Identification of a Putative Keratinase Gene and Analysis of a Peptidase S8 Family from a Hyperthermophilic Isolate, *Fervidobacterium* sp. Strain FC2004 in Thailand

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ABSTRACT

*A hyperthermophilic Fervidobacterium sp. strain FC2004 was previously isolated from a hot spring in Thailand. A putative gene encoding a proenzyme named in this study “ProA1” was identified from shotgun sequencing of genomic DNA of the strain FC2004. Predicted 3D-molecule contains a non-homologous signal peptide, a propeptide domain (PD), a catalytic domain (CD) with the typical catalytic triad residues of D169, H207 and S379, and a substrate binding domain (SD). Unlike fervidolysin and islandisin, the ProA1 completely lacks the SD2 domain. Phylogenetic analysis suggests that the ProA1 might be an intermediate isoform between high molecular weight and low molecular weight peptidase S8_subtilases. Although, the strain FC2004 is able to degrade pieces of native feather at high temperature, whether or not the mature ProA1 is active on keratin, the SD plays a role in hydrolyzing keratin substrate and the strain FC2004 might carry a second serine protease with two SDs remains to be investigated.*
**Keywords:** Fervidobacterium, Hyperthermophile, Keratinase, Serine protease, Thermostable enzyme

**INTRODUCTION**

*Fervidobacterium* sp. FC2004, a hyperthermophilic heterotroph belonging to order Thermotogales growing by peptide fermentation was isolated from a hot spring in Northern Thailand. The bacterium grew optimally at 75-80 °C, pH 7.5 and able to degrade native feather at growth temperature (Keawram and Kanoksilapatham, 2013). Proteases or peptide hydrolases can be categorized into four classes; cysteine, serine, metallo and aspartic proteases. Serine proteases consist of a serine residue in conserved catalytic triad which can be classified based on structures to be “chymotrypsin-like” and “subtilisin-like” (or known as “subtilase”). Superfamily of serine proteases has been divided into clans; subtilisin, thermitase, proteinase K, lantibiotic peptidase, kexin and pyrolysin (Siezen and Leunissen, 1997). The clan subtilisin comprises S8 (D, H and S) and S53 (E, D and S) subtilase families. The typical subtilisins are non-specific low molecular weight (20 - 45 kDa) enzymes and named in this study “small S8_subtilase”. Unlike other low molecular weight enzymes, molecular weight of keratinolytic enzymes from thermophilic *Fervidobacterium* spp. are approximately double in size or larger and named in this study “large S8_subtilase”. A cell bound keratinases (or fervidolysin) from a thermophilic *F. pennivorans* was revealed to be a serine protease with a molecular weight of 130 kDa (Friedrich and Antranikian, 1996). A 2.1-kb fls gene encoding fervidolysin, a 699 amino acids prosubtilase was cloned and expressed in *E. coli*. Autoproteolysis of subtilase precursor (73 kDa) was evidenced to a 58 kDa mature enzyme and a 14 kDa propeptide (Kluskens et al., 2002). A homomultimeric membrane bound keratinase (or islandisin)
native enzyme from *F. islandicum* (>200 kDa; 97 kDa subunits) was proved to be active optimally at 80 °C and very thermostable without detectable loss of activity after incubation at 80 °C for > 32 h (Nam et al., 2002; Godde et al., 2005).

In this study, a putative keratin-degrading gene was identified in the genomic DNA of a hyperthermophilic *Fervidobacterium* sp. strain FC2004. This study reports an intermediate size protease that is unique to this strain.

**MATERIALS AND METHODS**

**Organism and cultivation**

*Fervidobacterium* sp. strain FC2004 was employed. This strain is equivalent to strains JMC 18757, ATCC BAA-2483 and *Fervidobacterium thailandense* strain FC2004\(^\text{T}\) (Kanoksilapatham et al., 2016) Cultivation was performed in 480GM5 medium at 80 °C. Composition of 480GM5 medium is similar to the previously described 408G medium except that 5 g/L pancreatic digest of casein was used (Keawram et al., 2016).

**Gene amplification**

An ORF encoding a protease named in this study “ProtA1” was identified from shotgun sequencing of genomic DNA. Two specific primers, ProT1AF (5′ CATA TGCG TAGA CCCG TTAA CGTC 3′) and ProT1AR (5′ TTAC TATA GTTC AACT TCAA TTTG CAC 3′) were designed. PCR product sized of 1569 bp was amplified.

**Sequence similarities and protein structure**

A numbers of peptidase \_S8 family serine protease system are identified in the complete genome sequences of *Fervidobacterium pennivorans* DSM 9078\(^\text{T}\), *F. nodosum* Rt17-B1\(^\text{T}\), *F. islandicum* AW1 and *F. pennivorans* DYC that can be grouped in this study according to sizes (>680 and <450 amino acids) and named “large S8\_subtilase” and “small S8\_subtilase” mentioned previously. All protein sequences were obtained from
NCBI. Two protein ID of large S8_subtilases including WP_104451857.1 (699 aa) and WP_014451869 (697 aa) from *F. pennivorans* DSM 9078$^T$, WP_011994230.1 (682 aa) and WP_011994224.1 (710 aa) from *F. nodosum* Rt17-B1$^T$, WP_052107197.1 (701 aa) and WP_033191969.1 (699 aa) from *F. islandicum* AW-1, and WP_064012406.1 (695 aa) and WP_064012395.1 (688 aa) from *F. pennivorans* DYC were chosen. WP_014451703.1 (439 aa), WP_011993735.1 (440 aa), and WP_033191846.1 (439 aa) representing small S8_subtilase sequences were employed. 3D protein structure was generated using SWISS-MODEL software (http://swissmodel.expasy.org/).

**RESULTS**

**Amplification of keratinase gene**

Shotgun sequencing from genomic DNA of *Fervidobacterium* sp. strain FC2004 revealed an ORF containing a putative preprosubtilase (522 amino acids). Nucleotide sequence is shown in Figure 1. BlastP analysis revealed remarkably high similarity to peptidase S8 from *F. pennivorans* DSM 9078$^T$ (WP_064012406.1; 52 %), *F. islandicum* AW-1 (WP_052107197.1; 48 %) and *F. nodosum* Rt17-B1$^T$ (WP_011994230.1; 48 %) and subtilase from *Thermosipho africanus* (WP_004102466.1; 47%).
Figure 1. Nucleotide and deduced amino acid sequences of a proA1 from *Fervidobacterium* sp. strain FC2004. The ORF begins with a valine codon. Top lines represent nucleotide sequence. Bottom lines represent amino acid sequence. Underline represents primer binding site. Boxes represent catalytic triad amino acids. One letter abbreviation of amino acids is depicted.
3D structure analysis

Large S8_ and small S8_subtilases from *F. pennivorans*, *F. nodosum* and *F. islandicum* and ProA1 precursor were revealed for 3D-models. In general, all the large homologs (682 to 710 aa) including the fervidolysin (1r6v.1.A) contain a propeptide domain (PD), catalytic domain (CD) and two sandwich or substrate binding domains (SD1 and SD2) (Figure 2A, B, C, D, E, F). Unlike the fervidolysin/islandisin, all three small S8_subtilases were revealed lacking both sandwich domain (Figure 2 G, H and I). Interestingly, 3D-model of ProA1 contains an atypical SD lacks completely of the SD2 (Figure 2J).

Figure 2. 3D-models of Peptidase_S8 family serine proteases (A) Model of the Ir6v.1.A template. The template is x-ray crystal structure of the mutant fervidolysin (A200/H200) (Kim et al., 2004). (B) - (F) Models of large S8_subtilases. (G) - (I) Models of small S8_subtilases. (J) Model of ProA1 (522 aa). Symbols: Species and protein IDs are as indicated in the images. Numbers in parenthesis indicate sizes (amino acids). PD represents propeptide domain, CD represents catalytic domain, SD represents sandwich or substrate binding domain.
Phylogenetic tree of S8_subtilase sequences categorized proProA1 within a branch formed by the large S8_homologs from *Fervidobacterium* rather than the small ones (Figure 3). Whether or not this is an intermediate peptidase_S8 family typical for this particular strain of *Fervidobacterium* in Thailand lineage or a truncated mutant of a large S8_homologs in genome of this strain is unclear.

**Figure 3.** Phylogenetic tree of pro1A and other peptidase_S8 family serine proteases from *Fervidobacterium*. Abbreviations: FN, *Fervidobacterium nodosum*; FP, *F. pennivorans*; FI, and *F. islandicum*. 

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Figure 4. Alignment of signal peptide and propeptide sequences. Symbols: yellow represents nonpolar amino acids and F, red represents aspartic and glutamic acids, pink represents glycine, grey represents cysteine, green represents tyrosine, dark green represents polar amino acids (S, T, N, and Q), W and H and blue represents positively charged amino acids (K and R) and P. Dashes indicate spacing generated by alignment process. Numbers represent numbers of amino acids in a particular protein ID.

Multiple alignments of N-terminal segments revealed a non-homologous prepeptide sequence (approximately 20 aa-residues) that was aligned corresponding to the nonpolar amino acid rich signal peptides. PD of ProA1 precursor was contained many conserved stretches of nonpolar amino acids (Figure 4). Moreover, amino acid residues in PD of proProA1 (from P115 to L118) were identified associating with active grooves in CD region. This is an unambiguous characteristic reported on
molecules of many propeptidases including the fervidolysin and the protein ID of WP_014451703.1, a small subtilase (Figure 5). In Figure 5C, catalytic triad residues of D169, H207 and S379 were revealed in CD of the ProA1.

Figure 5. Catalytic triads and binding of C-terminal PD peptides. (A) Fervidolysin (1r6v.1.A) from *F. pennivorans*. (B) A small peptidase S8 serine protease (WP_014451703.1) from *F. pennivorans*. (C) ProA1. D169, H207 and S379 are identified as the catalytic triad.

Alignment of C-terminal sequences of ProA1 (A431 to Y495 segment) to large S8_subtilases suggest a homologous sequence to SD1 and thus named “SD-like” in this study. However, the ProA1 lacks the second SD2 (Figure 6A). Modeling experiment using predicted SD-like region (G422 to Q502) reveals that models constructed from G442 to I498 and N441 to M497 are similar to the partial segment of a collagen adhesion protein (3kpt.1) and fibronectin binding protein (2x5p.1.A).
Superimposed images from the 1r6v.1.A, 3kpt.1 and 2x5p.1.A templates (Figures 6B and 6C) reveal good fit models suggesting a protein-protein binding property of the SD-like. Although the strain FC2004 can degrade native feather, whether or not the mature ProA1 is active on keratin and the SD plays a role in hydrolyzing keratin substrate remained to be investigated.

**Figure 6.** (A) Alignment of C-terminal regions of large S8 subtilases and ProA1. (B) 3D-model of ProA1 constructed using 3kpt.1.A (collagen binding protein) and 2x5p.1.A (fibronectin binding protein like domain). (C) Superimposed images constructed on three templates (1r6v.1.A, 3kpt.1.A and 2x5p.1.A). Symbols: yellow represents nonpolar amino acids (A, V, L, I, M) and F, red represents aspartic and glutamic acids, pink represents glycine, grey represents cysteine, green represents tyrosine, dark green represents polar amino acids (S, T, N,
DISCUSSION & CONCLUSION

An ORF encoding a putative thermostable peptidase S8 family serine protease named in this study “ProA1” is identified in genome of a hyperthermophilic *Fervidobacterium* sp. FC2004 (Figure 1). ProA1 (522 amino acids) contains a signal peptide and a propeptide in molecule suggesting the ProA1 might be a membrane bound or extracellular subtilase precursor (Godde et al., 2005; Kim et al., 2004; Kluskens et al., 2002). Unlike keratinases and other S8_subtilases, mature ProA1 contains a SD-like (Figures 2, 6A). In Figure 3, phylogenetic analysis grouped ProA1 within the branch of fervidolysin-like molecules. The results imply that the ProA1 precursor is more closely related to the large S8_subtilases. Typical catalytic triad residues of D169, H207 and S379 in CD were identified suggesting ProA1 is an S8 family serine protease (Figure 5C). The possible function as substrate binding domain of the SD-like in mature ProA1 is evidenced (Figure 6). Whether or not the proProA1 is a truncated mutant peptidase S8_subtilase or an intermediate homolog (between the large and small S8) remained to be investigated. Comparative mapping of these alleles in the complete genome sequences of *F. pennivorans* DSM 9078^T^ (CP003260.1), *F. nodosum* Rt17-B1^T^ (CP000771.1) and *F. islandicum* AW1 (CP014334.1) might provide a clue. It is likely that strain FC2004 might contain other S8_subtilase homologs with two SDs and without SD in molecules. Although, strain FC2004 has ability to degrade native feather, the SD-like sequence might or might not be related to this phenotype.
In conclusion, the ProA1 of strain FC2004 is unique and have never been locating in the three complete genomes of *Fervidobacterium* (CP003260.1, CP000771.1 and CP014334.1). The enzyme ProA1 has potential applications in leather industries, textile processing and waste treatment from poultry industry. Also feather waste treatment at high temperature by strain FC2004 is considered a green process that possibly prevents contamination by dermatophytic fungi.

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