High-Throughput Assessment and Genetic Investigation of Vegetative Compatibility in *Verticillium dahliae*

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Keywords

complementation tests, heterokaryon incompatibility, *nit* mutants, population biology, VCGs

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Abstract

Classification of isolates into vegetative compatibility groups (VCGs) using nitrate-non-utilizing (*nit*) mutants has been widely used for the characterization of *Verticillium dahliae* populations. However, certain methodological limitations prevent its application on a large scale. Furthermore, systematic investigations into the genetics underlying complementation tests between *nit* mutants of fungal isolates (i.e. heterokaryon formation) are lacking for *Verticillium* species. In this work, a diverse collection of 27 *V. dahliae* isolates – including representatives of all VCGs, both mating types, and heterokaryon self-incompatible isolates – was employed for the development and optimization of (i) a protocol for the rapid generation of *nit* mutants of *V. dahliae* isolates using UV-irradiation and (ii) a reproducible high-throughput procedure for complementation tests between *nit* mutants in liquid cultures using 96-well microplates. The genetic analysis of selected heterokaryons demonstrated that the frequently encountered ‘weak’ cross-reactions between VCGs and their subgroups can be actually heterokaryotic, implying the absence of strict genetic barriers between VCGs. In conclusion, we provide in this work an optimized method for the high-throughput VCG assignment of *V. dahliae* populations and a genetic analysis of heterokaryons that may have serious implications for the interpretation of VCG classification data. These advancements in the available methodology and the genetic background of vegetative compatibility grouping may contribute to a better understanding of the population biology of *V. dahliae* and possibly other mitosporic fungi.

Introduction

*Verticillium dahliae* is a soilborne phytopathogenic ascomycete that causes vascular wilt in more than 400 plants. The limited host specialization of the fungus, its vascular habitat, the longevity of its resting structures and the shortage of resistant host plant varieties have hindered control of the notorious *Verticillium* wilt disease (Klosterman et al. 2009). A thorough understanding of the genetic diversity and the relationships within and between *V. dahliae* populations is predicted to facilitate effective disease management (López-Escudero and Mercado-Blanco 2011).

Although mating-type genes have been described in *V. dahliae* (Usami et al. 2009), and population genetics analyses have revealed some putative signs of sex (Atallah et al. 2010; Inderbitzin et al. 2011), a sexual stage of the fungus has never been found so far, and *V. dahliae* populations are considered to be clonally propagating for the most part (Klosterman et al. 2009; Gurung et al. 2014). Apart from the accumulation of mutations (Anderson and Kohn 1995) and chromosomal rearrangements (de Jonge et al. 2013), genetic variation may be generated with restricted exchange and reassortment of genetic information through heterokaryosis and the subsequent parasexual cycle (Puhalla and Mayfield 1974; Typas...
and Heale 1978). However, heterokaryon formation is often prevented by vegetative incompatibility systems (Typas and Heale 1976), similar to what is observed in most fungi (Leslie 1993). Different methods have been employed for the classification of V. dahliae isolates according to their compatibility behaviour (Typas and Heale 1976; Puhalla and Hummel 1983; Joaquim and Rowe 1990). Among them, complementation tests of nitrate-non-utilizing (nit) mutants have been widely adopted in the last two decades for the characterization of V. dahliae populations. According to this method, V. dahliae isolates are assigned to the same vegetative compatibility group (VCG) if viable heterokaryons with prototrophic-like growth are formed when complementary nit mutants of the isolates are confronted on minimal medium (Joaquim and Rowe 1990). Five main VCGs are generally recognized today for V. dahliae populations, that is VCGs 1–4 and 6, with each of VCGs 1, 2 and 4 being subdivided into A and B subgroups, depending on the extent of heterokaryon formation; VCGs 3 and 6 are only rarely encountered (Katan 2000; Bhat et al. 2003; Jiménez-Díaz et al. 2006; Papaioannou et al. 2013b).

Extensive research in model fungi, such as Neurospora crassa and Podospora anserina, has revealed that incompatibility is governed by a set of heterokaryon (het) or vegetative incompatibility (vic) loci, with hyphal compartmentalization and programmed cell death being triggered in any heterokaryon that contains nuclei with different alleles at any one of these loci (Hutchison and Glass 2012). A similar genetic control of incompatibility is predicted for other fungi, although the relevant gene repertoire is still unknown for almost all of them. Based on this assumption, it is inferred that all members of a VCG of an asexual fungus are identical at all het loci, which may probably result from common descent. Thus, each VCG can be often regarded as a clonal lineage that is isolated from other VCGs. Members of distinct VCGs have been traditionally hypothesized to differ in several physiological and pathogenicity-related characteristics, rendering VCG analysis a convenient and informative tool for the characterization of fungal populations (Leslie 1993; Katan 2000).

A number of recent studies have revealed a previously unexpected complexity regarding molecular relationships within V. dahliae VCGs and between their subgroups, by employing different molecular tools (Collado-Romero et al. 2006, 2008; Martin 2010; Papaioannou et al. 2013a,b). Remarkably, main VCGs 2 and 4 were overall demonstrated to comprise molecularly distinct subgroups, with each of them being more closely related to a subgroup of the other VCG than its ‘sister’ subgroup (i.e. a VCG 2A-4B and 2B-4A pattern of relatedness was observed), and VCG 2B was shown to be highly heterogeneous and polyphyletic (Collado-Romero et al. 2008; Papaioannou et al. 2013a,b; Jiménez-Gasco et al. 2014). Despite these complications, however, and although elaborate molecular methods and tools for the characterization of fungal populations at a high resolution are increasingly available today (Martin 2010; Davey et al. 2011; Papaioannou et al. 2013a), traditional VCG analysis is still important for the study of V. dahliae populations. A relationship between certain VCGs and the taxonomic position of the original host of V. dahliae isolates has been recorded, for example for VCGs 1 and 2 with cotton and VCG 4 with solanaceous plants (Joaquim and Rowe 1990; Daayf et al. 1995). Furthermore, certain VCGs or subgroups have been correlated in several cases with virulence on specific hosts such as tomato, potato, cotton, olive, eggplant, sweet pepper and artichoke (e.g. Joaquim and Rowe 1991; Strausbaugh et al. 1992; Daayf et al. 1995; Korolev et al. 2000; Tsror (Lahkim) et al. 2001; Bhat et al. 2003; Collins et al. 2005; Jiménez-Díaz et al. 2006). In addition, significant differences in morphology and physiology have been reported between different VCGs (Korolev et al. 2000; Zeise and von Tiedemann 2001; Papaioannou et al. 2013b). The phytopathological significance of VCG analyses for V. dahliae populations is reflected in the unabated frequency of relevant reports in the literature (some recent examples include Dung et al. 2013; El-Bebany et al. 2013; Iglesias-Garcia et al. 2013; Papaioannou et al. 2013b).

Although VCG classification using nit mutants proved sufficiently convenient to allow research-level investigations in V. dahliae, it appears impractical for application on a larger scale (Leslie 1993). Microplate-based complementation tests on MM agar have been previously used with Fusarium species (Klittich and Leslie 1988), but an equivalent high-throughput method has never been reported for Verticillium. Such methodological limitations hinder VCG classification of extended populations of the fungus, which may be essential for the determination of significant correlations between the VCG system and the fungal phenotype, as well as for large-scale molecular analyses of relationships on the VCG level. Furthermore, a significant variability in the extent of prototrophic growth has generally been observed among different pairings of nit mutants, ranging from reactions that resemble wild-type prototrophic growth (‘strong’ reactions) to less vigorous interactions with sparse aerial hyphae and slow growth rates (‘weak’ reactions); marginal
reactions have also been detected, usually characterized by dots of microsclerotia along the contact line of the confronted *nit* mutants (Joaquim and Rowe 1990; Daayf et al. 1995; Korolev et al. 2000; Göre 2009; Papaioannou et al. 2013b). The traditional VCG classification system for *V. dahliae* depends mostly on ‘strong’ and reproducible reactions for the assignment of isolates into VCGs, although a detailed genetic comparison of the different types of interactions has never been reported. This analysis may advance our understanding of heterokaryon incompatibility in *V. dahliae* and contribute towards reconsidering the ‘traditional’ way VCG classification data is interpreted.

The aims of this study, therefore, were (i) to expedite VCG classification of *V. dahliae* isolates by applying UV-mutagenesis for the rapid generation of *nit* mutants and developing a novel method for their consistent and high-throughput complementation testing and (ii) to compare genetically for the first time the two types of complementation (i.e. ‘strong’ and ‘weak’ reactions) that are frequently observed during VCG assignment tests.

**Materials and Methods**

**Fungal isolates, growth media and culture conditions**

A total of 27 *V. dahliae* isolates from various hosts and geographic origins were employed in this study (Table 1). Members of all VCGs – as well as three heterokaryon self-incompatible (HSI) isolates – and representatives of both mating types of this species were included in the collection. Potato dextrose agar (PDA) and Czapek-Dox agar (Typas and Heale 1976) were used as the complete and minimal medium (MM), respectively, except for the characterization of *nit* mutants, which was performed on minimal media with different nitrogen sources (Correll et al. 1987). Monoconidial cultures were obtained from all isolates and were stored as conidial suspensions (at concentrations of at least $10^6$ conidia/ml) in potato dextrose broth (PDB) with 20% glycerol, at –80°C. Prior to each experiment, active cultures of the isolates were obtained directly from glycerol stocks by plating aliquots on PDA plates and incubating them for 1 week at 24°C, in the dark.

**Production and characterization of *nit* mutants**

*Nit* mutants of *V. dahliae* isolates were obtained either by the standard procedure involving selection of resistant sectors of agar inocula (Joaquim and Rowe 1990) on the chlorate-amended medium WAC (Korolev and reactions have also been detected, usually characterized by dots of microsclerotia along the contact line of the confronted *nit* mutants (Joaquim and Rowe 1990; Daayf et al. 1995; Korolev et al. 2000; Göre 2009; Papaioannou et al. 2013b). The traditional VCG classification system for *V. dahliae* depends mostly on ‘strong’ and reproducible reactions for the assignment of isolates into VCGs, although a detailed genetic comparison of the different types of interactions has never been reported. This analysis may advance our understanding of heterokaryon incompatibility in *V. dahliae* and contribute towards reconsidering the ‘traditional’ way VCG classification data is interpreted.

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Katan 1997) or by exposing conidia to ultraviolet (UV) irradiation (Typas and Heale 1976). In the latter method, approximately $10^5$ conidia of each isolate were spread on WAC medium plates prior to exposure at 20 cm from a germicidal UV-C lamp emitting 95% of the radiation at 254 nm. At this distance, the radiation intensity was estimated at approximately $110 \mu W/cm^2$, and the UV dosage was adjusted to a 5% survival rate by exposing conidia for 20 s. Plates were scored for the emergence of chlorate-resistant colonies up to 10 days post-UV-irradiation. Phenotype identification of all nit mutants produced by either method was performed by growth assays on MM with different nitrogen sources (Correll et al. 1987). The reversion rates of nit mutants were calculated by spreading dense conidial suspensions of each isolate on MM and viable counting on PDA, after appropriate serial dilutions.

**Complementation tests and heterokaryon analysis**

Complementation tests between nit mutants on solid MM were performed according to standard procedures (Joaquim and Rowe 1990), except that plates were scored for prototrophic growth up to 5 weeks after inoculation. All pairings were performed in three independent repetitions. Heterokaryon analysis was also carried out according to previously described methods (Typas and Heale 1976; Puhalla and Spieth 1983). Ninety-six-well microplates were routinely used for complementation tests between nit mutants in liquid MM, and several methods of inoculation were tested in a preliminary investigation. The optimal procedure that was selected for all subsequent experiments involved (i) mixing 25-µl aliquots of conidial suspensions (with concentrations ranging from $10^5$ to $10^6$ conidia/ml) of two isolates per pairing in each well of the microplate, (ii) addition of liquid MM to a final volume of 100 µl in each well and (iii) static incubation of plates at $24^\circ C$, in the dark, for 10 days. Following incubation, compatibility of nit mutants in each well was assessed by recording the amount of prototrophic mycelial growth (an example is provided in Fig. 1). At least two independent nit mutants of each isolate were tested, and all pairings were performed in four to eight repetitions each. To rule out the possibility of reversion, the mycelial mats of selected pairings were transferred onto MM agar plates, and their conidia were checked for nit phenotype or prototrophy. To test the congruence between the results of the new method and the traditional complementation tests of nit mutants on solid MM, all pairings were repeated on MM agar, in three repetitions each, according to standard procedures (Joaquim and Rowe 1990), and results were recorded up to 5 weeks after inoculation. When sterile cellophane sheets were interposed between the inocula of selected pairings, no growth was observed, suggesting that complementation was not due to cross-feeding (syntrophism).

**Results**

**Rapid production of nit mutants with UV-mutagenesis**

A total of 785 chlorate-resistant mutants (Table S1) were produced from the 27 V. dahliae isolates of all

![Fig. 1 Example of high-throughput complementation testing of several nit mutants of selected Verticillium dahliae isolates on a 96-well microplate. Liquid MM (100 µl), inoculated with mixed conidial suspensions of different pairs of nit mutants, is contained in each well, and the microplate has been incubated at 24°C, in the dark, for 10 days. Considerable mycelial growth in a well, usually accompanied by significant pigment production, denotes complementation between the tested mutants in that well.](image-url)
VCGs (Table 1) with the agar plug sectoring and UV-irradiation methods. Remarkably, UV-treatment resulted in high frequency of chlorate-resistant colonies, reaching an average of approximately $10^{-2}$, at a 5% survival rate, 10 days after UV-irradiation. Almost half of all chlorate-resistant colonies produced by both methods (45.4% in total) were classified as stable nit mutants (i.e. no reversion to prototrophy was observed after at least three subculturings on MM). Both methods were equally successful for the generation of nit1 (51.2 and 53.1% of spontaneous and UV-induced mutants, respectively) and nitM (48.8 and 46.9%, respectively) mutant classes. Although some variability in the mutants’ growth characteristics was observed on nitrite, no actual nit3 mutants were recovered, as intra-isolate complementation tests of all ‘suspected’ mutants allowed their assignment to the nit1 class. Independent nitM mutants of some isolates could be assigned to three different complementation groups, which generally complemented each other more vigorously than complementation in intra-isolate pairings between nit1 and nitM mutants (data not shown).

A significant proportion of chlorate-resistant colonies produced by both methods (47.1% in total) were able to grow on MM with nitrate as the nitrogen source, after one to three subculturings on MM. Furthermore, 5.5% of all chlorate-resistant colonies exhibited wild-type growth characteristics on nitrate and nitrite but were unable to grow on hypoxanthine.

Table 2 Results from the new method for complementation testing of nit mutants produced from Verticillium dahliae isolates of all VCGs, in all possible combinations within and among VCG subgroups, in liquid MM microcultures$^{a,b}$

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$^a$All V. dahliae isolates have been matched to a one-letter code [A to Z; Table 1] for the sake of data presentation.

$^b$The vigour of complementation reactions is symbolized as ‘+’ (for ‘strong’ reactions leading to dense prototrophic mycelial growth, usually accompanied by extended microsclerotia production), ‘+/+’ (for ‘weak’ reactions characterized by slight mycelial growth or only limited microsclerotia production), or ‘–’ (the absence of macroscopically visible mycelial growth).

$^c$The full VCG assignment of isolates L and M, which are known to complement tester isolates of more than one VCG subgroups, is 2B (2A) and 2AB (1, 4A), respectively (secondary VCG affinities are provided in brackets; Table 1). HSI: heterokaryon self-incompatible.

$^d$Isolate D – V6611 (328 v-1) – was originally reported as heterokaryon self-incompatible (HSI), but vegetatively compatible with isolates of VCG 1B (Collado-Romero et al. 2006).
Although these mutants were recovered by both methods, they were more abundant among the UV-induced and significantly more frequent among the chlorate-resistant mutants of *V. dahliae* isolates PCW and CA26 (21.4 and 37.5%, respectively). Similarly, ten mutants unable to grow normally on any of the three nitrogen sources, but growing on PDA and on MM with ammonium tartrate as the sole nitrogen source, were identified mainly among the UV-induced mutants. Finally, no *nit* mutants were recovered from a single isolate (TO-0), as it appeared inherently resistant to chlorate at the concentration that was used in this work.

To examine the genetic stability of *nit* mutants, the reversion rates of 46 randomly selected mutants were determined (Table S2). Among these mutants, stability varied remarkably in a very wide range of reversion rates from approximately $10^{-3}$ to $10^{-8}$. Notably, the reversion rates were associated with individual isolates in an isolate-specific manner, with intra-isolate variation among independent mutants never exceeding two orders of magnitude (Table S2). On the other hand, no correlation with VCGs, method of mutants’ production or mutant class was detected.

A high-throughput procedure for complementation testing of *nit* mutants

The rapid and high-throughput method developed here for the classification of *V. dahliae* isolates into VCGs involved standard 96-well microplates containing liquid MM, in which *nit* mutants were paired in all possible combinations, in four to eight repetitions of each pairing. With the exception of isolate TO-0, from which no *nit* mutants were recovered, 26 isolates of all VCGs were tested (Table 1; an example is illustrated in Fig. 1). Results were clear-cut and highly reproducible (averaging 97.5% between repetitions of each pairing) 10 days after inoculation, and they are summarized in Table 2. After recording results in microplate wells, the mycelial mats of 24 randomly selected pairings were transferred to the surface of solid MM, where they all retained slow growth ability, while their conidia could not grow on MM, suggesting that they were produced by actual heterokaryons rather than revertants. The results from the new method were in excellent agreement with the traditional complementation tests (an overall congruence value of 96.0% was recorded). Furthermore, the effect of *nit* mutants’ stability on complementation testing was studied. Notably, when mutants with reversion rates higher than $10^{-6}$ were used (Table S2), false-positive results were frequently obtained with both methods. This was presumably due to reversion to prototrophy and subsequent growth on MM, underlining the need for careful selection of *nit* mutants prior to complementation testing with either method.

Except for VCGs 1A and 6, which never exhibited cross-reactions with other VCGs, several ‘bridging’ reactions were recorded with both methods between all other VCGs and their subgroups, including pairings with some of the widely used international VCG tester isolates, for example PH, 115, 70-21, BB and S39 (Table 2). The extent of prototrophic growth in these cross-reactions ranged from limited mycelial development (‘weak’ reactions) to the production of small dots of microsclerotia at the contact zone (marginal interactions) (some examples are shown in Fig. 2). Five of the tested *V. dahliae* isolates that had been previously assigned to VCGs (Table 1), that is V607I and V66I (originally characterized as VCG 1B and 1B HSL, respectively), V830 and Dvd-E6 (VCG 4A), and Cf.162 (VCG 6), generally failed to complement other isolates from the collection tested. Complementary self-pairings of these isolates were also consistently unsuccessful, even though all of their available *nit* mutants were tested in several repetitions and all possible combinations. These isolates were, therefore, characterized here as heterokaryon self-incompatible (HSI).

Heterokaryon analysis

To investigate the genetic differences between ‘strong’ and ‘weak’ complementation interactions, ten presumed heterokaryons from each of the ‘strong’ and ‘weak’ types were transferred to solid MM. When sterile cellophane sheets were interposed between the two inocula of each pairing, no growth was observed, suggesting that complementation was not due to cross-feeding (symbiosis). It was also observed that heterokaryons deriving from originally ‘weak’ interactions had generally slower growth rates than those from ‘strong’ reactions, were highly irregular in the shape of the colony and pigment distribution, occasionally produced fast-growing sectors and were frequently dominated by one of the parental types (most often the *nitM* type; data not shown) after one to three subculturings on MM (examples are shown in Fig. 2). In contrast, most heterokaryons from ‘strong’ reactions were stable in culture after at least five subculturings.

For further analyses, three heterokaryons of each of the ‘strong’ and ‘weak’ types – including heterokaryons between different VCGs or subgroups – were
randomly selected (Table 3). Mycelial agar block transfers from the centre of all six heterokaryons always grew on MM, giving rise to characteristic compact colonies with a wide outer margin of thin mycelium, which resembled the growth of nit mutants (Fig. 2), and slower growth rates than the

Table 3 Conidial and hyphal tip analyses of *Verticillium dahliae* heterokaryons

<table>
<thead>
<tr>
<th>Heterokaryon components</th>
<th>Nuclear A:B ratio among conidia (No. tested)</th>
<th>No. of hyphal tips of nuclear type</th>
<th>Mean colony diameter (mm)a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A (nit1) B (nitM)</td>
<td>A (nit1) B (nitM)</td>
<td></td>
</tr>
<tr>
<td>Heterokaryons derived from initially ‘strong’ reactions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ls.17.10S (VCG 2B)</td>
<td>Ls.17.15S (VCG 2B)</td>
<td>1.0: 2.9 (525)</td>
<td>56</td>
</tr>
<tr>
<td>T9.8U (VCG 1A)</td>
<td>V44.15S (VCG 1A)</td>
<td>1.0: 3.8 (489)</td>
<td>24</td>
</tr>
<tr>
<td>70-21.7U (VCG 3)</td>
<td>PCW.20U (VCG 3)</td>
<td>1.9: 1.0 (505)</td>
<td>58</td>
</tr>
<tr>
<td>Heterokaryons derived from initially ‘weak’ reactions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>115.1S (VCG 2B)</td>
<td>C138.42S (VCG 2B)</td>
<td>1.0: 5.4 (516)</td>
<td>38</td>
</tr>
<tr>
<td>V320I.5U (VCG 2A)</td>
<td>Ls.17.15S (VCG 2B)</td>
<td>1.0: 8.6 (435)</td>
<td>6</td>
</tr>
<tr>
<td>530-1.23S (VCG 2AB)</td>
<td>S39.45S (VCG 4B)</td>
<td>3.4: 1.0 (412)</td>
<td>41</td>
</tr>
</tbody>
</table>

*aMean heterokaryon diameter after incubation for 8 days at 24°C, in the dark.*
parental \textit{nit} mutants on appropriately supplemented MM. Conidia of 4-week-old heterokaryons invariably retained auxotrophy when spread on MM, and when analysed for their genotypes, they showed ratios ranging from approximately 1:1 to 1:9, most often being skewed towards the \textit{nitM} type. These ratios were more disproportionate for ‘weak’ rather than for ‘strong’ heterokaryons (Table 3). When 0.5-mm-wide hyphal tips from the margins of heterokaryons were transferred to MM, they were homokaryotic for either one or the other \textit{nit} marker, with one predominating (Table 3).

\textbf{Discussion}

Although \textit{nit} mutants-based VCG complementation studies have been widely used for the characterization of \textit{V. dahliae} populations for more than two decades, the available methods are insufficient for use in large-scale analyses (Leslie 1993). In this work, UV-mutagenesis was applied for the generation of \textit{nit} mutants, and these were inoculated into liquid microcultures for the rapid, consistent and high-throughput VCG assignment of \textit{V. dahliae} isolates. With our optimized protocols, we demonstrate that both steps can be reliably accomplished in considerably less time than with the traditional procedures and used for the simultaneous testing of hundreds of independent pairings in each experiment, without the need for special equipment, extra storage space or additional applicant’s expertise. Notably, under the mutagenesis conditions that were used, the phenotype distribution, genetic stability and complementation behaviour of UV-induced \textit{nit} mutants were essentially identical with those deriving from spontaneous chlorate-resistant sectors. These features, together with the high reproducibility and excellent congruence with the traditional VCG classification procedures, render the new method a highly suitable candidate for the large-scale VCG profiling of extended populations of \textit{V. dahliae} and probably other fungi.

Great variability in the extent of prototrophic growth between different pairings, or even between independent \textit{nit} mutants of the same pair of isolates, has been observed in most VCG classification studies (Joaquim and Rowe 1990; Korolev et al. 2000; Jimenez-Diaz et al. 2006; Gore 2009). To avoid misinterpretations, ‘strong’ reactions, resembling the wild type of the fungus in the amount of prototrophic growth achieved, were almost exclusively used for the classification of isolates into the VCG system. However, ‘weaker’ reactions accounting for the cross-VCG (or ‘bridging’) interactions between different VCGs and their subgroups were also frequently reported, thus underlining the complexity of the genetics of the VCG system (Joaquim and Rowe 1990, 1991; Elena and Paplomatas 1998; Hiemstra and Rat-aj-Guranowska 2003; El-Bebany et al. 2013; Papaioannou et al. 2013b). In this study, we performed a genetic analysis and comparison of the two types of complementation interactions, and we demonstrated that (i) ‘weak’ reactions within and between VCGs can be due to actual heterokaryosis, similar to the ‘strong’ responses, and (ii) heterokaryons deriving from ‘weak’ interactions are unstable, with an uneven distribution of genetically different nuclei tending to segregate into novel homokaryotic mycelia. According to the general assumption underlying VCG analysis, that is that each VCG consists of genetically similar isolates with identical \textit{het} genotypes, heterokaryon compatibility has been predicted to be restricted to the members of the same VCG; different VCGs, on the other hand, have been regarded as incompatible groupings of isolates due to differences at one or more \textit{het} loci (Leslie 1993; Katan 2000). Notwithstanding this hypothesis, the ‘weak’ inter-VCG heterokaryosis observed in this and other studies implies the absence of strict genetic barriers between certain \textit{V. dahliae} VCGs. This may involve serious implications for the interpretation of traditional VCG classification data, especially if actual genetic recombination between different VCGs is demonstrated in future studies. Based on the available data, an attractive hypothesis would be that a spectrum between highly stable and more unstable heterokaryosis exists in \textit{V. dahliae} instead of clear-cut compatibility or incompatibility. Alternatively, the unstable ‘weak’ heterokaryosis might merely be an artefact of the traditional VCG assignment method that involves testing compatibility under forced conditions. These possibilities have to be addressed in future genetic investigations for the clarification of the situation.

Until a more comprehensive understanding of the \textit{het} gene system underlying compatibility of \textit{V. dahliae} is attained, population analyses are used to shed some light on the genetic relationships within and between VCGs. Several recent studies have challenged the traditional principle of VCG clonality, showing altogether that (i) not all VCGs or their subgroups correspond to clonal lineages; for example, VCG 2B is heterogeneous and polyphyletic, comprising isolates of more than one distinct molecular types that are grouped with different VCGs, and (ii) main VCGs 2 and 4 consist of genetically unrelated subgroups which are similar to subgroups of other VCGs (Collado-Romero et al. 2008; Martin 2010; Papaioannou...
et al. 2013a,b; Jiménez-Gasco et al. 2014). These findings may be possibly related to observations of ‘weak’ heterokaryosis made previously and in this work. Marked heterogeneity has been observed in the compatibility behaviour of isolates of the polyphyletic VCG 2B, with certain representatives complementing several but not all other VCG 2B members, and various others exhibiting usually weak ‘bridging’ ability with other VCGs (Jiménez-Díaz et al. 2006; Papaioannou et al. 2013b; present work). Furthermore, ‘bridging’ isolates between most of the known VCG subgroups have been frequently detected (Joaquín and Rowe 1990, 1991; Elena and Paplomatas 1998; Hiemstra and Rataj-Guranowska 2003; El-Bebany et al. 2013; Papaioannou et al. 2013b; present work), including, for example, the pair of VCG subgroups 2A and 4B, which have been shown to be indistinguishable at several molecular characteristics (Martin 2010; Papaioannou et al. 2013a,b; Jiménez-Gasco et al. 2014). The hypothesis that the usually ‘weak’ heterokaryons formed between these ‘bridging’ pairs of isolates may permit exchange of genetic material between VCGs could explain some of the previously unexpected phylogenetic affinities between certain VCG subgroups, for example 2B and 4A of the example mentioned above. In addition, the observation in the present study that VCGs 1 and 6 never exhibited cross-reactions with other VCGs is also in concordance with phylogenetic data that show them to be rather clonal entities (Collado-Romero et al. 2008; Martin 2010; Papaioannou et al. 2013a; Jiménez-Gasco et al. 2014) (considering the revised classification of VCG 6 isolates; Papaioannou et al. 2014). The possibility of parasexual genetic recombination between VCGs is further supported by the previous demonstration that the exchange of genetic material between incompatible isolates can be induced experimentally in Verticillium through ‘weak’ heterokaryosis under appropriately forced conditions (Typas 1983).

Although the sexual state of V. dahliae has never been found, mating-type idiomorphs are present in this species (Usami et al. 2009), and population analyses have revealed putative signs of recombination (Atallah et al. 2010; Inderbitzin et al. 2011). Based on these findings, it cannot be ruled out that part of the observed variation and the deviation from clonality according to the VCG system may result from rare or even ancient sexual recombination events. Although parasexuality has been experimentally established in Verticillium (Puhalla and Mayfield 1974; Typas and Heale 1978), its importance in natural populations of fungi has never been unequivocally demonstrated to date. Therefore, sexual recombination might explain some of the observed patterns of variation in V. dahliae, and future research in this direction is especially welcome.

Another interesting finding of this work is the variety of chlorate-resistant mutants, especially after UV-treatment. Mutants unable to grow on all nitrogen sources except for ammonium may be defective in the major nitrogen regulatory gene areA (Lenouvel et al. 2001), while additional rare phenotypes were also recorded. Interestingly, a remarkably broad and isolate-specific range of reversion rates for individual nit mutants was detected, and an impact of their stability on the results of complementation tests was demonstrated. These data underline the need for careful selection of appropriate nit mutants prior to complementation studies. Furthermore, we found evidence for a selective advantage of the nitM type over nitI mutations in V. dahliae heterokaryons. The possible relevance of this genetic imbalance to the reliability of complementation tests merits further examination. Finally, it was observed that five of the tested V. dahliae isolates that had previously been assigned to VCGs, appeared consistently heterokaryon self-incompatible in this study and unable to complement any other isolate of the collection. This phenomenon could be explained by the hypothesis of Jacobson and Gordon (1990), who tried to explain self-incompatibility in Fusarium oxysporum as a mutant artefact that arises in fungi maintained for long periods in artificial culture, although further study is necessary for a solid conclusion.

In conclusion, our work provides (i) an optimized method for the high-throughput VCG classification of V. dahliae isolates using nit mutants on a large scale and (ii) a genetic analysis of heterokaryons between nit mutants of V. dahliae isolates, which implies the absence of strict genetic barriers between VCGs. The results suggest that a systematic re-evaluation of vegetative incompatibility in Verticillium and its biological significance is essential for the unravelling of the population biology of the fungus. Novel methods suitable for large-scale application, such as the one provided here, are hopefully expected to facilitate this endeavour.

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References


Inderbitzin P, Davis RM, Bostock RM, Subbarao KV. (2011) The ascomycete *Verticillium longisporum* is a hybrid and a plant pathogen with an expanded host range. PLoS ONE 6:e18260.


de Jonge R, Bolton MD, Kombrink A, van den Berg GCM, Yadeta KA, Thomma BPHJ. (2013) Extensive


Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Nit mutants of Verticillium dahliae isolates that were produced and characterized in this work.

Table S2. Reversion rates of 46 nit mutants of Verticillium dahliae isolates produced by either UV-induction or selection of spontaneous chlorate-resistant sectors.