

**Expression of multiple foreign epitopes presented
as synthetic antigens on the surface
of *Potato virus X* particles**

K. Uhde¹, R. Fischer^{1,2}, and U. Commandeur¹

¹RWTH Aachen, Institute for Molecular Biotechnology, Aachen, Germany

²Fraunhofer Institute for Molecular Biology and Applied Ecology (IME),
Schmallenberg, Germany

Received March 10, 2004; accepted June 8, 2004

Published online October 20, 2004 © Springer-Verlag 2004

Summary. We describe the construction of recombinant *Potato virus X* (PVX) vectors expressing two different epitopes, ep4 and ep6, from *Beet necrotic yellow vein virus* (BNYVV). The seven-amino-acid epitopes were expressed as N-terminal coat protein fusions and were displayed on the surface of PVX particles. Particle assembly into full virions was successful even though no wild type coat protein subunits were present, and the epitopes could be detected in crude extracts and purified virus preparations with appropriate antibodies. A construct containing both epitope sequences in tandem was also prepared. The resulting PVX particles could be detected by antibodies against ep4 and ep6, either individually or simultaneously, showing that both epitopes were accessible. In addition mixed infections with PVX vectors containing the individual ep4 and ep6 sequences were carried out. This resulted in the formation of PVX particles displaying ep4 alone, ep6 alone, or both epitopes. These experiments demonstrate for the first time that PVX can be utilized to present multiple epitopes, either tandemly on every coat protein subunit or as heteromultimeric assemblies, both of which could be useful vaccination strategies. The production of epitope-presenting viruses in which every coat protein subunit contains a foreign epitope allows the high-level expression of defined numbers of foreign antigen sites, making such viruses useful standards for immune detection.

Introduction

Viruses are widely used as vectors for the transient expression of foreign proteins in plants [27]. While the production of full-length proteins is often the objective of such expression experiments, another important application is the presentation of

foreign epitopes on the surface of viral particles [23]. After fusing the appropriate coding sequence to a highly expressed structural gene, such as a coat protein (CP) gene, transient expression of foreign proteins or peptides can be achieved at very high levels. The foreign coding region may be inserted so that the heterologous peptide is added to the end of the coat protein or located within an internal loop, since these regions tend to be displayed on the surface of the viral particle. As long as the inserted peptide is not too large, then neither the replication nor the assembly of the recombinant virus is impaired.

Epitope-presentation systems have been developed based on several different plant viruses, but the most popular are *Cowpea mosaic virus* (CPMV) [7, 26, 37] and *Tobacco mosaic virus* (TMV) [33, 35] since their structures have been solved at atomic resolution by X-ray crystallography. More recently, *Potato virus X* (PVX) has been investigated as a possible presentation system [9, 24, 25, 32] although much of the structural information available for this virus has been inferred rather than determined empirically. It is assumed that the N-terminal part of the CP is presented on the particle surface, so this is the ideal position to insert heterologous peptides [2, 18].

PVX, the type species of the genus *Potexvirus*, infects many solanaceous species [19]. The virus normally spreads by mechanical contact, and artificial infection with PVX is achieved by mechanical inoculation of host plants such as *Nicotiana benthamiana*, which become systemically infected. The RNA genome has five open reading frames (ORFs), the most 3' of which is controlled by a subgenomic promoter and encodes the 236-residue coat protein. PVX has been used as an expression vector for full-length proteins by incorporating an additional subgenomic promoter [8, 16]. It has also been widely used to study virus induced gene silencing [4, 10].

The major advantage of flexuous filament particles such as PVX for epitope presentation is that the capsid has no intrinsic size or packaging constraints, and many copies of the target epitope can potentially be displayed on the particle surface. The first experiment describing protein-presentation on PVX particles involved an N-terminal CP fusion with green fluorescent protein (GFP) [9]. Experimental investigations showed that proteins from 8.5 to 31 kD in size were only expressed on the intact PVX particle surface if wild type coat protein was also present. This was achieved by the insertion of a ribosomal skip site, the 2A peptide of *Foot-and-mouth disease virus* (FMDV) [11, 12, 30], between the heterologous sequence and the CP sequence [9, 30]. More recently, PVX has been used to express coat protein fusions with a single-chain variable fragment (scFv) antibody [32], a rotavirus major capsid protein (VP6) [25] and an epitope from glycoprotein 41 of *Human immunodeficiency virus* (HIV) [24]. In the present study we demonstrate that PVX can be used to display two synthetic antigens derived from BNYVV, either alone or in combination. We propose that synthetic antigens can be used as a standard reference for immune detection. Because of their particular structure, such antigens should also be useful as immunogens for vaccination.

Materials and methods

Expression constructs

The GFP-CP fusion vector CXI (a kind gift from S. Santa-Cruz, Horticulture Research International (HRI), East Malling, United Kingdom) contains a GFP-2A-CP gene fusion under the transcriptional control of a subgenomic CP promoter (Fig. 1) [9]. This was modified by replacing the 2A and GFP sequences with BNYVV sequences, resulting in the expression of BNYVV epitopes as 5'-end in-frame CP fusions (Fig. 1). BNYVV epitope sequences were amplified using primers that incorporated restriction sites suitable for cloning into CXI (Table 1). Part of the 5'-end of the PVX CP was also amplified using primers incorporating an *EagI* restriction site and an additional coding sequence for the respective foreign BNYVV epitopes (Table 1). Following amplification, the PCR products were cloned into the pCR2.1 vector (Invitrogen), released using restriction enzymes *EagI* and *SpeI*, and inserted into the CXI vector, which had been prepared using the same enzymes. This produced recombinant vectors pPVX^{ep4}CP, pPVX^{ep6}CP and pPVX^{ep4ep6}CP, respectively

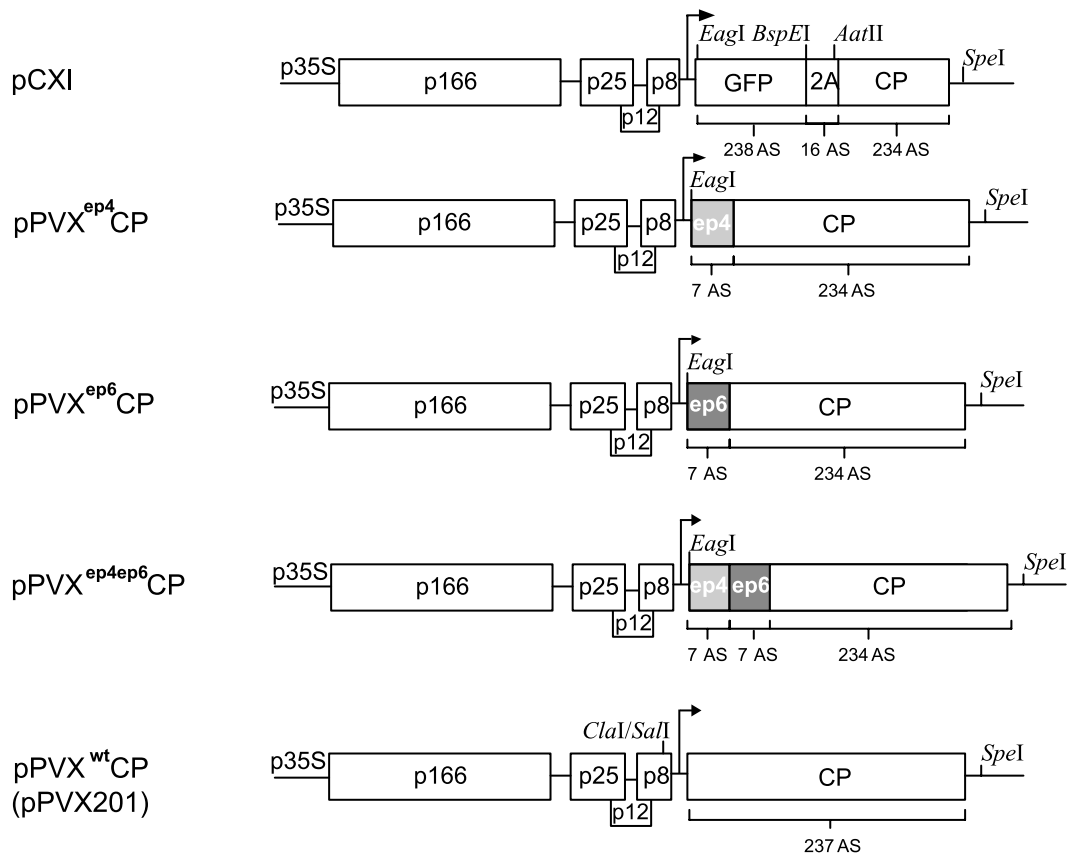


Fig. 1. Structure of PVX-derived vectors used for expression of wild type and recombinant coat proteins. Major features include the RNA-dependent RNA polymerase gene (166K), the triple gene block (25K, 12K, 8K), the virus coat protein gene (*CP*), the CaMV 35S promoter (*p35S*) and the additional BNYVV epitope sequences (*ep4* and *ep6*)

Table 1. Oligonucleotide primers used for preparative PCR and sequencing. *EagI* restriction sites are underlined

Primer name	Nucleotide sequence (5'–3')
CX-ep4	5'- <u>AAACGGCCGATGAGT</u> AGAACAAAGCCCACCAGGACAACCCGCGAGCACA ACACAGC-3'
CX-ep6	5'- <u>AAACGGCCGATGAGT</u> AGCGCTAACGTCAGAAGAGACCCCGCGAGCACA ACACAGC-3'
CX-ep4ep6	5'- <u>AAACGGCCGATGAGT</u> AGAACAAAGCCCACCAGGACAAAGCGCTAACGTCAGAA GAGACCCCGCGAGCACAACACAGC-3'
CX1	5'-TTGAAGAAGTCGAATGCAGC-3'
CX2	5'-CTAGATGCAGAAACCATAAG-3'
CX3	5'-ATAGCAGTCATTAGCACTTC-3'
CX4	5'-CGGGCTGTACTAAAGAAATC-3'

(Fig. 1). Plasmid DNA used for infection was amplified in the general *Escherichia coli* cloning strain DH5 α .

Plant infection and PVX purification

Nicotiana benthamiana plants were inoculated with plasmid pPVX201 (a kind gift from D. Baulcombe, The Sainsbury Laboratory, Norwich, United Kingdom) which carries the wild-type PVX CP gene [3]. Further inoculations were carried out with plasmids pPVX^{ep4}CP, pPVX^{ep6}CP and pPVX^{ep4ep6}CP, containing recombinant PVX genomes incorporating BNYVV epitopes (Fig. 1). Inoculation was achieved by gentle abrasion of the surfaces of three leaves per plant with carborundum and 5–10 μ g of plasmid DNA. After inoculation, the surface of the leaves was rinsed with water in order to remove carborundum and excess DNA. Plants were maintained with a 16-h photoperiod (25,000–30,000 lux, 25 °C/20 °C temperature regime) and 60% humidity. Upon full systemic infection (about 14 days after inoculation) the correct expression of the foreign constructs was confirmed by enzyme-linked immunosorbent assay (ELISA), western blot and electron microscopy as discussed below.

PVX particles were purified according to a modified method of Koenig et al. [17]. Briefly, 50 g of leaf material stored at –80 °C was homogenized in two volumes of ice-cold 0.5 M boric acid solution (pH 7.8) in a Waring blender. After filtration through three layers of gauze, the pH was adjusted to 6.5 with HCl, and 0.2% (w/v) ascorbic acid and 0.2% (w/v) sodium sulfite were added. Cellular debris were removed by centrifugation (5500 \times g for 20 min at 4 °C) and the supernatant was supplemented with 0.15 volumes of 0.5% silver nitrate and left at room temperature in the dark for 2–3 h. The solution was centrifuged as above and the second-stage supernatant was supplemented with one fifth of its volume of 1 M NaCl, 20% PEG in 0.5 M boric acid pH 7.8. Virus particles were sedimented by overnight incubation at 4 °C. The virus particles were pelleted by centrifugation (8000 \times g for 30 min at 4 °C) and resuspended in 10 ml 0.5 M boric acid buffer (pH 7.8) containing 0.5 M urea and 0.1% β -mercaptoethanol. The solution was purified by centrifugation (8000 \times g for 30 min at 4 °C) and the supernatant, containing the virus, was loaded onto a 30% sucrose cushion. After 150 min centrifugation at 72,500 \times g (4 °C), the virus pellet was dissolved in 5 ml 0.5 M boric acid buffer (pH 7.8). The supernatant was clarified by centrifugation for 15 min at 6000 \times g (4 °C), loaded onto a sucrose density gradient (10–45%, w/v in 0.5 M boric acid buffer pH 7.8) and centrifuged at 90,000 \times g for 1 h at 4 °C using a Beckman SW28

rotor. Sucrose gradient fractions containing PVX particles, verified by sodium dodecylsulfate polyacrylamide gel electrophoresis (SDS-PAGE), were combined and dialyzed against PBS (pH 7.4). The concentration of virus was calculated using the PVX extinction coefficient (2.97) for the extinction values at 260 nm.

Enzyme-linked immunosorbent assay (ELISA)

The presence of the foreign epitope on recombinant PVX particles in the sap of infected plants was confirmed using an enzyme-linked immunosorbent assay. Microtiter plates (Greiner, Germany) were coated for 2 h at 37 °C with 100 µl of anti-PVX polyclonal antibody (pAb) (DSMZ, Braunschweig, Germany) diluted 1:2000 in carbonate buffer (pH 9.6). After washing with phosphate buffered saline plus 0.05% (v/v) Tween-20 pH 7.4 (PBST), the plates were blocked with 2% skimmed milk powder in PBS for 1 h at 37 °C. Infected plant tissue was harvested and homogenized in PBS. Extracts were then added (100 µl per well) and plates were incubated at 37 °C for 2 h. After washing the plates, recombinant PVX particles were detected using monoclonal antibody (mAb) SCR84 or SCR86 (kindly provided by L. Torrance, SCRI, Dundee, United Kingdom) followed by incubation with an alkaline phosphatase (AP)-conjugated goat anti-mouse mAb (Dianova). Signal detection was achieved using *p*-nitrophenylphosphate (Sigma) as the substrate at a concentration of 1 mg ml⁻¹ in substrate buffer (50 mM Tris-HCl, pH 9.6, containing 150 mM NaCl and 2 mM MgCl₂). The plate absorbance was read with an ELISA reader (Spectra Max 340) at 405 nm.

SDS-PAGE and western blot analysis

The protein profiles of crude sap extracts (1:3 dilution) or purified virus preparations, were analyzed by SDS-PAGE [22] using a 12% resolving gel and a 4% stacking gel, with samples incubated at 100 °C for 5 min in Laemmli loading buffer prior to loading. The separated proteins were transferred onto ImmobilonP nitrocellulose transfer membranes (Millipore) using the Mini Trans-Blot Cell (BioRad). Western blot analysis was performed using mAb SCR84, mAb SCR86 or pAb PVX as primary antibodies and AP-conjugated goat anti mouse or goat anti rabbit mAb (Dianova) as secondary antibodies.

Electron microscopy

Electron microscope grids with a Pioloform-carbon support film were floated for 5 min on drops of virus-infected plant sap, washed and blocked afterwards with 0.1% bovine serum albumin (BSA; Sigma) in phosphate buffer (PB, pH 7.2). Monoclonal antibodies recognizing BNYVV epitopes (SCR84, SCR86) were bound to the sample by incubating the grid in a PBST dilution of the appropriate antibody for 1.5 h. After washing (PBST), samples were decorated with gold-labeled goat anti-mouse IgG (15 nm, 10 nm or 5 nm gold particles, Biocell, United Kingdom) by floating the grid on a 1:50 PBST dilution of the antibody for 2 h or overnight. The grids were extensively washed (once with PBST, and twice with distilled H₂O) and the sample stained with five drops of 1% uranyl acetate. Electron microscopy was carried out using a Zeiss 906 transmission electron microscope.

Sequencing

Sequencing was performed using an ABI Prism 3700 DNA Analyzer (Applied Biosystems), primers CX2 and CX4 (Table 1), and the ABI Prism BigDye Terminator v3.0 Ready Reaction Cycle Sequencing Kit (Applied Biosystems).

Results

Production and detection of hybrid viruses

Chimeric CP molecules were produced by adding nucleotide sequences encoding different BNYVV epitopes to the 5'-end of the PVX coat protein gene, resulting in the three recombinant PVX constructs shown in Fig. 1 (PVX^{ep4}CP, PVX^{ep6}CP, PVX^{ep4ep6}CP). Plasmid DNA containing the full-length wild type cDNA of PVX (pPVX201; Fig. 1) or PVX with modified CP sequences (PVX^{ep4}CP, PVX^{ep6}CP, PVX^{ep4ep6}CP; Fig. 1) was used to inoculate *N. benthamiana* plants. One to two weeks post inoculation (p.i.), all plants showed systemic infections and symptoms identical to those of plants infected with wild-type PVX (Fig. 2). Systemically infected leaves were collected, and the expression of foreign sequences was verified by ELISA (data not shown) and western blot (Fig. 3).

Virus particles were purified from the systemically infected leaves and yields of up to 115 μg per gram of leaves were achieved (data not shown). Total protein extracts from infected leaves as well as purified virus preparations were separated by SDS-PAGE. As shown in Fig. 3, the wild type CP and the three recombinant

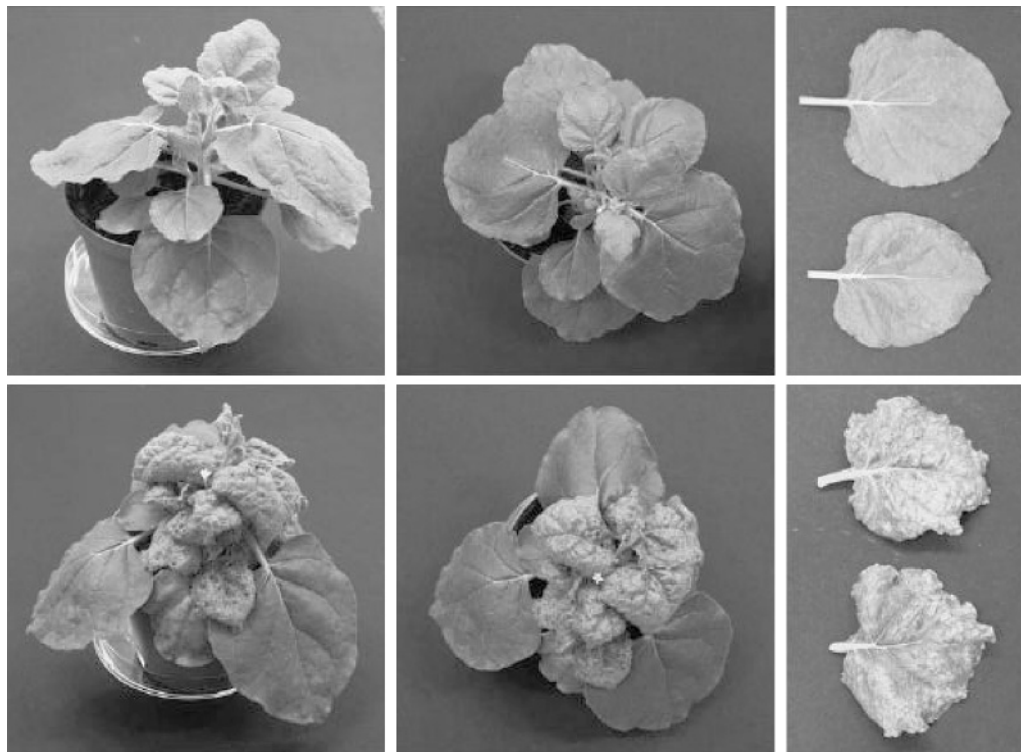


Fig. 2. PVX infection of *N. benthamiana* plants, showing whole plants viewed from the side and the top, and two sample leaves. Top row: Non-infected control plant. Bottom row: Infected plant 14 days after inoculation of three leaves with pPVX^{ep6}CP (5–10 μg per leaf)

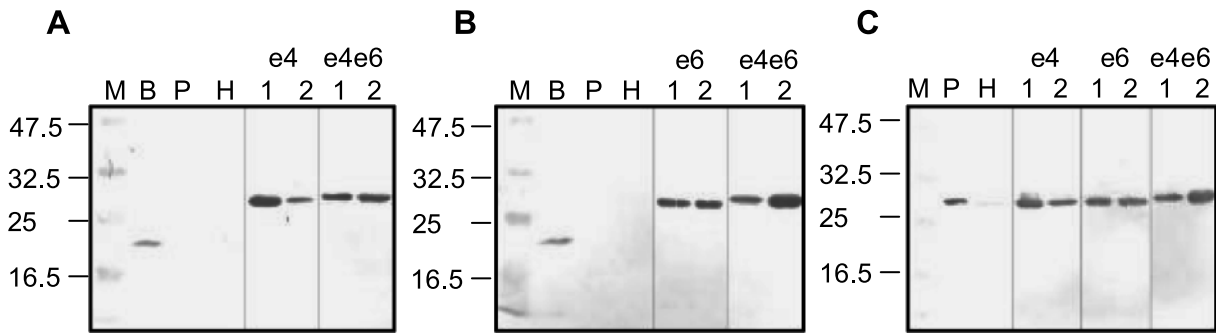


Fig. 3. Western blot analysis of PVX^{wt}CP and PVX^{ep}CP infected *N. benthamiana* plant extracts (24 days p.i.) diluted 1:3 in PBS. Antibody detection: **A:** mAb SCR86 + goat anti-mouse-alkaline phosphatase (GAM^{AP}). **B:** mAb SCR84 + GAM^{AP}. **C:** pAb PVX + goat anti-rabbit-alkaline phosphatase (GAR^{AP}). Lane designations: *M* = prestained broad range marker (New England Biolabs); *B* = BNYVV infected *Chenopodium quinoa* plant sap; *H* = non-infected control plant; *P* = PVX^{wt}CP control; *e4* = PVX^{ep4}CP; *e6* = PVX^{ep6}CP; *e4e6* = PVX^{ep4ep6}CP; *1, 2*: two different PVX^{ep}CP infected plants

CPs migrated close to their calculated positions with molecular masses of 25.05, 25.77, 25.85 and 26.57 kDa, respectively.

Immunogold labeling of hybrid PVX particles

Immunogold electron microscopy using monoclonal antibodies specific for each of the two BNYVV epitopes revealed binding to the surface of purified PVX particles as shown in Fig. 4A–D. This confirmed that the foreign epitopes were accessible and exposed on the surface of the PVX virions. Furthermore, viruses expressing both epitopes in a tandem array construct were detected by both antibodies either singly or in combination (Fig. 5A–D). This showed that the tandemly arranged epitopes were individually accessible on the surface of the particles, and could also be accessed simultaneously by different antibodies labeled with different-sized gold particles (Fig. 5C and D). Double labeling of the PVX^{ep4ep6}CP particles was accomplished by incubation with the primary ep6-specific mAb and a 5 nm gold-labeled secondary mAb followed by incubation with a mixture of the primary ep4-specific mAb and a 10 nm-gold labeled secondary mAb. The possibility that double labeling with different-sized gold grains was obtained by exchange of secondary antibodies was excluded by the results of a mixed infection described in the following section.

PVX infection with a mixture of two different DNA constructs

To establish the number of epitopes presented in mixed infections, *N. benthamiana* plants were inoculated with a 1:1 mixture of constructs pPVX^{ep4}CP and pPVX^{ep6}CP. In such infections, ^{ep6}CP and ^{ep4}CP could be synthesized in the same cell and virus particles could be formed with both epitopes presented on

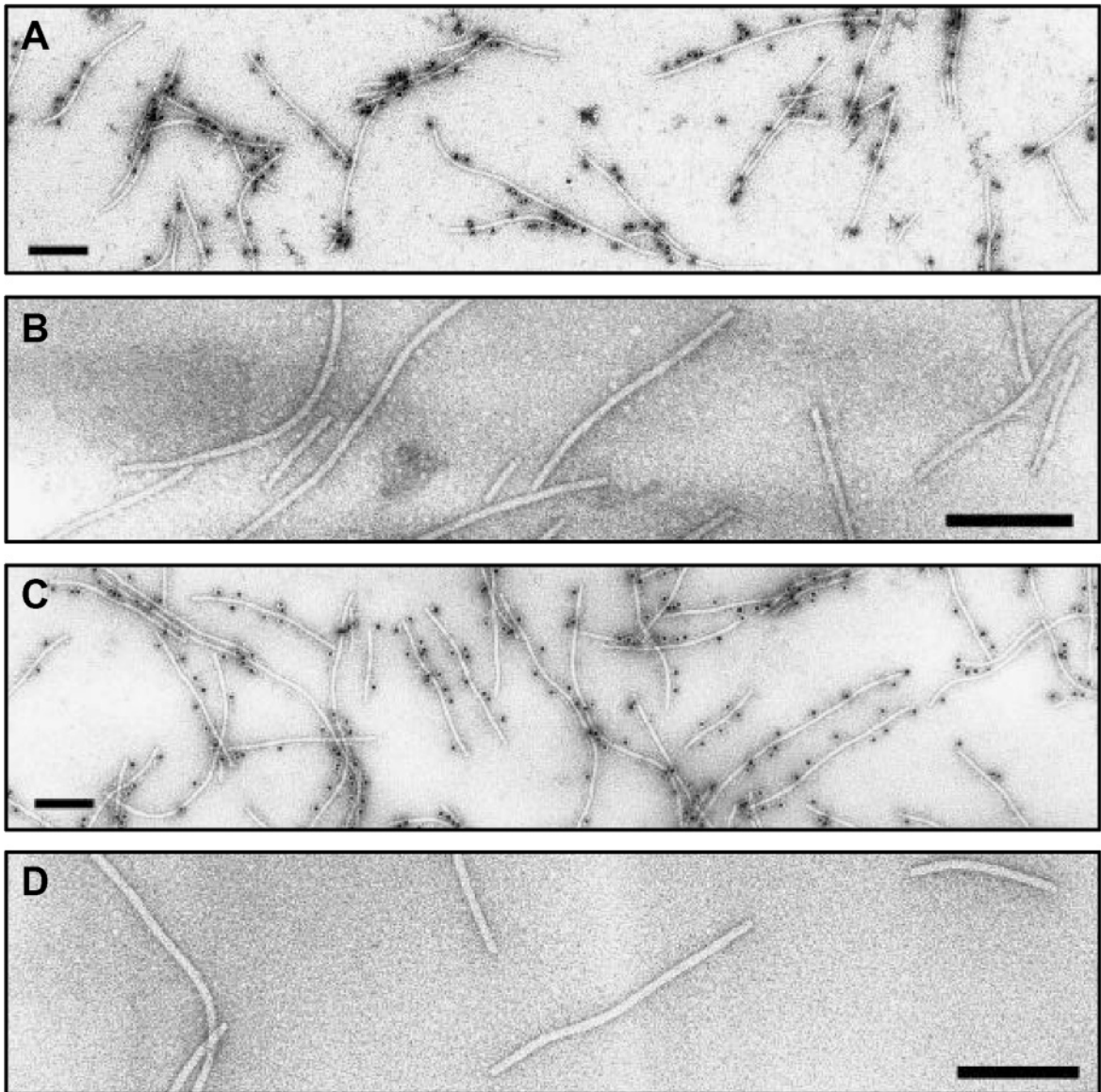


Fig. 4. Immunogold labeling of PVX^{ep}CP particles from infected *N. benthamiana* plants
A: PVX^{ep4}CP decorated with mAb SCR86 specific for BNYVV epitope 4 (1:5 dilution in PBST), detection with 15 nm goat anti-rabbit (GAR) gold conjugate (1:50 dilution in PBST).
B: Negative control, PVX^{ep4}CP particles, primary mAb SCR84 specific for BNYVV epitope 6 (1:10 dilution in PBST), secondary pAb 15 nm GAR gold conjugate (1:50 dilution in PBST).
C: PVX^{ep6}CP particles decorated with mAb SCR84 specific for BNYVV epitope 6 (1:10 dilution in PBST), detection with 15 nm gold conjugate GAR (1:50 dilution in PBST).
D: Negative control, PVX^{ep6}CP particle, primary mAb 3H12 specific for BNYVV epitope 4 (1:10 dilution in PBST), secondary pAb 15 nm gold conjugate GAR (1:50 dilution in PBST).
 Bar = 200 nm

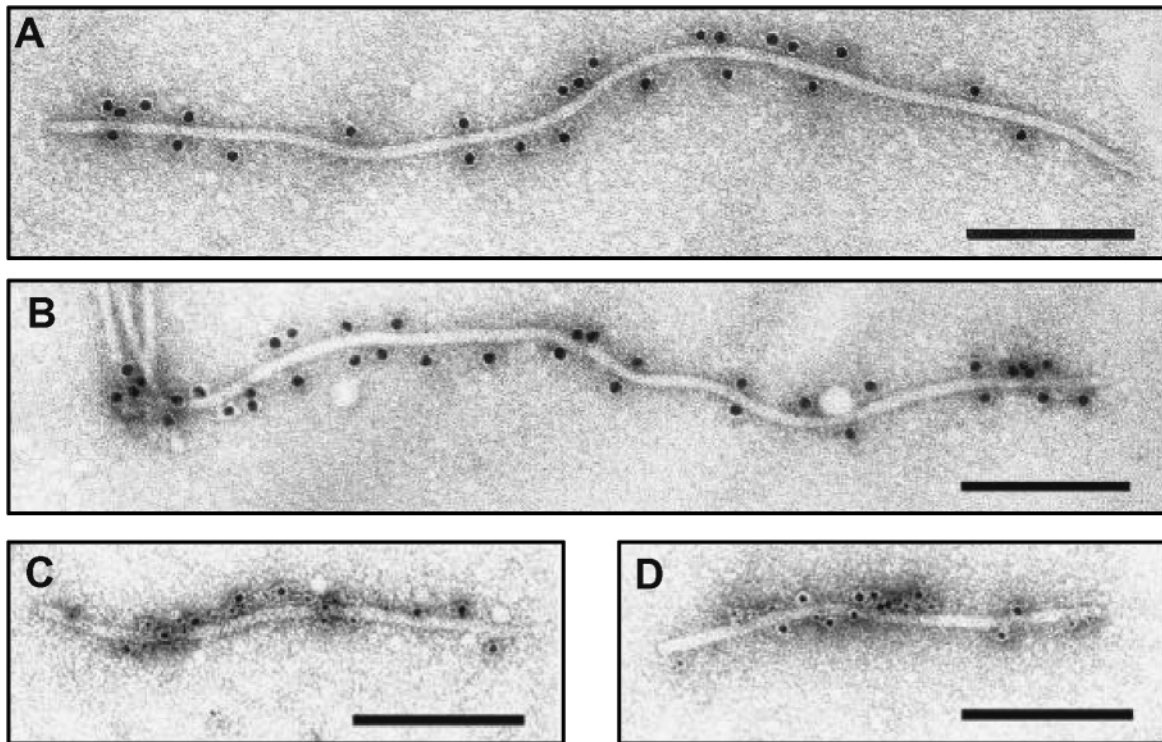


Fig. 5. Immunogold labeling of PVX^{ep4p6}CP-particles (tandem epitope) from *N. benthamiana*. **A:** PVX^{ep4p6}CP-particles decorated with mAb SCR86 specific for BNYVV epitope 4 (1:5 dilution in PBST), detection with goat anti-rabbit (GAR) 15 nm gold conjugate (1:50 dilution in PBST). **B:** PVX^{ep4p6}CP particles decorated with mAb SCR84 specific for BNYVV epitope 6 (1:10 dilution in PBST), detection with GAR 15 nm gold conjugate (1:50 dilution in PBST). **C and D:** PVX^{ep4p6}CP particles decorated with mAb SCR84 (1:10 dilution in PBST) and goat anti-mouse (GAM) 5 nm gold conjugate (1:50 dilution in PBST) and mAb SCR86 (1:5 dilution in PBST) saturated with GAM 10 nm gold (1:50 dilution in PBST). Bar = 200 nm

the surface. Purified PVX particles from mixed infections were initially incubated with the primary mAb specific for one of the two epitopes, and binding was detected with a 15 nm gold-labeled secondary antibody. This analysis showed that such preparations contained both gold-labeled virions and a fraction of completely undecorated virions (Fig. 6A and B). Double labeling was then carried out as described in the previous section. Incubation with the primary ep6-specific mAb and a 5 nm gold-labeled secondary mAb was followed by incubation with a mixture of the primary ep4-specific mAb and a 10 nm-gold labeled secondary mAb. As shown in Fig. 6D, PVX virions were identified that were labeled either with 5 or with 10 nm gold particles alone, but there were also virions that were labeled simultaneously with gold particles of both sizes (Fig. 6C and E).

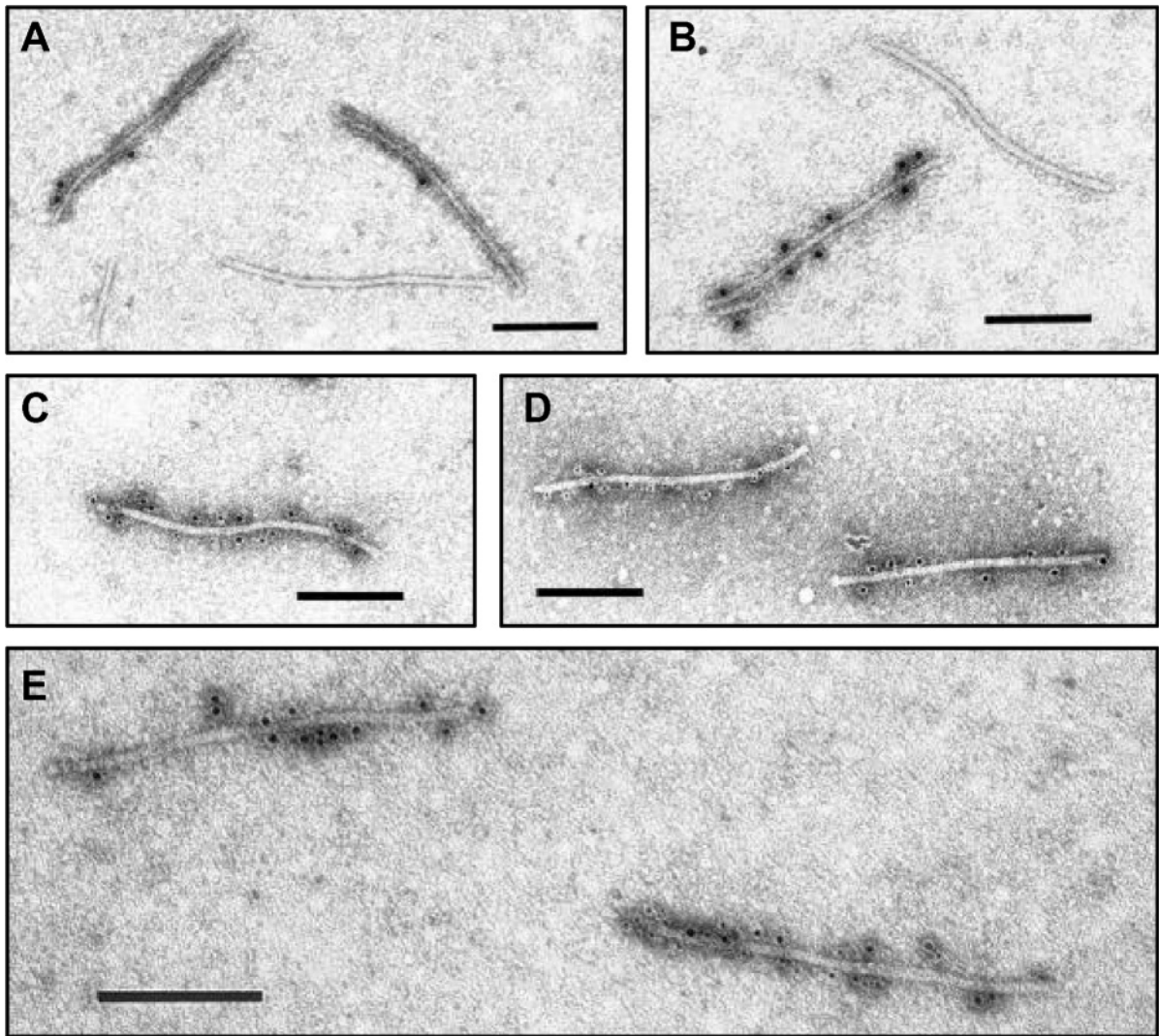


Fig. 6. Immunogold labeling of PVX^{ep4}CP and PVX^{ep6}CP particles from mixed infections. **A:** PVX^{ep4}CP + ^{ep6}CP particles decorated with mAb SCR86 specific for BNYVV epitope 4 (1:5 dilution in PBST) detected with goat anti-rabbit (GAR) 15 nm gold conjugate (1:50 dilution in PBST). **B:** PVX^{ep4}CP + ^{ep6}CP-particles decorated with mAb SCR84 against BNYVV epitope 6 (1:10 dilution in PBST), detected with GAR 15 nm gold conjugate (1:50 dilution in PBST). **C–E:** PVX^{ep4}CP + ^{ep6}CP-particles decorated first with mAb SCR84 (1:60 dilution in PBST) and goat anti-mouse (GAM) 5 nm gold conjugate (1:50 dilution in PBST) and then with mAb SCR86 (1:5 dilution in PBST) saturated with GAM 10 nm gold conjugate (1:50 dilution in PBST). Bar = 200 nm

Discussion

The use of plants as bioreactors for the synthesis of proteins has the potential to reduce the cost of protein therapeutics, and is especially useful for vaccine production and delivery [36]. Vaccines can be expressed as full length polypeptides

in transgenic plants, but epitope presentation on the surface of virus particles is a useful alternative which is quicker and does not require genetic transformation [23]. PVX is a good candidate carrier molecule for foreign epitopes because it self-assembles in an ordered manner, and accumulates to high levels in infected tissues. It also lacks the size and packaging constraints of icosahedral viruses like CPMV.

In this study we have used PVX vectors to express BNYVV epitopes fused to the N-terminus of the PVX coat protein. The seven-amino-acid epitopes ep4 and ep6 [39], as well as the 14-amino-acid tandem epitope ep4ep6, were expressed successfully in systemically infected *N. benthamiana* plants and were exposed on the surface of the recombinant PVX particles. Therefore, these additional sequences appear not to interfere with virus assembly nor with cell-to-cell movement, even though the PVX CP is known to be required for systemic spread during natural infections [34].

The assembly of recombinant PVX particles containing a six-amino-acid HIV epitope fused to the N-terminus of the CP has been reported [24]. To the best of our knowledge, however, we are the first to demonstrate the assembly of virions displaying tandem epitopes and the first to demonstrate the display of multiple epitopes on single virus particles following mixed infections. In each case, we have shown that both epitopes can be accessed and recognized by their cognate antibodies either individually or simultaneously, suggesting that either strategy could be useful for the simultaneous presentation of multiple antigens in vaccination programs. While recombinant viruses expressing the tandem construct display both epitopes, mixed infections result in the production of virions displaying ep4 alone, ep6 alone or both epitopes (Fig. 6).

The first experiments with PVX CP fusion proteins established that PVX assembly is blocked by the presence of foreign proteins [9]. For this reason, the FMDV 2A peptide [29] has been inserted between the heterologous sequence and the CP open reading frame in previous studies to allow a degree of co-translational cleavage and the production of some wild type CP subunits [15, 30]. Recently, it has been shown that this process is not proteolytic, but instead involves ribosomal skipping on the RNA template [11, 12]. Consequently the production of recombinant and wild type coat protein is accomplished, infectious particles are formed and proteins of 8.5 to 46 kD can be presented on the virus surface [9, 25, 32]. One disadvantage of this “overcoat” principle, however, is the unpredictable ratio of wild type to recombinant CP, which may vary between 1:1 and 1:5 [21, 25]. If peptide-presenting viruses are to be used as standards in diagnosis or vaccination, the presentation of epitopes on all CP subunits is advantageous, since this assures an exactly defined and high concentration of desired antigenic sites.

Our ultimate goal is to produce protein standards presenting a) several different epitopes concurrently or b) several short epitopes, either combining epitopes from different viruses in one standard or using different epitopes from the same virus to design a standard recognized by many different antibodies available against this virus. The latter would be particularly advantageous for simplification and reduction of standard production costs. Furthermore, the presentation of several epitopes on plant virions would also be very interesting for vaccine development,

since antibodies that are produced against a single pathogen epitope are often not sufficient to provide complete and long lasting protection against an infection [1, 20, 31].

The size of the longest peptide presented on TMV virions was 23 residues [5] and on PPV and CPMV particles up to 30 and 50 residues, respectively [13, 26, 28]. Although this capacity would potentially allow for the presentation of several shorter epitopes, there have been only few reports about the presentation of multiple peptides on these and other plant virions [13, 14, 38]. None of these studies addressed the accessibility of the individual epitopes, instead their efficiency to elicit protective antibodies was investigated. The construction of a CPMV chimera with an insert of *Pseudomonas* epitopes 18 and 10 in tandem within the L or S subunit of the virus elicited antibodies that were directed only toward the epitope 10 or 18, respectively [14]. Obviously the CPMV CP subunit into which the peptide was inserted greatly influenced the nature of the antibodies elicited [6]. PPV chimeric virions presenting different forms of an antigenic peptide (single and tandem repetition) from the VP2 capsid protein of canine parvovirus (CPV) showed a high antigenicity [13]. However no substantial difference was observed in the quality of immune response induced by single or tandem presentation of the CPV peptide.

The expression of peptides as coat protein fusions can lead to the production of large amounts of antigen. However, in order to prevent any undesirable changes in the aa sequence of the epitope through mutations, the recombinant particles should be isolated from infected plants after a relatively short period of time (2–3 weeks). This helps to prevent the accumulation of mutated viruses in which the epitope has been altered or even deleted, as observed after serial passages in *N. benthamiana* (Uhde K. & Commandeur U., unpublished results). Therefore, it is important to verify the presence and accessibility of foreign epitope sequences using appropriate antibodies.

Acknowledgements

We thank Martina Koch and Oliver Krohn for excellent technical assistance, Dr. D.-E. Lesemann (BBA, Braunschweig) for his help with electron microscopy and for critical reading of this paper, Christina Maas for development of the electron microscopic pictures and Dr. S.S. Cruz for supplying the CXI vector. This work was funded by the EU project SMTP4-CT98-2246.

References

1. Awram P, Gardner RC, Forster RL, Bellamy AR (2002) The potential of plant viral vectors and transgenic plants for subunit vaccine production. *Adv Virus Res* 58: 81–124
2. Baratova LA, Grebenshchikov NI, Shishkov AV, Kashirin IA, Radavsky JL, Jarvekulg L, Saarma M (1992) The topography of the surface of potato virus X: tritium planigraphy and immunological analysis. *J Gen Virol* 73: 229–235
3. Baulcombe D, Chapman S, Santa Cruz S (1995) Jellyfish green fluorescent protein as a reporter for virus infections. *Plant J* 7: 1045–1053
4. Baulcombe DC (1999) Fast forward genetics based on virus-induced gene silencing. *Curr Opin Plant Biol* 2: 109–113

5. Bendahmane M, Koo M, Karrer E, Beachy RN (1999) Display of epitopes on the surface of tobacco mosaic virus: impact of charge and isoelectric point of the epitope on virus-host interactions. *J Mol Biol* 290: 9–20
6. Brennan FR, Jones TD, Gilleland LB, Bellaby T, Xu F, North PC, Thompson A, Staczek J, Lin T, Johnson JE, Hamilton WD, Gilleland HE Jr (1999) *Pseudomonas aeruginosa* outer-membrane protein F epitopes are highly immunogenic in mice when expressed on a plant virus. *Microbiology* 145: 211–220
7. Brennan FR, Jones TD, Hamilton WD (2001) Cowpea mosaic virus as a vaccine carrier of heterologous antigens. *Mol Biotechnol* 17: 15–26
8. Chapman S, Kavanagh T, Baulcombe D (1992) Potato virus X as a vector for gene expression in plants. *Plant J* 2: 549–557
9. Cruz SS, Chapman S, Roberts AG, Roberts IM, Prior DA, Oparka KJ (1996) Assembly and movement of a plant virus carrying a green fluorescent protein overcoat. *Proc Natl Acad Sci USA* 93: 6286–6290
10. Dalmay T, Hamilton A, Mueller E, Baulcombe DC (2000) Potato virus X amplicons in *Arabidopsis* mediate genetic and epigenetic gene silencing. *Plant Cell* 12: 369–379
11. Donnelly ML, Hughes LE, Luke G, Mendoza H, ten Dam E, Gani D, Ryan MD (2001) The ‘cleavage’ activities of foot-and-mouth disease virus 2A site-directed mutants and naturally occurring ‘2A-like’ sequences. *J Gen Virol* 82: 1027–1041
12. Donnelly ML, Luke G, Mehrotra A, Li X, Hughes LE, Gani D, Ryan MD (2001) Analysis of the aphthovirus 2A/2B polyprotein ‘cleavage’ mechanism indicates not a proteolytic reaction, but a novel translational effect: a putative ribosomal ‘skip’. *J Gen Virol* 82: 1013–1025
13. Fernandez-Fernandez MR, Martinez-Torrecedrada JL, Casal JI, Garcia JA (1998) Development of an antigen presentation system based on plum pox potyvirus. *FEBS Lett* 427: 229–235
14. Gilleland HE, Gilleland LB, Staczek J, Harty RN, Garcia-Sastre A, Palese P, Brennan FR, Hamilton WD, Bendahmane M, Beachy RN (2000) Chimeric animal and plant viruses expressing epitopes of outer membrane protein F as a combined vaccine against *Pseudomonas aeruginosa* lung infection. *FEMS Immunol Med Microbiol* 27: 291–297
15. Halpin C, Cooke SE, Barakate A, El Amrani A, Ryan MD (1999) Self-processing 2A-polyproteins – a system for co-ordinate expression of multiple proteins in transgenic plants. *Plant J* 17: 453–459
16. Hendy S, Chen ZC, Barker H, Santa Cruz S, Chapman S, Torrance L, Cockburn W, Whitlam GC (1999) Rapid production of single-chain Fv fragments in plants using a potato virus X episomal vector. *J Immunol Methods* 231: 137–146
17. Koenig R, Tremaine JH, Shepard JF (1978) In situ degradation of the protein chain of potato virus X at the N- and C-termini. *J Gen Virol* 38: 329–337
18. Koenig R, Torrance L (1986) Antigenic analysis of potato virus X by means of monoclonal antibodies. *J Gen Virol* 67: 2145–2151
19. Koenig R, Lesemann DE (1989) Potato virus X, potexvirus group. In: Murrant AF, Harrison BD (eds) *AAB descriptions of plant viruses* No. 354, Association of Applied Biologists, Warwick, pp 1–5
20. Koprowski H, Yusibov V (2001) The green revolution: plants as heterologous expression vectors. *Vaccine* 19: 2735–2741
21. Lacomme CLS, Wilson MA (1998) Genetic engineering and the expression of foreign peptides or proteins with plant virus-based vectors. In: Setlow JK (ed) *Genetic engineering*, vol 20. Plenum Press, New York, pp 225–237
22. Laemmli UK (1970) Cleavage of structural proteins during the assembly of the head of bacteriophage T4. *Nature* 227: 680–685

23. Lomonossoff GP, Hamilton WD (1999) Cowpea mosaic virus-based vaccines. *Curr Top Microbiol Immunol* 240: 177–189
24. Marusic C, Rizza P, Lattanzi L, Mancini C, Spada M, Belardelli F, Benvenuto E, Capone I (2001) Chimeric plant virus particles as immunogens for inducing murine and human immune responses against human immunodeficiency virus type 1. *J Virol* 75: 8434–8439
25. O'Brien GJ, Bryant CJ, Voogd C, Greenberg HB, Gardner RC, Bellamy AR (2000) Rotavirus VP6 expressed by PVX vectors in *Nicotiana benthamiana* coats PVX rods and also assembles into virus like particles. *Virology* 270: 444–453
26. Porta C, Spall VE, Loveland J, Johnson JE, Barker PJ, Lomonossoff GP (1994) Development of cowpea mosaic virus as a high-yielding system for the presentation of foreign peptides. *Virology* 202: 949–955
27. Porta C, Lomonossoff GP (2002) Viruses as vectors for the expression of foreign sequences in plants. *Biotechnol Genet Eng Rev* 19: 245–291
28. Porta C, Spall VE, Findlay KC, Gergerich RC, Farrance CE, Lomonossoff GP (2003) Cowpea mosaic virus-based chimaeras. Effects of inserted peptides on the phenotype, host range, and transmissibility of the modified viruses. *Virology* 310: 50–63
29. Ryan MD, King AM, Thomas GP (1991) Cleavage of foot-and-mouth disease virus polyprotein is mediated by residues located within a 19 amino acid sequence. *J Gen Virol* 72: 2727–2732
30. Ryan MD, Drew J (1994) Foot-and-mouth disease virus 2A oligopeptide mediated cleavage of an artificial polyprotein. *EMBO J* 13: 928–933
31. Sette A, Livingston B, McKinney D, Appella E, Fikes J, Sidney J, Newman M, Chesnut R (2001) The development of multi-epitope vaccines: epitope identification, vaccine design and clinical evaluation. *Biologicals* 29: 271–276
32. Smolenska L, Roberts IM, Learmonth D, Porter AJ, Harris WJ, Wilson TM, Santa Cruz S (1998) Production of a functional single chain antibody attached to the surface of a plant virus. *FEBS Lett* 441: 379–382
33. Sugiyama Y, Hamamoto H, Takemoto S, Watanabe Y, Okada Y (1995) Systemic production of foreign peptides on the particle surface of tobacco mosaic virus. *FEBS Lett* 359: 247–250
34. Tamai A, Meshi T (2001) Cell-to-cell movement of Potato virus X: the role of p12 and p8 encoded by the second and third open reading frames of the triple gene block. *Mol Plant Microbe Interact* 14: 1158–1167
35. Turpen TH, Reinl SJ, Charoenvit Y, Hoffman SL, Fallarme V, Grill LK (1995) Malarial epitopes expressed on the surface of recombinant tobacco mosaic virus. *Biotechnology (NY)* 13: 53–57
36. Twyman RM, Stoger E, Schillberg S, Christou P, Fischer R (2003) Molecular farming in plants: host systems and expression technology. *Trends Biotechnol* 21: 570–578
37. Usha R, Rohll JB, Spall VE, Shanks M, Maule AJ, Johnson JE, Lomonossoff GP (1993) Expression of an animal virus antigenic site on the surface of a plant virus particle. *Virology* 197: 366–374
38. Yusibov V, Hooper DC, Spitsin SV, Fleysh N, Kean RB, Mikheeva T, Deka D, Karasev A, Cox S, Randall J, Koprowski H (2002) Expression in plants and immunogenicity of plant virus-based experimental rabies vaccine. *Vaccine* 20: 3155–3164
39. Commandeur U, Koenig R, Mantuffel R, Torrance L, Lüddecke P, Frank R (1994) Location, size, and complexity of epitopes on the coat protein of beet necrotic yellow vein virus studied by means of synthetic overlapping peptides. *Virology* 198: 282–287

Author's address: Dr. Ulrich Commandeur, RWTH Aachen, Institute for Molecular Biotechnology, Worringerweg 1, 52074 Aachen, Germany; e-mail: commandeur@molbiotech.rwth-aachen.de