Detection and frequency of recombination in tomato-infecting begomoviruses of South and Southeast Asia

HC Prasanna* and Mathura Rai

Address: Indian Institute of Vegetable Research, P B 5002, P O-B H U, Varanasi, Uttar Pradesh, 221005, India
Email: HC Prasanna* - prasanahc@yahoo.com; Mathura Rai - mathura.rai@gmail.com
* Corresponding author

Abstract

Background: Tomato-infecting begomoviruses are widely distributed across the world and cause diseases of high economic impact on wide range of agriculturally important crops. Though recombination plays a pivotal role in diversification and evolution of these viruses, it is currently unknown whether there are differences in the number and quality of recombination events amongst different tomato-infecting begomovirus species. To examine this we sought to characterize the recombination events, estimate the frequency of recombination, and map recombination hotspots in tomato-infecting begomoviruses of South and Southeast Asia.

Results: Different methods used for recombination breakpoint analysis provided strong evidence for presence of recombination events in majority of the sequences analyzed. However, there was a clear evidence for absence or low Recombination events in viruses reported from North India. In addition, we provide evidence for non-random distribution of recombination events with the highest frequency of recombination being mapped in the portion of the N-terminal portion of Rep.

Conclusion: The variable recombination observed in these viruses signified that all begomoviruses are not equally prone to recombination. Distribution of recombination hotspots was found to be reliant on the relatedness of the genomic region involved in the exchange. Overall the frequency of phylogenetic violations and number of recombination events decreased with increasing parental sequence diversity. These findings provide valuable new information for understanding the diversity and evolution of tomato-infecting begomoviruses in Asia.

Background

Begomoviruses are an important group of whitefly (Bemisia tabaci) transmitted viruses in the family Geminiviridae. They inflict significant economic losses in many dicotyledonous crops including beans, cassava, cotton, melon, pepper, potato and tomato [1-7]. Tomato yellow leaf curl virus (TYLCV) and Tomato leaf curl virus (ToLCV) are the begomoviruses severely constraining tomato production in many tomato-growing regions of the world.

Begomovirus genomes are composed of either one (monopartite) or two (bipartite) single stranded DNA molecules ranging in size between 2500 and 2800 nucleotides [8]. Most TYLCV of the old world and almost all known new world begomoviruses viruses are bipartite with genomes comprising DNA A and DNA B molecules. Monopartite old world begomoviruses, which are now believed to be the predominant begomovirus form, have only a DNA-A like genome component. The virion-sense strand of DNA A encodes the viral coat protein (AV1, V1...
or cp) and, in old-world begomoviruses [9], an AV2 or V2 gene that is necessary for virus accumulation and symptom development [10]. The complementary-sense strand of DNA-A encodes genes responsible for viral replication (AC1, C1 or rep), replication enhancer (AC3, C3 or ren), regulation of gene expression (AC2, C2 or trap) and AC4 or C4 involved in host range determination, symptom determination, symptom severity, and virus movement [11-13]. The DNA B of bipartite begomoviruses encodes two proteins, BV1 (a nuclear shuttle protein or NS) and BC1 (a movement protein or MP) involved in intra- and inter-cellular movement within the plant [14].

Begomoviruses exhibit a great deal of geographic dependent but host-independent genomic variation [15-17]. Recombination, especially interspecific homologous recombination, is a key contributor to the genomic diversification and evolution of begomoviruses [17]. To date, many natural begomoviruses recombinants have been reported [17-20]. Although the biological significance of begomovirus recombination is not clearly understood, in many parts of the world epidemics associated with the emergence of recombinant begomoviruses have been reported. These include the devastating cassava mosaic disease epidemic caused by recombinant East African cassava mosaic viruses in Uganda and neighbouring countries [18,21], the currently emerging pathogenic recombinant, tomato yellow leaf curl Malaga virus, in Spain [22] and the cotton leaf curl disease epidemic in Pakistan caused by a species complex including a variety of mostly recombinant begomovirus species [23]. Besides the apparent importance of recombination in begomovirus evolution the marks that it has left on currently sampled begomovirus genome sequences also have major implications when we attempt to use these sequences to infer the evolutionary histories of begomoviruses [24,25]. Consequently, the detailed characterization of recombination amongst tomato-infecting begomoviruses is a prerequisite for understanding how these important pathogens are evolving.

Although a few specific recombination events have been described so far in tomato-infecting begomoviruses [26-29], a full accounting of recombinants, recombination breakpoints and recombination hotspots in tomato begomovirus species and strains is lacking. For example, it is currently unknown whether there are differences in the number and quality of recombination events that are occurring among different tomato infecting begomovirus species. It is also currently unknown whether sequences in particular parts of the begomovirus genomes are more or less exchangeable between different species than sequences in other parts of these genomes. Such variations in recombination frequencies and patterns have been clearly observed in RNA viruses [30]. In this study we employ a variety of recombination analysis methods to characterize recombination in South and Southeast Asian tomato-infecting begomoviruses. We map recombination hotspots and provide evidence that not all tomato-infecting begomoviruses are equally prone to recombination and that specific characteristic of particular recombination events are reliant on both the relatedness of the recombining viruses and the genomic region involved in sequence exchanges.

Results and discussion
In this study, we sought to characterise recombination in South and Southeast Asian viruses using a different approach to those used previously: (1) By studying a different set of viruses to those studied previously; (2) Making use of a combination of recombination analysis methods that are both powerful and have low false positive rates; (3) by mapping and estimating the frequency of recombination events in begomoviruses.

The neighbor-net analysis revealed clear evidence of phylogenetic conflicts within the analysed sequences (Fig. 1). Notably, every sequence represented within the tree was implicated as a potential recipient of horizontally acquired sequences at some time in its evolutionary past. Unsurprisingly, the PHI test strongly supported the presence of recombination in these sequences (p < 0.0001).

Different methods used for recombination breakpoint analysis also provided strong evidence for presence of past recombination events in most of the sequences analysed. For each of the 32 potential recombinant sequences identified, possible breakpoint positions, sequence fragments and parental genotypes are listed in Table 1. Tomato leaf curl virus strains including ToLCBV, ToLCBV-[Ban4] and ToLCBV-[Ban5] from Bangalore, south India appeared to be the most complex recombinants carrying evidence of seven and six recombination events respectively. On the opposite end of the spectrum, Tomato leaf curl virus strains including ToLCNDV-[Mld] and ToLCNDV-[Luc] from New Delhi, ToLCNDV-[Svr] [Jes] from Bangladesh, and TYLCNV-[Y38] from China each carried evidence of only a single recombination event. In addition, viruses from geographically well separated regions appeared to have recombined at some time in the past. For example, tomato leaf curl virus strains from Bangalore and Gujarat in India contained sequences closely resembling those found in a ToLCITWV isolate from Taiwan. Also, Chinese viruses contained fragments of sequence closely resembling those found in sequences sampled in Thailand, Taiwan, Bangladesh and South India. Further, we used the TtreeOrderScan method [31] to investigate the phylogenetic evidence for recombination in the sequence alignment. This analysis revealed major deviations in the branching order of sequences within
trees constructed from different portions of the multiple sequence alignment (Fig. 2). Frequent tree order changes were observed at the region of rep and AC4. Importantly, most of the viruses detected as recombinants in the breakpoint analysis exhibited deviations in their branching order indicating that they were most likely correctly identified as recombinants. In addition, the TreeOrderScan analysis also provided evidence for gene flow amongst viruses in geographically separated regions. For instance, sequences found in southern Indian viruses grouped with those found in Thailand and Bangladesh virus positions from 2335–2652. Thai viruses contained sequences resembling those of Chinese viruses between 300–490 and 590–2372, but Indian viruses between 2472–2743. The recombination observed between geographically separated species/strains probably represents older events as they presumably occurred before their present separation [19]. Movement of vectors and/or infected plant materials may also have contributed to the gene flow observed between these widely separated locations [32]. Alternatively, it is possible that current sampling of Asian begomovirus diversity is so sparse that we do not yet fully appreciate the geographical range of many of the species studied here.

Interestingly, our breakpoint analysis indicated that three north Indian viruses (ToLCNDV-[Pkt1/8], ToLCNDV-Syr and ToLCNDV-[Pkt5/6]) were not detectably recombinant and three other north Indian viruses namely ToLCNDV-Mld, ToLCNDV-[Luc] and ToLCNDV-[Luf] were simple recombinants with only evidence of a single detectable recombination event involving a virus resembling ToLCPV sampled in the Philippines. While TreeOrderScan analysis also revealed an absence of recombination in two north Indian viruses, ToLCNDV-[Pkt1/8] and ToLCNDV-[Luf] (indicated by a horizontal line across the graph in Fig. 2). In addition, there was no phylogenetic support for inter-group recombination event reported for ToLCNDV-[Luc]. Thus there appears to be no or few recombination events in viruses reported from North India, signifying that certain begomovirus species may not recombine as readily as others. There are a
Table 1: Breakpoint analysis of tomato-infecting begomoviruses and their putative parental sequences.

| Species/Strain | ORF      | Breakpoint   | Possible parent                      | Method*     |
|---------------|----------|--------------|--------------------------------------|-------------|
| ToLCNDV-Mld   | AC1      | 2119–2211    | Unknown                              | gc, rdp     |
| ToLCNDV-Svr  | AC1, AC4 | 1953–2507    | ToLCPV-[LB]                          | RDP, gc     |
| ToLCNDV-[Luc]| AC1, AC4 | 1960–2514    | ToLCPV                                | RDP, gc     |
| ToLCBV-[Ban4]| AV1, AV2 | 481–894      | ToLCKV                               | GC, rdp, mc |
|               | AC1, AC2, AC3 | 1185–1784 | Unknown                              | gc          |
|               | AC1      | 1793–1894    | Unknown                              | rdp         |
|               | AC1, AC4 | 2076–2632    | ToLCGV-[Vad]                         | RDP         |
| ToLCBV-[Ban5]| AV1, AV2 | 481–894      | ToLCKV                               | GC, rdp     |
|               | AC1, AC2, AC3 | 1185–1784 | Unknown                              | gc          |
|               | AC1      | 1793–1894    | Unknown                              | rdp, mc     |
|               | AC1, AC4 | 2144–2734    | ToLCBV-[Kol]                         | GC          |
| ToLCBV-[Kol] | AV1, AV2 | 481–894      | Unknown                              | rdp         |
|               | AC1, AC2, AC3 | 1185–1784 | Unknown                              | gc          |
|               | AC1      | 1793–1894    | Unknown                              | rdp         |
|               | AC1, AC4 | 2144–2727    | ToLCGV                               | RDP         |
| ToLCBV        | AV1, AV2 | 481–894      | ToLCKV                               | RDP, GC     |
|               | AC1, AC2, AC3 | 1185–1784 | Unknown                              | gc          |
|               | AC1      | 1793–1894    | Unknown                              | rdp         |
|               | AC1, AC4 | 2144–2734    | ToLCBV                               | GC          |
| ToLCGV-[Kel] | AV1, AV3 | 598–1214     | TYLCTHV-[Y72]                        | RDP, gc     |
|               | AC1, AC2, AC3 | 1183–1782 | Unknown                              | gc          |
|               | AC1, AC4 | 2160–2514    | ToLCGV                               | RDP, GC     |
| ToLCGV-[Var] | AV1, AV3 | 603–1219     | TYLCTHV-[Y72]                        | RDP, gc     |
|               | AC1, AC2, AC3 | 1183–1782 | Unknown                              | gc          |
|               | AC1, AC4 | 2165–2519    | ToLCGV                               | RDP, GC     |
| ToLCGV-[Vad] | AV1, AV3 | 598–1214     | TYLCTHV-[Y72]                        | RDP, gc     |
|               | AC1, AC4 | 2160–2514    | ToLCGV                               | RDP, GC     |
| TYLCCNV-Tb [Y36]| AV1, AV2 | 451–924    | ToLCBV                               | GC          |
| TYLCCNV-Tb [Y38]| AV1, AV2 | 451–924    | ToLCBV                               | GC          |
| TYLCCNV-[Y64]| AV1      | 525–927      | ToLCBV                               | RDP, gc     |
|               | AC1, AC4 | 451–924      | ToLCBV                               | GC          |
| TYLCCNV       | AV1, AV2 | 451–924      | ToLCBV                               | RDP, gc     |
|               | AC1, AC4 | 2051–2210    | ToLCBV                               | GC, RDP     |
| TYLCCNV-Tb [Y10]| AV1, AV2 | 450–923    | ToLCBV                               | GC, RDP     |
| TYLCCNV-Tb [Y11]| AV1, AV2 | 450–923    | ToLCBV                               | RDP, GC     |
| TYLCHKV-[2]   | AV1, AV2 | 296–1197     | ToLCBV                               | RDP, gc     |
|               | AC1, AC4 | 2200–2360    | ToLCBV                               | GC          |
|               | AC1, AC4 | 2390–2630    | ToLCBV                               | RDP, GC     |
| TYLCHKV-[1]   | AV1, AV2 | 305–1206     | ToLCBV                               | RDP, gc     |
|               | AC1, AC4 | 2203–2363    | ToLCBV                               | GC          |
| TYLCHKV-[MM]  | AV1, AV2 | 157–1058     | ToLCBV                               | RDP, GC     |

*Method: gc - gap closing, rdp - reverse primer de novo, mc - mismatch correction, RDP - reverse primer de novo followed by gap closing, GC - gap closing followed by reverse primer de novo, MC - mismatch correction followed by reverse primer de novo.
number of prerequisites for recombination between begomoviruses. These include shared host ranges (possibly influenced by the emergence of B whitefly biotype), the ability to co-infect the same cells [33-35], high levels of viral replication [36], and overlapping geographical ranges. If all of these prerequisites are met for the tomato-infecting begomoviruses in South and Southeast Asia then one would expect there to be frequent and invariable recombination amongst all of these viruses. However, fitness disadvantages may be associated with some sequence exchanges that would lead to the selective elimination of many newly produced recombinants.

The recombination sites distributed non-randomly along the genome. The recombination breakpoints were detected in all the six reading frames of south Indian viruses and viruses from eastern and western India. The breakpoints in the Chinese and Thai viruses were located in AV1, AV2, AC1 and AC4, whereas ORFs AV1 and AV2 were identified to be cold spots in the Bangladeshi viruses.

The frequency and locations of recombination events measured as topological differences between trees constructed from different parts of the alignment were visualised as a half-diagonal compatibility matrix (Fig. 3). Each X and Y coordinate in the matrix is a gross estimate of the number of topological modifications needed to convert the tree constructed using sequences at position X into that constructed using sequences at position Y [31,37]. It was apparent from this matrix that recombination events are probably not randomly distributed throughout begomovirus genomes. The highest frequency of recombination apparently occurs in the portion of the C1/AC1 ORF encoding the N-terminal portion of Rep. For example, the matrix indicates that there are an excess of 0.16 phylogeny violations per clade when trees constructed using sequences between alignment positions 351 and 1251 are
compared with those constructed using sequences between alignment positions 2451 and 2951. This analysis also indicated the probable absence in certain regions of begomovirus genomes of recombination events that had any substantial phylogenetic effect. For example, all phylogenetic trees constructed using coat protein gene sequences were all in good agreement with one another indicating a relative absence of recombination breakpoints within the CP gene.

We examined phylogeny violations and number of recombination events in our data set from the perspective of parental sequence relatedness. We noted that in general phylogeny violations clustered around the genetic distance 0.30. The observed frequency of phylogeny violations were inversely correlated (r = -0.36 p < 0.05) to the pairwise distances of the fragments involved in exchange (Fig. 4A). In addition, the number of recombination events was also inversely correlated (r = -0.35 p < 0.05) to the diversity between the exchanged fragments (Fig. 4B), we used only identified parental sequences to estimate the genetic distance between horizontally transferred fragments and the sequences that they replaced. Overall the frequency of phylogenetic violations and number of recombination events decreased with increasing parental sequence diversity. In a study with artificial and natural geminivirus recombinants Martin and co-workers [38] demonstrated that the degree of similarity between a horizontally inherited sequence and the sequence it replaces is an important determining factor of recombinant fitness. Rather than the non-random distribution of break points observed here being due to higher recombination rates in some genome regions than others [39], the distribution seems to have been created by natural selection only allowing the survival of recombinants with high fitness. In the more diverse genome regions where recombination events are not detected it is possible that these regions would not function properly when transferred into foreign genetic backgrounds.

**Conclusion**

Finally, the variable recombination and diversity-dependent distribution of recombination hotspots in tomato-infecting begomoviruses is valuable new information that has emerged from this study. Perhaps this is the first report of variable recombination reported among tomato-infecting begomoviruses found in the same region. Further, recombinant forms, recombination hot spots and frequency of recombination documented in this study would provide new information for understanding the diversity and evolution of tomato-infecting begomoviruses in Asia. In addition to evolutionary considerations, understanding the implications of recombination observed in these viruses on efforts to develop resistant tomatoes through conventional breeding and genetic engineering are important and attempts should be focused on these issues for developing effective disease management strategies. Given that the N-terminal portion
of rep is highly recombinogenic it is perhaps worrying that so many virus derived transgenic resistance strategies are focusing on this portion of the geminivirus genome [40-43]. It may be wiser to develop virus derived resistance strategies using genome regions that are less recombinogenic as this will make it more difficult for viruses to overcome resistance by simply replacing targeted genome regions with variants that are not targeted.

Methods

Sequence data

The study sequences comprised 35 publically available (as on June 2006) complete Indian, Pakistani, Chinese, Bangladeshi, Sri Lankan, Malaysian, Thai, Philippine and Taiwanese tomato-infecting begomovirus DNA-A and DNA-A-like components (Table 2). These sequences were aligned using the CLUSTAL W [44] using gap open and extension penalties- of 10.

Phylogenetic network and pairwise homoplasy test

Phylogenetic evidence for recombination was detected with Splits-Tree version 4.3 [45] using the neighbor-Net method [46]. Neighbor-net depicts conflicting phylogenetic signals in the data that are caused by recombination as cycles within unrooted bifurcating trees. Although, we report evidence for reticulate evolution in such phylogenetic graphs obtained using parsimonious sites, pairwise Hamming distances and no gaps, we obtained similar results with other distance measures and settings.

We statistically verified the presence of recombination identified visually in phylogenetic graphs using the pairwise homoplasy test (PHI) implemented in Splits Tree 4.3. PHI has been shown to powerfully identify the presence/absence of recombination within a wide range of sequence samples with a low false positive rate [47].

Detection of recombination breakpoints

The recombination breakpoint analysis was carried out using Recombination detection program RDP [48], GENECONV [19] and MAXIMUM CHI SQUARE [49], selected following the conclusions of studies on evaluation of different methods of recombination detection [50,51]. All these methods are implemented in RDP2 [52,53]. Default RDP2 settings were used throughout (P-value cut-off = 0.05 and the standard Bonferroni correction was used), other than that sequences were considered as circular, consensus daughters were found and breakpoints were polished. We used principally the information inferred by more than one method, as evaluation of the performance of these recombination detection methods using simulated and empirical data indicated that one should not rely too heavily on the results of a single method (Posada, 2002). In RDP analysis, the length of the window was set to 10 variable sites, and the step size was set to one nucleotide. P values were estimated by randomizing the alignment 1,000 times. For GENECONV analysis, the g-scale parameter was set to 1 and the number of permutations was set to 10,000.

Phylogenetic congruence

To examine phylogenetic support for each identified recombination event in the breakpoint analysis, we used the retained sequence position version of the TreeOrder Scan method [31] implemented in Simmonics2005 (Version1.4) package. TreeOrder Scan records the position of each sequence in a series of phylogenetic trees produced by sets of overlapping fragments across the
genomes. Deviations in the tree order of individual sequences and of group of sequences between fragments of defined length indicate conflicting phylogenetic relationships. Alternatively, individual non-recombinant sequences show constant tree order (position) across the genome. In the present analysis, we recorded the changes in the phylogenetic relationships of clades supported by 70 per cent bootstrap values for sequential 300 base sequence fragments at 100 nucleotide intervals.

**Frequency and mapping of recombination**

Estimation of the frequency and mapping of the locations of recombination events was achieved by phylogeny compatibility analysis using the TreeOrder Scan method. First, the TreeOrder Scan program produces optimally ordered neighbor-joining trees for fragments of definite length along an alignment. In the next step, a pairwise comparison is made between trees constructed from each sequence fragment along the alignment. Then a phylogenetic compatibility value is computed as the number of times the phylogeny of one tree has to be violated to match the tree order observed in other trees constructed along the length of an alignment. In our case we assigned sequences to predefined groups based on their geographical origin and a bootstrap value of 70 per cent was used as threshold for scoring phylogeny violations. All pairwise compatibility values were calculated using trees constructed for 300 nucleotide sequence fragments separated by 100 nucleotides across the length of the analysed alignment. These compatibility values were then plotted on a phylogenetic compatibility matrix.

**Competing interests**

The author(s) declare that they have no competing interests.

| Species/strain | Genbank accession | Abbreviation |
|----------------|-------------------|--------------|
| Tomato leaf curl Bangalore virus | Z48182 | ToLCBV |
| Tomato leaf curl Bangalore virus-[Ban4] | AF165098 | ToLCBV-Ban4 |
| Tomato leaf curl Bangalore virus-[Ban5] | AF295401 | ToLCBV-Ban5 |
| Tomato leaf curl Bangalore virus-[Kolar] | AF432855 | ToLCBV-Kol |
| Tomato leaf curl Bangladesh virus | AF188481 | ToLBDV |
| Tomato leaf curl Gujarati virus-[Kellow] | AF449999 | ToLCGV-Kel |
| Tomato leaf curl Gujarati virus-[Vadodara] | AF413671 | ToLCGV-Vad |
| Tomato leaf curl Gujarati virus-[Varanasi] | AF190290 | ToLCGV-Var |
| Tomato leaf curl Joydebpur virus-Mild | A1875159 | ToLCJMV-Mld |
| Tomato leaf curl Karnataka virus | U38239 | ToLCKV |
| Tomato leaf curl Malaysia virus | AF327436 | ToLCMV |
| Tomato leaf curl New Delhi virus-Mild | U15016 | ToLCNDV-Mld |
| Tomato leaf curl New Delhi virus-Severe | U15015 | ToLCNDV-Svr |
| Tomato leaf curl New Delhi virus-Severe [Jessore] | A1875157 | ToLCNDV-Svr-Jess |
| Tomato leaf curl New Delhi virus-[Lucknow] | Y16421 | ToLCNDV-Luc |
| Tomato leaf curl New Delhi virus-[Luffa] | AF102276 | ToLCNDV-Luf |
| Tomato leaf curl New Delhi virus-[Pkt1/8] | AF448059 | ToLCNDV-Pkt1/8 |
| Tomato leaf curl New Delhi virus-[Pkt5/6] | AF448058 | ToLCNDV-Pkt5/6 |
| Tomato leaf curl Philippines virus | AB050597 | ToLCPV |
| Tomato leaf curl Philippines virus-[LB] | AF136222 | ToLCPV-LB |
| Tomato leaf curl Sri Lanka virus | A274349 | ToLCSLV |
| Tomato leaf curl Taiwan virus | U88692 | ToLCTWV |
| Tomato yellow leaf curl China virus | AF311734 | TYLCCCNV |
| Tomato yellow leaf curl China virus-[Y64] | A1457823 | TYLCCCNV-Y64 |
| Tomato yellow leaf curl China virus-[Y10] | A1319675 | TYLCCCNV-Y10 |
| Tomato yellow leaf curl China virus-[Y11] | A1319676 | TYLCCCNV-Y11 |
| Tomato yellow leaf curl China virus-[Y36] | A1420316 | TYLCCCNV-Y36 |
| Tomato yellow leaf curl China virus-[Y38] | A1420317 | TYLCCCNV-Y38 |
| Tomato yellow leaf curl China virus-[Y5] | A1319674 | TYLCCCNV-Y5 |
| Tomato yellow leaf curl China virus-[Y8] | A1319677 | TYLCCCNV-Y8 |
| Tomato yellow leaf curl China virus-[Y25] | A1457985 | TYLCCCNV-Y25 |
| Tomato yellow leaf curl Thailand virus-[1] | X63015 | TYLCTHV-1 |
| Tomato yellow leaf curl Thailand virus-[2] | A141922 | TYLCTHV-2 |
| Tomato yellow leaf curl Thailand virus-[MM] | AF206674 | TYLCTHV-MM |
| Tomato yellow leaf curl Thailand virus-[Y72] | A1495812 | TYLCTHV-Y72 |
Authors’ contributions
HCP conceived and designed the study; HCP, MR executed the study and wrote the paper. Both the authors read and approved the final manuscript.

Acknowledgements
We gratefully acknowledge Dr D P Martin for critical reading of the manuscript, and very useful suggestions and comments. We thank Dr Peter Simmonds for inputs provided during the analysis, reading of the manuscript and encouraging comments.

References
1. Brown JK, Bird J: Whitefly-transmitted geminiviruses and associated disorders in the Americas and the Caribbean basin. Plant Dis 1992, 76:220-225.
2. Moriones E, Navas-Castillo J: Tomato yellow leaf curl virus, an emerging virus complex causing epidemics worldwide. Virus Res 1998, 58:1-134.
3. Polston JE, Anderson PK: The emergence of whitefly-transmitted geminiviruses in tomato in the western hemisphere. Plant Dis 1997, 81:1338-1369.
4. Simone GW, Brown JK, Hiebert . Cullen RE: New geminivirus epidemics in Florida tomatoes and peppers. Phytopathology 1999, 80:1063.
5. Varma A, Malathi VG: Emerging geminivirus problems: A serious threat to crop production. Ann Appl Biol 2003, 142:145-164.
6. Morales FJ, Anderson PK: The emergence and dissemination of whitefly-transmitted geminiviruses in Latin America. Arch Virol 2001, 146:415-441.
7. Broidd RW, Markham PG: Cotton leaf curl disease. Virus Res 2000, 71:151-159.
8. Harrison BD: Advances in geminivirus research. Annu Rev Phytopathol 1985, 23:55-82.
9. Harrison BD, Robinson DJ: Green shoots of geminivirology. Physiol Mol Plant Pathol 2002, 60:215-218.
10. Padidam M, Beachy RN, Faquett CM: The role of AV2 ("precoat") and coat protein in viral replication and movement in tomato leaf curl geminivirus. Virolgy 1996, 224:390-404.
11. Jupin I, De Kouchkovsky F, Jouanneau F, Gronenborn B: Movement of tomato yellow leaf curl geminivirus (TYLCV): Involvement of the protein encoded by ORF C4. Virolgy 1994, 204:82-90.
12. Laufs J, Traut W, Heyraud F, Matzev T, Roger SG, Schell J, Gronenborn B: In vitro cleavage and joining at the viral origin of replication by the replication initiator protein of tomato yellow leaf curl virus. Proc Natl Acad Sci USA 1993, 92:3879-3883.
13. Wantig L, Heuwer A, Noris E, De Kouchkovsky F, Jouanneau F, Gronenborn B, Jupin I: Genetic analysis of monopartite tomato yellow leaf curl geminivirus: roles of V1, V2, and C2 ORFs in viral pathogenesis. Virolgy 1997, 228:132-140.
14. Pita JS, Fondong VN, Sangare A, Otun-Nape GW, Ogwal S, Faquett CM: Recombination, pseudorecombination and synergism of geminiviruses are determinants keys to the epidemic of severe cassava mosaic disease in Uganda. J Gen Virol 2001, 82:655-665.
15. Monci F, Sanchez-Campos S, Navas-Castillo J, Moriones E: A natural recombinant between the geminiviruses Tomato yellow leaf curl Sardinia virus and Tomato yellow leaf curl virus exhibit a novel pathogenic phenotype and is becoming prevalent in Spanish populations. Virolgy 2002, 303:317-326.
16. Zhou X, Liu Y, Robinson DJ, Harrison BD: Four DNA A variants among Pakistani isolates of cotton leaf curl virus and their affinities to DNA A of geminivirus isolates from okra. J Gen Virol 1998, 79:915-923.
17. Schierup MH, Hejn J: Consequences of recombination on traditional phylogenetic analysis. Genetics 2000, 156:879-891.
18. Posada D, Crandall KA: The effect of recombination on accurate estimation of phylogeny. Mol Biol Evol 2004, 21:1008-1021.
19. Delate H, Martin DP, Naze F, Golbach R, Reynaud B, Peterschmitt M, Let J M: South West indian ocean islands tomato begomovirus populations represent a new major monopartite begomovirus group. J Gen Virol 2005, 86:1533-1542.
20. Garcia-Andres S, Monci F, Navas-Castillo J, Moriones E: Begomovirus genetic diversity in the native plant reservoir Solanum nigrum: Evidence for the presence of a new virus species of recombinant nature. Virolgy 2006, 350:433-42.
21. Andrade EC, Manhani GG, Allenas PF, Calegario RF, Fontes EPB, Zerbini FM: Tomato yellow leaf curl geminivirus from Brazil with a closer relationship to viruses from Sida sp., forms pseudorecombinants with begomoviruses from tomato but not from Sida. J Gen Virol 2006, 87:3687-3696.
22. Garcia-Andres S, Tomas DM, Sanchez-Campos S, Navas-Castillo J, Moriones E: Frequent occurrence of recombinants in mixed infections of tomato yellow leaf curl disease-associated begomoviruses. Virolgy 2007, 365:210-219.
23. Worobey M, Holmes EC: Evolutionary aspects of recombination in RNA viruses. J Gen Virol 1999, 80:2535-2544.
24. Simmonds P, Midgley S: Recombination in the genesis and evolution of Hepatitis B virus genotypes. J Virol 2005, 79:15467-15476.
25. Reijas MR, Hagen C, Lucas WJ, Gilbertson RL: Exploiting chinks in plant’s armor: Evolution and emergence of geminiviruses. Annu Rev Phytopathol 2005, 43:361-394.
26. Sanchez-Compos SJ, Navas-Castillo R, Camero C, Soria JA, Diaz . Moriones E: Displacement of tomato yellow leaf curl virus (TYLCV)-Sr by TYLCV-Is in tomato epidemics in Spain. Phytomol 2009, 89:1038-1043.
27. Chowda Reddy RV, Munipappa V, Colvin J, Seal SE: Diversity and distribution of begomoviruses infecting tomato in India. Arch Virol 2005, 150:845-867.
28. Ribeiro SG, Ambrozewicz B, Avil AC, Bezzerra IC, Calegario RF, Fernandes JM, Lima MF, de Mello RN, Rocha H, Zerbini FM: Distribution and genetic diversity of tomato-infecting begomoviruses in Brazil. Arch Virol 2003, 148:281-295.
29. Accotto GP, Mullineaux PM, Brown SC, Marie D: Digitaria streak geminivirus replicative forms are abundant in S-phase nuclei of infected cells. Virolgy 1993, 193:257-259.
30. Simmonds P, van der Walt E, Posada D, Rybicki EP: The evolutionary value of recombination is constrained by genome modularity. PLoS Genet 2005, 1:475-479.
31. Faquett CM, Sawyer S, Idris AM, Brown JK: Sequence analysis and classification of apparent recombination begomoviruses infecting tomato in the Nile and Mediterranean basins. Phytomol 2005, 95:549-555.
32. Noris E, Accotto GP, Tavazza R, Brunetti A, Crespi S, Tavazza M: Resistance to tomato yellow leaf curl geminivirus in Nicotiana benthamiana has arisen by interspecific recombination. J Gen Virol 1997, 78:2101-2111.
33. Padidam M, Beachy RN, Faquett CM: Classification and identification of geminiviruses using sequence comparisons. J Gen Virol 1995, 76:249-263.
34. Swanson MH, Brown JK, Poulos BT, Harrison BD: Genome affinities and epitope profiles of whitefly-transmitted geminiviruses from Americas. Ann Appl Biol 1992, 121:285-296.
35. Harrison BD, Robinson DJ: Natural genomic and antigenic variation in whitefly-transmitted geminiviruses (begomovi- viruses). Annu Rev Phytopathol 1999, 37:369-398.
36. Zhou X, Liu Y, Calvert L, Munoz C, Otun-Nape GW, Robinson DJ, Harrison BD: Evidence that DNA-A of a geminivirus associated with severe cassava mosaic disease in Uganda has arisen by interspecific recombination. J Gen Virol 1997, 78:2101-2111.
37. Padidam M, Sawyer S, Faquett CM: Possible emergence of new geminiviruses by frequent recombination. Virolgy 1999, 265:218-225.
38. Sanz AL, Fraile A, Gallego JM, Malpica JM, Garcia-Arenal F: Genetic variability of natural populations of cotton leaf curl geminivirus, a single-stranded DNA virus. J Mol Evol 1999, 49:672-681.
39. Fauquet CM, Sawyer S, Idris AM, Brown JK: Sequence analysis and identification of apparent recombination begomoviruses infecting tomato in the Nile and Mediterranean basins. Phytopathol 2005, 95:549-555.
40. Noris E, Accotto GP, Tavazza R, Brunetti A, Crespi S, Tavazza M: Resistance to tomato yellow leaf curl geminivirus in Nicotiana benthamiana plants transformed with a truncated viral C1 gene. Virolgy 1996, 224:130-138.
41. Brunetti A, Tavazza M, Noris E, Tavazza R, Caciagli P, Ancora G, Crespi S, Accotto GP: High expression of truncated viral Rep protein confers resistance to tomato yellow leaf curl virus in...
transgenic tomato plants. *Mol Plant Microbe Interact* 1997, 10:571-579.
42. Yang Y, Polston JE, Patte CP, Hiebert E, Sherwood TA: *Use of Tomato yellow leaf curl virus (TYLCV) Rep gene sequences to engineer TYLCV resistance in tomato*. *Phytopathology* 2004, 94:490-496.
43. Antignus Y, Vunsh R, Lachman O, Pearlsman M, Maslenin L, Hananya U, Rosner A: *Truncated Rep gene originated from Tomato yellow leaf curl virus-Israel [Mild] confers strain-specific resistance in transgenic tomato*. *Ann Appl Biol* 2004, 144:39-44.
44. Thompson JD, Higgins DG, Gibson TJ: *CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice*. *Nucl Acids Res* 1994, 22:4673-80.
45. Huson DH, Bryant D: *Application of phylogenetic networks in evolutionary studies*. *Mol Biol Evol* 2006, 23:254-267.
46. Bryant D, Moulton V: *NeighborNet: An agglomerative method for the construction of phylogenetic networks*. *Mol Biol Evol* 2004, 21:255-265.
47. Bruen TC, Philippe H, Bryant D: *A simple and robust statistical test for detecting the presence of recombination*. *Genetics* 2006, 172:2665-2681.
48. Martin D, Rybicki E: *RDP: detection of recombination amongst aligned sequences*. *Bioinformatics* 2000, 16:562-563.
49. Maynard Sj: *Analyzing the mosaic structure of genes*. *J Mol Evol* 1992, 34:126-129.
50. Posada D, Crandall KA: *Evaluation of methods for detecting recombination from DNA sequences: computer simulations*. *Proc Natl Acad Sci USA* 2001, 98:13757-13762.
51. Posada D: *Evaluation of methods for detecting recombination from DNA sequences: Empirical data*. *Mol Biol Evol* 2002, 19:708-717.
52. Martin DP, Williamson C, Posada D: *RDP2: recombination detection and analysis from sequence alignments*. *Bioinformatics* 2004, 21:260-262.
53. *Recombination Detection Program (RDP) [http://Darwin.uvigo.es/rdp/rdp.html]*