Regulation of plasmodesmal transport by phosphorylation of tobacco mosaic virus cell-to-cell movement protein

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Cell-to-cell spread of tobacco mosaic virus (TMV) through plant intercellular connections, the plasmodesmata, is mediated by a specialized viral movement protein (MP). In vivo studies using transgenic tobacco plants showed that MP is phosphorylated at its C-terminus at amino acid residues Ser258, Thr261 and Ser265. When MP phosphorylation was mimicked by negatively charged amino acid substitutions, MP lost its ability to gate plasmodesmata. This effect on MP-plasmodesmata interactions was specific because other activities of MP, such as RNA binding and interaction with pectin methylesterases, were not affected. Furthermore, TMV encoding the MP mutant mimicking phosphorylation was unable to spread from cell to cell in inoculated tobacco plants. The regulatory effect of MP phosphorylation on plasmodesmal permeability was host dependent, occurring in tobacco but not in a more promiscuous Nicotiana benthamiana host. Thus, phosphorylation may represent a regulatory mechanism for controlling the TMV MP-plasmodesmata interactions in a host-dependent fashion.

Keywords: cell-to-cell movement/cell wall associated kinase/phosphorylation/plasmodesmata/tobacco mosaic virus

Introduction

Katherine Esau first postulated that viruses moved throughout the plant via plant intercellular connections: the plasmodesmata (Esau, 1948). Plasmodesmata are structurally complex plasma membrane-lined pores that, in most cases, limit molecular transport between plant cells to small molecules and metabolites with a molecular mass of up to 1 kDa (reviewed by Lucas et al., 1993). This plasmodesmal size exclusion limit can be dynamically increased to allow intercellular movement of large endogenous proteins (Lucas et al., 1995) or invading viruses (Wolf et al., 1989). In the case of plant viruses, an increase in plasmodesmal permeability and the subsequent viral spread between host cells is mediated by virus-encoded cell-to-cell movement proteins (MPs) (reviewed by Lucas and Gilbertson, 1994; Carrington et al., 1996; Lazarowitz and Beachy, 1999; Rhee et al., 2000; Tzifra et al., 2000).

The best characterized MP is that of tobacco mosaic virus (TMV). Several biological activities have been attributed to TMV MP: interaction with plasmodesmata to increase their permeability (Wolf et al., 1989; Ding et al., 1992; Waigmann et al., 1994), binding to single-stranded RNA (Citovsky et al., 1990, 1992), phosphorylation by a cell wall-associated protein kinase (Citovsky et al., 1993) and interaction with cytoskeletal elements (Heinlein et al., 1995; McLean et al., 1995) and plant cell wall pectin methylesterases (PMEs) (Dorokhov et al., 1999; Chen et al., 2000). Based on these observations, MP was proposed to form complexes with the transported genomic TMV RNA, move these complexes throughout the cell using the cytoskeletal network, associate with the cell wall PMEs, increase plasmodesmal permeability and transverse the enlarged plasmodesmal channels (Ghoshroy et al., 1997; Rhee et al., 2000; Tzifra et al., 2000).

Potentially, MP activity within infected cells may be negatively regulated to prevent its continuous interference with the host plant intercellular communication. The molecular mechanism by which such regulation may occur is unknown. Here, we address this question by demonstrating that MP is phosphorylated in vivo at its C-terminus. Mimicking MP phosphorylation by negatively charged amino acid substitutions severely disturbs MP’s ability to interact with plasmodesmata and to promote viral cell-to-cell movement, suggesting that phosphorylation acts as a negative regulator of plasmodesmal transport. Interestingly, this regulatory effect on plasmodesmal permeability is host dependent and may contribute to the differential susceptibility of various host plants to TMV.

Results

MP phosphorylation in vivo and in vitro

Previously, we have shown that MP is phosphorylated in vitro by a cell wall-associated protein kinase at its C-terminal serine and threonine residues (Citovsky et al., 1993). Because in vitro modifications may not reflect the actual biological state of the protein, it was necessary to examine whether or not MP is phosphorylated in vivo in plant tissue. To this end, we generated transgenic tobacco (Nicotiana tabacum) plants that express either wild-type MP or one of its mutants with amino acid substitutions within the phosphorylation site identified previously in vitro (Citovsky et al., 1993). These plants were grown in the presence of radioactive phosphate, and cell walls, known to contain the majority of the accumulated MP (Deom et al., 1990; Citovsky et al., 1993), were purified
from their leaves. MP contained in these cell wall fractions was immunoprecipitated and analyzed by autoradiography.

Wild-type MP was indeed phosphorylated within the cell walls of the transgenic tobacco plants (Figure 1A, lane 1). It was then important to determine whether the in vitro phosphorylation site of MP, comprising Ser258, Thr261 and Ser265 (Citovsky et al., 1993), is faithfully utilized in vivo. Thus, we analyzed transgenic plants that express MP mutants in which one phosphorylation site residue was left intact while the other two were substituted with non-phosphorylatable alanine residues. Specifically, mutants sb-10, sb-11 and sb-12 retained Ser258, Thr261 and Ser265 residues, respectively. Figure 1A shows that each of these MP mutants was phosphorylated in vivo. The phosphorylation intensity of sb-10, sb-11 and sb-12 (Figure 1, lanes 2, 3 and 4, respectively), however, was lower than that of the wild-type MP in which all three residues are presumably phosphorylated (Figure 1A, lane 1). Substitution of all three residues, sb-13 (Figure 1A, lane 5), abolished MP phosphorylation. These observations indicate that MP associated with cell walls in tobacco leaves is phosphorylated at amino acid residues Ser258, Thr261 and Ser265.

To confirm further that MP is indeed phosphorylated within host cell walls during actual viral infection, we inoculated wild-type tobacco plants with TMV and grew them in the presence of radioactive phosphate followed by MP immunopurification and polyacrylamide gel electrophoretic analysis. Figure 1A shows that MP accumulated in cell walls of TMV-infected plants became phosphorylated (lane 6) whereas no such radioactively labeled protein species was detected in cell walls from uninfected tobacco plants (lane 7). Figure 1B demonstrates that all transgenic plants indeed contained similar quantities of immunopurified MP (lanes 1–5). The amount of MP accumulated within cell walls of TMV-infected plants was slightly lower (Figure 1B, lane 6) and, as expected, no MP was detected in uninfected plants (Figure 1B, lane 7).

Nicotiana tabacum cell wall fraction was then used to characterize its protein kinase activity in vitro using MP as substrate. Figure 2 shows that this cell wall-associated enzyme functioned most efficiently at a slightly basic pH (Figure 2A) and required the presence of Mg²⁺ cations (Figure 2B). The optimal Mg²⁺ concentration was 5 mM. In contrast, the activity in the presence of 0.01–20.6 mM Ca²⁺ was negligible (Figure 2C), suggesting that MP-phosphorylating kinase is not responsible for the Ca²⁺

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**Fig. 1.** MP phosphorylation in vivo. (A) PhosphorImage of MP immunopurified from plant cell walls and analyzed by SDS-PAGE as described in Materials and methods. Transgenic plants expressed the following MP derivatives: lane 1, wild-type MP; lane 2, sb-10; lane 3, sb-11; lane 4, sb-12; lane 5, sb-13. MP phosphorylation during TMV infection: lane 6, cell walls of TMV-infected plants; lane 7, cell walls of uninfected plants. The numbers on the left indicate molecular mass standards in thousands of daltons. An arrowhead indicates the position of TMV MP. (B) Western blot analysis of MP amounts used in the experiment described in (A).

**Fig. 2.** Effects of pH, Mg²⁺ and Ca²⁺ on MP phosphorylation. (A) Effect of pH. The cell wall preparation was dialyzed against 0.1 M citrate buffer pH 4.0–5.5, HEPES pH 6.0–7.5, Tris–HCl pH 8.0–9.0 or borate pH 9.5–10.5 supplemented in each case with 5 mM MgCl₂. Each dialyzed sample was then assayed for its ability to phosphorylate MP. Inset (from left to right): MP phosphorylated at pH 5.0, 8.0 and 11.0, respectively. (B) Effect of MgCl₂. MP phosphorylation was assayed in the pH 8.0 reaction buffer supplemented with the indicated concentrations of MgCl₂. (C) Effect of CaCl₂. MP phosphorylation was assayed in the pH 8.0 reaction buffer supplemented with the indicated concentrations of CaCl₂. For a positive control, 5 mM MgCl₂ was used. The degree of MP phosphorylation was quantified based on analysis of the gel area corresponding to the phosphorylated protein band, using a Molecular Dynamics phosphorimagery.
dependent general protein kinase activity of plant cell walls (Epel, 1994; Yahalom et al., 1998). Mn$^{2+}$ and Zn$^{2+}$ cations were also ineffective (data not shown).

**Mimicking of phosphorylation by negatively charged substitutions blocks MP-induced increase in plasmodesmal permeability in a host-dependent fashion**

One of the major biological functions of MP is to elevate the plasmodesmal size exclusion limit from 1 to 10 kDa (Wolf et al., 1989) or even 20 kDa (Waigmann et al., 1994). The effect of phosphorylation on MP’s ability to increase plasmodesmal permeability was directly tested by microinjection into two different *Nicotiana* hosts, *N. tabacum* (tobacco) and *Nicotiana benthamiana*. *Nicotiana tabacum* is a relatively restrictive host whereas *N. benthamiana* is well known for its promiscuity to viral infection (Gibbs et al., 1977; Dawson and Hilf, 1992).

Purified MP and its derivatives that either cannot be phosphorylated or mimic phosphorylation were co-injected with 10 kDa fluorescently labeled dextran into spongy mesophyll cells of *Nicotiana* leaves (Waigmann et al., 1994; Kragler et al., 1998). Movement of 10 kDa dextran from the injected cell to other, non-injected cells was determined by fluorescence microscopy. Figure 3 and Table I show that 10 kDa dextran microinjected alone generally remained confined to the injected cell in *N. tabacum* as well as in *N. benthamiana* (Figure 3A and E, respectively; Table I), consistent with the known plasmodesmal size exclusion limit of this tissue (Wolf et al., 1989; Kragler et al., 1998). In contrast, microinjection of the wild-type MP promoted an extensive cell-to-cell movement of the dextran in both plant species in most microinjections (Figure 3B and F, respectively; Table I), indicating size exclusion limit-enhancing activity of MP. In addition, movement of fluorescent 10 kDa dextrans into the cells not directly connected to the injected cell indicates, as previously suggested (Waigmann et al., 1994), that MP itself moves into these distant cells to affect their plasmodesmal permeability.

To assess MP activity in the absence of phosphorylation, we used del 7, an MP derivative lacking amino acids 225–268 and thus the entire C-terminal phosphorylation site (Citovsky et al., 1990, 1992). Both in *N. tabacum* (see also Waigmann et al., 1994) and in *N. benthamiana*, del 7 promoted efficient intercellular movement of 10 kDa dextran molecules from the injected cell (arrows) into numerous surrounding cells in most injections (Figure 3C and G; Table I), indicating the active role of del 7 in the increase of plasmodesmal permeability observed in these experiments.

Next, MP phosphorylation was mimicked by substituting Ser258, Thr261 and Ser265 with negatively charged aspartate residues (sh-9 mutant), known to reproduce the electrostatic effects of phosphorylation (Dean and Koshy, 1990). Unlike del 7, sh-9 was unable to mediate cell-to-cell movement of dextran molecules in *N. tabacum* in nearly all injections (Figure 3D; Table I), indicating that

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**Table I. Summary of microinjection experiments**

<table>
<thead>
<tr>
<th>Host plant and injected material</th>
<th>Microinjections</th>
<th>$n$ (positive)/$n$ (total)$^a$</th>
<th>Movement competence</th>
</tr>
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<tbody>
<tr>
<td><em>Nicotiana tabacum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 kDa F-dextran alone</td>
<td>1/12</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>MP + 10 kDa F-dextran</td>
<td>7/9</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>del 7 + 10 kDa F-dextran</td>
<td>13/14</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>sh-9 + 10 kDa F-dextran</td>
<td>1/23</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td><em>Nicotiana benthamiana</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 kDa F-dextran alone</td>
<td>1/10</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>MP + 10 kDa F-dextran</td>
<td>5/6</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>del 7 + 10 kDa F-dextran</td>
<td>6/8</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>sh-9 + 10 kDa F-dextran</td>
<td>16/20</td>
<td>+</td>
<td></td>
</tr>
</tbody>
</table>

$a$($n$ (positive), number of positive injections that are defined by movement of the fluorescent dextran from the injected cell into multiple surrounding cells; $n$ (total), number of total injections.)
MP and del 7 elevated plasmodesmal permeability of the injected cell to allow efflux of a significant portion of the microinjected dye. Microinjection of sb-9, however, resulted in the loss of only 6% of the injected dextran in *N. tabacum* (Figure 4A) but induced an efflux of 73% of the fluorescent signal in *N. benthamiana* (Figure 4B). Statistical evaluation of these results by the unpaired two-tailed Student’s t-test confirmed that, in *N. tabacum*, signal values after co-injection of dextran with wild-type MP or del 7 were significantly different from those obtained after microinjection of dextran alone, indicating that MP and del 7 mediated cell-to-cell movement of the dye. In contrast, sb-9 did not promote a statistically significant dextran movement in *N. tabacum*. Micro-injection of the wild-type MP or its del 7 and sb-9 derivatives in *N. benthamiana* resulted in a statistically significant increase in plasmodesmal permeability compared with that observed after microinjections of 10 kDa dextran alone. The results of this analysis further support that sb-9 enhances plasmodesmal permeability in *N. benthamiana* but not in *N. tabacum*. Collectively, our analyses of the extent of plasmodesmal permeability changes induced by sb-9 suggest that intercellular transport machineries of *N. tabacum* and *N. benthamiana* responded differently to mimicking of MP phosphorylation.

**Nicotiana benthamiana contains a protein kinase activity that phosphorylates MP**

Since plasmodesmal transport in *N. benthamiana* was not sensitive to mimicking of MP phosphorylation, it was necessary to determine whether such phosphorylation occurs in this plant. Thus, we examined the ability of *N. benthamiana* cell walls to phosphorylate TMV MP at its C-terminal Ser258, Thr261 and Ser265 residues. Figure 5 shows that wild-type MP is efficiently phosphorylated by a protein kinase activity within *N. benthamiana* cell walls (lane 1). As expected, del 7, which lacks the entire phosphorylated region, was not phosphorylated (Figure 5, lane 2). Each of the sb-10, sb-11 and sb-12 mutants of MP was phosphorylated but to a markedly lesser degree than the wild-type protein (Figure 5, compare lanes 3, 4 and 5, respectively, with lane 1), suggesting that the substituted amino acid residues reside within the MP phosphorylation site. Finally, sb-9 completely failed to phosphorylate (Figure 5, lane 6). Thus, *N. benthamiana* is capable of phosphorylating TMV MP at the same amino acid residues as those utilized in *N. tabacum*. Taken together with the microinjection data, these results suggest that although MP is likely to be phosphorylated in *N. benthamiana*, the
plasmodesmal transport machinery of this host does not respond to this regulatory mechanism of MP activity.

Mimicking phosphorylation does not impair MP binding to RNA or PME
If mimicking phosphorylation specifically affects MP interaction with plasmodesmata, it should not interfere with biological activities of MP that are not directly related to plasmodesmal gating. Indeed, Figure 6A shows that sb-9 (lane 2) interacted in vitro with the 38 kDa tobacco PME thought to represent a potential cell wall receptor for TMV MP (Dorokhov et al., 1999; Chen et al., 2000). Importantly, the degree of sb-9–PME interaction was comparable to that observed with wild-type MP (Figure 6A, lane 1) or del 7 (lane 3). Furthermore, sb-9 binding to RNA, one of the functional hallmarks of TMV MP activity (Citovsky et al., 1990), was also indistinguishable from that of wild-type MP or del 7 (Figure 6B). These results suggest that the major protein structural characteristics were preserved in both sb-9 and del 7 derivatives of TMV MP. On the other hand, unlike wild-type MP, neither sb-9 nor del 7 was phosphorylated by the tobacco cell wall-associated protein kinase (Figure 6C), demonstrating that these mutations indeed targeted the C-terminal phosphorylation site of the protein. Taken together, our results suggest that mimicking MP phosphorylation did not compromise most biological activities of MP. Thus, the severely reduced plasmodesmata-gating activity of sb-9 observed in N. tabacum may indeed be due to a specific regulatory effect of phosphorylation on MP–plasmodesmata interactions, suggesting that it may represent a plant endogenous means to control TMV MP activity and that this regulatory mechanism is host dependent.

Mimicking MP phosphorylation by negatively charged substitutions blocks cell-to-cell movement of TMV
If phosphorylation indeed functions as a negative regulator of MP function, sb-9, which mimics the phosphorylated state of MP, must be unable to promote viral cell-to-cell spread in vivo. To test this hypothesis, sb-9 was introduced into the infectious cDNA clone of TMV in place of the wild-type MP. First, the wild-type and sb-9-encoding TMV cDNAs were transcribed in vitro and the resulting viral RNAs tested for their ability to replicate and accumulate genomic and subgenomic (sg) TMV RNA within inoculated protoplasts derived from N. tabacum and N. benthamiana hosts. Figure 7 shows that, in both hosts, the sb-9 mutation did not affect viral replication as judged from accumulation of positive- and negative-stranded TMV RNA in the infected protoplasts (Figure 7A and B, respectively). In addition, the appearance of the replication intermediate, negative-stranded TMV RNA, indicates that viral RNA accumulation was due to replication rather than input transcripts that persisted in the inoculated protoplasts.

Figure 7A also shows accumulation of 3′ co-terminal sgRNAs transcribed during the replication cycle and encoding the viral MP and coat protein (CP) gene products (Ishikawa et al., 1991). Similarly to the genomic TMV RNA, no significant differences in accumulation of MP or CP sgRNAs were found between transcripts encoding MP.
or sb-9 in either of the host species. The CP and MP sgRNAs produced were biologically functional, generating their respective protein products both in *N. tabacum* and *N. benthamiana* protoplasts (Figure 7C and D). Furthermore, these results indicate that the sb-9 mutation did not affect the stability and overall accumulation levels of MP in the infected protoplasts of both host plants (Figure 7D).

Next, the effect of mimicking MP phosphorylation on local and systemic TMV movement was examined following inoculation of *N. tabacum* plants with TMV RNA encoding either wild-type MP or its sb-9 derivative. The spread of the virus was assessed from the appearance of CP, a known hallmark of viral infection (Larney et al., 1997; Ghoshroy et al., 1998). Figure 8A shows that inoculation of *N. tabacum* with TMV RNA containing the intact MP gene resulted in efficient viral infection and accumulation of CP (lanes 3 and 4). Furthermore, CP was found both in the inoculated (Figure 8A, lane 3) and uninoculated leaves (lane 4), indicating cell-to-cell (within the inoculated leaf) as well as systemic (from the inoculated leaf into uninoculated leaves) viral movement. In contrast, neither cell-to-cell movement nor systemic spread of the virus occurred when plants were inoculated with TMV RNA encoding the sb-9 derivative of MP (Figure 8A, lanes 5 and 6; compare with uninoculated plant, lane 2). However, sb-9 TMV RNA became infectious both locally (Figure 8A, lane 7) and systemically (lane 8) on transgenic *N. tabacum* plants expressing the wild-type MP, indicating that its failure to inhibit wild-type *N. tabacum* is the result of non-functional MP. These results suggest that mimicking MP phosphorylation blocks viral cell-to-cell and systemic spread in wild-type *N. tabacum* but does not interfere with replication of viral genomes or their ability to spread when the functional MP is provided in trans in MP-transgenic plants.

Unlike *N. tabacum*, inoculation of *N. benthamiana* with TMV RNA encoding the sb-9 derivative of MP still produced efficient accumulation of viral CP both in the inoculated and systemic leaves (Figure 8B, lanes 3 and 4) as compared with *N. benthamiana* plants inoculated with the wild-type TMV RNA (lanes 5 and 6). As expected, no TMV CP band was observed in uninoculated *N. benthamiana* plants (Figure 8B, lane 2). Thus, the regulatory effect of mimicking MP phosphorylation on TMV movement is host dependent, occurring in *N. tabacum* but not in *N. benthamiana*.

**Discussion**

Recent evidence suggests that plasmodesmata function to transport large molecules such as viral MPs and MP–nucleic acid complexes (reviewed by Carrington et al., 1996; Lazarowitz and Beachy, 1999; Rhee et al., 2000; Tzfira et al., 2000), transcription factors (Lucas et al., 1995) and gene silencing signals (Palauqui et al., 1997; Vaucheret et al., 1998). Plasmodesmal transport of these biologically active factors should be tightly regulated. No such regulatory mechanism, however, has been described to date. Here, we utilized one of the best characterized proteins known to traffic from cell to cell, TMV MP, to show that its interaction with plasmodesmata is controlled by phosphorylation.

Although MP phosphorylation has been demonstrated in plant protoplasts (Watanabe et al., 1992; Haley et al., 1995), these cells lack cell walls and their cognate plasmodesmata, which represent the biological sites of action for MP (Tomenius et al., 1987; Atkins et al., 1991; Ding et al., 1992). Thus, it was still unknown whether MP phosphorylation takes place *in vivo*, within plant tissues, where MP is biologically active. Our present results show that phosphorylation of MP contained in the cell walls, presumably within plasmodesmata (Ding et al., 1992), occurs *in vivo* in transgenic plants. The MP phosphorylation site utilized *in vivo*, i.e. C-terminal amino acid residues Ser258, Thr261 and Ser265, is identical to that
determined in vitro using purified cell wall fractions (Citovsky et al., 1993). So far, this MP phosphorylation has been detected only in *N. tabacum*; here, we show that a similar MP kinase activity is also present in *N. benthamiana*. Furthermore, we provide evidence that MP produced during the wild-type TMV infection of tobacco plants is also phosphorylated. C-terminal phosphorylation of MP is carried out by a serine/threonine-specific protein kinase associated with the cell wall of the host cell. This enzyme activity requires Mg²⁺ but not Ca²⁺ cations. Independence from Ca²⁺ distinguishes it from several known protein kinases that associate with plant cell walls (He et al., 1996; Yahalom et al., 1998).

The biological function of MP C-terminal phosphorylation was examined using negatively charged amino acid substitutions within the phosphorylation site. Substitutions with aspartate or glutamate are known to reveal the electrostatic effects of phosphorylation (Dean and Koshland, 1990). For example, replacement of serine by aspartate in the HPt protein has been shown to cause shifts in two-dimensional NMR spectra similar to those elicited by phosphorylation (Wittekind et al., 1989). Also, inactivation by phosphorylation of Ser113 in isocitrate dehydrogenase is mimicked when aspartate is substituted at this site (Thorsness and Koshland, 1987). Our studies show that substituting Ser258, Thr261 and Ser265 with aspartate residues (sb-9 derivative of MP) inactivated the MP–plasmodesmata interactions leading to gating of these channels. Specific interference of MP phosphorylation with plasmodesmal transport is supported by the observation that mimicking this protein modification did not affect several other known activities of MP in vitro nor did it compromise MP stability in vivo. Interestingly, the negative regulation of MP interactions with plasmodesmata was host dependent. In *N. tabacum*, mimicking phosphorylation blocked MP’s ability to interact efficiently with plasmodesmata, whereas in *N. benthamiana* the same MP derivative remained fully active as compared with non-phosphorylated MP. This result, taken together with the observation that both *N. tabacum* and *N. benthamiana* are competent for MP phosphorylation, suggests that the plasmodesmal transport machineries of the two hosts differ. Specifically, the plasmodesmal transport machinery of *N. benthamiana* is likely to be able to mediate movement of phosphorylated and non-phosphorylated MP, whereas the plasmodesmal transport machinery of *N. tabacum* may support only movement of non-phosphorylated MP. Thus, the inactivation strategy by phosphorylation, highly successful in *N. tabacum*, is not effective in *N. benthamiana*, consistent with the known fact that *N. benthamiana* is one of the most susceptible hosts for plant viruses (Gibbs et al., 1977; Dawson and Hilf, 1992).

Besides TMV MP, few plant viral MPs have been tested for phosphorylation. However, recently reported phosphorylation of the 17 kDa MP of potato leafroll virus has also been proposed to regulate negatively the function of this transport protein (Sokolova et al., 1997). Thus, it is tempting to speculate that the regulatory activity of MP phosphorylation or other strategies to inactivate this viral protein (Hughes et al., 1995) may underlie the well known but previously unexplained differences between various tobacco species in their susceptibility to viral infection (Gibbs et al., 1977; Dawson and Hilf, 1992).

Because C-terminal phosphorylation of MP blocks its interaction with plasmodesmata, it should also inhibit the ability of the virus to spread from cell to cell in vivo. Indeed, TMV encoding the sb-9 derivative of MP within its genomic RNA was unable to move locally and systemically within *N. tabacum*. The mutant virus was fully capable of movement in transgenic plants expressing wild-type TMV MP, indicating that the mutation did not interfere with replication and translation of viral gene products. Indeed, the sb-9 mutation did not affect replication of viral RNA, production of MP and CP sgRNAs, and expression and stability of MP and CP in *N. tabacum* and *N. benthamiana* protoplasts. Thus, mimicking MP phosphorylation blocks TMV movement through plasmodesmata, probably by preventing the MP-induced increase in permeability of these intercellular channels. Similarly to microinjection experiments, the inhibitory effect of mimicking MP phosphorylation on TMV movement in vivo was host dependent, occurring in *N. tabacum* but not in a more promiscuous *N. benthamiana* host. These results were supported by direct evaluation of the cell-to-cell movement capacity of wild-type TMV MP, sb-9 and del 7, which were expressed as green fluorescent protein (GFP) fusion proteins following microinoculation of their encoding genes into epidermal cells of mature leaves from *N. tabacum* or *N. benthamiana*. del 7–GFP and wild-type TMV MP–GFP showed significant movement in both hosts. In contrast, cell-to-cell transport of sb-9–GFP was comparable to that of wild-type TMV MP in *N. benthamiana*, but was severely reduced in *N. tabacum* (K.Trutyneva, R.Bachmaier and E.Waigmann, unpublished).

Negative regulation of MP effects on plasmodesmata makes biological sense. Unlike other TMV-coded proteins expressed throughout the course of viral infection, MP is synthesized only transiently (reviewed in Hull, 1989). However, the produced protein does not appear to have a rapid turnover; instead, it persists in the infected cells, accumulating within their plasmodesmata (Tomenius et al., 1987; Ding et al., 1992). The continuous presence of active MP may elevate plasmodesmal permeability and alter intercellular communication, an important biological process. Potentially, phosphorylation minimizes this MP interference with plasmodesmal transport during viral infection. Thus, viral cell-to-cell movement may involve plasmodesmal gating by the newly synthesized, unphosphorylated MP followed by phosphorylation of MP that has already performed its function within plasmodesmata, preventing it from further action. This model suggests that MP is most active in freshly infected cells where it is known to be produced vigorously (Dawson and Lehto, 1990) while MP contained within cells at later stages of the infection is expected to become permanently inactivated. Indeed, recent observations demonstrate that plasmodesmal gating is restricted to the outer edge of an expanding TMV infection site even though MP itself is also present in plasmodesmata of cells located in the center of this infected area (Oparka et al., 1997).

Because viruses often adapt existing cellular machinery for their own needs, TMV is likely to employ an endogenous pathway for cell-to-cell transport of proteins and nucleic acids. Thus, MP phosphorylation may represent a general mechanism by which plants regulate intercellular transport of macromolecules.
Materials and methods

Plant lines
Nicotiana tabacum cv. Turk and N. benthamiana were used as hosts. Transgenic N. tabacum plants expressing MP, sb-10, sb-11, sb-12 and sb-13 were generated as described (Citovsky et al., 1992).

Mutagenesis and expression of MP
Substitution mutants sb-9, sb-10, sb-11, sb-12 and sb-13 of MP were constructed using oligonucleotide-directed mutagenesis (McCray et al., 1989) and confirmed byideoxynucleotide sequencing (Kraft et al., 1988). Generation of deletion mutant del 7 of MP was described previously (Citovsky et al., 1990, 1992). MP and its del 7, sb-9, sb-10, sb-11 to sb-12 derivatives were expressed in Escherichia coli using the T7 RNA polymerase system (Studier et al., 1990), purified to near homogeneity (95–98% pure as determined by silver-stained SDS–polyacrylamide gels), and verified by western blot analysis as described (Citovsky et al., 1990, 1992, 1993; Waigmann et al., 1994).

Infection of plants and detection of TMV in plant extracts
TMV cDNA clones encoding wild-type MP or sb-9 in the pTMV304 construct were linearized with KpnI. Infectious transcripts were produced from the SP6 promoter and capped with 7m-diguanosine triphosphate using the Ribomax large scale RNA Production System kit according to the manufacturer’s instructions (Promega). Nicotiana tabacum or N. benthamiana plants were mechanically inoculated with 10 μl of the resulting 100 μg/ml RNA solution supplemented with 3 μg/ml cellulose. Four individual plants were inoculated with each viral transcript.

Ten days after inoculation, leaf samples (50 mg fresh weight) were removed and homogenized in 100 μl of extraction buffer (75 mM Tris–HCl pH 6.8, 9 M urea, 4.3% SDS and 7.5% β-mercaptoethanol). The homogenate was heated at 100°C for 5 min, centrifuged (10 000 g for 10 min) to remove all insoluble material, and 10 μl of the supernatant were analyzed by SDS–PAGE (Laemmli, 1970) on 12.5% gels. Following electrophoresis, gels were stained with Coomassie Blue and photographed using a Fotodyne gel documentation system. The position of TMV CP was verified by western blot analysis (data not shown). No significant differences between individual plants inoculated with the same viral construct were observed.

Plant cell wall fractions
Cell wall fractions were prepared as described (Citovsky et al., 1993). Briefly, fresh plant tissue derived from mature leaves of wild-type or transgenic N. tabacum plants was ground to a fine powder and homogenized at 4°C in 1 vol. of buffer H (0.1 M HEPES pH 7.4, 100 mM NaCl, 10 mM EDTA, 5 mM dithiothreitol, 1 mM phenylmethylsulfonyl fluoride, 1 mM leupeptin, 1 mM pepstatin). The homogenates, which represented the total plant cell extract, were centrifuged (1000 g for 5 min at 4°C) to give a crude cell wall pellet and supernatant. Cell walls were further purified by homogenization at 4°C in 10 vols of buffer H with 2% Triton X-100 and centrifugation. This procedure was repeated twice, followed by six washes (1000 g for 5 min at 4°C) in buffer H with 2% Triton X-100 and two washes in buffer H without the detergent. The resulting white insoluble material was resuspended in 0.5 vols of buffer H, quick-frozen in liquid nitrogen and stored at –70°C until use.

MP phosphorylation in vivo
Transgenic plants expressing MP, sb-10, sb-11, sb-12 and sb-13 were grown from seeds in sealed Magenta jars on 0.7% agar in a low phosphate MS medium formulated in-house by replacing potassium phosphate from the MS recipe (Murashige and Skoog, 1962) with potassium chloride and supplementing it with 10 μCi (1 Ci = 37 GBq) of 32P. When the plants reached 6 cm in height, one of their leaves was harvested and used for isolation of cell walls as described above except that buffer H was supplemented with 10 μM okadaic acid, 50 mM sodium fluoride and 0.2 mM sodium orthovanadate to inhibit cellular phosphatases. The cell wall fraction was extracted in 1 vol. of 75 mM Tris–HCl pH 6.8, 9 M urea, 7.5% β-mercaptoethanol, centrifuged (15 000 g for 15 min at 4°C), and the supernatant dialyzed for 16 h at 4°C against buffer H. MP was then immunoprecipitated with affinity-purified rabbit anti-MP antibodies (McLean et al., 1995) using standard procedures (Ausubel et al., 1987) and analyzed on 12.5% SDS–polyacrylamide gels, followed by autoradiography. Equal loading of all lanes was confirmed by western blot analysis using mouse anti-MP antibodies followed by alkaline phosphatase detection.

MP phosphorylation in vitro
Cell wall fractions (10 μg of protein in 25 μl of buffer H) were mixed with 5 μl of 10× kinase buffer (0.5 M Tris–HCl pH 8.0, 50 mM MgCl2) and 20 μl of buffer H containing 2 μg of MP or its derivatives. The reaction was started by addition of 0.5 μl of [γ-32P]ATP (equal to 5 μCi, 3000 Ci/mmol), continued for 15 min at room temperature, and stopped with 50 μl of loading buffer (75 mM Tris–HCl pH 6.8, 9 M urea, 4.3% SDS, 7.5% β-mercaptoethanol). The phosphorylated protein was resolved on a 12.5% SDS–polyacrylamide gel and detected by autoradiography.

MP binding to RNA
The 300 bp fragment of the spinach chloroplast plastidic electron transport D gene was used as template for T7 RNA polymerase to generate a radioactively labeled RNA probe with a specific activity of 107 c.p.m.μg (Stern and Gruissem, 1987). A gel-purified RNA probe (1.0 ng) was incubated for 15 min at 4°C in 20 μl of buffer H with 0.1 μg of purified MP or its derivatives. RNA–protein complexes were cross-linked by UV-light irradiation and analyzed on 12.5% SDS–polyacrylamide gels as described (Citovsky et al., 1990, 1992).

MP binding to PME
A detergent blot overlay assay (Chen et al., 2000) was used to examine binding of MP or its derivatives to PME. Briefly, PME (1 μg) purified from tobacco leaves (Chen et al., 2000) was electrophoresed on a 12.5% SDS–polyacrylamide gel, electrophoblotted onto a PVDF membrane, depleted of SDS by guanidine hydrochloride extraction, renatured, blocked with 1% bovine serum albumin, and incubated with 10 μg/ml purified MP, del 7 or sb-9 as described (Chen et al., 2000). Following incubation, the membrane was washed and protein binding was detected using anti-MP antibody and the ECL western blotting kit (Amersham) (Chen et al., 2000).

Replication and translation of TMV RNA encoding MP or sb-9 in N. tabacum and N. benthamiana protoplasts
Protoplasts were prepared from mature leaves of N. tabacum and N. benthamiana plants as described (McLean et al., 1995), electroporated with 10 μl of the infectious transcripts of TMV cDNA clones encoding wild-type MP or sb-9 (see above), and incubated for 20 h at 25°C as described (Lewandowski and Dawson, 1998). For analysis of viral RNA, total RNA from 106 protoplasts was extracted, resolved on 0.9% agarose gels and analyzed by northern blotting as described (Lewandowski and Dawson, 1998). Strand-specific digoxigenin-labeled riboprobes were used to detect TMV-derived RNAs. Positively-stranded full-length and sg TMV RNAs were detected using a probe complementary to TMV nucleotides 5545–5845 and negative-stranded RNA was detected with a probe identical to the same TMV sequence. Because negative-strand TMV RNA is produced at much lower levels than positive-strand RNA (Eskikawa et al., 1991), 20-fold larger RNA samples were used for its detection. For protein analysis, 106 protoplasts were resuspended and boiled in the gel loading buffer (Laemmli, 1970), and the resulting total cell protein extracts resolved on 12.5% SDS–polyacrylamide gels followed by western blot analysis using affinity-purified anti-CP or anti-MP antibodies.

Microinjection, image analysis and fluorescence quantification
A mixture containing ~0.4 μg/ml TMV MP, sb-9 or del 7 and 0.5 mg 10 kDa fluorescein isothiocyanate-labeled dextran (FITC–dextran; Sigma-Aldrich) was microinjected into N. tabacum cv. Turk or N. benthamiana leaf mesophyll cells essentially as described (Ding et al., 1992; Waigmann et al., 1994). Before use, FITC–10 kDa dextran was separated from free dye by centrifugation through a 10 kDa cut-off spin column (Ultracel-MC; Millipore, MA). Intercellular movement of FITC–10 kDa dextran was observed and documented with a cooled slow scan, high resolution CCD camera (Quantix-G2; Photometrics, Tucson, AZ). Each experiment was repeated numerous times (Table I).

For quantitation of dextran fluorescence on a per cell level, 5 and 30 min after injection the signal was recorded with the CCD camera and quantified using the ‘analyse’ function of the IPlab software program (Scanalytics, Virginia, VA). In this function, all signal values within a
region of interest, i.e. the area of the injected cell (S), can be summed. Then, the system background (B) was obtained by combining the values in an area identical in size to that of the injected cell but in a region containing non-injected cells. Intracellular fluorescence values (F) for each time point were calculated as F = S – B. Finally, the movement of fluorescent dextran out of the injected cell was expressed as a percentage of signal input calculated as the ratio of the signal retained within the injected cell 30 min after injection to that measured 5 min after injection, i.e. F_{30min}/F_{5min} × 100. The 5 min period was chosen as the first time point at which the image could be reliably recorded in all experiments whereas the 30 min time point represented a standard observation time during which MP-mediated dextran movement is known to occur (Waightmann et al., 1994). For the evaluation of standard deviations, measurements were calculated based on 5–8 independent microinjection experiments, i.e. number of measured cells n = 5–8. To determine whether any two sets of fluorescent signal measurements were statistically different from each other, data were subjected to the unpaired two-tailed Student’s t-test. For this test, a combined n = 10–15 (derived from the two sets of fluorescent signal measurements to be compared) was used. 2P values <0.01, corresponding to the statistical probability of >99%, were considered statistically significant.

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References


Opara,K.J., Prior,D.A.M., Santa-Cruz,S., Padgett,H.S. and Beachy,R.N. (1997) Gating of epidermal plasmodesmata is restricted to the leading

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