BRIEF REPORT

High molecular diversity of full-length genome sequences of zucchini yellow fleck virus from Europe

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Abstract
Zucchini yellow fleck virus (ZYFV), genus Potyvirus, is the causal agent of a disease of cucurbits. The genome sequences of seven ZYFV isolates of different origin were determined, two of which were reconstructed from a squash (Cucurbita sp.) collected in 2017 in Greece, while the others, accessions from the DSMZ Plant Virus Collection, were from samples collected in Italy, Greece, and France in the 1980s and 1990s. A high level of molecular diversity, well dispersed along the genome, was observed, but this was within the limits for assignment of the virus isolates to the same species. P1 was the most diverse gene, and isolates from squash contained an insertion in this gene.

The genus Potyvirus (family Potyviridae) includes many economically important viruses of major crops, causing diseases associated with leaf mosaic, fruit discoloration and deformation, yellowing, and necrosis [1]. Epidemics are driven by aphid vectors, which transmit the viruses in a non-persistent manner. Potyvirus particles have a filamentous morphology, are 680-900 nm long, and contain a single-stranded RNA molecule, approximately ~10 kb in length, coding for a polyprotein that is proteolytically processed into at least 10 functional proteins [2]. For taxonomy purposes, the sequence of the complete polyprotein is used for species demarcation, and identity thresholds (below 76% and 82% at nt and aa level, respectively) are used to establish species within the genus [2, 3].

Zucchini yellow fleck virus (ZYFV) was first described in 1981 in zucchini in Italy [4], and it has since been reported in other countries of the Mediterranean basin, including Greece [5], Lebanon and Syria [6], Israel [7], and France [8], and, more recently, in Iran [9]. Given its transmission by Myzus persicae and Aphis gossypii [4, 8], which are major pests and virus vectors of cucurbits, further spread of the virus and serious impact was expected. However, the occurrence of ZYFV infection has remained localized, and there have never been reports of significant outbreaks of the virus. Despite its description as a member of a distinct potyvirus species [10], molecular characterization of ZYFV has been so far limited to sequences of a short region of the genome encompassing the CP [10, 11], due to a lack of complete genome sequences.

We conducted studies to provide an inventory of common virus diseases of cucurbits in Greece, testing for watermelon mosaic virus (WMV), papaya ringspot virus (PRSV), Moroccan watermelon mosaic virus (MWMV), and zucchini yellow mosaic virus (ZYMV), which are known to be endemic in Greece [12–14] and spread by several aphid species [15, 16]. When using high-throughput sequencing (HTS) to analyze a squash (Cucurbita sp.) sample with virus symptoms that was collected in 2017 in central Greece and had tested positive by ELISA for a potyvirus [17], we discovered two distinct isolates of ZYFV. A comparison with full-length genome sequences of ZYFV isolates from the DSMZ Plant Virus Collection with diverse geographic origin and collection dates (Table 1) revealed a high level of molecular diversity. Notably, the Greek ZYFV isolates from squash contain a distinctive insertion in the P1 coding region, whose possible biological function remains to be determined.
During field surveys conducted in 2017 in Farsala (Larissa Prefecture, Central Greece), leaf samples from a symptomatic squash showing fleck and yellow pinpoint spots on old and young leaves (Supplementary Fig. S1) were collected, and they tested positive by ELISA using a potyvirus group-specific antibody (DSMZ RT-0573/1) [17]. Further screening with antisera against WMV (DSMZ RT-0203), PRSV (DSMZ RT-0805), ZYMV (DSMZ RT-0234), and MWMV (DSMZ RT-0747) gave negative results. We therefore employed HTS followed by virus discovery analysis using the workflow established at the DSMZ Plant Virus Department [18]. We sequenced the original squash sample for a primary virome investigation and a zucchini plant that was infected mechanically with homogenates of the original sample. Total RNA was extracted from symptomatic leaves, using an RNeasy Plant Mini Kit (QIAGEN) or a Spectrum Plant Total RNA Kit (Sigma-Aldrich). Ribosomal RNA was depleted, followed by random cDNA synthesis and second-strand synthesis using random octamer primers. Nextera XT libraries (Illumina) were prepared according to the manufacturer’s instructions and sequenced on a MiSeq or NextSeq instrument as paired-end reads. The reads were assembled de novo using Geneious Prime v. 2022.0.1 (Biomatters) and were screened by BLAST against a custom reference virus database to reveal significant hits against potyvirus sequences assigned to ZYFV. To study the genome diversity of ZYFV, the sequences of five virus accessions from the DSMZ Plant Virus Collection were also determined following the same workflow. The analysis included ZYFV isolates from the 1980s and 1990s collected from zucchini in Italy (DSMZ PV-0379, PV-1256, PV-1257) and Greece (DSMZ PV-1258), and from Ecballium elaterium in France (DSMZ PV-1263) (Table 1).

Bioinformatic analysis allowed the reconstruction of seven ZYFV genome sequences (Supplementary Table S1), two of which (PV-1248 isolate #1 and #2) were assembled from the zucchini that had been infected mechanically with homogenates from the squash sample collected in 2017. The assembled genome sequences ranged in length from 10,161 to 10,325 nt and contained an open reading frame (ORF) encoding a polyprotein of 3320-3338 aa in length, and they exhibited a genome organization typical of potyviruses (Fig. 1a). The 5´- and 3´-terminal ends of the genome of isolates PV-1256 and PV-1257 were verified by RACE-PCR assays and direct sequencing of the amplification products. Sequence alignment revealed that two isolates, PV-0379 from zucchini from Apulia and PV-1258 from Greece (from 1986), were ~99% identical, indicating introduction and/or exchange of the virus between these two geographically close areas. Given the similarity of the two sequences, we refer to the two isolates as one entity. A high degree of sequence diversity among the isolates was found, evenly distributed over the entire length of the genome. This confirmed the high molecular divergence reported previously for the 3´-terminal region of the genome [10]. Alignment of the polyprotein sequences revealed 75.4-82.8% and 83.6-92.9% identity at the nt and aa level, respectively (Supplementary Table S2), which, according to current guidelines, supports the assignment of these viruses to the same species. The isolate PV-1263 from squirting cucumber (E. elaterium) from France had the most divergent sequence (Supplementary Table S2), which may correlate with adaptation to this host, a wild perennial gourd plant. A phylogenetic analysis of the ZYFV genome sequences with those of other potyviruses showed that the ZYFVs form a distinct clade that is well separated from the clade including papaya ringspot virus (PRSV) and zucchini tigré mosaic virus (ZTMV) (Fig. 2). This striking diversity among the ZYFV sequences exceeded the diversity of PRSV isolates, forming a less-dispersed cluster. Pairwise comparison with the coding regions of PRSV revealed an extreme diversification of P1 (~42 and ~28% nt and aa identity, respectively), with highest similarity in the NIb (~69 and ~76%, respectively) followed by HC-Pro (~68 and ~75%, respectively) (Supplementary Table S1). Compared to ZTMV, the P1 gene of ZYFV showed ~43 and ~31% nt and aa identity, respectively, and highest identity was observed with NIb and HC-Pro (Supplementary Table S2).

The first mature protein, the serine protease P1, is predicted to cleave itself from HC-Pro at residues Tyr/Ser or Tyr/Thr (Supplementary Table S3). Its diversity in sequence and size among potyviruses is well known [19], and P1 was indeed found to be the most variable gene among the ZYFVs isolates. The predicted P1 protein contained a highly

| DSMZ accession | Country of origin | Year of collection | Host | GenBank accession no. |
|----------------|------------------|--------------------|------|-----------------------|
| PV-0379        | Italy, Apulia    | 1980               | Cucurbita pepo | ON456303              |
| PV-1256        | Italy, Sicily    | 1990               | Cucurbita pepo | ON456306              |
| PV-1257        | Italy, Sardinia  | 1995               | Cucurbita pepo | ON456307              |
| PV-1258        | Greece           | 1986               | Cucurbita pepo | ON456308              |
| PV-1263        | France           | 1987               | Ecballium elaterium | ON456309              |
| PV-1248        | Greece, Farsala  | 2017               | Cucurbita sp. | ON456304, ON456305    |
variable region at position 200-220, with the highest diversity found in the sequences from squash, which included an insertion with the duplication of an “NSSVEM” motif located closely upstream (Fig. 1b). Verification of the ZYFV genome sequence by Sanger sequencing of RT-PCR products (Supplementary Table S4) confirmed this unique feature in the two isolates from squash. Future investigations are needed to clarify whether the insertion is a characteristic motif of current ZYFV populations in Greece and to investigate its possible effect on biological functions. A functional role of the P1 and its diversity in determining the host range has been demonstrated for the potyviruses plum pox virus [20, 21] and tobacco vein mottling virus [24], MWMV [25], and pepper vein banding virus [26]. The PTK motif, which is also involved in aphid transmission [27], is located at position 309-311. The motifs IGN and CCC, which have been implicated in replication and movement of potyviruses [28–30], are located at aa positions 249-251 and 291-293, respectively.

The remaining viral proteins are produced by cleavage by the cysteine protease Nla-Pro with the typical sequence motifs V-X-[F;H]-[Q;E]/[A;G;S] flanking the cleavage sites (Supplementary Table S3). The proteins 6K1, CI, 6K2, VPg, and Nla-Pro are predicted to be of the same size in all ZYFV isolates. As observed in a previous analysis on aa preferences of Nla-Pro protease in the polyprotein of potyviruses [31], a glutamic acid residue is conserved at position -1 of the VPg/Nla-Pro junction. A duplicated cleavage motif at the Nlb/CP junction [10], with the sequence VYHQ/S, is present in all isolates, with an inter-motif variable region of 7 aa (Fig. 1c). A putative second Nlb/CP cleavage site has also been reported for ZTMV [32] and PRSV, although in the latter case, the motif is not completely conserved [10]. The biological significance of this feature of the polyprotein remains to be determined.

Pairwise comparisons of the CP sequence showed that PV-1263 from France was the most divergent virus, with
~87-90% aa sequence identity to the other isolates. As has been observed in the CPs of other potyviruses [33–35], most of the divergence was observed in the N-terminal region of the protein, at positions 13-15 and 24-32, while aa residues at positions 121 to 277 were identical among all sequences; a deletion of 3 aa at position 29-31 was unique to PV-1263. A phylogenetic analysis with other CP sequences available in the NCBI/GenBank database showed that the Greek isolates from squash grouped in a cluster with the isolate from Apulia/Greece, while the CP of the DSMZ accessions from Sicily and Sardinia and two sequences from melon collected in 2006 in Sicily [11] formed a separate sequence cluster (Supplementary Fig. S2). All ZYFV genomes encoded a “DAA(A)” motif in the N-terminal region of the CP, instead of the more common “DAG(X)” implicated in aphid transmission [36]. This motif was also reported in the potyviruses onion yellow dwarf virus and peanut mosaic virus [37].

The PIPO ORF, embedded within the P3 region of the polyprotein and translated in the +2 reading-frame [38], was annotated and found to contain the highly conserved $G_{1-2}A_{6-7}$ motif in the form GAAAAAA in all sequences.

To assess the role of natural selection at the molecular level, the rate of non-synonymous substitutions per non-synonymous site (dN) and the rate of synonymous substitutions per synonymous site (dS) were calculated for each virus gene using DnaSP v. 6.12.03 [39]. The results revealed that the ZYFV genome is under negative selection, with strong selection in HC-Pro and weak selection in P1 and PIPO (Supplementary Table S5). This is

Fig. 2 Phylogenetic tree constructed using the seven zucchini yellow fleck virus genome sequences assembled in this study and other potyvirus sequences obtained from the GenBank database. The evolutionary history was inferred using the maximum-likelihood method and the general time-reversible model implemented in MEGA [42]. The scale bar represents a value of 0.01 substitutions per site.
in accordance with common observations for potyviruses [40].

Since recombination is shaping potyvirus populations [40], we performed a recombination analysis with RDP5 [41] on a MUSCLE alignment of the ZYFV genome sequences and included a set of PRSV and ZTMV sequences (Supplementary Table S6). One recombination event in P1 of PV-1256 from Sicily was traced to PV-1248 isolate #1 as the major parent, but the minor parent is unknown (Supplementary Table S6). Two recombination events involved ZYFV and potyviruses of other species, but the recombination signals were only weakly supported. We can thus assume that the ZYFV isolates in this study had not undergone genetic recombinaton.

In summary, the complete genome sequences of ZYFV isolates from different European countries were determined. Sequence comparisons revealed that the ZYFV isolates from squash from Greece contained a unique motif in the P1 region, the biological role of which remains to be determined.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00705-022-05558-9.

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Data availability The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

Statements and declarations

Conflict of interest The authors have no relevant financial or non-financial interests to disclose.

Ethical approval This article does not contain any studies with human participants performed by any of the authors.

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