Race 3, a New and Highly Virulent Race of *Fusarium oxysporum* f. sp. *niveum* Causing Fusarium Wilt in Watermelon

X. G. Zhou, Texas A&M University System, AgriLife Research and Extension Center, Beaumont 77713; K. L. Everts, University of Maryland, Lower Eastern Shore Research and Education Center, and University of Delaware, Georgetown 19947; and B. D. Bruton, United States Department of Agriculture—Agricultural Research Service, South Center Agricultural Research Laboratory, Lane, OK 74555

**ABSTRACT**


Three races (0, 1, and 2) of *Fusarium oxysporum* f. sp. *niveum* have been previously described in watermelon (*Citrullus lanatus*) based on their ability to cause disease on differential watermelon genotypes. Four isolates of *F. oxysporum* f. sp. *niveum* collected from wilted watermelon plants or infested soil in Maryland, along with reference isolates of races 0, 1, and 2, were compared for virulence, host range, and vegetative compatibility. Race identification was made on the watermelon differentials Sugar Baby, Charleston Gray, Dixielee, Calhoun Gray, and PI-296341-FR using a root-dip, tray-dip, or pipette inoculation method. All four Maryland isolates were highly virulent, causing 78 to 100% wilt on all differentials, one of which was PI-296341-FR, considered highly resistant to race 2. The isolates also produced significantly greater colonization in the lower stems of PI-296341-FR than a standard race 2 reference isolate. In field microplots, two isolates caused over 90% wilt on PI-296341-FR, whereas no disease was caused by a race 2 isolate. All four isolates were nonpathogenic on muskmelon, cucumber, pumpkin, and squash, confirming their host specific pathogenicity to watermelon. The Maryland isolates were vegetatively compatible to each other but not compatible with the race 2 isolates evaluated, indicating their genetic difference from race 2. This study proposes that the Maryland isolates belong to a new race, race 3, the most virulent race of *F. oxysporum* f. sp. *niveum* described to date.

*Fusarium oxysporum* f. sp. *niveum* (E.F. Sm.) W.C. Snyder & H.N. Han., the casual agent of Fusarium wilt of watermelon (*Citrullus lanatus* (Thunb.) Matsum. & Nakai), is widespread in watermelon-growing regions of the world, and frequently is the major factor limiting triploid (seedless) watermelon production. In the United States, the disease was first described by Smith in 1894 in South Carolina and Georgia (33). *F. oxysporum* f. sp. *niveum* is host specific and causes disease only on watermelon, although a few exceptions have been reported in studies conducted under greenhouse conditions (21,40).

Differences in virulence among isolates of *F. oxysporum* f. sp. *niveum* have long been recognized (25,30) and isolates are subdivided into three races on the basis of virulence on watermelon cultivars that vary in their level of resistance (2,6,21,27,37). In 1963, Crall (8) described two physiologic races of *F. oxysporum* f. sp. *niveum* in Florida according to virulence on cultivars such as Florida Giant (susceptible), Charleston Gray (moderately resistant), and Summit (highly resistant) but did not provide a race designation. In 1972, Cirulli (6) also differentiated two races in Italy and designated them as race 0 and race 1 according to disease reactions on Sugar Baby (susceptible), Charleston Gray, and Calhoun Gray (highly resistant). Race 0 caused wilt only on the susceptible cultivar, while race 1 caused wilt on cultivars classified as moderately resistant to Fusarium wilt. In 1973, Netzer and Dishon (28) reported a highly aggressive isolate and classified it as a new race. The race was virulent toward race-1-resistant cultivars such as Calhoun Gray, Summit, and Smo kylee (27,28). They designated this isolate as race 2 following the numbering system of Cirulli (6). In 1985, Martyn (19) reported two isolates of race 2 from Texas, both of which caused severe wilt in all known resistant commercial watermelon cultivars (20). Since then, race 2 was reported in Oklahoma in 1988 (5), in Florida in 1989 (22), and in Maryland and Delaware in 2001 (36). In 2003, Zhou and Everts (37) reported that race 2 was widely distributed in Maryland and Delaware, present in 22% of the watermelon production fields surveyed. More recently, race 2 was found in Georgia (9), Indiana (10), and South Carolina (A. P. Keinath, personal communication). Race 2 also has been reported in China (9) and in the eastern Mediterranean region, including Cyprus (14), Greece (27), Spain (11), and Turkey (27).

Martyn and Netzer (24) proposed inclusion of PI-296341-FR (*C. lanatus var. citroides*) as a differential for classifying races of *F. oxysporum* f. sp. *niveum*, along with Sugar Baby, Charleston Gray, and Calhoun Gray (Table 1). By their system, race 0 causes wilt only on Sugar Baby; race 1 causes wilt on Sugar Baby and Charleston Gray but not on Calhoun Gray and PI-296341-FR; and race 2 causes wilt on all differentials except PI-296341-FR. Therefore, PI-296341-FR is the key to differentiating any new and more virulent race of *F. oxysporum* f. sp. *niveum* from race 2.

However, difficulties in differentiating races of *F. oxysporum* f. sp. *niveum* have been encountered in some previous studies (1,17,22). This is because races of *F. oxysporum* f. sp. *niveum* are not defined as clearly as in many other formae speciales of *F. oxysporum*, where races are based on a gene-for-gene relationship on differential cultivars (17,31). Therefore, races within *F. oxysporum* f. sp. *niveum* were differentiating by disease reaction of watermelon genotypes used to differentiate races of *Fusarium oxysporum* f. sp. *niveum*.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sugar Baby</td>
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<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td>Charleston Gray</td>
<td>R</td>
<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td>Calhoun Gray</td>
<td>R</td>
<td>S</td>
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<td>S</td>
</tr>
<tr>
<td>PI-296341-FR</td>
<td>R</td>
<td>R</td>
<td>S</td>
<td>S</td>
</tr>
</tbody>
</table>

* Race classification system is developed from Cirulli (6), Netzer (27), Martyn (21), Martyn and Netzer (24), and Zhou et al. (42).

* Assignments of disease reaction ratings are based on incidence of Fusarium wilt on the differential genotypes, where R = resistant (<33% wilt) and S = susceptible (≥33% wilt), following the criteria established by Martyn and Bruton (22) and used by other studies (9,37).

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were collected from colonies on K's Fusa-
counties in Maryland. All four isolates
from race 2 and could be classified as a
determine whether they were different
and vegetative compatibility, and
isolates by pathogenicity in greenhouse
Source of isolates. Eleven isolates of F.
oxysporum f. sp. niveum were used in this
study (Table 2). In 2000, MD-ZE085
rium-selective medium (16), single-spore
cultured, and stored in soil tubes at 4°C.
Identification of F. oxysporum f. sp.
niveum was according to fungal morphol-
yogy (26) and pathogenicity of each isolate
on the wilt-susceptible watermelon cv.
Sugar Baby. Seven isolates that were pre-
viously categorized to race (one race 1,
one race 0, and five race 2) (Table 1) were
included as reference isolates in this study.
ATCC-44293 and TX-XID1 were pur-
chased from the American Type Culture
Collection (ATCC), Manassas, VA.

Inoculum preparation. To prepare in-
oculum, colonized agar plugs (2 mm in
diameter) from a 7- to 10-day-old colony
onto-dextrose agar (PDA) were trans-
ferred into a liquid mineral salt medium
(27) and grown for 10 to 14 days on a
rotary shaker at 128 rpm at room tempera-
ture. Colonized liquid medium was filtered
through four layers of sterile cheesecloth
and the suspension of spores (>95% mi-
croconidial) was adjusted to approximat-
ely 1 × 10⁶ microconidia/ml using a hemacy-
tometer.

Greenhouse pathogenicity test. Three
methods were used to inoculate seedlings
of watermelon in separate experiments.
The first method was the standard root dip
(18,20). At the first true leaf stage, seed-
lings raised in soilless potting mix consist-
ing of vermiculite-peat moss (2:3, vol/vol)
(Redi-Earth, Scotts-Sierra Inc., Marysville,
OH) were uprooted and the roots washed
gently in water. Seedling roots were
dipped into an inoculum for 30 s and
transplanted into plastic pots (10 by 10 by
8 cm) filled with the same soilless potting
mix with 10 plants per pot. The second
method was tray-dip inoculation (18,22).
Lettuce seedlings were sown in plastic trays
with cells (3 by 3 by 5 cm, 128 cells/tray)
containing the soilless potting mix described
above and then placed inside a larger plastic
flat filled with the same potting mix. After
emergence, seedlings were thinned to one
plant per cell. At the first true leaf stage,
at which time the roots of the seedlings had
grown through the drainage holes of the
trays and into the potting mix in the larger
flat, the trays were lifted and rinsed briefly
with water. The roots of the seedlings were
uniformly trimmed to about 3 cm. The
entire tray was placed in a shallow plastic
flat filled with 300 ml of spore suspension
of the inoculum for 20 min and then re-
turned to the original larger flat. The third
method was the pipette inoculation tech-
nique (18). Seedlings were raised in the
potting mix in plastic pots (6 by 6 by 5
cm) of 10 plants each. At the first true leaf
stage, 3 ml of inoculum was delivered to
the potting mix around each seedling with
a pipette (Labsystems, Helsinki, Finland).

Race was identified based on the re-
action of the following differentials rec-
ommended by previous studies (2,17,20–
22,37). Sugar Baby, Charleston Gray, Dix-
ilee, Calhoun Gray, PI-296341-FL (FL),
PI-296341-FR (OK), and PI-296341-FR
(MD). Two sources of PI-296341-FL, which were originally released in 1991 by
R. D. Martyn and D. Netzer (24), were
included. They were provided by B. Carle
in Florida in 1999 and by B. D. Bruton in
Oklahoma in 2004 as PI-296341-FL (FL)
and PI-296341-FR (OK), respectively. Two
sources of PI-296341-FL were included
because segregation for susceptibility to
race 2 was observed in progeny of the FL
and OK increases of PI-296341-FL to
produce the seed used in our initial tests.
To improve race 2 resistance in the PI
lines, a third PI line, designated as PI-
296341-FR (MD), was developed at the
University of Maryland Lower Eastern
Shore Research and Education Center in
Salisbury by inoculating seedlings of PI-
296341-FR (FL) and PI-296341-FR (OK)
with two race 2 isolates of F. oxysporum f.
sp. niveum in a greenhouse, transplanting
the surviving plants into a field naturally
infested with a mixture of races 1 and 2,
and harvesting seed from watermelon fruit
on plants with no sign of Fusarium wilt.
All other watermelon cultivars were ob-
tained from commercial seed suppliers.

Three evaluations to determine level of
virulence of the Maryland isolates were
conducted. The first evaluation was done
in 2002 using all three inoculation methods
for the two Maryland isolates (MD-ZE085 and MD-ZE125) on the FL increase of the PI line, PI-296341-FR (FL). The second evaluation was conducted in 2005 using only the standard root-dip method, in which the second PI line PI-296341-FR (OK) was included for comparison with PI-296341-FR (FL) and an additional assessment for colonization by *F. oxysporum* f. sp. *niveum* in lower stems of inoculated plants also was made for selected genotypes. Stem colonization was determined based on the procedure described previously (38). At the conclusion of the greenhouse experiment, three plants per replicate were randomly selected from the remaining live plants, and their lower stem sections (5 cm in length starting at the soil line) were collected. The stem sections were washed in tap water, surface disinfected with 0.6% NaOCl for 1 min, and rinsed in tap water four times. The stem samples were further cut into short pieces (about 10 mm long), weighed, and homogenized in sterile water in a blender. The resulting suspensions were filtered through four layers of sterilized cheesecloth and the filtered suspensions were pipetted onto K’s medium for culturing *Fusarium* colonies. After 7 to 10 days at room temperature, colonies on plates were counted and counts converted into CFU per gram of fresh tissue. There were three plates for each of three replicates.

In 2008, the root-dip method was used to evaluate the reaction of Sugar Baby, Charleston Gray, and Calhoun Gray along with the improved race 2 resistant PI line PI-296341-FR (MD). Two additional Maryland isolates (MD-ZE622 and MD-ZE632), obtained in 2007, were included along with the four race-2 isolates (F-032-1, F-100-2, F-17B-R-2, and F-17B1-10) of *F. oxysporum* f. sp. *niveum* for a total of five race 2 reference isolates tested.

In each experiment, each combination of genotype and isolate was represented by three replicate pots, with 10 plants per pot. Water-inoculated plants for each genotype were included as the controls. Seedlings in all experiments were maintained in a greenhouse with the temperature ranging from 19 to 28°C. After inoculation, the percentage of seedlings showing typical symptoms of Fusarium wilt (yellowing, stunting, or wilting) was assessed weekly and the percentage of wilt or death at 4 weeks after inoculation was recorded. To compare disease reactions for each genotype and isolate treatment, a genotype giving mean wilt incidence of ≥33% was considered susceptible and <33% was considered resistant (Table 1). This rating criterion followed the system established by Martyn and Bruton (22), which was later used in other studies for race determination of *F. oxysporum* f. sp. *niveum* (9,37). Each experiment was conducted twice.

**Field microplot test.** Two Maryland isolates, MD-ZE085 and MD-ZE125, were further evaluated for virulence on the two watermelon differentials Calhoun Gray and PI-296341-FR (MD) in field microplots during summer 2008. A third isolate with known race 2 designation (F-100-2) was also included for comparison. The Norfolk “A” loamy sand soil (pH 6.3) used in microplots (76 cm in diameter and 61 cm deep, surrounded by fiber glass) was fumigated with metam-potassium (K-Pam) at approximately 42 m²/m³ under black plastic film. The film was removed after 4 days, and the plots were aerated by turning the soil with a rake twice a week for 4 weeks prior to incorporating inoculum. Inoculum for each isolate was prepared as described earlier and added to the top 20 cm of soil. After 4 weeks, when inoculum density in the soil stabilized at approximately 6,500 CFU/g soil as determined by plating diluted soil on K’s medium, five 4-week-old plants of Calhoun Gray were transplanted in one half of each microplot, and the other half was transplanted with five 4-week-old plants of PI-296341-FR (MD). Microplots not infested with either isolate served as controls. Four replicate microplots for each isolate–genotype combination were arranged in a randomized complete block design. A repeat of this experiment was initiated 2 weeks after transplanting and the plots were placed into the first set of microplots. The second experiment was conducted in a second set of microplots that were located within the same field. The percentage of plants showing symptoms of Fusarium wilt (yellowing, stunting, and/or wilting) was assessed and recorded at 6 weeks after transplanting. Transplant dates for the first and second experiment were 1 and 14 June 2008, respectively. Average air temperature, minimum temperature during the 6-week study period were 23.9, 36.1, and 17.6°C for the first experiment and 23.6, 33.7, and 10.6°C, respectively, for the second experiment.

**Host range test.** In all, 19 cultivars representing seven plant species, including 16 cultivars within two genera of Cucurbitaceae (excluding *Citrullus* spp.), were tested for susceptibility to all the 11 isolates of *F. oxysporum* f. sp. *niveum* in the greenhouse. These were 10 muskmelon (*Cucumis melo* L.) cultivars (Ananas, Honeydew Greenflesh, Honeydew Orange-flesh, Imperial 4-50, Israeli [Old Original], Planter’s Jumbo, Super Star, Tam-Dew [Honey Dew], Top Mark, and Yellow Canada), two cucumber (*Cucumis sativus* L.) cultivars (Vlaspick and Orient Express), two squash (*Cucurbita* spp.) cultivars (Zucchini Elite and Goldbar), two pumpkin (*Cucurbita pepo* L.) cultivars (Wizard and Magic Lantern), a tomato (*Lycopersicon esculentum* Mill.) cultivar (Campbell 1327FR), a spinach (*Spinacia oleracea* L.) cultivar (Seven R), and a corn (*Zea mays* L.) cultivar (Bonus). The wilt-susceptible watermelon cv. Sugar Baby was included as a positive control. Water-inoculated plants were included as a negative control. Inoculations were conducted at 2 to 3 weeks after seeding, when most plants reached the first true leaf stage, using the same root-dip inoculation procedure as described before. Each combination of cultivar and isolate had three replicate pots (10 by 10 by 8 cm) filled with the same soilless potting mix as described before, and each pot contained 10 seedlings. The plants were maintained in a greenhouse with temperatures ranging from 18 to 30°C. Inoculum preparation and disease assessment were the same as greenhouse pathogenicity tests. The experiment was conducted twice.

**Vegetative compatibility test.** Vegetative compatibility tests were performed following the method developed by Pu-halla (29) and Correll et al. (7). Nitrate nonutilizing (nit) mutants were generated on both minimal salts medium (MM) and PDA containing 1.5% KClO₃. Portions of the fast-growing sectors emerging from restricted colonies were transferred to both MM and PDA. Those that produced thin, expansive growth with no aerial mycelium on MM but showed wild-type growth with aerial mycelium on PDA were considered to be nit mutants. The nit mutants were generated from each isolate until two complementary nit mutants were obtained. Complementary mutants were recognized by the robust growth at the interface of the two colonies (a visual indication of heterokaryon formation) when paired on MM under a daily 12-h fluorescent light for 5 to 14 days at 22 to 25°C. Mutants generated from each isolate were characterized as one of three physiological phenotypes (nit1 nitM, nit1 nit3, and nit1 nitM nit3), with the type of growth on MM supplemented with different nitrogen sources (7). In all, 6 to 10 mutants were recovered from each isolate, and three nit mutants (nit1, nitM and nit3) from each isolate were chosen as testers. The tests of each isolate were paired in all combinations with testers of other isolates. If a mutant from one isolate formed a heterokaryon with at least one of the three testers of another isolate, both isolates were considered compatible and placed in the same vegetative compatible group. Each pairing in all combinations with vegetative compatibility group testers was conducted at least three times.

**Data analysis.** All statistical analyses were conducted using the SAS software package (version 8.2, SAS Institute Inc., Cary, NC). Analysis of variance was performed on data with the general linear model procedure. Means for treatments were compared using Fisher’s protected least significant difference at P = 0.05. Disease incidence data were arcsine-square root transformed and CFU/g data transformed into log₁₀ (CFU/g + 1) for analysis. The results of two repeated experiments were combined for presentation if no sig-
significant experiment–treatment interactions were present. The linear regression procedure of SAS was performed to characterize the relationships among stem tissue colonization and incidence of Fusarium wilt.

RESULTS

Greenhouse pathogenicity test. In the first evaluation conducted in 2002, the use of the root-dip, tray-dip, and pipette inoculation methods resulted in similar results (Table 3). The two Maryland isolates, MD-ZE085 and MD-ZE125, were highly virulent and caused more than 78% wilt on all five differentials tested, including PI-296341-FT (FL) that is considered resistant to race 2. In contrast, the race 2 reference isolate TX-X1D produced a high incidence of wilt (≥67%) only on Sugar Baby, Charleston Gray, Dixielee, and Calhoun Gray but not on PI-296341-FT (FL). Both Maryland isolates caused susceptible reactions on PI-296341-FT (FL) whereas TX-X1D caused resistant reactions (Table 3). The small amount of wilt showing on the PI genotype inoculated with TX-X1D resulted from the small percentage of plants segregating to a race-2-susceptible phenotype. The reference isolates of races 0 (TX-471) and 1 (ATCC-44293) produced expected disease reactions. TX-471 caused severe wilt on Sugar Baby but was avirulent to all other genotypes possessing wilt-resistant genes; ATCC-44293 produced a high level of wilt on Sugar Baby, a moderate level of wilt on Charleston Gary, and little or no disease on all other differentials.

In 2005, a similar evaluation was conducted in which a second source of the PI differential, PI-296341-FO (OK), was included for comparisons with PI-296341-FT (FL). In this test, colonization by F. oxysporum f. sp. niveum in lower stems of inoculated plants also was determined in the Dixielee, Calhoun Gray, and both PI lines, which are resistant to races 0 and 1. The two Maryland isolates produced comparably high levels of wilt and stem colonization on all genotypes tested, including both PI-296341-FT (FL) and PI-296341-FO (OK) (Table 4). At 4 weeks post inoculation, both Maryland isolates produced high incidence of wilt in excess of 97% on PI-296341-FT (FL) and PI-296341-FO (OK). However, these two PI lines were resistant to TX-X1D, which induced wilt in less than 33% of the tested plants. The two Maryland isolates colonized stems of both PI lines to a significantly greater extent than TX-X1D. There were no significant (P > 0.05) differences between the Maryland isolates and TX-X1D in wilt incidence and stem colonization on all other differentials except on Dixielee, where Maryland isolates caused more stem colonization than TX-X1D. When regression analysis was made over all isolates and genotypes (Table 4), incidence of Fusarium wilt (Y, percent wilt) was strongly correlated to stem colonization (X, CFU/g tissue), with the regression equation of $Y = 37.35 \log(X + 1) - 7.15(R^2_{adj} = 0.90, P < 0.0001)$. The results of the evaluations conducted in 2002 and 2005

### Table 3. Incidence of Fusarium wilt of five differential watermelon genotypes and disease reaction ratings on PI-296341-FT (FL) inoculated with isolates of *Fusarium oxysporum* f. sp. *niveum* using root-dip, tray-dip, and pipette methods in 2002

<table>
<thead>
<tr>
<th>Inoculation method, isolate</th>
<th>Sugar Baby</th>
<th>Charleston Gray</th>
<th>Dixielee</th>
<th>Calhoun Gray</th>
<th>PI-296341-FT (FL)</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Root-dip</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>TX-471</td>
<td>100 b</td>
<td>7 a</td>
<td>0 a</td>
<td>0 a</td>
<td>5 a</td>
<td>R</td>
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<tr>
<td>ATCC-44293</td>
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<td>67 b</td>
<td>0 a</td>
<td>0 a</td>
<td>0 a</td>
<td>R</td>
</tr>
<tr>
<td>TX-X1D</td>
<td>100 b</td>
<td>97 c</td>
<td>97 b</td>
<td>92 b</td>
<td>30 b</td>
<td>R</td>
</tr>
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<td>MD-ZE085</td>
<td>98 b</td>
<td>100 c</td>
<td>100 c</td>
<td>100 c</td>
<td>98 d</td>
<td>S</td>
</tr>
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<td>MD-ZE125</td>
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<td>100 c</td>
<td>97 bc</td>
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<td>S</td>
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<td>67 b</td>
<td>68 b</td>
<td>20 c</td>
<td>R</td>
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<td>MD-ZE085</td>
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<td>82 c</td>
<td>80 c</td>
<td>81 d</td>
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<td>92 d</td>
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<td>85 b</td>
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<td>92 c</td>
<td>78 c</td>
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</table>

5 Data were averaged over two experiments of three replicated pots or trays (10 plants/pot or tray) each. Incidence of Fusarium wilt was rated at 4 weeks after inoculation. Means in column under each inoculation method followed by the same letter are not significantly different (P = 0.05) according to Fisher’s protected least significant difference test. Wilt incidence data were arcsine-square root transformed prior to analysis.

6 Ratings of disease reaction on PI-296341-FT (FL) were assigned for the isolates based on incidence of Fusarium wilt on the differential genotype, where R = resistant (≤33% wilt) and S = susceptible (≥33% wilt), following the criteria established by Martyn and Bruton (22).

### Table 4. Incidence of Fusarium wilt and stem colonization of six differential watermelon genotypes and disease reaction ratings on the FL and OK increases of PI-296341-FT inoculated with isolates of *Fusarium oxysporum* f. sp. *niveum* using the root-dip method in 2005

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Sugar Baby</th>
<th>Charleston Gray</th>
<th>Dixielee</th>
<th>Calhoun Gray</th>
<th>PI-296341-FT (FL)</th>
<th>PI-296341-FT (OK)</th>
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</thead>
<tbody>
<tr>
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<td>10 b</td>
<td>5 a</td>
<td>15 b</td>
</tr>
<tr>
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<td>100 b</td>
<td>98 c</td>
<td>321 b</td>
<td>427 c</td>
<td>25 b</td>
<td>35 b</td>
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<tr>
<td>MD-ZE085</td>
<td>100 b</td>
<td>100 b</td>
<td>1067 c</td>
<td>477 c</td>
<td>97 c</td>
<td>293 c</td>
</tr>
<tr>
<td>MD-ZE125</td>
<td>100 b</td>
<td>100 b</td>
<td>806 c</td>
<td>674 c</td>
<td>97 c</td>
<td>343 c</td>
</tr>
</tbody>
</table>

5 Data were averaged over two experiments of three replicated pots (10 plants/pot) each. Incidence of Fusarium wilt was rated at 4 weeks after inoculation. Stem colonization (CFU/g fresh tissue) by *F. oxysporum* f. sp. *niveum* was assayed by the following the same experiment by placing serially diluted, homogenized lower stem tissues on K’s medium (16). Means in column followed by the same letter are not significantly different (P = 0.05) according to Fisher’s protected least significant difference test. Prior to analysis, wilt incidence and CFU data were arcsine-square root- and log10(CFU/g + 1)-transformed, respectively. ... = not assayed.

Ratings of disease reaction on PI-296341-FT (FL) or PI-296341-FT (OK) were assigned for the isolates based on mean wilt incidence on the two differential genotypes, where R = resistant (≤33% wilt) and S = susceptible (≥33% wilt), following the criteria established by Martyn and Bruton (22).
with the two Maryland isolates provided the first indication of a possible new, more aggressive race.

In 2008, a third evaluation was conducted using the PI line PI-296341-FR (MD) that had been selected and developed for improved race 2 resistance. Two additional Maryland isolates, MD-ZE622 and MDZE632, obtained in 2007, were also included (Table 5). There was a clear difference between the four Maryland isolates and the five race 2 reference isolates based on disease reaction ratings on PI-296341-FR (MD). Inoculations with the Maryland isolates resulted in equally severe levels of wilt (over 95%) on the genotype while inoculations with the five race 2 isolates produced little or no disease. All four Maryland isolates had identical disease reaction and were capable of causing substantial wilt (over 98%) on Sugar Baby, Charleston Gray, and Calhoun Gray. The five race 2 isolates reacted similarly and produced comparatively severe levels of wilt on these differential cultivars. No disease was produced from inoculations with either TX-471 (race 0) or ATCC-44293 (race 1) on PI-296341-FR (MD) or Calhoun Gray. These results clearly demonstrated that the four new Maryland isolates were a distinct group and separate from race 2 isolates and races 0 and 1 isolates on the basis of disease reactions on PI-296341-FR (MD).

**Field microplot test.** In both experiments, MD-ZE085 and MD-ZE125 produced high incidence of wilt (over 90%) on PI-296341-FR (MD) whereas the race-2 reference isolate F-100-2 caused no disease on PI-296341-FR (MD) (Table 6). All three isolates caused disease in Calhoun Gray, with the incidence of wilt ranging from 95 to 100%.

**Host range test.** None of the 16 cultivars (non-Citrullus spp.) within Cucurbitaceae inoculated with any of the four Maryland isolates developed symptoms of Fusarium wilt (data not shown). The non-Citrullus cultivars included those from four species of Cucurbitaceae: cucumber, muskmelon, pumpkin, and squash. In contrast, the four new Maryland isolates were capable of causing 100% wilt on the watermelon cv. Sugar Baby. No disease was observed on tomato, corn, or spinach inoculated with the Maryland isolates.

**Vegetative compatibility test.** All isolates used in the present study were self-compatible. The four new Maryland isolates were vegetatively compatible with each other but all were incompatible with the reference isolates of races 0, 1, and 2 evaluated.

### Table 5. Incidence of Fusarium wilt of four differential watermelon genotypes and disease reaction ratings on PI-296341-FR (MD) inoculated with isolates of *Fusarium oxysporum* f. *sp. niveum* using the root-dip method in 2008

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Race</th>
<th>Sugar Baby</th>
<th>Charleston Gray</th>
<th>Calhoun Gray</th>
<th>PI-296341-FR (MD)</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX-471</td>
<td>0</td>
<td>63 a</td>
<td>2 a</td>
<td>0 a</td>
<td>0 a</td>
<td>R</td>
</tr>
<tr>
<td>ATCC-44293</td>
<td>1</td>
<td>98 bc</td>
<td>73 b</td>
<td>0 a</td>
<td>0 a</td>
<td>R</td>
</tr>
<tr>
<td>TX-XID</td>
<td>2</td>
<td>100 c</td>
<td>100 d</td>
<td>95 bc</td>
<td>95 bc</td>
<td>R</td>
</tr>
<tr>
<td>F-032-1</td>
<td>2</td>
<td>100 c</td>
<td>95 c</td>
<td>92 bc</td>
<td>92 bc</td>
<td>R</td>
</tr>
<tr>
<td>F-100-2</td>
<td>2</td>
<td>100 c</td>
<td>100 d</td>
<td>98 dc</td>
<td>98 dc</td>
<td>R</td>
</tr>
<tr>
<td>F-17B-R-2</td>
<td>2</td>
<td>95 b</td>
<td>95 c</td>
<td>90 b</td>
<td>90 b</td>
<td>R</td>
</tr>
<tr>
<td>F-17B-1-10</td>
<td>2</td>
<td>100 c</td>
<td>95 c</td>
<td>97 dc</td>
<td>97 dc</td>
<td>R</td>
</tr>
<tr>
<td>MD-ZE085</td>
<td>3</td>
<td>100 c</td>
<td>100 d</td>
<td>98 dc</td>
<td>98 dc</td>
<td>R</td>
</tr>
<tr>
<td>MD-ZE125</td>
<td>3</td>
<td>100 c</td>
<td>100 d</td>
<td>98 dc</td>
<td>98 dc</td>
<td>R</td>
</tr>
<tr>
<td>MD-ZE622</td>
<td>3</td>
<td>100 c</td>
<td>100 d</td>
<td>98 dc</td>
<td>98 dc</td>
<td>R</td>
</tr>
<tr>
<td>MD-ZE632</td>
<td>3</td>
<td>100 c</td>
<td>100 d</td>
<td>98 dc</td>
<td>98 dc</td>
<td>R</td>
</tr>
</tbody>
</table>

7 Data were averaged over two experiments of three replicated pots (10 plants/pot) each. Incidence of Fusarium wilt was rated at 4 weeks after inoculation.

8 Means in column followed by the same letter are not significantly different (P = 0.05) according to Fisher’s protected least significant difference test. Wilt incidence data were arcsine-square root transformed prior to analysis.

### Table 6. Incidence of Fusarium wilt of two differential watermelon genotypes and disease reaction ratings on PI-296341-FR (MD) transplanted in field microplots infested with isolates of *Fusarium oxysporum* f. *sp. niveum*

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Race</th>
<th>Calhoun Gray</th>
<th>PI-296341-FR (MD)</th>
<th>Calhoun Gray</th>
<th>PI-296341-FR (MD)</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>F-100-2</td>
<td>2</td>
<td>100 a</td>
<td>0 a</td>
<td>95 a</td>
<td>95 a</td>
<td>R</td>
</tr>
<tr>
<td>MD-ZE085</td>
<td>3</td>
<td>100 a</td>
<td>90 b</td>
<td>95 a</td>
<td>95 a</td>
<td>S</td>
</tr>
<tr>
<td>MD-ZE125</td>
<td>3</td>
<td>100 a</td>
<td>95 b</td>
<td>100 a</td>
<td>100 a</td>
<td>S</td>
</tr>
</tbody>
</table>

7 Data were the mean of four replicated microplots containing five plants of each of the watermelon genotypes Calhoun Gray and PI-296341-FR (MD). Incidence of Fusarium wilt was rated at 6 weeks after plants were transplanted in *F. oxysporum* f. *sp. niveum*-infested microplots. Control watermelon plants in noninfested microplots remained healthy and are not presented in the table. Means in a column followed by the same letter are not significantly different (P = 0.05) according to Fisher’s protected least significant difference test. Wilt incidence data were arcsine-square root transformed prior to analysis.

8 Ratings of disease reaction on PI-296341-FR (MD) were assigned for the isolates based on mean incidence of Fusarium wilt on the differential genotype over two experiments, where R = resistant (<33% wilt) and S = susceptible (≥33% wilt), following the criteria established by Martyn and Bruton (22).

### DISCUSSION

Three races (0, 1, and 2) of *F. oxysporum* f. *sp. niveum* have been described in watermelon (2,6,8,21,27,28). The results of this study demonstrated that the four isolates from Maryland belong to a fourth race which is more virulent than race 2. The Maryland isolates showed consistent virulence toward PI-296341-FR, a genotype which is considered resistant to race 2, in greenhouse and field microplot experiments. The Maryland isolates also exhibited a higher level of virulence toward PI-296341-FR than a race 2 isolate as measured by stem colonization by *F. oxysporum* f. *sp. niveum*. All four Maryland isolates were pathogenic on watermelon and nonpathogenic on non-Citrullus hosts examined. This indicated that they were host specific and that their valid formae specialis designation is *F. oxysporum* f. *sp. niveum*. The Maryland isolates, which were vegetatively compatible with each other, were not compatible to the reference isolates of races 0, 1, or 2 tested, indicating that the Maryland isolates are a different genetic strain of *F. oxysporum* f. *sp. niveum* than others evaluated here. Based on these results, a fourth race is designated as race 3 of *F. oxysporum* f. *sp. niveum* following the established numbering sys-
stem for races of *F. oxysporum* f. sp. *niveum* (6). The new race 3 was found 37 years after the highly virulent race 2 was first reported by Netzer and Dishon in 1973 (28).

The discovery of race 3 in Maryland is important but not surprising. It may have evolved due to the high selection pressure placed on the pathogen by the intensive cultivation of race-1-resistant watermelon cultivars in the fields where the race 3 isolates were originally collected. The fields had been cropped to watermelon of various cultivars, most of which were race 1 resistant, for many years (up to 22 years), resulting in the highest soil populations of *F. oxysporum* f. sp. *niveum* among watermelon production fields surveyed (37). These fields also contained the highest ratios of pathogenic to total populations of *F. oxysporum* (more than 80% of total isolates) and the highest proportions of race 2 (more than 76% of total isolates) (37). However, the possibility of introduction of race 3 from a foreign source along with infected seed or seedlings or the evolution of race 3 from nonpathogenic *F. oxysporum* cannot be excluded (20,34).

The potential origin of races 0, 1, and 2 of *F. oxysporum* f. sp. *niveum* in the Mid-Atlantic region has been discussed in a previous study (40).

Differences in determining races of *F. oxysporum* f. sp. *niveum* have been recognized for many years. This is partly due to the fact that the results of pathogenicity tests can be greatly affected by many factors such as inoculum concentration, age of inoculated plants, environmental conditions, inoculation method, and source of differential genotypes (1,18,23). The results of these tests also demonstrated the difficulties in race identification in two aspects. First, we observed apparent segregation in the populations of PL-296341-FL (FL) or PL-296341-OK (OK) that were used to differentiate race 3 from race 2 in the first two evaluations of this study. In these two tests, inoculations with a race 2 strain (TX-X1D) resulted in a range of 20 to 32% wilt on either FL or OK populations of PL-296341-FL. This confirmed that the genotypes of PL-296341-FL were not completely fixed (12,24,35) and that the race-2-susceptible portion of the populations may have increased with the propagation of the PI lines. Second, there is a concern about potential reduction in virulence of the reference race 2 strain (TX-X1D). We observed this during the evaluations conducted in 2002 and 2005. Therefore, it was necessary to include multiple known race 2 reference strains and more sources of the key differential PL-296341-FL for greater precision in the separation of race 3 from race 2 (17,20,23).

Because strains of *F. oxysporum* f. sp. *niveum* frequently exhibit a continuum in virulence on the differentials (2,17,21,37), quantitative wilt values used to assign isolates to a specific race were not provided in some studies (1,6,14,17,20). Furthermore, the wilt values used in other studies to classify the races have often been inconsistent (22,23,27,37). To standardize the criteria used to distinguish races, based on the results of our current study and previous studies (9,37), we supported the use of the system proposed by Martyn and Bruton (22) to characterize disease reactions on the differential genotypes for race determination (Table 1). In this system, a mean wilt incidence of ≥33% was rated as susceptible and <33% was rated as resistant. The categories of resistance and susceptibility established in this system also closely followed the significant (P = 0.05) separations of percentage wilt in our study and previous studies (17,37). This system has already been used for race determination in some previous studies (9,37). Appropriate disease values that could be used to differentiate races of *F. oxysporum* f. sp. *niveum* are critical to the studies of biology, epidemiology, and management of Fusarium wilt in watermelon.

The extent of the distribution of the new race in the Mid-Atlantic region has not been investigated but its occurrence in four farms located in two counties in Maryland indicates that it may exist throughout the Mid-Atlantic region. The existence of race 3 represents an additional threat to watermelon production in the region and increases the difficulty of managing *Fusarium* wilt. Race 2, to which no resistance exists in current watermelon cultivars, is already reported in 8 states (3,5,10,17,19,22,36; A. P. Keinath, personal communication) of which are in the top 10 watermelon production states in the United States. This situation is especially problematic in watermelon production areas such as Maryland and Delaware, where a high percentage of watermelon acreage is triploid (seedless) watermelon. Triploid watermelon cultivars are generally more susceptible to *Fusarium* wilt than diploid (seeded) cultivars. Triplid watermelon cultivars are generally more susceptible to *Fusarium* wilt than diploid (seeded) cultivars, and almost all currently grown triploid watermelon cultivars (with a few exceptions such as Seedless Sangria and Revolution) are susceptible to race 1 (41). Long-term crop rotation (more than 5 years) and chemical fumigation (not methyl bromide since 2005) may be the best available options for management of *Fusarium* wilt caused by race 3. Both control methods could significantly reduce the inoculum levels of the pathogen in soil. However, land availability and economic issues will limit implementation of these two tactics. Other approaches, such as grafted transplants (4), soil amendment (39), or integrated use of multiple methods, may be needed for effective management of *Fusarium* wilt caused by this new race in watermelon.

LITERATURE CITED