Short Communication

Disruption of gingipain oligomerization into non-covalent cell-surface attached complexes

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Abstract

RgpA and Kgp gingipains are non-covalent complexes of endoprotease catalytic and hemagglutinin-adhesin domains on the surface of Porphyromonas gingivalis. A motif conserved in each domain has been suggested to function as an oligomerization motif. We tested this hypothesis by mutating motif residues to hexahistidine tag or hexahistidine tag to disrupt the motif within the Kgp catalytic domain. All modifications led to the secretion of entire Kgp activity into the growth media, predominantly in a form without functional His-tag. This confirmed the role of the conserved motif in correct posttranslational proteolytic processing and assembly of the multidomain complexes.

Keywords: cysteine protease; limited proteolysis; periodontitis; Porphyromonas gingivalis; posttranslational processing.

Arguably, periodontitis is the most prevalent infection-driven chronic inflammatory disorder of mankind and Porphyromonas gingivalis is considered the pivotal aetiologic organism responsible for the development and progression of the disease (Albandar and Rams, 2002; Cobb et al., 2009). This anaerobic, asaccharolytic Gram-negative bacterium is well armed with virulence factors and the gingipain cysteine protease family plays a key role in P. gingivalis pathogenesis (Guo et al., 2010). Except for one laboratory strain of P. gingivalis (HG66), which secretes soluble gingipains, in all other strains, the gingipains are associated with the bacterial outer membrane (Potempa et al., 1995). While Arg-specific gingipain B (RgpB) is a single chain protease (Mikołajczyk-Pawlinska et al., 1998), membrane-associated Lys-specific gingipain (Kgp) and Arg-specific gingipain A (RgpA) are assembled into very large (>300 kDa), multifunctional (Pathirana et al., 2008; Tam et al., 2008, 2009; O’Brien-Simpson et al., 2009), non-covalent complexes composed of two individual unique protease domains (RgpAcat and Kgpcat) and several hemagglutinin-adhesin domains (RgpAHA1, KgPHA1, RgpAHA2, KgPHA2, RgpAHA3, and KgPHA3) (Bhogal et al., 1997) (Figure 1). All domains are generated by limited proteolysis of a single, nascent polypeptide chain encoded by rgpA or kgp genes, with the hemagglutinin-adhesin (HA) domains being located in the C-terminal extension of the translation products of the two genes (Pike et al., 1994; Pavloff et al., 1995, 1997). In addition, some adhesin domains in the complexes are derived from HagA (Pathirana et al., 2006), a large protein encompassing up to 4 HA-domain repeats (Kozarov et al., 1998). With the exception of Lys-Pro peptide bond being cleaved downstream of the hemoglobin-binding domain (HA2), the initial proteolytic processing of other domains occurs at Arg-Xaa peptide bonds (Figure 1). The initial cleavage is followed by progressive truncation of the C-terminus at Lys-Xaa sites by Kgp and removal of the resulting C-terminal Lys by a surface located carboxypeptidase (Figure 1) (Veith et al., 2002, 2004). Individual domains generated in this sequence of proteolytic events remain tightly associated via non-covalent, protein-protein interactions.

A conserved sequence (PVxNLT) present on the catalytic, HA1 and HA3 domains referred to as the adhesin-binding motif (ABM); was suggested to be responsible for oligomerization and assembly of multidomain complexes (Slakeski et al., 1998). To test this hypothesis, we have disrupted this motif in Kgpcat by insertion of hexahistidine residues or by mutation of six consecutive residues of ABM into histidine residues (Figure 1). The accuracy of these genetic manipulations was confirmed by DNA resequencing of the entire kgp gene and secretion of gingipains by different mutants
Figure 1  Schematic representation of preproKgp posttranslational proteolytic processing into catalytic and hemagglutinin/adhesin (HA) domains in *P. gingivalis* strain W83.

The nascent translation product of the *rgpA* gene has the same subdomain structure as preproKgp and the polypeptide is processed in the same way as proKgp (Pavloff et al., 1995, 1997). Parts of the polypeptide chain with sequence identity over 95% in Kgp and RgpA are highlighted red. Initial proteolysis releases individual domains from the polypeptide (A) is followed by progressive truncation of the C-terminus at indicated Lys-Xaa peptide bonds in the catalytic, HA1 and HA2 domains (B). Subsequently, C-terminal lysines are removed by a carboxypeptidase (Veith et al., 2002, 2004). The cleavage sites at the Kgp-HA1 junction are enlarged in panel C. Lowercase sequence indicates the region absent in the non-covalent complex of Kgp cat and HA domains. Blue shaded regions denote the adhesin binding motifs (ABM), which have been proposed to mediate protein-protein interaction between the various domains. Verification of the ABM role in oligomerization of gingipain domains was tested by disruption of the ABM motif located in the catalytic domain by insertion of hexahistidine tag or replacement of ABM residues with hexahistidine (D). In ABM1 and ABM2 mutants, hexahistidine tag was inserted after Pro683 and Pro685, respectively. In mutant ABM3, the sequence V689 SNLTA was replaced with hexahistidine. The proteolytic processing sites identified at the C-termius of purified Kgp variants by the nanoL-ESI-MS/MS analysis of trypic digests are indicated by empty arrowheads. ABM1, ABM2, and ABM3 mutants were obtained by genomic integration of modified plasmid constructs using site-directed mutagenesis (QuickChange II XL Site Directed Mutagenesis Kit, Agilent Technologies, Santa Clara, CA, USA (Smith et al., 1990). Briefly, master plasmid, pKgp/tetQ, used for site-directed mutagenesis was constructed based on the pUC19 vector. The TetQ cassette was amplified from plasmid pNFD13-2 using tetQSalIF (AAT A GT CGA C AAC GAA TTA TCT CCT TAA CGT) and tetQSphIR (TTC GCA TGC TTT TAT TGC CAA GTT CTA ATG CT) primers and inserted into SalI and SphI sites in pUC19. A 5′ fragment for homologous recombination included sequence corresponding to most of the *kgp* gene was amplified by PCR (Accuprime Pfx polymerase, Invitrogen) using *P. gingivalis* W83 genomic DNA as a template and primers KgpFrCXbaIF (GG T CTA GAG GTT CGT ATG CTT GTT GTT G) and KgpFrCSalIR (TAA TA G TCG AC C GAG TCC AAG ACA GAT T) and inserted to XbaI and SalI sites in the plasmid. Similarly, a 1.2-kb 3′ flanking region of the *kgp* gene was amplified using primers KgpFrBSphI (TA G CAT GC G GAC TCG GAG ACT TTG TG) and KgpFrBHindR (ACC T AAG CTT CAA AGC ATC ACA GTA TAA G) and ligated into SphI and HindIII sites. The *kgp* gene sequence encoding the ABM in the master plasmid pKgp/tetQ was modified by QuickChange II XL site-directed mutagenesis method using three sets of oligonucleotide primers: ABM1_F: CAGGTAGTTGAGGCTCTACCATCATTACCATCATACACAGGCCCCAGGC and ABM1_R: GGCGCTGGTAGGGCGCTGTGATGGTTGATGAGGCGGTACGCTACCT; ABM2_F: GTAGGTGAGCCACTACCATCATTACCATCATACACAGGCCCCAGGC and ABM2_R: CAAGTTGAAAACGGGCTGTGATGGTGATGATGGTAGTGGGCGTGAGGCGCTGTGATGGTTGATGAGGCGGTACGCTACCT; ABM3_F: AGCCCTACCGAGCCCATACCATCATTACCATCATACACACAGGCGGTACGCTACCT and ABM3_R: CTTCTTGACCCCTGCGCTGTGATGGTGATGATGGTAGTGGGCGTGAGGCGCTGTGATGGTTGATGAGGCGGTACGCTACCT, each containing the desired mutation and PfuUltra high-fidelity (HF) DNA polymerase, according to manufacturer’s instructions. Obtained pKgp/ABM1, pKgp/ABM2 and pKgp/ABM3 plasmids were used for generation of isogenic mutants via homologous recombination in *P. gingivalis*. Genomic integration by a double crossover event was characterized in comparison to the wild-type parental strain.

The *kgp* gene mutation had no effect on Rgp activity and subcellular distribution. In the mutants, the vast majority of Rgps are cell-associated as in the wild-type strain. However, ABM mutants produced approximately only one-third total Kgp activity of the parental strain and in stark contrast to the wild-type strain, this activity was found almost exclusively in a soluble form in the growth medium regardless of the age of the culture (Figure 2). This decreased production of Kgp was not related to hindered gene expression since qRT-PCR analysis revealed the same level the *kgp* gene transcription was characterized in comparison to the wild-type parental strain.
in mutants and in the parental strain (data not shown). As disintegration of bacteria by sonication did not release any additional Kgp activity, we assumed that lower level of Kgp activity produced by the ABM mutants is due to disturbed folding and impaired maturation of Kgp due to added hexahistidine motif. To assess this possibility, we have subjected the mutants to Western blot analysis.

In accordance with the cellular distribution of Kgp activity in the ABM mutants, a single band representing the Kgp catalytic domain (Kgp cat ) was located exclusively in the media fraction but absent from cell extracts (Figure 3A) in Western blot using Kgp cat -specific mAbs. Interestingly, cell extracts of P. gingivalis cultured only for 24 h revealed the presence of additional weak but clear bands of molecular mass higher than the Kgp cat in the ABM mutants (Figure 3B). The highest band (200 kDa) in ABM2 cell extract may represent a full-length proKgp while the others are proKgp-derived fragments containing Kgp cat. No such bands were found in the wild-type P. gingivalis regardless of the culture age. Therefore, the presence of intermediate forms of Kgp in cell extracts of ABM mutants suggests that indeed, a portion of mutated kgp translation products were misfolded and were inefficiently processed explaining why mutants produce less Kgp activity than the wild-type P. gingivalis strain. In concordance with unaffected Rgp activity and cellular distribution, mutations in Kgp had no effect on Rgps band patterns (Figure 3C).

The partially processed Kgp was also detected in 24 h culture medium of ABM2 and ABM3 mutants (Figure 3B). Western blots with anti-hexahistidine antibodies revealed strong immunoreactive bands only in the media from ABM1 and ABM3 mutants and barely visible band in the ABM2 mutant (Figure 3D) arguing that proKgp polypeptide proteolytic processing occurs downstream of hexahistidine tag in the former two mutants and mostly within or upstream of the tag in the latter strain. Similarly, two higher molecular mass (60- and 70-kDa) forms of Kgp bearing the hexahistidine tags were detected in the media of ABM3 mutant cultured for 24 h (Figure 3E).
Figure 3 Western blot analysis of media and the whole cell fractions derived from wild-type P. gingivalis W83 and isogenic ABM mutants cultivated for 24 h (B, E) or 48 h (A, C, D) for the presence of Kgp (A and B), Rgps (C), and hexahistidine tag (D and E).

For fraction separation, see Figure 2 legend. Proteins in 20 μl of original samples were resolved on 4–12% Bis-Tris SDS-PAGE gels and electrotransferred onto nitrocellulose membranes. After blocking non-specific binding sites, membranes were probed with mAb anti-Kgp (panels A and B), rabbit pAb anti-Rgp (panel C), rabbit pAb anti-6His tag (GenScript, Atlanta, GA, USA) (panels D and E), then secondary antibodies conjugated with alkaline phosphatase and specific bands were visualized with AP conjugate Substrate Kit (BioRad, Hercules, CA, USA). Arrowhead in panel B indicates full length proKgp.

The presence of His-tags on Kgp secreted by ABM1 and ABM3 mutants allows for their affinity purification on Ni-Sepharose (Figure 4A). Surprisingly, only a small fraction (6–7%) of the total Kgp activity was recovered from Ni-Sepharose after ABM1 and ABM3 media were applied onto Ni-chelating matrix. The bulk of Kgp was found unbound in the Ni-Sepharose flow-through fraction. Western blot analysis of bound and unbound Kgp forms separated on the affinity column clearly showed that the latter sample did not interact with anti-hexahistidine antibody (Figures 4B and 4C). The lack of interaction with anti-6His antibody and Ni-Sepharose was found to be due to the lack of hexahistidine motif as inferred from nanoL-ESI-MS/MS analysis of the unbound forms of Kgp. Tryptic-digests of Ni-Sepharose bound Kgp forms returned QIQVGEPSYPHPHHPHHHPHH and QIQVGEPSYPQPHPHHHH peptides in proteins produced by ABM1 and ABM3 mutants, respectively. In stark contrast, in the unbound Kgp variants from all three mutants, no peptides containing the hexahistidine motif were identified. Interestingly, peptides bearing mono- and/or dihistidine at the C-terminus [QIQVGEPH(H), QIQVGEPPSH(H) and QIQVGEPSYPQPH(H)] were found in unbound Kgp produced by ABM1, ABM2, and AMB3 mutants, respectively. In addition, peptides with tetrahistidine (QIQVGEPSYPHHHHH and QIQVGEPSYPQPHHHHH) were also identified in unbound Kgp from the ABM1 and AMB3 mutants. Results of the nanoL-ESI-MS/MS analysis are in agreement with reactivity of these forms with polyclonal Abs reactive with tetra-, penta-, and hexahistidine (Figure 4B) but not with monoclonal Abs specific only for the hexahistidine tag (Figure 4C).

All forms of Kgp obtained by affinity-chromatography and purified by ion-exchange and gel-filtration chromatography from Ni-Sepharose non-binding fraction were subjected to Edman degradation. Regardless which mutant the 50 kDa band was derived from, the same N-terminal sequence of the mature Kgp catalytic domain was found (DVYTDHGDLNYT). Conversely, no sequence was detected in the 60- and 70-kDa forms isolated from the ABM3 mutant, suggestive of a blocked N-terminus. Therefore, 50-, 60-, and 70-kDa protein bands were analyzed by MS to verify protein identity. In tryptic digests of the 70 kDa band, peptides derived from the entire Kgp profragment and catalytic domain were identified. One hundred and ten N-terminal residues of the profragment were missing in the 60 kDa form. Finally, no peptides derived from the profragment were found in the 50 kDa form. Together, it is clear that Kgp isoforms purified from the ABM3 mutant media represent the Kgp catalytic domain alone (50 kDa), with intact (70 kDa) and...
Figure 4  SDS-PAGE (A), Western blot analysis (B and C) and active site probing (D) of soluble forms of Kgp isolated from culture media of isogenic ABM mutants of P. gingivalis cultivated for 24 h. Growth media were dialyzed against binding buffer (20 mM Na-Phosphate, 500 mM NaCl, 20 mM imidazole, 0.02% NaN 3 , pH 7.4) and incubated with Ni 2+ -Sepharose 6 Fast Flow (GE Healthcare, Pittsburgh, PA, USA) slurry with continuous rotation overnight at 4°C. Ni 2+ -Sepharose beads were collected by centrifugation (1000 g, 10 min), washed with the binding buffer and bound protein eluted with 500 mM imidazole. The supernatant was used to purify remaining bulk of Kgp activity. To this end, the supernatant was passed through DE-52 to remove hemin then the enzyme was further purified by gel filtration on Superdex 75 followed by ion-exchange chromatography on a Mono Q column (ACTA). In obtained fractions protein content was determined by the BCA method using bovine serum albumin as a standard. The high molecular weight Kgp complex isolated from strain HG66 (1 μg) and indicated amounts of protein (ABM-U, Kgp non-binding to Ni 2+ -Sepharose and purified by conventional chromatography; ABM-B, Kgp purified on Ni 2+ -Sepharose) were subjected to SDS-PAGE (Panel A), Western blot analysis (5 μg) (Panels B and C) and active-site probing (Panel D). For the latter analysis samples were pretreated with AcKbio-Y-hex-KAOMK (Kato et al., 2005) at 10 μM final concentration for 30 min at room temperature. The reaction was stopped by addition of TLCK to 10 mM concentration. All samples were treated with reduced sample buffer, denatured at 100°C and resolved on 4–12% Bis-Tris SDS-PAGE gels. For Western blot and activity probing resolved proteins were electrotransferred onto nitrocellulose membranes. After blocking, membranes were probed with rabbit pAb anti-6His tag (GenScript, Santa Clara, CA, USA) reacts with tetra-, penta- and hexa-histidine) (Panel B), mAb anti-6His tag (Roche, Indianapolis, IN, USA detects exclusively hexahistidine) (Panel C) and streptavidin-alkaline phosphatase conjugate (Panel D). Blots were developed with AP conjugate Substrate Kit (BioRad, Hercules, CA, USA).

The kinetic constants for substrates turnover by the multidomain Kgp purified from strain HG66 and the catalytic domain alone produced by the ABM1 mutant.

<table>
<thead>
<tr>
<th>Enzymes</th>
<th>Ac-Lys-pNA</th>
<th>Z-Lys-pNA</th>
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<tr>
<td></td>
<td>kₐₑₛ (s⁻¹)</td>
<td>Kₑ (μM)</td>
</tr>
<tr>
<td>Kgp HG66</td>
<td>1.65±0.08</td>
<td>140.1±8.3</td>
</tr>
<tr>
<td>Kgp ABM1</td>
<td>0.99±0.02</td>
<td>98.7±2.2</td>
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n=3; ±SD.

The kinetic constants kₐₑₛ, Kₑ and kₐₑₛ/Kₑ were determined for the N-ε-acetyl-L-Lysine-p-nitroanilide (Ac-Lys-pNA, Bachem) and N-ε-carbobenzoxy-L-Lysine-p-nitroanilide (Z-Lys-pNA, Novabiochem). Kgp (20–40 nm) was incubated in assay buffer (200 mM Tris, 5 mM CaCl₂, 150 mM NaCl, and 0.02% NaN₃, pH 7.6) supplemented with 10 mM L-cysteine. After 10 min at 37°C, chromogenic substrates Ac-Lys-pNA or Z-Lys-pNA were added at various concentrations (0–500 μM) and the release of p-nitroanilide was monitored spectrophotometrically at 405 nm (Spectramax M5 spectrophotometer plate reader, Molecular Devices). kₐₑₛ and Kₑ were calculated by non-linear regression using Graphpad Prism software version 5 (GraphPad Software, La Jolla, USA). The specific activity of Kgp ABM1 and other Kgp forms produced by ABM1 and ABM2 mutants and purified on the analytical scale was practically the same. Therefore, we assumed that substrate hydrolysis by these Kgp variants occurs with the same kinetic constants as those determined for Kgp ABM1.
nanol-ESI-MS/MS analysis of the purified Kgp forms, the aberrant promiscuous proteolysis apparently occurred at a number of sites directly after (Ni-Sepharose bound forms) and within the hexahistidine sequence (Ni-Sepharose unbound forms) (Figure 1D). Generated in this way, C-terminally truncated Kgp variants have molecular weight indistinguishable one from the other in SDS-PAGE but clearly lower than that of wild-type Kgp as proteolysis occurred at least several residues upstream from the native cleavage site in the wild-type protein (Figure 1). Subsequently, the major part of the oligomerization motif was lost and the Kgp catalytic domain was released into the medium together with the truncated or intact profragment. Apparently, the remaining N-terminal fragment of the oligomerization motif GEESP and GEESPYQP in ABM2 and ABM3, respectively, is insufficient to mediate oligomerization of the catalytic domain to the hemagglutinin/adhesin domains. Of note, the mode of association of the Kgp/RgpA complex on the organism. The presence of the conservative C-terminal anchors the remaining domains to the outer membrane of the organism. Nevertheless, it can be hypothesized that glycosylated HA4 oligomerization of the catalytic domain to the hemagglutinin/adhesin domains. Of note, the mode of association of the Kgp/RgpA complex on the surface is unclear.

Purification of the Kgp catalytic domain alone opens the possibility to solve the crystal structure of this gingipain and investigate the role of HA domains in Kgp function as the major virulence factor of P. gingivalis.

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References


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