Alstroemeria yellow spot virus (AYSV): a new orthotospovirus species within a growing Eurasian clade

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Abstract
An orthotospovirus distinct from all other orthotospoviruses was isolated from naturally infected alstroemeria plants. Disease symptoms caused by this virus mainly consisted of yellow spots on the leaves based on which the name alstroemeria yellow spot virus (AYSV) was coined. A host range analysis was performed and a polyclonal antiserum was produced against purified AYSV ribonucleoproteins which only reacted with the homologous antigen and not with any other (established or tentative) orthotospovirus from a selection of American and Asian species. Upon thrips transmission assays the virus was successfully transmitted by a population of Thrips tabaci. The entire nucleotide sequence of the M and S RNA segments was elucidated by a conventional cloning and sequencing strategy, and contained 4797 respectively 2734 nucleotides (nt). Simultaneously, a next generation sequencing (NGS) approach (RNAseq) was employed and generated contigs covering the entire viral tripartite RNA genome. In addition to the M and S RNA nucleotide sequences, the L RNA (8865 nt) was obtained. The nucleocapsid (N) gene encoded by the S RNA of this virus consisted of 819 nucleotides with a deduced N protein of 272 amino acids and by comparative sequence alignments to other established orthotospovirus species showed highest homology (69.5% identity) to the N protein of polygonum ringspot virus. The data altogether support the proposal of AYSV as a new orthotospovirus species within a growing clade of orthotospoviruses that seem to share the Middle East basin as a region of origin.

Introduction
Members of the Tospoviridae continuously cause significant losses in both mono- and dicot crops worldwide [1, 2]. Although exact numbers indicating the economic impact of orthotospoviruses are not available, annual yield losses have been estimated to exceed at least 1 billion US$ for the type species Tomato spotted wilt virus (TSWV) only [3]. Due to this TSWV ranks high among plant viruses of economic importance worldwide [1, 4–6]. Orthotospoviruses are widespread and transmitted by at least 14 different thrips species [2, 7] in a circulative and propagative manner. Virus particles are membrane-bound with a quasi-spherical morphology of about 80-120 nm in diameter and contain a tripartite ssRNA genome of negative and ambisense polarity [6]. The L RNA is of entire negative polarity and codes for only one gene in the viral complementary (vc) sense, the RNA-dependent RNA polymerase (RdRp; L protein) required for viral replication. The M and S RNA both have an ambisense gene arrangement and contain two ORFs separated by a non-coding intergenic region (IGR). The M RNA encodes a non-structural (NSM) protein, in the viral (v) sense, that is involved in cell-to-cell movement and a glycoprotein precursor (GN-GC) in the vc-sense required for (receptor-mediated) entry into the midgut-epithelium of thrips and vector-specificity [8–13]. The S RNA similarly encodes the non-structural protein (NSs) in the v-sense that
acts as suppressor of RNA silencing and supports efficient virus transmission by the vector, and the major structural nucleocapsid protein (N) in vc-sense [14–18]. Although host range, vector specificity and serology of the N protein are often used to identify and/or characterize orthotospoviruses, they only become assigned as a new species if their N protein exhibits less than 90% sequence homology to any of the established orthotospovirus species [2, 19, 20].

Ever since TSWV re-emerged in the 1980's and started its world wide spread by the introduction and spread of *Frankliniella occidentalis*, new viral species have been discovered. This has led to the identification of a current total of 29 orthotospovirus species of which 11 are officially recognized by the ICTV while the remaining, listed as tentative species [2, 20–26]. During the past two decades many viruses have been observed and identified in ornamental crops of which some cause significant economical losses [27].

Currently, alstroemeria is one of the most important and successful cut flowers and its global production is still increasing [28, 29]. However, viral infections place a large constraint on the production levels and limit global trading [30–32]. To date 18 viruses have been reported from alstroemeria plants (Table S1) [29, 32–42] and among those are isolates from five different orthotospovirus species *i.e.*, TSWV, impatiens necrotic spot virus (INSV), iris yellow spot virus (IYSV) [32], tomato yellow ring virus (TYRV) [35] and alstroemeria necrotic streak virus (ANSV) [34]. From these, TSWV has the largest (natural and experimental) host range, exceeding more than 1350 species. INSV infects at least 300 host species, and mainly ornamentals [7], while IYSV and TYRV have a smaller host range and are frequently found in onion, leek and a few ornamentals [12, 35, 43, 44]. ANSV has been isolated most recently from alstroemeria in Colombia and, together with TSWV and INSV, clusters within the clade of American/New-World orthotospoviruses [2].

In 2000 an alstroemeria plant showing only chlorotic and yellow spots on the leaves (Figure 1), and reminiscent of a TSWV infection, was received by the Netherlands Inspection Service for Horticulture. Preliminary serological tests with available antisera to a wide range of different orthotospoviruses were negative and indicated the possible emergence of a new orthotospovirus species. Here data is presented on the biological, serological and molecular characterization of this Als-2000 virus isolate to support the proposal of a new tentative orthotospovirus species for which the name alstroemeria yellow spot virus (acronym AYSV) is proposed. The nucleotide sequence of the entire genome was elucidated using a simultaneous conventional and NGS cloning/sequencing approach.

**Materials and methods**

**Virus source, virus isolates and host range**

Leaf material of suspected virus-infected (Als-2000) alstroemeria was homogenized in PBS buffer (pH 7.0) with 0.01% sodium sulphite and mechanically inoculated to *Nicotiana benthamiana*. Orthotospovirus isolates TSWV BR-01 [19], tomato chlorotic spot virus (TCSV) BR-03 [19], groundnut ring spot virus (GRSV) SA-05 [19], iris yellow spot virus (IYSV-NL) [45] and TYRV-t [46] used in serological studies, were also maintained on *N. benthamiana*. For a host range analysis, sap from Als-2000-infected alstroemeria leaf material was mechanically inoculated on 29 plant species (at least 2 plants/species) and monitored during 3-4 weeks for symptom expression.

**Ribonucleocapsid purification, SDS-PAGE and serological analysis**

Ribonucleocapsid protein (RNP) was purified from Als-2000-infected *N. benthamiana* plants [47] and subjected
to 25-45% CsSO₄ gradient centrifugation. RNP fractions collected were used to immobilize rabbits for the production of a polyclonal antiserum. Prior to this, RNP fractions were subjected to a quality/purity check, in comparison to INSV and TSWV RNP preparations, on a 12% SDS-PAGE after denaturation in Laemmli buffer [48]. Rabbits were immunized 2-3 times with 50-100 μg of purified RNP emulsified with incomplete Freund’s adjuvant (1:1, w/v), during intervals of two weeks. Serum was collected two weeks after the last injection and used for the purification of IgG and as the preparation of an IgG-alkaline-phosphatase conjugate. For orthotospovirus differentiation double-antibody sandwich–enzyme-linked immunosorbent assays (DAS-ELISA) were performed using a panel of polyclonal antisera raised against Als-2000, TSWV, TCSV, GRSV, IYSV and TYRV [45-47]. Sap of infected and healthy N. benthamiana plants was diluted 1:30 and used as the antigen source and negative control, respectively.

**RNA purification and RT-PCR**

Viral RNA was purified either from purified RNPs using 1% SDS and hot phenol extraction followed by ethanol precipitation [9] or from infected plant material using Trizol buffer according to the manufacturer’s instructions (Invitrogen).

In a first approach to obtain Als-2000 viral sequences only a single primer “Asian Termini” (AT; 5′-CCC GGA TCCAGAGCAATCGAG-3′, containing a BamHI restriction site (underlined) and matching the 3′ terminal sequences of Asian orthotospoviruses (italics), was used for first-strand cDNA synthesis using M-MLV Reverse Transcriptase (Promega), and a subsequent PCR amplification [46] using Phusion High-Fidelity DNA Polymerase (New England Biolabs). Since the sequence of the amplicon obtained using the AT primer showed highest homology with the NS₅ gene of IYSV, a second approach was applied in which Als-2000 sequences were additionally RT-PCR amplified using primers IY1 (5′-CCCAGGGA TCCATGCTAACCCTAGGG-3′) and IY2 (5′-CCCGA GGATCCAAAATTAATATCTACTTCTTTG-3′) corresponding to IYSV N gene sequences (in italics) combined with the universal hairpin primer (UHP) (5′-CACT GGATCCTTTGTTTTTGTGTGTG-3′) as described before [45, 46]. Based on newly obtained sequences for Als-2000, specific primers were designed to verify/obtain additional sequences to complete the entire S RNA nucleotide sequence. Primers Als2000-NS₅-G-R (5′-GCT TGTTTTCTTTTTCTTCC-3′) and Als2000-G-R (5′-CCTCTAACAATATGACTTTCCC-3′) were used to amplify the 5′ ends of the NS₅ respectively G₅/G₆ glycoprotein precursor ORFs in combination with the AT termini primer. In analogy to the approach for the S RNA segment, new and specific primers were designed to verify and/or obtain sequences for the intergenic region of the M RNA segment. All PCR fragments obtained were cloned into pJET blunt1.2 vector (Fermentas) and their sequences determined (Eurofins).

**Sequence analysis of S and M RNA segments**

Contigs for the M and S RNA segments were assembled from obtained sequences using DNAStar software and used to search the GenBank database using BLASTn and BLASTp. Multiple sequence alignment and phylogenetic analysis were performed using MEGA 7 software [49]. The G₅-G₆ sequence was analyzed for the presence of N- and O-linked glycosylation sites by prediction using the NetNGlyc 1.0 and NetO GLyc 3.1 Servers (http://www.cbs.dtu.dk/services/NetNGlyc/; http://www.cbs.dtu.dk/services/NetOGLyc-3.1/). Potential signal cleavage sites within the glycoprotein were predicted using the SignalP 4.1 server (http://www.cbs.dtu.dk/services/SignalP/). Transmembrane regions within the G₅-G₆ precursor protein were predicted using the TMHMM Server v.2.0 program (http://www.cbs.dtu.dk/services/TMHMM/).

**Sequences analysis of Als-2000 by Next Generation Sequencing**

For the sequencing of the entire Als-2000 genome using next generation sequencing (NGS) technology, total RNA was isolated from systemically infected N. benthamiana leaves with the Qiagen RNeasy plant mini kit according the manufacturers’ procedures (Qiagen). Samples were prepared for HighSeq sequencing according to Verbeek et al. [50]. Sequence data were analyzed using the CLC Genomics Workbench 8.5.1 (Qiagen) as earlier described [51]. To obtain and verify the full length sequence of Als-2000 the 5′- and 3′-ends of the viral RNA segments and regions from which only few reads were obtained, were RT-PCR amplified with specific primers and directly sequenced by Sanger sequencing (Macrogen).

**Thrips transmission**

To determine the potential of Thrips tabaci for transmission of Als-2000, 10 first and early second instar larvae of T. tabaci were given an acquisition period of 24h in Als-2000 infected Emilia sonchifolia and next transferred to healthy E. sonchifolia plants. Plants were kept under cages at room temperature and monitored for symptom expression for two weeks. The presence of Als-2000 in plants was tested by RT-PCR using primers IY1 and IY2. Fragments of expected
sizes were collected and the AYSV sequence confirmed by sequence analysis.

Results

Host range analysis

From 31 plant species mechanically inoculated with Als-2000, from here onwards referred to as alstroemeria yellow spot virus (AYSV), 17 species did not show any sign of infection, 9 species only showed local symptoms while 5 species displayed local and systemic symptoms (Table 1). Back-inoculation of the isolate from *N. benthamiana* on the alstroemeria cultivar Dimention did not induce any symptoms. The data indicated that the virus had a relatively narrow host range and among the plants tested only dicots showed local or systemic symptoms.

N protein electrophoresis and DAS-ELISA

To comparatively analyze AYSV and serologically differentiate this virus from other orthotospoviruses, AYSV RNPs were purified and analyzed by SDS-PAGE. A major band of approximate *M*~w~ 30 kDa migrated at a position in between the INSV and TSWV N proteins and likely represented the AYSV N protein (Figure 2, left panel). Polyclonal antibodies produced against the AYSV RNPs were used to analyze its serological relationship to a set of orthotospoviruses. In DAS-ELISA, only leaf extracts of AYSV-infected plants reacted positively with polyclonal antiserum against AYSV, while no positive reaction was observed with antisera raised against other orthotospovirus species (Figure 2, right panel).

Sequence analysis and characterization of the S RNA, M RNA and L RNA

The entire sequence of AYSV S RNA consisted of 2734 nucleotides (nt) with 5′ terminal (72 nt) and 3′ terminal (71 nt) untranslated regions (UTR) that contain the highly conserved 8 terminal nucleotides present in all orthotospovirus RNA genome segments. AYSV terminal sequences show almost entire base pair complementarity of the first 18 nt with only 3 base pair mismatches (5′-AGAGCA AUCGNNGNAUAA-3′). This complementarity continues, although to a lesser extent, to ~ 60 nucleotides of the UTR sequences and assists in the panhandle formation of the S RNA segment.

The TYRV S RNA contains a vORF starting with an AUG at nucleotide position 73 and terminating with a UAG codon at position 1404, coding for 443 residues of the NS~S~ protein with a predicted molecular mass of 50.1 kDa. The

| Plant family and species | Local | Systemic |
|--------------------------|-------|----------|
| **Alstroemeriacae**      |       |          |
| Alstroemeria sp. cv. Dimension | - | - |
| **Amaranthaceae**        |       |          |
| Gomphrena globosa | - | - |
| **Amaryllidaceae**       |       |          |
| Allium porrum | - | - |
| Hippeastrum sp. cv. Orange Souvereign | - | - |
| Hippeastrum sp. cv. Red Lion | - | - |
| **Asteraeae**            |       |          |
| Emilia sonchifolia | CS, NS | CS, NS |
| Zinna elegance | - | - |
| **Balsaminaceae**        |       |          |
| Impatiens sp. | NL,NR,N | - |
| **Chenopodiaceae**       |       |          |
| Chenopodium amaranticolor | - | - |
| Chenopodium quinoa | - | - |
| **Cucurbitaceae**        |       |          |
| Cucumis sativus cv. Hokus | - | - |
| Cucurbita pepo cv. Black Beauty | - | - |
| **Fabaceae**             |       |          |
| Arachis hypogea | - | - |
| Vicia faba cv. Wikiem Majort | NL | - |
| Vigna unguiculata cv. Black Eye | NL | - |
| Vigna unguiculata cv. Early Red | CL,NL | S |
| Phaseolus vulgaris cv. Pinto | NL | - |
| **Iridaceae**            |       |          |
| Iris hollandicum cv. Blue Magic | - | - |
| **Solanaceae**           |       |          |
| Capsicum annuum cv. Westlandse Grote Zoete | - | - |
| Datura metel | NL | - |
| Datura stramonium | NL | - |
| Physalis floridana | - | - |
| Pisum sativum | - | - |
| Nicotiana benthamiana | CL | CS, NS, C,RG |
| Nicotiana clevelandii | C,N | C,N,RG |
| Nicotiana glutinosa | NL | - |
| Nicotiana hesperis-67A | CL,NL | NL |
| Nicotiana occidentalis-PJ | NL | - |
| Nicotiana rustica | - | - |
| Nicotiana tabacum cv. White Burley | NL,NR | - |
| Solanum lycopersicum cv. Money Maker | - | - |

C chlorosis; LC leaf curling; CL Chlorotic lesions; CS Chlorotic spot; N necrosis; NL necrotic lesions; NR necrotic rings; NS necrotic spots; RG retardation in growth; S symptomless infection
Characterization of a new orthotospovirus species from alstroemeria

The vcORF starts with an AUG at nucleotide position 2663 and terminates with a UAA stop codon at nucleotide position 1845, coding for 272 residues of the N protein with a predicted molecular mass of 30.1 kDa. A noncoding intergenic region (IGR) encompasses nucleotide position 1405 to 1844 and is highly rich in stretches of A and U residues, enabling the putative formation of a hairpin structure (data not shown).

Upon multiple sequence alignment of the AYSV N protein to other orthotospoviral N proteins, AYSV N showed highest identity (69.5 %) with polygonum ringspot virus (PolRSV). This homology is well below the threshold for demarcation of new orthotospovirus species and indicated that AYSV tentatively presents a new distinct species. In a phylogenetic analysis based on the N protein, AYSV clusters together with members initially found in Europe and Asia that belong to the so-called Eurasian clade. A similar phylogenetic analysis using AYSV NSM revealed 88.3 % identity to that of TYRV, and supports the relationship of AYSV to the Eurasian clade (Figure 3).

The complete nucleotide sequence of AYSV M RNA consists of 4797 nt, and contains a 5'- (62 nt) and 3'-UTR (51 nt) that is able to form a panhandle structure due to sequence complementarity. The UTRs show almost entire sequence complementarity for 29 nt with only 3 mismatches (5'- AGAGCAAUCCGGUGCAANAAUCAANUNU-3'). The vcORF starts with an AUG at nucleotide position 63 and terminates with an UAA codon at position 989, coding for the NSM protein with a predicted molecular mass of 34.2 kDa. The AYSV NSM protein shows conserved motifs that belong to the 30K movement protein superfamily [52, 53], including D_{155}, G_{211} and P/D-L-X (amino acid residues D_{106}S_{103}L_{104}), and motifs for a phospholipase A2 catalytic site and PLA2 (amino acid residues CMQLNLTS_{197-204}).

The vcORF codes for the glycoprotein precursor and starts with an AUG codon at nucleotide position 4746 and terminates with UGA at nucleotide position 1300. The M RNA encoding v- and vc-ORFs flank an IGR of 310 nt that runs from nucleotide position 990 to 1299 and, like the S RNA, consists of nucleotide stretches highly rich in A- and U residues. Folding predictions of the IGR sequence reveals the formation of a stable hairpin structure (data not shown).

Using BLASTp, the highest identity of AYSV NSM was found with TYRV NSM (88.7%) whereas its glycoprotein precursor (GP) showed highest identity with that of TYRV GP (81.0%). Phylogenetic analysis based on the NSM protein and the glycoprotein precursor support a classification of AYSV within the Eurasian clade (Figure 3).

The surface glycoprotein of AYSV was analyzed and compared to those of other Eurasian orthotospovirus species for the presence of N-, O-, N-glycosylation, peptide cleavage sites and transmembrane domains (Figure 4). Two putative N-glycosylation sites, Asn_{319} and Asn_{575} and eight O-glycosylation sites i.e. Thr_{80}, Thr_{82}, Thr_{84}, Thr_{99}, Thr_{100}, Thr_{101}, Thr_{102} and Thr_{105} were predicted. Two signal peptide sequences with predicted cleavage sites were found at amino acid positions Leu_{23} and Leu_{462}. Five (hydrophobic) transmembrane domains were found in the G N-GC precursor. Both of the predicted cleavage sites located within the first and fourth transmembrane domains, whereas none of the predicted glycosylation sites mapped within transmembrane domains (Figure 4). A comparison to the G N-GC precursor

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**Fig. 2** Analysis of the major structural N protein from purified AYSV RNPs and serology. Purified AYSV RNPs were comparatively analyzed to those of TSWV and INSV on SDS-PAGE (left panel). Proteins were stained with Coomassie Brilliant Blue. Size marker is indicated at the left side of the gel. Serological differentiation between AYSV and five orthotospoviruses in DAS-ELISA (right panel) was performed using polyclonal antisera raised against respective N proteins and the extracts from infected plants as antigen source.
of other Eurasian orthotospovirus species revealed that the AYSV precursor contains the lowest number of O-glycosylation sites and the highest number of O-glycosylation sites. Based on the predicted cleavage sites, the AYSV G_N-G_C precursor is processed into the mature G_N and G_C glycoproteins of 50.3 and 77.4 kDa, respectively (Figure 4). The AYSV G_N-G_C precursor does not contain a Arg-Gly-Asp (RGD) sequence, a motif earlier found in the TSWV precursor and putatively involved in virus cell attachment [9].

The entire sequence of AYSV L RNA consists of 8865 nt with a 5' terminal (214 nt) and 3' terminal (32 nt) UTR that contain the highly conserved 8 terminal nucleotides present in all orthotospoviral RNA segments. AYSV terminal sequences show almost entire base pair complementarity for the first 16 nt with only one base pair mismatch (5'-AGA-GCA AUCNAGCAAC-3').

The L RNA contains a single ORF of 8619 nt in the v-c strand and coding for the putative viral RNA-dependent RNA polymerase (RdRp, L protein) of 2873 aa with a predicted molecular mass of 316.0 kDa. The vcORF starts with an AUG at nucleotide position 8833 and terminates with an UAA stop codon at nucleotide position 212. The L gene of AYSV shares highest aa (87.3 %) identity with the one from TYRV. Analysis of the AYSV L protein revealed the presence of all conserved motifs earlier reported and indicative for the RdRp [54, 55]. The highly conserved region, shared among all characterized orthotospoviral L proteins, is found at aa position 1,275–1,559 of the AYSV L protein. This region contains the essential motifs of viral RdRp proteins from negative strand RNA viruses and includes motif A (aa 1,365–1,378), motif B (aa 1,452–1,471), motif C (aa 1,490–1,499), motif D (aa 1,534–1,544), motif E (aa 1,547–1,559), and motif F (aa 1,275–1,301) [54, 55]. The AYSV RdRp motif sequences are 100% identical to those of TYRV. Phylogenetic analysis of the RdRp shows a clustering of AYSV with TYRV, PolRSV, hippeastrum chlorotic ringspot virus (HCRV) and IYSV (Figure 3).

A summary and comparative analysis of the AYSV L, M and S RNA sequence to those of other Eurasian species is shown in Table 2. While the L RNA sequence has not been elucidated for all Eurasian species, the structural features of the M and S RNA of AYSV show highest similarity to those from PolRSV, the closest related species, with the exception of a differently sized IGR within the S segment (AYSV-IGR478 vs. PolRSV-IGR183).

The complete L, M and S RNA sequences of AYSV are available under GenBank accession no. MF469033, MF469034 and MF469035 respectively.

Vector transmission

Healthy *E. sonchifolia* plants exposed to *T. tabaci*, that in their larval stages were given access to AYSV-infected plants, revealed chlorotic spots similar to those observed earlier during the host range analyses (Table 1). The presence of AYSV was confirmed and successful transmission by viruliferous thrips was demonstrated by RT-PCR amplification of (partial) AYSV N gene sequences from infected leaf material and subsequent sequence analysis.

Discussion

Here a tentative new orthotospovirus species has been identified and characterized from alstroemeria, and for which the name alstroemeria yellow spot virus (AYSV) has been proposed. The virus causes yellow spots on alstroemeria, has a narrow experimental host range limited to a few dicots and is transmitted by the vector *T. tabaci*. A polyclonal antiserum has been raised against AYSV RNPs and only reacts positively with the homologous virus and not with any other established orthotospovirus species tested. The nucleotide sequence of the genomic L, M and S RNA segments has been elucidated (8619, 4797 and 2734 nt, respectively) and contains 5 ORFs coding for six mature proteins i.e., the L protein (viral RdRp, 316 kDa), the cell-to-cell movement protein (NS_M, 34.2 kDa), the precursor (127.7 kDa) to the glycoproteins G_N and G_C, the suppressor of RNA silencing (NS_p, 50.1 kDa) and the nucleocapsid protein (N, 30.1 kDa). The AYSV sequence shows all structural features typical for orthotospovirus RNA genomes with highest homology between its proteins and those from TYRV. Phylogenetic analysis indicates that AYSV, together with PolRSV, TYRV, HCRV and IYSV, belongs to a growing clade of...
orthotospoviruses for which the geographical origin seems to be based in the Middle-East [46]. Although the origin of the virus-infected alstroemeria plant could not be indicated with certainty, it was provided by a grower that at that time had simultaneously received shipments of alstroemeria plants from Colombia and Iran. Considering the relation of AYSV to the orthotospoviruses from the Middle-East cluster makes it most likely that Iran is the site of origin of the collected sample.

With the identification of this new orthotospovirus species, the family of Tospoviridae is still expanding and has currently reached the number of 30 tentative and established species ([2] and, this report). While this new species has only a small (experimental) host range, its natural host species so far is limited to alstroemeria. With the availability of a specific polyclonal antiserum to AYSV, field surveys could help to identify other natural (cultivated and non-cultivated) plant hosts of this virus. Considering that nothing is known on the prevalence and distribution of AYSV in agricultural and horticultural crops and ornamentals these surveys will also indicate the economic importance of this virus.

During this study a combination of conventional and NGS sequencing approaches has helped to elucidate the entire AYSV genome sequence. The speed and complete data sets

**Table 2** Comparison of structural features of the L, M and S RNA segments of AYSV to those of other Eurasian orthotospoviruses

| Virus acronym | AYSV | PolRSV | TYRV | IYSV | HCRV |
|---------------|------|--------|------|------|------|
| S RNA*        | 2,734| 2,484  | 3,061| 3,105| 2,744|
| 5’ UTR*       | 72   | 72     | 71   | 70   | 73   |
| NSs**         | 443  | 443    | 443  | 443  | 445  |
| IGR*          | 440  | 183    | 762  | 811  | 437  |
| N**           | 272  | 274    | 274  | 273  | 274  |
| 3’ UTR*       | 71   | 72     | 71   | 70   | 71   |
| M RNA*        | 4,797| 4,710  | 4,786| 4,838| 4,741|
| 5’ UTR*       | 62   | 62     | 62   | 63   | 47   |
| NSm**         | 308  | 308    | 308  | 311  | 308  |
| IGR*          | 310  | 263    | 354  | 379  | 330  |
| Gc-Gc**       | 1,148| 1,135  | 1,130| 1,136| 1,129|
| 3’ UTR*       | 51   | 50     | 50   | 49   | 47   |
| L RNA         | 8865 | 8893   | 8877 | 8880 | 8908 |
| 5’ UTR*       | 211  | 230    | 223  | 225  | 253  |
| RdRp**        | 2,873| 2,876  | 2,873| 2,873| 2,873|
| 3’ UTR*       | 32   | 32     | 32   | 33   | 33   |

* Number of nucleotide residue; ** Number of amino acid residue
obtained by the NGS approach, combined with lowering costs, favor usage of NGS for the elucidation of viral RNA/DNA genomes. However, during this study and our earlier sequence analysis of chrysanthemum stem necrosis virus (CSNV) [51] we have observed that the NGS approach may not always provide the entire 5’ and 3’ terminal sequences and sufficient reads to span the entire A- and U-rich intergenic region. Assembly of the latter reads into one contig is also hampered by the fact that they are relatively short and subsequently fail to properly assemble due to sequence stretches rich in A- and U-residues. Verification of the termini and intergenic region thus still needs to be done by conventional RT-PCR / 5’ and 3’ RACE amplification and Sanger sequence analysis of the amplicons.

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Compliance with ethical standards

Conflict of interest The authors declare to have no conflict of interest.

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

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