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Morphological and molecular characterization of a new microsporidian (Protozoa: Microsporidia) isolated from *Spodoptera litura* (Fabricius) (Lepidoptera: Noctuidae)

S. JOHNY^{1,2}, S. KANGINAKUDRU¹, M. C. MURALIRANGAN² and J. NAGARAJU^{1*}

 ¹Laboratory of Molecular Genetics, Centre for DNA Fingerprinting and Diagnostics, ECIL Road, Nacharam, Hyderabad 500 076, India
 ²G.S. Gill Research Institute, Guru Nanak College, Chennai 600 042, India

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SUMMARY

A microsporidium was isolated from larvae of Spodoptera litura (Fabricius) collected from Tamil Nadu, India. This microsporidian species is monomorphic, disporous and develops in direct contact with the cytoplasm of the host cell. The nuclear configuration of merogonic and sporogonic stages was diplokaryotic. The merogonic proliferative stage was unusual that normal development with 1, 2 and 4 binucleated forms were common, while large multinucleate meronts containing 8 and 12 small compact horseshoe-like diplokaryotic nuclei were also observed. The fresh spores were typically ovocylindrical in shape, with a mean size of $3.91 \times 1.91 \,\mu\text{m}$ and the polar filament length was $\sim 90 \,\mu\text{m}$. Infection was systemic with mature spores produced in the midgut, nervous tissue, muscles, labial glands, gonads, tracheae, epidermis, Malpighian tubules and, most extensively, fat body tissues. The new isolate was highly pathogenic to S. litura larvae. Host specificity tests performed on 37 non-target hosts of 5 different insect orders revealed that the new isolate is pathogenic only to lepidopteran insects. We sequenced the 16S small subunit rRNA (SSU rRNA) gene of the isolate and compared it with 72 non-redundant microsporidian sequences from the GenBank. Based on the light microscopic studies and phylogenetic analyses, the new isolate is assigned to the genus Nosema. Significant differences in the SSU rRNA sequence were identified when compared with the type species Nosema bombycis and other closely related species viz., Nosema spodopterae. Structural differences were also observed in the 16S SSU rRNA between the new isolate and the two above-mentioned microsporidian pathogens. We conclude that the microsporidian isolate reported here is distinctly different from the other known species and is likely to be a new species.

Key words: Microsporidia, Nosema, Spodoptera litura, 16S SSU rDNA, phylogeny.

INTRODUCTION

Microsporidia are obligate, intracellular parasites belonging to the phylum *Microsporidia*, traditionally placed in the Kingdom Protista. However, evidence from phylogenetic analyses using protein coding genes, especially α - and β -tubulins (Keeling and Fast, 2002; Keeling, 2003), and on LSU rRNA sequences (Van de Peer *et al.* 1998) now suggest that microsporidia share a common origin with fungi. The phylum *Microsporidia* consists of approximately 143 genera and over 1200 species (Wittner and Weiss, 1999), which infect a wide range of vertebrate and invertebrate taxa (Didier *et al.* 2000). Currently, almost half of the described genera of microsporidia have an insect as the host (Becnel and Andreadis, 1999). They may infect the gut, fat body,

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reproductive, muscle, nervous, and silk gland tissues and Malpighian tubules, or can be systemic (Becnel and Andreadis, 1999; Solter and Becnel, 2000). There is renewed interest in studying microsporidia because some species cause opportunistic infections in immunocompromised humans, mainly human immunodeficiency virus-positive subjects (Desportes, Le Charpentier and Galian, 1985; Franzen and Muller, 1999; Kotler and Orenstein, 1999; Molina et al. 2000), but also in travellers, the elderly, and malnourished children (Weber et al. 1994; Muller et al. 2001). Other microsporidia are important in agriculture as pathogens of beneficial insects such as silkworm and honeybees (Becnel and Andreadis, 1999; Nageswara Rao et al. 2004), or as biological control agents for use against medical or agricultural pests (Lockwood, Bomar and Ewen, 1999; Solter and Becnel, 2000; Briano and Williams, 2002). The members of the genus Nosema are often considered the most important and widely distributed group of microsporidia, particularly in the Lepidoptera (Tsai, Lo and Wang, 2003) and most 'Nosema' isolates from hosts of other insect orders are not closely related

^{*} Corresponding author: Laboratory of Molecular Genetics, Centre for DNA Fingerprinting and Diagnostics, ECIL Road, Nacharam, Hyderabad 500 076, India. Tel: +91 40 27171427. Fax: +91 40 27155610. E-mail: jnagaraju@cdfd.org.in

genetically and genus designations are being changed (Slamovits, Williams and Keeling, 2004; Sokolova *et al.* 2005).

The tobacco caterpillar, Spodoptera litura is a polyphagous noctuid with high reproductive potential and the ability to migrate long distances as adults. These factors contribute to the role of S. litura as a major pest of many agricultural crops throughout its geographical range (Armes et al. 1997). In India, it has attained a major pest status on agricultural crops such as cotton, groundnut and cauliflower. Failure to control S. litura in the major cotton and groundnut growing regions of Andhra Pradesh and Tamil Nadu states of India has resulted in an unexpected population outbreak and development of resistance to several insecticides (Armes et al. 1997; Johny and Muralirangan, 2000). In order to avoid resistance problems, as well as to address environmental concerns regarding chemical pesticide use, it is important to identify and develop suitable alternate control strategies. In this context, pathogens may be key biological control agents against major agricultural pests because they are eco-friendly and may be manipulated to effectively control insect pests. In an earlier study, Johny (2002) reported a microsporidian pathogen from the field population of S. litura. In this paper, we provide a developmental and molecular characterization of this microsporidian isolate from S. litura. We have studied the developmental stages including merogony and sporogony, host specificity data, pathogenicity, and the small subunit rRNA (SSU rRNA) gene sequence. The phylogenetic status of the new isolate based on SSU rRNA analyses is also presented.

MATERIALS AND METHODS

Isolation of spores

The microsporidian spores were originally collected from S. litura larvae on 18 January 2000 from a tobacco field in the Dindigul District (10.22°N 78.00°E) of Tamil Nadu, India. The spores were harvested and propagated in S. litura larvae reared on castor leaves. The spores were purified from infected fat body tissues of dead or moribund S. litura larvae collected from the field. The tissues were homogenized in sterile deionized water and the homogenate was filtered through cheesecloth and also cleaned in several cycles of water washes followed by centrifugation. The concentrated spore pellet was further purified by 90% Percoll gradient centrifugation (Undeen and Vavra, 1997). The pelleted spores were collected, washed with sterile distilled water to remove the Percoll content and resuspended in distilled water, quantified by haemocytometer, and stored at 4 °C until further use.

Life-cycle study

S. litura larvae, maintained on castor leaves as a microsporidian-free stock, were used for the present study. Castor leaf discs, each 7 mm diameter, were smeared with $10 \,\mu$ l of water/spore suspension containing 2.4×10^3 spores and allowed to dry at room temperature. Third-instar larvae were starved for 6 h and each of them was fed with 1 leaf disc. The larvae that consumed the entire disc only were retained. Thirty larvae were used for treatment and were maintained separately on fresh castor leaves in plastic Petri dishes at 28 ± 2 °C, light regime 14L:10D.

To study the microsporidian life-cycle, 1 or 2 larvae were dissected every 24 h following the initial infection, up to the completion of the life-cycle. The gut and fat body tissues were removed and placed in a drop of physiological saline (Yeager's solution) on a glass slide, macerated with forceps, fixed in absolute methanol for 2 min, air dried and stained with 10% (v/v) Giemsa staining solution (1 ml of Giemsa stock solution mixed with 9 ml of phosphate buffer, pH 7.4) for about 20-25 min at room temperature. The slides were then rinsed in running water for 5 min, air dried and then observed using a Leitz compound microscope under bright-field conditions for the presence of different stages of the microsporidia. Fresh spores were spread on water agar (Undeen and Vavra, 1997) and measured using an ocular micrometer under phase-contrast microscopy. All the measurements are presented in micrometers as mean values followed parenthetically by range, standard deviation and sample size.

Isolation of DNA, PCR amplification and sequencing of 16S SSU rDNA

The spores of the microsporidian isolate were suspended in 100 mM NaCl, pH 9.5, glycine-NaOH buffer (10 mM) at 30 °C for 15 min to elicit germination (Undeen and Cockburn, 1989). After germination, the emptied content of the spores was used for DNA extraction by standard phenol-chloroform method. The 16S small subunit (SSU) rRNA gene was amplified using the primers 18f 5'-CACCAG-GTTGATTCTGCC-3' and 1537r 5'-TTATGA-TCCTGCTAATGGTTC-3' designed by Baker et al. (1995). PCR amplification was carried out in $200 \,\mu$ l microfuge tubes, using 10 ng DNA, 5 pmol of each primer, 0.2 mM of each dNTP, 2 mM MgCl₂, and 1 U of Taq Polymerase (MBI Fermentas, USA). The amplification conditions used were: 94 °C denaturing for 3 min, followed by 30 cycles of 94 °C for 30 s, 54 °C for 45 s, and extension at 72 °C for 90 s, with a final extension of 10 min on a thermal cycler (Eppendorf). PCR products were directly cloned into PCR 2.1 vector (Invitrogen). Three clones were sequenced using Big dye terminator chemistry on a ABI PRISM 377 automated sequencer.

Phylogenetic analyses

Seventy-two non-redundant microsporidian 16S SSU-rRNA gene sequences were obtained from GenBank (Table 1) aligned with Clustal-X programme (Thompson et al. 1997) and manually edited using the programme GeneDoc (Nicholas and Nicholas, 1997). To identify the best-fit model of nucleotide substitution, MODELTEST programme (Posada and Crandall, 1998) was run as implemented in HYPHY programme (Sergei et al. 2005). Based on the output, GTR+G model was selected as the best-fit model. A maximum likelihood (ML) tree was constructed using the programme DNAml of Phylip package (Felsenstein, 1989) version 3.64, using a gamma shape parameter (alpha) value of 0.892351. The sequence of Amblyospora connecticus was used as an outgroup.

Structural analyses of 16S SSU rRNA

The secondary structure of 16S SSU-rRNA was predicted using the online software 'mfold' (Zuker, 2003) for 3 accessions, L39111 (*N. bombycis*), AY211392 (*N. spodopterae*) and the new isolate reported here. Structures were predicted using the default parameters and the thermodynamically stable and most conserved structure amongst all possible outputs were selected and compared.

Host range

Insects belonging to different orders, collected from both the agroecosystem and other ecosystems (Table 2), were exposed to $10 \,\mu$ l of water/spore suspension containing $2 \cdot 4 \times 10^3$ spores per insect by contaminating their diet. The inoculated larvae and adult insects were dissected from 10 to 15 days postinoculation (p.i.), and the tissues were examined microscopically for the presence of vegetative stages and spores in wet smears as well as in Giemsa-stained specimens.

Pathogenicity

Freshly moulted third-instar larvae of *S. litura*, maintained on castor leaves as a microsporidian-free stock, were inoculated with $10 \,\mu$ l of spore suspension containing $2 \cdot 4 \times 10^2$, $2 \cdot 4 \times 10^3$ and $2 \cdot 4 \times 10^4$ spores, as described under 'Life-cycle study'. Thirty larvae were used for each treatment and were maintained individually in plastic Petri dishes at 28 ± 2 °C, light regime of 14L : 10D, and mortality was recorded daily up to 20 days p.i.. Larval mortality in each treatment was corrected using Abbott's (1925) formula. The spore dosage required to cause 50% mortality (LD₅₀) was calculated by Probit Analysis (Finney, 1971).

Life-cycle of Nosema sp.

Hereafter referred to as NSD (*Nosema* sp. isolated from *S. litura* of Dindigul population.

First merogony and sporogony cycle

The first merogonial stage was observed in midgut epithelial cells at 24 h post-inoculation (p.i.) and was characterized by spherical binucleate meronts measuring $1.88 \,\mu m (1.6 - 2.3, \pm 0.2; n = 10)$ in diameter (Fig. 1A). Growth of the meronts was followed by a nuclear division, resulting in quadrinucleate meronts, measuring 5.94 μ m (5.3–6.6, ± 0.45 ; n = 10) in diameter (Fig. 1B). These meronts have diffused nuclei and an additional division resulted in an octanucleate form with a diameter of $7.09 \,\mu\text{m}$ $(6\cdot6-7\cdot5, +0\cdot34; n=10)$. This stage eventually divided into tetra- and binucleate daughter diplokarya (Fig. 1C-E). Division of meronts was by binary fission of tetranuleate forms. After 48 h p.i., stages with 1, 2 or 4 diplokarya were the most common in the midgut, and nuclei were large and irregular and occupied about half of the area of the meronts (Fig. 1F-H). Giemsa stained the cytoplasm of meronts moderately blue and the nuclei purple. We were unable to identify sporonts, sporoblasts and primary spores with light microscopy in the midgut tissues.

Second merogony and sporogony cycle

After 72 h p.i., large multinucleate meronts with small compact horseshoe-like nuclei of uniform size and shape were observed in the fat body tissues (Fig. 1I, J). These forms developed by repeated karyokinesis without immediate cytokinesis, resulting in the production of 8 and 12 nucleated forms. They subsequently gave rise to binucleated daughter meronts which developed into sporonts. In addition, bi- and tetranucleate meronts similar to that of primary merogony cycle were also observed. This probably represents a second phase of the merogony since we did not observe any infection in the fat body tissues at 48 h p.i.. After 96 h p.i., binucleate sporonts measuring $4.84 \,\mu\text{m}$ ($4.05-5.84, \pm 0.54; n=10$) in diameter, and with deeply stained compact nuclei that occupied a central position in the cells, were observed (Fig. 1K). Further nuclear and cytoplasmic divisions resulted in tetranucleated sporonts of size $5.59 \,\mu\text{m}$ (4.64–6.52, ± 0.70 , n = 10) (Fig. 1L). Subsequently, the tetranucleate sporonts divided into 2 sporoblasts measuring $5.4 \,\mu\text{m}$ (4.52-6.82, ± 0.45 , n = 10) in length and $2.07 \,\mu\text{m}$ (1.84–2.34, ± 0.25 , n=10) in width, characterized by the presence of 2 small intensely stained and centrally placed nuclei (Fig. 1M). After 144 h p.i., binucleate sporonts, sporoblasts and immature spores were

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Table 1.	16S SSU	rRNA se	quences o	of Microsporidia	a used for	phylogenetic	analyses
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Microsporidian name	Host name	Class: Order	Acc. no.
Amblyospora connecticus	Aedes cantator	Insecta: Diptera	AF025685
Antonospora locustae	Grasshopper (Melanoplus)	Insecta: Orthoptera	AY376351
Brachiola algerae	Anopheles stephensi	Insecta: Diptera	AF069063
Culicosporella lunata	Culex pilosus	Insecta: Diptera	AF027683
Encephalitozoon cuniculi 1	Homo sapiens	Mammalia: Primates	L17072
Encephalitozoon cuniculi 2	Homo sabiens	Mammalia: Primates	L39107
Encephalitozoon cuniculi 3	Mus musculus	Mammalia: Rodentia	X98467
Encephalitozoon cuniculi 4	Canis familiaris	Mammalia: Carnivora	X98469
Encephalitozoon cuniculi 5	Orvetolagus cuniculus	Mammalia: Lagomorpha	Z19563
Encephalitozoon hellem 1	Homo sapiens	Mammalia: Primates	AF118142
Encephalitozoon hellem 2	Homo sabiens	Mammalia: Primates	AF118143
Encephalitozoon hellem 3	Homo sapiens	Mammalia: Primates	AF338365
Encephalitozoon hellem 4	Homo sapiens	Mammalia: Primates	AF338366
Encephalitozoon hellem 5	Homo sapiens	Mammalia: Primates	L39108
Encephalitozoon intestinalis 1	Homo sapiens	Mammalia: Primates	L19567
Encephalitozoon intestinalis 2	Homo sapiens	Mammalia: Primates	L39113
Endoreticulatus bombycis	Bombyx mori	Insecta: Lepidoptera	AY009115
Endoreticulatus schubergi	Homo sapiens	Mammalia: Primates	L39109
Endoreticulatus sp. 1	Bombyx mori	Insecta: Lepidoptera	AF240355
Endoreticulatus sp. 2	Ocinara lida	Insecta: Lepidoptera	AY502944
Endoreticulatus sp. 3	Lymantria dispar	Insecta: Lepidoptera	AY502945
Enterocytozoon bieneusi	Homo sapiens	Mammalia: Primates	L16868
Enterocytozoon salmonis	Oncorhynchus tshawytscha	Actinoptervgii: Salmoniforms	U10883
NIK-3 h	Bombyx mori	Insecta: Lepidoptera	AY017212
NIK-4 m	Bombyx mori	Insecta: Lepidoptera	AY017213
Nosema apis 1	A bis mellifera	Insecta: Hymenoptera	U26534
Nosema apis 2	A bis sp	Insecta: Hymenoptera	X73894
Nosema hombi	Rombus terrestris	Insecta: Hymenoptera	AY008373
Nosema hombycis 1	Bombyx mori	Insecta: Lepidoptera	AB093008
Nosema hombycis ?	Bombyx mori	Insecta: Lepidoptera	AB093009
Nosema hombycis 3	Bombyx mori	Insecta: Lepidoptera	AB093010
Nosema hombycis 4	Spodoptera exigua	Insecta: Lepidoptera	AB093011
Nosema bombycis 5	Bombyx mori	Insecta: Lepidoptera	AB093012
Nosema bombycis 6	Bombyx mori	Insecta: Lepidoptera	AB097401
Nosema bombycis 7	Bombyx mori	Insecta: Lepidoptera	AB125666
Nosema bombycis 8	Bombyx mori	Insecta: Lepidoptera	AF240347
Nosema bombycis 9	Bombyx mori	Insecta: Lepidoptera	AY017210
Nosema bombycis 10	Bombyx mori	Insecta: Lepidoptera	AY017211
Nosema bombycis 11	Helicoverpa armigera	Insecta: Lepidoptera	AY259631
Nosema bombycis 12	Bombyx mori	Insecta: Lepidoptera	L39111
Nosema carpocapsae	Cydia pomonella	Insecta: Lepidoptera	AF426104
Nosema ceranae	Apis cerana	Insecta: Hymenoptera	U26533
Nosema furnacalis	Ostrinia furnacalis	Insecta: Lepidoptera	U26532
Nosema granulosis	Gammarus duebeni	Eumalacostraca: Amphipoda	AJ011833
Nosema oulemae	Oulema melanopus	Insecta: Coleoptera	U27359
Nosema sp. isolate 1	Antheraea mylitta	Insecta: Lepidoptera	AB009977
Nosema sp. isolate 2	Noctuid moth	Insecta: Lepidoptera	AF141130
Nosema sp. isolate 3	Calospilos suspecta	Insecta: Lepidoptera	AF240348
Nosema sp. isolate 4	Bombyx mori	Insecta: Lepidoptera	AF240349
Nosema sp. isolate 5	Bombyx mori	Insecta: Lepidoptera	AF240350
Nosema sp. isolate 6	Bombyx mori	Insecta: Lepidoptera	AF240351
Nosema sp. isolate 7	Phyllobrotica armata	Insecta: Coleoptera	AF240353
Nosema sp. isolate 8	Pieris rapae	Insecta: Lepidoptera	AF240354
Nosema sp. isolate 9	Pieris rapae	Insecta: Lepidoptera	AY383655
Nosema sp. isolate 10	Bombyx mori	Insecta: Lepidoptera	D85501
Nosema spodopterae	Spodoptera litura	Insecta: Lepidoptera	AY211392
Nosema tyriae	Tyria jacobaeae	Insecta: Lepidoptera	AJ012606
Nosema sp.	Pierris rapae	Insecta: Lepidoptera	AF485270
Pleistophora sp.	Bombyx mori	Insecta: Lepidoptera	D85500
Vairimorpha cheracis	Cherax destructor destructor	Malacostraca: Decapoda	AF327408
Vairimorpha imperfecta 1	Plutella xylostella	Insecta: Lepidoptera	AJ131645
Vairimorpha imperfecta 2	Plutella xylostella	Insecta: Lepidoptera	AJ131646
Vairimorpha lymantriae 1	Lymantria dispar	Insecta: Lepidoptera	AF033315
Vairimorpha lymantriae 2	Lymantria dispar	Insecta: Lepidoptera	AF141129
Vairimorpha necatrix 1	Apis cerana	Insecta: Hymenoptera	U11051
Vairimorpha necatrix 2	Pseudaletia unipuncta	Insecta: Lepidoptera	Y00266

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Microsporidian name	Host name	Class : Order	Acc. no.
Vairimorpha sp. 1	Solenopsis richteri	Insecta: Hymenoptera	AF031539
Vairimorpha sp. 2	Plutella xylostella	Insecta: Lepidoptera	AF124331
Vairimorpha sp. 3	Bombyx mori	Insecta: Lepidoptera	D85502
Vairimorpha sp. 4	Bombyx mori	Insecta: Lepidoptera	L39114
Vavraia oncoperae	Wiseana sp.	Insecta: Lepidoptera	X74112
Vittaforma corneae	Homo sapiens	Mammalia: Primates	U11046
NSD*	Spodoptera litura	Insecta: Lepidoptera	DQ323510

* The new isolate reported in the present study.

simultaneously observed in the fat body tissues. The cytoplasm of the sporoblasts stained lightly with Giemsa.

Mature spores

At 168 h p.i., sporoblasts were observed in the fat body. The binucleate condition was apparent in immature spores (Fig. 1N), while in the mature environmentally resistant ('environmental') spores nuclei were not distinguishable (Fig. 1O). However, on hydrolysis with 1 M HCl at 60 °C for 1 min, the binucleate state became clear (Fig. 1P). In fresh smears of infected tissues, different stages of spore development were observed. Immature spores were relatively large, with a conspicuous vacuole at one end. After 10 days p.i., environmental spores were observed in large numbers in the cytoplasm of fat body cells. They were highly refractive, ovocylindrical in shape, measuring $3.91 \,\mu m$ (3.8–4.0, ± 0.0123 , n = 10 in length and $1.91 \,\mu m$ (1.8–2.0, ± 0.0143 , n = 10) in width. Mature spores completely replaced the fat body tissue at 15 days p.i.. The polar filament of the spore was extruded upon rehydration of air-dried spores (Kellen and Lindegren, 1969; Watanabe, 1976) and the mean length of the extruded polar filaments was found to be approximately 90 µm.

Sequencing and phylogenetic analyses of 16S SSU-rRNA gene

The 1231 bp 16S SSU rRNA gene of NSD was sequenced using the primers 18f and 1537r. The sequences of 72 non-redundant microsporidian isolates were downloaded from the GenBank and aligned along with the sequence of NSD and the best-fit model of evolution was identified using MODELTEST, which selected the 'TIM+G' model as the best-fit model of nucleotide substitution based on Akaike Information Criterion (AIC) (Akaike, 1974). However, because this model was not available in the Phylip package, we used the nearest alternative GTR+G model (AIC=33501.762) for phylogenetic analyses. The 50% consensus ML tree using gamma correction resulted in a single large cluster of Nosema/Vairimorpha clades (Fig. 2). The SSU rRNA gene sequence of NSD strongly suggested that this species is closely related to the typical Nosema subgroup within the Nosema/ Vairimorpha clade as identified by Baker et al. (1994). The phylogenetic analyses using the sequence data of 72 microsporidia (Table 1) indicated that NSD is closely related to N. bombycis and N. spodopterae. The sequence of NSD differed by 18 bases from N. bombycis (GenBank Accession no. L39111). Altogether there were 2 insertions, 5 deletions, 8 transitions and 3 transversions. NSD also differed by 11 bases from N. spodopterae (AY211392), including 2 insertions, 3 deletions and 6 transitions.

Structure analyses of 16S SSU rRNA

Secondary structure prediction tool, mfold was used to obtain the folding patterns of the 16SSU-rRNA sequences. The default parameters were used for obtaining the most feasible structure at a minimum free energy (ΔG value of -260 ± 4). The result indicated that the secondary structures of *N. bombycis* (L39111) and *N. spodopterae* (AY211392) were similar to each other whereas the structure of the NSD differed from those of both *N. bombycis* and *N. spodopterae* (Fig. 3). Such a difference is obvious because of a 3 nucleotide deletion in the new isolate at the position 918.

Host range

Cross-transmission tests to insects of 5 different insect orders (Table 2) showed that NSD was infective to all the 9 lepidopteran species tested namely, *Bombyx mori* (L.), *Chilo infuscatellus* Sn, *Earias* vitella (F.), Spodoptera exigua (H.), Helicoverpa armigera (H.), Leucinodes orbonalis Gn., Pericallia ricini (Fab.), Plutella xylostella (L.), and Tricoplusia ni (Hb.), although lower percentages of some species were infected. B. mori, S. litura, and P. ricini showed 100% infection. However, other insects of Orthoptera, Coleoptera, Hemiptera, and Diptera did not develop any infection.

Order	Insect species	Stage of inoculation $2 \cdot 4 \times 10^3$ spores/insect	Percentage infected (N*)
Orthoptera	Acrida exaltata (Walk.) Ailopus t. thalassinus (Fab.) Atractomorpha crenulata (Fab.) Curtocomtheories t. textening (Ling.)	Adult Adult Adult	$\begin{array}{c} 0.0 \ (5) \\ 0.0 \ (3) \\ 0.0 \ (10) \\ 0.0 \ (4) \end{array}$
	Diabolocatantops pinguis (Walk.) Epistaurus sinetyi Bol. Eyprepocnemis a. alacris (Serv.) Heteracris pulcher (Bol.) Orthacris maindroni Bol. Oxya fuscovittata (Marsh.)	Adult Adult Adult Adult Adult Adult Adult	$\begin{array}{c} 0.0 & (1) \\ 0.0 & (5) \\ 0.0 & (4) \\ 0.0 & (10) \\ 0.0 & (3) \\ 0.0 & (5) \\ 0.0 & (10) \end{array}$
	Oxya nitidula (Walk.) Spathosternum p. prasiniferum (Walk.) Trilophidia annulata (Thunb.) Tristria pulvinata (Uv.) Poekilocerus pictus (Fab.)	Adult Adult Adult Adult Adult	$\begin{array}{c} 0 \cdot 0 & (10) \\ 0 \cdot 0 & (5) \\ 0 \cdot 0 & (5) \\ 0 \cdot 0 & (3) \\ 0 \cdot 0 & (5) \end{array}$
Coleoptera	Rhizopertha dominica (F) Tribolium castaneium (Herbst) Sitophilus oryzae (L) Atactogaster finitimus Fst Mylabris pustulata (Thump.)	Adult Adult Adult Adult Adult	$\begin{array}{c} 0.0 \ (10) \\ 0.0 \ (15) \\ 0.0 \ (10) \\ 0.0 \ (5) \\ 0.0 \ (3) \end{array}$
Hemiptera	Pyrilla perpusilla Wlk. Nephotettix nigropictus Stal Nilaparvata lugens Stal. Spilostethus hospes (Fab.) Spilostethus pandurus (Scopoli) Dysdercus cingulatus (F.)	Adult Adult Adult Adult Adult Adult	$\begin{array}{c} 0 \cdot 0 \ (4) \\ 0 \cdot 0 \ (7) \\ 0 \cdot 0 \ (10) \\ 0 \cdot 0 \ (5) \\ 0 \cdot 0 \ (5) \\ 0 \cdot 0 \ (5) \end{array}$
Lepidoptera	Bombyx mori (L.) Chilo infuscatellus Sn. Earias vitella (F.) Helicoverpa armigera (Hubner) Leucinodes orbonalis Gn. Pericallia ricini (Fab.) Plutella xylostella (L) Spodoptera litura (Fab) Spodoptera exigua (H.) Trichoplusia ni (Hb.)	Larva Larva Larva Larva Larva Larva Larva Larva Larva Larva	$100 (10) \\ 57.1 (7) \\ 60.0 (10) \\ 88.8 (9) \\ 80.0 (9) \\ 100 (10) \\ 40.0 (10) \\ 100 (30) \\ ++ \\ ++$
Diptera	Anopheles sp.	Larva	0.0 (25)

Table 2. Host range of the new isolate, Nosema sp. from Spodoptera litura

* Number of insects tested; ++ Known to cause disease (Solter, personnel communication).

Pathogenicity

The inoculation of NSD spores to third-instar larvae at 3 different concentrations of $2 \cdot 4 \times 10^2$, $2 \cdot 4 \times 10^3$, and $2 \cdot 4 \times 10^4$ spores/larva resulted in larval mortality rates of 13.33%, 46.66%, and 73.33% respectively. The LD₅₀ was 3885 spores/larva and the 95% limit lies between 2558.51–6110.06 spores/larva.

DISCUSSION

The microsporidium isolated from *S. litura* from the Dindigul, Tamil Nadu State, India is an obligatory intracellular parasite, multiplying in host cells in the form of small binucleate meronts and sporonts. Heavy infection resulted in change of larval colour to pink, frequently accompanied by swelling caused by hypertrophy of infected fat body cells. Occasionally, the body of the insect appeared deformed. The

spores were found in the faecal material of the infected larvae, which probably provide the primary means of transmission to the healthy individuals due to contamination of the food source. Though cannibalistic behaviour is uncommon in S. litura, the parasitized larvae feed on the dead diseased larvae in the laboratory, suggesting that transmission of disease could also be due to cannibalism. Although experiments were not conducted to establish the transovarial transmission, presence of spores in ovarian tissues suggested that the parasite is probably incorporated into the eggs as most of the lepidopteran Nosema, for example N. pyraustae (Andreadis, 1987), N. heliothidis (Brooks, 1968) N. bombycis (Hatakeyama and Hayasaka, 2002) and also Vairimorpha pathogenic to lepidopterans (Haque, Canning and Wright, 1999; Nageswara Rao et al. 2004). The new microsporidian isolate appears to be apansporoblastic since the spores were in direct



Fig. 1. Giemsa-stained light micrograph of merogonic and sporogonic stages of *Nosema* sp. in *Spodoptera litura*. Scale bar on A applies also to B–P. (A) Early diplokaryotic meront. (B,C) Diplokaryotic meronts with 2 and 4 nuclei. (D) Dividing octanucleate meront undergoing cytoplasmic division. (E) Meront with 6 nuclei. (F) Binucleate meront. (G, H) Dividing meronts. (I) Octanucleate meront with horseshoe-like diplokaryotic nuclei. (J) Meront with 12 horseshoe-like diplokaryotic nuclei. (K) Binucleated sporont. (L) Tetranucleate sporont. (M) Early sporoblast. (N) Sporoblast (arrow). (O) Mature spores. (P) Spores after treatment with 0.1N HCl showing diplokarya.

contact with the host cell cytoplasm possessing no sporophorous vesicle. All developmental stages were observed to be diplokaryotic. NSD shares all the salient features of the genus *Nosema* namely, apansporoblastic, and diplokaryotic developmental stages.

Sprague and Vernick (1971) characterized *Nosema* as having sporonts that divide into 2 sporoblasts and

distinguished Nosema from the genera Glugea and Encephalitozoon by their nuclear arrangement. In the present study, the sporonts were binucleate and developed into 2 sporoblasts. Our observations show the NSD to be very similar in its development to N. bombycis, as described by Ishihara (1969), and therefore we placed it in the genus Nosema. The



Fig. 2. The 50% majority consensus maximum likelihood tree of 73 non-redundant microsporidian 16S SSU rRNA gene sequences. *Amblyospora connecticus* was included as an outgroup. Numbers on the nodes indicate the number of times a particular branch was recorded per 100 bootstrap replications following 1000 replicates. The new *Nosema* sp. isolate (NSD) is shown in bold.

frequent occurrence of multinucleate meronts with 12 nuclei, however, easily distinguished NSD from *N. bombycis*.

A few reports have described the microsporidia from *S. litura*. Watanabe (1976) reported a *Nosema* sp. (NSW) from *S. litura* with development of



Fig. 3. Comparison of the 16S SSU rRNA structure of (A) Nosema sp. from Spodoptera litura (NSD) (B) Nosema bombycis (L39111) and (C) Nosema spodopterae. Only a portion of the structure (875–950) corresponding to the deleted region in NSD is shown in the figure. For all the species, the most conserved structure with similar minimum free energy ($\Delta G = -260 \pm 4$ Kcal/mol) was used.

binucleated meronts very similar to NSD. But the monosporous nature (sporont developed into single sporoblast) and uninucleate meronts of NSW distinctly differentiate it from NSD. Hsu, Hsu and Yen (1991, 1992), isolated a microsporidian species from S. litura in Taiwan and named it as Nosema spodopterae since they observed dimorphism of sporoblasts in their transmission electron microscopic study. The authors reported that one type of sporoblast possessed a plasmalemma inside of plasma membrane, and the other appeared as an amoeba-like shape without a plasmalemma within the spores. In general, the plasma membrane and plasmalemma are two names of the same structure. The internal layer of the spore wall is always referred to as a plasma membrane. The sporoblast derives its plasma membrane from the membranes of the polaroplast during the process of eversion through the polar filament during germination. No microsporidia have an amoeba-like stage in the spores. The sporoblast is the stage most difficult to fix properly for electron microscopy and electron microscopy fixation often destroys the shape of the sporoblast (Larsson, personal communication) leading to a star-like form in fixed tissues (Sokalova and Lange, 2002). Hence the

two kinds of sporoblasts reported by Hsu et al. (1992) in N. spodopterae are probably an artifact arising due to fixation. Though the size of NSD sporoblasts ranges from 4.52 to 6.52 μ m, the shape of the sporoblasts did not vary. The fresh spore size of NSD $(3.91\pm0.012\times1.91\pm0.014\,\mu\text{m})$ was similar to that of N. spodopterae $(4.0 \pm 0.26 \times 1.9 \pm 0.12 \,\mu\text{m})$. Li and Wenn (1987) described a microsporidium, Nosema liturae, obtained from S. litura collected from China. The fresh spore size of N. liturae $(4.34 \pm 0.27 \times$ $1.99 \pm 0.14 \,\mu\text{m}$) exceeds the size of NSD ($3.91 \pm$ $0.012 \times 1.91 \pm 0.014 \,\mu$ m). Bombyx mori is refractive to N. liturae, while the NSD is highly pathogenic to the same host species. The multinucleated meronts with up to 12 compact horseshoe-like nuclei reported in NSD were not reported in N. spodopterae and N. liturae.

Since many attempts to amplify large subunit rDNA did not succeed across all microsporidian species (Vossbrinck *et al.* 1993; Kent *et al.* 1996), 16S SSU rDNA has become the standard sequence for molecular characterization of microsporidia (Weiss and Vossbrinck, 1999; Vossbrinck and Debrunner-Vossbrinck, 2005). Here we aligned the SSU rRNA gene sequence of NSD with those of other known microsporidia that infect lepidopteran as well as nonlepidopteran hosts. The results lead us to confirm that NSD is the member of true *Nosema* namely, *N. bombycis* complex and belongs to the genus *Nosema* Nageli, 1857.

Comparison of the 16S rRNA gene sequence of NSD with the corresponding sequence from several Nosema species showed that the parasite is closer to N. bombycis. We used N. bombycis 'L39111' as a reference strain in the present study since it is the most studied isolate for phylogenetic analyses (Baker et al. 1995; Canning et al. 2002; Choi et al. 2002; Vossbrinck and Debrunner-Vossbrinck, 2005). In NSD, the absence of 3 nucleotides at position 918 as compared to L39111 and other N. bombycis suggested a possible difference between these two isolates. The published sequence of N. spodopterae is, in fact, closer to that of L39111 than to NSD, suggesting that the new isolate is possibly a new species of Nosema from S. litura. Consistently, the secondary structure of N. bombycis and N. spodopterae 16S SSU rRNA was identical (Huang et al. 2004; Tsai, Huang and Wang, 2005) whereas the structure of NSD was different. The significance of these structural differences needs to be experimentally verified, which is out of purview of the present manuscript. These observations compel us to consider that the microsporidian isolate NSD as a new species, confirmation of which awaits further ultrastructural characterization.

The pathogenicity of NSD appears to be proportional to the inoculation dosage. The LD₅₀ for larvae inoculated in third-instar was calculated to be 3.88×10^3 spores/larva at 20 days p.i. and is comparable with the published reports of other *Nosema* isolates from *S. litura*. Tsai and Wang (2001) obtained an LD₅₀ of 1.13×10^4 spores/larva after 21 days when *S. litura* third-instar larvae were treated with *N. spodopterae* spores. The LD₅₀ of Taiwan isolate in third-instar *S. litura* larvae was lower, 1.78×10^3 spores/larvae at 30 days p.i. (Tsai *et al.* 2003).

Host specificity is an important consideration for pathogens that are used for biological control. Microsporidia usually do not infect species in other insect orders when larvae are fed spores, but interordinal infections have been reported in laboratory infectivity studies (Fantham and Porter, 1958; Undeen and Maddox, 1973). All the lepidopteran hosts tested were susceptible to NSD indicating that this species appear to have a wider host range under laboratory conditions. Previous host specificity studies under lab conditions (Undeen and Maddox, 1973; Solter, Maddox and McManus, 1997; Solter and Maddox, 1998) suggest that terrestrial microsporidia have a broader laboratory host range within the taxonomic order of the natural host, but atypical microsporidia/host interactions suggested that the pathogens often do not reproduce optimally in

non-target hosts, and data from surveys of microsporidian pathogens of various native and introduced insect hosts from field collections suggest that the ecological host ranges of entomopathogenic microsporidia are relatively narrow (Campbell and Podgwaite, 1971; Andreadis *et al.* 1983; Siegal *et al.* 1988; Jeffords *et al.* 1989). In the present study, lower percentages of infection rate observed in some lepidopteran species like *P. xylostella* and *C. infuscatellus* also reflected that these may be nontarget hosts to NSD under field conditions. Hence, the use of NSD as a biological control agent against *S. litura* should be investigated.

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