue (25.5% identical, 59.6% similar; 255-residue overlap; Lalign56) of EcPBP5 in another Gram-negative bacterium. It shows more reactivity \( (k_{cat}/K_m) \), in general, with the peptide panel than the \( E. coli \) enzyme does but still not significant affinity \( (K_m) \) for any particular peptide (Table 1). The enzyme is comparably reactive with the N-terminally extended peptides 7, 8, and 10 as with 5, the peptide with the free N-terminus. These results are in accord with those from boronate inhibition (Table 3) where NgPBP4 shows greater affinity for 11 and 12 than does EcPBP5, but no discrimination between the two. NgPBP4 also appears as unenthusiastic about an extended leaving group [e.g., peptide \( 13 \) (Table 1)] as for the extended N-terminus. This pattern of reactivity rather mirrors that of NgPBP3, although NgPBP3 and NgPBP4 are structurally quite different enzymes from different subgroups. Boronates \( 11 \) and \( 12 \), however, give no evidence of a strong N-terminal specificity of NgPBP4 as they do for NgPBP3. It is interesting to note here that the amino acid sequence of NgPBP4 is more similar to that of EcPBP7 (42.9% identical, 240-residue overlap; Lalign56) than that of EcPBP5, and EcPBP7 is thought to be an endopeptidase;57 perhaps PBP4 in \( N. gonorrhoeae \) covers the roles of both PBP5 and PBP7 in \( E. coli \).

SpPBP3 is similar to NgPBP4 in general reactivity with peptides, although slightly different in detail (Table 2). This enzyme is also a member of the LMMA group but, as noted above, is derived from a Gram-positive organism with a stem peptide containing lysine\(^{22} \) rather than diaminopimelic acid.\(^{22} \) Although SpPBP3 does not show high affinity for any of the group of peptides \( 14 - 19 \), it does have comparable reactivity with \( 13 \), the peptide with the free N-terminus, as with the N-terminally extended peptides \( 14, 15, 18, \) and \( 19 \), where the latter two contain a significant number of the elements of a peptidoglycan dimer. The presence of this extended N-acetylation in \( 18 \) and \( 19 \) does not enhance the reactivity of the peptide or its affinity for the enzyme. SpPBP3 does, however, show distinct aversion to extended leucine group peptides 16 and 17, thus displaying the distinct signature of a carboxypeptidase (unlike NgPBP4).

SpPBP3 does bind boronates \( 20 \) and \( 21 \) quite tightly but shows little discrimination between them. Thioesters are also substrates of LMMA enzymes, but little substrate specificity with respect to peptidoglycan structure has been noticed, e.g., with substrates of LMMA enzymes, but little substrate specificity with EcPBP5, and SpPBP3 displays the distinct signature of a carboxypeptidase (unlike NgPBP4).

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The results described above tend to divide the LMM DD-peptidases into three groups on the basis of their behavior toward peptidoglycan-mimetic ligands. First, the soluble LMMB and LMMC enzymes, the R61 and R39 DD-peptidases, respectively, can be distinguished from a pair of LMMC enzymes, BsPBP4a and NgPBP3. The latter pair, like the former, apparently has a well-defined and specific binding site for a peptidoglycan-mimetic substrate, but unlike with the former group, its presence is not reflected in substrate turnover kinetics in vitro. The latter enzymes are, however, inhibited strongly and specifically by a peptidoglycan-mimetic boronate. This transition-state analogue apparently has the ability to induce peptidoglycan specificity in the enzyme active site. Finally, the LMMA enzymes, EcPBP5, NgPBP4, and SpPBP3, show no marked affinity for peptidoglycan-mimetic substrates or boronate inhibitors, and thus, these ligands are unable to induce a specific active site structure under the conditions employed. Crystal structures have not, to date, revealed a specific binding site on these enzymes beyond that accommodating D-alanyl-D-alanine.\(^{62} \) Although the details of the in vivo function of all of the enzymes mentioned above are not well understood, it is interesting to note that the soluble enzymes have no defined role in peptidoglycan synthesis and maintenance while the membrane-bound/associated enzymes have at least tentatively assigned roles.\(^{48, 82} \)

One thus gets the impression that a number of these enzymes are present in solution in less than optimally specific and/or reactive conformations and that these cannot efficiently be induced into specific or reactive forms by small peptidoglycan-mimetic peptide fragments. In some cases (LMMC), although not in others (LMMA), they can be so induced by tighter binding transition-state analogues. A crystal structure of one of the LMMC enzymes, currently available in apo form (e.g., BsPBP4a), bound to a specific boronate would thus be of considerable interest. \( \beta \)-Lactams, transition-state analogues themselves,\(^{65} \) are also able to induce reactivity with many of these enzymes. One must conclude that the in vivo environment must also be able to induce specific activity by interactions yet unknown. The same conclusion applies to HMM PBPs, which, 85.6% similar, 387-residue overlap\(^{56} \), although the role of PBP6 in vivo may be different from that of PBP5.\(^{58-61} \) A crystal structure of PBP6 has recently been described and, notably, also one with an extended stem peptide analogue bound.\(^{62} \)

A striking feature of the latter structure is that the N-terminus of the peptide (lysine) does not interact with the protein at all and extends directly out into solution. The structure is certainly in accord with the kinetic results for EcPBP5 in this paper, if it is assumed that a complex of an analogous peptide with EcPBP5 would have a similar structure. There is also little evidence of strong productive interaction between EcPBP5 or EcPBP6 and the NAG-NAM sugars of the peptidoglycan monomer.\(^{52, 60} \) A crystal structure of a complex of PBP5 with a \( \beta \)-lactam bearing a 0-α-aminopimelyl side chain showed the side chain to be so mobile that it cannot be observed.\(^{59} \) One possibility is that, in vivo, the extended N-terminus interacts with either the peptidoglycan or another protein, which then induces specificity. There does not appear to be convincing evidence of complexes of EcPBP5 with other PBPs or cell cycle proteins,\(^{6} \) although there is a report of a complex among PBP1b, PBP2, PBP4, and PBP5, isolated by chemical cross-linking, in \( Haemophilus influenzae \), another Gram-negative bacterium.\(^{64} \)

### CONCLUSIONS

The results described above tend to divide the LMM DD-peptidases into three groups on the basis of their behavior toward peptidoglycan-mimetic ligands. First, the soluble LMMB and LMMC enzymes, the R61 and R39 DD-peptidases, respectively, can be distinguished from a pair of LMMC enzymes, BsPBP4a and NgPBP3. The latter pair, like the former, apparently has a well-defined and specific binding site for a peptidoglycan-mimetic substrate, but unlike with the former group, its presence is not reflected in substrate turnover kinetics in vitro. The latter enzymes are, however, inhibited strongly and specifically by a peptidoglycan-mimetic boronate. This transition-state analogue apparently has the ability to induce peptidoglycan specificity in the enzyme active site. Finally, the LMMA enzymes, EcPBP5, NgPBP4, and SpPBP3, show no marked affinity for peptidoglycan-mimetic substrates or boronate inhibitors, and thus, these ligands are unable to induce a specific active site structure under the conditions employed. Crystal structures have not, to date, revealed a specific binding site on these enzymes beyond that accommodating D-alanyl-D-alanine.\(^{62} \) Although the details of the in vivo function of all of the enzymes mentioned above are not well understood, it is interesting to note that the soluble enzymes have no defined role in peptidoglycan synthesis and maintenance while the membrane-bound/associated enzymes have at least tentatively assigned roles.\(^{48, 82} \)
as solubilized forms in vitro, also do not in general turn over small peptidoglycan-mimetic substrates at rates sufficient to support bacterial growth.\textsuperscript{34,35,66} Larger model substrates may be needed.\textsuperscript{67,68} There is some evidence of alternative conformations of particular HMM PBPs from crystal structures \textsuperscript{69–71} and solution studies.\textsuperscript{72–74} It is thus clear that an understanding of the molecular details of catalysis in vivo by the most important of these enzymes has not yet been achieved.

**ASSOCIATED CONTENT**

**Supporting Information**

Synthetic details for the preparation of peptides \textsuperscript{6–9} and boronates \textsuperscript{12, 20, and 21}. This material is available free of charge via the Internet at http://pubs.acs.org.

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**ABBREVIATIONS**

BsPB4a, \textit{B. subtilis} penicillin-binding protein 4a; EcPB5, \textit{E. coli} penicillin-binding protein 5; HMM, high-molecular mass; LMM, low-molecular mass; NAG, N-acetylglucosamine; NAM, N-acetylmuramic acid; NgPB3 and NgPB4, \textit{N. gonorrhoeae} penicillin-binding proteins 3 and 4, respectively; NMR, nuclear magnetic resonance; SpPB3, \textit{S. pneumoniae} penicillin-binding protein 3.

**ADDITIONAL NOTE**

“It should be noted that peptidoglycan in \textit{S. pneumoniae} typically contains both unbrided (direct D-Ala-D-Lys) and bridged (D-Ala-D-\textit{l}-Ala-D-\textit{l}-Lys) or (D-Ala-D-\textit{l}-Ala-D-\textit{l}-Ala-D-\textit{l}-Lys) cross-links; in wild-type strains, >50% of the N-termini are simply D-Lys.\textsuperscript{36–38} The elaborated N-termini are found to predominate in \beta-lactam-resistant strains. \textit{S. pneumoniae} transpeptidases must therefore catalyze nucleophilic attack on an acyl–enzyme intermediate by the \alpha-amino of lysine and also by the terminal \alpha-amino of an \textit{l}-Ala peptide, the latter in an interesting and, as yet, not well-characterized \textit{dL}-transpeptidase reaction. This paper deals with the former reaction, expected to be catalyzed by “wild-type” transpeptidases, but not the latter.

**REFERENCES**


