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Acylpeptidyl Oligopeptidase: Porphyromonas gingivalis Periplasmic Novel Exopeptidase Releases N-Acylated Di- and Tri-peptides from Oligopeptides*

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*Running title: Acylpeptidyl oligopeptidase in P. gingivalis

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ABSTRACT

Exopeptidases including dipeptidyl- and tripeptidyl-peptidase are crucial for the growth of Porphyromonas gingivalis, a periodontopathic asaccharolytic bacterium that incorporates amino acids mainly as di- and tri-peptides. In this study, we identified a novel exopeptidase, designated acylpeptidyl oligopeptidase (AOP), composed of 759 amino acid residues with active Ser⁶¹⁵ and encoded by PGN_1349 in P. gingivalis ATCC 33277. AOP is currently listed as an unassigned S9-family peptidase or prolyl oligopeptidase. Recombinant AOP did not hydrolyze a Pro-Xaa bond. In addition, though sequence similarities to human and archaea-type acylaminoacyl peptidase sequences were observed, its enzymatic properties were apparently distinct from those, as AOP scarcely released an N-acyl-amino acid as compared to di- and tri-peptides, especially with N-terminal modification. The $k_{cat}/K_m$ value against benzoxycarbonyl-Val-Lys-Met-4-methylcoumaryl-7-amide, the most potential substrate, was $123.3 \pm 17.3 \mu M^{-1} sec^{-1}$, optimal pH was 7-8.5, and the activity was decreased with increased NaCl concentrations. AOP existed predominantly in the periplasmic fraction as a monomer, while equilibrium between monomers and oligomers was observed with a recombinant molecule, suggesting a tendency of oligomerization mediated by the N-terminal region (Met¹⁶-Glu¹⁰¹). The three dimensional modeling revealed the three domain structures: residues Met¹⁶-Ala¹²⁵, which has no similar homologue with known structure, residues Leu¹²⁷-Met⁴⁹⁵ (β-propeller domain) and residues Ala⁴⁹⁶-Phe⁷³⁶ (α/β hydrolase domain), and further indicated the hydrophobic S1 site of AOP in accord with its hydrophobic P1 preference. AOP orthologues are widely distributed in bacteria, archaea, and eukaryotes, suggesting its importance for processing of nutritional and/or bioactive oligopeptides.

Porphyromonas gingivalis, a Gram-negative obligate anaerobe, has been implicated as the causative agent of chronic periodontal disease (1, 2), which is a major reason for permanent tooth loss (3-5). Recently, much attention has been paid to this bacterium due to its association with systemic diseases, such as cardiovascular disorders (6), decreased kidney function (7), and rheumatoid arthritis (8). P. gingivalis does not ferment carbohydrates, but rather metabolizes amino acids to produce ATP by a putative respiratory chain of fumarate respiration without oxygen (9). In addition, it incorporates nutritional
amino acids not as single amino acids, but preferentially as di- and tri-peptides and short oligopeptides, possibly via inner-membrane-associated oligopeptide transporters (9, 10).

In *P. gingivalis*, extracellular nutritional proteins are initially digested to oligopeptides by potent cysteine endopeptidases, such as Arg-gingipain (Rgp) (C25.001 in MEROPS classification) (11) and Lys-gingipain (Kgp) (C25.002) (12), and subsequently degraded into di- and tri-peptides by exopeptidases, i.e., dipeptidyl-peptidases (DPPs) (13-16) and prolyl tripeptidylpeptidase A (PTP-A) (17), respectively. Therefore, the topology of the sub-cellular localization of these peptidases, i.e., extracellular and outer membrane-bound Rgp/Kgp, and DPPs/PTP-A in periplasmic space, appears to be suitable for an ordered degradation of proteinaceous substrates and their incorporation into cells. Through amino acid metabolism, the organism excretes sulfide, ammonia, butyrate (18, 19), and methyl mercaptan (20) as end-products, which have been suggested to cause host tissue damage (21-23). Accordingly, DPPs, PTP-A, and gingipains are considered to play crucial roles not only in cell growth but also bacterial pathogenicity.

Thus far, four DPPs, i.e., DPP4, DPP5, DPP7, and DPP11, have been identified in *P. gingivalis*. These are serine peptidases belonging to either the S9 or S46 family, with DPP4 (S09.003) showing a preference for Pro at P1 (13, 14) and DPP5 (S09.012), the first entity identified in bacteria, showing a preference for hydrophobic P1 residues and no specificity at the P2 position (24). Furthermore, DPP7 (S46.001) has a hydrophobic preference for the P1 (15, 25) as well as P2 (26) positions, while we discovered that DPP11 (S46.002) is a unique DPP specific for acidic P1 residues (Asp and Glu) (27). In addition to these DPPs, *P. gingivalis* possesses a metallopeptidase encoded by the gene PGN_1645, which was identified as DPP3 (M49.001) and specific for P1 Arg. However, DPP3 appears to be localized in the cytosol, while the Arg-specific DPP activity of Rgp plays a role in extracellular substrate processing (24). We also found Lys-specific DPP activity in Kgp (24). DPPs do not cleave polypeptides with Pro at the third position from the N-terminus, while PTP-A (S09.017) is able to release the N-terminal tripeptide Xaa1- Xaa2-Pro3 (16, 17). Therefore, most extracellular polypeptides, at least those without N-terminal modification, should be sequentially and completely degraded into di- or tri-peptides in *P. gingivalis* by the cooperative activities of the four DPPs, PTP-A, and gingipains (24).

On the other hand, our previous observations of *P. gingivalis* NDP212, a *dpp4*-5.7-11-knockout strain, suggested the existence of an unidentified DPP responsible for Met-Leu-MCA hydrolysis, since the activity was markedly elevated in the mutant strain as compared to a *dpp4*-5.7-11-knockout strain (24). In order to define this entity, we focused on and studied the remaining uncharacterized three putative S9-family proteins, i.e., PGN_1349, PGN_1542, and PGN_1878, of *P. gingivalis* in the present study. We found peptidase activity exclusively in PGN_1349 and, interestingly, most exopeptidase activities remaining in NDP212 were explained by the activity of PGN_1349. The observed peptidase properties of PGN_1349 indicated it as a novel oligopeptidase, designated acylpeptidyl oligopeptidase (AOP), with a preference for hydrophobic amino acids at the P1 position of its substrates, especially those with N-terminal modification.

### EXPERIMENTAL PROCEDURES

**Materials**—pQE60 (Qiagen Inc., Chatsworth, CA) and pTrcHis2-TOPO (Invitrogen, Carlsbad, CA) were used as expression vectors. Restriction enzymes and DNA-modifying enzymes were purchased from Takara Bio (Tokyo, Japan) and New England Biolabs (Ipswich, MA), respectively. Quick Taq HS DyeMix and KOD-Plus-Neo DNA polymerase came from Toyobo (Tokyo, Japan). Met-Leu-4-methycoumaryl-7-amide (MCA) was from Bachem (Bubendorf, Switzerland), and Leu-Asp-, Leu-Glu-, Lys-Met-, benzylxycarbonyl-(Z)-Lys-Met-, and Z-AVKM-MCA were synthesized by Thermo Fisher Scientific (Ulm, Germany) and Scrum (Tokyo, Japan). Oligonucleotide primers were synthesized by FASMAC (Atsugi, Japan). Low-molecular-weight markers, full-range rainbow molecular weight markers, rabbit muscle aldolase, egg white ovalbumin, and a Sephacryl S-200 High Resolution and Superdex 200 Increase 10/300 columns were from GE Healthcare (Little Chalfont, UK). Bovine liver
catalase and bovine milk α-lactalbumin were obtained from Sigma-Aldrich (St. Louis, MS). Lysozyme from egg white and formyl cellulofine were from Seikagaku Biobusiness Corp. (Tokyo, Japan).

**Culture Conditions—**P. gingivalis strains ATCC 33277, KDP136 (28), NDP212 (24), and NDP600 were grown anaerobically (80% N₂, 10% CO₂, 10% H₂) in enriched brain heart infusion broth (Becton Dickinson, Franklin Lakes, NJ) supplemented with 5 µg/ml of hemin and 0.5 µg/ml of menadione. Ampicillin (10 µg/ml) for NDP600 and appropriate antibiotics (ampicillin, erythromycin, tetracycline, chloramphenicol) were added to the cultures of KDP136 and NDP212, as previously described (24, 27). Bacterial cells were suspended in phosphate-buffered saline (PBS) at pH 7.4, then centrifuged at 6000 x g for 15 min at 4 °C. The cell pellet was washed once with PBS, re-suspended in PBS to adjust absorbance to 2.0 at 600 nm, and then used in the following experiments.

**Expression and Purification of Recombinant Proteins**—Oligonucleotides and plasmids used in this study are listed in Table 1. DNA fragments encoding the mature forms (Met₁⁶-Lys₅⁷⁹) of PGN_1349 (29) (GenBank Accession Number 6329856, MEROPS code MER034614) and Leu₆-Glu²⁷⁹ of PGN_1542 (MER110015) were amplified by PCR using sets of primers (1349-5F1 and 1349-3R2comp), then the obtained DNA fragments were ligated with pTrcHis2-Topo according to the manufacturer’s protocol, resulting in production of pTrcHis2-PGN1349 and -PGN1542, respectively. A DNA fragment encoding Ile₆-Leu⁴⁷₉ of PGN_1878 (MER109588) was PCR amplified using a set of primers (5PGN1878-16Bgl and 3PGN1878-L473Bgl). After restriction cleavage by BglII, the fragment was inserted into the BamHI site of pQE60, resulting in production of pQE60-PGN1878. A deletion mutation of Met₁⁶-Gly⁰₁⁵ (designated pTrcHis2-PGN1349-N102) was constructed using a PCR-based technique with primers 5PGN1349-N102 and 3pTrcHisTopo-L-1, with the substitution of Ser⁶₁⁵ by Ala, (designated pTrcHis2-PGN1349-S615A and pTrcHis2-PGN1349-N102-S615A) constructed from pTrcHis2-PGN1349 and pTrcHis-PGN1349-N102 using the primers 5PGN1349-S615A and 3PGN1349-A614, respectively. Mutations were confirmed by DNA sequencing.

**Disruption of P. gingivalis Genes—**P. gingivalis NDP212, with deletion of four DPP genes (dpp4, dpp5, dpp7, dpp11) (24), and KDP136, with deletions of kgp, rgpA, and rgpB (28), were previously reported. To construct the PGN_1349 gene deletion mutant, DNA fragments of each 5'- and 3'-part of the PGN_1349 gene were PCR amplified with a set of primers (1349-5F1 and cep-comp-1349-5R for the 5' part, cep3F-1349-3F1 and 1349-3Rcomp for the 3' part). A cepA fragment was amplified using primers (1349-5F1-cepF, 1349-3Fcomp-cep3R-comp) and pCR4-TOPO as a template. Nested PCR was performed with a mixture of these three fragments using primers (1349-5F2, 1349-3R2comp), then the obtained DNA fragment (3,440 bp) was introduced into P. gingivalis by electroporation, resulting in NDP600 (pgn1349-cepA).

**Measurement of Peptidase Activity**—Peptidase activity was measured using peptidyl-MCA as previously reported (24, 27). Briefly, the reaction was started by addition of recombinant proteins (5-100 ng), a periplasmic cell fraction (1-5 μl), or P. gingivalis cell suspensions (1-5 μl) in a reaction mixture (200 μl) composed of 50 mM sodium phosphate (pH 7.5), 5 mM EDTA, and 20 μM peptidyl MCA. After 30 min at 37 °C, fluorescence intensity was measured with excitation at 380 nm and emission at 460 nm. In some experiments, pH values varied from 5.5-9.5 with 50 mM phosphate (pH 5.5-8.5) or Tris-HCl (pH 7.0-9.5), and NaCl concentrations varied from 0-1.6 M. To determine enzymatic parameters, recombinant proteins were incubated with various concentrations of peptidyl MCA. Obtained data were analyzed using a nonlinear regression curve fitted to the Michaelis-Menten equation with the GraphPad Prism software program (San Diego, CA). Values are shown as the average ± S.D. and calculated from 4 independent measurements.

**Subcellular Fractionation**—All procedures
were carried out at 4 °C according to a previously reported method (30), with a slight modification. Briefly, a 20-ml culture of *P. gingivalis* in the log-phase was centrifuged at 6000 x g for 15 min, then the extracellular fraction was obtained from the supernatant by filtration with a 0.20-µm membrane filter. Bacterial cells were washed with ice-cold PBS, resuspended in 4 ml of 0.25 M sucrose in 5 mM Tris-HCl (pH 7.5), and then left on ice for 10 min. Cells were precipitated at 12,500 x g for 15 min, resuspended in 4 ml of 5 mM Tris-HCl, and mixed gently for 10 min to disrupt the outer membrane. The supernatant was obtained by centrifugation at 6000 x g for 15 min and collected as the periplasmic fraction containing the outer membrane. The obtained spheroplasts as precipitate were resuspended in PBS and then disrupted in an ice-water bath by sonication pulse 10 times for 10 seconds each with 2-second intervals. The cytosol and inner membrane fractions were separately prepared by ultracentrifugation, as previously described (24).

**Size Exclusion HPLC**—Recombinant proteins and the periplasmic fraction were subjected to size exclusion HPLC using ÄKTA explorer 10S (GE Healthcare) with a Superdex200 increase 10/300 column (1.5 X 30 cm) equilibrated with 20 mM Tris-HCl (pH 8.0) containing 1 mM EDTA. They were separated with the identical buffer at a rate of 0.75 ml/min containing 1 mM EDTA. They were separated into fractions, each of which was collected. Aliquots of the fractions were subjected to a peptidase assay with Z-VKM-MCA, as well as SDS-PAGE or native PAGE.

**Immunoblotting Analysis**—Recombinant PGN_1349 (Met<sup>16</sup>-Lys<sup>759</sup>) was purified by use of the Talon affinity chromatography and further subjected to a Sephacryl S200 HR column (1.5 X 30 cm) equilibrated with 20 mM ammonium bicarbonate (pH 8.5). Rabbit anti-PGN_1349 (AOP) antiserum was prepared as previously reported (24, 27). For immunoblotting, separated proteins on polyvinylidene difluoride membranes (Millipore) were incubated with anti-AOP antiserum (10<sup>-3</sup> fold dilution), followed by alkaline phosphatase-conjugated anti-rabbit IgG. Finally, specific bands were visualized with 5-bromo-4-chloro-3-indolyl phosphate and nitro blue tetrazolium. Similarly, immunoblotting against *P. gingivalis* DPP5 and DPP7 was performed as previously reported (24, 25).

**N-Terminal sequencing of proteins**—Proteins separated on SDS-PAGE and transferred to a Sequi-Blot membrane (Bio-Rad) were stained with Coomassie Brilliant Blue, and subjected to the protein sequencing by use of a model Procise 49XLc protein sequencer (Applied Biosystems).

**3D Homology Modeling**—In order to search for an ideal template for homology modeling, *P. gingivalis* AOP sequence was submitted to Psipred (31) and GenTherader (32) server for domain and fold assignment. Following, a Blast search (33) against the Protein Data Bank was performed to identify the closest homologue to *P. gingivalis* AOP with known 3D structure. The model was generated via the server Phyre2 (Protein Homology/Analogy Recognition Engine) (34), using the tool “one-to-one threading”, in the expert mode. The coordinates of *Pyrococcus horikoshii* acylaminoaoyl peptidase (AAP) (35) (PDB code: 4HXE) and *P. gingivalis* AOP sequence were submitted to structural alignment and further model building. The final model covered around 85% of the sequence, sharing 20% identity with the template.

**RESULTS**

**PGN_1349 Responsible for Remaining Activity in NDP212**—We previously reported that Met-Leu-MCA hydrolyzing activity was markedly decreased in a dpp4-5-7 triple knockout *P. gingivalis* strain (NDP211). However, unexpectedly, that activity was reversed by introduction of a fourth dpp-gene disruption to NDP211, resulting in NDP212, a *dpp4-5-7-11* quadruple knockout strain (24). These results indicated that the bacterium additionally expresses a peptidase related to Met-Leu-MCA hydrolysis in addition to DPP5 and DPP7, and suggested an up-regulation mechanism to compensate for the loss of di-peptide production.

To identify this entity, the peptidase activity of NDP212 was reexamined in detail with various MCA substrates (portion of results shown in Fig. 1A). NDP212 showed the most efficient hydrolysis of Met-Leu-MCA, followed by benzylxoycarbonyl (Z)-LLL- and succinyl (Suc)-LLVY-MCA, suggesting that the candidate is not a DPP, but likely an oligopeptidase with a hydrophobic P1 preference. These activities were observed in a cell-associated form and not with the culture supernatant, and were efficiently inhibited by diisopropyl fluorophosphates, suggesting a serine peptidase, the same as previously reported periplasmic DPPs (data not
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shown).

Taking into consideration these properties, along with information from the KEGG and MEROPS databases (29, 36), we focused on three uncharacterized peptidases among the seven S9-family members, i.e., PGN_1349 (KEGG definition: DPP), PGN_1542 (esterase), and PGN_1878 (hypothetical protein), and expressed them as C-terminal His<sub>6</sub>-tagged proteins in E. coli. Purified PGN_1349, PGN_1542, and PGN_1878 migrated as 85-<i>kDa</i>, 35-, and 50-<i>kDa</i> species, respectively, in accordance with their calculated molecular masses (Fig. 1B). Subsequent peptidase analysis using 80 peptidyl MCAs available in our laboratory revealed that PGN_1349 hydrolyzed several substrates. In contrast, attempts to detect peptidase activities in PGN_1542 and PGN_1878 have not been successful to date.

PGN_1349 is currently annotated as a DPP, though was found to be hydrolyzed by both N-terminally blocked and unblocked substrates, such as Met-Leu-MCA, Z-LLL-, Z-VKM-, and AAF-MCA, to a small extent (Fig. 1C). Other substrates showed much lower activities, including glutalyl (Glt)-AAF- (4.21 ± 0.05 pmol/min/µg, mean ± S.D., n = 3), Suc-AAA- (1.01 ± 0.05), N-methoxysuccinyl [Suc(OMe)]AAP- (0.38 ± 0.02), GP- (0.28±0.01), acetyl (Ac)-Ala- (0.12 ± 0.00), Lys-Ala- (0.10 ± 0.00), Ac-Met- (0.05 ± 0.002), Z-MEK- (0.026 ± 0.005), Gly-Gly- (0.017 ± 0.004), Z-LRGG- (0.015 ± 0.000), t-butyloxycarbonyl-[25]-2-amino-3-(benzyloxycarbonyl) propionyl]- (Boc)-FSR- (0.013 ± 0.012), Z-LLE- (0.009 ± 0.002), Ala- (0.009 ± 0.000), and Leu-Asp-MCA (0.004 ± 0.002). These results indicate that PGN_1349 is not a DPP, but rather an oligopeptidase that is able to liberate N-terminally blocked peptides with a P1 preference for hydrophobic residues. Interestingly, its substrate-preference profile resembles that of NDP212, except for no activity of PGN_1349 against Suc-LLVY-MCA. In addition, NDP212 scarcely hydrolyzed Z-VKM-MCA, while PGN_1349 showed efficient hydrolysis. We speculated that the reason for the low level of Z-VKM-MCA hydrolysis by NDP212 was due to its Kgp activity, which preceded the hydrolysis of Z-VKM-MCA into Z-Val-Lys and Met-MCA, in which the latter was scarcely degraded by PGN_1349 (Fig. 1C). In fact, the level of hydrolysis of Z-VKM-MCA was even higher than that of Z-LLL-MCA by the gingipain-null strain KDP136 (data not shown). At this time, we are unable to explain the activity against Suc-LLVY-MCA in NDP212, though it may be due to an unidentified peptidase. Taken together, we concluded that most hydrolyzing activities observed in NDP212 can be mainly attributed to PGN_1349.

Distribution of PGN_1349 Orthologues in Organisms—PGN_1349 in P. gingivalis ATCC 33277 and its homologues in other <i>P. gingivalis</i> strains have been annotated under various names, such as DPP (ATCC 33277) (29), prolyl oligopeptidase (W83) (10), putative DPP (TDC60) (37), and peptidase S9 A/B/C family catalytic domain protein (JCV1 SC001) (38). In the MEROPS peptidase database, PGN_1349 is classified into Clan SC, family S9, subfamily C of the peptidase S09.A77, for which the dipetidyl-peptidase family member 6 (DPF-6) from <i>Caenorhabditis elegans</i> (UniProt #P34422) is a representative. An orthologue search of PGN_1349 produced a long list of S9 family peptidase members (Supplemental Table S1). This family comprises at least 142 genera of organisms, in which most orthologues are distributed in both Gram-negative and -positive bacteria, such as the phyla <i>Actinobacteria</i>, <i>Proteobacteria</i>, <i>Bacteroidetes</i>, <i>Cyanobacteria</i>, and <i>Firmicutes</i>, but are also found in archaea (the phyla <i>Euarchaeota</i> and <i>Crenarchaeota</i>) including <i>Aeropyrum pernix</i> and eukaryotes (the phylum <i>Nematoda</i>) including <i>C. elegans</i>. The average amino acid length of the 477 DPF-6 members is 652 ± 62 (mean ± S.D.).

DPF-6 from <i>C. elegans</i> is composed of 740 amino acids (39) and postulated to have serine-type peptidase activity, while no actual activity in any family members has yet been reported. Alignment of the amino acid sequences indicates that the Ser-Asp-His catalytic triad and their adjacent sequences are highly conserved (Fig. 2). The peptidase S9 family motif is predicted to be present in the C-terminal half region in both PGN_1349 and DPF-6 by the Pfam database (40). The N-terminal part corresponding to Met<sup>16</sup>-Lys<sup>142</sup> of PGN_1349 is absent in <i>C. elegans</i> DPF-6. Importantly, this sequence identity between PGN_1349 and <i>C. elegans</i> DPF-6 (27.3%, INT/OPT score = 132/653) is even higher than that with <i>C. elegans</i> DPF-5/AAP (23.3%, 82/292), which releases acylaminoacyl amino acids, or <i>E. coli</i> oligopeptidase B (19.8%, 24/123), which releases acyloligopeptides from the N-terminus of
These findings strongly suggest that the enzymatic activities of DPF-6 family members are represented by that of P. gingivalis PGN_1349, and enzymatically distinct from DPF-5/AAP and oligopeptidase B.

**PGN_1349 Is a Novel Exopeptidase, Designated Acylpeptidyl Oligopeptidase (AOP)**—The peptidase activity of PGN_1349 is re-arranged based on the residue numbers of 26 MCA substrates. It showed predominant release of tripeptides with N-terminal blockage. Other substrates, such as acylaminoacyl-, dipeptidyl-, and N-terminally-blocked tetrapeptidyl-MCA, were slowly hydrolyzed, while aminoacyl-MCA was scarcely hydrolyzed (Fig. 3A). To more precisely evaluate the substrate length preference, we prepared a set of substrates with altered lengths, i.e., Z-Met-, Z-Lys-Met-, Z-VKM-, and Z-AVKM-MCA, and then determined the activities (Fig. 3B). PGN_1349 most potently hydrolyzed Z-Lys-Met-MCA, followed by Z-VKM-MCA. The hydrolysis of Z-AVKM-MCA was lower as compared to the former and Z-Met-MCA was scarcely hydrolyzed. Taken together, we concluded that PGN_1349 is neither a DPP nor AAP, though it has acyl-oligopeptide-releasing activity. In order to more precisely evaluate the influence of N-terminal blockage on its enzymatic activity, the hydrolysis of four pairs of substrates, with the N-terminal either blocked or un-blocked, were examined. As shown in Table 2, PGN_1349 consistently hydrolyzed the acylated substrates more efficiently than the substrates without N-terminal modification. Hence, we designated PGN_1349 as acylpeptidyl oligopeptidase (AOP).

The P1-position amino acid preference of AOP was further examined using synthetic dipeptidyl MCA or N-terminally modified tripeptidyl MCA with different P1 amino acids. When the activities for these substrates were plotted against P1 amino acid residual hydrophobicity indexes (41), a close positive relationship with P1-position hydrophobicity was demonstrated with both the di- and tri-peptidyl MCA substrates (Fig. 3C). These results clearly indicate that AOP prefers hydrophobic residues at the P1 position.

The pH profile of AOP revealed an optimum pH of 8.5 for Met-Leu-MCA and 7.0 for Z-VKM-MCA. The activity was gradually decreased as the concentration of NaCl increased. Therefore, to achieve direct comparison with various substrates, the activity was simply measured in 50 mM sodium phosphate buffer, pH 7.5, containing 5 mM EDTA in the absence of NaCl in the following experiments. We also tested the effects of reducing reagents, i.e., dithiothreitol, reduced glutathione, and oxidized glutathione, since it has been reported that they modulate the peptidase activity of oligopeptidase B (42). However, none (up to 1.6 mM) affected the peptidase activity of AOP (data not shown), again indicating characteristics distinct from oligopeptidase B.

The enzymatic parameters of AOP for the three substrates indicate that the $k_{cat}$ value was maximal for Z-Lys-Met-MCA ($1155 \pm 77 \text{ sec}^{-1}$), while $K_m$ was smallest for Z-VKM-MCA ($4.9 \pm 1.0 \mu M$) and $k_{cat}/K_m$ maximal for Z-VKM-MCA ($123.3 \pm 17.0 \mu M^{-1} \text{ sec}^{-1}$) (Table 3). When the values were compared with those of P. gingivalis DPPs for their best dipeptidyl-MCA substrates, the $k_{cat}/K_m$ values of AOP ($58 \sim 123 \mu M^{-1} \text{ sec}^{-1}$) were shown to be comparable with those of DPP4, DPP5, DPP7, and DPP11 ($11 \sim 548 \mu M^{-1} \text{ sec}^{-1}$), suggesting that AOP and the DPPs are able to function under similar substrate concentrations in the bacterium.

**Peptidase Activities in aop-Gene Disrupted Strain**—An aop-disrupted strain, NDP600, was constructed and absence of AOP in the strain was confirmed by immunoblotting (Fig. 4A). Cell-associated peptidase activity of NDP600 was determined in the presence of 50 µM N-tosyl-lysine chloroketone (TLCK) and 3 µM E-64, under which the activity of Kgp toward Z-HEK-MCA was completely abolished in both the wild-type and NDP600 strains (Fig. 4B), while addition of these gingipain inhibitors markedly enhanced Z-Lys-Met-MCA hydrolysis in the wild type. In contrast, this activity was greatly reduced in NDP600 both with and without the inhibitors (Fig. 4C). These results clearly demonstrated that AOP is the main peptidase responsible for this hydrolysis. Similarly, the hydrolysis of Z-LLL-MCA was significantly reduced in NDP600 (Fig. 4D). However, hydrolyses of Z-VKM- and Z-AVKM-MCA were modestly reduced in NDP600, presumably because of incomplete protection of the substrates from Kgp. On the other hand, that of Met-Leu-MCA was not affected by disruption of the aop-gene, as this substrate was predominantly hydrolyzed by DPP5 (24) and DPP7 (25), which are active in NDP600. Taken together, these results suggest that AOP is primarily responsible for hydrolysis of Z-Lys-Met-MCA in P. gingivalis.
for release of di- and tri-peptides with P1 hydrophobic residues from N-terminally-blocked oligopeptides in *P. gingivalis*.

**Active Serine and Endopeptidase Activity**—Recombinant AOP was mainly purified as an 85-kDa species with the N-terminal sequence of Ala²-Ala⁶-Met¹⁶-Thr-Val-His-Ala-Gln-Lys-Ile²³ ( Ala²-Ile¹ was derived from the vector sequence) due to release of the initial met³ (Figs. 1 and 2). In addition, a minor 75-kDa species with the N-terminal sequences of Thr⁸⁹-Phe-Lys-Gly-Thr-Leu⁹⁴ and Asn¹⁰²-Val-Glu-Gln-Met-Gly¹⁰⁷ was occasionally obtained, indicating cleavage at the Phe⁸⁸-Thr⁸⁹ and Gly¹⁰¹-Asn¹⁰² bonds, respectively (Figs. 2 and 5A). These results suggested that the N-terminal region at around the 100th residue is protease sensitive. Similarly, 75- and 70-kDa species were produced by limited proteolysis with chymotrypsin at the Phe¹⁰⁸-Ser¹⁰⁹ and Phe¹¹⁷-Glu¹¹⁸ bonds, respectively (data not shown).

We compared the peptidase activities of several forms of AOP, in which either N-terminal 101 residues were truncated (Asn102 form), the potentially essential Ser at position 615 was substituted by Ala (S615A form), or both. The S615A substitution completely abolished the activity (Fig. 5B). These results indicate that Ser⁶¹⁵ is essential, as previously proposed (36). The N-terminal truncated Asn102 form maintained one-third of the activity, suggesting that N-terminal 101 residues are not essential, though have affects on peptidase activity.

Interestingly, 75-, 52-, and 42-kDa species emerged in enzymatically-active forms and were increased by incubation at 37 °C for 6 days (Fig. 5A), suggesting an auto-proteolytic activity of AOP. In accord with the P1 hydrophobic preference of AOP activity, the cleavage sites of the 52- and 42-kDa species were Leu⁵⁹⁰-Tyr⁵⁹¹ and Ala⁴¹⁴-Tyr⁴¹⁵, respectively. Endopeptidase activity of AOP was further confirmed, as it degraded 14.3-kDa α-lactalbumin into 7.5- and 8.5-kDa species at the Gln²⁵⁸-Asn²⁵⁹ and Gln⁵₈-Ala⁵₉ bonds, respectively (data not shown), and was estimated to be less than 1/100th of that of α-chymotrypsin when assessed with α-lactalbumin as a substrate.

**Oligomerization and Subcellular Localization**—We also examined the molecular structure of AOP in an aqueous solution using size-exclusion HPLC (Fig. 6). The hydrolyzing activity of purified AOP was split into a major peak larger than 440 kDa and a minor 85-kDa peak. Fractions 14-17 of the first HPLC consisted solely of the 85-kDa species, as shown by SDS-PAGE, while the second peak at fractions 22-25 consisted of both the 85- and 75-kDa entities. Native PAGE confirmed that the 85-kDa species in the first peak formed oligomers, and that the 85- and 75-kDa species in the second peak were monomers (m1 and m2, respectively). Furthermore, the second HPLC of the large peak sample (fraction 15 of the first HPLC) consisting of the 85-kDa species reproduced the 2-peak profile. These findings clearly indicated an equilibrium of oligomeric and monomeric forms of the 85-kDa AOP in solution, and comparable specific activities of these two forms. Since the 75-kDa species starting at Thr⁸⁹ or Asn¹⁰² was a monomer, the N-terminal region (Met¹⁶-Glu¹⁰¹) appears to be indispensable for AOP oligomerization.

In *P. gingivalis* KDP136 cells, the apparent molecular weight of native AOP was found to be 78 kDa (Fig. 7), suggesting loss of some region as compared with 85- and 75-kDa AOP, though the N-terminus of the 78-kDa form could not be determined. AOP at 78 kDa was predominantly detected in the periplasm/outer membrane and cytosol fractions. When the periplasmic/outer membrane fraction was subjected to size-exclusion HPLC, 78-kDa AOP was detected as a monomer at fractions 23-26. Since the cytosolic preparation contained a considerable amount of periplasmic components, *P. gingivalis* AOP appears to be primarily distributed in periplasm and present mainly as a monomer in cells. In accord with these findings, a previous proteome analysis also found that PG_1004, an orthologue of PGN_1349 in *P. gingivalis* W83, was mainly recovered in cytosol/periplasm and periplasm fractions (43).

**Modeling of 3D Structure**—Psipred and GenThreader results indicate that AOP possesses three domains: Met¹⁶-Ala¹²⁶, which has no similar homologue with known 3D structure. Leu¹⁷-Met⁴⁹⁵ and Ala⁴⁹⁶-Phe⁷³⁶ form β-propeller and α/β hydrolase domains, respectively. The 3D model comprises residues Ala¹³⁸ to Lys⁷⁵⁹ and bears the domain organization of S9 family members, as indicated by GenThread. The α/β hydrolase domain hosts the catalytic triad (Ser⁶¹⁵, Asp⁷⁰², and His⁷³⁴) and is responsible for the enzymatic activity, while the β-propeller provides enzyme specificity (Fig. 8A and B).
In agreement with the hydrophobic P1 preference, the S1 site is formed by the hydrophobic residues: Pro$^{536}$, Ile$^{583}$, Tyr$^{619}$, Val$^{641}$, Ile$^{644}$, Phe$^{647}$, and Trp$^{666}$ (Fig. 8C). Differently from DPPs, that have an N-anchor region determining the protein specificity (44, 45), our model indicates a more open active site without any steric impediment for its oligopeptidase feature. The auto-proteolytic site (Gly$^{101}$-Asn$^{102}$) associated to the production of the 75-kDa truncated AOP is localized in the N-terminal domain. It suggests that this region is very likely exposed to the solvent, reasonably explaining the site accessibility by another AOP molecule.

To estimate the evolutionary conservation of amino acid positions in AOP, we performed a Psi-Blast (33) search with AOP sequence, followed by multi alignment in Cobalt (46), and mapped their conservation onto the AOP structure, using the server Consurf (47). This analysis revealed that the α/β-hydrolase domain is more conserved than the β-propeller domain (Fig. 8D), in agreement with the observation that the β-propeller domain is responsible for enzyme specificity. Therefore, β-propeller domain is expected to present more variation among the proteins than the α/β-hydrolase domain, which provides the catalytic activity shared by all members. Furthermore, because the oligomer interface region lies in the N-terminus, we hypothesize that AOP might display a novel oligomer formation process. In this way, determining the 3D structure of AOP by an experimental method (for instance, X-ray crystallography) could shed light into its mechanism of action.

**DISCUSSION**

In the present study, we identified a novel exopeptidase encoded by the gene PGN_1349 in the *P. gingivalis* strain ATCC 33277. This enzyme, designated acylpeptidyl oligopeptidase (AOP), was found to possess active Ser$^{615}$, most preferentially degrade peptide bonds between Yaa$^{2}$-Xaa$^{3}$ and Yaa$^{3}$-Xaa$^{4}$ (Yaa is a hydrophobic amino acid) of oligopeptides with N-terminally blocked oligopeptides, and exist mainly in periplasmic space as a monomer. Furthermore, AOP showed a tendency of oligomerization at the N-terminal 100-amino acid region, thus it may bind to other molecules via interactions in this region. Our 3D modeling results also indicated a conserved structure of S9 peptidases coinciding with the hydrophobic P1 preference of AOP.

Peptidase activities of three of seven members categorized as *P. gingivalis* S9-family proteins have been identified in previous studies, *i.e.*, DPP4, DPP5, and PTP-A (14, 15, 24, 27), while the present study identified the fourth, AOP. Whether the remaining three, *i.e.*, PGN_1542 (annotated as esterase), PGN_1694 (Ala-DPP), and PGN_1878 (hypothetical protein), possess hydrolyzing activity remains unknown (24, this study), however, we confirmed here that they could not release N-acylated amino acids from MCA substrates available in our laboratory. Therefore, to the best of our knowledge, AOP is a unique exopeptidase that hydrolyzes nutritional N-terminally blocked peptides in *P. gingivalis*. The activity of AOP may provide substantial benefits for efficient utilization of amino acids by the bacterium. Notably, when considering its habitat in sub-gingival space, the bacterium frequently utilizes serum proteins contained in gingival crevicular fluid, some of which are N-terminally blocked by acyltransferase or glutamyl cyclase by processing into their secretory forms (48).

Our comparison of amino acid sequences supported the kinship of AOP with *C. elegans* DPP-6 (S9.A77). On the other hand, sequence homologies were rather lower among AOP, *C. elegans* DPF-5/AAP, and *E. coli* oligopeptidase B, and their peptidase characteristics are obviously distant. Although both AOP and DPF-5/AAP exhibit a hydrophobic P1 preference, a typical feature of mammalian (S09.004) and *Aeropyrum*-type (S09.070) AAP is removal of an N-acylated hydrophobic amino acid from oligopeptides with various N-terminal acyl groups (49, 50). Because of this activity, AAP is also termed an acylamino acid-releasing enzyme. In contrast, AOP was shown to scarcely release acylamino acids, but preferably released di- and tri-peptides (Fig. 1). Furthermore, the P1 hydrophobic preference of AOP is clearly distinct from oligopeptidase B, which targets basic amino acid residues such as Arg (51). On the other hand, both share common features in that these exopeptidases poorly exhibit aminopeptidase activity and possess elevated activities against N-acylated substrates (52, this study). Therefore, DPF-6 family members including *P. gingivalis* AOP should be considered as the third exopeptidase group in the S9 family responsible for N-terminally blocked peptidyl substrates. Although AOP possessed a weak endopeptidase
activity, this activity was also reported on AAP (53).

In Supplemental Table S1, Bacteroides peptidases, such as Bacteroides fragilis (BF9343_3151, 23.4% amino acid identity), Bacteroides vulgatus (BVU_4083, 21.6%), and Bacteroides thetaiotaomicron (BT_1838, 23.8%) are listed as homologues of P. gingivalis AOP (PGN_1349). Among them, BT_1838 and BVU_4083 are annotated as Ala-DPP in the S9 family, of which enzymatic activity remains unknown (24). The amino acid identities of P. gingivalis Ala-DPP (PGN_1694) to BF_9343_3151 from B. fragilis, BVU_4083, and BT_1838 are 39.1%, 39.4%, and 40.2%, respectively, which are substantially higher than those to P. gingivalis AOP. In fact, we cloned BF_9343_3151 and expressed its recombinant form in E. coli as an 80-kDa protein. It did not show any peptidase activity (data not shown), as P. gingivalis Ala-DPP did not (24). Therefore, we conclude that all members of Supplemental Table S1 are not AOP, especially those with lower similarities. The final annotation of AOP should be carefully done in future studies. In addition, a study on peptidase activity of Ala-DPP is now underway in our laboratory.

Subcellular fractionation experiments demonstrated that P. gingivalis AOP is localized in periplasmic space as a monomer. It has been reported that S9 family members are present as various molecular forms (50), as oligopeptidase B and prolyl oligopeptidase were found to exist as monomers (42), and DPP4 (54) as well as PTP-A as a dimer (17), while the dimeric structure of human DPP4 is indispensible for peptidase activity (54). In addition, mammalian and Aeropyrum pernix AAPs are present as a tetramer (55) and dimer (56), respectively. P. gingivalis DPP5 also forms a homo-dimer (24). In this study, we found that recombinant AOP exists in equilibrium between a large oligomer and monomer. It is interesting to note that two forms of AOP demonstrated peptidase activity and the specific activity of both seems to be equal. Furthermore, our findings indicated that the N-terminal region is indispensible for oligomerization, though the biological meaning of AOP oligomerization is obscure at present. This region might be related to heteromeric interactions with periplasmic components, such as other peptidases and peptidoglycan.

Genome and biochemical analyses of P. gingivalis demonstrated the presence of two oligopeptidase transporters (9, 10) and a Ser/Thr transporter (57). Previous studies reported that di- and tri-peptides are predominantly incorporated into cells (18, 19), thus they appear to be major cargoes for the oligopeptide transporters and the contribution of the Ser/Thr transporter toward nutritional amino acid incorporation may be limited. Regarding this point, there is a critical difference between P. gingivalis and E. coli, a well-studied Gram-negative rod that does not possess DPPs and predominantly incorporates single amino acids (58, 59). It is possible that DPPs/PTP-A and oligopeptides transporters co-evolved, which may explain, at least in part, why novel exopeptidases, i.e., DPP11 (27), DPP5 (24), and AOP (this study), have recently been discovered in P. gingivalis.

Based on our findings, we describe the metabolism of extracellular oligopeptides in P. gingivalis as follows. P. gingivalis produces oligopeptides from extracellular polypeptides by potent cysteine peptidases, Kgp and Rgp, located in the outer membrane and extracellular space. Subsequently, oligopeptides are converted in periplasmic space into di- and tri-peptides by DPPs and PTP-A, respectively, with DPP5 and DPP7 functioning with oligopeptides with hydrophobic residues at the second position from the N-terminus. Additionally, the P2-position hydrophobic residue enhances the activities of DPP7 (26). Those with Pro at the second and third positions are cleaved by DPP4 and PTP-A, respectively. Oligopeptides with the penultimate acidic residues Asp and Glu are metabolized by DPP11, while those with the basic residues Lys and Arg are metabolized by gingipains. Moreover, N-terminally blocked oligopeptides are segregated by AOP into acylated di- and tri-peptides, and N-terminally un-blocked oligopeptides, which are then processed by DPPs and PTP-A. These di- and tri-peptides are promptly incorporated via oligopeptide transporters in the inner membrane. Efficient utilization of extracellular proteins as carbon and energy sources is made possible by coordination of a series of exopeptidases and nutritional oligopeptides transporters in P. gingivalis, which may have inevitably evolved to choose and adapt its anaerobic sub-gingival habitat where carbohydrates are consumed mainly by oral streptococci.

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Conflict of interest—The authors declare that they have no conflicts of interest with the contents of this article.

Author Contributions—T. K. N. and Y. O.-N. designed and performed the experiments. Y. S. and S. K. performed amino acid sequencing. G. A. B. performed 3D modeling. T. K. N. and Y. O.-N. wrote the first draft. All authors contributed and approved the final version of the manuscript.
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Acylpeptidyl oligopeptidase in P. gingivalis

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FOOTNOTES

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2Supported by the Vienna International Postdoctoral Program (VIPS).

3The abbreviations used are: Rgp, Arg-gingipain; Kgp, Lys-gingipain; DPP, dipeptidyl-peptidase; PTP-A, prolyl tripeptidylpeptidase A; AOP, acylpeptidyl oligopeptidase; AAP, acylaminoacyl peptidase; MCA, 4-methcoumaryl-7-amide; Z-, benzoyloxycarbonyl-; Suc-, succinyl-; Glt-, glutaryl-; Suc(OMe)-, N-methoxysuccinyl-; Ac-, acetyl; Boc-, t-butyloxycarbonyl-[(2S)-2-amino-3-(benzyloxycarbonyl)propionyl-].
Acylpeptidyl oligopeptidase in \textit{P. gingivalis}

FIGURE 1. Peptidase activities remaining in \textit{dpp4-5-7-11} knockout strain and recombinant \textit{PGN\_1349}.

\textit{A}, The peptidase activity of NDP212 cells (\textit{Adpp4-5-7-11}) was determined with peptidyl MCA. \textit{B}, Purified proteins (0.5 µg) of \textit{PGN\_1349} (lane 1), \textit{PGN\_1542} (lane 2), and \textit{PGN\_1878} (lane 3) were separated on SDS-PAGE gels and detected by Coomassie blue staining. Lane M, molecular-weight marker. \textit{C}, The peptidase activity of recombinant \textit{PGN\_1349} was determined with peptidyl MCA. Values are shown as the mean ± S.D. (n = 3).

FIGURE 2. Comparison of amino acid sequences between \textit{PGN\_1349} and \textit{C. elegans} DPF-6.

The amino acid sequence of \textit{PGN\_1349} (PgAOP) was compared with that of \textit{C. elegans} DPF-6 (CeDPF-6). Identical amino acids are indicated by asterisks. Potential sets of 3 amino acids (Ser\textsuperscript{615}, Asp\textsuperscript{702}, His\textsuperscript{734}) forming the essential triad of serine peptidases are indicated by red bold letters. Putative signal sequences are shown in bold italic and peptidase S9 family domains (residues 544-759 on PgAOP and 445-662 on CeDPF-6) by green boxes. The starting amino acids (Met\textsuperscript{16} and Asn\textsuperscript{102}) of recombinant proteins expressed in this study (see Fig. 5) are boxed in yellow. N-terminal sequences and the auto-proteolytic cleavage sites producing 75-kDa species in the purified sample, 75- and 52-kDa species appearing after incubation at 37 °C for 6 days (Fig. 5), are indicated by blue arrows and arrowheads, respectively. Chymotryptic cleavage sites are indicated by open arrowheads (see text). Lower panel, Percent homology of amino acid sequences among PgAOP, CeDPF-6, CeDPF-5/AAP, and \textit{E. coli} (Ec) oligopeptidase B were determined using Genetyx software.

FIGURE 3. Properties of \textit{PGN\_1349} (AOP).

\textit{A}, Peptidase activities of AOP with MCA substrates were arranged according to the numbers of amino acids and N-terminal states. 1, aminoacyl-MCA (Ala-, Leu-, Met-, and Phe-MCA); N+1, N-terminal-modified aminoacyl MCA (Ac-Ala- and Ac-Met-MCA); 2, dipeptidyl-MCA (Met-Leu-, Gly-Pro-, Gly-Phe-, Lys-Ala-, Ser-Tyr-, Gly-Gly-, Leu-Asp-, and Thr-Ser-MCA); N+3; N-terminal-modified tripeptidyl-MCA (Z-VKM-, Z-LLL-, Glr-AAF-, Suc-AAA-, Suc-IIW-, Suc(OMe)-AAP-, Z-MEK-, Boc-FSR-, and Z-LLL-MCA); N+4, N-terminal-modified tetrapeptidyl MCA (Suc-LLVY- and Z-LRGG-MCA). \textit{B}, The activity of \textit{PGN\_1349} was determined with Z-Met-, Z-Lys-Met-, Z-VKM-, and Z-AVKM-MCA. \textit{C}, Activities of dipeptidyl-MCA (open square, broken line, “2” in panel \textit{A}) and N-terminal-modified tripeptidyl-MCA (closed square, solid line, “N+3”) were re-plotted against the hydrophobicity index of the P1 residues. Data for Gly-Pro- and Suc(OMe)AAP-MCA were omitted from this plot because of the peculiarity of the imino acid, Pro. \textit{D}, The activity of recombinant AOP was determined with Met-Leu- (open circle) and Z-VKM-MCA (closed circle) in 50 mM sodium phosphate (pH 5.5-8.5) or 50 mM Tris-HCl (pH 7.0-9.5). The highest activities in phosphate buffer (pH 8.5) for Met-Leu-MCA (48.6 ± 0.9 pmol/min/µg) and in Tris-HCl (pH 7.0) for Z-VKM-MCA (367.9 ± 8.0 pmol/min/µg) were used as 100%. \textit{E}, The activity of \textit{PGN\_1349} for Z-VKM-MCA was determined at 0-1.6 M NaCl in 50 mM Tris-HCl buffer, pH 8.0. Values are shown as the mean ± S.D. (n = 3).

FIGURE 4. Effect of gene disruption of AOP.

\textit{A}, Cell suspensions (10 µl) from ATCC 33277 (wt) and an AOP-knockout strain (NDP600) were subjected to immunoblotting analysis with anti-AOP, DP5, and DPP7 antibodies. \textit{B}, Z-HEK-MCA- and \textit{C}, Z-Lys-Met-MCA-hydrolyzing activities were determined with cell suspensions of wt and KD600 in the absence or presence of 60-µM TLCK and 3-µM E-64. \textit{D}, Peptidase activities were determined using wt and NDP600 in the presence of TLCK and E-64.

FIGURE 5. Effects of N-terminal truncation and substitution of Ser\textsuperscript{615} in AOP.

\textit{A}, Recombinant AOP and its derivatives (10 µg) were heated in 100 µl of SDS-sample buffer just after purification (-) or incubated at 37 °C for 6 days (+). Then, aliquots (1 µg) were subjected to SDS-PAGE. Lane M, molecular-weight markers. Apparent molecular masses of the major products are indicated on the right. \textit{B}, Activities of recombinant AOP and its derivatives were determined with...
Z-Lys-Met-MCA (mean ± S.D., n = 3).

FIGURE 6. Molecular status of recombinant AOP.
A, AOP (10 mg/0.3 ml) and fraction 15 of the first HPLC (0.3 ml) were subjected to size-exclusion HPLC (first and second, respectively) with a Superdex200 10/300 column, as described in “EXPERIMENTAL PROCEDURES.” Relative activities (first, FU/µl, and second, FU/5 µl) were determined with Z-VKM-MCA. Molecular markers run in parallel were as follows: blue dextran 2000 (2000 kDa), ferritin (440 kDa), aldolase (158 kDa), ovalbumin (43 kDa), and cytochrome C (12 kDa). B, Aliquots (2.5 µl for SDS-PAGE and 10 µl for native PAGE) of each fraction of the first HPLC were separated on PAGE gels under denaturing (SDS-PAGE) and non-denaturing (Native PAGE) conditions. sm, starting material (2 µg with SDS-PAGE, 16 µg with Native PAGE). Positions of oligomers and monomers (m1 and m2) are indicated. Fraction 15 of the first HPLC (f15) and each fraction (5 µl) of the second HPLC were separated on SDS-PAGE gels. Lanes M, molecular-weight markers.

FIGURE 7. Molecular status of endogenous AOP.
A, Recombinant AOP (50 ng) and sub-cellular fractions of KDP136 (2 µg protein) were separated on SDS-PAGE gels, and subjected to immunoblotting. Lane M, full-length rainbow molecular weight marker: lane 1, recombinant wt AOP; lane 2, Asn102; lane 3, KDP136 whole cells; lane 4, cytosol; lane 5, inner membrane; lane 6, periplasm/outer membrane. B, The periplasm/outer membrane fraction of KDP136 (0.3 ml) was loaded onto an HPLC column, as described in Fig. 6. Aliquots (5 µl) of separated fractions were subjected to SDS-PAGE, then visualized by immunoblotting with the anti-AOP antibody. p/o, periplasm/outer membrane fraction (5 µl).

FIGURE 8. 3D homology model of AOP.
A, Cartoon representation of AOP 3D structure, β-propeller domain (Ala138-Met495) is shown in blue and α/β hydrolase domain (Ala496-Lys759) is in green. B, Surface representation of AOP with the catalytic triad region (red). C, Stick representation of catalytic triad (yellow) and the residues forming the S1 subsite (magenta). D, Surface representation of AOP colored in terms of evolutionary conservation of its amino acids. Psi-Blast, Cobalt and Consurf were used to calculate the conservation scores. Left panel: Side view of AOP. Middle panel: top view of α/β-hydrolase domain, β-propeller was removed for better visualization. Right panel: top view of β-propeller domain. The color code ranges from blue (non-conserved) to red (conserved). The figure was prepared using the program PyMOL (http://www.pymol.org).
**TABLE 1**

**Primers, plasmids, and DNA fragments used in this study**

Restriction and mutated sites are shown as *italics* and underlined, respectively.

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TABLE 3
Enzymatic parameters of *P. gingivalis* exopeptidases

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<th>$k_{cat}$ (sec$^{-1}$)</th>
<th>$K_m$ (µM)</th>
<th>$k_{cat}/K_m$ (µM$^{-1}$sec$^{-1}$)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>AOP</td>
<td>Z-Lys-Met-</td>
<td>1155 ± 77</td>
<td>17.8 ± 2.0</td>
<td>65.0 ± 3.7</td>
<td>this study</td>
</tr>
<tr>
<td></td>
<td>Z-VKM-</td>
<td>592 ± 40</td>
<td>4.9 ± 1.0</td>
<td>123.3 ± 17.3</td>
<td>this study</td>
</tr>
<tr>
<td></td>
<td>Z-AVKM-</td>
<td>399 ± 13</td>
<td>6.9 ± 0.6</td>
<td>57.5 ± 3.7</td>
<td>this study</td>
</tr>
<tr>
<td>DPP4</td>
<td>Gly-Pro-</td>
<td>6917 ± 1253</td>
<td>100.9 ± 20.1</td>
<td>66.8 ± 2.0</td>
<td>this study</td>
</tr>
<tr>
<td>DPP5</td>
<td>Lys-Ala-</td>
<td>1948 ± 165</td>
<td>185 ± 21</td>
<td>10.5</td>
<td>24</td>
</tr>
<tr>
<td>DPP7</td>
<td>Met-Leu-</td>
<td>394 ± 79</td>
<td>39.6±16.0</td>
<td>10.6 ± 2.5</td>
<td>25</td>
</tr>
<tr>
<td>DPP11</td>
<td>Leu-Asp-</td>
<td>10707 ± 140</td>
<td>19.5 ± 0.4</td>
<td>547.4 ± 6.3</td>
<td>this study</td>
</tr>
<tr>
<td></td>
<td>Leu-Glu-</td>
<td>13587 ± 577</td>
<td>80.9 ± 5.7</td>
<td>167.7 ± 5.3</td>
<td>this study</td>
</tr>
</tbody>
</table>
Acylpeptidyl oligopeptidase in P. gingivalis

FIGURE 1

A

B

C

NPD212

PGN1349
**FIGURE 2**

<table>
<thead>
<tr>
<th>Acylopeptidyl oligopeptidase in <em>P. gingivalis</em></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FIGURE 1</strong></td>
<td></td>
</tr>
</tbody>
</table>

**%Homology of amino acid sequences** (INT/OPT Score)

<table>
<thead>
<tr>
<th></th>
<th>PgAOP 100</th>
<th>CeDPF-6 100</th>
<th>CeDPF-SAAP 25.0</th>
<th>Ec oligopeptidase B 19.8</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PgAOP</strong></td>
<td>23.3 (132/653)</td>
<td>25.0 (79/204)</td>
<td>100</td>
<td>24.0 (25/103)</td>
</tr>
<tr>
<td><strong>CeDPF-6</strong></td>
<td>27.3 (82/292)</td>
<td>25.0 (93/292)</td>
<td>100</td>
<td>24.0 (25/103)</td>
</tr>
</tbody>
</table>
FIGURE 3

A

B

C

D

E

Acylpeptidyl oligopeptidase in P. gingivalis
FIGURE 4

Acylpeptidyl oligopeptidase in P. gingivalis

A

B

C

D

Activity (pmol min⁻¹ μL⁻¹)

Activity (pmol min⁻¹ μL⁻¹)

Activity (pmol min⁻¹ μL⁻¹)

Activity (pmol min⁻¹ μL⁻¹)

kDa

Z-HEK-MCA

Z-Lys-Met-MCA

Z-Lys-Met

Z-AVKM

Z-VKM

Z-LLL

Met-Leu

wt

NDP600
FIGURE 5

A

B

Activity (pmol min⁻¹ µg⁻¹)

37˚C, 6 days

M          wt         S615A      Asn102     Asn102

M          wt       S615A   Asn102

Acylpeptidyl oligopeptidase in P. gingivalis
FIGURE 6

A

B

SDS-PAGE of 1st HPLC

Native PAGE of 1st HPLC

oligomer

m1

m2

SDS-PAGE of 2nd HPLC

Fraction no.

Relative activity (FU/1 or 5 µl)

Fraction no.
FIGURE 7

A

B

Acylpeptidyl oligopeptidase in P. gingivalis
FIGURE 8

Acytpeptidyl oligopeptidase in P. gingivalis