Brome mosaic virus defective RNAs generated during infection of barley plants

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Brome mosaic virus (BMV) purified from systemically infected barley leaves 8 weeks post-inoculation (p.i.) contained defective RNAs (D-RNAs). The D-RNAs were detected in total and virion RNAs extracted from infected plants at 8 weeks p.i. or later, but not before, when barley plants had been inoculated with virions either containing or lacking D-RNA. The D-RNAs were derived from genomic RNA3 by double or mainly single deletions in the 3a protein ORF, and formed a heterogeneous population. By using in vitro transcripts of D-RNA synthesized from full-length cDNA clones, the D-RNAs were shown to replicate in a helper virus-dependent manner and to be packaged into virions in barley protoplasts. Subgenomic RNA4 was produced from the D-RNA and the coat protein was also expressed. Existence of the D-RNAs together with BMV genomic RNAs in inoculated protoplasts decreased the accumulation of 3a protein but it had no apparent effect on the accumulation of BMV genomic RNA3 or the coat protein. This is the first report of naturally occurring D-RNAs generated during prolonged infection with BMV.

Introduction

Defective RNAs (D-RNAs) are deleted RNAs containing portions of the parental virus genome. They maintain cis-acting elements that contain virus replication and encapsidation signals (Roux et al., 1991; White et al., 1991) and accumulate only in mixed infection with non-defective helper virus, which supplies essential components in trans (Roux et al., 1991). D-RNAs that interfere with the replication of the helper virus genome are referred to as defective interfering RNAs (DI-RNAs). Interference with the parental helper genome may result in an increase (Romero et al., 1993) or a decrease in symptom severity (Roux et al., 1991; Hillman et al., 1987; Burgyan et al., 1989). DI-RNAs have been found in association with animal virus infections (Perrault, 1981; Holland, 1987) and have proved useful in identifying sequence elements involved in virus functions such as encapsidation and replication (Levis et al., 1986; Weiss et al., 1989; Schlesinger, 1988). They have also been implicated as important components in driving virus evolution (Steinhauer & Holland, 1987).

In recent years, D- and DI-RNAs have been found and well characterized in many plant viruses, including several tombusviruses (Hillman et al., 1987; Burgyan et al., 1989; Rochon, 1991; Rochon & Johnson, 1991), carmoviruses (Li et al., 1989), potexviruses (White et al., 1991), broad bean mottle bromovirus (BBMV) (Romero et al., 1993; Pogany et al., 1995) and cucumber mosaic cucumovirus (CMV) (Graves & Roossinck, 1995).

Brome mosaic bromovirus (BMV) is a small, spherical plant virus that infects cereals, including barley (Lane, 1981). The genome of BMV consists of three species of messenger-sense single-stranded RNA, 1, 2 and 3 (Ahlquist, 1992). RNA1 (3.2 kb) and RNA2 (2.9 kb) encode the 1a and 2a proteins, respectively, which are required for virus RNA replication (French et al., 1986; Kibertis et al., 1981). RNA3 encodes the 3a protein, which is required for cell-to-cell movement of the virus (Schmitz & Rao, 1996). Subgenomic RNA4, which encodes the coat protein (CP), is synthesized by the virus replicase from a promoter present in the (−)-strand of RNA3 (Miller et al., 1985).

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The EMBL accession number of the sequence reported in this paper is X58459.
We have found D-RNAs in purified virions of wild-type BMV and in several CP mutants after prolonged infection (8 weeks) of barley plants. The D-RNAs were derived from RNA3 by single or double deletions in the 3a protein gene. Here we report on the molecular characterization of two D-RNA clones that were derived from a CP mutant of BMV.

Methods

- **Virus strains.** Strains of BMV used were ATCC66, which has been propagated in our laboratory (Mise et al., 1992), and ATCC PV-47, ATCC PV-178, ATCC PV-180 and M1 (Ahlquist et al., 1984).

- **Plasmid clones.** Plasmids pBTF1, pBTF2 and pBTF3W contain the full-length cDNAs of BMV RNA1, RNA2 and RNA3, respectively (Mori et al., 1991; Mise et al., 1992). Progeny virus derived from infected plants inoculated with in vitro transcripts from these plasmids was referred to as KU2 strain (Nagano et al., 1997).

  A plasmid pBF3WSSSR25 was constructed as follows. pBTF3W was digested with Sall and SacI and the resulting 0.2 kb fragment was exchanged with the corresponding fragment of pAT3J5, a plasmid that contains the CP gene of BMV strain ATCC PV-47 (accession number X58459), to create pBF3WSSS. The nucleotide sequences of the Sall/SacI region differed between pBTF3W and pAT3J5 at six sites [A1312G, T1314C, A1323G, C1350A, A1356G and G1374A, where the bold letters indicate amino acid substitutions created by site-directed mutagenesis (Kunkel et al., 1987). Six amino acids contained in the resulting cDNA clones of BMV RNA1 and RNA2 were named R25.

- **Cloning and sequencing of D-RNA cDNA.** Full-length cDNAs of BMV RNAs or D-RNA were synthesized by RT–PCR with a set of primers; 5′-primers that contained a PstI site and corresponded to the 5′ end of BMV RNAs 1 and 2 or RNA3, and a 3′ primer that contained an EcoRI site and was complementary to the 3′ end of the BMV RNAs. The RT–PCR was conducted under conditions described previously (Nagano et al., 1997). The amplified cDNA products were separated by agarose gel electrophoresis, cloned directly into a TA cloning vector (pCR II, Invitrogen) or cloned into pUC118 (Vieira & Messing, 1987) at the PstI/EcoRI sites after digestion of the cDNA with these enzymes. To ascertain the heterogeneity of D-RNA, cDNA was partially synthesized from D-RNA and amplified by using the 5′ primer specific to BMV RNA3 and a 3′ primer, B6, the sequence of which is complementary to nucleotides 1324–1338 of RNA3 (Mise et al., 1992). The amplified products were digested with PstI and Aar5IHI and ligated into pUC118 (Vieira & Messing, 1987) at the PstI/Smal sites.

  Nucleotide sequences of two full-length and several partially synthesized cDNA clones of D-RNA were analysed with an automated DNA sequencer (Applied Biosystems, model 373A) according to the manufacturer’s recommendations.

- **In vitro transcription, inoculation, purification of virus and RNA extraction.** All plasmids were linearized with EcoRI and used as templates for in vitro transcription. Capped full-length transcripts were synthesized in vitro by using T7 RNA polymerase (Mori et al., 1991).

  Barley (Hordeum vulgare L. cv. Gose-shikoku) plants were grown under conditions described previously (Fujita et al., 1996). Six-day-old seedlings were used for inoculation. Virions and virion RNA were purified from infected plants as described previously (Okuno & Furusawa, 1979).

  Isolation of protoplasts of barley (cv. Hinode-hadaka) and inoculation of in vitro transcripts and virion RNAs were performed as described previously (Okuno & Furusawa, 1978; Kroner & Ahlquist, 1992). Total RNAs were extracted and virion fractions were obtained by PEG precipitation (Kroner & Ahlquist, 1992) from infected protoplasts at 24 h after inoculation.

- **Northern blot analysis.** Total or virion RNA was denatured and separated in a 1.5% agarose gel containing formaldehyde and MOPS and transferred to a nylon membrane (Hybond-N+, Amersham). (+)- and (−)-strand RNAs were detected by using 32P-labelled SP6 transcripts from HindIII-linearized pBSPL10 (Kaido et al., 1995) and EcoRI-linearized pBSML10 (Mori et al., 1993), respectively. The RNA signals were quantified with a digital radioactive imaging analyser (Fuji BAS 2000, Fuji).

- **Western blot analysis.** Proteins were extracted from barley protoplasts with sample buffer (Laemmli, 1970) and separated by electrophoresis on 15% polyacrylamide gels containing formaldehyde and MOPS and transferred to a nylon membrane (Hybond-N+, Amersham). (+)- and (−)-strand RNAs were detected by using 32P-labelled SP6 transcripts from HindIII-linearized pBSPL10 (Kaido et al., 1995) and EcoRI-linearized pBSML10 (Mori et al., 1993), respectively. The proteins were detected with alkaline phosphatase-conjugated anti-lg secondary antibody, followed by a colour reaction with 5-bromo-4-chloro-3-indolyl phosphate in combination with nitro blue tetrazolium. Protein bands were scanned by an Epson Scan II (Seiko Epson) and protein accumulation was quantified with the Quantity One program (PDI) version 3.0.

Results

Occurrence of BMV D-RNA

Barley seedlings were inoculated with BMV RNA3 transcripts together with transcripts of RNAs 1 and 2 (inoculum

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### Table 1. Summary of amino acid differences among BMV RNA3 cDNA clones

<table>
<thead>
<tr>
<th>Plasmid*</th>
<th>Amino acid encoded at the indicated site in the CP gene†</th>
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<tbody>
<tr>
<td></td>
<td>21</td>
</tr>
<tr>
<td>pBF3W</td>
<td>His</td>
</tr>
<tr>
<td>pBF3WS5</td>
<td>Arg</td>
</tr>
<tr>
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<tr>
<td>pBF3WG25</td>
<td>His</td>
</tr>
<tr>
<td>pBF3WR21G25</td>
<td>Arg</td>
</tr>
</tbody>
</table>

* Plasmids pBF3WR21, pBF3WS5H21, pBF3WS55R25, pBF3WG25 and pBF3WR21G25 were constructed on the basis of the amino acid differences between pBF3W and pBF3WS5.

† Bold letters indicate amino acid substitutions created by site-directed mutagenesis.
Fig. 1. (a) Agarose gel electrophoresis of RNAs extracted from virions 2 and 12 weeks p.i. RNAs were analysed on a 1% agarose gel in TBE buffer followed by ethidium bromide staining. The additional RNA is indicated by an arrowhead (left) and positions of RNAs 1, 2, 3 and 4 are indicated on the right. (b) Electrophoresis in a 0.8% agarose gel of RT–PCR products of R25 lacking D-RNA (lanes 2 and 3) and R25 containing D-RNA (lanes 4 and 5). Lane 1 shows a λ DNA marker digested with HindIII. Sizes of the markers are indicated on the left (kbp). P1/2 (lanes 2 and 4) and P3 (lanes 3 and 5) are primers specific to the 5′ termini of RNAs 1 and 2 and RNA3, respectively. The 3′-terminal primer used is complementary to all BMV RNAs. The RT–PCR product of D-RNA is indicated on the right (D).

Table 2. Time-course analysis of generation and maintenance of D-RNA in infected barley plants

Seven-day-old barley seedlings were inoculated with virions lacking (−D) or containing (+D) D-RNA. Total and virion RNAs were extracted as described in Methods. Detection of D-RNA, assessed by Northern blot analysis of both total and virion RNAs extracted from infected leaves, is scored as positive (+) or negative (−). Similar results were obtained by RT–PCR with 3′ and 5′ primers specific for BMV RNA3.

<table>
<thead>
<tr>
<th>Inoculum</th>
<th>Time p.i. (weeks)</th>
<th>1</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>10</th>
<th>12</th>
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<td>−D</td>
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<td>−</td>
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<td>−</td>
<td>−</td>
<td>+</td>
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<td>+</td>
</tr>
</tbody>
</table>

R25) and virions were purified from systemically infected leaves at 2 and 12 weeks post-inoculation (p.i.). Analyses of virion RNA by agarose gel electrophoresis showed that an additional RNA that was not detectable in the 2 weeks p.i. virion sample was present in the 12 weeks p.i. virion sample (Fig. 1a). The additional RNA migrated between RNA3 and RNA4 and appeared to be approximately equimolar with each genomic RNA.

To test the maintenance and generation of the additional RNA, barley seedlings were inoculated with either the 2 weeks p.i. or 12 weeks p.i. virions. Total and virion RNA obtained from infected leaves at various time p.i. up to 12 weeks were analysed. An additional RNA was first detected in each inoculation by at least 8 weeks p.i., but not before (Table 2). It was also detected by Northern blot analysis with a probe complementary to the 3′ conserved sequence of BMV RNAs. This suggested that the additional RNA was not maintained or generated in the early stages of infection but was generated naturally during infection between 6 and 8 weeks p.i. To test whether the additional RNA could generally be observed in BMV infection, several BMV strains and BMV CP mutants derived from the KU2 strain were inoculated to barley seedlings and total RNAs obtained from systemically infected leaves at 8 weeks p.i. were analysed by Northern blot analysis. Similar additional RNAs corresponding to that observed in the R25 infection were found to be associated with KU2, ATCC66, ATCC PV-47, ATCC PV-178, ATCC PV-180 and M1 strains and also with the BMV-KU2 CP mutants SS5, R21, G25, H21 and R21G25 (Table 1) (data not shown).
Fig. 3. (a)–(b) Northern blot analyses of progeny virus RNA in barley protoplasts inoculated with a mixture of in vitro transcripts containing or lacking D-RNA. Total RNA was extracted from infected protoplasts 24 h p.i., separated by electrophoresis in a 1–5% agarose gel, transferred to a nylon membrane and detected with probes for (+)-strand (a) or (−)-strand (b) BMV RNAs. Inocula: lanes 1, water (mock); 2, transcript of D2 RNA; 3, transcripts of BMV RNAs 1 and 2 and D2 RNA; 4, transcripts of BMV RNAs 1, 2 and 3 and D2 RNA; 5, transcripts of BMV RNAs 1, 2 and 3. Identities of the RNAs are indicated on the left. (c) Western blot analysis of the accumulation of 3a and coat proteins in barley protoplasts inoculated with a mixture of in vitro transcripts containing or lacking D-RNA. Protein was extracted from 4 × 10^4 protoplasts 24 h p.i. and was loaded onto a 15% SDS–PAGE gel, transferred to a PVDF membrane and detected with anti-BMV 3a or anti-BMV antibody. Inocula: lane 1, water (mock); 2, transcripts of BMV RNAs 1 and 2, and D2 RNA; 3, transcripts of BMV RNAs 1, 2 and 3 and D2 RNA; 4, transcripts of BMV RNAs 1, 2 and 3. (d) Northern blot analysis of the encapsidation of D-RNA in barley protoplasts. Virions were obtained at 24 h p.i. by PEG precipitation. RNAs were extracted and analysed with a probe for (+)-strand BMV RNAs. Inocula: as (a) and (b).

To determine the origin of the additional RNA, virion RNA of R25 containing the additional RNA was amplified by RT–PCR with the 3′ primer common to all BMV genomic RNAs and the 5′ primer specific to BMV RNAs 1 and 2 or the 5′ primer specific to RNA 3 (Fig. 1b). cDNA fragments corresponding to the additional RNA and RNA3 were amplified successfully by using the 5′ primer specific to BMV RNA3, while only RNA1 and RNA2 cDNAs were amplified by using the 5′ primer specific to BMV RNAs 1 and 2. No cDNA fragments corresponding to the additional RNA were amplified in virion RNA obtained from the 2 weeks p.i. samples (data not shown). These results suggest that the additional RNA was a defective RNA derived from BMV RNA3. Therefore, it will be referred to as D-RNA.

Two full-length cDNA clones of the D-RNA were obtained and their nucleotide sequences were determined. These clones had sequences similar to that of R25 RNA3. However, one clone (D1) had a single large deletion (500 bp) in the 3a ORF (Fig. 2). The other clone (D2) had a similar 500 bp deletion and an additional small deletion (66 bp) (Fig. 2), both in the 3a ORF. There was no deletion in the CP ORF or the non-coding region (sequence data not shown) in these clones.

Replication and encapsidation of D-RNA in protoplasts

The two full-length cDNA clones (D1 and D2) of the D-RNA were tested for their ability to replicate and to interfere with BMV RNA replication in barley protoplasts. In vitro
Fig. 4. (a) Schematic representation of restriction enzyme map of BMV R25 RNA3. The enzymes are identified above the RNA3 diagram with the nucleotide position of cleavage. (b) Nucleotide sequence heterogeneity of BMV D-RNA with varied deletion junction sites. RT–PCR products from the D-RNA were cloned into a plasmid vector and sequenced. Deleted regions are drawn as horizontal lines. Individual clones are indicated alphabetically on the left. Clones a–e possess an FbaI site as indicated. R25, partial nucleotide sequence of the parental virus (Mise et al., 1994).

transcripts of the D-RNAs from each clone were inoculated into barley protoplasts together with transcripts of BMV RNAs 1 and 2 or with those of BMV RNAs 1, 2 and 3. Total RNA from infected protoplasts was examined by Northern blot analyses. Fig. 3 shows the results obtained with clone D2. Similar results were obtained with clone D1 in other independent experiments (data not shown). Regardless of whether RNA3 transcript was included in the inoculum, the D-RNA was detected by both probes for (+) (Fig. 3a) and (−) strands (Fig. 3b) of BMV RNAs. Both (+)- and (−)-strand D-RNAs were also detectable when protoplasts were inoculated with BMV virion RNA containing D-RNA (data not shown). These results indicate that D-RNAs can replicate in protoplasts. When D-RNA transcripts were inoculated with transcripts of BMV genomic RNAs 1 and 2, accumulation of subgenomic RNA4 was detected (Fig. 3a). However, when D-RNA was inoculated without BMV RNAs 1 and 2, no accumulation of D-RNA was detected (Fig. 3a), indicating that the replication of D-RNA was helper-dependent.

The effects of the presence of D-RNA on the accumulation of virus RNAs and proteins were examined. The level of RNA3 accumulation relative to RNA1 + 2 was reduced to 91%, while that of RNA4 increased to 117% [on the basis of measurements from Fig. 3(a) and three other experiments]. The presence of D-RNA increased the accumulation of CP to 114% [on the basis of measurements from Fig. 3(a) and three other experiments]. In contrast, the presence of D-RNA reduced the accumulation of protein 3a to 54%. These results suggest that the presence of D-RNA does not interfere or interferes only slightly with the replication of RNA3 but apparently interferes with the synthesis of protein 3a in protoplasts.

Encapsidation of D-RNA was tested in barley protoplasts. The virion fraction was prepared from infected protoplasts and packaged RNA was analysed by Northern blot analysis. The D-RNA was detected together with BMV genomic RNAs (Fig. 3d, lanes 3 and 4), indicating that D-RNA was encapsidated into virions.

Heterogeneity of D-RNA sequence
To analyse any heterogeneity that may be present in the D-RNA population, virion RNA containing D-RNA was extracted at 8 weeks p.i. and full-length cDNAs were amplified by RT–PCR as described above. The PCR products of RNA3 and D-RNA were separated and purified after low-melting-point agarose gel electrophoresis and digested with restriction enzymes FokI, Scal, ClaI, Fbal, Aor51HI or SacI (Fig. 4a). Digested cDNAs were analysed by agarose gel electrophoresis. Full-length cDNA of R25 RNA3 used as control was completely digested by all the enzymes tested, resulting in the expected fragments corresponding to the enzyme cutting sites (Fig. 4a). The cDNA products of D-RNA were completely digested by FokI, Aor51HI and SacI. After Aor51HI digestion, D-RNA cDNA as well as RNA3 cDNA produced a 1.0 kbp fragment corresponding to the 3′ half of RNA3, indicating that there were no deletions in that region, which includes most of the intercistronic region, the CP ORF and the 3′ non-coding region. However, the cDNA of the D-RNAs was not digested by Scal or ClaI (central part of the 3a ORF), indicating that those sites did not exist in the D-RNA. Since the cDNA of the D-RNA was only partially digested by Fbal (3′ region of 3a ORF), the D-RNA population was assumed to be heterogeneous. This restriction enzyme mapping also confirms the sequencing result (Fig. 2) that a deletion occurred around the central region of 3a ORF. However, the cDNA of the D-RNA was completely digested by FokI, which has no recognition site.
in the D2 sequence, suggesting that D2 RNA could be a minor species.

For further examination of D-RNA heterogeneity, we synthesized partial cDNA of the D-RNA by using the 5′ primer specific to BMV RNA3 and the 3′ primer Be6. We obtained 20 cDNA clones and determined their nucleotide sequences. In this experiment, all the D-RNA clones contained single deletions in the 3a ORF and no D-RNA contained two deletions, such as were found in D2 (Fig. 2). The sequence data also confirmed the results of restriction enzyme mapping, showing that both the SspI and ClaI sites were absent from all the clones and that only a portion of the clones (5 of 20) contained an FspI site. The 5′ borders of the deleted region were roughly the same among the clones, though the exact junction sites varied (Fig. 4b), resulting in deletions ranging from 477 to 500 bp.

Discussion

The structure of the additional RNAs described here suggests that they are the first reported naturally occurring D-RNAs associated with BMV infection, although artificial DI-RNAs constructed from BMV RNA2 were reported previously (Marsh et al., 1991). A number of features of the BMV D-RNAs distinguish them from previously characterized plant virus D/DI-RNA. First, smaller RNAs with electrophoretic mobilities similar to that of the D-RNAs of R25 were detected in several BMV strains or BMV-KU2 CP mutants. This suggests that the presence of D-RNAs is a general occurrence after prolonged BMV infection of barley plants. It also suggests that mutations in the 5′-terminal region of the CP gene did not affect the generation of D-RNA.

Second, the D-RNAs were generated by either single or double deletions, exclusively in the 3a ORF (Figs 2 and 4b). The deleted regions are nt 369–868 of D1 RNA and nt 201–266 and 366–865 of D2 RNA, suggesting that the regions retained are essential for the accumulation of BMV D-RNA in planta. The D-RNA clones with single deletions had different deletion junctions (Fig. 4b), which resulted in sizes of deletions from 477 to 500 bp. In barley protoplasts inoculated with in vitro transcripts of D-RNA with either one or two deletions, the D-RNAs were replicated and encapsidated into virions when co-inoculated with BMV genomic RNAs. When present together with the genomic RNAs, the D-RNA reduced the accumulation level of protein 3a (Fig. 3c). This could be the result of competition for ribosomes between D-RNA and wt RNA3 in the synthesis of the truncated and wt 3a proteins.

The third specific feature of the BMV D-RNAs is their generation. As previously reported, repeated passage at high m.o.i. is required for the generation of DI-RNAs in animal viruses (Holland, 1990) and D-/DI-RNA in some plant viruses (Morris & Hillman, 1989; Knorr et al., 1991; Graves & Roossink, 1995). However, the BMV D-RNAs were generated in rather unique circumstances. Generation of D-RNA was demonstrated by inoculation of barley seedlings with virion inocula either containing or lacking D-RNA. In both cases, the D-RNA was not detected at 1–6 weeks p.i., but was detected after prolonged infection (8 weeks p.i.) (Table 2).

These features raise interesting questions; why isn’t the D-RNA with deletions in the 3a ORF maintained even in the initially inoculated leaves, and why does the 3a ORF become a target for deletions after prolonged infection? Explanations for these observations could be: (i) since D-RNAs replicate efficiently in protoplasts (Fig. 3a) and are encapsidated into virions (Fig. 3d), the lack of cell-to-cell movement rather than replication may be responsible for the lack of D-RNA maintenance in initially inoculated leaves. Because the 3a gene has a crucial role in virus cell-to-cell movement (Schmitz & Rao, 1996), RNA3 with a truncated 3a gene may be less advantageous for further cell-to-cell movement than that with an intact 3a gene. Therefore, the D-RNA may fail to accumulate to detectable levels even in initially inoculated leaves. (ii) It is possible that there may be a unique interaction between either BMV strains or BMV CP mutants and 8-week-old barley plants. Physiological changes in old barley may interact with and/or alter the 3a gene. Alternatively, the intact 3a gene might be dispensable in old barley plants. Therefore, the dispensable region in the 3a gene might be deleted and regions that are presumed to be necessary for RNA replication and encapsidation in plants could be retained.

Previously, D- and DI-RNAs have been found to be associated with CMV (Graves & Roossink, 1995) and BBMV (Romero et al., 1993; Pogany et al., 1995), respectively. Both viruses, as well as BMV, belong to the family Bromoviridae and their genomic organization is similar. D-RNAs of CMV and BMV originate from parental genomic RNA3, while DI-RNAs of BBMV are from RNA2. On the other hand, the generation processes are quite different. The CMV D-RNAs are produced upon serial passage of the wild-type Fry strain and are maintained by virus even after additional passage, while the BBMV DI-RNAs occur naturally in strains Tu and Mo (Romero et al., 1993) and are generated de novo by serial passage at high m.o.i. (Pogany et al., 1995). In contrast, BMV D-RNAs are produced during prolonged infection of BMV and cannot be maintained even in the initially inoculated leaves. This might result from young-barley-mediated inhibition of D-RNA encapsidation or an increased instability of D-RNA-containing virions, as previously reported in the case of BBMV DI-RNAs in pea (Romero et al., 1993). Further studies to find a favourable host that could support the maintenance of BMV D-RNAs are needed to investigate whether the existence of D-RNAs has any effect on the symptom development induced by the helper virus.

The mechanism of formation of BMV D-RNA is unknown. However, short regions showing sequence similarity and/or complementarity were found at the 5′ and 3′ junction sites (Figs 4b and 5). These local complementarities could juxtapose two molecules of (+)-strand RNA3, which might allow the
replicate to switch from one template to another (Fig. 5a), as
proposed previously for recombination events among BMV RNAs (Bujarski & Dzianott, 1991; Bujarski et al., 1994). Such a template-switching mechanism has also been largely accepted to be responsible for the generation of D-RNA molecules (Bujarski & Dzianott, 1991; Bujarski et al., 1994).

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