Maturation of subtilisin-like protease NbSLP1 from microsporidia Nosema bombycis

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Microsporidia are obligate intracellular parasites and possess a unique way of invading hosts, namely germination. Microsporidia are able to infect almost all animal cells by germination. During the process, the polar tube extrudes from the spores within, thus injecting infectious sporoplasm into the host cells. Previous studies indicated that subtilisin-like protease 1 (NbSLP1) of microsporidia Nosema bombycis were located at the polar cap of germinated spores where the polar tube extrusion. We hypothesized that NbSLP1 is an essential player in the germination process. Normally, SLP need to be activated by autoproteolysis under conditions. In this study, we found that the signal peptide of NbSLP1 affected the activation of protease, two self-cleavage sites were involved in NbSLP1 maturation between Ala104Asp105 and Ala124Asp125 respectively. Mutants at catalytic triad of NbSLP1 confirmed the decreasing of autoproteolysis. This study demonstrates that intramolecular proteolysis is required for NbSLP1 maturation. The protease undergoes a series of sequential N-terminal cleavage events to generate the mature enzyme. Like other subtilisin-like enzymes, catalytic triad of NbSLP1 are significant for the self-activation of NbSLP1. In conclusion, clarifying the maturation of NbSLP1 will be valuable for understanding the polar tube ejection mechanism of germination.

KEYWORDS microsporidia, Nosema bombycis, subtilisin-like protease, autoproteolysis, germination
Introduction

Microsporidia, a group of obligate intracellular and unicellular fungi alike (Keeling and Fast, 2002), can infect almost all kinds of invertebrates and vertebrates, including immuno-competent and immuno-deficient people (Didier and Weiss, 2011) as well as economic animals, like silkworm, shrimp and honey bees (Ma et al., 2020). The life cycle of the parasite divides into three parts, namely the infective phase (environmental phase), proliferative phase, and sporogenic phase (Franzen, 2004). In the infective phase, the only extracellular stage, mature spores can be activated by some stimulus, then eject the polar tube and transport infective sporoplasm into host cells, i.e., germination (Han and Weiss, 2017). Germination of microsporidia is a significant infectious process for the parasites, although the mechanism has not been unrevealed up to now.

Previous studies have shown that subtilisin-like protease (SLP) widely exist in microsporidia. In Pararosea locustae, the microsporidia infecting migratory locust, SLP play roles in physiology of the spores instead of host-parasites interaction (Ronnebaumer et al., 2014). SLP of Encephalitozoon cuniculi may be involved in spor development regulation (Ronnebaumer et al., 2006). Three subtilisin homologs, namely NbSLP1, NbSLP2-1, and NbSLP2-2, were found in Nosema bombycis, the first identified microsporium which infects Bombyx mori. NbSLP2 localized around spore walls, and could interact with cytoskeletal protein Nbsquitin2 (Liu et al., 2017), while NbSLP1 showed apical localization features, and was activated after germination (Dang et al., 2013), which may imply the function in germination.

As a biological regulator, protease is involved in all sorts of biological processes in parasites. For SLP, these proteases always act as virulence factors in pathogenic microorganisms, involving in the formation of infective organelle, egress, and invasion in the host cells. In parasites, Toxoplasma and Cryptosporidium share the similar subtilisin SUB1, which acts as a pioneer for the parasite’s infection (Wanyiri et al., 2009; Saouros et al., 2012), while Plasmodium falciparum SUB1 processes membrane-anchored protein MSP1, which interacts with Red Blood Cell spectrin cytoskeleton, playing a role in the egress of merozoite from erythrocytes (Das et al., 2015). Besides, SUB2 and SUB3 play different roles in Plasmodium when infecting host (Barale et al., 1999; Withers-Martinez et al., 2004; Alam et al., 2013). In fungi, Kex2 of Blastomyces albicans is involved in pathogenesis when infecting host cells; SPM1 (subtilisin in Magnaporthe oryzae) is involved in multifunction infection, including infection-related germination, invasion, and growth (Oh et al., 2008; Saioh et al., 2009). Subtilisin in bacteria is a significant virulence effector, which was demonstrated by knock-out of CylA in Enterococcus faecalis and IvaP in Vibrio cholerae (Howell et al., 2019; Tang et al., 2019).

Subtilisin is produced as a large precursor, containing a pro-peptide domain and catalytic domain, in which the pro-peptide domain served as an intramolecular chaperone to ensure the protease fold correctly, and as a temporary inhibitor to inhibit the protease activity (Takagi et al., 2001; Yabuta et al., 2001). In some biological processes, subtilisin is triggered by certain conditions, then autoprocesses to release the pro-peptide domain and become a mature enzyme.

The present research explores self-activation of heterologously-expressed NbSLP1 and identifies the two autoproteolytic sites and three active sites. It is hoped that this research will contribute to a deeper understanding of NbSLP1 maturation and help to explore the polar tube extrusion.

Materials and methods

Plasmid construction

Full length of Nbslp1 (FL-Nbslp1) and a signal peptide deleted fragment of Nbslp1 (Pro-Nbslp1) were amplified by PCR with the genome of Nosema bombycis preserved in our lab. Primers designed based on Nbslp1 sequence from N. bombycis database (https://silkpathdb.swu.edu.cn/) are listed in Table 1. Recombinant plasmid pET30a-FL-Nbslp1 and pET30a-Pro-Nbslp1 were constructed by inserting BamHI and Sall restriction endonuclease digested pET30a vector.

Heterologous expression

Recombinant plasmids pET30a-FL-Nbslp1 or pET30a-Pro-Nbslp1 were transformed into E. coli BL21(DE3). The positive clones were inoculated in 5 mL LB medium containing 50 µg/mL of kanamycin at 37°C for more than 8 h. 300 µL of the cell suspension were transferred into 30 mL culture medium, shaken with 180 rpm at 37°C for 2-3 h, until reaching A600~0.6. Expression of His6-FL-Nbslp1-His6 or His6-Pro-Nbslp1-His6 was induced by adding 0.2 mM isopropyl β-D-thiogalactopyranoside (IPTG) to the culture medium, prior to inoculating at 16°C, 130 rpm for 20-24 h. 5 mL of culture were lysed by ultrasonication. Soluble and inclusion body fractions were separated by centrifugation at 13000 rpm for 10 min. Protein samples were subjected onto the SDS-PAGE and analyzed by Western blot.

Protein Purification

The cultured medium of recombinant bacteria was collected (300 mL of culture medium). The supernatant was then ultrasonically fractured in 20 mL. Binding buffer (2 mM KH₂PO₄, 10 mM Na₂HPO₄, 140 mM NaCl, 3 mM KCl, 20 mM imidazole,
TABLE 1 Primers for Nbslp1 and mutants.

| Primer ID    | Primer sequence (5'-3')                      | Length | Fragment          |
|--------------|----------------------------------------------|--------|-------------------|
| FL-Nbslp1-F  | CGgactecATGAATCTGGAACATCAATCCAAT             | 1476 bp| FL-Nbslp1         |
| FL-Nbslp1-R  | GCgtgacATTTAATTGTTACACTTTCA                 | 1395 bp| Pro-Nbslp         |
| Pro-Nbslp1-F | CGgactecGATAATTATATAGGTAGTTTAAAA             | 1188 bp| Nbslp1-mutA-1     |
| Pro-Nbslp1-R | GCgtgacATTATTGTTACACTTTCA                  | 245 bp |                   |
| Nbslp1-MRIA104-F | AAGCAAAAGTGCAGCTGTTACATTTTTAT             | 1182 bp| Nbslp1-mutA-2     |
| Nbslp1-MRIA104-R | TAAAATTTAAAATACACGACATCTTGTTCT           | 253 bp | Nbslp1-mutB       |
| Nbslp1-DFNL108-F | AGCTTGTTACAGCTGAGTTACATTT             | 308 bp |                   |
| Nbslp1-DFNL108-R | GACAGCTGAGTTACATTTGTC                  | 917 bp | Nbslp1-D192A      |
| Nbslp1-D192A-F | AGTTTTAGTTATACACCACG                    | 497 bp |                   |
| Nbslp1-D192A-R | TGATACCTGTTACGAAAC                      | 818 bp | Nbslp1-H224A      |
| Nbslp1-H224A-F | ATGCAAAAGGAGCCAGGAACT                    | 600 bp |                   |
| Nbslp1-H224A-R | AGCTTGTTACAGCTGAGTTACATTT             | 247 bp | Nbslp1-S413A      |
| Nbslp1-S413A-F | AGCTTGTTACAGCTGAGTTACATTT             | 1116 bp| ProFound Lysis Buffer | incubated on ice for 30 min, then centrifuged to clarify crude lysate. 50 μL immobilized cobalt chelate into Handee™ Spin Column was washed 5 times with wash solution (1:1 wash solution of TBS : ProFound™ Lysis Buffer to which 4 M Imidazole Stock Solution was added to a final concentration of 40 mM imidazole). Polyhistidine-tagged fusion protein lysates were incubated with settled cobalt chelate resin at 4°C for 2 h with a gentle rocking motion on a rotary mixer. After 5 washes, 250 μL of 290 mM Imidazole Elution Buffer (63 μL of 4 M Imidazole Stock Solution added to 937 μL of wash solution per 1 mL. Elution Buffer) was added into each spin column, then incubated at 4°C for 5 min, and finally elution was collected by centrifugation at 1250 rpm for 30 sec.

**Western blot**

Protein samples were loaded on SDS-PAGE gels and transferred to PVDF membranes with Tris-Gly Transfer Buffer. The membranes were blocked with 5% skim milk in TBST buffer at room temperature for 1 h, and then incubated overnight at 4°C with primary antibody and anti-His antibody (Roche, 1:1000 dilution in blocking solution). After 3 washes with TBST buffer, the membranes were incubated for 1 h with Peroxidase Conjugated Goat anti-Mouse IgG, (H+L) (Sigma-Aldrich, 1:5000 dilution). The blots were detected with ECL western blot detection kit (Thermo Fisher).

**His-tag pull-down assay**

To enable us to identify the protease and digestion site after self-shearing, His pull-down was performed following the instructions (ThermoFisher Scientific) to purify the recombinant proteins. After washing with TBS, E. coli pellets that express the recombinant proteins were suspended in 1 mL of ice-cold TBS and 1 mL of ProFound™ Lysis Buffer and incubated on ice for 30 min, then centrifuged to clarify crude lysate. 50 μL immobilized cobalt chelate into Handee™ Spin Column was washed 5 times with wash solution (1:1 wash solution of TBS : ProFound™ Lysis Buffer to which 4 M Imidazole Stock Solution was added to a final concentration of 40 mM imidazole). Polyhistidine-tagged fusion protein lysates were incubated with settled cobalt chelate resin at 4°C for 2 h with a gentle rocking motion on a rotary mixer. After 5 washes, 250 μL of 290 mM Imidazole Elution Buffer (63 μL of 4 M Imidazole Stock Solution added to 937 μL of wash solution per 1 mL. Elution Buffer) was added into each spin column, then incubated at 4°C for 5 min, and finally elution was collected by centrifugation at 1250 rpm for 30 sec.

**Mass spectrometry analysis**

Protein bands captured from Pull-down assay were collected in 1.5 mL centrifuge tubes. After reduction and alkylation treatment, Trypsin (mass ratio: 1:50) was added to the test samples for enzymatic hydrolysis at 37 conditions for 20 h. The enzymatic hydrolysis samples were desalted and lyophilized, then dissolved in 0.1%FA solution and stored at -20°C for later use. Mass spectrometry analysis: Solution A was an aqueous solution of 0.1% formic acid, and Solution B was an acetonitrile aqueous solution of 0.1% formic acid (84% acetonitrile). After the chromatographic column was balanced with 95% of Solution A, the sample was loaded from the automatic sampler to TRAP column. Mass spectral data collection: the mass charge ratio of polypeptides and polypeptide fragments was collected by the following methods: 20 fragment maps were collected after each full scan (MS2 scan); Mascot 2.2 software was used to retrieve the corresponding database for the original file of mass
spectrometry test; and finally the identified protein result was obtained. The search database parameters are listed in Table 2.

N-terminal sequence analysis

After being pre-run for 1 hour, protein captured from Pull-down assay was subjected onto an SDS-PAGE gel and then transferred to PVDF membrane (Roche) using CAPS Buffer (10 mM CAPS, pH 11). Target protein band was visualized with Coomassie Bright Blue R-250, and then cut off from the membrane and tested by the PPSQ Protein Sequencer (SHIMADZU). Raw data and chromatogram were analyzed by PPSQ DataProcessing/Labsolutions.

Enzymatic activity

To assess the enzymatic activity of purified recombinant protein, In-gel zymography was conducted (Lantz and Ciborowski, 1994; Seawell and Bose, 2021). Subtilisin purchased from Sigma (Sigma) was used as positive control. Protease samples in nonreducing sample buffer (2.5% SDS) was prepared without boiling. Electrophoresis of the samples on 12% SDS-polyacrylamide gel containing casein (1 mg/mL) was carried out on ice. After 2 washes with 2.5% TritonX-100 diluted in distilled water and 2 washes with 2.5% TritonX-100 diluted in TB buffer (50 mM Tris-HCl buffer, pH 7.4) for 10 minutes respectively, the gel was washed with TB buffer to remove SDS and TritonX-100. After incubation for 24-48 hours at 37 °C in TB buffer containing 10 mM CaCl$_2$, then the gel was cut off from the membrane and stained by Coomassie blue R-250 for 30 minutes. Regions with enzyme activity appear as a clear colorless region against the blue background.

Mutants

To conform the two autocleavage sites which identified recombinant protein, NbSLP1 mutants were designed. Before the experiments in the lab, homology modelling of NbSLP1 and its mutants was conducted to confirm the structures of the proteins with little variation. The target amino acid sequences were submitted in Swiss-Model (https://swissmodel.expasy.org/). With Pro-subtilisin E (PDB: 3whi.1) being the template of NbSLP1 and its mutants, 3D protein models were automatically generated and used to analyze the variation in position. After that, NbSLP1 mutants (NbSLP1mut) were generated by PCR site-directed mutagenesis. The strategy of mutation is shown in Figure 4A and primers are listed in Table 1. Briefly, NbSLP1mut were generated by overlap extension PCR with mutant fragments as templates. Then recombinant plasmids pET30a-NbSLP1mut were constructed by the same methods as above.

Results

Identification of NbSLP1 autoproteolysis and cleavage sites

In order to identify the autoproteolysis of NbSLP1, pET30-Pro-NbSLP1 and pET30-FL-NbSLP1 expression vector with a N-terminal His$_6$-tag and a C-terminal His$_6$-tag were constructed and transformed into E. coli BL21 (DE3) respectively (Figure 1A). Western Blotting analysis showed that His$_6$-FL-NbSLP1-His$_6$ was expressed and purified in soluble form with a molecular weight of ~61.83 kDa, indicated by a black arrow and light band at ~44.55 kDa, indicated by a red arrow, while His$_6$-Pro-NbSLP1-His$_6$ was expressed and purified in soluble form with a molecular weight of ~44.55 kDa, indicated by a red arrow, while inclusion bodies formed with two bands with a molecular weight of ~58.77 kDa and ~44.55 kDa (Figures 1B–D), which share a similar size with predicted Pro-NbSLP1 and mature-NbSLP1, respectively. To further affirm autoproteolysis of NbSLP1 in the prokaryotic expression system, pull-down assay was carried out to obtain putative His$_6$-inhibitor$_{I9}$ and mature-NbSLP1-His$_6$ (Figures 2A, B), then they were identified by mass spectrum. The results showed that the two bands at ~44.55 kDa were both identified as Subtilisin-like serine protease, NbSLP1, which means that NbSLP1 could autoproteolysis twice in E. coli BL21.

In order to assess the enzymatic activity of the purified NbSLP1, in-gel zymography was conducted. The results showed that Pro-NbSLP1 and FL-NbSLP1 had no clear colorless bands, indicating no enzymatic activity of NbSLP1 or that the activity was too low for detection by this method (Figure 2C). Gel filtration chromatography (HiLoad™ 16/600 Superdex™ 200 pg purchased from GE Healthcare) was adopted to separate mature-NbSLP1-His$_6$ from mixture with His$_6$-Inhibitor$_{I9}$, but the results showed that it failed to remove His$_6$-Inhibitor$_{I9}$ even in high-resolution fractionation (Figure S1), which implied that Inhibitor$_{I9}$ may bind to mature NbSLP1 in a non-covalent manner, providing a reasonable clue about low enzymatic activity.

N-terminal sequence of mature-NbSLP1 was conducted to confirm autoproteolysis sites of NbSLP1 in E. coli. The sequencing results showed that N-terminus of mature NbSLP1 and NbSLP1mut were both identiﬁed as Subtilisin-like serine protease, NbSLP1, which means that NbSLP1 could autoproteolysis twice in E. coli BL21.

TABLE 2 The retrieval parameter table of database.

| Parameters           | condition                        |
|----------------------|----------------------------------|
| Enzyme               | Trypsin                          |
| Database             | uniprot_Noemoa_bambuscis_4624_20191014.fasta |
| Fixed modification   | Carbamidomethyl (C)              |
| Variable modification| Oxidation (M)                    |
| Missed Cleavage      | 2                                |
| Peptide Mass Tolerance| 20 ppm                          |
| Fragment Mass Tolerance| 0.1 Da                          |
| Filter by score      | >=20                             |
is NH₂-Asp-Phe-Asn-Leu-Ser-Asp-Tyr-Arg-Lys-Lys (DFNLSDYRKK), demonstrating that the cleavage sites is MRIA₁₀⁴↓D₁₀⁵FNL. To further explore autoproteolysis of NbSLP1 in E. coli, pellets at different timepoints post induction by IPTG were collected and sampled to identify autoproteolysis. We found that Pro-NbSLP1 cleaved itself as soon as the protease was detected, 2 hours post induction (Figure 2D). According to homologous modeling of NbSLP1 against subtilisin E (3whl.2.A) with SWISS MODEL, cleavage sites were around catalytic triad D₁₉₂H₂₂₄S₄¹₃ in the predicted 3D structure (Figure 3). Taken together, these imply autoproteolysis is influenced by catalytic function of the enzyme.

**Mutation in cleavage sites inhibiting autoproteolysis of NbSLP1**

To further demonstrate autoproteolysis sites of NbSLP1, the expression vector pET-30a containing the site-directed mutagenesis in cleavage sites (M₁₀¹RIADFNL₁₀₈ replaced with A₁₀¹AAAAAA₁₀₈ named NbSLP1-mut-A) was constructed, and expressed in E. coli BL21, while the Western Blotting analysis showed that no effect was observed for the cleavage (Figure 4).

Due to a similar sequence of mature-SLP1 N-terminus found as M₁₋₁₂₃ADFNLSD₁₋₁₃₀, combined with the purification results, we suspect that SLP1 may cleave itself more than once. Therefore, another two recombinant plasmids with mutagenesis at sites 1₂₅₋₁₂₈ (DFNL) to A₁₂₅AAA₁₂₈, named E T ₃₀ - NbSLP1-mut-B, and both sites 1₀₁₋₁₀₈ and 1₂₅₋₁₂₈ to A (pET30-NbSLP1-mut-AB) were constructed. Western blotting results showed that autoproteolysis of NbSLP1 was inhibited by mutagenesis at two putative cleavage sites, while the homologous model predicted by SWISS MODEL showed no significant shift happened (Figure 4).

**Mutation in catalytic triad sites inhibiting autoproteolysis of NbSLP1**

Mutations in catalytic triad of NbSLP1 were constructed to verify that autoproteolysis of NbSLP1 was induced by catalytic function. NbSLP1 sequence analysis has been performed before, showing that conserved catalytic triad sites of NbSLP1 are D₁₉₂, H₂₂₄ and S₄¹₃, respectively. Homologous models of these mutants predicted by SWISS-MODEL were compared with the model of NbSLP1, showing that there were some changes at catalytic triad. The mutant proteins were expressed at soluble form. Western
Identification of Pro-NbSLP1 autoproteolysis in E. coli. (A) SDS-PAGE analysis of Pro-NbSLP1 by pull-down assay. Lane M in (A) and (C): premixed protein marker; Lane M in (B) and (D): ECL marker; Lane I: crude lysate extracted from induced E. coli BL21 (DE3) transformed with pET50-Pro-Nbslp1; Lane L: Flow-through after combination with cobalt chelate resin; Lane E: Elution of Pro-NbSLP1 by Pull-down assay; Lane S8: NbSLP1-Peptidase_S8 peptide; Pro-NbSLP1 are indicated by black arrow; mature NbSLP1 are indicated by red arrows and Pro-peptide/I9 inhibitor are indicated by blue arrows. (B) Western blot analysis of Pro-NbSLP1 by pull-down assay. (C) Identification of enzymatic activity by casein zymography. Subtilisin was purchased from Sigma as positive control. Pro-NbSLP1 and FL-NbSLP1 adding PMSF (phenylmethylsulfonyl fluoride) were as inhibitor control. (D) Pro-NbSLP1 autoproteolysis at different hours post induction by IPTG.

MS analysis and autoproteolysis sites in sequence and predicted 3D structure of NbSLP1. (A) Protein sequence of Full-length NbSLP1, peptides of ~14 kDa (band in Figure 2A indicated by blue arrow) detected by MS are showed in red font, peptides of ~45 kDa (band in Figure 2A indicated by red arrow) highlighted in yellow, N-terminus sequence of mature NbSLP1 is showed in blue frame; (B) Predicted 3D structure of NbSLP1. Pro-peptide is showed in red ribbon; mature NbSLP1 is showed in yellow ribbon; autoproteolysis site is showed between red and blue ribbon; NbSLP1 catalytic triad Asp192-His224-Ser413 is showed in ball and stick structure.
blotting analysis showed that no cleavage was observed in D193A and H224A mutants, while S413A mutant showed decreased autoproteolysis (Figure 5). Among the three catalytic triads, serine nucleophilic attacks the substrate, which is crucial for catalytic activity. The S413A mutant partially maintained the autoproteolytic ability, demonstrating that catalytic function was the key factor of maturation, but S413A can be cleaved through a mechanism other than autoproteolysis.
**Discussion**

**NbSLP1 activated after germination hints at the role in this process**

Microsporidia have a unique infective mode as pathogens. However, the molecular mechanisms of spore germination have not been revealed yet. Subtilisin-like protease 1 of *Nosema bombycis* (NbSLP1) can be activated by some conditions after germination, processing itself to release mature-SLP1. Dang *et al.* found that NbSLP1 may have different forms before and after spore germination by dimensional electrophoresis (Dang *et al.*, 2013), implying the role of NbSLP1 in the germination process, while our results shows that NbSLP1 without signal peptide

**FIGURE 5**

Autoproteolysis analysis of Pro-NbSLP1 mutants of active sites and predicted 3D structure. (A) Western blot analysis of Pro-NbSLP1 mutants in active sites. Supernatant from induced *E. coli* BL21 (DE3) [pET30-pro-Nbslp1] (WT), *E. coli* BL21 (DE3) [pET30-pro-Nbslp1-D193A], *E. coli* BL21 (DE3) [pET30-pro-Nbslp1-H224A], *E. coli* BL21 (DE3) [pET30-pro-Nbslp1-S413A] were analyzed by immunoblotting using His antibody. Details were showed below. Predicted 3D structure of NbSLP1 and its mutants against Pro-subtilisin E are showed in B–E; unmutated NbSLP1 is showed in red ribbon (B), NbSLP1-D193A is showed in green ribbon (C), NbSLP1-H224A is showed in blue ribbon (D), NbSLP1-S413A is showed in yellow ribbon (E), catalytic triads were showed in bottom right of each structure.
heterologously expressed in *E. coli* can autoprocess itself without any external factor, sharing a similar expression feature with mirolase, the subtilisin-like protease in *Tamberelli forsythia* (Ksiazek et al., 2015), whereas full-length NbSLP1 can be expressed as precursor without any other forms, meaning that signal peptide of NbSLP1 or other factors in *N. bombycis* may work in the enzyme activation during spore germination.

**Activation of NbSLP1 may be related to some stimulus**

NbSLP1 localizes in mature spores mostly as a zymogen. We know that activation of zymogen is always related to some environmental factors, such as pH, temperature, and irons. For instance, Furin, the homolog of bacterial subtilisin in mammals, is activated by the pH-dependent mechanism, in which His69 in furin propeptide served as pH sensor (Feliciangeli et al., 2006). Inactive furin undergoes auto-cleavage twice, at Arg107 and Arg75, to cleave and release propeptide, which occurs in the endoplasmic reticulum (ER) and mildly acidic environments (trans-Golgi network), respectively (Anderson et al., 1997; Dillon et al., 2012). Considering the process of expression of this heterogeneous NbSLP1 in *E. coli* cells, there were no such factors that might contribute to activation of NbSLP1. It is reasonable to infer that the cleavage of the enzyme happened spontaneously (auto-cleavage). However, while there was no visible enzymatic activity detected after purification, extra factors might be needed for the full maturation of the protease in spores, such as post-translation regulation (PTM) (Dewpura et al., 2008) and other protein regulations. Meanwhile, there also some stimulus or stability factors related to spore germination, environmental factors (Leitch and Ceballos, 2008), and some proteins like aquaporins (Ghosh et al., 2006) or Septin 2 (Liu et al., 2020), which can also be considered as potential associators of NbSLP1.

**Different forms of mature NbSLP1 may imply various functions**

In this study, we focused on the autoproteolysis sites of NbSLP1, and found the cleavage sites at Asp105 and Asp125, which means at least two forms of mature NbSLP1 expressed *in vitro*, compared with three potential forms of protease after germination *in vivo* (Dang et al., 2013), which demonstrated the variants of NbSLP1. In the process of NbSLP1 autoproteolysis, we speculated that NbSLP1 can cleave at Asp105 and Asp125.
simultaneously, or otherwise the form cleaved at Asp\textsuperscript{105} can promote the second cleavage (Figure 6).

Cleavage sites at different locations of the amino acid sequence have been reported in caspase and other subtilisin (Gauthier et al., 2015). Sometimes, different forms of certain protease may lead to other roles. In Caspase-4, there are three cleavage sites at Asp\textsuperscript{59}, Asp\textsuperscript{270}, and Asp\textsuperscript{289}, leading to four forms, named p31, p24, p12 and p10, in which only the p33/p10 form is critical for protease activation to induce pyroptosis, while other forms are of little significance in this process (Wang et al., 2020). In view of localization signal and expression pattern of NbSLP1, the protease may work on various substrates in parasites and/or host cells.

**NbSLP1 with propeptide provide foundation for active NbSLP1**

Previous studies about NbSLP1 show that it is difficult to produce soluble mature NbSLP1 by the E. coli expression system, yeast expression system, and even baculovirus expression vector system. There is some research related to active protease preparation with a subtilisin with its own propeptide (Xie et al., 2019). In this study, we constructed a NbSLP1 vector with its own propeptide to produce the soluble protein. This is because inhibitor_19 of SLP1 promotes proper folding of mature enzyme, while the pro-peptide also keeps the proenzyme inactive as a temporal inhibitor of SLP (Yabuta et al., 2001). However, the soluble mature NbSLP1 we prepared showed low catalytic activity, which was caused by the rate-limiting process of propeptide released from inactive propeptide-NbSLP1 complex. We are planning to explore the substrate of NbSLP1 to further identify the role of this protease in the parasites spore germination.

In this study, we identified the autoproteolysis and cleavage sites of recombinant Pro-NbSLP1 in position Ala\textsuperscript{104} and Ala\textsuperscript{124}, while full-length NbSLP1 displayed a low cleavage rate. We found that catalytic triad (D\textsuperscript{193}H\textsuperscript{224}S\textsuperscript{413}) and cleavage sites were both significant to NbSLP1 activation, provided that the 3D structure of NbSLP1 and its mutants are less differentiating at some level.

**Data availability statement**

The original contributions presented in the study are included in the article. Further inquiries can be directed to the corresponding authors.

**Author contributions**

RW and JC designed the project, performed the experiments, analyzed the data, and drafted the manuscript. RW and QL performed most of the experiments. XD, FL, QS, and XS gave input on method refinement. MH performed part of the plasmid construction. B JL review and edit the manuscript. GP and ZZ provided overall study supervision and grant support. All authors read and approved the final manuscript.

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**Conflict of interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2022.897509/full#supplementary-material

**SUPPLEMENTARY FIGURE 1**

Gel filtration analysis of purified NbSLP1. (A) Elution profile of protein samples taken from an absorbance of 280 nm. (B) SDS-PAGE analysis of protein samples taken from peak 1, 2 and 3. Mature NbSLP1 are indicated by red arrows and Pro-peptide (Inhibitor_19 domain) is indicated by blue arrow.
References

Alam, A., Bhatnagar, R. K., Relan, U., Mukherjee, P., and Chaunhan, V. S. (2013). Proteolytic activity of plasmodium falciparum subtilisin-like protease 3 on parasite profilin, a multifunctional protein. *Mol. Biochem. Parasitol.* 191 (2), 58–62. doi: 10.1016/j.molbiopara.2013.09.006

Anderson, E. D., VanZyle, J. K., Thulin, C. D., Jean, F., and Thomas, G. (1997). Activation of the furin endoprotease is a multiple-step process: requirements for acidification and internal propeptide cleavage. *EMBO J.* 16 (7), 1508–1518. doi: 10.1093/emboj/16.7.1508

Barale, J. C., Blasnick, T., Fujisaka, H., Altar, P. M., Aikawa, M., Braun Breton, C., et al. (1999). Plasmodium falciparum subtilisin-like protease 2, a merozoite candidate for the meronzoite surface protein 1-42 maturase. *Proc. Natl. Acad. Sci. U.S.A.* 96 (11), 6445–6450. doi: 10.1073/pnas.96.11.6445

Dang, X., Pan, G., Li, T., Lin, L., Ma, Q., Gong, L., et al. (2013). Characterization of a subtilisin-like protease with apical localization from microsporidian *nosema bombycis*. *J. Invertebr. Pathol.* 112 (2), 166–174. doi: 10.1016/j.jip.2012.10.009

Das, S., Hertrich, N., Perrin, A. J., Withers-Martinez, C., Collins, C. R., Jones, M. L., et al. (2015). Processing of plasmodium falciparum meronzoite surface protein MSP1 activates a spectrin-binding function enabling parasite egress from RBCs. *Cell Host Microbe* 18 (4), 433–444. doi: 10.1016/j.chom.2015.09.007

Dewpura, T., Raymond, A., Hamelin, J., Seidah, N. G., Mbikay, M., Chretien, M., et al. (2008). PCSK9 is phosphorylated by a golgi casein kinase-like kinase ex vivo and circulates as a phosphoprotein in humans. *FEBS J.* 275 (3), 3489–3493. doi: 10.1111/j.1742-4689.2008.06495.x

Dikker, E. S., and Weiss, L. M. (2011). Microsporidiosis: not just in AIDS patients. *Curr. Opin. Infect. Dis.* 24 (5), 490–495. doi: 10.1515/hsz-2014-0256

Dillon, S. L., Williamson, D. M., Efferich, J., Radler, D., Joshi, R., Thomas, G., et al. (2012). Propeptides are sufficient to regulate organelle-specific pH-dependent activation of furin and proprotein convertase 1/3. *J. Mol. Biol.* 423 (1), 47–62. doi: 10.1016/j.jmb.2012.06.023

Feliciangeli, S. F., Thomas, L., Scott, G. K., Subbian, E., Hung, C. H., Molloy, S. S., et al. (2006). Identification of a pH sensor in the furin propeptide that regulates enzyme activation. *J. Biol. Chem.* 281 (23), 16108–16116. doi: 10.1074/jbc.M600760200

Franzen, C. (2004). Microsporidia: how can they invade other cells? *Trends Parasitol.* 20 (6), 275–279. doi: 10.1016/j.pt.2004.04.009

Gauthier, M. S., Persutte, J. R., Awan, Z., Bouchard, A., Tessier, S., Champagne, J., et al. (2015). A semi-automated mass spectrometric immunoassay coupled to selected reaction monitoring (MSRA-SRM) reveals novel relationships between circulating PCSK9 and metabolic phenotypes in patient cohorts. *Methods* 81, 66–73. doi: 10.1016/j.ymeth.2015.03.003

Ghosh, K., Cappiello, C. D., McBride, S. M., Occi, J. L., Cali, A., Tavkorian, P. M., et al. (2006). Functional characterization of a putative aprotinin from *encephalitozoon cuniculi*, a microsporidia pathogenic to humans. *Int. J. Parasitol.* 36 (1), 57–62. doi: 10.1016/j.ijpara.2005.08.013

Han, B., and Weiss, L. M. (2017). Microsporidia: Obligate intracellular pathogens within the fungal kingdom. *Microbiol. Spectr.* 5 (2). 10.1128/microbiolspec.MUSP-0018-2016. doi: 10.1128/microbiolspec.MUSP-0018-2016

Howell, M., Dumitrescu, D. G., Blankenship, L. R., Herkert, D., and Hatzios, S. (2004). Subtilisin-like protease in the microsporidium paranosema locustae. *Parazitologiia* 48 (5), 337–347.

Wang, K., Sun, Q., Zhong, X., Zeng, M., Zeng, H., Shi, X., et al. (2020). Structural mechanism for GSDMD targeting by autoprocessed caspases in pyroptosis. *Cell* 180 (5), 941–955.e920. doi: 10.1016/j.cell.2020.02.002

Wanyiri, J. W., Techasintana, P., O’Connor, R. M., Blackman, M. J., Kim, K., and Ward, H. D. (2009). Role of CgSUBL, a subtilisin-like protease, in cryptosporidium parvum infection in vitro. *Eukaryot. Cell* 8 (4), 470–477. doi: 10.1128/EC.00306-08

Withers-Martinez, C., Jean, L., and Blackman, M. J. (2004). Subtilisin-like proteases of the malaria parasite. *Mol. Microbiol.* 53 (1), 55–63. doi: 10.1111/j.1365-2958.2004.04144.x

Xie, G., Shao, Z., Zeng, L., Li, X., Cong, D., and Huo, R. (2019). Heterologous expression and characterization of a novel subtilisin-like protease from a thermophilic thermus thermophilus HB8. *Int. J. Biol. Macromol.* 138, 528–535. doi: 10.1016/j.ijbiomac.2019.07.101

Yatabu, Y., Takagi, H., Inouye, M., and Shinde, U. (2001). Folding pathway mediated by an intramolecular chaperone: propeptide release modulates activation precision of pro-subtilisin. *J. Biol. Chem.* 276 (48), 44427–44434. doi: 10.1074/jbc.M107573200