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OPEN A 2-Cys peroxiredoxin in response to oxidative stress in the pine wood nematode, Bursaphelenchus xylophilus

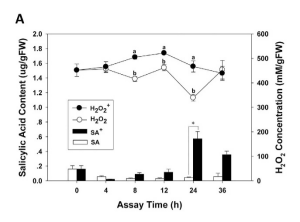
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The pine wood nematode, Bursaphelenchus xylophilus, is the causal agent of pine wilt disease that has devastated pine forests in Asia. Parasitic nematodes are known to have evolved antioxidant stress responses that defend against host plant defenses. In this study, the infestation of whitebark pine, Pinus bungean, with B. xylophilus led to a significant increase in plant hydrogen peroxide (H₂O₂) and salicylic acid levels. Correspondingly, the expression of an antioxidative enzyme, 2-Cysteine peroxiredoxin (BxPrx), was elevated in B. xylophilus following the H2O2 treatments. Recombinant BxPrx, a thermal stabile and pH tolerant enzyme, exhibited high level of antioxidant activity against H₂O₂, suggesting that it is capable of protecting cells from free radical attacks. Immunohistochemical localization study showed that BxPrx was broadly expressed across different tissues and could be secreted outside the nematode. Finally, the number of BxPrx homologs in both dauer-like and fungifeeding B. xylophilus were comparable based on bioinformatics analysis of existing EST libraries, indicating a potential role of BxPrx in both propagative and dispersal nematodes. These combined results suggest that BxPrx is a key genetic factor facilitating the infestation and distribution of B. xylophilus within pine hosts, and consequently the spread of pine wilt disease.

Pinewood nematode, Bursaphelenchus xylophilus (Steiner and Buhrer) Nickle, is a pathogenic parasite of the devastating epidemic pine wilt disease in Asia¹ and Europe², and has caused serious damage to pine forests, especially in Japan³ and China⁴. In addition to the economic importance as a notoriously effective plant pathogen, B. xylophilus also has drawn attention for its complex ecological traits centered around the tritrophic interactions among insect vectors, plant parasitic nematodes, and pine/fungi hosts⁵. To complete its life cycle, B. xylophilus switches from a propagative to a dispersal stage⁶. In the propagative portion of the lifecycle, B. xylophilus grow and develop through four juvenile stages and reach adulthood by feeding on the parenchymal cells of the healthy pine trees. This feeding by nematodes leads to the disruption of water transportion and subsequent death of affect tissues in pine trees⁷. When the host plant becomes seriously infected, B. xylophilus enters the mycophagous phase wherein nematodes feed on the blue stain fungi colonized in the dying tree until the food sources become limited. The dispersal life cycle starts with the formation of the pre-dauer juveniles that aggregate around the pupal chambers of the pine sawyer beetles, Monochamus alternatus Hope. Finally, M. alternatus beetles feed on healthy pine wood after emergence, and incidentally brought B. xylophilus dauer with them8. Subsequently, the dispersal of M. alternatus facilitates the spread of beetles and pathogens. The transmission of pine wilt disease would fail unless the three factors, the pine host, parasitic nematode, and insect vector, work in sync⁷.

To ensure successful infestation, B. xylophilus needs to break through the pine host defense system. Unlike other plant parasitic nematodes, such as cyst and root-knot nematodes which are sedentary and feed on host plant cells, B. xylophilus, a migratory endoparasitic nematode, has to invade and move through the pine host to feed. To counteract host defense, B. xylophilus has evolved multiple measurements9. On the other hand, host plants activate their defense mechanisms locally and systemically in response to pathogenic attacks¹⁰ as well as nematode infestations¹¹. Reactive oxygen species (ROS), including superoxide anion $(O_2 - \cdot)$, hydrogen peroxide (H_2O_2) , and hydroxyl radicals (·OH), are considered to be the first line of defense in plants¹². Reactive oxygen

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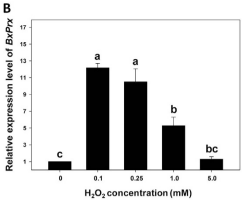


Figure 1. Interactions between pine hosts and nematodes involving BxPrx. (**A**) Comparative analysis of H_2O_2 and SA levels between pine seedlings inoculated with *B. xylophilus* (with "+" sign) and control pine inoculated with M9 buffer (without "+" sign). H_2O_2 and SA levels were documented at 0, 4, 8, 12, 24, and 36 h post-inoculation. (**B**) After incubated with 0, 0.1, 0.25, 1 and 5 mM H_2O_2 for 5 min, BxPrx expression was recorded.

species oxidize DNA, proteins, and lipids, which causes damage to organelles and inhibits cell functions in plant attackers¹³. H_2O_2 can cross the plasma membrane and activate signaling molecules to induce defensive genes and enzymes. Specifically, H_2O_2 acts as a marker in pathogen recognition, causal trigger for hypersensitive reaction and apoptosis of host plants. Reactive oxygen species, particularly H_2O_2 , is an important factor for regulation of host-nematode interactions and partly govern the success or failure of disease^{14,15}. Salicylic acid (SA) is an important signaling molecule involved in systemic resistance by inducing the expression of pathogenesis-related (PR) proteins in plants¹⁶. A feature of SA signaling is its interaction with ROS in plants¹⁷. The increase of H_2O_2 titer at the site of nematode penetration could induce the accumulation of SA. *Vice versa*, SA could regulate H_2O_2 level through manipulating ROS-scavenging systems^{18,19}. Durner *et al.* propose that H_2O_2 might play a role in the activation of PR genes by interaction with SA¹⁰. Therefore, breaking down ROS defense is not only important to the initial invasion, but also facilitate its ongoing and persistent infestations by weakening the resistant in host plants. Neutralizing ROS is essential for parasitic nematodes to overcome plant defenses²⁰. Consequently, the survival of nematodes is directly linked to the antioxidant enzymes²¹.

In many parasitic nematodes, 2-Cysteine peroxiredoxins (Prxs) are overexpressed and involved in H_2O_2 detoxification²². In animal parasitic nematodes, the secretion of Prxs is considered an evolutionary adaptation to protect nematodes against host $H_2O_2^{23}$. Onchocerca volvulus, the parasitic nematode for 'river blindness', excreted a 2-Cys Prx that localized within the hypodermis and cuticle in the infective larvae²⁴. Among plant parasitic nematodes, 2-Cysteine peroxiredoxin (Prx2) has been found on the cuticle surface of infective and post-infective juvenile Globodera rostochiensis^{25,26}, and excreted by Meloidogyne incognita when in close contact with plant cells¹⁴. Furthermore, silencing of Prx2 by RNAi led to the impairment of M. incognita development inside the host¹⁴. These results suggested that Prxs, especially Prx2, may play a critical role in plant-nematode interactions by counteracting plant defensive $H_2O_2^{27}$.

Previously, a 2-Cys *Prx* gene from *B. xylophilus*, *BxPrx*, was cloned and its expression profile was analyzed²⁸. Using an integrative approach, we comprehensively characterized *BxPrx* to gain a better understanding of its ecological and biological functions in the co-evolution of pine-nematode arm race. The combined results demonstrated that *BxPrx* plays a key role in the dispersal of *B. xylophilus*, acting as a scavenger for host-derived ROS attack in the propagative cycle, and may protect against its own metabolic H₂O₂ in the dispersal cycle.

Results

Nematode, plant defensive compounds and BxPrx. The roles of plant secondary metabolites, including H_2O_2 and SA, have been investigated in this study. H_2O_2 titer in pine hosts were significantly increased after inoculation with *B. xylophilus* for 8 h, peaked at 12 h post inoculation, and high level of H_2O_2 was sustained for about 16 h before it dropped back to the level of control pines (Fig. 1A). Similarly, SA titer in pine was significantly induced after inoculation with *B. xylophilus*, and it peaked at 24 h post-inoculation (Fig. 1A).

The stability of four candidate reference genes, including $EF \, 1\alpha$ (eukaryotic translation elongation factor 1α , accession number: GU130132), α -tubulin (GU130151), β -tubulin (AB500150), and β -actin (EU100952), under an array of hydrogen peroxide treatments were examined by quantitative real-time PCR (qRT-PCR) analysis. GeNorm analysis demonstrated β -actin (0.400) and $EF \, 1\alpha$ (0.493) had the lowest M-values, suggesting that this pair was the most stably expressed reference genes in B. xylophilus under oxidative stress (Table S1)²⁹. BxPrx expression in B. xylophilus was found significantly induced in respond to elevated H_2O_2 titer (0.1 ~ 1.0 mM) in the environment, although the expression of BxPrx started to decrease when H_2O_2 concentration reached 1.0 mM (Fig. 1B).

Immunohistochemical localization study showed that BxPrx was abundantly expressed in *B. xylophilus* and predominantly distributed in excretory canal, and the two lateral glands (Fig. 2B). BxPrx also could be secreted

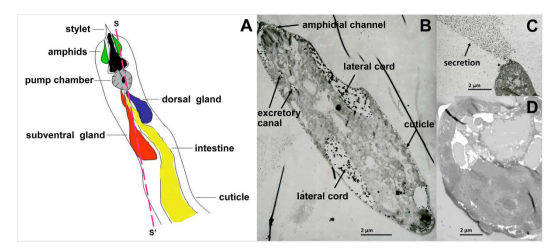


Figure 2. Immunohistochemical localization of BxPrx in *B. xylophilus*. (**A**) Schematic drawing of the anterior section of *B. xylophilus*. It is modified from Vanholme *et al.*⁶⁰, and the dash line indicates the sample section in *B. xylophilus*; (**B**) BxPrx expression was detected under cuticle, in amphid, dorsal and subventral gland, and under cuticle; (**C**) BxPrx was secreted outside the nematode through stylet; (**D**) Preimmune serum showed no detectable binding on the nematode samples.

outside the nematode through stylet (Fig. 2C). Preimmune serum showed no detectable binding to nematodes (Fig. 2D).

Antioxidant activity of recombinant BxPrx. Recombinant BxPrx exhibited limited thione peroxidase (TPx) activity in a reaction system containing both thioredoxine (produced from *E. coli*, EcTx) and thione reductase (produced from *E. coli*, EcTx). In contrast, in a reaction system containing L-glutathione and glutathione reductase (GR), BxPrx showed substantial GPx activity (563.8 ± 41.6 units/mg protein) against H_2O_2 , however, it did not reduce cumene hydroperoxide ($C_9H_{12}O_2$) (Fig. 3A). In optimal condition assays, BxPrx exhibited GPx activity between pH 5 and 9, with pH 7and 8 recorded the highest activities, whereas pH 3 and 10, respectively, led to a dramatic reduction of the antioxidant activity (Fig. 3C). BxPrx activity was stable across a wide range of temperature (4-60°C), however, its antioxidant activity was significantly decreased when temperature reached 80 °C (Fig. 3D).

The conformational change of BxPrx from monomeric to dimeric protein was induced by the addition of H_2O_2 (Fig. 3B). The rate of dimerization is positively correlated with H_2O_2 concentration. Consequently, BxPrx is a typical 2-Cys Prx with the formation of an inter-molecule dimer to complete the catalytic cycle.

Recombinant BxPrx showed significant tolerance to the toxicity of H_2O_2 oxidation (30% lower susceptibility than a blank, P < 0.05, Fig. 4A,B). DNA cleavage protection test demonstrated that BxPrx protected plasmid DNA against the free radical attack, and this protection weakened when the amount of BxPrx proteins declined. During a 4h-observation, supercoiled plasmid DNA was best protected at a BxPrx concentration of $200 \,\mu\text{g}/\mu\text{l}$. The damaged DNAs, such as nicked and slowly moving linear DNAs, became evident when less BxPrx was added into the reaction system. When BxPrx concentration reduced to $25 \,\mu\text{g}/\mu\text{l}$, only the linear DNAs were detected (Fig. 4C).

Antioxidants among dauer-like and fungi-feeding *B. xylophilus*. Genes encoding antioxidants, including *SOD* (*superoxide dismutase*), *GPX* and *CAT* (*catalase*), were extracted from EST libraries generated from dauer-like and fungi-feeding *B. xylophilus*, respectively³⁰. In general, fungi-feeding nematodes had a greater number of antioxidants than dauer-like *B. xylophilus*³⁰. *BxPrx* homologs, however, showed relatively stable expression between fungi-feeding (Genbank ID: CJ984042, CJ984159, CJ984424, and CJ984735) and dauer-like *B. xylophilus* (Genbank ID: CJ989294, CJ989141 and CJ988426) in comparison to expressions of housekeeping genes, *Act1* and *Hsp70* (Table 1).

Discussion

Prx2s in nematodes with different ecological niches. Peroxiredoxin was first detected in the yeast *Saccharomyces cerevisiae*, where its expression was induced by exposure to oxidative stress³¹. In free living bacterivore, *Caenorhabditis elegans*, expression of *CePrx2* was induced by short-term exposure to t-butyl hydroperoxide $(t\text{-BOOH})^{32}$. In an animal parasitic nematode, *Haemonchus contortus*, HcPrx2, exhibited high *Km* toward H_2O_2 , suggesting that HcPrx2 can protect the nematode against the host immune response²³. Similarly, in a plant parasitic nematode, *G. rostochiensis*, a 2-cys peroxiredoxin could breakdown H_2O_2 and was located on the surface of invasive and post-infective juveniles, suggesting that Prx may play a role in protection against plant defensive responses²⁵. MiPrx2, a Prx2 from the other plant parasitic nematode, *M. incognita*, was located in tissues in close proximity with plant cells during parasitism. The development of nematode was impaired after the silencing of *MiPrx2*¹⁴. *Bursaphelechus xylophilus* has a complex life cycle and unique feeding behavior in comparison to other plant parasitic nematodes which feed primarily on plant tissues for nutrients. It resides two distinctly different habitats: 1) insect vectors for dispersal and foraging, and 2) pine hosts for food, shelter, and proliferation.

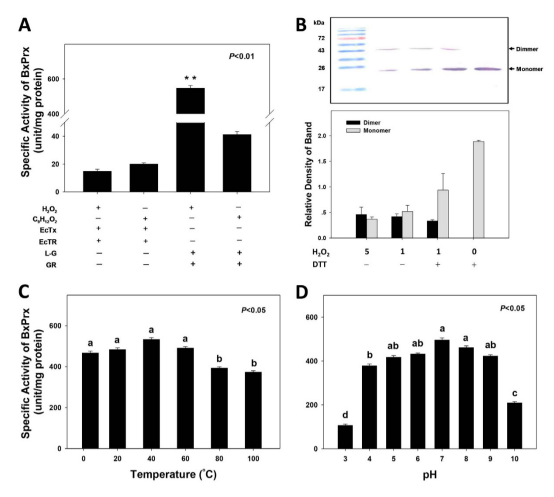


Figure 3. Characterization of BxPrx activity. All assays were done in triplicates, with $1.5\,\mu g$ of purified recombinant BxPrx in each $500\,\mu l$ reaction system. (A) TPx activity was test with EcTrx and EcTR in the reaction; GPx activity was test with L-G and GR in the reaction; H_2O_2 and $C_9H_{12}O_2$ were used as substrates separately. (B) Immunoblot assay of total protein extract from *B. xylophilus* incubated with 0, 1 mM and 5 mM H_2O_2 by polyclonal antiserum against BxPrx. $80\,\mu g$ protein extract was loaded in each lane separately, and the change from monomer to dimer was detected. Monomer was showed below $26\,k$ Da, and dimer was around $43\,k$ Da. Band density analysis resulted from three replicates. (C) GPx activity of BxPrx after incubated at different temperature for $30\,m m$, and the tests were performed in the reaction system of pH 7.0. (D) GPx activity of BxPrx under different pH conditions, the tests were performed at room temperature.

Moreover, *B. xylophilus* also exploits a wide range of fungi as food source which occupy the dying pine hosts caused by the infestation of *B. xylophilus*⁷. Thus, we hypothesize that genetic factors which suppress or counter the innate immune responses from the pine hosts are essential for the infestation and distribution of *B. xylophilus*. BxPrx, a conserved 2-Cys peroxiredoxin protein with notable antioxidant activity²⁸, is likely the genetic factor facilitating *B. xylophilus* to circumvent the host immune response to ensure a successful infestation.

Host innate immune response to B. xylophilus infestation. H_2O_2 titers and SA levels were induced in pine seedlings against the infestation of B. xylophilus. The induction of SA was lagging behind the release of H_2O_2 , indicating that ROS was likely to be the first line of defense in pine host innate immune response to B. xylophilus infestation. As one of the ROS species, H_2O_2 is an early stress signal interacting with a network of signal transduction in plants, including SA pathway to promote defensive responses and systematic acquired resistance (SAR) against pathogens^{33,34}. Following B. xylophilus infestation, elevated H_2O_2 level led to the induction of SA pathway and the initiation of SAR in pine hosts. Van Camp et al. reported that H_2O_2 , not SA, might induce the expression of a resistant gene AoPR-1 in tobacco, indicating that certain defense genes in plant genomes could be activated directly by $H_2O_2^{35}$. Researchers also suggested that H_2O_2 and SA constitute a self-amplifying system, in which H_2O_2 induces SA accumulation and, in turn, SA can increase H_2O_2 level³⁶. These studies demonstrate that H_2O_2 and SA are important plant defensive compounds to deter invading pathogens and to initiate SAR.

Antioxidant activity of BxPrx. BxPrx has significant antioxidant activity toward H_2O_2 in a reaction system containing L-glutathione and GR, but limit activity in a system composed of EcTx and EcTR or on organic oxidants. Some human and animal parasites can use both reaction systems for regeneration, such as *Schistosoma mansoni*³⁷ and *Plasmodium falciparum*³⁸, although HcPrx could not utilize the glutathione system directly²³. Prxs

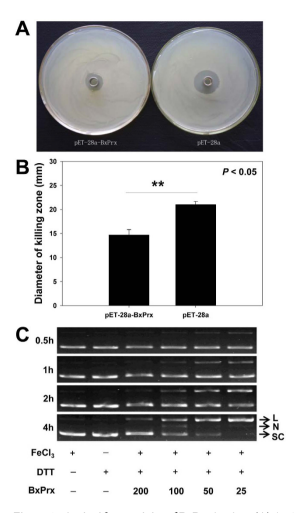


Figure 4. Antioxidant activity of BxPrx in vitro. (**A**) Antioxidant activity of BxPrx transformants was investigated using a discs assay. (**B**) Difference of susceptibility to H_2O_2 between E. coli with pET-28a-BxPrx and blank pET-28a transformed were compared by the killing zone diameter. (C) DNA protection assay. Free oxidant radical attack was generated due to the presence of Fe³⁺ and DTT. PUC19 plasmid was incubated for 0.5, 1, 2, and 4 h respectively. After attacked by the radical, supercoiled plasmid (SC) was degraded to slower-moving nicking (N), and then slowest linear DNA (L).

in the potato cyst nematode G. rostochiensis and plant parasitic root-knot nematode M. incognita exhibited notable antioxidant activity in a system that used both trypanothione and thiocyanate as thione donors ^{14,25}. MiPrx2, CePrx2 and HcPrx2 all exhibited strong activities for the reduction of H_2O_2 and organic oxidants. Similar to BxPrx, Prx in G. rostochiensis was not active against organic oxidants²⁵. The different substrate specificity exhibited by these Prxs may be correlated with the habitats occupied by these nematodes. In animals, acquired immunity shows its major capacity in suppressing pathogen infection, however, plants mainly rely on innate immune responses, including ROS and accompanied signal molecules to initiate $SAR^{39,40}$. Prxs, with strong antioxidant activity to breakdown H_2O_2 and protect DNA from oxidative damage, facilitate B. xylophilus infestation by down-regulating the defensive responses in pine hosts.

Bursaphelenchus xylophilus, an invasive nematode with a wide range of distribution 41 , shows strong tolerance toward cold and heat stresses 5,42 (11 to 32 °C). In this study, recombinant BxPrx were active against H_2O_2 under a broad range of temperature and pH, and protected DNA from oxidative damages. Based on the catalytic mechanisms and the presence of either one or two highly conserved cysteine residues, Prxs are categorized into 1-Cys, typical 2-Cys, and atypical 2-Cys Prxs 43 . The primary structure of BxPrx contains two highly conserved active sites (Cp and Cr). Catalytic reactions of the typical 2-Cys are accomplished through the formation of intermolecular disulfide. In contrast, inner-molecular disulfide is the result of the reaction catalyzed by the atypical 2-Cys Prx 44,45 . The conformational change of BxPrx from monomer to inter-molecular dimer was detected after it was oxidized by H_2O_2 . The rate of dimerization and H_2O_2 concentration was positively correlated, indicating that BxPrx is a typical 2-Cys Prx.

BxPrx in the propagative and dispersal cycle of B. xylophilus. The combined results in this study demonstrated that oxidative burst (H_2O_2) and secondary metabolites (SA) were inducible when pine hosts were infested with B. xylophilus. Following B. xylophilus infestation, excessive amount of ROS generated in pine hosts contributes to the development of disease symptoms^{46,47}. Effective antioxidant activities are an adaptation in pathogens to neutralize host innate defenses to enhance their survivorship and infestation. Previous studies

	Transcript ^a	Description	Number of ESTs	
			Fungi-feeding	Dauer-like
Antioxidant	BXC00823	SOD-1	1	0
	BXC04177	SOD-3	1	0
	BXC07336	SOD-3	0	1
	BXC03446	SOD-4	1	0
	BXC03943	GPX	2	0
	BXC00081	GPX	2	3
	BXC01784	GPX	1	0
	BXC02367	CAT-1	2	0
	CJ984042	Prx2	1	0
	CJ984159	Prx2	1	0
	CJ984424	Prx2	1	0
	CJ984735	Prx2	1	0
	CJ989294	Prx2	0	1
	CJ989141	Prx2	0	1
	CJ988426	Prx2	0	1
Housekeeping gene	CJ984848	Actin1	1	
	CJ984265	Actin1	1	
	CJ984341	Actin1	1	
	CJ984728	Actin1	1	
	CJ985146	Actin1	1	
	CJ984832	Actin1	1	
	CJ984026	Actin1	1	
	CJ985003	Actin1	1	
	CJ985085	Actin1	1	
	CJ990129	Actin1		1
	CJ988716	Actin1		1
	CJ990734	Actin1		1
	CJ989041	Actin1		1
	CJ988766	Actin1		1
	CJ990669	Actin1		1
	CJ989163	Actin1		1
	CJ984710	Hsp70	1	
	CJ984856	Hsp70	1	
	CJ989482	Hsp70		1
	CJ988689	Hsp70		1

Table 1. Antioxidative transcripts extracted from fungi-feeding and dauer-like *B. xylophilus* EST libraries. aE -value $< E^{-10}$.

demonstrated that virulent B. xylophilus isolates survived better than avirulent counterparts under the oxidative stress 9,48 . The bacteria symbiont Serratia spp 9 , and elevated catalase expression in the intestine of B. $xylophilus^{49}$ contribute their tolerance towards the oxidative stress and facilitate the infection of pine wood disease. In this study, BxPrx expression was significantly induced by the increase of environmental H_2O_2 titer. Recombinant BxPrx showed high levels of antioxidant activity to breakdown H_2O_2 and protect DNAs from oxidative radical attacks. Vice versa, decreased H_2O_2 level also reduce the accumulation of SA and the subsequent host plant innate immune responses. In addition, BxPrx were broadly expressed under cuticle surface where is in close contact with pine hosts, in excretory canals and two lateral glands which are the main secretory organs in nematodes. BxPrx could also be secreted outside the nematode through stylet. These combined results suggest that BxPrx plays a critical role in protecting B. xylophilus against pine host-derived H_2O_2 .

Prx2 is also essential for the development and reproduction of nematodes. Silencing of *CePrx2* in *C. elegans* resulted in a stunted development and a significantly reduced brood size³². Similarly, knocking down *BxPrx* significantly reduced the propagation in *B. xylophilus*⁵⁰. Our previous study found that anti-serum of BxPrx could only detect protein samples extracted from virulent *B. xylophilus* but not *B. mucronatus*²⁸, suggesting that BxPrx could be used as a molecular marker to identify the virulent *B. xylophilus*⁵⁰. A shorter life cycle and higher proliferation rate are the fundamental differences between avirulent and virulent *B. xylophilus*, therefore, BxPrx could play a role in the pathogenicity of *B. xylophilus*. Different gene clusters were identified between the EST libraries derived from dauer-like and fungi-feeding *B. xylophilus*³⁰. Many homologs of antioxidant genes, including *SOD*, *GPX* and *CAT*, which were involved in longevity in *C. elegans*³², were found with notably reduced number in dauer-like EST library compared to fungi feeding *B. xylophilus*³⁰. In contrast, *BxPrx* isoforms were similar between dauer-like and

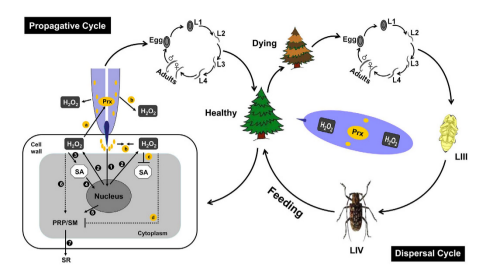


Figure 5. Proposed role of BxPrx in the pine-nematode-pine wilt disease tritrophic interactions. In the propagative cycle, BxPrx protects *B. xylophilus* from the damage of host ROS defense, helps the nematode for a successful infestation into a healthy pine, and then *B. xylophilus* finish the life cycles with egg, four instars of larvae and adults (1) The stylet of *B. xylophilus* injected into pine cell for feeding. (2) Infestation of *B. xylophilus* could induce H₂O₂ defense in pine (3) The accumulation of H₂O₂ could lead to the increase of SA in host plant. (4–7) The accumulation of H₂O₂ and SA induced the expression of pathogenesis-related protein (PRP) and signal molecular (SM) which lead to the following systematic resistance (SR) reactions in pines. Correspondingly, (a) Expression of BxPrx in *B. xylophilus* could be induced by the H₂O₂ from host defense reaction. (b) Prx could be secreted to the cuticle surface of nematodes and into host from stylet to neutralize H₂O₂ produced by host. (c,d) The neutralization of H₂O₂ by BxPrx and the following inhibition of the SA accumulation, all would lead to the retard of SR reactions in host. After dramatic reproduction of *B. xylophilus*, host pines become dying, and the dispersal cycle of the nematodes startup. In the dispersal cycle, dauer LIII and LIV nematodes appear and are distributed by the feeding of intermediate insect *M. alternatus*. BxPrx protects *B. xylophilus* against the ROS damage generated from its own metabolism until new infestation.

fungi-feeding EST libraries. *BxPrx* expression was consistent in the dauer stage, a life stage that is structurally and physiologically adapted for the long-term survival and dispersal without much food sources, suggesting that BxPrx may play an important role against the nematode's own metabolic ROS for survival during its dispersal cycle.

Summary. To summarize our finding, we schematically proposed the multiple functions of BxPrx in the nematode-pine arm race (Fig. 5). During the propagative cycle, the infestation of B. xylophilus can induce the production of H_2O_2 in pine hosts. Besides direct toxicity to B. xylophilus, the accumulation of H_2O_2 triggers the elevation of SA. It will also induce the expression of resistance genes and initiate the SAR pathway in pine hosts. In the meantime, however, the elevated H_2O_2 level in pine hosts likely induces BxPrx expression in nematodes. Synchronized secretion of BxPrx, in turn, reduces H_2O_2 level at the micro-environment surrounding B. xylophilus in pine hosts and thus protecting nematodes from oxidative damages. This can also reduce the SAR response in pine hosts through the suppression of H_2O_2 to a level that the induction of SA pathway is arrested. Therefore, in the propagative cycle, BxPrx plays a critical role in protecting B. xylophilus against ROS attack from hosts.

After infestation and proliferation of *B. xylophilus*, hosts show symptoms of wilt and nematodes turn to feed on a variety of fungi occupied in the decaying pine materials. When fungal food sources become scarce, the pre-dauer LIII nematodes congregate around the *M. alternatus* pupae and latch onto the newly emerged adults. The dauer LIV nematodes complete the dispersal cycle through the maturation of the adult pine sawyer beetle on healthy trees. ROS generated from the metabolism can threaten the survivorship of nematodes in the dispersal period. In the dauer stage, antioxidative BxPrx protects nematodes from their own metabolic attacks for a longer life span and a broader distribution during the dispersal cycle.

In conclusion, we suggest that BxPrx is a key genetic factor that facilitates the infestation and distribution of *B. xylophilus* within pine hosts, and consequently the spread of pine wilt disease. With the advent of functional genomics tools, including RNA interference (RNAi)⁵¹ and transgenic approach⁵⁰, genetic basis governing the transmission and infection of pine wilt disease is a logic choice for future research. A better understanding of how nematodes overcome oxidative stresses generated from their own metabolic processes and from the innate immune responses in pine hosts will provide novel targets for the long-term, sustainable management of this devastating pine disease.

Materials and Methods

Biological materials. The *B. xylophilus* isolate JSZJ1-6 (Sampled from Jiangsu, Zhejiang province¹) was maintained as described previously⁵². Seedlings of 1-2 year-old whitebark pine, *Pinus bungean* (20-40 cm in

height and 5-8 cm in diameter), were maintained in a green house with a long-day photoperiod (light/dark 16:8) at 25 °C for 1.5 months until new shoots start to grow.

Interactions between the pine hosts and nematodes. Change of H_2O_2 and SA level in pine host responding to B. xylophilus infestation. The innate resistance of P. bungean was investigated by an artificial infestation (Fig. S1). Pine seedlings were inoculated with 20,000 B. xylophilus per plant⁵³ using M9 buffer (KH₂PO₄ 30 g/L, K₂HPO₄ 60 g/L, NaCl 50 g/L) as a control. Leaves were sampled at 0, 4, 8, 12, 24, and 36 h after inoculation, and snap frozen at -80 °C for the subsequent H_2O_2 and SA analysis.

 H_2O_2 concentration was determined as described previously by Brennan and Frenkel⁵⁴. In brief, 0.5 g of frozen pine needles were ground in 1 ml acetone and centrifuged at 3,000 g for 10 min. After mixing the supernatant with 20% TiCl₄-HCl solution and ammonium hydroxide, the resultant pellet was harvested and resuspended in 2 M H_2SO_4 . H_2O_2 concentration was measured by A_{410} of the titanium-peroxide complex. Free SA was extracted as described previously⁵⁵. A total of 0.5 g of frozen needles were homogenized in 3 ml of 90% methanol and kept soaking at -20 °C for overnight. After centrifugation at 4 °C, 8,000 rpm, for 30 min, the supernatants were collected and dried in a speed vacuum with heat (~40 °C). The residue was resuspended in 2.5 ml of 5% trichloroacetic acid and sonicated for 10 min. Free SA was then separated from conjugated SA through organic extraction with 2 volumes of ethyl acetate-cyclohexane-isopropanol (50:50:1). The organic phase containing free SA was then dried using a speed vacuum, resuspended in 400 μl of acetonitrile, filtered, and measured by reverse-phase HPLC (HP1100 Agilent, USA). An Agilent-C18 column (5 μm, 250 mm × 4.6 mm), with a detection wavelength of 302 nm and elution condition of V (acetonitrile): V (H_2O containing 1.5% acetic acid), 3:97 (10 min), 38.8: 61.2 (20 min), 38.8: 61.2 (26 min), was used in the HPLC analysis. The amount of free SA was determined based on the standard curve of a SA reference (Sigma-Aldrich, USA). Paired-samples T test (SPSS17.0, SPSS Inc., Chicago, USA) was used to analyzed changes of H_2O_2 and SA levels between pines inoculated with or without *B. xylophilus*.

BxPrx expression profile under oxidative stresses. To select the suitable reference gene(s) for the subsequent qRT-PCR analysis, the stability of four housekeeping genes, EF 1 α (accession number: GU130132), α -tubulin (GU130151), β -tubulin (AB500150), and β -actin (EU100952), under the hydrogen peroxide treatments were evaluated by geNorm²⁹. The end point measurement of this conventional statistical software is M-value, the variation of a gene compared to all other candidates. Based on the pair-wise comparison, geNorm automatically eliminates the gene with the highest M-value, and repeats the process until there is only two genes left. The last pair remaining is considered as the optimum pair of reference genes²⁹. The PCR primers were listed in Table S2, and the PCR was performed with 40 cycles of 94 °C (30 s), 58 °C (45 s), and 72 °C (30 s).

Bursaphelenchus xylophilus of mixed developmental stages were incubated with various concentrations of H_2O_2 for 5 min⁵⁶. Total RNA extraction and cDNA synthesis were carried out as described previously²⁸. BxPrx (GenBank accession number: EU095848) expression profile under oxidative stresses was investigated using qRT-PCR with primers listed in Table S2. qRT-PCR was performed on a 7300 Real-Time PCR System (Applied Biosystems, Foster City USA). The $2^{-\Delta\Delta Ct}$ method was used to calculate the expression level of BxPrx. SPSS Statistics 17.0 (SPSS Inc., Chicago, USA) was used for the statistical analyses. Since the data did not fit homoscedasticity, BxPrx expression level under oxidative stress was assessed by one-way ANOVA and followed by a Games-Howell test.

Immunohistochemical localization of BxPrx. *Bursaphelenchus xylophilus* of mixed developmental stages were collected by centrifugation at 2,000 rpm for 1 min, and washed with M9 buffer for 3 times. The resultant nematode samples were fixed, dehydrated, cut to ultrathin sections, hybridized with antiserum, and observed in transmission electron microscope²⁸.

Antioxidant activity of recombinant BxPrx. Biochemical characterization of recombinant BxPrx. Recombinant BxPrx protein was expressed in a Novagen's pET-28a expression vector (Merck KGaA, Darmstadt, Germany) and purified with Ni-NTA agarose (Qiagen, Hamburg Germany)²⁸. Optimal conditions for BxPrx activity assay, including substrate specificity, pH tolerance, and temperature range, were determined enzymatically. The capability of recombinant BxPrx on the reduction of H_2O_2 and $C_9H_{12}O_2$ was measured, respectively, at OD₃₄₀⁵⁷. The reaction system contained 5 mM potassium phosphate (pH 7.0), 1 mM EDTA, 0.1 mM NADPH, and with 16 μ M EcTx (Sigma) and 0.23 μ M EcTR (Sigma) for the TPx activity assay; or 1 mM L-glutathione (Sigma) and 0.1 unit/ml GR (Sigma) for the GPx activity assay. To determine the optimum pH, BxPrx activity was measured between pH 3 and 10 at room temperature. For optimal temperatures, recombinant BxPrx was incubated for 30 min at a temperature gradient of 4, 20, 40, 60, 80, and 100 °C, respectively, before engaging in a BxPrx activity assay at pH 7. The activities of BxPrx were compared by one-way ANOVA and followed by a Tukey test using SPSS Statistics 17.0 (SPSS Inc., Chicago, USA).

Immunoblot analysis. Bursaphelenchus xylophilus was treated with $\rm H_2O_2$ and harvested for the assay as previously described by Oláhová⁵⁸. Nematodes were incubated in 0, 1 and 5 mM $\rm H_2O_2$ solution separately for 5 min, and then washed with M9 buffer for 3 times. Nematodes were then harvested in 20% trichloroacetic acid (TCA) before snap-frozen in liquid nitrogen. Thawed pellets were washed with acetone to remove TCA, dried and resuspended in SDS-PAGE loading buffer. The resultant samples were then separated through 12% SDS PAGE and analyzed by Western blotting using an anti-BxPrx serum³⁵. Band intensity was analyzed using Quantity One 4.6.2 (Bio-Rad, USA).

Antioxidant activity assay. Antioxidant activity of BxPrx transformants was assayed following Isermann³² with minor modifications. Escherichia coli was spread onto the top of a LB (Lysogeny Broth) agar plate (LB medium

supplemented with a 2 mm of 0.8% agarose layer below, 1 mM IPTG, and $100\,\mu\text{g/ml}$ ampicillin inside), and then incubated at 37 °C for 3 h. Subsequently, oxford disc (1 cm in diameter) was inserted into the middle of the LB medium and positioned upon the agarose layer. A total of $100\,\mu\text{l}$ of 1 mM H_2O_2 was then added onto the disc. The resultant plates were placed at the room temperature for 16 h without moving until all the H_2O_2 infiltrated into the LB medium, and then plates were transferred to 37 °C. After 24 h of incubation, the diameters of the killing zones were measured. The viability of *E. coli* transformats with or without BxPrx was compared using a paired-samples T test (SPSS17. 0, SPSS Inc., Chicago, USA).

DNA cleavage protection assay. According to $Stacy^{59}$, various concentrations of recombinant BxPrx was first incubated in a reaction mixture containing 250 mM Hepes, 5 mM EDTA, 100 mM DTT, and 600 μ M FeCl₃, at 37 °C for 30 min, and then 250 ng of pUC19 plasmid was introduced into the reaction system (Tiangen, Beijing, China). The status of pUC19 with different incubation conditions was resolved by agarose gel electrophoresis (1%).

Transcripts abundance of antioxidant enzymes in B. xylophilus. Transcripts abundance of antioxidant enzymes was analyzed by searching and comparing the number of homologs between EST libraries generated from dauer-like and fungi feeding B. xylophilus. The sequence information of the two EST libraries was downloaded from NCBI database, and the retrieval of the homologs was conducted using software BioEdit (Ibis Biosciences, Carlsbad, CA).

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Author Contributions

Z.L. conceived the experiments; Z.L., Q.W.Z. and X.Z. designed the study; Z.L. and X.Z. analyzed the data; and Z.L., Q.W.Z. and X.Z. wrote the manuscript. All authors read and approved the final version of manuscript.

Additional Information

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