INTERACTION OF ROOT-KNOT NEMATODE
(MELOIDOGYNE JAVANICA) AND TOMATO AS
AFFECTED BY HYDROGEN PEROXIDE

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Abstract: The effect of hydrogen peroxide (H$_2$O$_2$) on root-knot nematode (RKN, Meloidogyne javanica) in tomato was investigated. Soil drenching with exogenous H$_2$O$_2$ was done using seven H$_2$O$_2$ concentrations (1, 10, 100, 250, 500, 750 and 1000 mM) at different application times (24 hours before the time of plant inoculation with the RKN (T1), at the time of inoculation (T2), and 24 hours after the inoculation time (T3)). The nematode reproduction rate (eggs/g fresh root) was significantly reduced in all H$_2$O$_2$ treatments compared with the untreated control. The lowest reduction in nematode reproduction occurred at 10 mM H$_2$O$_2$. The application times T1 and T2 were significantly higher in reducing the reproduction rate than T3 at 250, 750 and 1000 mM H$_2$O$_2$. The content of endogenous H$_2$O$_2$ in the treated plants was significantly higher than in the non-treated plants. Some phytotoxicity was apparent at the higher concentrations of H$_2$O$_2$ (≥ 500 mM) in the treated plants due to the accumulation of the endogenous H$_2$O$_2$. The treatments with 1 and 10 mM H$_2$O$_2$ did not differ from the untreated control in plant chlorophyll content while the content was significantly reduced at the higher concentrations. Exogenous application of H$_2$O$_2$ may have a direct effect on the nematode reproduction and an indirect effect on the treated tomato plants that can be elicited by H$_2$O$_2$ to resist the nematode infection.

Key words: Host resistance, nematode reproduction, host-pathogen interaction.

INTRODUCTION

Root-knot nematodes (RKNs, Meloidogyne spp.) attack a wide range of crop species. Annually, about 5% of the world crop production is destroyed by Meloidogyne species (Sasser et al. 1983; Barker et al. 1985; Sasser 1987).

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Three Meloidogyne species (M. javanica (Teurb) Chitwood, M. incognita (Kofoid and White) Chitwood, race 1 and 2, and M. arenaria (Neal) Chitwood, race 2) were reported to occur in Jordan, with a predominance of M. javanica (Abu-Gharbieh 1982; Atieh 1986; Karajeh et al. 2005). The average annual losses of irrigated vegetable crops cultivated in the Jordan Valley due to RKNs has been estimated for nearly 15% (Abu-Gharbieh 1994).

Hydrogen peroxide (H$_2$O$_2$) functions as a stress signal in plants, mediating adaptive responses to various stresses. Exposure to various abiotic and biotic stresses results in the accumulation of H$_2$O$_2$ (Desikan et al. 2001). It is already known that H$_2$O$_2$ can induce the expression of genes involved in antioxidant defense (Levine et al. 1994; Karpinski et al. 1999; Morita et al. 1999; Lopez-Huertas et al. 2000; Jaiti et al. 2004). Many studies have reported the killing capacity of H$_2$O$_2$ produced by some bacteria or applied exogenously on nematodes e.g. Caenorhabditis elegans (Gustin et al. 2002; Jansen et al. 2002; Bolm et al. 2004).

To our knowledge, no previous studies were conducted to explore the effect of H$_2$O$_2$ on plant-parasitic nematodes except for its use for clearing cyst walls and for increasing egg hatching when it is applied with irrigation to increase soil aeration (Goodey 1963). Hydrogen peroxide could affect the nematodes directly by its toxicity and/or indirectly as an elicitor triggering the host-plant defense. Therefore, this study was done to investigate the influence of H$_2$O$_2$ applied exogenously on M. javanica and its interaction with tomato.

MATERIALS AND METHODS

Source of RKN Inoculum

Inoculum of M. javanica was obtained from a population that was naturally infecting eggplant plants in the central part of Jordan Valley. Galled root samples were washed thoroughly under tap water for 5 minutes. Small-galled roots were excised and examined under a dissecting microscope for finding developmental stages of the RKNs. Identification of M. javanica was done by observing female perineal patterns and measuring length of second-stage juveniles (Barker et al. 1985) and confirmed by species-specific SCAR-PCR (Karajeh 2004). The nematode eggs were extracted from galled roots by using 0.5% sodium hypochlorite (Hussey and Barker 1973).

Plant Material and Treatments

Seeds of tomato cultivar Speedy were sown in nursery polystyrene trays filled with a pasteurized mixture of peatmoss, perlite and clay soil (1:2:1) in the greenhouse. The tomato plants were transplanted to pots and were transferred to a growth chamber (25 ± 3°C air temperature and 16 h day). Twenty-nine treatments were arranged in a completely randomized design with 10 replicates. A technical grade of H$_2$O$_2$ was used. The treatments were soil drenching with H$_2$O$_2$ at 7 concentrations (1, 10, 100, 250, 500, 750 and 1 000 mM) in plant rhizosphere at three different application times: 24 hours before the time of plant inoculation with the RKN (T1), at the time of inoculation (T2), and 24 hours after the inoculation (T3). Treatments were repeated on un-inoculated plants (NI) and some inoculated plants were not treated with H$_2$O$_2$, used as controls. The seedlings were inoculated with 3 000 eggs of M. javanica per pot. The experiment was ended after sixty days. The nematode eggs were re-extracted
from treated plants by using 0.5% NaOCl. The number of eggs/g root fresh weight was then determined.

Analysis of data
Data were statistically analyzed using the general linear model (GLM) procedure of the system of analytical statistics (SAS). Least significant difference (LSD) test was used for mean separation at 0.05 probability (Steel and Torrie 1986).

Determination of $H_2O_2$ Content
Plant content of endogenous hydrogen peroxide was determined five days after the exogenous application according to Velikova et al. (2000). Briefly, plant material (500 mg fresh weight) was homogenized in 2 ml trichloroacetic acid (TCA) solution (1 g/l). After centrifugation at 12000Xg for 15 minutes, 0.5 ml of the supernatant was added to the reaction mixture containing 0.5 ml of 10 mM potassium phosphate buffer (pH 7.0) and 1 ml of 1 mM KI. Absorbance was determined at 390 nm using a UV spectrophotometer (Helios Alfa, Thermo Electron Corporation, USA) and the amount of $H_2O_2$ was evaluated using a standard curve under the same conditions.

Determination of Chlorophyll Content
Chlorophyll content was determined at the end of the experiment according to Sestak et al. (1971). Samples (30 mg of fresh leaves) were immersed in 5 ml of 96% ethanol at 80°C for 10 min to extract the pigments. The absorbance of extracts was measured at 663 and 647 nm using the UV spectrophotometer. The amount of chlorophyll (chlorophyll a + chlorophyll b) was evaluated using a standard curve determined under the same conditions.

RESULTS AND DISCUSSION
Compared with the untreated control, the RKN reproduction, expressed by the number of eggs/g fresh root was significantly reduced in all $H_2O_2$ treatments. The lowest reduction in RKN reproduction occurred at 10 mM $H_2O_2$ concentration used (Fig. 1). Regarding the application time of exogenous $H_2O_2$, T1 and T2 were definitely significantly higher in reducing the nematode reproduction than T3 at 250, 750 and 1000 mM $H_2O_2$. The reduction in the RKN reproduction rate could be correlated with the direct killing effect of $H_2O_2$ of the nematode eggs and hatched juveniles. Hydrogen peroxide vapor was reported to have the ability to kill the eggs of *C. elegans* in vitro (Gustin et al. 2002). Streptococcus bacteria are able to kill *C. elegans* and this killing is only mediated by hydrogen peroxide. Streptococci can produce sufficient amounts of hydrogen peroxide to kill *C. elegans*, with killing kinetics similar to those of equimolar concentrations of pure hydrogen peroxide (Jansen et al. 2002; Bolm et al. 2004).

Content of endogenous $H_2O_2$ in plants increased at higher concentrations of exogenous $H_2O_2$ (≥ 500 mM) in treated plants compared with the control plants regardless the time of application (Fig. 2). There were significant differences between the treatments with exogenous $H_2O_2$ used and the untreated control of uninoculated plants in the content of endogenous $H_2O_2$ (Fig. 2). In general, chlorophyll content of treated plants was reduced with the increase in the concentration of exogenous $H_2O_2$ used except for 1 and 10 mM where there was no difference (Fig. 3). At the higher concen-
tration of exogenous H$_2$O$_2$ treated plants were yellowish (low in chlorophyll content); some plants were stunted with relatively small root systems and some plants died before the end of the experiment. No significant differences were found among T1, T2 and T3 in plant contents of endogenous H$_2$O$_2$ and chlorophyll (Fig. 2 and 3).

The use of exogenous H$_2$O$_2$ at low concentrations (1–100 mM) did not affect plant growth. As a result of the application of exogenous H$_2$O$_2$, there were elevated levels of endogenous H$_2$O$_2$ in the treated plants. Hydrogen peroxide is known as mobile and reactive compound in plants and in fact, tomato plants are known to accumulate relatively high levels of H$_2$O$_2$ (Orozco-Cárdenas and Ryan 1999). Hydrogen peroxide has

![Fig. 1. Effect of soil drenching with hydrogen peroxide on the reproduction of Meloidogyne javanica at three application times (DBI (T1): one day (24 hours) before the time of plant inoculation with the RKN, AI (T2): at the time of inoculation, and DAI (T3): one day (24 hours) after the inoculation). Columns followed by the same letters are not significantly different according to least significant (LSD) test at 0.05 probability level.](image1)

![Fig. 2. Plant content of hydrogen peroxide five days after treatment with exogenous hydrogen peroxide at different concentrations in plants uninoculated (NI) or inoculated with Meloidogyne javanica (DBI (T1): one day (24 hours) before the time of plant inoculation with the RKN, AI (T2): at the time of inoculation, and DAI (T3): one day (24 hours) after the inoculation). H$_2$O$_2$ content means (LSD$_{0.05} = 0.25$) are not significant at 1, 10, 100 and 250 mM H$_2$O$_2$ concentrations.](image2)
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Fig. 3. Chlorophyll content of hydrogen peroxide five days after treatment with exogenous hydrogen peroxide at different concentrations in plants uninoculated (NI) or inoculated with *Meloidogyne javanica* (DBI (T1): one day (24 hours) before the time of plant inoculation with the RKN, AI (T2): at the time of inoculation, and DAI (T3): one day (24 hours) after the inoculation). Chlorophyll content means (LSD$_{0.05} = 0.65$) are not significant at 1 and 10 mM H$_2$O$_2$ concentrations.

been reported to play an important role in the development of disease resistance in plant species (Grant et al. 2000). Its accumulation in specific tissues, and in the appropriate quantities, is of benefit to plants and can mediate cross-tolerance toward other stresses (Bolwell 1999). The exogenous application of H$_2$O$_2$ and ethephon protected tomato seedlings from chilling injury (Al-Haddad et al. 2002). The response to H$_2$O$_2$ in tomato plant may be associated with the defense of plants against both herbivores and pathogens (Orozco-Cardenas and Ryan 1999). Hydrogen peroxide in plant cells has been shown to regulate the hypersensitive response and cell death as a defense response against pathogen attack (Klessig et al. 2000). At the higher concentrations of H$_2$O$_2$ (≥ 500 mM), the apparent toxicity in the treated plants might be due to the direct exposure of plant roots to the exogenous H$_2$O$_2$ within few days after the application or due to the extreme accumulation of the endogenous H$_2$O$_2$ in plant leaf tissues. As a conclusion, the reduction of RKN reproduction may be attributed to the direct effect of the exogenous H$_2$O$_2$ on the nematode eggs and hatched juveniles and indirectly to the H$_2$O$_2$ effect on the treated tomato plants that could be elicited by H$_2$O$_2$ to resist the nematode infection.

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REFERENCES


**POLISH SUMMARY**

**WSPÓLDZIALENIE MATWIKA KORZENIOWEGO MELOIDOGYNE JAVANICA I ROŚLIN POMIDORA WYWOŁANE NADTLENKIEM WODORU**

Badano wpływ nadtlenku wodoru \( (H_2O_2) \) na porażenie korzeni pomidora matwiakiem *Meloidogyne javanica*. W tym celu nasączono ziemię roztworami nadtlenku wodoru o zróżnicowanym stężeniu \( (1, 10, 100, 250, 500, 750 \) i \( 1000 \) mM \( H_2O_2 \)\) w trzech terminach: T1 – 24 h przed inokulacją patogenem; T2 – w czasie inokulacji; T3 – 24 h po inokulacji. We wszystkich traktowanych kombinacjach uzyskano istotne zmniejszenie reprodukcji matwika (jaja/g świeżej masy korzeni) w porównaniu do nietraktowanej kombinacji kontrolnej. Najniższy stopień zmniejszenia reprodukcji nicienia stwierdzono przy użyciu stężenia \( 10 \) mM \( H_2O_2 \). Przy stosowaniu stężeń \( 250, 750 \) i \( 1000 \) mM \( H_2O_2 \) w terminach T1 i T2 wystąpiło istotnie wyższe ograniczenie tempa reprodukcji szkodnika, niż w terminie T3. Zawartość endogennego \( H_2O_2 \) w traktowanych roślinach była istotnie niższa niż w roślinach nietraktowanych. W przypadku wyższych stężeń \( (\geq 500 \) mM \( H_2O_2 \) \) stwierdzono na traktowanych roślinach fitotoksyczność, co było wynikiem nagromadzenia endogennego \( H_2O_2 \). W kombinacjach za stężeńiami \( 1 \) i \( 10 \) mM \( H_2O_2 \) rośliny nie różniły się pod względem zawartości chlorofilu od roślin w kombinacji kontrolnej, podczas gdy przy użyciu stężeń wyższych zawartość chlorofilu była istotnie niższa. Stwierdzono, że zewnętrzne zastosowanie \( H_2O_2 \) może wywierać bezpośrednie działanie na reprodukcję *M. javanica*, oraz pośrednie działanie na rośliny pomidora, które stają się mniej wrażliwe na zakażenie tym patogenem.