Infectious clones of Tomato leaf curl Palampur virus with a defective DNA B and their pseudo-recombination with Tomato leaf curl New Delhi virus

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Abstract

Background: Tomato leaf curl Palampur virus (ToLCPMV) is a bipartite begomovirus which has been reported from India and Iran but infectious clones have not been obtained. We have previously shown the association of Zucchini yellow mosaic virus (ZYMV), a potyvirus, with severe leaf curl disease of muskmelon in Pakistan. However, the severity of symptoms in the field and yield losses led us to believe that some other agent, such as a begomovirus, could be associated with the disease.

Results: A bipartite begomovirus associated with a severe yellow leaf curl disease on muskmelon in Pakistan has been characterized. Analysis of the complete nucleotide sequence of the DNA A and DNA B components of the begomovirus showed that it has the highest DNA sequence identity with ToLCPMV. However, the gene encoding the nuclear shuttle protein (NSP) was truncated in comparison to previously characterised isolates. Agrobacterium-mediated inoculation of Nicotiana benthamiana with the ToLCPMV clones obtained here did not result in symptoms. However, inoculation of plants with the DNA A component of ToLCPMV and the DNA B component of Tomato leaf curl New Delhi virus (ToLCNDV) lead to systemic infection with leaf curl symptoms. This suggested that the lack of infectivity of the ToLCPMV clones was due to the defect in DNA B. The DNA B of ToLCPMV was able to move systemically when inoculated with DNA A of the either virus. Agro-infiltration of muskmelon with the DNA A and DNA B components of ToLCPMV did not lead to symptomatic infection whereas inoculation with the DNA A with the DNA B of ToLCNDV resulted in a hypersensitive response (HR) along the veins. Additionally, agro-infiltration of muskmelon with a construct for the expression of the NSP gene of ToLCNDV under the control of the cauliflower mosaic virus 35S promoter induced a HR, suggesting that this is the gene causing the HR.

Conclusions: Both ToLCPMV and ZYMV are associated with muskmelon leaf curl disease in Pakistan. However, the ToLCPMV variant identified in association with ZYMV has a defective NSP. The results suggest that a variant with a defective NSP may have been selected for in muskmelon, as this protein is an avirulence determinant in this species, and possibly that infection requires the synergistic interaction with ZYMV.

Background

Whitefly-transmitted geminiviruses have emerged as major pathogens of food and fiber crops throughout the world, particularly in tropical and sub-tropical regions [1]. Viruses of the family Geminiviridae have circular single-stranded (ss) DNA genomes, encapsidated in characteristic twinned isometric particles. These viruses are classified into four genera, Mastrevirus, Curtovirus, Topocuvirus and Begomovirus [2] depending upon their insect vector, host range and genome organization. Begomoviruses are exclusively transmitted by the whitefly Bemisia tabaci and infect only dicotyledonous plants. They may be monopartite or bipartite (having genomes consisting of one or two circular single-stranded DNA molecules) [3]. Since the identification of the betasatellites [4], it has become apparent that the majority of begomoviruses in the Old World are monopartite and associate with betasatellites, asymptom modulating ssDNA satellite [5-7].

Several factors, including multiple infections, recombination and interspecific synergism, appear to be the major cause of the increased importance of begomoviruses to
agriculture in recent years. Intriguingly, begomovirus-betasatellite complexes do not appear to be a problem in cucurbits; only a single defective betasatellite isolated from a cucurbit has been reported so far [8]. The DNA A and DNA B components of begomoviruses encode genes both in the virion and complementary-sense orientations. Old World begomoviruses encode four open reading frames (ORFs) in the complementary-sense that are involved in viral DNA replication and control of gene expression, while two ORFs in the virion-sense are involved in virus encapsidation and movement. The two proteins encoded by the DNA B component are the nuclear shuttle protein (NSP) and the movement protein (MP) that are involved in nuclear transport and cell-to-cell movement of viral DNA, respectively [9].

Both watermelon and muskmelon form an important part of diet in Pakistan, particularly in early summer. These crops fit nicely in the cropping pattern where vegetables are grown in the winter months when land is available before the start of cotton growing season. However, the traditional muskmelon-growing areas have recently been hit by a severe viral disease. We have previously shown that a potyvirus, *Zucchini yellow mosaic virus* (ZYMV), is associated with the disease of muskmelon [10]. Here, we have characterized a bipartite begomovirus associated with severe leaf curl disease on muskmelon in Pakistan.

Methods

Collection of samples and DNA extraction

Samples were collected as described earlier [10]. Total DNA was isolated from leaf samples using CTAB method [11].

Detection and cloning of virus components

PCR-mediated detection of begomoviruses in DNA samples extracted from symptomatic leaves was done by using two sets of universal primer pairs (Begomo1/Begomo2 and Begomo3/Begomo4; Table 1) designed for amplification of different length fragments [12]. Full-length PCR product of DNA A was amplified by using primers Melonfor and Melonrev (Table 1). Oligonucleotide primers used for amplification of complete DNA B were TLCVB F and TLCVB R (Table 1). PCR was performed under standard conditions as described previously [13]. Primer pairs used for amplification of MP, NSP and full-length DNA B are also shown in Table 1. Samples were also tested for the presence of DNA B of begomoviruses using a PCR product (produced using primer pair MPF/MPR; Table 1), encompassing the MP gene of *Tomato leaf curl New Delhi virus* (ToLCNDV-IN[N:ND:Svr:92]) (acc. no U15017), as a probe for ToLCNDV DNA B [14]. The probe was prepared using a Hexalabeling DNA labeling kit (Fermentas) and [α-32P]dCTP (Amersham, UK). Hybridization was carried out overnight at 65°C in a hybridization oven. Post-hybridization washes were conducted at medium-stringency.

Sequencing and sequence analysis

The complete nucleotide sequences of clones were determined by dideoxynucleotide chain-termination sequencing using the PCR-based BIG DYE kit (Perkin-Elmer Cetus) and specific internal primers. Sequencing products were resolved commercially (Macrogen, Korea). Sequence information was stored, assembled and analyzed using the Lasergene sequence analysis package (DNASTar Inc., Madison, WI, USA) running on an IBM-compatible PC. Phylogenetic analyses were conducted on matrices of aligned sequences using the neighbour-joining and bootstrap options of Phylip (ver. 3.5c) running on an IBM-compatible personal computer. Sequence alignments were produced using CLUSTAL X [15]. Phylogenetic dendrograms were viewed, manipulated and printed using Treeview [16].

Agroinoculation of cloned DNA

Partial direct repeat constructs for the *Agrobacterium*-mediated inoculation of the full-length DNA A and DNA B clones were produced in the binary vector pGreen0029 [17]. An approximately 1680 bp fragment of the DNA A and an approximately 823 bp fragment of the DNA B clones were released from the full-length pTZ57/RT clones by digestion with *Kpn*I and *Bam*HI or *Hind*III and *Mlu*I, respectively, and ligated into suitably digested pGreen0029 vector. The resulting pGreen0029 constructs were then linearised using *Kpn*I and *Hind*III, respectively, and the full-length fragments (release from the pTZ57/RT clones by digestion with *Kpn*I and *Hind*III, respectively) ligated into these to yield the partial, direct repeat constructs. The production of constructs in the binary vector pBin19 for inoculation of ToLCNDV has been described previously [18]. These binary vector constructs were transformed into *Agrobacterium tumefaciens* GV1301 and inoculated to plants by infiltration as described previously [18].

Results

A bipartite begomovirus is associated with severe disease on melon

As described earlier, a very severe yellow leaf curl disease was observed in several districts of central and southern Punjab, including Vehari, Sahiwal and Khanewal, and was shown to be associated with *Zucchini yellow mosaic virus* (ZYMV) [10]. Total DNA isolated from symptomatic plants was used as template in PCR with universal primers designed for the amplification of the partial sequences of begomovirus DNA A. The amplified product was cloned and sequenced and it showed the highest levels of nucleotide sequence identity to isolates of
**Table 1** Primers used to amplify partial and full length genome sequences of begomovirus components

| Primer   | Sequence                        | Expected fragment size (kb) | Target Component/gene |
|----------|---------------------------------|-----------------------------|-----------------------|
| Begomo1  | CCGTGCTGCTGCCCCCATTGTCCGCAGTAC | 1.1                         | DNA A                 |
| Begomo2  | CTGACCAACCTGGGTTACCCGACAGGG    | 1.1                         | DNA A                 |
| Begomo3  | GTTCCTCTGGCTGGAATTCTCAGGCGTGG  | 1.7                         | DNA A                 |
| Begomo4  | TTTTTGGGCAGGCAGCAATGGGAGGAGACA | 1.7                         | DNA A                 |
| Melonfor | GGTACCTAAGGACCTGGGTTCTG        | 2.8                         | DNA A                 |
| Melonrev | GGTACCTGATATGCTGTTGTTATAG      | 2.8                         | DNA A                 |
| TLCVB F  | GCTAAGCTTCTGCTGAACTGGAGGAAA   | 2.7                         | DNA B                 |
| TLCVB R  | CAGAGCTAGGCAGTGAGGAGTATAG     | 2.7                         | DNA B                 |
| MLU F    | ATACCGCTAGGGAAATCTTGTGAGAACAC | 2.7                         | DNA B                 |
| MLU R    | CTTACCGCTATATTGTGTGAGATT      | 2.7                         | DNA B                 |
| Pst F    | GCTCTGAGACATCTTTGTGAGGG       | 2.7                         | DNA B                 |
| Pst R    | CTGCTGCAAGTTACCTTGTATATAG    | 2.7                         | DNA B                 |
| MP F     | CACCATGGCAAAGAATATGCAGATATGGG |                | MP                   |
| MP R     | AAGGATCTCTCTATTGTTCAATTATTGG  |                | MP                   |
| NSF F    | GGCATCGAGGGTTTCCCGTGGCACAC   |                | NSP                  |
| NSF R    | GGGTGCACATTACCCAATTTTTGAATT  |                | NSP                  |

**Tomato leaf curl Palampur virus** (ToLCPMV), a bipartite begomovirus. To confirm the bipartite nature of virus from muskmelon, a PCR product encompassing the movement protein (MP) gene of ToLCNDV was used as a specific probe for Southern hybridisation. All the muskmelon samples found positive for DNA A were also found positive for DNA B (results not shown). Samples were similarly examined for the presence of begomoviruses by Southern blot hybridization using the DNA A of ToLCNDV as a general probe for begomoviruses. The probe hybridized with all sixty samples collected from symptomatic plants that were found positive by PCR.

**Cloning and complete nucleotide sequence of DNA A**

Based upon the partial sequences obtained, back-to-back primers for amplification of full-length DNA A (MelF and MelR; Table 1) were designed to the coat protein region on a naturally present, unique KpnI restriction site. A full-length product (approximately 2.7 kb) was obtained and cloned in pTZ57RT. Restriction of the clone with KpnI resulted in a single band on the gel whereas restriction of clone with Smal linearized the clone and a 5.6 kb band appeared on the gel. The potential full-length clone was sequenced in both orientations. A sequence similarity search (Blast) was performed by comparing the sequence to other begomovirus sequences in the database. The results showed that the sequence has a high level of identity to begomoviruses from Asia. The DNA A comprised of 2756 bp and showed the highest levels of nucleotide sequence identity (97%) to a ToLCPMV isolate from India (accession no. AM884015) and 96% identity to an isolate from Iran (accession no. FJ660444). The sequence identity with ToLCPMV isolates ranged from 96-97% whereas to isolates of ToLCNDV the highest sequence identity (86.7%) was with an isolate from Rahim Yar Khan, Pakistan (accession no. DQ116885). On the basis of DNA A sequence identity, it is clear that the muskmelon infecting begomovirus in Pakistan is an isolate of ToLCPMV, a begomovirus species previously reported infecting tomato in India and Iran [19,20]. Sequence analysis showed that it is a typical Old World begomovirus and encodes four open reading frames (Rep, REn, TrAP and AC4) in the complementary-sense and two (V2 and coat protein) in the virion-sense (Table 2), separated by an intergenic region (IR) (Figure 1A). The IR contains the conserved nonanucleotide motif (TAATATTAC), forming part of the loop of a predicted stem-loop structure found in all geminiviruses, a TATA motif (a sequence found in the promoter region and binding site for transcription factors) [21] (2664-2667 nt) and iteron sequences (GGTGTC) 5` of the TATA motif [21,22]. The proteins predicted to be encoded by the DNA A component were of the sizes expected for begomovirus DNA A encoded proteins (Table 2) [14].

**Cloning and sequence analysis of DNA B**

The partial sequences of DNA B obtained were used to design back-to-back primers (TLCVB F and TLCVB R; Table 1) for the amplification of the full-length DNA B within the sequence of the movement protein (MP) gene. The desired sized fragment (2.7 kb) of DNA B was cloned in pTZ57RT. The clone was fully sequenced and shown to consist of 2724 bp. The DNA B showed 98% nucleotide sequence identity to an isolate of ToLCPMV recorded from India (accession no. AM992534) whereas the identity
to the ToLCPMV isolate from Iran was 90% (accession no. FJ660442). Among the DNA B components of ToLCNDV available in the databases the highest levels of identity (82%) were to an isolate from India obtained from pumpkin (accession no. AM286435). In ORFs, NSP showed 99% sequence identity with ToLCPMV (accession no. AM992534) and it ranged from 90-99%. Whereas it was 73.3% identical to ToLCNDV (accession no AY150305). The maximum sequence identity to ToLCPMV isolate originating from India was 99% but to the isolates originating from Iran it was 95%. Overall among ToLCPMV isolates, sequence identity ranged from 92-99%. ToLCPMV MP was 86% identical to MP of ToLCNDV (accession no AY150304). Thus, DNA B might be a recombinant molecule where the MP is derived from DNA B of ToLCNDV while the intergenic region and NSP are derived from yet an unknown virus. The intergenic region of ToLCPMV shows 86% nucleotide sequence identity to Indian isolate (accession no. AM992534) whereas the identity to Iranian isolate is 76% (accession no. FJ660442). NSP has a size of 555 nucleotides (184 amino acids) and it is truncated at the N-terminus, since first 84 amino acids are missing in comparison to the ToLCNDV sequences available in the database (Figure 1B). The MP has a predicted size of 884 bases (294 amino acids) (Table: 2). Non-coding IR sequences of DNA B are distinct from already published sequences. The intergenic regions of DNA A and DNA B show only 38.2% identity but share a 34-bp potential stem-loop forming region (GGCCATTCGTATAATATTACCGAATGGCCGCGGT). This sequence has the conserved nonanucleotide sequence (TAATATTAC). The iterated elements (iterons) were close to a TATA box in the common region and were identified as GGTGTC (Figure 2) being the same as those identified in DNA A of ToLCPMV and ToLCNDV [22].

Efforts were made to identify a DNA B in muskmelon having no truncation of the NSP gene. Three set of primers were designed in the NSP and non-coding sequences of DNA B of ToLCNDV (Table: 1). None of the primer sets was able to amplify a product of approximately 2.8 kb. The ability of these sets of primers to amplify full-length DNA B was confirmed by the use of these primers on tomato samples infected experimentally with ToLCNDV. All sets of primers

| Component | Gene | Position of start codon | Position of stop codon | Size (number of nucleotides) | Predicted size of encoded product (kDa) |
|-----------|------|-------------------------|------------------------|-------------------------------|----------------------------------------|
| DNA A     | AV2  | 120                     | 458                    | 338                          | 13.2                                   |
|           | CP   | 280                     | 1050                   | 771                          | 29.8                                   |
|           | REn  | 1457                    | 1047                   | 411                          | 16.0                                   |
|           | TrAp | 1596                    | 1177                   | 410                          | 15.9                                   |
|           | Rep  | 2602                    | 1499                   | 1104                         | 41.6                                   |
|           | AC4  | 2445                    | 2269                   | 177                          | 6.7                                    |
| DNA B     | NSP  | 681                     | 1235                   | 555                          | 20.9                                   |
|           | MP   | 2146                    | 1301                   | 884                          | 32.0                                   |

Figure 1 Genome organization of ToLCPMV DNA A and DNA B. The positions and orientations of predicted ORFs are shown with arrows. The common region (CR) of each component of the virus is indicated by a grey box. The position of the stem-loop structure, containing the conserved nonanucleotide loop sequence (TAATATTAC), is indicated by a black circle.
amplified products of the expected size. To further rule out the possible presence of other DNA B, phi29 polymerase was used which has the ability to amplify circular molecules. The amplified product was restricted with 
PstI endonuclease (for which there are two restriction sites in DNA A) yielding two bands of the expected sizes (969 bases and 1786 bases). The DNA B of ToLCPMV has two 
PstI restriction sites that are separated by 62 bases. The restriction analysis suggested the presence of a single type of DNA B of ToLCPMV and the absence of DNA B of ToLCNDV (results not shown).

Phylogenetic analysis of ToLCPMV DNA A and DNA B
A phylogenetic analysis, based upon an alignment of the complete DNA A sequence of ToLCPMV with selected other begomoviruses is shown in Figure 3A. This analysis shows ToLCPMV to segregate with ToLCPMV isolates reported from India [19] and Iran [20] followed by Cucumber leaf curl virus (CuLCuV-[BD:06](acc. no. EF450316); a begomovirus recently identified in Bangladesh). These two viruses group with, and are basal to, the ToLCNDV isolates but distinct from Squash leaf curl China virus (SLCCNV) isolates. This is well supported by bootstrapping. A phylogenetic analysis based upon an alignment of the complete DNA B sequence of ToLCPMV with selected other begomoviruses is shown in Figure 3B. In this analysis ToLCPMV is again basal to the ToLCNDV isolates which form a clade distinct from the SLCCNV isolates. It is also noteworthy that the DNA B of Tomato leaf curl Gujarat virus (ToLCGV-[IN:Var:01]) is included in this analysis and shows it to have a ToLCNDV-like DNA B, as discussed later. The position of ToLCPMV in the tree is well supported by bootstrapping.

Infectivity of ToLCPMV clones from muskmelon
The infectivity of ToLCPMV clones was assessed by Agrobacterium-mediated inoculation to Nicotiana benthamiana (Figure 4 and Table: 3). Plants inoculated with ToLCPMV DNA A alone (Figure 4A) or ToLCPMV DNA A and DNA B did not develop symptoms (Figure 4B), even 2 months post-inoculation. However, diagnostic PCR showed the presence of both ToLCPMV DNA A and DNA B in systemic leaves. In contrast, plants inoculated with ToLCNDV DNA A and DNA B developed upward leaf curl symptoms and vein swelling within 14 days of inoculation (Figure 4E). These symptoms are typical of ToLCNDV in N. benthamiana [18]. The inability of ToLCPMV clones to cause disease symptoms in N. benthamiana indicates that they are competent for systemic infection but unable to produce disease symptoms.

ToLCNDV DNA B will support symptomatic infection of ToLCPMV DNA A
To ascertain which of the components was responsible for the lack of infectivity, the ToLCPMV clones were inoculated as pseudo-recombinants with the components of ToLCNDV. Inoculation of twenty N. benthamiana plants with ToLCNDV DNA A and ToLCPMV DNA B did not develop begomovirus symptoms (Figure 4F) but
seven plants were found positive in PCR for the presence of DNA B. In contrast, all the plants inoculated with ToLCPMV DNA A and ToLCNDV DNA B developed symptoms typical of ToLCNDV including upward leaf curl and vein swelling symptoms 14 days post-inoculation (Table 3; Figure 4C). These results demonstrate that the defect(s) in symptom induction by ToLCPMV clones resides on DNA B. The earlier sequence analyses indicated that ToLCPMV DNA B contains a truncation of the Ns gene due to a premature stop codon. This is the likely cause of the lack of ability of ToLCPMV DNA B to support symptoms development. The ability of ToLCPMV to trans-replicate and to establish a systemic infection with ToLCNDV also indicates that the Rep encoded by ToLCPMV DNA A is able to interact with the origin of replication of ToLCNDV. The sequence analysis showed that the predicted iteron sequences of ToLCPMV are the same as iteron of ToLCNDV [14] and these elements are crucial in the interactions required for Rep to initiate rolling-circle replication [24].

Inoculation of ToLCPMV DNA A with ToLCNDV DNA B induces a hypersensitive response in muskmelon

Our data on infectivity in *N. benthamiana* showed that ToLCPMV DNA A in association of DNA B of ToLCNDV induced typical disease symptoms. The same combination was tried in muskmelon to establish infectivity in this host. Inoculated as well as systemic leaves showed hypersensitive response (Figure 5A). In systemic leaves this appeared as leaves necrosis near the veins (Figure 5B). We have previously shown that the expression of NSP of ToLCNDV in transgenic plants induced hypersensitive response which was dependant on growth stage of plant [25]. We agroinfiltrated NSP of ToLCNDV cloned under 35S promoter while MP of ToLCNDV cloned under 35S promoter was used as control (Figure 6B). No effect on infiltration of 35S driven MP was found but the expression of NSP developed necrotic lesions that spread out of infiltration point (Figure 6B).

**Discussion**

Phytopathogenic viruses are major constraints to agricultural productivity throughout the world. It is an unfortunate fact that Pakistan, in common with all other countries of southern Asia, is home to members of virtually all taxonomic groups of plant-infecting viruses [26-31] which, at least in part, explains the low agricultural productivity in the country. Prime amongst these viruses are the begomoviruses that have appeared as dominant pathogens on several crops. We show here that...
muskmelon samples previously found positive for ZYMV are co-infected with a begomovirus. The complete nucleotide sequence of DNA A of begomoviruses is sufficient for identification and classification of begomoviruses. Viruses that share more than 90% sequence identity are considered as strain of the same virus while DNA sequence identity below 89% is considered as new species [32]. Based on this threshold value, the begomovirus identified in muskmelon is an isolate of ToLCNDV. The high levels of nucleotide sequence similarity, of both DNA A (86%) and DNA B (73%), between ToLCPMV and ToLCNDV indicate that they have a recent common ancestor and that ToLCPMV represents a lineage that has diverged from ToLCNDV. ToLCNDV has frequently been identified in cucurbits [33-38]. However, Koch’s postulates for ToLCNDV causing diseases of cucurbits have not been satisfied. ToLCNDV is a cosmopolitan species that occurs throughout southern Asia, having been reported from Pakistan, India, Thailand and Taiwan [39-43]. However, since no efforts were made in these reports to inoculate clones on cucurbits, it is difficult to know whether an RNA virus was co-infecting these cucurbits.

The genome organization of ToLCPMV is the typical of Old World begomoviruses (having an AV2 gene) and phylogenetic analysis (Figure 3) show it and ToLCNDV to be most similar to (to group with) other begomoviruses with their center of diversity and origins lying in southern Asia [39,44,45-47]. Recombination contributes to the genetic diversification of geminivirus populations and has been related to the emergence of some serious plant diseases, as discussed earlier [44-48]. The prerequisite for recombination to occur is co-infection of the same cell of a host plant [49]. ToLCPMV DNA A has high levels of sequence identity to ToLCNDV-IN [PK:Kha:Chi:04] (accession number DQ116880) in the virion-sense (including the intergenic region) and to ToLCNDV-IN[IN:ND:AVT1] (AY428769) in the complementary-sense, indicating that this molecule is an intraspecific recombinant. The geminiviruses evolve/adapt by a number of mechanisms. Contrary to expectations, geminiviruses have nucleotide substitution rates that are comparable to those of RNA viruses [50]. Such high substitution rates would not be expected from a virus that utilizes host-encoded DNA polymerases with error correction. Additionally, recombination and component exchange (known as pseudo-recombination) are the major processes of geminivirus evolution [46,51-54].

A prime example of this is the ongoing pandemic of cassava mosaic disease that originated in northern Uganda, spread across eastern Africa and continues to spread throughout central and western Africa. The severe cassava mosaic disease of the pandemic is attributed to a recombinant strain of East African cassava mosaic virus known as the “Uganda Variant” [44].

Like the DNA A component, the DNA B component of ToLCPMV is a recombinant. The complementary-sense sequences (containing the MP gene) originate from ToLCNDV (showing the highest sequence identity with isolate ToLCNDV-IN [PK:Isl:T1:8:00; accession no. AY150304]) while the intergenic region and virion-sense sequences are derived from an as yet undiscovered (or extinct) virus; having no high sequence similarity to any sequences in the databases. The DNA B of ToLCPMV is a mutant with a truncation of the NSP gene. It is unable to support symptoms development when inoculated to N. benthamiana in the presence of ToLCPMV DNA A. Mutational analysis suggests that the N-terminal
sequences of NSP are involved in nuclear localization while C-terminal sequences are required for interaction with the MP [55]. Thus, truncation of ToLCPMV NSP at the N-terminus likely abolished the nuclear localization of the protein resulting in a defective molecule which cannot support infection or the protein is not produced at all since it is prematurely truncated. The NSP of ToLCNDV virus is a pathogenicity determinant where N-terminal sequences of protein are required for pathogenicity [25]. PCR amplification with universal and specific DNA B primers failed to show the presence of an intact DNA B in muskmelon samples, indicating that this defective molecule is the only DNA B present in muskmelon. We propose that truncated DNA B was selected to show the presence of an intact DNA B in muskmelon samples, indicating that this defective molecule is the only DNA B present in muskmelon. A recent example of a begomovirus associated with resistance breakdown in cotton lacked an intact TrAP, probably because NSP was recognized by host defense system. Thus, DNA B with truncated NSP was maintained on muskmelon. A recent example of a begomovirus associated with resistance breakdown in cotton lacked an intact TrAP, probably to avoid host defense targeting the TrAP [56].

The integrity of the genomes of bipartite begomoviruses is maintained by them having compatible Rep binding sequences in the CR. Thus, the DNA A-encoded Rep is able to recognize and initiate the replication of both components [57-59,23]. Iteron sequences are usually species-specific, thus in most cases, the Rep of one virus would not be expected to recognize the iteron sequences of a distinct species [60-62]. The iteron sequences of ToLCNDV have been identified experimentally [63]. ToLCNDV and ToLCPMV have the same Rep binding sites and this likely explains our experimental finding that ToLCPMV DNA A is able to trans-replicate ToLCNDV DNA B and initiate a productive, symptomatic systemic infection of plants. ToLCNDV is a part of viral complex consisting of several DNA As that exchange DNA Bs readily. Here evidence is provided that, in addition to ToLCNDV and Tomato leaf curl Gujarat virus [64], the complex also encompasses ToLCPMV. This ability to readily form genomic reassortments (pseudo-recombination) is probably an adaptation that gives the virus complex an evolutionary advantage. Since DNA B components are involved in movement in plants [65,66], and thus host range determination [67-69], the ability to readily interact with distinct DNA B components will undoubtedly allow the viruses of the complex to alter/adapt their host ranges to take advantage of new niches.

For bipartite begomoviruses DNA B is considered as an integral part of the genome (a genomic component), rather than a satellite, due to the presence of the conserved region (CR) which is present in both components. Satellites are defined as molecules which share no significant sequence similarity with their helper viruses but require them for replication and movement in host plant [70]. Although clearly a DNA B, the lack of appreciable sequence similarity between the DNA A and DNA B components of ToLCPMV, with the exception of the

| Inoculum                         | Infectivity (no. of plants showing symptoms/no. of plants inoculated) |
|----------------------------------|---------------------------------------------------------------------|
| ToLCPMV DNA A + ToLCPMV DNA B   | 0/20                                                                |
| ToLCPMV DNA A + ToLCNDV DNA B    | 20/20                                                               |
| ToLCNDV DNA A + ToLCPMV DNA B    | 0/20                                                                |
| ToLCNDV DNA A + ToLCNDV DNA B    | 20/20                                                               |
| ToLCPMV DNA A                     | 0/20                                                                |
| ToLCNDV DNA A                     | 0/20                                                                |

Table 3 Infectivity of ToLCPMV and ToLCNDV clones to N. benthamiana

![Figure 5](http://www.virologyj.com/content/8/1/173)

**Figure 5** Inoculation of muskmelon plant with ToLCPMV DNA A ToLCNDV DNA. Cell death along the veins of systemic leaves of a muskmelon plant agroinoculated with ToLPMV DNA A and DNA B (A). Necrotic lesions along the veins of systemic leaf of a muskmelon plant micrografted with leaf discs of a N. benthamiana plant infected with ToLCPMV DNA A and ToLCNDV DNA B by agroinoculation (B).
iteron sequences and nonanucleotide sequence, means that the ToLCPMV DNA B component could be deemed a satellite. This contention is strengthened by the ability of the DNA A to readily exchange the component, a feature in common with satellites such as DNA betasatellite. Likely, in this case, the ToLCPMV DNA B is maintained as a satellite (although the question remains as to whether the NSP gene is expressed). Possibly then, in muskmelon, the ToLCPMV DNA B is maintained as a poorly-functional satellite which is not required for the infection. A recent analysis of DNA A and DNA B sequences has suggested that the two components differ in their evolution and DNA B may be considered as a satellite [71]. Further investigation will be needed to address this question. A possible explanation is that NSP being a pathogenicity determinant is the target of host defense as has been demonstrated previously [25]. Since all plants where ToLCPMV is found were earlier reported to be infected with ZYMV [10], there is a possibility that the defect in NSP is complemented by ZYMV. The presence of this defective DNA B in many plants and in geographically distant fields of muskmelon, as well as in various cucurbit species, however, suggests that this molecule may play an important part. A satellite that provides no selective advantage to a virus would be expected to be lost quite rapidly.

The data presented here show that ToLCPMV is widespread in the region that includes Iran, Pakistan and India and therefore an emerging virus on several crops. Unfortunately, the earlier reports have not demonstrated infectivity analysis and therefore, it is not possible to assess whether DNA B cloned from India and Iran was infectious. The role of ToLCPMV in muskmelon leaf curl disease remains to be demonstrated. We suggest that dual infection of a begomovirus with a defective DNA B and ZYMV may be important for disease symptoms. The lack of availability of infectious clones of ZYMV precludes us satisfying Koch’s postulates for the disease in muskmelon. Our earlier results showed that mechanical inoculation of sap from infected muskmelon to healthy melon plants developed disease symptoms [10]. However, the possibility that begomovirus was also transmitted by sap inoculation cannot be ruled out. Our results show that ToLCPV is infectious to experimental hosts when inoculated with DNA B of ToLCNDV. Our recent results from cotton show that the begomovirus isolated from resistant cotton lack TrAP [56]. We hypothesized that begomoviruses may adopt to resistant host by truncation of an important gene. The maintenance of the defective DNA B in all field samples suggests a role for DNA B. It therefore appears that a synergistic interaction with ZYMV may have compensated for the defective NSP. We are trying to understand the possible synergistic interaction by cloning of helper component proteinase of ZYMV in a RNA virus vector such as PVX.

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Authors’ contributions
AHM performed the experiments. AHM, RWB and SM were involved in data analysis experimental design and wrote the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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