Coat protein expression strategy of oat blue dwarf virus

Michael C. Edwards *, John J. Weiland

USDA-Agricultural Research Service, Cereal Crops Research Unit, Northern Crop Science Laboratory, 1605 Albrecht Blvd. N., Fargo, ND 58102-2765, USA

A R T I C L E   I N F O

Article history:
Received 1 October 2013
Returned to author for revisions 19 October 2013
Accepted 15 December 2013
Available online 14 January 2014

Keywords:
MaraVirus
Tymoviridae
OBDV

A B S T R A C T

Oat blue dwarf virus (OBDV) is a member of the genus Marafivirus whose genome encodes a 227 kDa polyprotein (p227) ostensibly processed post-translationally into its functional components. Encoded near the 3′ terminus and coterminal with the p227 ORF are ORFs specifying major and minor capsid proteins (CP). Since the CP expression strategy of marafaviruses has not been thoroughly investigated, we produced a series of point mutants in the OBDV CP encoding gene and examined expression in protoplasts. Results support a model in which the 21 kDa major CP is the product of direct translation of a sgRNA, while the 24 kDa minor CP is a cleavage product derived from both the polyprotein and a larger ~26 kDa precursor translated directly from the sgRNA. Cleavage occurs at an LXG[G/A] motif conserved in many viruses that use papain-like proteases for polyprotein processing and protection against degradation via the ubiquitin-proteasome system.

Introduction

Members of the genus Marafivirus are alpha-like plant viruses (Goldbach et al., 1991; Rozanov et al., 1992) belonging to the family Tymoviridae in the order Tymovirales. These small, isometric, positive stranded RNA viruses have approximately 6.3–6.8 kb genomes that are similar in organization to those of the tymoviruses and encode large polyproteins with methyltransferase, helicase, and polymerase motifs (reviewed in Dreher et al., 2011). While the tymovirus coat protein (CP) is encoded in a separate open reading frame (ORF) just downstream of the polyprotein ORF, marafavirus CP-encoding sequences are nested within, and are 3′ co-terminal with, the ORF encoding the polyprotein. The tymovirus genome expression strategy has been shown to employ a papain-like protease to process the large precursor polyprotein, while the CP is translated directly from a subgenomic RNA (Dreher et al., 2011). A highly conserved core sequence known as the tymobox serves as a promoter for transcription of this sgRNA (Ding et al., 1990).

Precise details of marafavirus genome expression have not been demonstrated, although Edwards et al. (1997) proposed a model for expression of the oat blue dwarf virus (OBDV) genome based in part on its genomic similarities with tymoviruses. Marafaviruses possess a sequence analogous to the highly conserved tymobox subgenomic promoter sequence, known as the marabox (Izadpanah et al., 2002), which is presumed to have a similar function. Virions of marafaviruses, however, contain a major CP of 21 kDa and a minor CP of 24 kDa that differ only by an amino terminal extension present in the minor CP, whereas tymoviral particles contain a single CP. Because these CPs likely have a role in transmissibility of leafhopper-borne marafaviruses and the fact that there are structural differences between marafavirus and tymovirus genomes, the CP expression strategy of marafaviruses is an important feature to characterize. We now report the use of mutation analysis with an infectious OBDV clone to dissect and analyze CP expression of OBDV as a first step toward a better understanding of marafavirus gene expression. We also map the initial nucleotide at the 5′-terminus of the sgRNA encoding the CPs and show that it is consistent in position relative to the tymobox/marabox with that of sgRNA termini of the tymoviruses.

Results and discussion

The OBDV genomic region targeted to investigate CP expression and the mutants used in this investigation are shown in Fig. 1. Our strategy employed mutation to create premature stop codons, disrupt proposed initiation codons and a potential protease cleavage site, as well as to disrupt the putative sgRNA promoter sequence. Inoculation of oat leaf protoplasts with OBDV wild type transcripts (OBDV-2r, GenBank #GU396990) resulted in the accumulation after 24 h of readily detectable amounts of viral CP and gRNA+sgRNA on western and northern blots, respectively. Infection of protoplasts by OBDV was confirmed using fluorescence microscopy to visualize a variant in which the entire major CP was
replaced by the enhanced green fluorescent protein gene (not shown).

Initiation point for sgRNA transcription and role of sgRNA in CP production

Alignment of various tymo/mara box sequences (a hallmark of the Tymoviridae and putative promoter for sgRNA synthesis of tymo- and maraviruses) reveals conservation of an adenine nucleotide ~10 nt downstream from the 3′-edge of the core sequence (Fig. 2). This is the known 5′-end nucleotide for the sgRNAs of turnip yellow mosaic virus (TYMV), ononis yellow mosaic virus (OYMV), and kennedya yellow mosaic virus (KYMV) (Ding et al., 1990; Guilley and Briand, 1978). To determine whether the conserved A in that position in OBDV (A5573) is the 5′-terminal nucleotide of the OBDV sgRNA, 5′-RACE was performed on size-fractionated RNA extracted from infected protoplasts and the resulting amplicons were sequenced (not shown). Sequence data showed that A5573 represents the 5′-end of the OBDV sgRNA encoding the viral CPs and is the likely start site of transcription for this 939 nt sgRNA (excluding the poly A tail). OBDV is the first maravirus for which initiation point and protease cleavage site codons are shown under the sequence. (B) Mutant description and nomenclature. The mutant name is followed by indication of nucleotide/codon substitutions and the cognate amino acid change. Mutant IJMN-9 represents a double mutant altering both the putative minor CP initiation codon and the potential cleavage site.

---

**Fig. 1.** Illustration of the OBDV genome, and the genomic region and mutants used in the investigation of coat protein (CP) expression. (A) Diagram of the OBDV genome and the nucleotide sequence in the region encoding the CPs. The genomic RNA possesses a 5′-cap structure (star) and a 3′-poly A tail (An). The single, large ORF encodes a polyprotein with domains specifying methyltransferase (mtr), protease (pro), helicase/NTpase (hel), and polymerase (pol) activities fused to the sequence encoding the CPs. The major and minor coat proteins map to the same CP sequence and size differences between them are potentially determined by translation initiation at two different start codons (minor, AUG5581, and major, AUG5710). Whereas the major CP is postulated to be translated primarily from sgRNA, the minor CP could be produced by translation directly from the sgRNA, by cleavage after Gly–Gly (▼) of either or both the large polyprotein or a protein initiated at AUG5581, or by both direct translation and cleavage. The putative promoter for sgRNA synthesis, the mara box, is underlined and the start site for sgRNA synthesis (this work) is indicated by the bent arrow. Sites where premature termination codons were introduced are indicated by asterisks. Mutant names are shown above the corresponding sequence locations, and amino acid changes in potential initiation and protease cleavage site codons are shown under the sequence. (B) Mutant description and nomenclature. The mutant name is followed by indication of nucleotide/codon substitutions and the cognate amino acid change. Mutant IJMN-9 represents a double mutant altering both the putative minor CP initiation codon and the potential cleavage site.
to those known to severely reduce production of sgRNA in TYMV (Schirawski et al., 2000). Mutation of two key nucleotides (mutant AB15-25) reduced transcription of the sgRNA to an undetectable level even after 40 h of incubation, with a concomitant large reduction in accumulation of both forms of CP (Fig. 2). Production of neither CP was completely eliminated, suggesting that either sgRNA production was not completely shut down or that a low level of translation from sgRNA occurred. An extraneous band migrating slightly faster than the minor CP appeared in some instances on western blots, perhaps representing a conformational variant (Edwards et al., 1997). Interestingly, gRNA accumulation was also significantly reduced relative to that of wild type. This reduction appeared to be due to the loss of the major CP and not to the marafibox mutations, a view supported by similar reductions in gRNA accumulation for two other mutants whose ability to produce major CP was severely compromised (GH1-7 premature termination mutant and KL1-3 initiation codon mutant; Fig. 2).

The predicted initiation codons for the major and minor CPs are functional in vivo

As we noted previously (Edwards et al., 1997), two initiation codons (AUG5581 and AUG5710) exist with the potential to encode the minor and major CPs, respectively. The possible roles of these codons in the production of the OBDV CPs were investigated using a combination of site-directed mutants designed to prevent translation either through introduction of premature stop codons or elimination of initiation codons AUG5581 and AUG5710.

Results with point mutants generating premature stop codons (CD3-1, EF2-2, and GH1-7) confirmed and expanded our previous model for the origins of the major and minor CPs of OBDV. When a stop codon (UAA5711) was placed downstream of the major CP initiation codon (mutant GH1-7), neither CP was detectable after 24 h (Fig. 3) and gRNA accumulation was greatly reduced (Figs. 2 and 3). However, a trace amount of major CP was detectable after 40 h (Fig. 2), possibly due to UAA suppression from this highly translated ORF (Lao et al., 2009). With the stop codon (UUA5579) placed upstream of the major CP initiation codon (mutant EF2-2), abundant major CP was produced without detectable minor CP (Fig. 3). That the accumulation of gRNA and major CP was restored to near wild type levels in mutant EF2-2 suggests that the reduction of gRNA accumulation in mutant GH1-7 was linked to the absence of major CP expression. Mutant CD3-1 (UAA5578) positions the stop codon upstream of both the candidate start codon for the minor CP as well as the start codon for the major CP and thus precludes translation of either CP as a part of the polyprotein; both a minor and major CP are produced by this mutant (not shown). Thus, complete translation of the entire polyprotein ORF is not an absolute requirement for the production of either CP species, and AUG5581 is the probable initiation codon for the production of at least some fraction of the minor CP pool despite its close proximity to the 5' terminus of the sgRNA. Although such a short leader (8 nt for OBDV, as close as 1 nt for other marafiviruses) is unusual, initiation of translation at AUG codons positioned very close to the 5' end has been observed in...
other biological systems (Li and Wang, 2004; Elfakess and Dikstein, 2008).

Additional mutants were made to confirm the use of AUG5581 and AUG5710 as initiation codons in the production of minor CP and major CP, respectively. Mutation of AUG5710 to UUA (KL1-3) or UUG (KL2-5) abolished accumulation of major CP in protoplasts, as expected, but also greatly reduced accumulation of viral RNA (Fig. 3). However, near wild-type levels of viral RNA and the major CP, along with reduced levels of the minor CP, accumulated in protoplasts inoculated with mutants IJ4-7 and IJ2-1, in which AUG5581 was replaced by GUA or GUG, respectively (Fig. 3, IJ2-1 not shown). This supports the notion that the minor CP of OBDV can be produced via proteolytic cleavage from the large polyprotein.

Proteolytic cleavage plays a role in formation of the minor CP species

Previous peptide sequence evidence indicated the presence of a potential papain-like protease cleavage site located immediately following Gly1834 of the OBDV polyprotein (Edwards et al., 1997). Taken together with the experimental evidence above, it is thus conceivable that functional minor CP might be produced via proteolytic cleavage from the large polyprotein precursor. To further investigate this possibility, the candidate cleavage site was changed from Gly–Val1834 to Val–Val1834 (mutant MN6-1; Fig. 4). Typical accumulation of major CP was observed for this mutant, but the apparent size of the minor CP increased slightly, suggesting production exclusively from translation initiating at AUG5581 (Fig. 4). A similar result was obtained when Gly–Gly1834 was changed to Val–Val1834 in mutant MN2-1 (not shown). When both the putative cleavage and initiation sites were knocked out by changing AUG5581 to GUA in the MN6-1 background (double mutants IJMN9A and B), only major CP was detected in infected protoplasts. When combined with the results from premature stop codon mutant CD3-1 and initiation codon mutant IJ4-7, it is clear the minor CP can be produced by cleavage of either or both the polyprotein and the minor CP originating from sgRNA translation.

A model for OBDV CP expression and its applicability to other marafviruses

The empirical evidence offered here, taken together with previous sequence analysis by Edwards et al. (1997), supports a model for OBDV CP expression in which the major CP is translated directly from the 939 nt sgRNA (excluding the polyA tail), while the minor CP is cleaved from both the polyprotein and a minor CP precursor translated from the sgRNA. The underlying reason for the existence of multiple means of expression of the minor CP is not evident. While cleavage from the replicase polyprotein provides a source of minor CP, it may be that production of minor CP exclusively through this mechanism doesn’t provide stoichiometric amounts necessary for virion assembly. It is also possible that viruses such as OBDV are in an evolutionary transition toward CP production solelyvia sgRNAs and that readthrough of the larger replicase polyprotein is vestigial. In any case, production of the minor CP through a cleavage mechanism provides a regulatory feature with probable functional significance for both replication and encapsidation. Ultimately, the complexity of CP expression relative to that of tymoviruses may relate in some way to the infection by OBDV of both plant and insect hosts.

To consider this model in the larger context of CP expression in the marafviruses as a group, amino acid sequences of known and proposed marafviruses were aligned and compared (Fig. 5). In all cases, putative initiator methionine residues for both minor and major CPs are evident. The putative initiator Met for the minor CP is 5–15 amino acids downstream of the highly conserved GELL motif that is encoded by the core sequence of the mara box, with the exception of maize rayado fino virus (MRFV), switchgrass mosaic virus (SwMV), and Bermudagrass etched-line virus (BELV). A conserved motif comprised of LXGG is present 7–8 amino acids downstream of the initiator Met. Since this incorporates the demonstrated cleavage site that creates the amino terminus of the OBDV minor CP, it is reasonable to speculate that production of these other marafvirus minor CPs also involves cleavage at this site. A similar motif is present immediately upstream of the putative initiator Met in the polyproteins of MRFV, SwMV, and BELV. The SwMV sequence encodes an alanine at the putative cleavage site, but that is consistent with the previous observations of Kadare et al. (1995) and Sulea et al. (2006), who noted the presence of G or A residues in the P1 and P2 positions of confirmed and predicted cleavage site sequences of tymoviruses and coronaviruses, respectively. Thus, we propose the consensus sequence at the predicted cleavage site associated with minor CP expression in marafviruses is LXG[G/A]. Intriguingly, to counter cellular antiviral strategies that utilize the ubiquitin-proteasome system, the protease of the closely-related TYMV cleaves ubiquitin conjugates from its RdRp at the same consensus sequence in addition to its role in polyprotein processing (Chenon et al., 2012). Moreover, this consensus sequence is targeted by other viral cysteine proteases with both deubiquitinating and polyprotein processing activities (Barretto et al., 2005; Clementz et al., 2010; Karpe and Lole, 2011; Wang et al., 2011).

The model for expression of OBDV CPs appears directly applicable to the majority of accepted and proposed marafviruses, while an interesting variation of this strategy likely occurs with SwMV, MRFV, and BELV. For the latter, the predicted cleavage sites lie upstream of the predicted minor CP initiation codons, necessitating further studies to determine the precise CP expression strategies of these viruses. The location of the predicted cleavage site

![Fig. 4.](Image) Accumulation of viral coat proteins (CPs) and RNA in oat protoplasts inoculated with putative polymerase/CP cleavage site mutants. Oat protoplasts were inoculated with capped transcripts of wild type clone pOBDV, putative cleavage site mutant MN6-1, and double mutant IJMN-9 (minor CP initiation codon mutation combined with putative cleavage site mutation; IJMN-9A and B represent two different inoculations) and incubated for 24 h. Methods for detection of viral protein and RNA were as described in the legend for Fig. 2. The locations of major CP (Maj. CP), minor CP (Min. CP), genomic (gRNA), and subgenomic RNA (sgRNA) on the blots are shown. Partially purified OBDV (OBDV prep) was included on the western blot for size comparison.
downstream from the initiator methionine of the minor CP would provide a consistent amino terminus for minor CP produced by cleavage of both the replicase polyprotein and a minor CP precursor translated directly from sgRNA. Irrespective of the role of protein cleavage in the formation of the minor CP N-terminus, an important function of the cleavage site may be to liberate the polymerase from the CP. The location of a putative protease cleavage site upstream of the initiator methionine for the minor CPs of SwMV, MRFV, and BELV is consistent with this view. Furthermore, the single CP of the closely-related but unclassified Poinsettia mosaic virus (PnMV) is fused to the PnMV replicase initiation codon.

Fig. 5. Amino acid sequence alignment of the encoded polyproteins for known and proposed marafiviruses and PnMV in the region surrounding the coat protein (CP) amino termini. In (A) the proposed initiator methionine residues for the minor (left) and major (right) marafiviral CPs are boxed, as are invariant amino acids encoded by the marafivirus. Underlined amino acids indicate conserved sequence around the proposed LXG[G/A] cleavage recognition site for the encoded papain-like protease. (B) Realignment of sequences to highlight the proposed conserved protease recognition sequence (boxed). The conserved LXG[G/A] motif resembles that targeted by cellular deubiquitinating enzymes (Ubiquitin). Proposed initiator methionine residues for the minor and major marafiviral CPs are underlined. Although GSyV and GVQ are the same species, the reported sequences for each isolate are not identical and therefore both are included here.

Materials and methods

Parental and mutant clone construction

A full-length cDNA clone of OBDV from which infectious transcripts can be derived was generated previously (Edwards and Weiland, 2010). This clone (pOBDV-2r; GenBank #GU396990) served as the parent plasmid in the generation of mutants of OBDV with alterations in the CP gene(s). General amplification primers OBDS140fw and OBDS901rev were used in conjunction with mutagenic primers (Fig. 1, Table 1) to produce overlapping amplicons. PCR reactions included 1 ng of linearized plasmid as template and conditions were as described in Higuchi (1990). Typical reactions contained 1X Platinum Taq buffer, 0.1 mM each d(GATC)TP, 30 ng of each oligonucleotide primer, and 0.5 units of Grand Island, NY USA) in a volume of 30 µl was transferred to a 1.5 ml microfuge tube and placed on ice for 30 min prior to inoculation. The resulting amplicon with SalI (located at nts 5213 and 5831) released a 618 bp fragment that was used to replace the homologous region in pOBDV-13B2 (a derivative of pOBDV-2r in which the MCS was altered to remove the homologous region in pOBDV-13B2 is designated pOBDV herein. Transcript RNA was adjusted to a concentration of 0.2 µg/µl, and 5 µl was transferred to a 1.5 ml microfuge tube and placed on ice for 30 min prior to inoculation.

Protoplast inoculation

Protoplasts were produced from oat (Avena sativa L. cv ‘Rodney’) as previously described (Weiland and Edwards, 1994). The procedure routinely produced 10^6 protoplasts per gram fresh weight tissue from 7 day-old oat seedlings. A modification of the procedure by Matsuda and Dreher (2005) was used to transfet protoplasts with capped transcript RNA. Isolated protoplasts prepared from 3 g fresh weight of oat leaves were collected from the surface of a sucrose pad after centrifugation and diluted in 0.55 M mannitol/0.1% MES pH 5.6 (MM) containing 5 mM CaCl₂ and 40 mM KCl (MMCK). Protoplasts were pelleted by centrifugation at 115 g for 8 min. and resuspended in 4 ml MMCK. Approx. 5 x 10^5 protoplasts (per inoculation) were transferred to a microfuge tube, spun to collect the protoplasts, and the cells resuspended in 0.7 ml MMCK. Tubes containing protoplasts were incubated on ice for 30 min prior to inoculation.

Transcript RNA was adjusted to a concentration of 0.2 µg/µl, and 5 µl was transferred to a 1.5 ml microfuge tube and placed on
ice. A 0.8 ml aliquot of incubation medium (IM) consisting of MM supplemented with micronutrients as previously described (Weiland and Edwards, 1994) was chilled on ice in a microfuge tube along with a 0.4 cm gap electroporation cuvette. Chilled protoplasts were transferred to the tube containing transcript RNA and the mixture was rapidly pipetted into the pre-chilled cuvette. Contents of the cuvette were subjected to an exponential decay pulse of 100 V, 2 mF, and 480Ω, yielding a pulse duration of ~110 ms (delivered by a BTX ECM-600 electroporation system, Harvard Apparatus, Holliston, MA, USA). Following transfer of the contents of the cuvette to 0.8 ml of chilled IM, samples were incubated on ice for 10 min. Protoplasts were collected by centrifugation, resuspended in 1 ml fresh IM containing 100 µg/ml cefotaxime, and incubated for 24 h or 40 h at 22°C under continuous fluorescent light. Each mutant was tested in protoplast inclusions at least 4 times and included parallel mock and pOBDV inclusions within each experimental replication.

**Northern and western blot analysis**

Samples were harvested and processed for the analysis of viral RNA and protein as previously described (Weiland and Edwards, 1994). Total protein (for Western blots) and nucleic acids (for Northern blots) from 10⁴ protoplasts were separated on 12% sodium dodecylsulfate polyacrylamide gels and 1% denaturing agarose gels, respectively, as indicated (Edwards and Weiland, 2010). Proteins were electrophoretically transferred from polyacrylamide gels to nitrocellulose membranes (Schleicher and Schuell, Keene, N.H., USA), BA-85, 0.2 µm), which subsequently were incubated in a 1:1000 dilution of rabbit anti-OBDV, followed by incubation in a 1:2000 dilution of alkaline phosphatase-conjugated goat anti-rabbit IgG (Product #A0418, Sigma-Aldrich, St. Louis, MO, USA). Protein complexes were detected on a Kodak Image Station 2000 MM following treatment of the blot with LumiPhos WB (Thermo Scientific, Lafayette, CO, USA). Nucleic acids transferred to positively-charged nylon membranes by capillary blotting (Roche Applied Science, Indianapolis, IN, USA) were probed with a digoxigenin-labeled, denatured dsDNA probe representing nucleotides 5929–6508 of OBDV–2r. After incubating blots with alkaline phosphatase-conjugated anti-digoxigenin IgG, CDP-Star (Roche Applied Science, Indianapolis, IN, USA) was added and viral RNAs were detected by chemiluminescence.

**Mapping of the 5' end of the sgRNA**

Total nucleic acids were prepared from OBDV-infected oat protoplasts as previously described (Edwards and Weiland, 2010). Due to substrate competition with the genomic RNA in the cDNA generation step of the 5'-RACE procedure, gel-purified RNA was used as template in the mapping of the 5'-end of the sgRNA. Thus, RNAs were size-fractionated by agarose gel electrophoresis (1% GTG agarose) and RNA migrating a distance consistent with the estimated size of OBDV sgRNA (~1.0 kb) was extracted and purified using the Zymoclean™ Gel RNA Recovery Kit (ZymoResearch, Irvine, CA, USA). Following the manufacturer’s recommendations, primer OBDS5901rev (Table 1) was used to prime cDNA synthesis in the presence of SmartScribe reverse transcriptase and anchor primers contained in the SmartRACE kit (Clontech, Mountain View, CA, USA). Following amplification of primers Term1–221 K(–) and Term2–221 K(+) ( Table 1), gel-purified cDNAs were sequenced (MWG Operon, Huntsville, AL, USA). CLUSTAL-W software was used to illustrate conservation of amino acid residues around the CP amino termini employed the same strategy to align the resulting nucleotide sequences and the sgRNA 5'-end determined; initial protein sequence alignments used to illustrate conservation of amino acid residues around the CP amino termini employed the same software.

**Virus names, acronyms, and sequences used**

Acronyms and GenBank accession numbers for viruses discussed in the text and figures are: Bermuda grass etch-line virus (BELV, AY040531), Blackberry virus S (BIVS, FJ915122), Citrus sudden death-associated virus (CSDaV, NC_006950), Grapevine asteroid mosaic-associated virus (GAMaV, AJ249357), Grapevine rupestris vein feathering virus (GRVFV, AY706994), Grapevine Syrah virus-1 (GSY-S1, NC_012484), Grapevine virus Q (GVQ, an isolate of GSVV-1, FJ977041); Maize rayado fino virus (MRFV, NC_002786), Oat blue dwarf virus isolate 2r (OBDV–2r, GU396990), Poinsettia mosaic virus (PmMV, NC_002164), Switchgrass mosaic virus (SWMV, GU396990).
References


