The gut commensal bacterium Enterococcus faecalis LX10 contributes to defending against Nosema bombycis infection in Bombyx mori

Xiancui Zhang,a Huihui Feng,a Jintao He,a Xili Liang,a Nan Zhang,a Yongqi Shao,a Fan Zhangb*, and Xingmeng Lu*a

Abstract

BACKGROUND: Microsporidia, a group of obligate intracellular fungal-related parasites, have been used as efficient biocontrol agents for agriculture and forestry pests due to their host specificity and transovarial transmission. They mainly infect insect pests through the intestinal tract, but the interactions between microsporidia and the gut microbiota of the host have not been well demonstrated.

RESULTS: Based on the microsporidia–Bombyx mori model, we report that the susceptibility of silkworms to exposure to the microsporidium Nosema bombycis was both dose and time dependent. Comparative analyses of the silkworm gut microbiome revealed substantially increased abundance of Enterococcus belonging to Firmicutes after N. bombycis infection. Furthermore, a bacterial strain (LX10) was obtained from the gut of B. mori and identified as Enterococcus faecalis based on 16S rRNA sequence analysis. E. faecalis LX10 reduced the N. bombycis spore germination rate and the infection efficiency in vitro and in vivo, as confirmed by bioassay tests and histopathological analyses. In addition, after simultaneous oral feeding with E. faecalis LX10 and N. bombycis, gene (Akrin, Cecropin A, Mesh, Ssk, DUOX and NOS) expression, hydrogen peroxide and nitric oxide levels, and glutathione S-transferase (GST) activity showed different degrees of recovery and correction compared with those under N. bombycis infection alone. Finally, the enterococcin LX protein was identified from sterile LX10 fermentation liquid based on liquid chromatography–tandem mass spectrometry (LC–MS/MS) analysis.

CONCLUSION: Altogether, the results revealed that E. faecalis LX10 with anti-N. bombycis activity might play an important role in protecting silkworms from microsporidia. Removal of these specific commensal bacteria with antibiotics and utilization of transgenic symbiotic systems may effectively improve the biocontrol value of microsporidia.

Supporting information may be found in the online version of this article.

Keywords: Bombyx mori; microsporidia; Enterococcus faecalis; anti-Nosema bombycis activity; protein

1 INTRODUCTION

The silkworm is an important model for studies on insect physiology and control of lepidopterous pests due to its relatively small body size and ease of rearing.1 Microsporidians are a group of widely distributed pathogens that can invade a broad range of hosts, including immunocompromised humans.2 Nosema bombycis is the first described microsporidium species recognized as the etiological agent of silkworm pebrine disease.3 To date, pebrine disease is the only disease with a mandatory quarantine in the silk production industry due to its destructive consequences.4 The microsporidian pathogens are also considered effective biological control agents for mosquitoes, gypsy moth, locust, and corn borer red imported fire ants.5–7 In 1980, Paranosema locustae became the first microsporidium to be registered and successfully commercially marketed in the United States (USEPA 2000). It is still being sold for bait-and-spray applications for Mormon cricket and grasshopper control due to the high grasshopper mortality associated with products, such as Nolo Bait™ and Semaspore™.8,9 Similar results have been reported in Argentina and China where government-subsidized, large-scale applications were deemed adequate by the growers because of wide host ranges and high transmission efficiency.10,11

As an important intestinal pathogen, N. bombycis can be transmitted horizontally through per os infection as well as vertically

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from adults to offspring. The pathogen *N. bombycis* is well adapted to the silkworm gut environment, where it germinates and penetrates the gut epithelium, absorbing nutrients and energy from the host and exporting cytotoxic compounds through the cell membrane. This species has a sophisticated and unique mechanism for infecting host cells, involving a rigid spore wall and a coiled polar tube joined to an anchoring disc at the apical part of the spore. With appropriate stimuli, mature microsporidian spores rapidly discharge the infective polar tube, come into contact with the surface of the cell membrane, and transfer the sporoplasm into the cell cytoplasm, where proliferation and the next round of spore production occurs. *N. bombycis* then proliferates inside the host, leading to chronic damage to larval organs and tissues, including the gut, muscles, silk glands, Malpighian tubules, and ultimately to death. Nevertheless, it is not surprising that silkworms have evolved effective anti-*N. bombycis* strategies, including its innate immune system as well as the chemical and physical barriers present in the gut, such as a strong alkaline digestion mechanism. *N. bombycis* infection of *Bombyx mori* larvae activates the JAK/STAT and Toll pathways in the midgut, which induce the up-regulation of the expression of innate immune-related defense proteins involved in humoral and cellular immunity, including moricins, lebocins and lysozymes, to resist invasion by microsporidia. In addition, the production of the prophenoloxidase (PPO) by the gut epidermal cells of the silkworm is also suppressed after microsporidian infection due to the induction of melanization, thus preventing pathogens from entering the surrounding habitat.

Similar to most lepidopteran larvae, silkworm larvae possess a relatively simple gut without specialized substructures; however, a sufficient gut bacterial population exceeding $10^7$ CFU/mL is clearly present. Recent surveys of the taxonomic composition of the gut bacteria reveal a relatively simple but specific community dominated by *Enterococcus*, *Lactococcus*, *Bacillus*, *Stenotrophomonas* and *Pseudomonas*. The indigenous microbiota is being increasingly recognized as an important component associated with biological fitness, detoxication, vitamin synthesis, reproductive regulation and pathogen resistance. Comparative metabolomics has revealed that *Stenotrophomonas* species provide essential amino acids to the host and hence increases host tolerance to organophosphate insecticides. A study confirmed that *Enterococcus* spp. decreased the toxicity of *Bacillus thuringiensis* δ-endotoxin in the silkworm larva gut. Some *Enterococcus* species have been identified in the gut of insects, fish, birds, and mammals, and many of these *Enterococcus* species have been utilized as unique livestock probiotics for decades. Additionally, these commensal intestinal bacteria can assist different insect hosts with pathogen resistance in different insects, such as by secretion of antimicrobial substances and activation of host defense pathways. For example, for *Aedes aegypti* and *Anopheles gambiae*, the creation of microbe-free aseptic strain with antibiotics renders the mosquitoes more susceptible to infection by the apicomplexan parasite *Plasmodium falciparum*. In addition, the intestinal bacterium *Pseudomonas aeruginosa* BGF-2 has been shown to decrease the mortality of cockroaches after treatment with *Beauveria bassiana* and to restrict conidial germination and hyphal growth by producing antifungal proteins. Moreover, a stable class Ia bacteriocin (mundtin KS) from the gut bacterium *Enterococcus mundtii* of *Spodoptera littoralis* suppressed the proliferation of pathogenic bacteria *in vitro*. Furthermore, 70% of pests are lepidopteran species, so exploration of the relationship between the commensal bacteria of silkworms and the fungal parasites *N. bombycis* in the lepidopteran model, *B. mori* is urgently needed. The aim of the current study was therefore to investigate the direct effects of *N. bombycis* infection on the community structure of the gut microbiota. We isolated the bacterial strain *E. faecalis* LX10 with anti-*N. bombycis* activity from the *B. mori* gut. Moreover, we investigated the effect of supplementation with *E. faecalis* LX10 on immune gene expression, hydrogen peroxide (H$_2$O$_2$) and nitric oxide (NO) levels and glutathione S-transferase (GST) activity of silkworm larvae challenged with *N. bombycis*.

## 2 MATERIALS AND METHODS

### 2.1 Silkworms and microsporidia

The hybrid *B. mori* strain HaoyuexJingsong was provided by the Silkworm Germplasm Bank of the Cathaya Group, Chun an, Zhejiang, China (118°71’N, 29°36’E). All insects were reared under standard conditions (25 ± 1 °C, 70 ± 5% humidity) in plastic boxes (50 cm × 25 cm × 10 cm) using fresh mulberry leaves collected from a mulberry field at Zhejiang University, China (30°18’N, 120°04’E). Additionally, to prepare germ-free (GF) larvae, silkworm eggs were surface sterilized using 1% sodium hypochlorite (NaClO) solution and then rinsed with sterile water. The eggs were subsequently immersed in 70% ethanol followed by sterile water. The GF larvae were reared with an autoclaved artificial diet. To validate the GF conditions, fecal samples were examined by Petri dish culture.

*N. bombycis* ZJU1 spores originally obtained from infected silkworms in Zhejiang, China, were produced and harvested at the Institute of Sericulture and Apiculture, Zhejiang University, and propagated and purified in laboratory-reared silkworms adults at a temperature of 25 ± 1 °C by centrifugation over Percoll (pH=7).

### 2.2 Susceptibility of silkworms to *N. bombycis* applied through feeding

HaoyuexJingsong silkworms were equally divided into four groups [control (CK), 10$^5$, 10$^4$, and 10$^3$] after the fourth exuviation (fifth-instar larvae). Newly molted silkworm larvae were challenged by feeding on mulberry leaves (15 cm × 15 cm) that were previously artificially besmeared with *N. bombycis* spores at different concentrations (10$^3$ spores/mL, 10$^4$ spores/mL, and 10$^5$ spores/mL; 25 μL/larvae). The CK group larvae were reared with fresh mulberry leaves with the same dosage of sterile water. Briefly, 15 samples (*n* = 15) were collected from each group for DNA extraction and gut microbiota analysis (a single individual insect for each sample) at four time points (0, 2, 4, and 6 days) post-infection (dpi). The gut samples were homogenized in a Precellys-24 homogenizer (Bertin Technologies, Aix-en, France) at 6000 x g for 120 s. Bacterial DNA was extracted using a MasterPure™ Purification Kit (Epicentre, Madison, WI, USA) following the manufacturer’s protocol. Moreover, a negative control without gut tissue was processed simultaneously with the same DNA extraction method to control for contamination with the reagent and other substances. The concentration and quality of the DNA were measured by a 2100 Bioanalyzer (Agilent, Palo Alto, CA, USA).

The absolute quantification of *N. bombycis* (*n* = 15) was conducted by real-time quantitative polymerase chain reaction (qPCR) analysis for each silkworm gut. The primers ssu1092F and ssu1227R were designed to amplify the *N. bombycis* small-subunit rRNA gene fragment from position 1092 to position
Enterococcus faecalis Inhibits Microsporidian Infection in Bombyx mori

1227 (GenBank accession no. EU864525.1). Then, a standard curve was constructed with pure N. bombycis DNA (1 × 10^6 to 1 × 10^10 copies/µL) as the standard substance. All runs were performed using SYBR qPCR Master Mix (Vazyme Biotech, Nanjing, China) and a Roche LightCycler 480 system (Roche, Basel, Switzerland). In addition, sterile water served as a negative control. The qPCR was conducted using the following program: 95 °C for 5 min and 40 cycles of 95 °C for 10 s, 60 °C for 10 s and 72 °C for 10 s. A melting curve (from 65 to 92 °C, 0.5 °C/s) was analyzed to ensure that the reaction produced a single product. The gene copies of N. bombycis were calculated by comparing Cq values to the standard curve.

An analytical balance (Mettler Toledo, Greifensee, Switzerland) was used to measure the larval mass at 2, 4, and 6 dpi for the experimental (N. bombycis treatment, NB) and CK groups (n = 15). The pH measurements were performed using a pH micro-electrode (Unisense, Aarhus, Denmark) with a tip diameter of 20 to 30 µm (Unisense). As soon as the larvae pupated, they were transferred into labeled 2 cm-wide and 5 cm-long sterilized cages at a temperature of 25 ± 1 °C and humidity of 70% ± 1%. The whole cocoon weight, cocoon length, cocoon width and cocoon shell weight were recorded after cocooning (n = 60).

2.3 Gut microbiota analysis by 16S rRNA gene sequencing

Gut DNA was extracted by following the method described in Section 2.2. To prepare an Illumina sequencing library, a region encompassing the V3–V4 hypervariable regions of the 16S rRNA gene was amplified from 1 µL of purified DNA using the universal primers 338F and 806R, which are shown in Supporting Information Table S2. Agarose gel electrophoresis was used to estimate the product size (~450 bp). All 16S rRNA products were purified and pooled together after gel purification by a MiniBEST Agarose Gel DNA Extraction Kit (Takara, Dalian, China) for the Illumina MiSeq platform (San Diego, CA, USA), and paired-end sequencing was performed by Shanghai Biotechnology (Majorbio, Shanghai, China). In total, 8,908,622 raw reads were obtained and preprocessed as previously described. Briefly, the demultiplexed paired-end sequence reads were assembled using DADA2. Thus, 165 rRNA gene sequencing generated 120,000–450,000 sequences per sample after preprocessing. Sequences assigned to chloroplasts and mitochondria were removed from the amplicon sequence variant (ASV) tables. We obtained high-quality processed sequences with a mean length of 410 bp and an average of 49,492 sequences per sample. To minimize bias due to sequencing depth, we normalized the sequence number to the lowest sequence number by rarefaction. High coverage values (average = 99%) were obtained for sequences in all samples, indicating that the sequencing depth was sufficient. To investigate the gut bacterial community structure, Shannon and Faith’s Phylogenetic Diversity (Faith’s PD) indices were estimated. Bray–Curtis dissimilarities were determined and visualized via principal coordinate analysis (PCoA) for community comparison. The relationships between gut microbiota structure and environmental variables (N. bombycis concentration, inoculation time) were determined with the use of multivariate redundancy analysis (RDA) and canonical correspondence analysis (CCA). Network analyses were used to explore the cooccurrence patterns of gut bacterial microorganisms in each niche. The Spearman’s correlation between two ASVs was considered statistically robust if the P value was < 0.01 and Spearman’s correlation coefficient (r) was > 0.6. The resulting adjacency matrices were converted to network objects by the R package igraph. Network summary statistics and node centrality measures were calculated to compare network properties across microbial species (degree, average degree, average path length, modularity, diameter, and transitivity). Networks were visualized using the interactive Gephi platform.

Receiver operator characteristic (ROC) curve analysis for sample classification was performed using the random forest algorithm. The area under the curve (AUC) was calculated to measure the predictive accuracy. Moreover, linear discriminant analysis (LDA) effect size (LEfSe) analysis was performed using the normalized ASV table. Variation analyses were performed using Fisher’s least significant difference (LSD) followed by one-way analysis of variance (ANOVA) or the non-parametric Kruskal–Wallis test. Finally, Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt2) was employed for stringent predictions of microbial functional metabolic pathways from the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis. Sequencing data were deposited in the National Center for Biotechnology Information (NCBI) Short Read Archive (SRA) under accession number PRJNA763083.

Absolute quantification of the number of bacteria was conducted by qPCR of 16S rRNA gene copies of total bacteria (n = 15) and Enterococcus (n = 15) using the DB200Q (all bacteria) and EN144Q (Enterococcus) primers. The number of copies was routinely deduced by comparing the Cq values to a standard curve generated from a ten-fold serial dilution series of the full-length 16S rRNA genes.

2.4 Enterococcus isolation and anti-N. bombycis effects in vitro and in vivo

Enterococcus bacteria were isolated from the normal fifth-instar B. mori larval gut using Enterococcus selective agar with 1% 2,3,5-triphenyl tetrazolium chloride (TTC) in the medium. Molecular analysis was performed by isolating genomic DNA and amplifying the bacterial 16S rRNA genes using universal primers LoFp forward primer, LoFp reverse primer, and 144Q (E. faecalis strain was cultured in de Man–Rogosa–Sharpe broth (Hopebio, Qingdao, China) at 37 °C for 24 h. The supernatant (500 mL) was collected after centrifugation at 10,000 × g for 10 min and filtered through a 0.22 µm polyvinylidene fluoride (PVDF) membrane (Millipore, Billerica, MA, USA) to obtain cell-free fermentation liquid. Crude proteins were precipitated with 20–80% saturated ammonium sulfate (NH₄)₂SO₄. The proteins were dialyzed with regenerated cellulose membrane tubing (3.5 kDa; Spectrum) in distilled water. Then, the crude proteins from E. faecalis LX10 supernatants (SMP) were concentrated via ultrafiltration with Amicon Ultra 3 kDa (< 3 K and < 3 K SMP) and 10 kDa (< 10 K and > 10 K SMP) cellulose ultrafiltration device (Millipore) to remove proteins < 3 KDa in size, according to the manufacturer’s instructions. Crude protein from MRS medium (MRS) was obtained in the same way as a negative control. The protein concentration was quantified using a BCA protein detection kit (Thermo Fisher Scientific, Waltham, MA, USA) and adjusted for equal loading. The effect of pH on the inhibitory substance was tested by adjusting each of the crude protein samples to pH 5.0 to 10.0 with 1 mol/L sodium hydroxide (NaOH) or 1 mol/L hydrochloric acid (HCl). The samples were readjusted to pH 6.0.
after 30 min of incubation and tested for antimicrobial activity. Moreover, the effect of temperature was tested by heating the crude protein at 40, 60, 80 and 100 °C for 30 min, and the residual activity was measured.

Anti-\textit{N. bombycis} activity assays were performed both \textit{in vitro} and \textit{in vivo}. In the \textit{in vitro} study, 20 \muL of \textit{N. bombycis} ZJU1 spore suspension (10^{7} spores/mL) was aliquoted into 1.5 mL centrifuge tubes, and 30 \muL of crude protein (2 mg/mL) with different molecular weight ranges (< 3 K, > 3 K, < 10 K, and > 10 K SMP) was added to the tubes in the treatment group. Additionally, 30 \muL of phosphate buffer (pH 6) was used as a control. The control and treatment group samples were treated with 150 \muL of GKK germination buffer ([0.05 mol/L glycine, 0.05 mol/L potassium hydroxide (KOH), and 0.375 mol/L potassium chloride (KCl); pH 10.5]), and the samples were incubated at 27 °C for 60 min. The germination rate was monitored by measuring the absorbance at 625 nm (OD_{625}). The calculation formula was as follows: germination percentage = [(germination initation liquid OD_{625} – germination termination liquid OD_{625})/germination initation liquid OD_{625}] × 100%.

For \textit{in vivo} assessment of anti-\textit{N. bombycis} activity, fifth-instar GF silkworm larvae were divided into four groups (CK, EF, NB, EF + NB) (n = 15). Each silkworm was fed 25 \muL of \textit{E. faecalis} LX10 cell suspension (1 \times 10^{7} CFU/mL) and 25 \muL of \textit{N. bombycis} suspension (1 \times 10^{6} spores/mL) (EF + NB); 25 \muL of \textit{E. faecalis} LX10 cell suspension (1 \times 10^{7} CFU/mL) and 25 \muL of sterile water (EF); or 25 \muL of \textit{N. bombycis} suspension (1 \times 10^{6} spores/mL) and 25 \muL of sterile water (NB), with GF larvae serving as a negative control (CK). After 6 days, randomly selected gut samples were processed for scanning electron microscopy (SEM) and histopathological analysis, whereas the remaining sample was homogenized for assessment of the \textit{N. bombycis} burden via qPCR after 2, 4, and 6 days. For SEM analysis, the gut samples were fixed in 25% glutaraldehyde at 4 °C overnight. Then several gut samples were washed in phosphate-buffered saline (PBS) three times and post-fixed in 1% osmium tetroxide for 2 h. Then, the tissue with serial dilutions of ethanol (30% to 100%) were dehydrated. The samples were dried using a critical point dryer, mounted, and sputter coated with gold/palladium. SEM was conducted on gold-sputtered samples with a Hitachi S8080 scanning electron microscope (Tokyo, Japan). Moreover, the gut epithelial samples were fixed in polyformaldehyde fixing solution for histopathological analysis. Periodic acid–Schiff (PAS) staining was performed to evaluate pathological injury.

### 2.5 Anti-\textit{N. bombycis} protein identification for LC–MS/MS

The active crude proteins (3–10 K) were analyzed by 15% sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) to determine the components of the proteins samples and to estimate their molecular masses. Protein binding was visualized by staining with Coomassie Blue 250. The gel bands were subjected to in-gel digestion followed by alkylation and trypsin digestion at 37 °C for 2 h. An aliquot of the tryptic digest (in 0.1% formic acid–0.1% acetonitrile in acetonitrile–water) was analyzed by liquid chromatography tandem mass spectrometry (LC–MS/MS) on a Q Exactive Orbitrap Fusion Trubrid mass spectrometer that was coupled to an Easy LLC system (Thermo Fisher Scientific, Waltham, MA, USA). All raw MS and MS/MS files were processed and analyzed using Mascot 2.2 software (Matrix Science, London, UK) against the Protein Homology/Analogy Recognition Engine version 2.0 (Phyre2) database. The search parameters were set as follows: variable modification, oxidation; fixed modification, carbamidomethylation; peptide tolerance, ± 20 ppm; number of missed cleavages, 2; fragment mass tolerance, ± 0.1 Da; and filtered by score ≥ 20. The MS/MS data of the anti-\textit{N. bombycis} proteins were retrieved and aligned against both the NCBI BLAST database and the protein sequence database of \textit{E. faecalis} (NCBI, Uniprot). The three-dimensional structure prediction models were illustrated using Phyre2 database (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index).

### 2.6 Gene expression analysis and enzyme activity assays

To evaluate the effect of \textit{N. bombycis} infection (CK, 10^{3}, 10^{4}, and 10^{5} spores/mL) on host immunity and whether \textit{E. faecalis} LX10 has a protective effect on silkworm infection, \textit{B. mori} larvae were orally inoculated with an \textit{E. faecalis} LX10 suspension (1 \times 10^{7} CFU/mL, Ef), \textit{N. bombycis} (1 \times 10^{5} spores/mL, Nb), or \textit{E. faecalis} LX10 and \textit{N. bombycis} (Ef + Nb), with healthy \textit{B. mori} GF larvae serving as a negative control (CK). Then, the expression levels of immune-related genes (cerecin, Akirin, Mesh, Ssk, DUOX, and NOS) were determined in the gut at 2, 4, and 6 days post-exposure (n = 5). Total RNA was extracted with a Promega Eastep® Total RNA Super Extraction Kit (Beijing Biotec. Co., Ltd, Beijing, China) according to the manufacturer’s protocol. Total RNA (1 μg) was reverse transcribed to complementary DNA (cDNA) using HiScript® II Q RT SuperMix R223-01 (Vazyme Biotech) according to the manufacturer’s guidelines. The reverse transcription qPCR (RT-qPCR) was performed as described earlier. The expression level of the housekeeping gene RP49 was used as an endogenous control. The sequences of gene-specific primers used for transcript quantification are listed in Table S2. The 2^{ΔΔC_{t}} method was used to analyze the relative transcript levels of the genes of interest.

The H_{2}O_{2} and NO levels and GST enzyme-mediated protective response in the silkworm gut were determined (n = 5). The silkworm gut was isolated and immediately homogenized in ice-cold PBS (10 mmol/L). The gut homogenates were centrifuged at 10 000 \times g for 10 min at 4 °C, and then, the supernatants were collected for different measurements. The H_{2}O_{2} and NO levels were assayed using a Hydrogen Peroxide Assay Kit (serial no. S0038; Beyotime Biotechnology, Shanghai, China) and Total Nitric Oxide Assay Kit (serial no. S0023) (Beyotime Biotechnology). The protein content was determined using a BCA kit according to the manufacturer’s protocol (Thermo Fisher Scientific, Waltham, MA, USA). GST activity was analyzed with a GST Activity Detection Kit (Sangon Biotech, Shanghai, China).

### 2.7 Statistical analysis

Statistical analyses were carried out using GraphPad Prism (version 9.0) and Statistical Package for the Social Sciences, version 20.0 (SPSS, Chicago, IL, USA). All the results are presented as the mean ± standard error (SE). Parametric/non-parametric statistical analyses were performed after the data were checked for normality and homogeneity of variance. One-way ANOVA, Tukey’s post hoc test or Student’s t test was performed to compare the absolute abundances of bacteria and \textit{N. bombycis}, larval mass, pH, whole cocoon weight, cocoon length, cocoon width, cocoon shell weight, relative germination rate, gene expression changes, and enzyme activities between the experimental and control groups. In each case, the type of test is stated before the P value, and P ≤ 0.05 was considered statistically significant. All experiments were repeated at least three times. The software used included GraphPad Prism 9.0 and Python and R packages, run as standalone software on Windows 10.
3 RESULTS

3.1 Susceptibility of silkworms to N. bombycis

The susceptibility of silkworms to N. bombycis treatment was both dose and time dependent, and this phenomenon was more pronounced when the silkworms were treated with high concentrations of N. bombycis (Fig. 1(A)). N. bombycis was detected in only the gut tissues of 10^6 and 10^9 spore/mL-treated silkworms but not in those in the CK group and 10^3 spore/mL-treated group. For example, at 6 dpi, the number of spores in the gut reached 0.89 × 10^6 and 5.25 × 10^8 spores/gut compared to that at 2 days (0 versus 5.37 × 10^4 spores/gut) at 10^6 and 10^9 spores/mL (t = −40.18, df = 14, P = 0.006 for 10^6, t = 18.68, df = 14, P < 0.001 for 10^9) (Fig. 1(A)).

The larval mass and pH are shown in Fig. 1(B,C). After treatment with N. bombycis, the larval mass decreased significantly (P < 0.05), especially at concentrations of 1 × 10^6 and 1 × 10^9 spores/mL (Fig. 1(B)). There was no significant difference in larval mass between the N. bombycis-fed and CK groups on Day 2 (P = 0.4916, F_{(3,56)} = 0.8138, Fig. 1(B)). With increasing N. bombycis concentrations and inoculation time, the difference gradually increased (P < 0.0001, F_{(3,56)} = 8.534 for 4 days, P < 0.0001, F_{(3,56)} = 27.74 for 6 days) (Fig. 1(B)). The gut pH showed a similar trend (F_{(11,48)} = 40.37, P < 0.0001) (Fig. 1(C)). The gut exhibited macroscopic white swelling and wrinkles at 6 days after the treatment Fig. 1(D). The fitness-related parameters reported in Fig. 1(E–H) show that the whole-cocoon weight, cocoon length, cocoon width, and cocoon shell weight decreased significantly after N. bombycis treatment (P < 0.05).

3.2 N. bombycis infection causes dysbiosis of the gut microbiota

A total of 180 intestinal samples were prospectively collected, and 18 samples failed to generate data and were removed from the following analyses. In total, 1081 species, 754 genera, 453 family, 261 order, 118 class, and 39 phyla were detected. To characterize the effect of N. bombycis on gut microbiome diversity, different measures of alpha diversity were applied based on 16S rRNA sequences. Briefly, the Shannon and Faith’s PD indices were higher in the non-infected silkworms (CK group) than in the N. bombycis-infected silkworms at 2, 4, and 6 dpi (P < 0.05). Overall, N. bombycis infection resulted in decreased gut microbial richness and diversity (Fig. 2(A,B)). With prolonged infection, PCoA showed that the gut bacterial communities of gut regions with different inoculation concentrations clustered separately (Fig. 2(C–E)). A random distribution of the samples was found despite the gut microbiota being somewhat similar among individuals at 2 days after inoculation Fig. 2(C).

We next investigated the variability in bacterial communities in relation to N. bombycis infection. The bacterial phyla Proteobacteria, Actinobacteria, Firmicutes, Bacteroidota, and Desulphobacterota, together accounting for up to 90% of sequences on average, were the five most dominant taxa (Fig. 3(A)). Compared with that in healthy individuals, the abundance of Firmicutes generally decreased in a time- and dose-dependent manner in the treatment groups. At 6 days after inoculation, the abundances of Firmicutes in the CK, 10^3, 10^6, and 10^9 groups showed a decreasing trend (58.33%, 58.51%, 56.29% and 17.41%) (P < 0.05, Fig. 3(A)). The qPCR confirmed that the total bacterial load in infected silkworms after infection (1.21 × 10^{11} copies/gut) was significantly higher than that in the CK group (0.58 × 10^{9} copies/gut) and that in silkworms in the 10^9 group at 6 dpi (t = 6.199, df = 28, P < 0.0001) (Supporting Information Fig. S1). Correlation analysis of the microbial abundance at the phylum level was used to construct networks, wherein the network parameters represented the microbial communities and the correlations of their abundances. Compared to the CK network, those of the treatment groups, especially the 10^9 group,
showed a higher average path length (6.10), modularity (0.80), and diameter (10.25) (Fig. S2(A)), while the CK network showed a higher degree (496), higher average degree (3.16), shorter average path length (3.37), and lower network diameter (6.99). These finding, together with the network global properties, indicated that the CK group exhibited much closer interconnections than the treatment group at the phylum level (Fig. S2(A)).

At the genus level, the most dominant genera detected within the gut microbiome were *Rhodococcus*, *Escherichia-Shigella*, *Enterococcus*, *Sphingomonas*, *Sediminibacterium*, *Ralstonia*, *Lactobacillus*, and *Sphingobium* and *Lactobacillus*, which accounted for 80% of the overall abundance of bacteria in the silkworm gut (Figs 3(B) and S2(B)). Coexistence and competition relationships between gut bacteria were found in the silkworm. The abundance of *Enterococcus* was negatively correlated with that of the most predominant genera (Fig. S2(C)). LEfSe analysis confirmed that the gut microbiota composition at different inoculation times was consistent, but the abundance differed. For example, the relative abundances of *Enterococcus*, *Escherichia-Shigella*, *Staphylococcus* and *Glutamicibacter* were higher at 6 dpi (Fig. S2(D)).

With prolonged infection, the abundance of *Enterococcus* increased gradually ($F_{(3,56)} = 40.04$, $P < 0.0001$ for 2 days; $F_{(3,56)} = 5.408$, $P < 0.0024$ for 4 days; $F_{(3,56)} = 25.35$, $P < 0.0001$ for 6 days). With treatment at a concentration of $10^9$ spores/mL (6 days), $10^{10}$ 16S rRNA gene copies of *Enterococcus* were observed in silkworm intestinal contents (Fig. 3(C)). There was a statistically significant positive correlation between the total *N. bombycis* number and the *Enterococcus* number in the gut ($R^2 = 0.14$, $P = 0.002$ for 2 days; $R^2 = 0.54$, $P = 0.8$ for 4 days; $R^2 = 0.77$, $P < 0.001$ for 6 days; Fig. 3(D)). The results of RDA and CCA of the environmental factors and the bacterial microbial community in all the treatments are shown in Fig. 3(E) and S3. The *Enterococcus* abundance showed a positive correlation with infection time ($R^2 = 0.4398$, $P = 0.001$) and *N. bombycis* concentration ($R^2 = 0.3121$, $P = 0.001$) (Fig. 3, Table S1). In addition, we explored the predicted functions of the gut microbiota across all groups in our study by PICRUSt analyses Fig. 3(F). Across all samples, the enriched KEGG pathways included carbohydrate metabolism, amino acid metabolism, and energy metabolism. Interestingly, almost all the KEGG pathways of the

**Figure 2.** Spatial structure of the *B. mori* gut microbiota. (A,B) Alpha diversity and (D,E) beta diversity were investigated. Alpha diversity (Shannon and Faith’s PD) of bacterial communities (based on ASVs) at different spore concentrations (CK, $10^3$, $10^6$, and $10^9$ spores/mL) and infection times (2, 4, and 6 days). Different letters describe significant differences at $P = 0.05$. One-way ANOVA with Tukey’s multiple comparisons test and unpaired Student’s $t$ test. PCoA plot showing variation in community structure among different infection concentrations (permutational multivariate analysis of variance (PERMANOVA) test with 999 permutations, $P > 0.05$) based on Bray–Curtis distance. Each point represents an individual sample.
N. bombycis treatment group were significantly down-regulated compared with those of the CK group ($P < 0.01$) (Fig. 3(F)). Finally, the Sloan neutral model showed a more niche-based process of the gut microbiome in N. bombycis-treated silkworm larvae (Fig. S4(A)). The AUC at 4 dpi (0.84) and 6 dpi (0.82) was significantly greater than that at 2 dpi (0.61) ($P < 0.001$) (Fig. S4(B–D)).

### 3.3 Anti-N. bombycis assay of Enterococcus

Amplification of the LX10 16S rRNA sequence yielded a 1200-bp PCR product, and the sequence (GenBank) showed 98% similarity with that of E. faecalis Feb-67 (MH385355) (Fig. S5). The proteins (3–10 K SMP) precipitated by (NH$_4$)$_2$SO$_4$ from the supernatant showed strong inhibitory activity (Fig. 4(A)). The protein concentration was quantified using the Bradford reagent and adjusted...
The inhibitory activity of *Enterococcus* showed that the spore germination rate was in the order CK (GKK) > proteinase K > (< 3 K SMP) > MRS > (> 10 K SMP) > (> 3 K SMP) > (< 10 K SMP) (Fig. 4(A)). The germination rate with the precipitated proteins (> 3 K SMP, < 10 K SMP) was significantly decreased to 10.44 ± 0.63% (t = 8.804, df = 10, P = 0.0009) and 9.45 ± 1.22% (t = 8.887, df = 10, P = 0.0009) compared to that of the CK group (42.06 ± 2.72%) (Fig. 4(A)). The inhibitory activity was absent in supernatants treated with proteinase K.

In the *in vivo* study, a significant decrease in the silkworm gut *N. bombycis* burden was evident in the *E. faecalis* LX10-treated (Nb) group relative to the group inoculated only with *N. bombycis* (Fig. 4(B)). DNA was extracted from the intestinal epithelium, and the *N. bombycis* burden was estimated by qPCR amplification after treatment with *E. faecalis* LX10 (10^7 spores/mL) (Ef), *N. bombycis* (10^9 spores/mL) (Nb), or *N. bombycis* and *E. faecalis* LX10 (Ef + Nb). Healthy silkworms with the same volume of sterilized water were used as the control group (CK). The gut of silkworms was excised for SEM and histological examination (PAS staining) of the CK (C1, C2), Ef (D1, D2), Nb (E1, E2), and Ef + Nb (F1, F2) group silkworms. The arrows indicate that the *N. bombycis* spores proliferated in epithelial cells. Cc, columnar cell; Gc, goblet cell; Lu, lumen; Mu, muscle layer; Mi, microvilli; Va, vacuoles. Statistically significant differences were calculated by using one-way ANOVA (*P < 0.05).

### 3.4 Antimicrobial protein identification

The molecular weight of the crude protein (enterococcin LX) (3–10 K) was determined by SDS-PAGE (Fig. S6(A)). Through matrix-assisted laser desorption/ionization-time-of-flight-mass spectrometry (MALDI-TOF-MS) analysis, the protein was confirmed as enterococcin EntV (*E. faecalis*) (NCBI reference sequence: QHN67019.1), an antimicrobial peptide of the streptococcin A-M57 family with a molecular mass of 7.2 KDa and translated from the EF1097 locus gene in *E. faecalis* (strain V583) (Fig. 5(A,D)). The bactericidal invasion in vivo, and epithelial cells still rested on muscle fibers and the basement membrane. Although the vacuoles increased in size, there were no signs of damage at this stage (Fig. 4(F1,F2)).
substance (3–10 K) retained its activity after being incubated over a wide pH range (5–10) \( (F_{6,14} = 35.53, P < 0.0001) \) and tolerated heat treatment (40–100 °C) \( (F_{4,10} = 29.83, P < 0.0001) \), even at 100 °C for 30 min (Fig. 5(B,C)). These characteristics reveal the high stability of this antimicrobial compound, suggesting that it can function under harsh gut conditions. The templates with the highest quality were then selected for model building, and a Phyre2 model for enterococcin LX showed sequence identity (44.4%) (Fig. S6(B)).

Figure 5. Identification of the \( E. faecalis \) LX10 protein (enterococcin LX) by LC–MS/MS. (A) Analysis of the amino acid sequence of the enterococcin LX protein by MS/MS. Amino acid sequence of the enterococcin LX protein deduced from the draft \( E. faecalis \) genome sequence. Peptide sequences matching the LC–MS/MS results are highlighted in red. (B) Effect of pH (5–10) on the activity of crude proteins (3–10 K). (C) The effect of temperature on the inhibitory substance was tested by heating the supernatant at 40, 60, 80, and 100 °C. (D) Peptides identified with LC–MS/MS.

| Protein FDR confidence | Coverage | Unique peptides | Protein group | MV | PI |
|-------------------------|----------|----------------|---------------|----|----|
| high                    | 80.8%    | 20             | 1             | 7219.58 | 9.84 |

Figure 6. Immune-related gene expression level. (A–F) The expression of each gene in the gut of infected larvae was compared to that in the gut of uninfected larvae (CK) after inoculation with different concentrations of \( N. bombycis \) (10³, 10⁶, and 10⁹ spores/mL). (G–L) Expression levels of each gene after treatment with \( E. faecalis \) LX10 (10⁵ spores/mL) (EF), \( N. bombycis \) (10⁹ spores/mL) (Nb), or \( N. bombycis \) and \( E. faecalis \) LX10 (EF + Nb). The healthy silkworms were fed the same volume of sterile water as a control (CK). The messenger RNA (mRNA) levels of Akirin (A, G), Cecropin (B, H), Mesh (C, I), Ssk (D, J), DUOX (E, K) and NOS (F, L) were quantified by RT-qPCR. Each bar represents the mean fold-change ± SE of five independent experiments. Different letters represent significant differences at \( P < 0.05 \) (one-way ANOVA with Tukey’s post hoc test).
3.5 Gene expression changes and enzyme activities

The results showed that the Akirin, Cecropin A, Mesh, and ssk genes were gradually significantly up-regulated after treatment with *N. bombycis* (*P* < 0.05) (Fig. 6(A–D)). Nevertheless, the expression levels of DUOX and NOS showed gradual up-regulation during the early stages of infection (2 days) and then gradual down-regulation in later stages of infection (4 and 6 days) (*P* < 0.0001) (Fig. 6(E,F)). After feeding with *E. faecalis* LX10, the gene expression level of the *N. bombycis*-treated silkworms showed different degrees of recovery and correction compared with the simple *N. bombycis*-infected silkworms (Fig. 6(G–L)).

We further investigated the host gut H$_2$O$_2$ and NO levels and GST enzyme activities. The H$_2$O$_2$ and NO levels displayed complex temporal profiles, initially increasing after 2 days (*F*$_{(3,8)}$ = 42.08, *P* = 0.001) for H$_2$O$_2$; *F*$_{(3,8)}$ = 9.548, *P* = 0.005 for NO) and then decreasing after 4 days (*F*$_{(3,8)}$ = 1.888, *P* = 0.2101 for H$_2$O$_2$; *F*$_{(3,8)}$ = 10.06, *P* = 0.0043 for NO) and 6 days (*F*$_{(3,8)}$ = 7.633, *P* = 0.009 for H$_2$O$_2$; *F*$_{(3,8)}$ = 36.47, *P* < 0.001 for NO) (Fig. 7(A,B)). Similarly, GST activity was clearly inhibited after 4 and 6 days by *N. bombycis* (*F*$_{(3,8)}$ = 43.10, *P* = 0.001 for 4 days, *F*$_{(3,8)}$ = 4.311, *P* = 0.0437 for 6 days) (Fig. 7(C)). Notably, after treatment with *E. faecalis* LX10, the enzyme activity level showed a certain degree of correction, but there was no significant difference compared with that in the group with *N. bombycis* infection alone (Fig. 7(D–F)).

4 DISCUSSION

This study showed that the sensitivity of the silkworm to *N. bombycis* treatment was both dose- and time-dependent. Similar results have been observed for *Plutella xylostella* and *Apis mellifera*. Moreover, the growth of silkworms and food consumption were significantly inhibited, leading to substantial loss in cocoon yield after infection with *N. bombycis*. Many studies have found negative impacts of microsporidian infections on foraging efficiency, wing deformity, lifespan, colony fitness, fecundity and survival in honeybees and bumblebees. Other studies have also explored whether microsporidian infection reduced the weight of the silk gland and growth rate due to the reduced intake of food and its utilization compared to those of healthy silkworms. Many factors, such as initial spore dose, silkworm species, physiological resistance, nutrition status and immune response, may influence the virulence of *N. bombycis* to silkworms, which may contribute to the detoxification of *B. mori*.

Comparative analyses of the silkworm gut microbiome revealed that the gut microbiota compositions and abundance were changed dramatically after *N. bombycis* infection, and these differences might be associated with the gut health of the host. The occurrence and codiversity pattern of the silkworm gut microbiota was significantly altered by *N. bombycis*. High microbial diversity can improve community stability, which may explain why a more intertwined and complex network was present in...
samples with healthy silkworms. Importantly, we also found a strong positive correlation of Enterococcus with inoculation time and infection concentration. It has been reported that the genus Enterococcus is the most dominant bacterial genus in the silkworm gut, implying that the resistance of the silkworm to infection with N. bombycis can be increased by the use of probiotics and optimization of the configuration of the gut bacterial microbiota. Enterococci isolated from the gastrointestinal tract of humans and other diseased animals have been considered important opportunistic pathogens with strong antibiotic resistance. However, Enterococcus sp. have also been found to be probiotic and beneficial in several biomedical models, including in protecting honey bees from American foulbrood disease and improving fish health, so we hypothesize that the gut microbiota might be an important factor driving differences in resistance to N. bombycis infection. Consistent with the KEGG prediction, the changes in gut bacteria caused by microspores may contribute to intestinal dysfunction.

E. faecalis LX10 not only reduced the spore germination rate of N. bombycis in vitro but also ameliorated gut injury in vivo, as confirmed by histopathological analyses. It is worth noting that E. faecalis LX10 reduced but did not eliminate N. bombycis colonization. Similar to our research, Suraporn and Tenerius demonstrated that feeding with Lactobacillus bacteria could increase the survival rate of silkworm larvae challenged by N. bombycis. In addition, the presence of E. faecalis could inhibit fungal morphogenesis and thereby prevent tissue damage caused by hypha expansion in nematode and mouse gut infection models. Using a liquid probiotic form of E. faecalis L3 in infants has a positive impact on overall health and can increase resistance to acute respiratory infections. Some species of Enterococcus produce lactic acid, and accumulation of lactic acid might reduce the pH of digestive juice in grasshoppers and silkworms. The highest germination rate of Nosema occurs at pH 9–10. Thus, the decrease in intestinal injury in silkworms treated with Enterococcus could be related to a reduction in pH in the gut, which possibly led to lowered infectivity of the spores. An alternative inhibitory mechanism of action that could affect pathogen infection might be competitive adhesion of E. faecalis to the intestinal tissue, which would inhibit N. bombycis adhesion. We speculated that in the presence of E. faecalis, N. bombycis reverts to a benign commensal interaction with the host. In addition, the antimicrobial protein enterococcin LX with anti-N. bombycis activities was confirmed as enterococcin EntV. Graham et al. identified that the E. faecalis bacteriocin EntV, produced from the entV (ef1097) locus, is both necessary and sufficient for the reducing in Candida albicans virulence and biofilm formation through the inhibition of hyphal formation, a critical virulence trait. Recent studies have also suggested that members of the gut bacterial genus Enterococcus that produce the peptidoglycan hydrolase SagA improve checkpoint inhibitor cancer immunotherapy in mouse melanoma models. Nevertheless, whether metabolic products of intestinal bacteria can also directly inhibit silkworm infection by pathogens deserves further exploration. We speculate that after the mixing of Enterococcus with N. bombycis, an enzymatic reaction occurs to produce enterococcin, which is an antimicrobial substance, thus limiting the germination of N. bombycis.

We also demonstrated that E. faecalis LX10 can alter host resistance by influencing host physiology, such as via the host immune system and enzyme activity. In our study, after simultaneous feeding with E. faecalis LX10 and N. bombycis, the gene expression level and enzyme activity showed different degrees of recovery and correction compared with those with simple N. bombycis infection. Recent research has revealed that the gut microbiota plays an important role in combating systemic Salmonella infection by inducing local immune responses in mice. In addition, the intestinal symbiotic genus Bacteroides has been found to directly reset the host’s innate immune system through surface carbohydrate structures, increasing tolerance toward pathogens. Modulation of the immune system by E. faecalis could potentially lead to the production of antimicrobial peptides and modulate key signaling pathways, such as the NF-κB and MAPK pathways, to confer an advantage to the host. Nishida et al. demonstrated that lactic acid-producing bacteria stimulated innate immunity in silkworms. In addition, the B. mori immune system quickly activates immune defense mechanisms after infection with N. bombycis, e.g., increased expression of Akirin, cecropinA, Mesh and Ssk. N. bombycis exposure has been shown to induce changes at the molecular level of the innate immune system. For example, the relative expression of the antibacterial peptides abaecin, hymenoptaecin, and defensin significantly increases after Nosema apis infection of honey bees. Moreover, conserved components of smooth septate junction transmembrane proteins (Mesh, Snakeskin: Ssk) in Drosophila and B. mori have been implicated in immune signaling functions and maintenance of the midgut barrier and homeostasis. In contrast, N. bombycis infection seems to suppress the immune response by reducing protection against oxidative stress genes (DUOX, NOS) with prolonged infection. During most gut–pathogen interactions, intestinal redox homeostasis mediated via infection-induced ROS generation by the dual oxidase enzyme and subsequent ROS elimination by immune-regulated catalase is critical for host survival.

In conclusion, the present study provides a clear profile of the complex interplay among N. bombycis, silkworms, and the gut microbiome. From the perspective of biological control, there have been many successful precedents based on the removal of certain commensal bacteria and transgenic symbiosis, especially for vector pests. For example, the symbiont, Rhodococcus rhodnii, has been genetically transformed to express cecropin A and a functional single-chain antibody in the intestinal track of the host vector, Rhodnius prolixus, thereby producing stable paratransgenic insects to eliminate or reduce the number of Trypanosoma cruzi. The present study paves the way for the expression of molecules with anti-microsporidia activity via genetically transformed Enterococcus symbionts of disease-transmitting insects, or direct construction of transgenic insects via mobile DNA elements, which will greatly improve the practical application of microsporidia in lepidopteran pest groups.

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CONFLICTS OF INTEREST
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DATA AVAILABILITY STATEMENT
The data that support the findings of this study are openly available in NCBI at https://www.ncbi.nlm.nih.gov/, reference number PRJNA763083.

SUPPORTING INFORMATION
Supporting information may be found in the online version of this article.

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