Identification of Regions Involved in Enzymatic Stability of Peptide Deformylase of *Mycobacterium tuberculosis*

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Sequence analysis of peptide deformylase of *Mycobacterium tuberculosis* revealed the presence of insertions (residues 74 to 85) and an unusually long carboxy-terminal end (residues 182 to 197). Our results with deletion mutants indicated the contribution of these regions in maintaining enzymatic stability. Furthermore, we showed that the region spanning the insertions was responsible for maintaining resistance to oxidizing agents, like H$_2$O$_2$.

The amino-terminal ends of all nascent polypeptides in eubacteria are formylated. As N-terminal peptidases are unable to utilize formylated peptides as substrates, their removal is a mandatory step during protein synthesis. Since the enzyme peptide deformylase (PDF) is known to deformylate the N-formyl group of the nascent polypeptide chains in the cytoplasm of bacteria (3), its importance has long been appreciated. Available genome sequencing data revealed the presence of a putative gene encoding the peptide deformylase (def) throughout the eubacterial lineage (12), including pathogens (1, 5, 9–11, 14). In this context, we concentrated on the peptide deformylase enzyme of the bacterial pathogen *Mycobacterium tuberculosis* (mPDF), which causes the dreadful disease tuberculosis.

Recently, we PCR amplified the def gene from *M. tuberculosis* and, following cloning in pET28c vector, expressed it as a histidine-tagged fusion protein (mPDF) in *Escherichia coli* strain BL21(DE3). The overexpressed protein was obtained as inclusion bodies and subsequently solubilized with 3 M urea. This was followed by dialysis at 4°C against 20 mM phosphate buffer, pH 7.4, and finally the mPDF protein was purified using a Ni-nitrilotriacetic acid (NTA) column (17). Although the available literature indicated that an extended C-terminal end is characteristic of type I PDFs (2, 4, 7), interestingly we found that mPDF possessed an unusual feature. The unusually long C-terminal end (amino acid residues 182 to 197) of mPDF was very striking compared to the sequences of type II members (Fig. 1) (17), and it is likely to form a helix by the sequence-based secondary-structure prediction program (6). Besides this, like other gram-positive bacteria (type II class), mPDF possessed insertions (amino acid residues 74 to 85) (Fig. 1) between conserved motifs 1 and 2, which are predicted to form a loop (6). We created two deletion mutants of mPDF, one at the insertion sequences (designated ID, where 6 amino acids, MTARRR, were deleted) and the other at the C-terminal end (named TD, where 16 amino acids, PG LSWLPGEPPDFGH, were removed), employing a PCR-based mutagenesis approach (17, 18) (primer sequences are given in Table 1). Like mPDF, they (both ID and TD) were obtained as inclusion bodies when overexpressed as histidine-tagged fusion proteins. The mutant proteins were solubilized with 3 M urea, refolded, and purified by using Ni-NTA columns, as described for mPDF (17). This was followed by assessment of enzyme activity to evaluate the contribution of these regions to the deformylation ability of mPDF. The enzyme activity of mPDF was determined in the presence of catalase and bovine serum albumin (BSA) by using N-formyl-Met-Ala as the substrate in the trinitrobenzenesulfonic acid (TNBSA) assay (17).

For overexpression of different proteins, BL21(DE3) cells harboring wild-type (WT) or mutant constructs in pET28c vector were grown and induced with 0.4 mM IPTG (isopropyl-β-D-thiogalactopyranoside) by following the method described earlier (17). Cells were harvested, suspended in lysis buffer (20 mM
phosphate buffer, pH 7.4, containing 5 mM dithiothreitol, 10 \mu g/ml of catalase, 1 mM phenylmethylsulfonyl fluoride, 1 \mu g/ml of pepstatin, and 1 \mu g/ml of leupeptin), and sonicated. The pellet fraction (12,000 \times 9262 g for 30 min at 4°C) was resuspended in lysis buffer containing 3 M urea and 2% Triton X-100. Following centrifugation, the supernatant fraction was dialyzed (14 h at 4°C against 20 mM phosphate buffer, pH 7.4) to remove urea and purified on an Ni-NTA column (17). Finally, mPDF or different mutants were eluted in elution buffer (20 mM phosphate buffer, pH 7.4, containing 300 mM NaCl, 250 mM imidazole, and 10 g/ml of catalase).

The expressed mutant proteins (ID and TD) were recognized by the anti-His-tag antibody, as evidenced by Western blotting (Fig. 2A). As reported earlier, deletion of TR (Fig. 1), resulting in the mutant TD, affected deformylase activity of mPDF (Fig. 2B, left panels) (17). Like TD, even with the use of an excess amount of protein (20 \mu g incubated with 5 mM of N-formyl-Met-Ala) in the assays, the ID mutant hardly showed any deformylase activity (Fig. 2B, left, panel 1). Loss in enzyme activity has often been correlated with its stability. To address this question, we mixed both of the mutant proteins (keeping the total protein concentration of stock the same for each mutant) with each other (ID:TD and TD:ID at 1:4, named ID:TD and TD:ID, respectively) after denaturation with urea. Following refolding (removing urea by slow dialysis [17]) and subsequent purification through Ni-NTA resins, their deformylation abilities were assessed as the function of increasing concentration of total proteins. Among them, the use of 20 \mu g of “cofolded” protein displayed marginal deformylation ability (for ID:TD, 11–16%, and for TD:ID, 25–4%) compared to the wild type (100%), suggesting the abilities of both mutants in partially complementing enzyme activity (Fig. 2B, left, panel 2). To further confirm this aspect, mutants were mixed with the wild type (WT:ID and WT:TD at 1:4, named WT:ID and WT:TD, respectively) and used to assess enzymatic activity. Interestingly, both WT:ID and WT:TD displayed the ability to deformylate N-formyl-Met-Ala more efficiently than the corresponding amounts of the wild-type protein in the mixtures (Fig. 2B, left, panel 3). To highlight the authenticity of such an observation, we carried out a similar experiment by using a C106S mutant of mPDF (mutation at the metal ion coordinating Cys at motif 2 of mPDF), which did not exhibit any deformylase activity (17). Unlike WT:ID and WT:TD, the C106S mutant (recognized by the anti-His-tag antibody) had 0% deformylation activity. To overcome this, we used the anti-His-tag antibody.

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\text{TABLE 1. List of PCR primers used in this study}^{a}
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<table>
<thead>
<tr>
<th>Primer name</th>
<th>Primer sequence (5' to 3')</th>
<th>Amplified gene or region</th>
</tr>
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<tbody>
<tr>
<td>CR26 (forward)</td>
<td>GGAATTCCATAATGCGGATGTCGTAACC</td>
<td>def</td>
</tr>
<tr>
<td>CR27 (reverse)</td>
<td>CCCAAGCTTTTATGACCGGAACCGG</td>
<td>def</td>
</tr>
<tr>
<td>CR23 (reverse)</td>
<td>TTAACGCGCCCACGCTAG</td>
<td>TD mutant of def</td>
</tr>
<tr>
<td>CR25 (internal)</td>
<td>ACCCGCCAGGTTGGTCTGTTAAT</td>
<td>ID mutant of def</td>
</tr>
<tr>
<td>CR24 (internal)</td>
<td>ACCACACCTGGCGGCTTCCG</td>
<td>ID mutant of def</td>
</tr>
</tbody>
</table>

\(^a\) CR26 and CR27 were used as external primers for the amplification of the def open reading frame along with mutation. GGAATTCATATGCGGATGTCGTAACC in CR26 and CCCAAGCTTTTATGACCGGAACCGG in CR27 do not correspond to the genome sequences but have been introduced to incorporate NdeI and HindIII sites at respective 5' and 3' ends of the PCR-amplified products. Details of PCR amplification, construction of recombinant plasmid, and generation of PDF mutants were described elsewhere (17).
Fig. 2A), when mixed with the wild type (WT:C106S at 1:4, named WT+/C106S), showed enzyme activity at the level corresponding to the amount of wild-type protein present in the mixture (Fig. 2B, right). These results argued that mPDF was catalytically active as a multimer, and it is therefore logical to presume that heteromonomeric units of the wild type and ID/TD exhibited cooperativeness among themselves for rendering enzymatic activity of cofolded proteins. As shown in Fig. 2C, the extent of deformylation with 800 ng of total protein (amounts of wild-type and TD/ID proteins were 160 ng and 640 ng, respectively) was increased by 58 to 75% compared to that of the wild-type control (160 ng protein). On the other hand, such an increase in enzyme activity was not evident with the WT+/C106S mixture (Fig. 2C). Therefore, the obvious explanation of such an observation was the contribution of either ID or TD towards the enzymatic activity of wild-type mPDF. Furthermore, this increase in the enzyme activity could not be seen if the purified mutant (ID or TD) or any other proteins (catalase/BSA) were mixed with the wild type at the same ratio during the assay (data not shown). Interestingly, following mixing with the wild type (WT+/ID or WT+/TD; amount of total protein used per assay was 70 ng), both of the mutants lost deformylation ability within 2 h (Fig. 3). Even the use of an increased amount of total protein (350 ng, where the amount of WT protein was 70 ng) in the assay yielded similar results (Fig. 3, inset, showing a representative experiment). Thus, all of this evidence argues that the loss in enzyme activity for the ID or TD mutant was the result of the enzymatic instability of the protein.

The activity as well as the stability of an enzyme often depends on its intermolecular association status. PDF from *E. coli* has been reported to be active as a monomer and that of *Leptospira interrogans* as a dimer (9, 16). To gain insight on this aspect, we determined the molecular masses of the purified recombinant mPDF and its deletion mutants (ID and TD) through dynamic light-scattering studies performed at 25°C on a Dyna Pro instrument (Protein Solutions) equipped with DYNAMICS V6 software by using a microcuvette. Samples (10 to 15 μl) used for this study were always filtered through a 0.1-M filtration device (Millipore). The majority of the wild-type mPDF in solution showed a molecular mass of 100 kDa, with a radius of the molecules of 4.3 nm (Table 2). Since the calculated molecular mass of the protein was 22.6 kDa (although by sodium dodecyl sulfate-polyacrylamide gel electrophoresis analysis, the expressed protein exhibited an anomalous migration of ~31 kDa [Fig. 2A] [17]), our results argued the multimeric nature of the catalytically active mPDF. Interestingly, in the ID mutant these parameters remained unaltered. On the
other hand, considering the deletion of 16 amino acids from mPDF, in the TD mutant both radius and molecular mass of the protein were less than those of the wild type, but the multimeric nature was evident (Table 2). By gel filtration chromatography (AKTAprime Plus chromatography system with a Sephacryl 200 column; GE Biosciences), mPDF exhibited two peaks, one at the void volume and the other at \(~44\) kDa (data not shown). We found that only the \(~44\) kDa protein had the deformylation ability (\(k_{cat}/K_m\) of \(~1,084\) M\(^{-1}\) s\(^{-1}\) as opposed to \(~1,220\) M\(^{-1}\) s\(^{-1}\) reported in reference 17 with the supernatant fraction obtained following low-speed centrifugation) and that further mixing of these fractions (high and low molecular masses) did not have any effect on the enzyme activity of mPDF. As expected, by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, the \(~44\) kDa protein showed a molecular mass of \(~31\) kDa and glutaraldehyde cross-linking (0.0125% at 25°C for 30 to 240 min) indicated the existence of high-molecular-mass (\(~100\) kDa) forms (data not shown). Notably, the dynamic light-scattering data suggested that the mPDF was predominantly tetramer, while gel filtration chromatography indicated its dimeric status. Since both methods rely on shape and hydrodynamic volume and do not always coincide, we prefer to leave this issue untouched for the time being. However, our results unequivocally established the multimeric nature of the active protein, and mutations at IR or TR did not affect the intermolecular association status of mPDF.

Analysis of the crystal structure of PDF from \(L.\) interrogans, which is a Zn\(^{2+}\)-containing metalloprotease, indicated that part of the C-terminal end was involved in anchoring of the substrate to the active site of the protein (20). A similar role of this region in mPDF is rather difficult to predict from our study, as insertion of the C-terminal end to the active site of the neighboring molecules might occur due to crystal packing. However, it is apparent from our results that the extended C-terminal end of mPDF definitely influenced enzyme stability. IR, on the other hand, has been predicted to form a loop in mPDF (6). For other gram-positive bacteria, this loop was found to be flexible but had no role in substrate binding (7).

![FIG. 3. Assessment of enzymatic stability of mPDF mutants. Overexpressed and purified proteins were obtained as described in the text. Enzyme activities were monitored with 17.5 ng of WT or 70 ng of WT+ID or WT+TD proteins in the presence of catalase and BSA by using 5 mM of N-formyl-Met-Ala as the substrate in the TNBSA assay (17). (Inset) Representative experiment showing the enzyme activity with 5 mM of N-formyl-Met-Ala using 70 ng of WT or 350 ng of WT+ID or WT+TD protein.](http://jb.asm.org/)

![FIG. 4. Effect of \(H_2O_2\) on deformylase activity of mPDF mutants. Overexpressed and purified proteins were obtained as described in the text. WT (17.5 ng/reaction), WT+ID, or WT+TD (each cofolded protein, 70 ng/reaction) proteins were preincubated with or without \(H_2O_2\) (final concentration, 500 mM) at 30°C for 15 min. This was followed by an enzyme assay in the presence of catalase and BSA with 5 mM of N-formyl-Met-Ala as the substrate, following the method described earlier (17).](http://jb.asm.org/)
The enzymatic stability, especially in Fe$^{3+}$-containing PDFs, is a perplexing issue. The iron was coordinated with histidine and cysteine (H148, H152, and C106S in the M. tuberculosis enzyme) residues in all PDFs (2, 4, 7, 15). The metal-coordinating cysteine S has been shown to undergo oxidation to cysteine-sulfonic acid and/or cysteine-sulfinic acid in E. coli, S. aureus, Streptococcus pneumoniae, and Thermotoga maritima, resulting in severe consequences or even complete loss of enzyme activity (7, 15). Thus, despite structural identity near the active site metal, stabilities of PDFs in different bacteria were very different (2). These findings led to the postulation that such variation might be due to protein dynamics and/or alteration in sequences beyond the conserved regions, which could affect the rate of oxidation of iron or amino acid side chains that are metal ion ligands (2).

We have recently reported that preincubation of mPDF with an oxidizing agent, like H$_2$O$_2$, had no significant effect on its deformylating ability, despite the presence of Fe$^{3+}$ at its metal binding core (17). This seems to be an important observation, considering the fact that M. tuberculosis has to cope with oxidative stress for its survival within the host as a successful pathogen. To know the contributions of IR or TR (Fig. 1) of the mPDF towards its resistance to oxidizing agents, we utilized WT and TD proteins (total protein, 70 ng) and monitored the effect of H$_2$O$_2$ (500 mM) on enzyme activity. As shown in Fig. 4, preincubation with H$_2$O$_2$ for 15 min exhibited a significant decrease in the deformylase activity of WT+TD compared to that of the wild type or WT+TD. Under our assay conditions, even the use of an increased amount of total protein displayed similar results (data not shown). Thus, our findings argued for a contribution of the IR region, specifically MTARRR sequences, in protecting Fe$^{3+}$ in the metal binding core of mPDF from oxidation to the Fe$^{3+}$ form. Interestingly, sequence analysis of known iron-containing PDFs revealed that three consecutive arginines in the IR of mPDF are typical of different mycobacterial species (M. avium, M. bovis, M. leprae, M. smegmatis, and M. tuberculosis). Furthermore, interaction of oxygen with the side chain of arginine has already been established (8) and, therefore, it could play a crucial role in preventing oxidation of Fe$^{3+}$ in mycobacterial PDFs. However, structure-function analysis needs to be carried out to unravel the mystery. Nonetheless, the results we present here unequivocally establish the association of IR and TR of mPDF in maintaining its enzymatic stability.

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