Extracts of Killed *Penicillium chrysogenum* Induce Resistance Against Fusarium Wilt of Melon

Hezhong Dong^{1,2} and Yigal Cohen^{1,*}

was mediated by induced resistance. The resistance induced by DME protected melon plants a race-non-specific resistance against Forn. Both DME and NDME significantly increased not only against race 1,2, but also against the three other races of the pathogen, indicating (0.5-5%) had any effect on fungal growth in vitro, which implied that disease control in vivo plants treated 0-6 h before inoculation or 0-48 h after inoculation. Neither DME nor NDME the disease compared with water-drenched, challenged seedlings. No protection was seen in drench with 2-5% DME applied 12-72 h before inoculation provided significant control of dry mycelium extract (NDME) were protected against challenge infection with Fom. A single days after emergence) treated with either acidic dry mycelium extract (DME) or neutralized before or after inoculation with Fusarium oxysporum f.sp. melonis (Fom). pharmaceutical industry, was extracted with water and applied to the roots of melon plants Dry fungal biomass of Penicillium chrysogenum (dry mycelium), a waste product of the in the defense mechanisms activated by DME or NDME. peroxidase activity but not with free L-proline content. Thus, peroxidase might be involved respectively. Resistance to Fusarium wilt was significantly associated with elevated levels of peroxidase activity and free L-proline content in seedlings 12 h and 48 h after soil drench Seedlings (4-6

induced resistance; peroxidase; L-proline. KEY WORDS: Penicillium chrysogenum; Fusarium wilt; Fusarium oxysporum f.sp. melonis:

INTRODUCTION

induction period necessary for the activation of the plant's defense mechanisms (20) in plants that are susceptible to a pathogen's attack and is normally achieved after an (7,20). In contrast to genetic resistance, acquired or induced resistance may be activated own defense system by certain chemical compounds has attracted attention in recent years control. Among these, the technology for disease control based on activation of the plant's environment has enhanced the need for development of alternative methods of disease resistant cultivars or fungicides (3). disease of melon. which makes it less likely to be overcome by the pathogens (20). long duration of expression. Induced resistance is based on multiple defense mechanisms Induced resistance is often characterized by a broad spectrum of protection and relatively Fusarium wilt caused by Fusarium oxysporum f.sp. melonis (Fom) is a devastating Current measures for control are based mostly on soil disinfection, The necessity of reducing pesticide levels in the

Received March 25, 2001; received in final form July 9, 2001; http://www.phytoparasitica.org posting Aug. 31 2001. *Author for correspondence [Fax: +972-3-5354133; e-mail: coheny@mail.biu.ac.il].

1-Faculty of Life Sciences, Bar-Ilan University: Ramat Gan 52900, Israel.

On sabbatical leave from Cotton Research Center, Shandong Academy of Agricultural Sciences, Jinan 250100

soil protected corn plants against Fusarium moniliforme. and improving soil fertility (13). Recently Gao et al. (10) reported that DM applied to the faba beans (17). When added to the soil, DM enhances plant growth by supplying nutrients that live P. chrysogenum is a potential biological control agent against Botrytis fabae in fungal biomass of Penicillium chrysogenum after extraction of penicillin. It was reported of SA (4,9,29). Some plant extracts (8,15) were also reported to protect plants against produced naturally in plants, is involved in signaling for induced resistance (20). Synthetic a group of mRNAs encoding PR-1, chitinases, β -1,3-glucanases and peroxidase (16). SA. reported by Cohen and co-workers to be effective in inducing local and systemic resistance against fungal diseases (including *Fom*) and nematodes (4,5,7,21). Zimmerli *et al.* (31) and certain pathogens. inducers such as INA, BTH and BABA have been reported to induce resistance downstream mosaic virus, respectively. Induced resistance is associated with increased transcription of Siegrist et al. (26) reported that 3-aminobutyric acid protects against bacteria and tomato Recently, a new class of compounds, belonging to 3-aminobutyric acids (BABA), has been pathogens without possessing any direct antimicrobial activity in vitro or in vivo (9,11,14). ester (BTH), are able to induce resistance in a number of plants against a wide range of 2,6-dichlorisonicotic acid (INA) and benzo[1,2,3]thiadiazole-7-carbothionic acid-S-methyl application of chemicals (2,4-6,11,14). Various chemicals, such as salicylic acid (SA) Induced resistance can be achieved by using biotic inducing agents or through Dry mycelium (DM) used in this study is made from the dry killed

against F.o. f.sp. melonis in melon, with particular reference to the relationship between resistance and peroxidase activity or L-proline content. induced resistance. biomass on peroxidase activity and L-proline accumulation, or on their relationship to to environmental stress (30). However, very little is known about the effect of dry fungal peroxidase, L-proline accumulation has also been correlated in many plants with resistance Peroxidase activity could also be a biochemical marker for genetic resistance (22-24). Like be directly associated with the increased ability of protected tissue to lignify Peroxidase catalyzes the final polymerization step of lignin synthesis, and may therefore mildew of wine grapes and result in a remarkable increase in soluble peroxidase activity. Reuveni et al. (22) noted that foliar application of phosphates could control powdery In this paper we describe how extracts of DM can induce resistance (12).

MATERIALS AND METHODS

DME (NDME, pH=6.8) was obtained by adding 0.1 M KOH. 10% DME and 10% NDME at 110°C and the pH was measured after cooling (10% DME, pH=2.6-2.8). and filtered through Whatman No. 1 filter paper. The filtrate was autoclaved for 30 min h at 100 rpm, and then stored for 22 h at room temperature. It was then briefly agitated 100 g of DM was suspended in 1000 ml distilled water (= 10% DME, w/v), shaken for 2 for 4 h at 110°C. Dry mycelium extract (DME) was prepared by the following procedure: from Biochemie Ltd., Kundl, Austria. This fungal biomass was dried by the manufacturer were stored as stock solutions at 4°C Preparation of extracts Penicillin-free DM (powder) of P. chrysogenum was obtained Neutralized

of perlite and peat (1:1, v/v) and allowed to grow in the greenhouse (18-28°C). After emergence (usually 8 days after sowing), ten seedlings of uniform size were left in each pot. Plants and DME (and NDME) treatment Twenty seeds of melon (Cucumis melo L.) 'En-Dor', which is susceptible to Fom, were sown in 0.5-l pots containing a mixture

Each pot was drenched with DME or NDME (20 ml per pot) of various concentrations (0.5water per pot, starting 3 days after treatment. 10%) using a pipette. Potted plants drenched with 20 ml distilled water served as controls unless indicated otherwise. Plants were watered every 2 days with an equal volume of

conidia per ml with a cytometer before inoculation. 15-day-old cultures into distilled water and their concentration was adjusted to 106-107 grown in 9-cm petri dishes on PDA at 25°C in the dark. Conidia were removed from 10-Fungal inoculum The four known races of Fom, race 0, race 1, race 2 and race 1,2, were

calculated. Each experiment was carried out with five replicates of ten plants each, and at 6-8 days after inoculation, and the percentage of diseased and protected plants was wilt symptoms 3-4 days after inoculation. The number of wilted seedlings was counted greenhouse under the conditions described above. Inoculated seedlings usually exhibited root tips were cut off, the root system dipped for 1 min in the conidial suspension, and then inoculation, seedlings were carefully removed from the soil, thoroughly washed with water, with race 1,2 of Fom 48 h before DME-treatment or 0-72 h after DME-treatment. For evaluate the effect of time of DME-treatment on disease control, plants were also inoculated of Fom 2-3 days after application of DME or NDME, unless indicated otherwise. was repeated at least twice. transplanted into fresh potting mixture. Inoculation and disease assessment Plants were inoculated with one of the four races Inoculated plants were allowed to grow in the

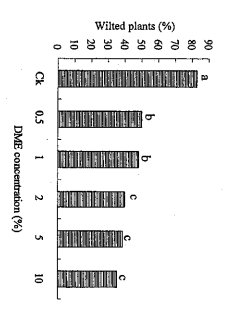
three mycelial plugs (2×4 mm) of race 1,2 of Fom per petri dish (three plates / extract / were measured at 2 and 4 days after inoculation. concentration). was autoclaved for 30 min at 110°C, poured into 9-cm petri dishes, and inoculated with **Fungitoxicity tests** The inoculated plates were kept at 25°C in the dark and colony diameters Czapek Dox agar supplemented with 0.5-5% (v/v) DME or NDME

supernatant was measured and expressed as the change in absorbance per minute per gram the suspension was centrifuged (10,000 g for 10 min at 4°C). Peroxidase activity of the or inoculation, thoroughly washed with water, blotted dry and weighed. One whole plant either treated or untreated with DME or NDME, were used for assessing peroxidase activity fresh weight (22). Four replicate plants per treatment were used in each experiment. and free L-proline content. Plants were removed from soil after DME (or NDME) treatment Peroxidase activity and L-proline content 0.5 g) was ground in 10 ml of cold 15 mM sodium phosphate buffer (pH 6), Non-inoculated or Fom-infected plants

homogenate was centrifuged (10,000 g for 5 min) The supernatant was treated with acetic (1). Briefly, samples (\sim 0.5 g each) were homogenized with 3% sulfosalicylic acid and the was extracted and its concentration was determined according to the method of Bates et al. acid and ninhydrin solution, boiled for 1 h, and the absorbance at 520 nm was determined Amounts of proline were expressed as ng/g fresh weight. Plant samples for L-proline assessment were obtained as described above. L-proline

peroxidase activity and L-proline content were calculated using Excel (Microsoft, Bothell some bearing on plant protection, Spearman's rank correlation coefficient (r) test (P=0.01) establish significant difference (P=0.05). To test if peroxidase activity and L-proline had was also performed, based on data in Figures 4, 5 and 6 WA, USA). Analysis of variance was performed by Duncan's Multiple Range Test to Means and standard deviations (SD) of diseased plants percentage

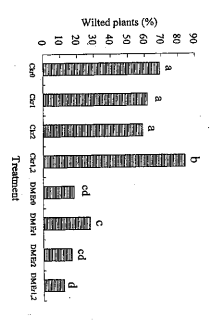
or NDME at up to 5% had no fungicidal activity against Fom. pathogen. Colony diameter (range) did not differ significantly between DME- or NDME-Dox agar media to study the possible fungitoxic activity of DME against race 1,2 of the wilt free (control) and DME- or NDME-amended plates (data not shown), indicating that DME Various concentrations (0-5%) of DME or NDME were added to Czapek



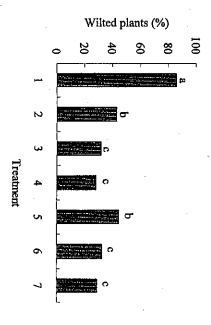
soil drench 2 days before inoculation. Plants drenched with water were used as control. Data were by dry mycelium extract (DME). Plants were treated with various concentrations of acidic DME by according to Duncan's multiple range test. recorded 8 days after inoculation. Columns with the same letter do not differ significantly (P=0.05) Fig. 1. Protection obtained in melon plants against Fusarium oxysporum f.sp. melonis (race 1,2)

inoculation, respectively but non-inoculated plants (data not shown). Protected plants started to wilt 15 days after enhancement of protection was observed, and phytotoxic symptoms were seen in treated against wilt 8 days after inoculation. A significantly higher level of protection (52%) inoculation. before inoculation to various concentrations of DME (0.5-5%) were significantly protected was obtained with 2% Disease control by DME Control and DME-treated plants (2%) were wilted at 10 and 21 days after DME than with 0.5-1% DME (40%). Data in Figure 1 show that melon plants exposed 2 days At 5-10% no further

protection against races 0, 1 and 2 was 74%, 55% and 71%, respectively. inoculation. Plants treated with 2% DME were 85% protected against this race. Percentage 1,2 was the most aggressive, with 85% of the untreated plants evincing wilt 7 days after Protection induced by DME is race-non-specific DME protected melon plants against Fom of race 0, 1, 2 or 1,2. Among the four races, race Figure 2 presents data showing that



differ significantly (P=0.05) according to Duncan's multiple range test. Ck = control inoculated before inoculation. Data were recorded 7 days after inoculation. Columns with the same letter do not Fig. 2. Protection obtained in melon plants by 2% dry mycelium extract (DME) against four races of Fusarium oxysporum f.sp. melonis (t0, r1, r2, and r1,2). 2% DME was applied by soil drench 3 days

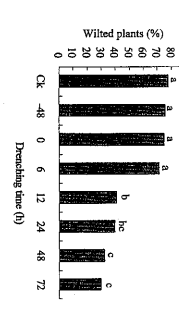


and neutralized DME (NDME) against Fusarium oxysporum f.sp. melonis (race 1,2). Plants were treated by soil drench of 0.5-2% DME or 0.5-2% NDME 2 days before inoculation, or with water as the control. Data were recorded 8 days after inoculation. 1, control; 2, 0.5% DME; 3, 2% DME; 4, 5% DME; 5, 0.5% NDME; 6, 2% NDME; 7, 5% NDME. Columns with the same letter do not differ significantly (P=0.05) according to Duncan's multiple range test. Fig. 3. Protection obtained in melon plants by various concentrations of dry mycelium extract (DME)

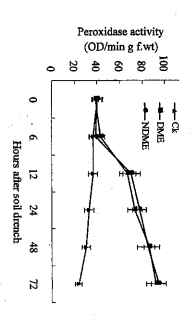
after inoculation both DME and NDME significantly reduced the percentage of wilted 62% and 66% protection, respectively. provided 50%, 63% and 67% protection, and NDME at those concentrations gave 48% wilted plants between the two extracts of equal concentrations. DME at 0.5%, 2% and 5% plants relative to the control. No significant difference was found in the percentage of Effect of NDME on induced protection Data presented in Figure 3 show that 8 days

h after inoculation or 0-6 h before inoculation did not suppress Fusarium wilt, whereas a recorded at 6 days after inoculation. Effect of time of DME application on induced protection A soil drench with 2%.DME applied either 0-48 Data in Figure 4 were

required for DME to control Fom effectively. done 12-72 h before inoculation. It seems that a root exposure period of at least 12 h was significant decrease in the percentage of wilted plants occurred when the soil drench was



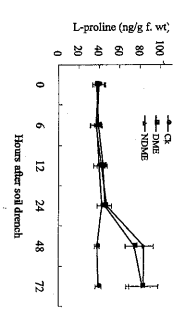
Data were recorded 6 days after inoculation. Columns with the same letter do not differ significantly either before (-) or after inoculation. 0 indicates that 2% DME was applied soon after inoculation. melon against Fusarium oxysporum f.sp. melonis. Soil drench with water (Ck) or 2% DME was done Fig. 4. Time-dependent efficacy of dry mycelium extract (DME) application on protection obtained in (P=0.05) according to Duncan's multiple range test.



2% dry mycelium extract (DME) or 2% neutralized DME (NDME). Ck = soil drench with water. Plants were treated 3 days after emergence. Bars represent ±SD. Fig. 5. Changes in peroxidase activity in healthy melon plants (not inoculated) after soil drench with

significant (P = 0.01) negative correlation between peroxidase activity and percentage of wilted plants (r = -0.98). peroxidase activity in both DME-treated and NDME-treated plants, relative to control Spearman's rank correlation coefficient test on the data in Figures 4, 5 and 6 showed a or NDME-treated plants did not accumulate significantly until 48-72 h after treatment. plants, occurred at 12-72 h after treatment. Data in Figure 6 show that L-proline in DMEequally low at 0-6 h after treatment. that peroxidase activity in water-treated, Effect of DME on peroxidase activity and L-proline content Significant enhancements (up to fourfold) in DME-treated and NDME-treated plants was Data in Figure 5 show

healthy plants. L-proline content was not measured in Fom-inoculated plants did water-treated inoculated plants exhibit a significantly higher peroxidase activity than treated healthy plants (not inoculated), whereas only at 6 days (3 days after inoculation) peroxidase activity was seen in DME-treated inoculated plants as compared with DME compared with water-treated plants. Four and 6 days after soil drench, significantly higher application, a fourfold increase in peroxidase activity was measured in DME-treated plants activity in both non-inoculated and Fom-inoculated plants. Other data presented in Figure 7 show that DME significantly increased peroxidase vity in both non-inoculated and *Fom*-inoculated plants. Three days after DME



Bars represent ±SD. Fig. 6. L-proline content in melon plants after application of 2% dry mycelium extract (DME) or 2% neutralized DME (NDME). Ck = soil drench with water. Plants were treated 3 days after emergence.

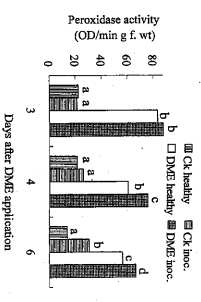


Fig. 7. Peroxidase activity in Fusarium oxysporum f.sp. melonis-inoculated (race 1,2) and non-inoculated melon plants after either water (Ck) or 2% dry mycelium extract (DME) treatment. For Duncan's multiple range test. each time interval, columns with the same letter do not differ significantly (P=0.05) according to

DISCUSSION

made from P. chrysogenum were protected against wilt caused by Fom. Data from several We report herein that melon plants growing in pots drenched with a water extract of DM

also against the three other races of the pathogen, indicating a race-non-specific resistance the disease control effected by DME or NDME was probably based on induced resistance DME had no negative effect on growth of the pathogen in vitro. neutralized DME (NDME, pH 6.8) was as effective as acidic DME against the disease control was observed. Water extract of DM is quite acidic (pH 2.6-2.8). We found that effect increased. However, at concentrations of 5% or higher, no further increase in disease experiments revealed that DME may provide up to 85% protection against Fusarium will relative to water-treated control. In general, as the concentration of DME rose, the control The resistance induced by DME was effective not only against race 1,2, but Thus, it is inferred that

leaves or not translocated to the root system. not protect against Fom (data not shown), indicating that DME was either not taken up by by the plant. Unlike other inducers such as BABA (4,5), DME spray on melon leaves did time was necessary for DME uptake by the root and/or activation of defense mechanisms applied 12 h or more before challenge inoculation with Fom, indicating that an interval of inoculation, implying that induced resistance by a single drench of DME could supply Additional treatment with DME might be needed to boost protection. DME had to be only a limited duration of relative to water-treated control. However, the surviving plants wilted 21 days after found that DME provided significant protection against Fom for 15 days after inoculation pressure often causes insufficient protection against the pathogen (19). In this study, we concentration and application mode of activators and infection pressure. species and pathogens (18,20). Expression of induced resistance is also associated with long duration of expression (20), but its expression is usually dependent on activators, plan Induced resistance is characterized by a broad spectrum of protection and relatively protection under high infection pressure (107 spores per ml) High infection

was closely associated with enhanced levels of peroxidase activity. We assume that the 12 h between DME application and the onset of resistance is required for this phenomenon enhanced peroxidase activity may contribute to induction of resistance. A time interval of reported (22-24,28) that infection with pathogens or application of inorganic chemicals host plants (18). The role of oxidative enzymes, such as peroxidase, and of their metabolic treatment and/or inoculation with Fom. peroxidase activity. Peroxidase activity was raised only after a lag period following DME genotypes of muskmelon, those with high peroxidase activity often had higher levels of enhances peroxidase activity, and this is often associated with resistance. Among different products in the defense mechanisms of infected plants, has also been studied. It has been the accumulation of PR-proteins and other biochemical and physiological changes in the formation of cell wall apposition at sites of attempted penetration by fungal pathogens. responses include processes of oxygen burst (26), lignification of host cell walls (25), the induced resistance were conducted with various plant species. In general, induced defense During the last decade a number of studies concerning the mechanisms responsible for The data presented in this paper demonstrate that DME increased It was found that induced resistance by DME

may function in the storage of energy, amino nitrogen and reducing power (27). We have (30). In addition to its role in osmoprotection, proline accumulation in plants under stress accumulation has been correlated in many organisms with adaptation to osmotic stress Environmental stress can increase accumulation of free proline in many plants. Proline

is needed before final conclusions are drawn. incidental physiological response to DME application rather than a factor in the induced DME significantly increased L-proline content in melon plants. accumulation and induced resistance is still unclear. We found that 48 h after application, foliar application (unpublished data). However, the direct relationship between proline found that BABA significantly increases L-proline content in tomato leaves 2 days after resistance. Nevertheless, further investigation into its relationship with induced resistance ~ 24 h later than that of peroxidase activity and no significant correlation with induced to Fom was found. It seems, therefore, that L-proline accumulation is an This increase occurred

resistance against root diseases such as Fom. Its application may have multiple effects nutrients to crop plants and improving humus in the soil, but also a good inducer of DME and DM powder against diseases under field conditions for both crop nutrition and crop protection. Future research should examine the efficacy of In conclusion, DM of P. chrysogenum is not only an organic fertilizer providing

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