Widespread distribution and a new recombinant species of Brazilian virus associated with cotton blue disease
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Abstract

Background: Cotton blue disease (CBD), an important global cotton crop pathology responsible for major economic losses, is prevalent in the major cotton-producing states of Brazil. Typical CBD symptoms include stunting due to internodal shortening, leaf rolling, intense green foliage, and yellowing veins. Atypical CBD symptoms, including reddish and withered leaves, were also observed in Brazilian cotton fields in 2007. Recently, a Polerovirus named Cotton leafroll dwarf virus (CLRDV) was shown to be associated with CBD.

Results: To understand the distribution and genetic diversity of CLRDV in Brazil, we analyzed 23 CBD-symptomatic plants from susceptible cotton varieties originating from five of the six most important cotton-growing states, from 2004–2007. Here, we report on CLRDV diversity in plants with typical or atypical CBD symptoms by comparing viral coat protein, RNA polymerase (RdRp), and intergenic region genomic sequences.

Conclusion: The virus had a widespread distribution with a low genetic diversity; however, three divergent isolates were associated with atypical CBD symptoms. These divergent isolates had a CLRDV-related coat protein but a distinct RdRp sequence, and probably arose from recombination events. Based on the taxonomic rules for the family Luteoviridae, we propose that these three isolates represent isolates of a new species in the genus Polerovirus.

Background

Cotton (Gossypium spp.) is one of the most economically important crops in the world. Among its biotic pathologies, cotton blue disease (CBD) plays an important role due to its worldwide distribution and the high-magnitude of its associated productivity losses. Cotton blue disease was first described in the Central African Republic in 1949. It is transmitted by the aphid Aphis gossypii and its symptoms include leaf rolling, intense green foliage, vein yellowing, and a severe to moderate stunting caused by internodal shortening. Similar symptoms have also been observed in cotton crops of several regions of Africa, Asia and the Americas [1]. In Brazil, CBD is a serious crop problem capable of reducing the productivity of susceptible varieties by up to 80%, if intensive insecticidal control is not properly performed during the growing season.
Losses of up to 1,500 kg·ha⁻¹ of cotton seed have been reported in some states [2].

The viral origin of CBD was only recently identified [3]. In Brazil, the disease is associated with a virus from the family Luteoviridae, genus Polerovirus, named Cotton leafroll dwarf virus (CLRDV) [3]. Viruses from the family Luteoviridae are icosahedral and have a positive-sense RNA genome of ~6 Kb. These viruses are transmitted by aphids in a circulative and persistent manner [4], and are restricted to the phloem cells of their hosts. The Luteoviridae family is comprised of three genera, Luteovirus, Polerovirus, and Enamovirus, that are divided based on differences in their RNA-dependent RNA-polymerase (RdRp) and structural proteins.

Polerovirus genomes consist of six open reading frames (ORFs), typically designated ORFs 0 to 6 [5]. Non-structural genes are located in the 5′ portion of the genome. ORF0 encodes a silencing suppressor protein, P0 [6], and ORFs1 and 2 encode proteins related to viral replication, including the viral RNA polymerase [5]. Between the non-structural and structural protein sequences exists an intergenic region (IR) involved in subgenomic RNA synthesis [5,7,8]. The major coat protein (CP) is encoded by ORF3 at the 3′ portion of the genome. The viral movement protein is encoded by ORF4, which lies within the CP gene sequence but in another reading frame [9]. ORF5 is expressed by occasional suppression of the CP termination codon [10] and encodes the read-through domain (RTD) required for efficient aphid transmission [11,12].

The genetic variability of plant viral populations is an important aspect of their evolution and epidemiology [13]. Variation among isolates may affect virulence, infectivity, transmission, and symptom severity, and therefore is important to consider when developing strategies for disease control [14]. To better understand the distribution and genetic diversity of CLRDV in Brazil, we analyzed CBD-symptomatic plants from susceptible cotton varieties obtained from five of the six most important cotton-growing states of the country. This work represents the first time that isolates of CBD-associated viruses have been analyzed at the molecular level.

**Materials and methods**

**Plant material**

Twenty one cotton plants belonging to five susceptible cultivars of Gossypium hirsutum and two plants of G. barbadense were collected from cotton fields in the states of Mato Grosso, Goiás, Minas Gerais, São Paulo, Paraná, and Federal District from 2004 to 2007 (Table 1 and Figure 1).

**RNA extraction and amplification**

The total RNA was extracted from leaf samples using the RNeasy Plant Mini kit (Qiagen) in combination with a borate extraction buffer, following a previously described procedure [15]. Approximately 2.5 μg of total RNA from each isolate were used to synthesize first-strand cDNA using the CLRDV-specific primer OSR2 (5′-GCAACCTTT-TATAGTCCTCACAAT-3′), which anneals in the middle of CLRDV ORF5. Two independent nested PCRs were carried out for each viral isolate, one to amplify the CP gene and the other to amplify part of the RdRp gene plus the complete IR.

To amplify the CP gene, an initial reaction with the primers PL2F [3] and OSR2 was performed. The obtained amplicon was used as a template for a second amplification with the nested primers CPF and CPR [3], generating a 650-nt fragment. To amplify the partial RdRp sequence plus the IR, the previously described degenerated primers PLF and PLR [3] were used in the first reaction. Following this, the internal primers PL2F and PL2R [3] were used in the second reaction, generating a 468-nt fragment. Reactions were carried out with a denaturation step of 5 min at 95°C followed by 40 cycles (for the pairs PLF, PLR and PL2F, PL2R) or 35 cycles (for the primers PL2F, OSR2 and CPF, CPR) at 95°C for 1 min, 52°C (for PL2F, OSR2), 65°C (for CPF, CPR and PLF, PLR), or 50°C (for PL2F, PL2R) for 1 min, and 72°C for 1 min, and a final extension step at 72°C for 10 min. To amplify the full-length fragment corresponding to the partial RdRp, IR, and CP regions in a single PCR product, PL2F and CPR were used with an annealing temperature of 52°C for 40 cycles. RT-PCR products were purified using the Wizard SV Gel and PCR Clean-UP System (Promega) following the instructions provided.

**Sequencing and sequence analysis**

Three amplicons of each viral fragment, derived from independent RT-PCR reactions, were purified and then directly sequenced in both directions in automated ABI sequencers (models 310 or 377) using dye terminator cycle sequencing for all isolates. The resulting sequences were compared with the GenBank database. Consensus sequences of the CP, partial RdRp, and IR were obtained through the Multalin program [16].

Multiple sequence alignments of nucleotide or deduced amino acid sequences were submitted to the GeneDoc program [http://www.psc.edu/biomed/genedoc](http://www.psc.edu/biomed/genedoc) for computational analysis of the identities between the sequences. Phylogenetic reconstructions were performed using the MEGA 4 software [17]. Trees were constructed by the neighbor-joining (NJ) method [18]. The pair-wise deletion option, excluding gaps from the sequence alignment, and p-distance matrix were adopted. Data sets were
bootstrapped (1,000 replicates) to assess the confidence values of the phylogenetic trees, and bootstrap values < 50% were omitted. *Turnip yellow virus* (TYV), which represents the species most closely related to CLRDV across the polymerase and coat protein sequences, was included as an out-group.

The following sequences from members of the family *Luteoviridae* were used in the phylogenetic analyses: *Carrot red leaf virus* (CRLV) AY695933, *Cereal yellow dwarf virus-RPV* (CYDV-RPV), L25299; *Cereal yellow dwarf virus-RPS* (CYDVRPS), AF235168; *Chickpea stunt disease associated virus* (CpSDaV), AY956384; *TYV*, X13063; *Beet mild yellowing virus* (BMYV), X83110; *Beet chlorosis virus* (BChV), AF352024; *Beet western yellows virus* (BWYV), AF473561; *Cucurbit aphid-borne yellows virus* (CAbYV), X76931; *Potato leafroll virus* (PLRV), D00530; *Soybean dwarf virus* (SbDV), AB038147; *Tobacco vein distorting virus* (TVDV), AF402621; *Bean leafroll virus* (BLRV), AF441393; *Barley yellow dwarf virus-PAV* (BYDV-PAV), X07653; *Barley yellow dwarf virus-PAS* (BYDV-PAS), AF218798; *Barley yellow dwarf virus-GAV* (BYDV-GAV), AY220739; *Barley yellow dwarf virus-MAV* (BYDV-MAV), D01213; *Sugarcane yellow leaf virus* (ScYLV), AF157029; and *Pea enation mosaic virus-1* (PEMV-1), L04573.

The occurrence of recombination events was investigated by the programs Simplot [19], Genetic Algorithms for Recombination Detection (GARD) [20], and bootscans implemented by Recombination Detection Program ver-

![Geographical map of Brazil indicating the districts where cotton samples were harvested. DF, Federal District; GO, Goiás; MT, Mato Grosso; MG, Minas Gerais; PR, Paraná; and SP, São Paulo.](image)
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Table 1: Locations, symptoms, and cotton species and cultivars of the 23 CLRDV isolate samples analyzed.

| Location1 | Cotton sp.       | Cultivar | Isolate | Date | Symptom2 | Nested PCR |
|-----------|------------------|----------|---------|------|----------|------------|
| Cascavel – PR | G. hirsutum L. | CD401xFM966 | Cas1* | 2004 | typical | +          |
| Cascavel – PR | G. hirsutum L. | FM966     | Cas2*  | 2004 | typical | +          |
| Cascavel – PR | G. hirsutum L. | CD406xFM966 | Cas3* | 2004 | typical | +          |
| Cascavel – PR | G. hirsutum L. | CD406xFM966 | Cas4* | 2004 | typical | +          |
| Sta. Helena de Goiás – GO | G. hirsutum L. | CNPA ITA 90 STG | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir1  | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir2  | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir3  | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir4  | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir5  | 2005 | typical | +          |
| Piracicaba – SP | G. hirsutum L. | CNPA ITA 90 | Pir6  | 2005 | typical | +          |
| Brasília – DF | G. barbadense L. | na DF1 | 2005 | typical | +          |
| Brasília – DF | G. barbadense L. | na DF2 | 2005 | typical | +          |
| Acreúna – GO | G. hirsutum L. | ST 474 Acr10 | 2006 | typical | +          |
| Presidente Olegário -MG | G. hirsutum L. | ST474 PO1 | 2007 | atypical | +          |
| Holambra – SP | G. hirsutum L. | nd Hol1 | 2007 | typical | +          |
| Primavera do Leste – MT | G. hirsutum L. | FM966 PL2 | 2007 | typical | +          |
| Primavera do Leste – MT | G. hirsutum L. | FM966 PL3 | 2007 | typical | +          |
| Primavera do Leste – MT | G. hirsutum L. | FM966 PL4 | 2007 | typical | +          |
| Campo Verde – MT | G. hirsutum L. | FM966 CV1 | 2007 | typical | +          |
| Campo Verde – MT | G. hirsutum L. | FM966 CV2 | 2007 | atypical | +          |
| Campo Novo do Parecis – MT | G. hirsutum L. | FM977 CNP1 | 2007 | typical | +          |

* Samples inoculated in the Coodetec Research Center greenhouse. na, not applied; nd, not determined. 1DF, Federal District; GO, Goiás state; MT, Mato Grosso state; MG, Minas Gerais state; PR, Paraná state; and SP, São Paulo state. 2Symptoms description: typical, internodal shortening, leaf rolling, intensive green foliage and vein yellowing; atypical, typical disease symptoms plus withered and reddish basal leaves.

To further analyze recombination events, sequences obtained from individual PCRs were concatenated in silico and used in the above-described software. For Simplot analysis, the two previously described sequences of CLRDV (AY758560 and AY758561) were used as queries against the reference sequences of the isolates PL3, CV2, and PO1 and a cluster of sequences composed of all other sampled CLRDV isolates. The identities were calculated in a sliding window of 120 bp, which was moved across the alignment in 10 bp steps. The parameters used in the GARD program for checking alignments of CLRDV and the divergent isolates were the multiple break points options and general discrete rate variation. A manual bootscan was implemented by RDP3 Beta 24, with window and step size parameters of 100 and 10 bp, respectively. A 70% bootstrap support was considered to be definitive.

Results

To analyze the ubiquity and diversity of CBD in Brazil, cotton plants showing typical symptoms were harvested from commercial and experimental fields in four of the six major cotton-producing states of Brazil, Mato Grosso (MT), Goiás (GO), São Paulo (SP), Paraná (PR), and at Federal District (DF), between 2004 and 2007 (Figure 1). Three plants with atypical CBD symptoms were also harvested in the states of Minas Gerais (MG) and MT. Plants with atypical symptoms had all of the typical CBD symptoms along with withered and reddish middle and basal leaves.

Using nested RT-PCR, we tested for the presence of CLRDV-related sequences in all 23 sampled plants. The CP sequence, part of RdRp, and the full IR were amplified in all the isolates tested (Table 1). All amplicons showed the expected size for CLRDV and were sequenced. At least three independent PCR reactions for each fragment from each sample were sequenced. The obtained sequences were aligned and a consensus nucleotide sequence for each isolate was made for the three viral fragments.

The CP sequences of all isolates were similar to the CP sequence from CLRDV deposited in the GenBank database (accession number AY758560). In addition, most of the RdRp sequences analyzed were also closely related to that of the previously reported isolate (accession number AY758561). However, three isolates (designated PL3, CV2, and PO1) showed best hit results with TVDV, an unclassified member of the family Luteoviridae, but not with CLRDV. This suggests that these three isolates may have resulted from recombination events. Interestingly, these three divergent viruses were found in plants with atypical CBD symptoms.
When we compared CP sequence identities from all the isolates, we observed a high degree of nucleotide identity (97–100%). However, when the partial RdRp sequences were analyzed, the sequence identities were lower (65–100%). Most of this diversity resulted from a high sequence divergence in the three isolates found in plants with atypical CBD symptoms. Isolates PL3, CV2, and PO1 shared identities of 68%, 68%, and 67%, respectively, with the original CLRDV RdRp sequence. The identities among PL3, CV2, and PO1, however, ranged from 93% to 95%. Excluding the divergent isolates from the analysis, identities ranging from 94% to 100% for the RdRp sequence were observed among the 20 newly-identified CLRDV isolates. Similar results were also obtained for the IR, where the identities ranged from 94% to 100% among the 20 CLRDV isolates. The three divergent isolates shared identities of 66% with the original CLRDV IR sequence.

**Phylogenetic relationships among isolates**

Phylogenetic relationships among the CLRDV isolates were determined using the NJ method. Since dendograms constructed from nucleotide or amino acid sequences produced similar results, only those derived from nucleotide sequences are shown (Figure 2). In general, CLRDV displayed a widespread distribution in Brazil. Two groups of CP phylogeny were found: a small cluster formed by isolates CV2 and PO1 (bootstrap of 96%) and a large cluster containing almost all of the other sampled isolates (bootstrap of 86%). PL3 appeared to be the most divergent isolate, since its CP sequence was not grouped with the others. Within the major cluster were clusters of viruses from the same locality. This was particularly observed in plants harvested in Cascavel, PR and in DF during the 2004 and 2005 harvests, and in those obtained from Primavera do Leste, MT in 2007 (Figure 2A).

The trees constructed from partial RdRp (Figure 2B) and IR (data not shown) sequences were congruent and revealed a clear segregation of the isolates into two monophyletic clusters. This dichotomy was supported by 100% bootstraps in trees constructed either on nucleotide- or amino acid-based alignments (data not shown). The first cluster contained the three divergent isolates, PL3, CV2, and PO1, while the second was comprised of all the remaining isolates. Again, there were clusters of isolates obtained from the same geographical region in the same harvest. The isolates from Cascavel that were harvested in 2005 grouped together, as did those from DF (2005) and Primavera do Leste (2007) (Figure 2B).

The three divergent isolates were clearly separated from the other 20 CLRDV isolates found. Although these three isolates were very close to the other isolates, the CP phylogenetic tree placed them in an independent cluster. Significant differences were found between RdRp sequences from the divergent isolates and those from viruses associated with typical CBD symptoms, suggesting the possibility of recombination events between the viral populations in the field.

**Characterization of the divergent isolates**

To characterize the divergent isolates, we first checked whether the sequences of two different viruses were misamplified in a co-infection context. Using a primer comb-
bination capable of amplifying a fragment comprised of the 3’ end of RdRp plus the IR and CP sequences. We obtained a single PCR product with the expected size for each divergent isolate (PL3, CV2, and PO1) (Figure 3A). The amplified fragments were cloned and sequenced, and the new sequences confirmed that each isolate corresponded to the amplification of a single viral genome.

The sequences obtained were aligned with the sequences of CLRDV and nine other randomly chosen isolates. All the viruses had almost identical CP sequences. Similarities were also observed in the 3’ end of the IR. However, the 5’ end of the IR and the 3’ portion of the RdRp sequence showed consistent differences (Figure 3B).

To better understand the taxonomic and evolutionary positions of isolates PL3, CV2, and PO1 within the family Luteoviridae, sequences of the three viruses were compared to well-characterized Luteoviridae members. When the deduced amino acid sequence of the CP gene was analyzed, identities of 97%, 79%, 78%, and 77% were observed with CLRDV, TYV, BMV, BChV, and BWV, respectively. Phylogenetic analysis grouped the three divergent isolates with CLRDV in the Polerovirus branch of the tree, together with other definitive members of this group (Figure 4A).

The partial RdRp sequence analyzed in this work encodes the 92 C-terminal residues of the viral protein. Although the RdRp sequences of the three recombinant isolates shared an almost 70% sequence identity with CLRDV, significant identities were also found with Polerovirus members, including TYV (68%) and BMV (66%), and with TVDV (66%), an unclassified Luteoviridae that has a Polerovirus-like RdRp. Identities with other known Luteoviridae viruses indicated that PL3, CV2, and PO1 are more related to members of the genus Polerovirus than to those of Luteovirus (data not shown). Phylogeny obtained from the partial RdRp amino acid alignment confirmed this result by grouping the three isolates with CLRDV in the Polerovirus branch (Figure 4B).

Together, the above results indicate that PL3, CV2, and PO1 should be regarded as definitive members of the family Luteoviridae, genus Polerovirus. Considering the high degree of sequence divergence in the RdRp region, these three isolates may also represent isolates of a new species in Luteoviridae and of a second CBD-associated virus in Brazil.

Recombination analysis

To identify recombination events that could explain the emergence of the PL3, CV2, and PO1 isolates, the RdRp, IR, and CP sequences from each of the 23 isolates were analyzed using Simplot. For the 20 CLRDV isolates, the analyses were carried out by in silico concatenation of the individual sequences obtained by PCR. Similar to the above results, a comparison of the RdRp sequences revealed that these three isolates had low sequence similarities to CLRDV and to the other 20 isolates (Figure 5). However, a pronounced increase in the similarities between PL3, CV2, and PO1 and the other CLRDV isolates was observed from the middle of the IR to the end of the CP sequence (Figure 5).

The hypothesis that the divergent isolates resulted from recombination events was reinforced by results obtained using GARD. A discordant phylogenetic signal in the alignment of PL3, CV2, PO1, the 20 other concatenated CLRDV isolates, and TYV nucleotide sequences indicated a clear breaking point at nucleotide (nt) 443 (Figure 6A). The proposed breakpoint-delimited phylogenetic tree revealed that the cluster formed by PL3, CV2, and PO1 was more divergent to CLRDV isolates than to those of the TYV out-group, when the 5’ portion of the alignment was considered (Figure 6B). However, for a segment downstream of the breakpoint, the phylogeny showed a grouping of the clusters formed by the divergent and CLRDV isolates

Nucleotide 443 is located within the IR, a well-known site of both intra- and inter-species recombination in the family Luteoviridae. To identify possible parental lines for the divergent isolates, a bootscan analysis was performed. Luteoviridae sequences corresponding to the partial RdRp, IR, and capsid gene were used to scan for homologue segments in PL3, CV2, and PO1. For the RdRp and IR sequences of PO1, the bootstrap values were low until ~nt 435 and did not support their clustering with CLRDV (Figure 7). However, no significant bootstrap values were reached to support the clustering of the divergent sequences with any other luteovirus. For CP and nearby sequences, the bootstrap values supported the grouping of the divergent isolates with CLRDV (Figure 7), as expected. Similar plots were obtained for isolates CV2 and PL3 (data not shown).

Discussion

Analyzing CLRDV distribution in cotton Brazilian fields, we were able to show that it has a widespread distribution and high sequence conservation. These conclusions arise from nucleotide sequence comparing part of the polymerase, the coat protein and the intergenic region of 23 new isolates and CLRDV isolate identified in 2005 in Primavera do Leste. Interestingly, we also found that three of them have enough divergence in their polymerase sequences to be considered as a new species.

Cotton blue disease is widely distributed in Brazil and throughout the world; however, due to the recent charac-
Amplification and alignment of the divergent nucleotide sequences.

(A) Amplification of a single amplicon (1074 bp) corresponding to the partial RdRp, IR, and CP regions of isolates PO1, PL3, and CV2. NI, non-infected plant; M, λ-DNA digested with PstI. (B) Alignment of isolates PO1, PL3, and CV2 with a CLRDV isolate from each of the Brazilian regions analyzed in this work. Black boxes represent residues present in all viruses in the alignment. Columns in the alignment with < 100% but > 60% conservation are shaded in gray. The arrow represents the break point indicated in the GARD analysis (see Figure 6). Sequences inside the black boxes are associated with subgenomic RNA formation [7].
Figure 4
Phylogenetic analysis of isolates PO1, PL3, and CV2 and other viruses from the family Luteoviridae. (A) The predicted amino acid sequences of the CP and (B) the C-terminal region of RdRp were aligned with Multalign and the trees were constructed with MEGA4. Bootstrap values are indicated at each node of the trees. GeneBank accession numbers for the sequences are listed in the Materials and Methods section. Luteovirus, Polerovirus, and Enamovirus members are distinguished with grey, orange, and blue circles, respectively. Carrot red leaf virus (CRLV) and Tobacco vein distorting virus (TVDV) are unclassified Luteoviridae members with Polerovirus-like RdRps.
terization of its causal agent at the molecular level, its viral population and/or isolate diversity have not previously been reported. We previously described a virus associated with CBD in Brazil [3]. This virus, CLRDV, was present in CBD-symptomatic plants from a crop field in Primavera do Leste, MT. Here we map for the first time the genetic diversity of CLRDV in Brazilian cotton fields covering the most important producer regions. Analyzing 23 new viral isolates, we observed a low diversity in the CP CLRDV gene. However, phylogenetic analysis of this sequence revealed a small genetic distance between CV2, PO1, and PL3 and the other CLRDV isolates, clustering them in different groups. These results may suggest a co-adaptation phenomenon, resulting from selection pressure, among the divergent and the other CLRDV isolates, giving rise very similar CP sequences. Indeed, evolutionary studies concerning the CP of Luteoviridae members show that this protein is subjected to various selection pressures [22]. The CP is directly associated with the success of infection, as it is involved in viral transmission, particle packaging, and viral accumulation within the plant [23,24]. Thus, a high degree of conservation in the CP protein sequence is expected.

_Luteoviridae_ members have varying degrees of sequence conservation along their RNA genome. A model has been proposed where the CP and RTD sequences are the most conserved, while the IR is the least. Accordingly, our
results showed that the partial RdRp CLRDV gene displayed a greater diversity than CLRDV CP. Our analysis of the IR, however, contradicted this model, and showed a low diversity in this region comparable to that of the RdRp sequence. The IR is a noncoding sequence, and should be more prone to sequence variation than other parts of the genome. However, its role as a promoter in the generation of the subgenomic RNAs could result in sequence conservation due to structural constraints [7,8].

Interestingly, the isolates amplified from plants displaying typical CBD symptoms were very similar with one another, but those associated with atypical symptoms (PL3, CV2, and PO1) displayed highly divergent RdRp
sequences compared to the previously published CLRDV sequence. A phylogenetic analysis of the partial RdRp region revealed a clear segregation between PL3, CV2, and PO1 and the other CLRDV isolates identified in this study, forming two monophyletic clusters. This segregation was partially observed in the phylogeny of the CP gene, although the dendograms for this sequence revealed a stronger relationship among the divergents and the other CLRDV isolates.

Taxonomic characterization of the divergent isolates using phylogenetic analyses indicated that PL3, CV2, and PO1 have a Polerovirus-like RdRp and a CLRDV-like CP. Moreover, the IR between the RdRp and CP sequences from divergent isolates have 187 nt, the same as that in CLRDV [3]. Thus, their IRs resemble those of Polerovirus members, which are ~100 nt long. Taken together, these data suggest that the three divergent isolates could result from recombination events between CLRDV ancestors and another member of the genus Polerovirus. Evidence of recombination was found using three different methods (Simplot, GARD, and Bootscan). However, we were unable to detect a parental sequence for PL3, CV2, or PO1. Thus, the Polerovirus member involved in recombination with CLRDV remains unknown.

Recombination events may play an important role in generating genome diversity. Inter-species recombination has frequently occurred in the evolution of Luteoviridae. RNA recombination event(s) probably created the divergence observed between the genera Luteovirus and Polerovirus [25]. Subsequent recombination events between polerovirus-polerovirus [26] and luteovirus-polerovirus [27,28]...
have occurred since the birth of these genera [5,25].

Recombination sites in the family appear to coincide with the starting points for subgenomic RNA synthesis in the IR [7]. Comparisons among the IR sequences of BWYV, PLRV, and BYDV RNAs revealed two spots that are apparently conserved in all viruses. These regions contain a repeated U1,A signature that is usually associated with the formation of subgenomic RNAs [7,8]. Our data suggest that the recombination event found in the three divergent isolates occurred downstream of the second recombination spot of the IR (Figure 3B). Consistent with this, crossover events at the same region have already been described for a bovine coronavirus [29].

Amino acid sequence comparisons between the three recombinant viruses isolated from plants with atypical CBD symptoms and the 20 analyzed CLRDV isolates revealed that they share identities of ~70% in the partial RdRp sequence and of ~97% in the CP region. The molecular criterion used for species discrimination [30] in the family Luteoviridae states that amino acid sequences from any viral gene product exceeding 10% divergence should be regarded as a new species. Furthermore, symptomatology differences are also an important criterion for species discrimination. Therefore, our results suggest that PL3, CV2, and PO1 should be considered isolates of a new species in the genus Polerovirus. Further studies on CBD are necessary to understand the taxonomic and epidemiological relationships between the viruses associated with this important disease.

Conclusion

Our data suggests a CLRDV low genetic diversity and widespread distribution in Brazilian states. Between the 23 samples analyzed, however, we identified three divergent isolates associated with atypical CBD symptoms. The analyses revealed that these isolates have a CLRDV-related coat protein but distinct RdRp sequences and probably arose from recombination events between CLRDV and an unidentified luteovirus. In agreement with molecular taxonomic criterions for the Luteoviridae, we can consider that these isolates represent a new specie in the genus Polerovirus.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

TFS carried out the molecular genetic studies, participated in the sequence alignment, the recombination analysis and drafted the manuscript. RLC participated in the design of the study, in the design of primers, in the sequence alignment. YC performed RT-PCR and NESTED assays. Sample identification and collection in field were carried out by PS and J-LB. They also participate in the design of the study. MF SV conceived of the study, and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

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References

1. Cauquil J: Études sur une maladie d’origine virale du cotonnier: la maladie bleue. Cat Fib Trop 1977, 32:259-278.
2. Freire EC: Doença azul tem solução. In Cultivar Volume 1. Pelotas; 1998:64-65.
3. Correa RL, Silva TF, Simeos-Araujo JL, Barroso PA, Vidal MS, Vaslin MF: Molecular characterization of a virus from the family Luteoviridae associated with cotton blue disease. Arch Virol 2005, 150:1357-1367.
4. Gray S, Gildow FE: Luteovirus-aphid interactions. Annu Rev Phytopathol 2003, 41:539-566.
5. Mayo MA, Ziegler-Graff V: Molecular biology of luteoviruses. Adv Virus Res 1996, 46:413-460.
6. Pfeffer S, Dunoyer P, Heim F, Richards KE, Jonard G, Ziegler-Graff V, Reutenauer B, Martin J, Van den Heuvel JF, Verbeek M: Nuclearotide sequence of potato leafroll luteovirus RNA. J Gen Virol 1989, 70:1037-1105.
7. Miller WA, Brown MC, Wang S: New punctuation for the genetic code: luteovirus gene expression. Seminars in Virology 1997, 8:3-13.
8. Tacke E, Schmitz J, Prufer D, Rohde W: Mutational analysis of the nucleic acid-binding 17 kDa phosphoprotein of potato leafroll luteovirus identifies an amphipathic α-helix as the domain for protein/interaction. Virology 1993, 197:274-282.
9. Mayo MA, Miller WA: The Luteoviridae. In The structure and expression of luteovirus genomes; Wallingford Edited by: Smith HG, BarKer H. CAB1: 1999:23-42.
10. Brault V, Heuvel JFVan den, Verbeek M, Ziegler-Graff V: P0 of beet Western yellow virus is a suppressor of posttranscriptional gene silencing. J Virol 2002, 76:6815-6824.
11. Mayo MA, Robinson DJ, Jolly CA, Hyman L: Nucleotide sequence of potato leafroll luteovirus RNA. J Gen Virol 1998, 79:1509-1522.
12. García-Arenal F, Fraile A, Malpica JM: Variability and genetics structure of plant virus populations. Annu Rev Phytopathol 2001, 39:29.
13. Hall G: Selective constraint and genetic differentiation in geographically distant barley yellow dwarf virus populations. J Gen Virol 2006, 87:3067-3075.
14. Yingru WU: A quick and easy method for isolating good-quality RNA from cotton (Gossypium hirsutum L.). Journal Molecular Biology Reporter 2002, 20:213-218.
15. Corpet F: Multiple sequence alignment with hierarchical clustering. Nucleic Acids Res 1988, 16:10881-10890.
16. Tamura K, Dudley J, Nei M, Kumar S: MEGA4: molecular evolutionary genetics analysis (MEGA) software version 4.0. Mol Biol Evol 2007, 24(8):1596-1599.
17. Saitou N, Nei M: The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol Biol Evol 1987, 4(4):406-425.
deficiency virus type 1 genomes from subtype c-infected seroconverters in India, with evidence of intersubtype recombination. J Virol 1999;152:160.

20. Pond SLK, Posada D, Gravenor MB, Woelk CH, Frost SDW: Automated phylogenetic detection of recombination using a genetic algorithm. Molecular Biology and Evolution 2006, 23:1891-1901.

21. Martin DP, Williamson C, Posada D: RDP2: recombination detection and analysis from sequence alignments. Bioinformatics 2005, 21(2):260-262.

22. Torres MW, Correa RL, Schrago CG: Analysis of differential selective forces acting on the coat protein (P3) of the plant virus family Luteoviridae. Genet Mol Res 2003, 4:790-802.

23. Peiffer ML, Gildow FE, Gray SM: Two distinct mechanisms regulate luteovirus transmission efficiency and specificity at the aphid salivary gland. J Gen Virol 1997, 78:495-503.

24. Brau H, Bergdoll M, Mutterer J, Prasad V, Pfeiffer S, Erdinger M, Richards KE, Ziegler-Graff V: Effects of point mutations in the major capsid protein of beet western yellows virus on capsid formation, virus accumulation, and aphid transmission. J Virol 2003, 77:3247-3256.

25. D'Arcy CJ, Mayo M: Proposals for changes in luteovirus taxonomy and nomenclature. Arch Virol 1997, 142:1285-1287.

26. Gibbs MJ, Cooper JI: A recombinational event in the history of luteoviruses probably induced by base-pairing between the genomes of two distinct viruses. Virology 1995, 206:1129-1132.

27. Rathjen JP, Karageorgos LE, Habili N, Waterhouse PM, Symons RH: Soybean dwarf luteovirus contains the third variant genome type in the luteovirus group. Virology 1994, 198:671-679.

28. Moonan F, Molina J, Mirkov TE: Sugarcane yellow leaf virus: an emerging virus that has evolved by a recombination between luteoviral and poleroviral ancestors. Virology 2000, 269:156-171.

29. Chang R-Y, Krishnan R, Brian DA: The UCUAAAC promoter motif is not required for high-frequency leader recombination in bovine coronavirus defective interfering RNA. J Virol 1966:2720-2729.

30. Fauquet CM, Mayo MA, Maniloff J, Desselberger U, Ball LA: Virus Taxonomy. Eighth report of the international committee on taxonomy of viruses 2005.