A Novel Spore Wall Protein from *Antonospora locustae* (Microsporidia: Nosematidae) Contributes to Sporulation

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**ABSTRACT**

Microsporidia are obligate intracellular parasites, existing in a wide variety of animal hosts. Here, we reported AlocSWP2, a novel protein identified from the spore wall of *Antonospora locustae* (formerly, *Nosema locustae*, and synonym, *Paranosema locustae*), containing four cysteines that are conserved among the homologues of several Microsporian pathogens in insects and mammals. AlocSWP2 was detected in the wall of mature spores via indirect immunofluorescence assay. In addition, immunocytochemistry localization experiments showed that the protein was observed in the wall of sporoblasts, sporonts, and meronts during sporulation within the host body, also in the wall of mature spores. AlocSWP2 was not detected in the fat body of infected locust until the 9th day after inoculating spores via RT-PCR experiments. Furthermore, the survival percentage of infected locusts injected with dsRNA of AlocSWP2 on the 15th, 16th, and 17th days after inoculation with microsporian were significantly higher than those of infected locusts without dsRNA treatment. Conversely, the amount of spores in locusts infected with *A. locustae* after treated with RNAi AlocSWP2 was significantly lower than those of infected locusts without RNAi of this gene. This novel spore wall protein from *A. locustae* may be involved in sporulation, thus contributing to host mortality.

MICROSPORIDIA are intracellular parasites of all major Insecta and Mammalia classes and have a described diversity of over 1,400 species, although their actual diversity is estimated to be much higher (Szumowski and Troemel 2015; Williams 2009). Identification of the molecular mechanisms for pathogenicity of Microsporidia to its hosts is increasing remarkably with the progress of microsporian genome sequencing. Although Microsporidia differ greatly in host range and cell type specificity, they share a similar mechanism for host cell invasion (Franzen 2005; Franzen et al. 2005). Infection involves the rapid expulsion of a polar tube from a dormant spore that pierces the host cell membrane and allows the direct transfer of the spore contents into the host cell cytoplasm (Williams 2009). A general viewpoint is that the spore wall is involved not only in this initial process, but also plays central roles in the complex interactions between Microsporidia and its host cells: including adherence, invasion, infection, and pathogenicity. Adherence of the microsporidian spores to host cells is the first step in the infection process (Southern et al. 2007) that occurs when spore wall proteins bind to sulfated glycosaminoglycans (GAGs) on the host cell surface (Hayman et al. 2005).

The microsporidian spore wall proteins identified so far, have very low or no sequence similarity with any other eukaryotic proteins: SWP1, SWP2, and EnP1 from *Encephalitozoon intestinalis* (Hayman et al. 2001; Southern et al. 2007); EnP1 from *Encephalitozoon ronalaee* (Pom-bert et al. 2012); EcSWP1, EcCDA, EnP1, EnP2, and SWP3 from *Encephalitozoon cuniculi* (Brosson et al. 2005; Peuvel-Fanget et al. 2006; Taupin et al. 2006; Xu et al. 2006); Ehswp1a and Ehswp1b from *Encephalitozoon hel-lern* (Polonais et al. 2010); as well as NbSWP7, NbSWP9, NbSWV16, NbHSWP11, SWP5, NbSWP12, SWP25, SWP26, SWP30, and SWP32 from *Nosema bombycis* (Chen et al. 2013; Li et al. 2009, 2012; Wang et al. 2015; Wu et al. 2008, 2009; Yang et al. 2014, 2015, 2017). All of these spore wall proteins can be localized to the exospore, endospore, or plasmalemma, and nearly all of these have functional binding sites similar to the heparin-binding motif (HBM), or modifications such as phosphorylation and glycosylation.
**Antonospora locustae** is an important pathogen, which has been commercialized and widely used for locust and grasshopper control (Brooks 1988; Henry 1971). Based on both molecular and morphological evidence, a change in the generic name of *Nosema locustae* to the genus *Antonospora* (*Paranosema*), as *A. locustae* n. comb. has been proposed (Slamovits et al. 2004; Sokolova et al. 2003). In particular, Microsporidia-specific proteins such as spore wall proteins and polar tube proteins have received further attention (Dolgikh et al. 2005; Polonais et al. 2013). The close relative of *A. locustae*, *Paranosema grylli* was identified to have one spore-wall protein via selective extraction of a major 40 kDa protein (Dolgikh et al. 2005). The spore-wall and polar-tube proteins are transported from the endoplasmic reticulum to the target membranes through these tubular networks (Beznoussenko et al. 2007). In aspects of host-parasite interactions, the localization of hexokinase secreted by *A. locustae* into infected host cells suggests that some of Microsporidia possess a broad set of enzymes and regulatory proteins that have the potential to alter metabolic processes and molecular programs of the host (Senderskyi et al. 2014; Timofeev et al. 2016). However, little is known about the molecular pathogenicity of *A. locustae*, a potentially intriguing model system for understanding the extremes of reductive parasite evolution and host cell manipulation (Williams 2009).

In this paper, we have identified a putative spore wall protein of *A. locustae* via MALDI-TOF mass spectrometry. Indirect immunofluorescence and immunoechemistry localization experiments showed that this protein was localized in the spore wall. Furthermore, RNAi treatment against AlocSWP2 indicated that this protein was involved in sporulation, thus contributing to host mortality.

**MATERIALS AND METHODS**

**Microsporidia and insects**

*Antonospora locustae* spores were provided by the Kay Lab for Biocontrol of the Ministry of Agriculture of China, China Agricultural University, and were purified from its host locust (*Locusta migratoria*) in the laboratory. *Antonospora locustae* spores were purified from infected locust abdomens on a discontinuous Percoll gradient (25%, 50%, 75%, and 100%, v/v) centrifuged at 14,000 g for 20 min. Then, spores were washed with ultrapure water at least thrice. The purified spores were stored at −20 °C until further use (Gatehouse and Malone 1998).

Locusts were raised in our department at 28–30 °C, a relative humidity of 60%, and a photoperiod of 18:6 h light:dark. Fresh corn leaves were provided daily. To infect the locusts, locusts at 2nd or 3rd day of the 3rd instar were selected and starved for 4 h, then fed with 10^7 purified spores of *A. locustae* on 20 mm × 5 mm corn leaves, then reared identical to healthy controls. The leaves should be completely consumed within 12 h, and locusts, which failed to do so, were not used in the vexperiment.

**Protein extraction, gel electrophoresis, and MALDI-TOF MS assay**

A small amount of the poorly soluble fraction of protein from *A. locustae* was extracted as follows. Briefly, using the Brosson method (Brosson et al. 2006), spores were disrupted in 200 μl of SDS extraction buffer, containing 100 mM DTT, 4% CHAPS and 0.2% SDS, by repeated cycles of freezing-thawing and sonication (Scientz-IIE, 300 W, 20–25 kHz) (“SDS extract”). The proteins from broken cells were extracted with a solution containing 7 M urea, 2 M thiourea, 100 mM DTT, 4% CHAPS, and 0.2% SDS for 6 h at room temperature. After clarification via centrifugation (12,000 g, 5 min), the supernatant or “urea extract” was collected and the sediment material treated with 30 mM NaOH overnight at 47 °C before centrifugation (as before) to collect the “NaOH extract” supernatant. The NaOH-insoluble material, containing a small amount of spore proteins, was finally boiled for 10 min in Laemmli solution containing 2.5% SDS, 0.125 M Tris-HCl pH 6.8, 20% glycerol, 2 mM EDTA, and 100 mM DTT (“Laemmli extract”).

Two-dimensional electrophoresis (2-DE) analysis was done as follows. Isoelectric focusing (IEF) was performed using linear immobilized pH gradient strips of 17 cm pH 3–10 (Bio-Rad, Berkeley, CA) in rehydration buffer (7 M urea, 2 M thiourea, 4% CHAPS, 2 mM tributylphosphine solution, and 0.5% ampholytes) using the IPGphor apparatus (GE). After SDS-PAGE on 12% polyacrylamide gels (18*18 cm), the strips were equilibrated with 50 mM Tris HCl pH 8.8, 6 M urea, 30% glycerol, 2% SDS, and 100 mM DTT, and then 135 mM iodoacetamide. After 2-DE separation, the protein spots were manually excised from Coomassie brilliant blue (CBB) stained 2-DE gels and transferred to a tube. The 2-DE gel analysis was performed using the PDQuest 6.2.1 software. The protein spot in the gel was excised and digested by trypsin and submitted to for commercial MALDI-TOF MS peptide mass finger printing analysis (Wu et al. 2008).

**5′, 3′ rapid amplification of cDNA ends analysis of full-length gene**

Based on the peptides sequences identified by MALDI-TOF MS analysis, RACE-PCR primers were designed amplify the putative full-length AlocSWP2 gene, and the gene amplified by 5′ and 3′ RACE PCR with the SMARTer RACE Amplification kit (Clontech, 634859; Mountain View, CA), according to the manufacturer’s protocol. Total RNA was extracted from spores using TRIzol (Invitrogen, 15596026; Waltham, MA). 5′-RACE-Ready cDNA and 3′-RACE-Ready cDNA were Synthesized using a 5′-CDS Primer or a 3′-CDS Primer A (included in the kit) respectively. The resulting cDNA was used as a template for PCR using an UPm (universal primer A mix) with the 5′ or 3′ AlocSWP2-specific primers (5′-RACE GSP: 5′-AAYTNG-CYTCYTNGTYTCNAGAA-A-3′, 3′-RACE GSP: 5′-ATGT-TRAAYAAYTYYAAYAGRGTGA-3′). Sequencing of the
RACE-PCR products thus confirmed amplification of the full-length AlocSWP2 gene.

**In silico analysis**

Signal peptides were predicted by SignalP 4.1 Server (http://www.cbs.dtu.dk/services/SignalP/). Other modifications, such as N- and O-glycosylation potential sites were predicted by NetOglyc (http://www.cbs.dtu.dk/services/NetOGlyc/) and NetNGlyc (http://www.cbs.dtu.dk/services/NetNGLyc/) servers. Phosphorylation site prediction was also performed through the website, http://www.dabi.temple.edu/disp/phos/pred/predict. The search for glycosylphosphatidylinositol (GPI)-anchorage was done using the DGPI algorithm (http://mendel.imp.ac.at/gpi/gpi_server.html), and also by UniProt (http://www.uniprot.org/) and InterProScan (http://www.ebi.ac.uk/interpro/scan.html).

**Recombinant protein expression, purification, antibody production, SDS-PAGE identification and Western analysis**

The gene encoding AlocSWP2 without signal coding sequence was amplified, via PCR or reverse transcription-PCR (RT-PCR), from A. locustae genomic DNA or total mRNA from infected locust by oligonucleotide primers AlocSWP2-F (5'-CGGATCTATCACAGCAGCGACA-3'), containing a BamHI restriction site (underlined), and AlocSWP2-R (5'-CGGAAATTCGACGAGTAAGCAGCAGCAGCAGCAGC-3'), containing an EcoRI restriction site (underlined) to facilitate cloning, were designed based on the predicted open reading frame XX255658. The amplified fragments were digested by BamHI and EcoRI and cloned into the corresponding restriction enzyme digested expression vector pGEX-4T-2. The recombinant plasmid was transformed into Escherichia coli BL21 (DE3). After induction of expression by IPTG (0.5 mmol/L) at 30 °C, total bacterial protein was extracted and detected in SDS-PAGE. The expressed protein, fused with GST-tag, was purified by affinity chromatography over a glutathione-Sepharose 4B column.

Monospecific polyclonal antiserum against the recombinant AlocSWP2 or GST was produced from rabbits using a standard shortened immunization protocol, conducted by a commercial facility (Vital River Laboratory Animal Technology Co., Ltd., Beijing, China). A rabbit was immunized via intradermal injection at the dorsum with 0.25 g of GST-AlocSWP2 fusion protein mixed with Freund’s adjuvant (1:1 [v/v]), followed by booster injections with incomplete Freund’s adjuvant (1:1[v/v]) every 2 wks. Sera were collected preimmunization 2 wks after the third injection and stored at −80 °C.

GST-AlocSWP2 fusion protein and A. locustae “SDS extract” samples were subjected to SDS-PAGE on 12% polyacrylamide gels. After electrophoresis, proteins were stained with CBB and transferred onto nitrocellulose filter membrane (Whatman, 10401196; Dassel, Germany) for western blot analysis. The membranes were blocked with 10 ml with 5%(w/v) nonfat milk in PBST (0.05% [v/v] Tween-20 in PBS) (blocking solution) for 2 h, and then incubated at room temperature for 1 h with 10 μg/ml rabbit anti-AlocSWP2 antibody in PBST. Membranes were washed thrice in PBST; the membranes were then incubated at room temperature for 1 h with goat anti-rabbit IgG conjugated with HRP (1:10,000) as second antibody in PBST with 5%(w/v) nonfat milk, and then finally washed with PBST, and reactivity was detected using enhanced chemiluminescence (ECL) reagent (thermo) in a Western blot analysis by FluorChemM system (ProteinSimple, San Jose, CA).

**Immunofluorescence assay**

A host cell binding assay was designed to determine which Spore wall protein interacted with the host cell surface according to Southern et al. (2007). The purified spores (5 × 10⁵) were washed with PBS and fixed with 80% cold acetone for 20 min at room temperature. Slides of fixed spores were permeabilized by PBS with 0.5% Triton X-100 for 15 min. After washing with PBS with 0.05% Tween-20 for three times, the samples were blocked in 5% nonfat milk for 30 min, followed by incubation with anti-AlocSWP2 antibody, diluted at 1:64 dilution of FITC-conjugated goat anti-rabbit IgG for 1 h. DAPI (4′,6-diamidino-2-phenylindole) (5 μg/ml) was used to stain DNA for 30 min at room temperature before the final wash. Finally, the slides were mounted with a glycerol solution, and visualized on an OLYMPUS IX81 fluorescence microscope (excitation wavelength of FITC-IgG: 495 nm; DAPI: 359 nm) (Accoceberry et al. 1999; Alfa Cisse et al. 2002).

**Immunocytochemistry localization**

Transmission electron microscopy immunolabeling (IEM) experiments were conducted with A. locustae-infected. The chemical fixation was done via immersion of locust fat bodies into a mixture of paraformaldehyde (4%) and glutaraldehyde (2%) in 0.1 M PBS (pH 7.4), followed by dehydration in an ethanol series. The samples were embedded in LR White resin (Taab, Aldermaston, Berks, U.K.) via polymerization at 60 °C in tightly closed gelatin capsules. Ultrathin sections were cut with a glass knife on an ultramicrotome and mounted on Formvar-coated grids. For immunocytochemistry, the grids were subsequently floated on 30 μl droplets of the following solutions, mainly adapted from Steinbrecht (Steinbrecht 1992, 1993): in brief, PBS containing 50 mM glycine, PBGT (PBS containing 0.2% gelatin, 1% bovine serum albumin, and 0.02% Tween-20), primary antibody diluted with PBGT, six washings with PBGT, secondary antibody in PBGT, two washings on each PBGT, PBS glycine, PBS, and water. Optional silver intensification (Danscher 1981) increased the size of the gold granules from 10 to about 40 nm; 2% uranyl acetate increased the tissue contrast for observation in transmission electron microscope (HITACHI H-7800; Tokyo, Japan).
Immunocytochemical labeling was done on sections of the fat bodies of three male and three female adult infected locusts. The following antibodies were used in this study: anti-GST-AlocSWP2 antibody and anti-GST antibody. The primary antibodies were diluted at 10 μg/ml in PBGT and incubated at 4 °C overnight. As a control, the primary antisera was replaced by serum from a healthy rabbit. The secondary antibody was anti-rabbit IgG, coupled to 10 nm colloidal gold (AuroProbe™ EM, GAR G10, Amersham, U.K.), diluted 1:20 and incubated at room temperature for 60–120 min.

RT-PCR analysis

Two or three days after the 3rd instar nymph locusts, which were starvation treatments for 4 h, they were fed with 10⁷ spores of A. locustae coated on 20 mm x 5 mm corn leaves and were used to detect when A. locustae entered the fat body and propagated. Locusts’ fat bodies were run on 1.2% agarose gels and visualized via ethidium bromide staining, then observed and photographed. Approximately 160 locusts were collected and transferred into individual mini-boxes. According to the previously described method, locusts were infected with A. locustae. At the 10th day postinfection, dsRNA was injected into the locust. Three days later, the mRNA was extracted from dissected tissues using Trizol. The cDNA was synthesized from 1 mg of total RNA with MLV reverse II RT-PCR system (Promega). Both AlocSWP2 and A. locustae actin sequences were amplified from the same cDNA (3–5 individuals per replicate). The actin sequence of A. locustae (GenBank accession number: AF031702.1) was used as an internal control, with the primers AlocActin-F: 5′-GGCATTCCCAAGGCACA AAGG-3′; AlocActin-R: 5′-ACAGAACACGCTGAATCGCA-3′. The mRNA from the AlocSWP2 gene was amplified using the AlocSWP2-F/R primers. Semi-quantitative PCR was performed using a LightCycler Nano (Roche, Basel, Switzerland), and the specificity of amplification was confirmed through melting curve analysis. Gene transcription difference was calculated by the 2-DDCt values method (Livak and Schmittgen 2001). For normalization between samples, the mRNA levels from the actin of A. locustae gene were validated experimentally for each generation and treatment, with the geometric mean then used for normalization according to the strategy described previously (Vandesompele et al. 2002). The PCR program was as following: 94 °C for 10 min, followed by 45 cycles for qRT-PCR (semi-quantitative PCR at 25 cycles), each cycle consisting of 94 °C for 15 s and 58 °C for 15 s and 72 °C for 15 s. At the end, samples were incubated to 4 °C for 10 min.

Production of dsRNA

To specifically reduce AlocSWP2 protein levels in A. locustae, we used RNAi to knockdown the AlocSWP2 expression. Primers for the full AlocSWP2 gene, without the signal coding sequence used were: AlocSWP2-F: 5′-TAATACGACTCATATAGGATGACACGGAGCGAGCGACAG-3′; AlocSWP2-R: 5′-TAATACGACTCATATAGGATGACACGGAGCGAGCGACAG-3′. AlocSWP2-F: 5′-ATGT TCTCGGGGCCACAC-3′. PCR reactions were performed under the following thermal program: 94 °C for 10 min; 30 cycles of 94 °C for 15 s, 58 °C for 15 s, 72 °C for 30 s; followed by one cycle at 72 °C for 10 min. PCR products were run on 1.2% agarose gels and visualized via ethidium bromide staining, then observed and photographed. The spores just entered into fat body to express AlocSWP2, dsRNA was injected into the locust. Three days later, the mRNA was extracted from dissected tissues using Trizol. The cDNA was synthesized from 1 mg of total RNA with MLV reverse II RT-PCR system (Promega). Both AlocSWP2 and A. locustae actin sequences were amplified from the same cDNA (3–5 individuals per replicate). The actin sequence of A. locustae (GenBank accession number: AF031702.1) was used as an internal control, with the primers AlocActin-F: 5′-GGCATTCCCAAGGCACA AAGG-3′; AlocActin-R: 5′-ACAGAACACGCTGAATCGCA-3′. The mRNA from the AlocSWP2 gene was amplified using the AlocSWP2-F/R primers. Semi-quantitative PCR was performed via MyCycler (Bio-Rad), qRT-PCR was performed using a LightCycler Nano (Roche, Basel, Switzerland), and the specificity of amplification was confirmed through melting curve analysis. Gene transcription difference was calculated by the 2-DDCt values method (Livak and Schmittgen 2001). For normalization between samples, the mRNA levels from the actin of A. locustae gene were validated experimentally for each generation and treatment, with the geometric mean then used for normalization according to the strategy described previously (Vandesompele et al. 2002). The PCR program was as following: 94 °C for 10 min, followed by 45 cycles for qRT-PCR (semi-quantitative PCR at 25 cycles), each cycle consisting of 94 °C for 15 s and 58 °C for 15 s and 72 °C for 15 s. At the end, samples were incubated to 4 °C for 10 min.

RNAi of AlocSWP2 in infected locusts

To observe the effect of RNAi treatment on AlocSWP2 transcription, semi-quantitative PCR and qRT-PCR experiments were performed according to the manufacturer protocol. On the 10th day after A. locustae infection, when

Semi-quantitative PCR and quantitative reverse transcription PCR analysis

The effect of RNAi treatment on AlocSWP2 transcription, semi-quantitative PCR and qRT-PCR experiments were performed according to the manufacturer protocol. On the 10th day after A. locustae infection, when...
infected control groups were supplemented with dsRNA buffer (control) and GFP-specific dsRNA (GFP RNAi), respectively. As previous treatment strategy, locusts were infected with A. locustae. At the 10th day postinfection, RNAi treatment was conducted. Samples of dead locusts were collected immediately and frozen in −80 °C, until at the 16th day postinfection, all locusts were obtained for spore load counts with a hemocytometer (Plishchuk et al. 2013). All statistical t-tests (and nonparametric tests) followed by two-tailed comparison tests were performed using GraphPad Prism version 6.00 for Windows, (GraphPad Software Inc., La Jolla, CA).

RESULTS

Identification and characterization of a spore wall protein from Antonospora locustae

About 20 protein spots were separated by 2-DE of the protein preparation extracted from A. locustae spore walls (Fig. 1A). A spore wall protein, with an approximate molecular weight and pl of 20 kDa and 5, respectively was extracted and digested by trypsin, and the protein sequence identified by peptide fingerprinting with MALDI-TOF MS and expressed protein sequence comparisons, as AlocSWP2 (Fig. 1B). RACE-PCR was then used to obtain the full-length gene (GenBank with accession number KX255658), with an open reading frame of 669 bp encoding for a 222-amino acid protein, with a predicted molecular mass of 25 kDa and pl of 5.16, consisting of a 203 amino acid mature protein and a 19 amino acid signal peptide. The predicted mass and pl were consistent with those observed for the protein isolated by 2-DE. The protein contains a potential GPI-modification site in the C-terminal (Ser, 206 amino acids site) and a heparin-binding motif (HBM) composed of LRKGRT (amino acids 43-48), which conforms to the consensus sequence for HBMs, “XBBXBX”. Most interestingly, there are four conserved cysteines (Fig. 1C, asterisks), located at 117, 146, 173, and 23, as well as several other conserved amino acids in these proteins, such as Leucine (128), Phenylalanine (155, 180), Proline (163), and glycine (119) although the proteins are from different microsporidian parasites whose hosts vary from Insecta to Mammalia. However, the phylogenetic analysis showed that AlocSWP2 fall into a specific single branch that is quite different from other proteins, both from insects and mammals in the phylogenetic tree (Fig. 1D). The purified recombinant protein (Fig. 2A) and the protein containing extract from A. locustae (Fig. 2B) were analysed by 12% SDS-PAGE followed by western blotting. The anti-GST-AlocSWP2 polyclonal antibody labeled the recombinant GST-fused protein that showed a molecular weight of ~40 kDa, consistent to the calculated molecular weight of the fusion protein (GST-AlocSWP2 without the signal peptide; Fig. 2C). A single ~25 kDa protein band was detected by anti-GST-AlocSWP2 polyclonal antibody in SDS extract (Fig. 2D). Preimmune serum was used as a negative control in all Western blots (data not shown) and nonspecific bands were detected. This proved that a polyclonal antibody was successfully produced in rabbits and had a strong activity against purified recombinant AlocSWP2 or endogenous AlocSWP2 in A. locustae.

AlocSWP2 was localized in the spore wall

AlocSWP2 was detected in the wall of mature spores via indirect immunofluorescence assay (IFA) (Fig. 3). Antonospora locustae spores were treated with the anti-GST-AlocSWP2 antibody, which labeled the spore wall (green fluorescent signal), while DAPI stained DNA was visible within the spores (Fig. 3A). No background green fluorescence was observed in spores treated with anti-GST (Fig. 3B). The results suggest that AlocSWP2 was localized on the surface of the spore coat.

Furthermore, the IEM experiments were used to further confirm that AlocSWP2 was a spore wall protein. The fat body of the host locust was used to provide more detail about the protein in spore development process. Dark granules in ultra-sections are anti-GST-AlocSWP2 antibody coupled to gold particles, and then amplified by silver staining, were observed in both the endospore and exospores wall, indicating that this protein is expressed in both (Fig. 4). The dark granules of anti-AlocSWP2 were also observed in the spore walls of the sporoblast and sporont, but much less intense than in mature spores, indicating that the protein expression begins with spore wall formation. We found that the anti-GST-AlocSWP2 antibody labeled the layers of spore wall during formation (Fig. 4E–J), and the thickness of this label increased with thickness of spore wall (Fig. 4F–J). After maturation, AlocSWP2 is mainly localized to the endospore (Fig. 4J), shows in magnified pictures (Fig. 4H and I). Notably, dark granules distributed in both sporoblast and sporont (Fig. 4F and G) in disarray; however, they were arranged in an orderly fashion in the mature spore (Fig. 4J). AlocSWP2 is differently distributed during the life cycle stages of the sporoblasts and mature spores of A. locustae. No labeling was observed in the negative controls labeled with anti-GST (Fig. 4A–D).

RNAi of AlocSWP2 reduced the mortality of locusts infected by Antonospora locustae through its sporulation

In order to determine when AlocSWP2 starts to be expressed in its host, RT-PCR of AlocSWP2 gene was performed after inoculation (Fig. 5A), showing no expression until 9–11 d postinoculation. For RNAi treatment, dsRNA was prepared and injected into the locust hemolymph on the 10th day postinoculation with spores of A. locustae. Total RNA was extracted 3 days later for semi-quantitative PCR and qRT-PCR experiments. AlocSWP2 transcription was reduced in both detection of RNAi by semi-quantitative PCR (Fig. S1) and in the fat body of infected locust after injection of AlocSWP2 dsRNA was significantly reduced approximately 26 fold compared to those in the mock-injected group (no RNAi; Fig. 5B and
Figure 1  A novel protein, AlocSWP2, isolated from Antonospora locustae spores. (A) Proteins profile extracted from A. locustae spores in 2 dimensions electrophoresis SDS PAGE. The target protein highlighted (dashed circle). M. protein molecular weight markers, pl. isoelectric point. (B) Amino acid sequence of AlocSWP2, showing the signal peptide (italicized), and the predicted HBM (box), the asparagine predicted to be N-glycosylated (underlined) and the potential GPI-modification (grey shade). (C) Sequence alignment of the amino sequence of AlocSWP2 with other 14 amino acid sequences of putative spore wall proteins of other species in Microsporidia. Identical and similar residues are highlighted in black and grey, respectively. The conservative cysteine sequences of these Microsporidia marked with an asterisk. Microsporidia and UniProt database ID: A. locustae (Q6E6F1); Encephalitozoon cuniculi (QBSVGB); Encephalitozoon hellllem (I6UB54); Encephalitozoon romalaei (I6ZS97); Enterocytozoon bieneusi (B7XHL8); Encephalitozoon intestinalis (E0SM53); Edhazardia aedis (J9DKY7); Nosema ceranae (C4VBU8); Nosema bombycis (R0KVG4); Nosema apis (T0KWN0); Vittaforma corneae (L2GPT0); Vavraia culicis (L2GDW6); Trachipleistophora hominis (L7JV52); Spraguea lophii (S7W7E5). (D) Molecular phylogenetic analysis of spore wall protein sequences from Microsporidia. The evolutionary history was inferred via the Maximum Likelihood method based on the JTT matrix-based model and conducted in MEGA6 (Tamura et al. 2013). The bootstrap consensus tree inferred from 1,000 replicates is taken to represent the evolutionary history of the taxa analyzed.
This proved that injection with dsRNA of AlocSWP2 can effectively depress expression of AlocSWP2.

To examine the effects of RNAi of AlocSWP2 on pathogenicity of the parasite to its host, survival rates were measured with and without the RNAi treatment by injection 10 d postinoculation with A. locustae spores. Mortality rates from five treatment groups were compared: locusts infected with A. locustae spores and treated with RNAi against AlocSWP2 (infected + AlocSWP2 RNAi); infected locust without RNAi against AlocSWP2 (infected control) and Infected + GFP control RNAi group; healthy locust (noninfected control); and healthy locusts injected with dsRNA against AlocSWP2 mRNA (noninfected + AlocSWP2 RNAi). Both survival proportions of Infected + AlocSWP2 RNAi group against Infected + GFP control RNAi group and the group of Infected control was significantly different. No deaths occurred during the examined period, exceptions happened due to natural and accidental causes within 10 days, as well as due to failed dsRNA injections. However, the mortalities of the infected locusts treated with RNAi against AlocSWP2 mRNA were significance lower on the 15th, 16th, and 17th days after inoculation with parasite spores than those of locusts in infected control group (Fig. 5C and Table S2).

The effects of treatment on A. locustae intensity (average spore load) are summarized in Fig. 5D. Among the AlocSWP2 RNAi treatments that received spore inoculations, significant differences were found in spore intensities ($P < 0.01$). There were no significant differences in dsRNA buffer (Control) and GFP-specific dsRNA (GFP RNAi) treatments ($P = 0.6131$). After silencing AlocSWP2, A. locustae failed to produce spores; consequently, the host suffered less from the parasite burden.
Figure 4 Immunocytochemistry localization of AlocSWP2 in fat bodies of locusts. (A–D) Negative control, A. locustae-infected locust fat cells treated with anti-GST antibody. (E–K) Sections of A. locustae-infected locust fat cells incubated with anti-GST- AlocSWP2 antibody. Ex = exospore; En = endospore; Pt = polar tube; Nu = nucleus; Sp = sporont; Sb = sporoblast; M = meront; S = mature spores; F = locust fat cell. Bars in D, H, I, 0.5 μm; Bars in A, C, F, G, J, 1 μm; Bar in B, 2 μm; Bar in E, 5 μm.
and demonstrated reduced mortality in the case of RNAi treatments.

**DISCUSSION**

Some spore wall proteins have been identified since the late 1990s. The genes and proteins of microsporidians can now be identified rapidly since the genomes of several microsporidians have been sequenced, such as *E. cuniculi*, *N. bombycis*. So far, only about 20 spore wall proteins have been reported both in the *Encephalitozoonidae* family and in *N. bombycis*. Typical spore wall proteins of microsporidian may have the following features (Table 1): (i) 15–20 amino acid signal peptide sequences; a predicted GPI-anchor site indicating that it is a membrane or spore wall protein; glycosylation and/or other posttranslational modification; glycine or serine rich; and have adherence domains such as HBM. (ii) These proteins are generally cysteine rich, with most spore wall proteins containing four conservative cysteine required disulfide bond formation, however, there is one such protein without any cysteines. (iii) The sequences of SWPs are highly diverse. In the same host, the sequences of amino acid of spore wall proteins may be similar, for example SWP25 and SWP26 in *N. bombycis*. However, no homologous proteins were found in *E. cuniculi* so far (Hayman et al. 2001; Li et al. 2009; Wu et al. 2009). In our study, four conserved cysteines were identified in *AlocSWP2* and other Microsporidia, which infect both insects and mammals. Furthermore, this feature may lead to them resembling tertiary structures or functions via disulfide-bonded linkages. Bohne et al. studied the spore wall protein *E. cuniculi*, and found that the cysteines and N-terminal signal peptide sequences were conserved, indicating similar tertiary structure or function (Bohne et al. 2000).

It has been reported that Microsporidia adhere to host cells that are dominated by GAGs on surface (Hayman et al. 2005), with some spore wall proteins acting as a ligand during spore adherence to the host (Brosson et al. 2005; Southern et al. 2007). Heparin-binding motifs (HBMs) also exist in the amino acid sequence of *AlocSWP2*. It was suggested that HBMs interact with extracellular GAGs, with the consensus sequence necessary for protein-heparin interaction characterized by “XBBXB,” “XBBBXXBX,” or “XBBXBXXBX” (“X” is any neutral amino acid and “B” is a positively charged basic amino acid) (Wu et al. 2008). Typical spore wall proteins from Microsporidia have features that include HBMs (Table 1), with one such motif identified in *AlocSWP2*. Recent reports showed that blocking either SWP16 or SWP11 using in vitro antibody treatment caused a 20% decrease in the adherence of *N. bombycis* spores to host cells for either case (Wang et al. 2015; Yang et al. 2014).

By using a host cell binding assay, recombinant SWP26 protein (with HBM) can inhibit *N. bombycis* adherence by 10%, resulting in decreased host cell infection. In contrast, mutant rSWP26 (without HBM) did not inhibit spore adherence (Li et al. 2009). The HBMs may be important
Table 1. The identified spore wall proteins of Microsporidia

| Protein        | Source                        | Mw (kDa) | Full length (nt)/amino acids (aa) | PI     | Phosphorylation/ Nglycosylation/ Oglycosylation | Subcellular localization | Subcellular localization | Functional domain | References/GenBank ID |
|----------------|-------------------------------|----------|-----------------------------------|--------|-----------------------------------------------|--------------------------|--------------------------|----------------------|-----------------------|
| EiSWP1         | Encephalitozoon intestinalis  | 41.5     | 1167/388 4.78                     | Yes/18 | 36/1/37                                       | External wall            | –                        | –                    | Hayman et al. (2001); AF355750.1 |
| EiSWP2         | E. intestinalis               | 107.2    | 3009/1002 3.68                    | Yes/18 | 83/1/1                                        | External wall            | HBM                     | –                    | Hayman et al. (2001); AF355749.1 |
| EiEnP1         | E. intestinalis               | 39.1     | 1047/348 8.84                     | Yes/16 | 28/3/1                                        | Internal wall/ external wall/polar membrane layer | –                        | HBM                  | Corradi et al. (2010) and Pombert et al. (2012); XM_003072248 |
| ErEnP1         | E. romaleae                   | 39.5     | 1047/348 9.12                     | Yes/16 | 15/3/3                                        | External wall            | –                        | HBM                  | Pombert et al. (2012); XM_009265627.1 |
| EcSWP1         | Encephalitozoon cuniculi      | 45.9     | 1353/450 4.96                     | Yes/18 | 66/0/52                                       | External wall            | –                        | HBM                  | Katinka et al. (2001); NM_001042141.6 |
| EcCDA          | E. cuniculi                   | 28.1     | 765/254 4.43                      | Yes/15 | 16/0/0                                        | Endospore plasma membrane | Glycoside hydrolase/ Deacetylase | –                      | Brosson et al. (2005); NC_003237.1 |
| EcEnP1         | E. cuniculi                   | 40.6     | 1074/357 9.07                     | Yes/16 | 20/1/3                                        | Endospore               | HBM                     | –                    | Katinka et al. (2001) and Peuvel-Fanget et al. (2006); XM_900824 |
| EcEnP2/SWP3    | E. cuniculi                   | 22.5     | 666/221 8.42                      | Yes/20 | 31/0/27                                       | Transmembrane plasma membrane | –                      | –                    | Katinka et al. (2001), Peuvel-Fanget et al. (2006), and Xu et al. (2006); NC_003242.2 |
| EhSWP1a        | Encephalitozoon hellem        | 54.9     | 1530/509 4.30                     | Yes/18 | 38/3/27                                       | Extracellular spores extracellular spores | –                      | –                    | Polonais et al. (2010); FJ870923 |
| EhSWP1b        | E. hellem                     | 57.9     | 1602/533 4.64                     | Yes/18 | 44/3/29                                       | Extracellular spores extracellular spores | –                      | –                    | Polonais et al. (2010); FJ870924 |
| NbSW16         | Nosema bombycis               | 22.5     | 666/221 8.42                      | Yes/15 | 34/2/29                                       | Exosporium               | HBM                     | –                    | Wang et al. (2015); KB908937.1 |
| NbHSWP11       | N. bombycis                   | 52.3     | 1341/446 9.27                     | No     | 32/0/1                                        | Internal wall/ external wall | DnaJ domain             | –                    | Yang et al. (2014); EF683111 |
| SWP5           | N. bombycis                   | 20.3     | 561/186 4.39                      | Yes/22 | 13/0/6                                        | Exosporium/polar tube   | –                        | –                    | Li et al. (2012); HQ881497 |
| NbSWP12        | N. bombycis                   | 26.6     | 687/228 6.78                      | No     | 13/1/0                                        | Internal wall/ external wall | BAR-2 domain           | –                    | Chen et al. (2013); KC193258 |
| SWP25/HSWP2    | N. bombycis                   | 30.7     | 807/268 8.45                      | Yes/25 | 15/2/6                                        | Endospore               | HBM                     | –                    | Wu et al. (2008, 2009); EF683102 |
| SWP26          | N. bombycis                   | 25.7     | 672/223 5.09                      | Yes/16 | 13/0/0                                        | Endospore               | HBM                     | –                    | Li et al. (2009); EU677842 |
| SWP30/HSWP1    | N. bombycis                   | 32.1     | 837/278 7.95                      | Yes/19 | 22/1/4                                        | Endospore               | –                        | –                    | Wu et al. (2008); EF683101 |
| SWP32/HSWP3    | N. bombycis                   | 37.4     | 951/316 7.29                      | Yes/18 | 12/1/0                                        | Exosporium               | –                        | –                    | Wu et al. (2008); EF683103 |
| NbSWP7         | N. bombycis                   | 32.3     | 864/287 4.78                      | Yes/19 | 7/0/1                                         | Exosporium/ endospore   | –                        | –                    | Yang et al. (2015); EOB13707.1 |

(continued)
for adherence of the spore wall to host cells during the passage of the spores through the gastrointestinal tract, facilitating invasion.

In our research, AlocSWP2 transcription in the fat body of infected locusts by injection of AlocSWP2-specific dsRNA was significantly reduced as detected by qRT-PCR. In fact, RNAi were demonstrated to be efficiently used to regulate microsporidian gene expression in honeybee or silkworm hosts (Paldi et al. 2010; Pan et al. 2017). Survival percentages of locusts that were infected with A. locustae without RNAi of AlocSWP2 were lower than those of the infected locusts that were injected with dsRNA of AlocSWP2 gene, suggesting that this protein may be involved in host pathogenicity. Further analysis revealed that the amount of spores of RNAi of AlocSWP2 in each infected locust was much lesser than those of infected locusts without RNAi of the gene. Therefore, considering the interference of RNA within host fat body, less expression of the protein caused fewer amount of sporonts and spores, resulting in lower host mortality. As we know, the pathogenicity of microsporidian mainly depends on the amount of spores, which cause an acute anaphylactic reaction (Selman 1982; Szumowski and Troemel 2015). Interestingly, here we found that locusts in the RNAi treatments specifically knockdown certain structural protein such as AlocSWP2, and consequently had lower A. locustae intensities, and higher survival. We further speculate that the expression of some of the spore wall proteins of microsporidian may contribute to the sporulation of spores or other functions, which influence the pathogenicity of this microsporidian to its host. This is the first demonstration that a spore wall protein from A. locustae contributes to the mortality of its host, and provides a novel spore wall protein of Microsporidia whose hosts can be found within both Insecta and Mammalia contributing to sporulation.

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**LITERATURE CITED**

Accoceberry, I., Thellier, M., Desportes-Livage, I., Achbarou, A., Biligui, S., Danis, M. & Datry, A. 1999. Production of monoclonal antibodies directed against the microsporidium Enterocytozoon bieneusi. J. Clin. Microbiol., 37:4107–4112.

Alfa Cisse, O., Ouattara, A., Thellier, M., Accoceberry, I., Biligui, S., Minta, D., Douombo, O., Desportes-Livage, I., Thera, M. A., Danis, M. & Datry, A. 2002. Evaluation of an immunofluorescent-antibody test using monoclonal antibodies directed against Enterocytozoon bieneusi and Encephalitozoon intestinalis for diagnosis of intestinal microsporidiosis in Barnako (Mali). J. Clin. Microbiol., 40:1715–1718.
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Beznoussenko, G. V., Dolgikh, V. V., Seliverstova, E. V., Seme
nov, P. B., Tokarev, Y. S., Trucco, A., Micaroni, M., Di Gian
domenico, D., Auinger, P., Sendersky, I. V., Skarlato, S. O., Snigirevskaya, E. S., Komnissarchik, Y. Y., Pavelka, M., De Mat
teis, M. A., Luini, A., Sokolova, Y. Y. & Mironov, A. A. 2007. Analogs of the Golgi complex in Microsporidia: structure and aves
cular mechanisms of function. J. Cell Sci., 120:1288–1298.

Bohne, W., Fersdgon, D. J., Kohler, K. & Gross, U. 2000. Devel
opmental expression of a tandemly repeated, glycine- and ser
ine-rich spore wall protein in the microsporidian pathogen Nosema locustae. Infect. Immun., 68:2268–2276.

Brooks, W. M. 1988. CRC handbook of natural pesticides. Vol. V. Microbial insecticides. Part A. Entomogenous protozoa and fungi. In: Ignoffo, C. M. (ed.), Entomogenous Protozoa. CRC Press, Florida, p. 1–49.

Brosson, D., Kuhn, L., Delbac, F., Garin, J., Vives, C. P. & Texier, C. 2006. Proteomic analysis of the eukaryotic parasite Encephalitozoon cuniculi (Microsporidia): a reference map for proteins expressed in late sporogonial stages. Proteomics, 6:3625–3635.

Brosson, D., Kuhn, L., Prensier, G., Vives, C. P. & Texier, C. 2005. The putative chitin deacetylase of Encephalitozoon cuniculi: a surface protein implicated in microsporidian spore-wall formation. FEMS Microbiol. Lett., 247:81–90.

Chen, J., Geng, L., Long, M., Li, T., Yang, D., Ma, C., Wu, H., Ma, Z., Li, C., Pan, G. & Zhou, Z. 2013. Identification of a novel chitin-binding spore wall protein (NbsSWP12) with a BAR-2 domain from Nosema bombycis (Microsporidia). Parasitology, 140:1394–1402.

Coman, R. S., Chen, Y. P., Schatz, M. C., Street, C., Zhao, Y., Desany, B., Egholm, M., Hutchison, S., Pettis, J. S., Lipkin, W. I. & Evans, J. D. 2009. Genomic analyses of the microsporidian Nosema ceranae, an emergent pathogen of honey bees. PLoS Pathog., 5:e1000466.

Corradi, N., Pombert, J. F., Farinelli, L., Didier, E. S. & Keeling, P. J. 2010. The complete sequence of the smallest known nuclear genome from the microsporidian Encephalitozoon intestinalis. Nat. Commun., 1:177.

Danscher, G. 1981. Light and electron microscopic localization of silver in biological tissue. Histochemistry, 71:177–186.

Dolgikh, V. V., Sementov, P. B., Mironov, A. A. & Bezouensenko, G. V. 2005. Immunocytochemical identification of the major exospore protein and three polar-tube proteins of the Microsporidia Nosema grylli. Protist, 156:77–87.

Franzen, C. 2005. How do Microsporidia invade cells? Folia Para
tiol. (Prah.), 52:36–40.

Franzen, C., Muller, A., Hartmann, P. & Salzberger, B. 2005. Cell invasion and intracellular fate of Encephalitozoon cuniculi (Microsporidia). Parasitology, 130:285–292.

Gatehouse, H. S. & Malone, L. A. 1998. The ribosomal RNA gene region of Nosema apis (Microsporid): DNA sequence for small and large subunit RNA genes and evidence of a large tandem repeat unit size. J. Invertebr. Pathol., 71:97–105.

Hayman, J. R., Hayes, S. F., Amon, J. & Nash, T. E. 2001. Develop
mental expression of two spore wall proteins during matura
tion of the microsporidian Encephalitozoon intestinalis. Infect. Immun., 69:7057–7066.

Hayman, J. R., Southern, T. R. & Nash, T. E. 2005. Role of sul
fated glycans in adherence of the microsporidian Encephalitozo
n intestinalis to host cells in vitro. Infect. Immun., 73:841–
848.

Henry, J. E. 1971. Experimental application of Nosema locustae for control of grasshoppers. J. Invertebr. Pathol., 18:389–394.

Katinka, M. D., Duprat, S., Cornillett, E., Metenier, G., Thomarat, F., Prensier, G., Barbe, V., Peyretailade, E., Brottier, P., Wincker, P., Delbac, F., El Alaoui, H., Peyret, P., Saurin, W., Gouy, M., Weissensenbach, J. & Vives, C. P. 2001. Genome sequence and gene compaction of the eukaryote parasite Encephalitozoon cuniculi. Nature, 414:450–453.

Li, Z., Pan, G., Li, T., Huang, W., Chen, J., Geng, L., Yang, D., Wang, L. & Zhou, Z. 2012. SWP5, a spore wall protein, inter
acts with polar tube proteins in the parasitic microsporidian Nosema bombycis. Eukaryot. Cell, 11:229–237.

Li, Y., Wu, Z., Pan, G., He, W., Zhang, R., Hu, J. & Zhou, Z. 2009. Identification of a novel spore wall protein (SWP26) from Microsporidia Nosema bombycis. Int. J. Parasitol., 39:391–398.

Livak, K. J. & Schmittgen, T. D. 2001. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods, 25:402–409.

Maori, E., Paldi, N., Shafir, S., Kalev, H., Tsur, E., Glick, E. & Sela, I. 2009. iAPV, a bee-affecting virus associated with Colony Col
apse Disorder can be silenced by dsRNA ingestion. Insect Mol. Biol., 13:55–60.

Nunes, F. M., Aleixo, A. C., Barchuk, A. R., Bomtorin, A. D., Groz
inger, C. M. & Simoes, Z. L. 2013. Non-target effects of Green Fluorescent Protein (GFP)-derived double-stranded RNA (dsRNA-GFP) used in honey bee RNA interference (RNAi) assays. Insects, 4:90–103.

Paldi, N., Glick, E., Oliva, M., Zilberberg, Y., Aubin, L., Pettis, J., Chen, Y. & Evans, J. D. 2010. Effective gene silencing in a microsporidian parasite associated with honeybee (Apis mellifera) colony declines. Appl. Environ. Microbiol., 76:5960–
5964.

Pang, Q., Wang, L., Dang, X., Ma, Z., Zhang, X., Chen, S., Zhou, Z. & Xu, J. 2017. Bacterium-expressed dsRNA downregulates microsporidia Nosema bombycis gene expression. J. Eukaryot. Microbiol., 64:278–281.

Pelín, A., Selman, M., Aris-Brosou, S., Farinelli, L. & Corradi, N. 2015. Genome analyses suggest the presence of polyploidy and recent human-driven expansions in eight global populations of the honeybee pathogen Nosema ceranae. Environ. Micro
biol., 17:4443–4458.

Pevuel-Fanget, I., Polonais, V., Brosson, D., Texier, C., Kuhn, L., Peyret, P., Vives, C. & Delbac, F. 2006. EnP1 and EnP2, two proteins associated with the Encephalitozoon cuniculi endos
drome, the chitin-rich inner layer of the microsporidian spore wall. Int. J. Parasitol., 36:709–715.

Pilschuk, S., Bardi, C. J. & Lange, C. E. 2013. Spore loads of Para
nosema locustae (Microsporidia) in heavily infected grasshoppers (Orthoptera: Acridoidea) of the Argentine Pampas andPatagonia. J. Invertebr. Pathol., 114:89–91.

Polonais, V., Belkorchia, A., Roussel, M., Peyretailade, E., Peyret, P., Diogon, M. & Delbac, F. 2013. Identification of two new polar tube proteins related to polar tube protein 2 in the microsporidian Antonospora locustae. FEMS Microbiol. Lett., 346:36–44.

Polonais, V., Mazet, M., Wawrzyniak, I., Texier, C., Blot, N., El Alaoui, H. & Delbac, F. 2010. The human microsporidian Encephalitozoon hellem synthesizes two spore wall polymorphic proteins useful for epidemiological studies. Infect. Immun., 78:2211–2230.

Pombert, J. F., Selman, M., Burki, F., Bardell, F. T., Farinelli, L., Sol
ter, L. F., Whitman, D. W., Weiss, L. M., Corradi, N. & Keeling, P. J. 2012. Gain and loss of multiple functionally related, horizontally transferred genes in the reduced genomes of two microsporidian parasites. Proc. Natl Acad. Sci. USA, 109:12638–12643.
Selman, B. J. 1982. Microbial control of pests and plant diseases 1970–1980. *Crop Prot.*, 1:124–125.

Sendersky, I. V., Timofeev, S. A., Seleverstova, E. V., Pavlova, O. A. & Dolgikh, V. V. 2014. Secretion of Antonospora (Paranosema) locustae proteins into infected cells suggests an active role of Microsporidia in the control of host programs and metabolic processes. *PLoS ONE*, 9:e93585.

Shormanovits, C. H., Williams, B. A. & Keeling, P. J. 2004. Transfer of Nosema locustae (Microsporidia) to Antonospora locustae n. comb. based on molecular and ultrastructural data. *J. Eukaryot. Microbiol.*, 51:207–213.

Selman, B. J. 1982. Microbial control of pests and plant diseases 1970–1980. *Crop Prot.*, 1:124–125.

Sendersky, I. V., Timofeev, S. A., Seleverstova, E. V., Pavlova, O. A. & Dolgikh, V. V. 2014. Secretion of Antonospora (Paranosema) locustae proteins into infected cells suggests an active role of Microsporidia in the control of host programs and metabolic processes. *PLoS ONE*, 9:e93585.

Williams, B. A. 2009. Unique physiology of host-parasite interactions in Microsporidia infections. *Cell. Microbiol.*, 11:1551–1560.

Wu, Z., Li, Y., Pan, G., Tan, X., Hu, J., Zhou, Z. & Xiang, Z. 2008. Proteomic analysis of spore wall proteins and identification of two spore wall proteins from *Nosema bombycis* (Microsporidia). *Proteomics*, 8:2447–2461.

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Additional Supporting Information may be found online in the supporting information tab for this article:

**Figure S1.** The results of semi-quantitative PCR for the detection of RNAi of *AlocSWP2*.

**Table S1.** Numerical data of qRT-PCR to check the results of RNAi of *AlocSWP2*.

**Table S2.** Raw data of the survival assay after RNAi of *AlocSWP2*.

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