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RESEARCH ARTICLE

Novel Mutations Detected in Avirulence Genes Overcoming Tomato Cf Resistance Genes in Isolates of a Japanese Population of Cladosporium fulvum

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Abstract

Leaf mold of tomato is caused by the biotrophic fungus Cladosporium fulvum which complies with the gene-for-gene system. The disease was first reported in Japan in the 1920s and has since been frequently observed. Initially only race 0 isolates were reported, but since the consecutive introduction of resistance genes Cf-2, Cf-4, Cf-5 and Cf-9 new races have evolved. Here we first determined the virulence spectrum of 133 C. fulvum isolates collected from 22 prefectures in Japan, and subsequently sequenced the avirulence (Avr) genes Avr2, Avr4, Avr4E, Avr5 and Avr9 to determine the molecular basis of overcoming Cf genes. Twelve races of C. fulvum with a different virulence spectrum were identified, of which races 9, 2.9, 4.9, 4.5.9 and 4.9.11 occur only in Japan. The Avr genes in many of these races contain unique mutations not observed in races identified elsewhere in the world including (i) frameshift mutations and (ii) transposon insertions in Avr2, (iii) point mutations in Avr4 and Avr4E, and (iv) deletions of Avr4E, Avr5 and Avr9. New races have developed by selection pressure imposed by consecutive introductions of Cf-2, Cf-4, Cf-5 and Cf-9 genes in commercially grown tomato cultivars. Our study shows that molecular variations to adapt to different Cf genes in an isolated C. fulvum population in Japan are novel but overall follow similar patterns as those observed in populations from other parts of the world. Implications for breeding of more durable C. fulvum resistant varieties are discussed.
Introduction

*Cladosporium fulvum* Cooke [syn. *Passalora fulva* (Cooke) U. Braun & Crous] is a biotrophic pathogen that causes leaf mold of tomato [1]. The fungus has been reported on tomato since the late 1800s [2]. The disease is primarily a problem in greenhouse-grown tomatoes and occurs worldwide in areas with high humidity and moderate temperatures. Infection begins with conidia germinating on the lower leaf surface and producing runner hyphae that enter the host through stomata. Subsequently, the fungus colonizes the intercellular space between mesophyll cells, and 10–14 days after penetration, conidiophores emerge from stomata producing large number of conidia that can re-infect tomato leaves [1, 3–5].

The *C. fulvum*-tomato interaction follows the gene-for-gene system indicating that each dominant pathogen avirulence (*Avr*) gene product is recognized by the product of a corresponding dominant host *Cf* resistance gene directly or indirectly [6]. To date, five *Avr* genes (*Avr2, Avr4, Avr4E, Avr5, and Avr9*) have been cloned and characterized from *C. fulvum* [7–13], and their encoded proteins trigger a hypersensitive response (HR) in host plants carrying the corresponding *Cf*-2, *Cf*-4, *Cf*-4E, *Cf*-5, and *Cf*-9 genes, respectively [14–18]. The different *Cf*-genes encoding leucine-rich receptor-like proteins, originate from wild *Solanum* species, and have been introduced into tomato cultivars currently grown worldwide [19]. However, by selection pressure imposed by *Cf* genes, new *C. fulvum* races evolved that overcome introduced *Cf* resistance genes. DNA modifications observed in *Avr* genes of new races resulted in frame-shift mutations or point mutations leading to amino acid substitutions in the encoded *Avr* proteins, whereas complete loss of an *Avr* gene and transposon insertions in *Avr* genes were also observed [11, 20–23].

In Japan, the first reports on the occurrence of tomato leaf mold date from the 1920s [24]. Initially, race 0 isolates carrying all known *Avr* genes were the only indigenous isolates identified and there were no reports of pathogen specialization until the 1960s [24]. In the USA, Canada and Western Europe, breeding of tomato cultivars resistant against leaf mold started in the 1930s [25, 26] and in Japan in the 1960s. In Japan, the *Cf*-2 gene was the first resistance gene to be introduced in commercial tomato lines in 1965 [27]. As a consequence race 2 isolates were identified in the late 1970s [28]. Soon after the introduction of the *Cf*-4 gene (and likely also the *Cf*-11 gene) in the 1990s, races 2.4 and 2.4.11 isolates appeared [29] followed by race 4 and 4.11 isolates in 2003 [30]. In the 2000s, new resistant cultivars carrying the *Cf*-9 gene were launched, and in 2008, the races 4.9, 4.9.11 and 2.9 isolates that overcome *Cf*-9-mediated resistance were identified [31, 32]. It is not known when the *Cf*-5 gene was introduced, but recently, in Japan, new races (2.5.9 and 4.5.9) were identified on a cultivar carrying the *Cf*-5 and *Cf*-9 genes, though the presence of these genes in this cultivar was then unknown [33]. Of the eleven races presently identified in Japan (races 0, 2, 2.4, 2.4.11, 2.9, 2.5.9, 4, 4.9, 4.5.9, 4.11, 4.9.11, 7) also occur elsewhere in the world [34], whereas races 2.9, 4.9, 4.5.9 and 4.9.11 are unique to Japan [31–33].

For many years *C. fulvum* was not a serious economic problem for tomato growers in Japan, but the recent appearance of new races [31–33] prompted us to perform a detailed study of the virulence spectrum of the fungal population in the whole country and to analyze the molecular basis of adaptation to the introduced *Cf* genes. Of all eleven races of *C. fulvum* in Japan nothing is known about DNA modifications in *Avr* genes that cause adaptation to the corresponding *Cf* resistance genes. To understand the molecular basis of adaptation of *C. fulvum* to the introduced *Cf* genes in Japan, we determined both the virulence spectrum and the DNA modifications present in the *Avr* genes of 133 isolates of a *C. fulvum* population collected between 1997 and 2013. Most of the new races appeared to be confined to particular regions of the country, whereas older ones have spread over the whole country. We identified many new DNA
modifications in Avr genes leading to virulence on plants with corresponding Cf genes that are unique to the Japanese C. fulvum population. We determined the effect of new DNA modifications on HR-inducing activities of Avr2, Avr4 and Avr5 genes present in Japanese races of C. fulvum on tomato plants carrying the corresponding Cf resistance gene. For some of the new races, the parental isolate(s) of new races could be inferred based on shared sequences in Avr genes.

Materials and Methods
Ethics Statement
Between 1997 and 2013, diseased tomato leaves, from which the C. fulvum were isolated, were collected from 22 of Japan’s 47 prefectures (Fig 1A). Sampling of tomato leaves was performed in private greenhouses under the permission by all owners. No specific permissions were

Fig 1. Distribution of tomato Cf resistance genes and Cladosporium fulvum races in Japan. Prefectures from where Cladosporium fulvum isolates were collected (A), the distribution of Cf genes employed in the different prefectures (B), and the virulence spectrum of the isolates collected in these prefectures (C) are presented. The surveys were conducted between 1997 and 2013 in prefectures highlighted in grey. The prefecture numbers correspond to those shown in S2 Table.

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required for the all locations. The surveys did not involve regulated, endangered, or protected species.

**Fungal isolates and virulence assays**

Collected leaves showed the typical leaf mold symptoms: pale-green or yellow spots on the upper side, and multiple velvet, olive-brown conidia on the lower side of the leaves. 133 single-spores isolated from these lesions were cultured on potato dextrose agar (PDA) for two weeks at 20°C. Isolates were collected from greenhouses at different locations in these prefectures. There is a slight over-representation of isolates collected from Gunma and Mie prefectures, where recent outbreaks of leaf mold were reported. When available, information on the Cf genes present in the tomato cultivars from which the isolates were collected was recorded. The virulence spectrum of the isolates was determined by inoculating them on a differential set of tomato cultivars including ‘Potentate’ (no Cf resistance gene), ‘Vetomold’ (Cf-2), ‘Purdue 135’ (Cf-4), ‘Moneymaker-Cf-5’ (Cf-5), ‘Ontario 7818’ (Cf-6), ‘Moneymaker-Cf-9’ (Cf-9), and ‘Ontario 7716’ (Cf-4 and Cf-11). Three four-week-old plants of each differential cultivar were spray-inoculated on the lower side of the leaves with a conidial suspension of 10⁴ spores/ml of each isolate. The inoculated plants were incubated in a moist chamber at 100% humidity and 25°C with a 16 h light/8 h dark photoperiod. After two to three weeks, the inoculated plants were analyzed and scored visually as either resistant or susceptible. Susceptible cultivars showed heavy sporulation, whereas resistant cultivars were immune and free of disease symptoms. A representative set of 120 isolates was submitted to the microorganism component of the GeneBank resources maintained by the National Institute of Agrobiological Sciences (http://www.gene.affrc.go.jp/index_en.php) under accession numbers MAFF 242495 to 242550, MAFF 242556 to 242575, and MAFF731146 to 731149.

**DNA manipulation and sequencing**

DNA was isolated from 133 isolates of *C. fulvum* for analysis of DNA modifications in Avr genes. In addition, the DNA sequence of Avr genes in four Japanese reference isolates CF5, CF9, CF44 and CF56 collected in Japan in 1973 was also determined [29, 35]. Mycelium from the isolates grown on PDA was collected, transferred to Eppendorf tubes, freeze-dried overnight and disrupted in liquid nitrogen using the Retsch Qiagen Tissue Lyser twice for 30 s with 30 oscillations per minute. DNA was extracted using the DNeasy Plant Mini Kit (Qiagen, USA) according to the manufacturer’s instructions. The DNA concentration was determined using a Nanodrop ND-1000 spectrophotometer (Thermo Scientific, USA).

The mating type in the collection of 133 isolates was examined by PCR amplification of partial genes (MAT1-1 or MAT1-2) using the primers shown in S1 Table. PCR reactions were performed in 25μl volumes containing 50 ng genomic DNA, 1× GoTaq PCR buffer, 0.2 mM dNTPs, 0.4 μM of each primer and 1 U of GoTaq DNA polymerase (Promega, USA). The PCR conditions were 95°C for 2 min, followed by 35 cycles of 95°C for 30 s, 60°C for 30 s and 72°C for 1 min, and a final extension at 72°C for 5 min. The PCR products obtained were separated by electrophoresis and visualized under UV-light.

PCR reactions for amplification of the Avr2, Avr4, Avr4E, Avr5 and Avr9 genes were performed with the primers shown in S1 Table. PCR reactions were performed in 25μl volumes containing 50 ng genomic DNA, 1× GoTaq PCR buffer, 0.2 mM dNTPs, 0.4 μM of each primer and 1 U of GoTaq DNA polymerase (Promega, USA). The PCR conditions were 95°C for 2 min, followed by 35 cycles of 95°C for 30 s, 60°C for 30 s and 72°C for 1 min, and a final extension at 72°C for 7 min. The PCR products were purified using Illustra GFX PCR DNA and Gel Band Purification Kit (GE Healthcare) according to the manufacturer’s instructions. After
DNA purification the DNA concentration was measured using a Nanodrop spectrophotometer. Purified PCR products were sequenced by Macrogen Inc. (Seoul, South-Korea). DNA sequences of the Avr genes were analyzed using the Lasergene package (DNASTAR, USA) and compared with the sequences present in four Japanese reference isolates collected in 1973 as well as those present in a worldwide collection of C. fulvum isolates [21]. The DNA sequence of the Avr genes in the four reference isolates were considered as wild-type.

PVX-mediated expression of Avr genes in tomato plants with corresponding Cf genes

In order to confirm that the DNA modifications observed in the collected isolates were the cause of overcoming a particular Cf-gene, we investigated the HR-inducing activities of wild-type and unique mutant versions of Avr2, Avr4 and Avr5 genes present in Japanese races of C. fulvum on tomato plants carrying the matching Cf resistance gene. To this end, we expressed them in tomato plants carrying corresponding Cf genes by agroinfection using pSfinx, a modified binary Potato Virus X (PVX)-based vector for transient expression of foreign genes into plants as described by Stergiopoulos et al. [36]. This system is based on Agrobacterium tumefaciens-mediated delivery of the recombinant PVX virus that enables targeted systemic production of Avr proteins into the apoplast of PVX-infected plants. The wild-type and mutant versions of Avr2, Avr4 and Avr5 genes were cloned into the PVX vector using primers presented in S1 Table. Ten-day-old tomato seedlings were agroinfected and analyzed by a method described by van der Hoorn et al. [37]. Photographs were taken at 20 days post inoculation. HR-inducing activities of Avr4E and Avr9 were not assayed as the observed mutations leading to virulence observed in these genes involved loss of the complete gene or a point mutation that was already reported to overcome a particular Cf gene (Cf-4E and Cf-9, respectively). Thus, only Avr genes with unique new mutations were tested for loss of their HR-inducing activity.

Results

The virulence spectrum of isolates of a Cladosporium fulvum population collected in Japan

Morphological characteristics of single-spore isolates and disease symptoms caused by them after inoculation onto susceptible tomato plants appeared similar to those described previously for C. fulvum [38]. In total 133 C. fulvum single-spore isolates were collected from diseased tomato plants grown in Japan from north to south (Fig 1A). For each isolate, its acronym, year of sampling, prefecture, and the Cf resistance gene present in the cultivar from which the isolate was collected, were recorded (S2 Table). In addition, the mating type (MAT1-1 or MAT1-2) of all isolates was determined because both are still present in worldwide populations although C. fulvum is supposed to be an asexual fungus [21, 39]. Both mating types are also present in Japan, with a slight bias for MAT1-2 (73%; S2 Table).

The virulence spectrum of the 133 C. fulvum isolates is shown in Table 1. In total twelve races with a novel virulence spectrum were identified: races 0, 2, 4, 9, 2.4, 2.9, 2.5.9, 4.9, 4.5.9, 4.11, 2.4.11, and 4.9.11. It is the first time a race 9 is reported in Japan (prefecture Gunma) and elsewhere in the world. The frequencies of the different races in the different prefectures vary obviously, which most likely reflects the different frequencies of the Cf-4, Cf-5 and Cf-9 genes employed in these prefectures (Fig 1B and 1C). Indeed, all isolates that overcome Cf-5 and Cf-9 resistance were isolated from Cf-5 and Cf-9 plants, respectively, introduced in prefectures Iwate, Fukushima, Tochigi, Gunma, Chiba and Saga (Fig 1B). Races 2.5.9 and 4.5.9 that
overcome Cf-5 are confined to one prefecture only (Fig 1C) reflecting the locations where they were identified for the first time. So far, races 2.5.9 and 4.5.9 have not migrated to other prefectures. Although the Cf-2 gene is no longer used in Japan, many race 2 isolates are still present throughout the country (Fig 1C), suggesting that these races are not outcompeted yet. From Cf-0 plants not only race 0 isolates but often also additional races overcoming different Cf genes were isolated, including races 2, 4, 4.11 and 4.9.11 (Fig 2).

Although Cf-2, Cf-4, Cf-5 and Cf-9 genes have all been employed in Japan, no isolates able to overcome all four Cf genes were identified (Table 1). Also no isolate overcoming the Cf-6 gene was identified, which is consistent with the absence of this resistance gene in cultivars used in Japan.

DNA modifications in Avr genes

The nucleotide sequences of the Avr genes of four race 0 isolates that served as a reference were considered to contain wild-type Avr genes [21]. The ability to overcome a particular Cf gene is likely attributed to non-synonymous DNA modification(s) in the ORF of a corresponding Avr gene or loss of the corresponding Avr gene. The Avr2 gene amplified from races 2, 2.4, 2.9, 2.5.9 and 2.4.11 showed mutations that allowed them to overcome the Cf-2-mediated resistance (S2 Table). These races contain in total five different mutations in the Avr2 ORF when compared with the wild-type reference Avr2 gene including (i) a mutation destroying the start codon (c.1A>G; 21 isolates), (ii) a nucleotide insertion and substitution (c.50insT; c.52A>C; 4 isolates) leading to a truncated Avr2 protein, (iii) a nucleotide change (c.242G>T; 2 isolates) leading to Cys63Phe amino acid substitution in the C-terminus of the Avr2 protein, (iv) an insertion of five As (c. (64_69) insA; 1 isolate) leading to a frame shift in the Avr2 protein, and (v) a multiple nucleotide deletion (c.56delCAGCAGCCAA; 1 isolate) also leading to a frame shift in the Avr2 protein. Finally, a transposon insertion was observed in three isolates, leading to production of a nonfunctional Avr2 protein (S2 Table). Several of the observed mutations in Avr2 are new, and have not been observed in C. fulvum populations elsewhere in the world (Table 2).

The Avr4 gene amplified from races 4, 2.4, 4.9, 4.5.9, 4.11, 2.4.11 and 4.9.11 contained in total five different mutations in the Avr4 ORF that allowed them to overcome the Cf-4-
mediated resistance (S2 Table). They include (i) c.118T>C (1 isolate), (ii) c.191G>T (48 times), (iii) c.191G>C (34 isolates), (iv) c.191G>A (2 isolates), mutations all causing the substitution of a cysteine residue by an arginine, serine, phenylalanine and tyrosine residue, respectively. The mutation c.318delG causes a frame shift in the Avr4 protein (p. ser107ValfsX4; 10 isolates). Several of the observed mutations in Avr4 gene are new and have not been observed in C. fulvum populations elsewhere in the world (Table 2).

The set of Cf differentials used to identify the virulence spectrum of the collection did not include a cultivar with the Cf-4E gene alone. Cf-4E is a paralog of the Cf-4 gene and is located close to Cf-4 gene, while effector genes including Avr4 and Avr4E are not linked in the C. fulvum genome [14–17, 41]. In 80 out of 133 isolates two mutations (c.244T>C; c.278T>C) were observed in the Avr4E ORF causing two amino acid substitutions in the Avr4E protein (p.Leu82Phe and p.Thr93Met) (S2 Table) enabling them to overcome Cf4-E-mediated resistance [12]. Remarkably, 22 isolates that can overcome Cf4-E-mediated resistance in the Japanese C. fulvum population lacked the Avr4E gene. Both mutations in the Avr4E gene have been observed in C. fulvum populations elsewhere in the world [21].

Only six out of 133 isolates in the Japanese C. fulvum collection overcome Cf-5-mediated resistance (races 2.5.9 and 4.5.9) (Table 1). In two isolates of race 4.5.9, the Avr5 gene contained a new mutation (c.268G>C) that had not been observed in C. fulvum populations elsewhere in the world (S2 Table). This new mutation involves the substitution of a glycine residue by a
arginine in the C-terminus of the Avr5 protein (Table 2). The remaining four isolates of races 2.5.9 and 4.5.9 that did overcome Cf-5-mediated resistance had lost the Avr5 gene (S2 Table).

Overcoming Cf-5-mediated resistance by deletion of the Avr5 gene has been reported before in C. fulvum [13]. Similarly, the Avr9 gene in 36 isolates that overcome Cf-9-mediated resistance (races 9, 2.9, 2.5.9, 4.9, 4.5.9 and 4.9.11) was absent from the genome (S2 Table). In addition, in 25 isolates not able to overcome the Cf-9 gene, the mutation c.23T>C in the Avr9 ORF was observed leading to a p.Val8Ala amino acid substitution in the signal peptide of the Avr9 protein which has,

| Avr gene | Allele | Frequency | Effect on protein | Remarks | Loss of Cf-mediated HR | Genotypes with independent events |
|----------|--------|-----------|-------------------|---------|------------------------|------------------------------------|
| Avr2     | Wild-type | 101       | No (Fig 3)        |         |                        |                                    |
| c.1A>G   | 21     | p.Met1Val | Disruption of start codon | Yes | 2 (G30, G37)          |                                    |
| c.50insT | (4)    | p.Ile18ThrfsX24 | Frameshift | Yes | 2 (G03, G35)          |                                    |
| c.52A>C  | (4)    | p.Ile18Leu | Mutation in signal sequence | No  |                       |                                    |
| c.56delCAGCAGCCAA | 1     | p.Ala19GlufsX38 | Frameshift | Yes | 1 (G31)               |                                    |
| c.64-69insA | 1   | p.Leu24TyrfsX24 | Frameshift | Yes | 1 (G11)               |                                    |
| c.242G>T | 2      | p.Cys63Phe | Disruption of S-bridge | Yes (Fig 3) | 1 (G08)           |                                    |
| Transpon insertion | 3    | no protein | Yes (Fig 3) | Yes | 1 (G01)               |                                    |
| Avr4     | Wild-type | 38       | No (Fig 3)        |         |                        |                                    |
| c.118T>C | 1      | p.Cys40Arg | Disruption of S-bridge | Yes (Fig 3) | 1 (G04)           |                                    |
| c.191G>T | 48     | p.Cys64Phe | Disruption of S-bridge | Yes (Fig 3) | 4 (G19, G20, G26, G32) |                                    |
| c.191G>C | 34     | p.Cys64Ser | Disruption of S-bridge | Yes (Fig 3) | 1 (G38)           |                                    |
| c.191G>A | 2      | p.Cys64Tyr | Disruption of S-bridge | Yes (Fig 3) | 1 (G08)           |                                    |
| c.318delG | 10   | p.Ser107ValfsX4 | Frameshift | Yes (Fig 3) | 2 (G03, G13) |                                    |
| Avr4E    | Wild-type | 31       | Yes               |         |                        |                                    |
| c.244T>C; | (80)  | p.Phe82Leu | Yes               |         |                        |                                    |
| c.278T>C | (80)  | p.Met93Tyr | Yes               |         |                        |                                    |
| Gene deletion | 22  | no protein | Yes | 1 (G27)               |                                    |
| Avr5     | Wild-type | 127      | No (Fig 3)        |         |                        |                                    |
| Gene deletion | 4   | no protein | Yes | 2 (G16, G18) |                                    |
| c.268G>C | 2      | p.Gly90Arg | Yes (Fig 3) | Yes | 1 (G15)               |                                    |
| Avr9     | Wild-type | 72       | No (Fig 3)        |         |                        |                                    |
| Gene deletion | 36  | no protein | Yes (Fig 3) | Yes | 5 (G06, G17, G24, G26, G40) |                                    |
| C.23T>C  | 25     | p.Val8Ala | Mutation in signal sequence | No  |                       |                                    |

Table 2. Overview of all DNA modifications present in the coding sequence of five avirulence genes in a population of 133 Cladosporium fulvum isolates collected in Japan.

aCodes for mutations at DNA level are according to den Dunnen and Antonarakis [40]. Mutations specific to isolates of the Japanese population of C. fulvum are highlighted in bold.

b The numbers refer to the number of isolates in the population carrying the allele. The DNA modifications in Avr2 (c.50insT and c.52A>C) and Avr4E (c.244T>C and c.278T>C) that always appeared together were showed in parentheses.

c Codes for mutations at protein level are according to den Dunnen and Antonarakis [40]. Mutations specific to isolates of the Japanese population of C. fulvum are highlighted in bold.

d The Avr alleles were expressed in the PVX expression system and analyzed in tomato plants carrying the corresponding Cf resistance gene. Representative pictures of the hypersensitive response (HR)-inducing activity of wild-type and mutant alleles are shown in Fig 3.

e Single independent mutation, transposon insertion or deletion events deduced from related genotypes.

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arginine in the C-terminus of the Avr5 protein (Table 2). The remaining four isolates of races 2.5.9 and 4.5.9 that did overcome Cf-5-mediated resistance had lost the Avr5 gene (S2 Table). Overcoming Cf-5-mediated resistance by deletion of the Avr5 gene has been reported before in C. fulvum [13].

Similarly, the Avr9 gene in 36 isolates that overcome Cf-9-mediated resistance (races 9, 2.9, 2.5.9, 4.9, 4.5.9 and 4.9.11) was absent from the genome (S2 Table). In addition, in 25 isolates not able to overcome the Cf-9 gene, the mutation c.23T>C in the Avr9 ORF was observed leading to a p.Val8Ala amino acid substitution in the signal peptide of the Avr9 protein which has,
however, no effect on its HR-inducing activity on Cf-9 plants [21]. Overcoming the Cf-9 resistance due to loss of the Avr9 gene in C. fulvum has been observed before in populations of this fungus elsewhere in the world [21]. Overall, this molecular analysis revealed diverse types of mutations in Avr genes that reflect adaptation of C. fulvum to specific Cf genes.

HR-inducing activity of novel mutant Avr proteins produced by new Japanese races overcoming Cf-mediated resistance

All Avr2, Avr4 and Avr5 genes in the Japanese collection carrying novel DNA modifications in their ORFs were assayed for their HR-inducing activity on tomato plants carrying the corresponding Cf gene using the PVX expression system. Avr alleles carrying DNA modifications that were reported before [36] to have lost HR-inducing activity due to absence of an Avr gene (Avr4E and Avr9) or coding for a strongly truncated Avr protein (Avr2) were not included in these assays. Avr genes with novel DNA modifications were cloned in the PVX vector and assayed for HR-inducing activity on tomato plants carrying the corresponding Cf genes [36]. Without an exception, all Avr genes with novel and unique DNA modification in their Avr coding sequence had lost HR-inducing activity on tomato cultivars carrying the corresponding Cf gene (Fig 3).

Loss of Cf-2-mediated HR is usually caused by frame shifts observed in the Avr2 gene [21], but here for the first time we observed an amino acid substitution in the Avr2 protein (p. Cys63Phe) that caused loss of HR-inducing activity on tomato plants carrying the Cf-2 gene (Fig 3A).

Most of the novel mutations found in the Avr4 gene led to cysteine substitutions in the Avr4 protein causing loss of HR-inducing activity on Cf-4 plants (Fig 3B). The Cys64Tyr substitution in Avr4 reported before [42] showed a weaker effect on destroying HR-inducing activity on Cf-4 plants than the novel Cys64Phe and Cys64Ser substitutions reported here. Also the new frame shift mutation (p. Ser107ValfsX4) caused loss of HR-inducing activity (Fig 3B).

Only the mutation (p. Pro4LeufsX18) in the signal peptide of the Avr5 gene causing a frame shift in the Avr5 protein was reported previously [13]. The amino acid substitution (p. Gly90Arg) in the Avr5 protein that caused loss of HR-inducing activity on tomato plants carrying the Cf-5 gene had not been reported before (Fig 3C).

Relationship between different Avr genotypes in C. fulvum isolates collected in Japan

In order to identify relationships between the C. fulvum isolates collected in Japan, we classified the different Avr genotypes based on the nucleotide sequence of Avr genes and presence of either the MAT1-1 or MAT1-2 mating type locus. In total 41 unique genotypes could be identified (G01 to G41; S2 Table). We subsequently tried to infer parental relationships between existing races and recently identified races 2.9, 4.9, 4.9.11 and 2.5.9 collected in same prefecture.

The race 2.9 isolates were first reported to occur in the Iwate prefecture [32] and most likely developed from a race 2 isolate present in Iwate. Race 2 isolate H-48/G07 from Iwate is most likely the parents of all race 2.9 isolates (G06) identified in Iwate (Table 3). Race 2 isolates occurring in other prefectures (Hokkaido, Akita, Fukushima, Gunma, Gifu, Mie, Shimane, Fukuoka, Nagasaki and Miyazaki) are excluded as the potential parents of race 2.9 isolates because they carry a different mating type locus or contain different mutations in ORFs, or low probability due to geographical distance (S2 Table; Fig 1). The other race 2.9 isolates were collected
in prefectures Tochigi and Saga, but never a race 2 isolate has been collected in these prefectures (S2 Table).

Race 4.9 isolates were first reported to occur in Gunma prefecture and race 4.9.11 isolates in prefectures Gunma, Chiba and Fukushima [31] and this study discovered race 4.9 isolates in Tochigi (S2 Table). No additional race 4.9 and 4.9.11 isolates were identified outside these prefectures. Thus, race 4.9 and 4.9.11 isolates have likely developed from parental race 4 and race 4.11 isolates, respectively. By comparing the genotypes of the new race 4.9 and race 4.9.11 isolates with potential parental race 4 and race 4.11 isolates, they all appear to carry MAT1-1 and all contain the c.158+26_158+28insTGA mutation in the \textit{Avr2} gene, c.36T>C mutation in the

Fig 3. \textit{Cf}-mediated hypersensitive responses (HR) triggered by wild-type and mutant Avr proteins. \textit{Avr}-wild-type and mutant genes were cloned in the PVX vector and the recombinant PVX virus was assayed for HR-inducing activity on \textit{Cf}-2, \textit{Cf}-4 or \textit{Cf}-5 tomato plants. (A) PVX-mediated expression of wild-type \textit{Avr2} protein causes strong HR-inducing activity and eventually kills \textit{Cf}-2 tomato plants, but mutant \textit{Avr2} protein (p. Cys63Phe substitution) present in isolate CF212 lost HR-inducing activity. (B) PVX-mediated expression of wild-type \textit{Avr4} protein causes strong HR-inducing activity and eventually kills \textit{Cf}-4 tomato plants. Five different mutant \textit{Avr4} proteins (substituted amino acid residues are indicated) all lost HR-inducing activity on \textit{Cf}-4 tomato plants. (C) PVX-mediated expression of wild-type \textit{Avr5} protein elicits a strong HR on \textit{Cf}-5 tomato plants, whereas mutant \textit{Avr5} protein (p.Gly90Arg substitution) lost HR-inducing activity. Plants were photographed at 3 weeks post inoculation.

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Table 3. Potential parental isolates of new *Cladosporium fulvum* races collected in same prefecture.

| Prefecture | Isolate  | Genotype \(^a\) | MAT\(^b\) | Race\(^c\) | Avr2\(^d\) | Avr4\(^d\) | Avr4E\(^d\) | Avr5\(^d\) | Avr9\(^d\) |
|------------|----------|-----------------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| (03) Iwate  | H-41     | G06 MAT1-2      | 2.9       | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | H-48\(^g\) | G07 MAT1-2      | 2         | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type |  |
| (09) Tochigi | Ohtawara1 | G14 MAT1-1      | 4.9       | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | Ohtawara2 | G14 MAT1-1      | 4.9       | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | Kaminokawa\(^e\) | G13 MAT1-1 | 4       | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
| (10) Gunma  | CF308    | G12 MAT1-1      | 4.9.11    | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | CF309    | G12 MAT1-1      | 4.9.11    | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | CF307\(^a\) | G20 MAT1-1      | 4.11      | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type |  |
| (09) Tochigi | Utsunomiya1 | G18 MAT1-1 | 2.5.9    | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | Utsunomiya2 | G18 MAT1-1 | 2.5.9    | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | Tochigi1\(^e\) | G17 MAT1-1 | 2.9     | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |
|            | Tochigi2\(^e\) | G17 MAT1-1 | 2.9     | c.1A>G    | c.-57G>A  | c*36T>C   | wild-type | gene deletion |

\(^a\) Genotype of isolates based on sequence of *Avr* genes and mating type loci.

\(^b\) Mating type of isolate; MAT1-1 or MAT1-2.

\(^c\) Virulence spectrum of isolate.

\(^d\) DNA modifications observed in *Avr* genes; Codes for mutations at DNA level are according to den Dunnen and Antonarakis (2000) [40]. Mutations different to potential parental isolate are highlighted in bold.

\(^e\) Potential parental isolate(s)
Avr4E gene and wild-type Avr5 gene. Race 4.9 stains (G14) from Tochigi were most likely derived from a race 4 isolate Kaminokawa/G13 identified from Tochigi (Table 3). For the new race 4.9.11 isolates collected in Gunma, the parent is most likely race 4.11 isolate CF307/G20 collected in same prefecture Gunma (Table 3).

Isolates that can overcome Cf-5-mediated resistance were recently collected and races 2.5.9 and 4.5.9 were identified only in Tochigi prefecture [33]. Thus, the new race 2.5.9 and 4.5.9 isolates have likely developed from regional parental race 2.9 and 4.9 isolates, respectively. Race 2.9 isolates (G17) from Tochigi are most likely the parents of race 2.5.9 stains (G18) identified in Tochigi based on mating type locus and nucleotide sequences of Avr genes (Table 3). Potential parental isolate of races 9 and 4.5.9 isolated from same prefecture could not be assigned.

Discussion

Strong selection pressure on C. fulvum population by tomato Cf resistance genes

C. fulvum is present in Japan since the 1920s, but introduction of most Cf resistance genes in tomato plants started later than elsewhere in the world [21, 34]. The Cf-2 gene has been introduced in tomato grown in Japan in the 1960s [27], the Cf-4 and Cf-9 genes in the last two decades, and the Cf-5 gene recently. Since the introduction of the latter Cf genes, no nation-wide survey has been performed in Japan; little is known about changes in the virulence spectrum of the C. fulvum population in Japan, whereas nothing is known about DNA modifications in Avr genes of isolates overcoming introduced Cf genes.

A total of 133 isolates collected between 1997 and 2013 belong to 12 different races (Table 1). Only four races (0, 2, 2.4, and 2.4.11) were reported to occur in Japan before 1998 [29]. Introductions of the Cf-4 gene in the late 1990s and the Cf-9 gene in the 2000s and very recently the Cf-5 gene has led to the appearance of eight new races that can overcome the Cf-2, Cf-4, Cf-5 or Cf-9 gene (races 4, 2.9, 2.5.9, 4.9, 4.5.9, 4.11 and 4.9.11 [30–33]; race 9 identified in this report). Of these twelve races, seven have a virulence spectrum also reported to occur elsewhere in the world [21, 34], whereas five have a virulence spectrum that is unique to Japan (races 9, 2.9, 4.9, 4.5.9 and 4.9.11). Most of the DNA modifications observed in the Avr genes of the races that can overcome Cf-2-, Cf-4-, Cf-5- or Cf-9-mediated resistance are unique to the Japanese population of C. fulvum (Table 2). Strikingly, mutations in Avr2 and Avr4 genes are very diverse (6 and 5 different types, respectively), suggesting that new races may have arisen independently from different parental isolates. Similarly, Avr9 deletion in races overcoming Cf-9 gene occurred at least five times independently, which is at much higher frequency than deletion of Avr4E and Avr5, or transposon insertion in Avr2 that occurred only once in the studied population (Table 2). The frequency of Avr gene loss is likely correlated with the presence of repeats flanking these Avr genes causing their instability [42].

Introduction of single Cf genes not only contributed to the appearance of new races, but also decreased the genetic diversity of the fungal population. The fact that race 4.11 isolates rather than race 4 isolates seem to have been selected recently by introduction of Cf-4 cultivars in central Japan suggests that growers also used cultivars that contained the Cf-11 gene (Fig 1). Surprisingly, although the race 2 isolates are still collected frequently, very few race 2.4 and 2.4.11 isolates were collected, suggesting that simultaneous adaptation to both the Cf-2 and Cf-4 might cause a fitness penalty.
New races of *C. fulvum* adapted to *Cf-2, Cf-4, Cf-5* and *Cf-9* genes carry unique mutations in the corresponding *Avr* genes

The majority (71%) of the collected isolates in the population can overcome the *Cf-4* gene, 50% the *Cf-11* gene, 27% the *Cf-9* gene, 24% the *Cf-2* gene and 4.5% the *Cf-5* gene. From *Cf-0* tomato cultivars, not only race 0 isolates but often also isolates adapted to different *Cf* genes were collected (Fig 2). This is surprising as *Avr* genes are supposed to encode virulence factors, and races with a complex virulence spectrum are supposed to be less viable on *Cf-0* plants than race 0 isolates and would be outcompeted by the latter in time [23, 43–45]. DNA modification in *Avr* genes leading to virulence would be beneficial for an isolate only when growing on cultivars carrying the corresponding *Cf* gene, except for modifications that avoid recognition by the corresponding *Cf* proteins without affecting their virulence function. The latter is true for isolates that overcome the *Cf-4* gene as they produce mutated versions of *Avr4* proteins that are no longer recognized by the *Cf-4* protein but can still bind to chitin and protect the fungus against the deleterious effects of plant chitinases [46–49]. Most of DNA modifications identified in the *Avr4* gene of Japanese isolates adapted to *Cf-4* plants are unique, lead to production of an unstable *Avr4* effector that still binds to chitin. This mechanism of avoiding *Cf-4* recognition seems to be under strong selection because the same position (c. 191G) in the *Avr4* gene showed three different nucleotide substitutions that occurred at least six times (Table 2). Another unique DNA modification observed in the *Avr4* gene (c.318delG) causes a frame shift at the 3’ of the gene leading to a mutant *Avr4* protein that is 25 amino acid residues shorter than the wild-type protein and is no longer recognized by the *Cf-4* protein (Table 2; Fig 3). Frame shift mutations in *Avr4* leading to a truncated *Avr4* protein have reported only once before [41], but it is not known whether these truncated versions of *Avr4* are still able to bind chitin. Dissociation between chitin binding and recognition by the *Cf-4* protein might explain why such a high frequency of isolates that can overcome the *Cf-4* gene occurs. These isolates are likely not quickly outcompeted by race 0 isolates on *Cf-0* tomato plants.

In race 2 isolates that can overcome *Cf-2*-mediated resistance, the mechanism might be different as they no longer produce a functional *Avr2* protein that can inhibit plant cysteine proteases (the supposedly intrinsic virulence function of *Avr2*) like Rcr3, Pip2, TD65 and aleurain present in the apoplast of tomato [50, 51]. The c.A>G mutation leading to disruption of the start codon, and the c.242G>T mutation destroying a disulfide bond at the C-terminus of the *Avr2* protein are most conspicuous. Interestingly, the latter mutation has been introduced artificially before in *Avr2*, and the shorter but stable *Avr2* protein lacking the C-terminal disulfide bond produced by this mutant was hundred fold less active than the wild-type *Avr2* protein in inhibiting Rcr3 cysteine protease and triggering *Cf-2*-mediated HR [50]. It is surprising that none of the cultivars from which isolates were collected contained the *Cf-2* gene, whereas still 24% of the collected isolates can still overcome the *Cf-2* resistance gene (Table 1). As the intrinsic function of the *Avr2* gene is lost in race 2 isolates, it is expected that on *Cf-0* plants race 2 isolates will gradually be outcompeted by race 0 isolates as they are supposed to be less viable.

Interestingly, all Japanese isolates that can overcome the *Cf-9* resistance gene lacked the entire *Avr9* gene which was also reported for this type of isolates collected elsewhere in the world [9, 21]. The genome of *C. fulvum* has recently been sequenced and the *Avr9* gene was shown to be flanked by repeats, which might explain *Avr9* instability leading to its loss [42]. Loss of the *Avr9* gene seems to occur even at a higher frequency than the accumulation of DNA modification in the *Avr9* gene as all *Avr9* genes present in isolates that could not overcome the *Cf-9* gene appeared identical. Although the biological function of the *Avr9* protein is not known, it is expected that races lacking the *Avr9* gene would also be outcompeted in time by race 0 isolates on *Cf-0* plants since the *Avr9* gene is supposed to encode virulence factor as well.
Similarly, deletion of the *Avr4E* gene was also observed and might be explained by its location in a repeat-rich area in the genome [42].

Likewise all isolates that overcome *Cf-5* resistance gene collected elsewhere in the world had lost the entire *Avr5* gene, except isolate IPO 1979 which has a frame shift mutation in the signal peptide leading to a pseudogene [13]. This study revealed that all isolates that could not overcome *Cf-5* resistance gene possessed a wild-type *Avr5* gene without any DNA modification, and two isolates of race 4.5.9 with a glycine to arginine substitution in C-terminus of *Avr5* protein caused loss of HR-inducing activity on tomato plants carrying the *Cf-5* gene (Fig 3). These results imply that accumulation of mutations in the *Avr5* gene occur less frequently than loss of the gene from the genome of *C. fulvum*. Like the *Avr9*, the *Avr5* gene in the genome of *C. fulvum* is also surrounded by repetitive elements, which might explain *Avr5* gene instability leading to its loss [13]. The *Avr5* gene functions as a virulence factor, but the intrinsic function of the *Avr5* protein is not known yet [13].

Although the Japanese population of *C. fulvum* has evolved independently from populations elsewhere in the world, it is striking that all unique DNA modifications observed in the *Avr* genes leading to adaptation to different *Cf* genes have similar biological effects on the encoded effector proteins [9, 12, 21].

Following similar overall mechanisms of adaptation in *C. fulvum* populations independently in different parts of the world might be related to the intrinsic functions of the *Avr* proteins, or their indirect or direct interaction with corresponding *Cf* proteins. *Avr2* interacts indirectly with the *Cf-2* protein through inhibition of *Rcr3*, which triggers a *Cf-2*-mediated HR [50–52]. *Avr9* is also expected to interact indirectly with the *Cf-9* protein [53]. A high affinity binding site is assumed to be the target of *Avr9* which is guarded by the *Cf-9* protein. Indirect interaction of an *Avr* protein with the corresponding *R* protein favors jettison of the *Avr*-encoding gene as has been observed for *Avr4E* and *Avr9* [12, 54, 55]. Point mutations in an *Avr* protein in order to overcome recognition by a matching *Cf* resistance protein might suggest direct interaction with that *Cf* protein as is suggested for the *Avr4* protein for which the *Cf-4* resistance protein is supposed to be the only host plant target [36].

**Prospects for protection of Japanese tomato cultivars against new races of *C. fulvum***

Based on the 41 genotypes identified in the Japanese population of *C. fulvum*, for only the most recently identified races the potential parental isolate could be inferred. The oldest races 0 and 2 isolates carry more diverse genotypes (16) and are more widespread in Japan than the new races that overcome the recently introduced *Cf* genes. The relatively new race 4, 4.11, 2.4 and 2.4.11 isolates overcoming the *Cf-4* gene that was introduced in the 1990s have spread over a large part of the country. Especially race 4.11 isolates have migrated into many different prefectures, suggesting that adaptation to both the *Cf-4* and *Cf-11* genes does not strongly affect viability of these isolates.

The very new race 9, 2.9, 2.5.9, 4.9, 4.5.9 and 4.9.11 isolates were mainly confined to one prefecture or a small region only. Overall, new races do not seem to spread very quickly, likely due to the fact that tomatoes are grown in greenhouses limiting the possibilities of conidia to escape.

Presently, most commercial tomato cultivars in Japan contain only single *Cf* genes, mainly the *Cf-4*, *Cf-5* or *Cf-9* resistance gene, but the high frequency of races overcoming these *Cf* genes has become a serious problem for tomato growers since 1990s [29–33]. The quick appearance of new race isolates is likely due to sequential introduction of single *Cf* genes in monocultures of tomato cultivars. In addition, as the use of agrochemicals in Japan has been
reduced significantly since 1990s, control of *C. fulvum* largely depends on resistance cultivars carrying *Cf* genes only. This has increased the selection pressure on this pathogen which has led to serious disease outbreaks caused by races with new virulence spectra [29–33]. Combining multiple *Cf* genes in one cultivar, as has occurred in most European countries, has slowed down the appearance of new races with complex virulence spectra. So far the new complex races mainly occur locally, introduction of multiple *Cf* genes (*Cf-2, Cf-4, Cf-5* and *Cf-9*) in one cultivar would slow down the spread of these races. These *Cf* genes could also be combined with the *Cf-6* gene as so far in Japan no race 6 isolates have been identified even elsewhere in the world. Also different *Cf-Ecp* genes could be introduced as worldwide no isolates of *C. fulvum* have been reported with *Avr6* or *Ecp* genes adapted to these *Cf* genes [45, 56, 57].

**Supporting Information**

S1 Table. Primers used in this study  
(DOCX)

S2 Table. Information on 133 isolates of *Cladosporium fulvum* collected in Japan.  
(DOCX)

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**Author Contributions**

Conceived and designed the experiments: YI IS RM JC PJGMW. Performed the experiments: YI PVH HB CM IS RM KF. Analyzed the data: YI PVH HB CM IS RM JC. Contributed reagents/materials/analysis tools: YI MK AN KF AB KAE. Wrote the paper: YI JC PJGMW.

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