Monoclonal Antibodies Detect a Single Amino Acid Difference Between the Coat Proteins of Soilborne Wheat Mosaic Virus Isolates: Implications for Virus Structure

Jianping Chen, Lesley Torrance, Graham H. Cowan, Stuart A. MacFarlane, Gerald Stubbs, and T. Michael A. Wilson

First author: Zhejiang Academy of Agricultural Sciences, Hangzhou 310021, People's Republic of China; second, third, fourth, and sixth authors: Department of Virology, Scottish Crop Research Institute, Invergowrie, Dundee DD2 5DA, United Kingdom; and fifth author: Department of Molecular Biology, Vanderbilt University, Nashville, TN 37235. Accepted for publication 9 December 1996.

ABSTRACT


Four monoclonal antibodies (MAbs) were prepared against an isolate of soilborne wheat mosaic furovirus from Oklahoma (SBWMV Okl-7). Three MAbs had different reactivities in tests on SBWMV isolates from Nebraska (Lab1), France, and Japan. One MAb (SCR 133) also reacted with oat golden stripe furovirus. None of the MAbs cross-reacted with other rod-shaped viruses including beet necrotic yellow vein furovirus, potato mop-top furovirus, and tobacco rattle tobravirus. Sequence analysis of nucleotides between 334 and 1,000 of RNA 2, the region that encodes the coat protein (CP) and the first 44 amino acids of a readthrough domain of the CP, revealed up to 27 base changes between geographically separate isolates of SBWMV. The results indicate that the MAbs detect changes in amino acids near the N- or C-termini of the CPs of the different isolates. Two further single amino acid changes were found at the beginning of the readthrough domain of the CP. Some of these amino acid changes could be discriminated by MAbs SCR 132, SCR 133, and SCR 134. Peptide scanning (Pepscan) analysis indicated that the epitope recognized by SCR 134 is located near the N-terminus of the CP. SCR 132 was deduced to react with a discontinuous CP epitope near the C-terminus, and SCR 133 reacted with a surface-located continuous epitope also near the C-terminus. Predictions of CP structure from computer-assisted three-dimensional model building, by comparison with the X-ray fiber diffraction structure of tobacco mosaic virus, suggested that the three CP amino acids found to differ between isolates of SBWMV were located near the viral surface and were in regions predicted to be antigenic. Additional keywords: coat protein sequence, ELISA, epitope mapping.

MATERIALS AND METHODS

Virus isolates, maintenance, and purification. Two SBWMV isolates from the United States were used for antiserum production and RNA sequencing. The SBWMV Okl-7 isolate, a stable RNA 2-deletion mutant, was obtained by serial mechanical inoculation (7–9) of a field isolate of SBWMV from Oklahoma (Okl-WT), which was kindly provided by J. L. Sherwood (Oklahoma State University, Stillwater). The SBWMV Lab1 isolate, another RNA 2-deletion mutant derived by successive manual inoculations of a Nebraska field isolate over 2 years (22), was generously provided by Y. Shirako (University of Tokyo). Both isolates were maintained by mechanical inoculation to wheat (Triticum aestivum) cv. Galahad (7). SBWMV was purified from infected wheat leaves by the method of Shirako and Brakke (21).

Two further SBWMV isolates and four other rod-shaped viruses (donors in parentheses) were used in MAb cross-reactivity tests: SBWMV from Japan (SBWMV-J; I. M. Roberts, Scottish Crop Research Institute [SCRI], Dundee, United Kingdom), France (SBWMV-F; M. J. Adams, Institute of Arable Crops Research, Rothamsted, Harpenden, United Kingdom), OGSV (M. J. Adams), potato mop-top virus (PMTV, isolate T [27]), beet necrotic yellow vein virus (BNYVV, strain H3; R. Koenig, Braunschweig, Germany), and tobacco rattle virus (TRV, strain PRN; D. J. Robinson, SCRI).

Polyclonal antiserum production. An antiserum to SBWMV Okl-7 was produced in a New Zealand white rabbit as follows: approximately 50 μg of purified virions emulsified in complete...
Freund’s adjuvant (CFA) were injected intramuscularly (half depo-
osed in each leg) on day 1; 50 µg of virions in incomplete
Freund’s adjuvant (IFA) were injected intramuscularly on day 14;
and 100 µg of virions were injected subcutaneously on day 126.
Antiserum was collected 2 weeks after the final injection. The
titer was 1/128 in microprecipitin tests against purified prepara-
tions of SBWMV Okl-7.

Hybridoma production, screening, and isotyping. The MAbs
secreting cell lines were derived from fusions between spleen
cells from immunized BALB/c mice (taken 3 days after the last
injection) and the mouse myeloma cell line X63/Ag 8.653. Immu-
nization schedules were as follows: fusion 1 (20 to 50 µg of
SBWMV Okl-7 virions per injection) on day 1, intraperitoneal in
IFA; day 28, intraperitoneal in IFA; day 107, intraperitoneal in
IFA; and day 321, intraperitoneal in PBS. The procedures for cell fusion,
cloning by limiting dilution, and maintenance of hybridomas are
described by Torrance (25). About 10 days after fusion, hybrido-
as were screened for specific antibody production by enzyme-
linked immunosorbent assay (ELISA) (see below).

ELISA. ELISAs were done essentially as described by Chen et
al. (9). In antibody-trapped antigen (ATA)-ELISA (24), plant sap
and conjugate were diluted in PBS containing 0.1% (vol/vol)
Tween-20 (PBS-T) and 0.1% (wt/wt) nonfat milk powder. Rabbit
antiserum to SBWMV Okl-7 was used at a dilution of 1:1,000 for
2 h at 37°C. Sap extracts of wheat infected with SBWMV Okl-7
or SBWMV Lab1 were used at a dilution of 1:10 and incubated at
4°C overnight. The undiluted culture supernatant fluids were
incubated at room temperature for 2 h and detected by the addition
of alkaline phosphatase-conjugated goat anti-mouse IgG (Sigma
Chemical Co., St. Louis).

Electron microscopy showed that SBWMV particles were dis-
rupted in 0.05 M sodium carbonate-bicarbonate, pH 10.5 (coating
buffer; J. Chen, unpublished data), and this buffer was used to
prepare sap extracts for plate-trapped antigen (PTA)-ELISA (4,
25). Therefore, PTA-ELISA was used to select MAbs that might
be specific for cryptotopes. Sap extracts from SBWMV Okl-7–
infected leaves (1 g/10 ml) were made in coating buffer and incu-
bated at 4°C overnight in the wells of microtiter plates. After
the removal of sap by rinsing the plates with PBS-T, the plates
were blocked with 5% (wt/vol) nonfat milk powder in PBS-T
for 1 h at room temperature. The procedures for antibody, en-
zyme conjugate, and substrate incubations were the same as for
ATA-ELISA.

Electron microscopy. The methods for immunosorbent elec-
tron microscopy and antibody coating were as described by
Roberts et al. (19).

Molecular biology. The methods for RNA extraction, reverse
transcribe-polymerase chain reaction (RT-PCR), cDNA clon-
ing, and sequencing were those described by Chen et al. (7,9). A
738-nucleotide region extending from RNA 2 genome coordinates
300 to 1,037 and including the entire CP gene was amplified from
first-strand cDNA of SBWMV isolates Okl-7, Lab1, and F using
primers JC28 (5'-ATTATTGCAAAGGTTA-3') and JC41 (5'-
ATGATAGACAGAACCTT-3') (23). The fragments were inserted into
pT7Blue PCR-cloning vector (Novagen, Inc., Madison, WI) and
sequenced using a kit (Amersham International, Amersham,
United Kingdom) according to the manufacturer’s instructions.
Sequence analysis was carried out using the Genetics Computer
Group programs (11). The EMBL sequence accession number for
SBWMV-F is X89078.

Peptide scanning (Pepscan). A set of overlapping octapeptides
was synthesized on cellulose paper using a SPOTs kit (Genosys
Biotechnologies Inc., Cambridge) and tested for reaction with
MAbs by the methods described by Pereira et al. (18). Seven spots
contained amino acids in the sequence 1-MA VNKGYTGYNKEL
from the N-terminus of SBWMV Okl-7 CP. A positive control
spot of sequence STVVVKGN, the epitope for MAb SCR 3 (26),
was also included.

TABLE 1. Some properties of monoclonal antibodies against soilborne wheat mosaic virus (SBWMV)

<table>
<thead>
<tr>
<th>Code name</th>
<th>Isotype</th>
<th>Titer</th>
<th>SBWMV Okl-7</th>
<th>SBWMV Lab1</th>
<th>H</th>
<th>SBWMV Okl-7</th>
<th>SBWMV Lab1</th>
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<td>Cloned lines</td>
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<tr>
<td>SCR 132</td>
<td>IgG&lt;sub&gt;2b&lt;/sub&gt;</td>
<td>262,144&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.921&lt;sup&gt;d&lt;/sup&gt;</td>
<td>1.831</td>
<td>0.067</td>
<td>0.844</td>
<td>0.680</td>
<td>0.032</td>
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<tr>
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<td>1.851</td>
<td>0.271</td>
<td>0.038</td>
<td>0.267</td>
<td>0.139</td>
<td>0.046</td>
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<tr>
<td>SCR 134</td>
<td>IgG&lt;sub&gt;2b&lt;/sub&gt;</td>
<td>64</td>
<td>1.830</td>
<td>0.063</td>
<td>0.046</td>
<td>0.515</td>
<td>0.039</td>
<td>0.023</td>
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<td>SCR 135</td>
<td>IgG&lt;sub&gt;3&lt;/sub&gt;</td>
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<td>0.770</td>
<td>0.568</td>
<td>0.071</td>
<td>0.322</td>
<td>0.279</td>
<td>0.100</td>
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<td>Uncloned lines</td>
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<tr>
<td>2/4</td>
<td>IgM</td>
<td>256</td>
<td>1.944</td>
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<td>0.034</td>
<td>0.469</td>
<td>0.080</td>
<td>0.034</td>
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<tr>
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<td>0.023</td>
<td>0.037</td>
<td>0.037</td>
<td>0.038</td>
</tr>
<tr>
<td>2/6D7</td>
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<td>0.028</td>
<td>0.031</td>
<td>0.035</td>
<td>0.036</td>
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</table>

<sup>a</sup> ATA-ELISA = antibody-trapped antigen–enzyme-linked immunosorbent assay.
<sup>b</sup> PTA-ELISA = plate-trapped antigen–enzyme-linked immunosorbent assay.
<sup>c</sup> + = strongly stained band; ++ = weak reaction; and – = no reaction.
<sup>d</sup> H = healthy plant sample.
<sup>e</sup> Reciprocal of greatest dilution of culture supernatant fluid that reacted with SBWMV Okl-7.
<sup>f</sup> Absorbance value (A<sub>405</sub>) after 1 h of substrate incubation.

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RESULTS

Characterization of MABs. MAbs SCR 135 and SCR 133 were selected from the first fusion and MAbs SCR 132 and SCR 134 were selected from the second fusion. Several other hybridoma lines were obtained from the second fusion that were unstable and stopped secreting antibodies after cloning, e.g., cell lines 2/4, 2/6D9, and 2/6D7. Some properties of the cell lines are given in Table 1.

All seven antibody preparations reacted with the homologous antigen SBWMV Okl-7 in A TA-ELISA, whereas only SCR 132, SCR 133 (weak), and SCR 135 reacted with SBWMV Lab1 (Table 1). A similar result was obtained in PTA-ELISA in which SCR 132, SCR 133, and SCR 135 reacted with both SBWMV Okl-7 and SBWMV Lab1, but SCR 134 and 2/4 reacted strongly only with SBWMV Okl-7. The two other uncloned lines did not work in PTA-ELISA (Table 1). Therefore, antibodies secreted by five out of the seven hybridoma lines obtained discriminated between the two American SBWMV isolates in ATA-ELISA.

In immunoblot experiments, the rabbit polyclonal antiserum reacted strongly with the CP and weakly with an approximately 40-kDa band (possibly a dimer of CP) in preparations from SBWMV Okl-7 and SBWMV Lab1 (Fig. 1). Of the seven MAAb preparations, only SCR 135 reacted strongly in immunoblots with the CP of both isolates (Table 1 and Fig. 1). SCR 133 reacted strongly with SBWMV Okl-7 and weakly with SBWMV Lab1, and SCR 134 reacted very weakly with SBWMV Okl-7, but not with SBWMV Lab1. SCR 132 did not react in this test.

Trypsin-treatment of SBWMV particles. Electron microscopy of purified particles of SBWMV Okl-7 after trypsin treatment for 60 min showed that the particles were intact with no evidence of disruption or degradation. However, SDS-PAGE, Ponceau S staining, and immunoblotting of trypsin-treated (5 and 20 min) virus showed that, in addition to intact CP (19 kDa by sequence [23], but of apparent gel mobility of approximately 22

### Table 2. Comparison of the effects of trypsin-treatment of soilborne wheat mosaic furovirus (SBWMV) particles on electron microscopy trapping and coating reactions obtained with monoclonal antibodies SCR 132 and SCR 133

<table>
<thead>
<tr>
<th>SBWMV isolate</th>
<th>Antibody</th>
<th>Experiment</th>
<th>Trypsin&lt;sup&gt;a&lt;/sup&gt;</th>
<th>U&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Trypsin&lt;sup&gt;a&lt;/sup&gt;</th>
<th>U&lt;sup&gt;b&lt;/sup&gt;</th>
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<td>Okl-7</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>PeAb&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Trapping</td>
<td>514&lt;sup&gt;d&lt;/sup&gt;</td>
<td>707</td>
<td>2,718</td>
<td>5,341</td>
<td></td>
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<tr>
<td>SCR 132</td>
<td>Trapping</td>
<td>133</td>
<td>465</td>
<td>15</td>
<td>391</td>
<td></td>
</tr>
<tr>
<td>SCR 133</td>
<td>Trapping</td>
<td>0</td>
<td>22</td>
<td>13</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Coating</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Normal serum</td>
<td>Trapping</td>
<td>0</td>
<td>0</td>
<td>9</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>Coating</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Trypsin = trypsin-treated (60 min) and U = untreated.
<sup>b</sup> SBWMV rabbit polyclonal antiserum.
<sup>c</sup> Number of particles in $4.5 \times 10^{-10}$ m<sup>2</sup>.
<sup>d</sup> For coating experiments, virions were first trapped using polyclonal antibody.
<sup>e</sup> + = Uniform coating along the sides of particles and – = no antibody coating.

![Fig. 1](image1.png)

Fig. 1. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis and immunoblots of healthy wheat leaf sap and purified preparations of soilborne wheat mosaic furovirus (SBWMV) isolates Okl-7 and Lab1. Immunoblots were probed with polyclonal rabbit antiserum (PeAb) or monoclonal antibodies (MABs) SCR 133, SCR 135, SCR 134, or SCR 132 against SBWMV. Protein size markers are shown on the left (K = kDa).

![Fig. 2](image2.png)

Fig. 2. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis of purified soilborne wheat mosaic furovirus (SBWMV) Okl-7 particles with or without trypsin treatment. Gels were stained (Ponceau S) or immunoblotted with monoclonal antibody SCR 133 or rabbit polyclonal antibody (PeAb). Healthy wheat sap is shown in lanes 1. Particles were untreated (lanes 2) or treated with trypsin for 5 or 20 min (lanes 3 and 4, respectively). Right arrows indicate the immunoreactive bands. Marker protein (lane M) sizes are given on the left.

![Fig. 3](image3.png)

Fig. 3. Electron micrographs of soilborne wheat mosaic furovirus (SBWMV) Okl-7 particles coated with A, rabbit polyclonal antiserum; B, monoclonal antibody (MAb) SCR 132; C, MAb SCR 133; or D, untreated control particles. Virions were first trapped using polyclonal antibody. Scale bar = 250 nm.
kDa), several polypeptide fragments were produced (Fig. 2). Two bands (CP and a 13-kDa species) reacted with rabbit polyclonal antiserum to SBWMV. MAb SCR 133 reacted only with the CP band and not with the smaller fragment. Therefore, the epitope with which SCR 133 reacts is probably located within 6 kDa (19 kDa minus 13 kDa) of one extremity of the linear CP sequence.

Immunosensor format electron microscopy trapping and coating tests (Table 2 and Fig. 3) of untreated or 60 min-trypsin--treated SBWMV Okl-7-- or SBWMV Lab1-infected sap showed that the rabbit polyclonal antiserum trapped and coated both untreated and trypsin-treated virus particles of both isolates. SCR 132 also trapped and coated untreated particles of both isolates, but many fewer untreated SBWMV Lab1 particles were trapped by SCR 132 than by the rabbit polyclonal antiserum. SCR 132 also trapped fewer trypsin-treated particles of SBWMV Okl-7, but only a very small number of trypsin-treated particles of SBWMV Lab1. However, SCR 132 coated both treated and untreated particles of both isolates. SCR 133 failed to trap appreciable numbers of any SBWMV particles and only coated untreated SBWMV Okl-7 particles.

Tests for cross-reactivity with other SBWMV isolates and rod-shaped viruses. SBWMV-J, SBWMV-F, OGSV, PMTV, BNYVV, and TRV-PRN were tested by ELISA. The viruses were trapped using homologous polyclonal antiserum and detected with homologous MAbs (positive controls) (data not shown) or with each of the four SBWMV MAbs.

The results of a typical experiment are presented in Table 3 and show that only SCR 133 reacted with all four SBWMV isolates, although consistently weaker with Lab1 compared with the others (Table 1). SCR 133 reacted strongly with OGSV. SCR 132 reacted with SBWMV Okl-7, SBWMV Lab1, and SBWMV-J (weakly), but not with SBWMV-F or OGSV. SCR 135 reacted weakly with SBWMV Okl-7 and SBWMV Lab1, but not with SBWMV-F, SBWMV-J, or OGSV. SCR 135 reacted in immunobots with SDS-denatured CP, and so probably detects a continuous epitope. However, no further tests were done with this MAb, because the cells stopped producing antibodies and the cell line could not be recovered. SCR 134 reacted only with SBWMV Okl-7. None of the MAbs reacted with PMTV, BNYVV, or TRV-PRN.

Sequence analysis of the CP and part of the readthrough domain of SBWMV isolates. RT-PCR cloning and sequencing of nucleotides 334 to 1,000 of SBWMV Okl-7, SBWMV Lab1, and fewer untreated SBWMV Lab1 particles were trapped by SCR 132 than by the rabbit polyclonal antiserum. SCR 132 also trapped fewer trypsin-treated particles of SBWMV Okl-7, but only a very small number of trypsin-treated particles of SBWMV Lab1. However, SCR 132 coated both treated and untreated particles of both isolates. SCR 133 failed to trap appreciable numbers of any SBWMV particles and only coated untreated SBWMV Okl-7 particles.

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The results of a typical experiment are presented in Table 3 and show that only SCR 133 reacted with all four SBWMV isolates, although consistently weaker with Lab1 compared with the others (Table 1). SCR 133 reacted strongly with OGSV. SCR 132 reacted with SBWMV Okl-7, SBWMV Lab1, and SBWMV-J (weakly), but not with SBWMV-F or OGSV. SCR 135 reacted weakly with SBWMV Okl-7 and SBWMV Lab1, but not with SBWMV-F, SBWMV-J, or OGSV. SCR 135 reacted in immunobots with SDS-denatured CP, and so probably detects a continuous epitope. However, no further tests were done with this MAb, because the cells stopped producing antibodies and the cell line could not be recovered. SCR 134 reacted only with SBWMV Okl-7. None of the MAbs reacted with PMTV, BNYVV, or TRV-PRN.

Sequence analysis of the CP and part of the readthrough domain of SBWMV isolates. RT-PCR cloning and sequencing of nucleotides 334 to 1,000 of SBWMV Okl-7, SBWMV Lab1, and

TABLE 3. Cross-reactions of monoclonal antibodies (MAbs) with four isolates of soilborne wheat mosaic furovirus (SBWMV) and other rod-shaped viruses

<table>
<thead>
<tr>
<th>MAb</th>
<th>SCR 132</th>
<th>SCR 133</th>
<th>SCR 134</th>
<th>SCR 135</th>
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<tbody>
<tr>
<td>SBWMV Okl-7</td>
<td>1.964a</td>
<td>2.059</td>
<td>1.048</td>
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<td>SBWMV Lab1</td>
<td>1.589</td>
<td>0.443</td>
<td>0.193</td>
<td>0.540</td>
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<tr>
<td>SBWMV-F</td>
<td>0.134</td>
<td>1.165</td>
<td>0.187</td>
<td>0.195</td>
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<tr>
<td>SBWMV-J</td>
<td>0.511</td>
<td>0.904</td>
<td>0.165</td>
<td>0.157</td>
</tr>
<tr>
<td>Healthy wheat</td>
<td>0.151</td>
<td>0.146</td>
<td>0.142</td>
<td>0.147</td>
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<tr>
<td>OGSV</td>
<td>0.209</td>
<td>1.108</td>
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<tr>
<td>Healthy oat</td>
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<td>TRV-PRN</td>
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<td>Healthy tobacco</td>
<td>0.066</td>
<td>0.061</td>
<td>0.059</td>
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a OGSV = oat golden stripe virus, PMTV = potato mop-top virus, BNYVV = beet necrotic yellow vein virus, and TRV-PRN = tobacco rattle virus strain PRN.
b A 405nm value in antibody-trapped antigen–enzyme-linked immunosorosent assay using homologous antibody preparations to coat microtiter plates. Value recorded after 1 h of substrate incubation.

TABLE 4. Nucleotide differences between soilborne wheat mosaic furovirus (SBWMV) RNA 2 isolates between nucleotides 334 and 1,000

<table>
<thead>
<tr>
<th>Base</th>
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<td>349</td>
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<td>G</td>
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<td>A*</td>
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** = Nucleotide caused amino acid changes in coat protein or coat protein-readthrough protein when compared with SBWMV Okl-WT and SBWMV Okl-7.

Base 663 is an A in Nebraska WT strain RNA2.
SBWMV-F RNA 2, which encode the CP and the first 44 amino acids of the CP-readthrough domain (CP-RT) showed that they differed from SBWMV Okl-WT at up to 27 loci (Table 4). However, most nucleotide changes did not alter the amino acid sequences of the proteins. The deduced amino acid sequences of the SBWMV Okl-7 CP and CP-RT domains were the same as those of SBWMV Okl-WT (7), but the SBWMV Lab1 and SBWMV-F CPs differed from SBWMV Okl-WT by one or three amino acids, respectively (Fig. 4). The CPs of SBWMV Lab1 and SBWMV-F both differed from SBWMV Okl-WT by one amino acid near the N-terminus (Gly to Ser at position 6). The SBWMV-F CP also differed from SBWMV Okl-WT towards the C-terminus (Tyr-159 and Thr-160 were changed to Ile and Ser, respectively; Fig. 4).

Analysis showed that Gly-180 (in the N-terminal part of the readthrough domain of the CP-RT protein) was conserved in all isolates except SBWMV Lab1, where it was changed to Asp-180. SBWMV Lab1 and SBWMV-F also differed from SBWMV Okl-WT in the CP-RT domain, where Val-221 was changed to Ile-221.

Pepscan. Pepscan analysis has been used successfully to identify continuous epitopes on BNYVV and PMTV CPs (10,18). SCR 133 and SCR 134 probably react with continuous epitopes, because they reacted with SDS-denatured CP in immunoblots (Fig. 1), although the reaction with SCR 134 was weak.

A set of seven overlapping octapeptides corresponding to amino acids at the N-terminus of the SBWMV Okl-WT CP (1-MAVNGYTGYNKEL) were synthesized on cellulose paper as described (18). In subsequent tests, SCR 133 and SCR 134 probably react with continuous epitopes, because they reacted with SDS-denatured CP in immunoblots (Fig. 1), although the reaction with SCR 134 was weak.

Computer-assisted model building. A three-dimensional structural model of SBWMV CP was constructed by considering the amino acid sequence similarity with TMV CP and the known structure of TMV CP (15,17). TMV CP was chosen as template, because of the sequence similarity between the two proteins and because of their functional and geometric similarities. Alignments of the CPs of several rod-shaped viruses have shown that key amino acid residues identified as having important structural roles are conserved among them (12). Model building based on amino acid sequence similarity is a well-established procedure (5). Although the 25% similarity between SBWMV and TMV CPs is close to the lower limit appropriate for such modeling, it extends over a large number of amino acids and, together with other similarities between the two proteins, makes structural similarities highly probable (13). The sequences were aligned (Fig. 5) according to the requirements that the four core alpha-helices should not contain any insertions or deletions, and that, wherever possible, residues conserved in all tobamoviruses (2) should also be conserved in SBWMV.

A speculative model of the SBWMV CP molecule was constructed using the program Discover (Biosym Technologies, San Diego, CA). In general, large insertions were added as loops projecting from the protein surface, while small insertions and deletions were constructed by replacing one or two adjacent residues with the appropriate slightly larger or smaller sequence. The structure was frequently adjusted by repeated cycles of energy minimization, which removed the structural strains induced by the modifications. A ribbon drawing of the model is shown in Figure 6A, together with the TMV CP (Fig. 6B) for comparison.

DISCUSSION

Four MAbs produced against an isolate of SBWMV from Oklahoma (Okl-7) discriminated between SBWMV isolates of different geographic origin in ATA-ELISA. MAb SCR 133 reacted with all four SBWMV isolates (Okl-7, -J, -F, and Lab1) and OGSV. In contrast, SCR 134 reacted only with SBWMV Okl-7; SCR 135 reacted only with SBWMV Okl-7 and SBWMV Lab1; and SCR 132 reacted with SBWMV Okl-7, SBWMV Lab 1, and SBWMV-J. None of the MAbs reacted with PMTV, BNYVV, or TRV-PRN.

The complete nucleotide sequences of the RNA 2 of SBWMV isolates Okl-WT, Okl-7, and Nebraska-WT have been published.
shown. Putative epitopes for monoclonal antibodies SCR 134 and SCR 132 are coat protein from isolates SBWMV Okl-7, SBWMV-F, and SBWMV Lab1. Fig. 7. Contains the amino acids Tyr-159 and Thr-160. Thus, it is possible that the SCR 132 epitope contains Tyr-159 and 160. The epitope with which SCR 132 reacts is probably discontinuous, because the MAb did not react with SDS-denatured CP. MAb SCR 132 was shown by electron microscopy to react at the surface and along the sides of virus particles. The epitope was not removed completely by treatment of particles with trypsin, but reactivity was diminished. SCR 132 reacted with three of the four isolates of SBWMV. It did not detect SBWMVF, which differs from SBWMV Okl-7 and SBWMV Lab1 at amino acid positions 159 and 160. Thus, it is possible that the SCR 132 epitope contains the amino acids Tyr-159 and Thr-160.

From our results, we deduce that MAb SCR 133 reacts with a continuous epitope located at one end of the linear CP sequence. The epitope was exposed on the surface along the sides of the particles and was readily removed by trypsin-treatment. Since MAb SCR 133 reacted with all isolates, it seems unlikely that it discriminates the Ser-6 for Gly-6 change in SBWMV-F and SBWMV Lab1. It is, therefore, probable that it reacts with an epitope near the C-terminus, the amino acid residues of which are identical in SBWMV Lab1, SBWMV Okl-WT, and SBWMV Okl-7. MAB SCR 134 reacted only with SBWMV Okl-7, and this can be explained by assuming that it reacts with an epitope that contains Gly-6 near the N-terminus of the CP. The Pepscan data support this conclusion.

The SBWMV CP amino acid sequence fitted easily into the TMV CP fold. The main modifications required were a 10-residue deletion near the N-terminus, an eight amino acid deletion at TMV CP residue 101, and four- to eight-residue insertions at TMV CP amino acid positions 32, 52, 66, 87, and 137. The model-building strategy, thus, demonstrates the feasibility of the protein fold suggested, although it does not prove its existence. The general topology and the approximate positions of the amino acids are probably correct, although fine details such as specific side-chain interactions cannot be predicted by models based on similarities of less than 50% (13). During virus assembly, the sites of the N-terminal deletion and the insertions at 52, 66, and 137 would interact spatially, as would the sites of the deletion at 101 and the insertions at 32 and 87. Significantly, Gly-6, Tyr-159, and Thr-160 are all near the viral surface in the model and are probably exposed (Fig. 6A), so that any modifications to them could affect antibody binding.

The MAbs produced in this work confirm the serological relationship previously found between OGSV and SBWMV (6). They will be useful for surveys to verify the health status of winter wheat crops in the United Kingdom and elsewhere and in further basic research on SBWMV.

**ACKNOWLEDGMENTS**

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**LITERATURE CITED**


