The Role of Microtubule Association in Plasmodesmal Targeting of *Potato mop-top virus* Movement Protein TGBp1

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Abstract: Cell-to-cell movement of *Potato mop-top virus* (PMTV) is mediated by three virus-encoded ‘triple gene block’ (TGB) proteins termed TGBp1, TGBp2 and TGBp3. TGBp1 binds virus RNAs to form viral ribonucleoprotein complexes (vRNPs), the transport form of viral genome. TGBp2 and TGBp3 are necessary for intracellular delivery of TGBp1-containing vRNPs to plasmodesmata. To analyze subcellular localization and transport of TGBp1 we used a single binary vector for agrobacterium-mediated co-expression of PMTV TGBp1 fused to green fluorescent protein and TGBp2/TGBp3. At two days post infiltration (dpi) TGBp1 was found in the nucleus and in association with microtubules (MTs). Similar localization pattern was revealed in cells expressing GFP-TGBp1 alone after particle bombardment. At 3 dpi, in addition to the nucleus and MTs, TGBp1 was detected in numerous granular bodies located both along the MTs and at the cell wall. The latter structures co-localized with plasmodesmata-associated callose depositions. At 4 dpi, GFP-TGBp1 was detected in cell wall-associated bodies and also in residual MTs, the nucleoplasm and large perinuclear inclusions resembling aggresomes. Therefore GFP-TGBp1 association with MTs preceded to its localization to plasmodesmata. Disassembly of cell MTs by colchicine prevented GFP-TGBp1 targeting to plasmodesmata and the MT-dependent aggresome formation. Deletion analysis also revealed a correlation between TGBp1 microtubule association and plasmodesmata targeting. We propose that TGBp1 interaction with MTs may be important for the formation of vRNP bodies destined for the transport to plasmodesmata as well as degradation of the excessive TGBp1.

Keywords: Movement protein, virus movement, plasmodesmata, microtubules, subcellular localization.

INTRODUCTION

Cell-to-cell transport of plant viruses occurs through plasmodesmata, the channels interconnecting neighboring cells in plant tissues, and requires dedicated virus-encoded movement proteins (MPs) [1-5]. In positive-stranded RNA viruses the number of encoded MPs can vary between distant virus groups, and they are often unrelated in their primary structure [1]. Nevertheless, viral MPs which represent dissimilar transport systems share three characteristic features. First, the RNA-binding activity is believed to be responsible for formation of complexes with viral genomic RNAs (vRNPs) which represent the transport form of viral genomes capable of cell-to-cell movement. Second, the ability of MPs to interact with cell membranes, in particular with the endoplasmic reticulum (ER), may direct MPs to specific pathway of intracellular transport to plasmodesmata. Third, MPs are able to accumulate in plasmodesmata or plasmodesmata-associated sites and to increase the size exclusion limit of plasmodesmata pores for efficient passage of macromolecules to neighboring cells [3-5].

A number of plant virus genera require for their cell-to-cell movement a ‘triple gene block’ (TGB), conserved gene module encoding three MPs [6]. The TGB position relative to other genes vary in genomes of different viruses, however the order of genes within the TGB and the properties of the encoded proteins are conserved [7, 8]. Based on differences in the structure of TGB proteins and the requirement of viral capsid protein for cell-to-cell movement two types of TGB are distinguished: ‘hordei-like’ TGB in rod-shaped viruses (family *Virgaviridae* and genus *Benyvirus*) and ‘potex-like’ TGB in filamentous viruses (family *Flexiviridae*) [7, 8].

Studies of hordei-like TGB carried out in the recent years and employed as models *Barley stripe mosaic virus* (BSMV, genus *Hordeivirus*), *Poa semilatent virus* (genus *Hordeivirus*) and *Potato mop-top virus* (PMTV, genus *Pomovirus*) revealed characteristic biochemical properties...
and functions of individual TGB proteins, as well as their physical and functional interactions involved in virus cell-to-cell movement [8, 9]. TGBp1, the largest of TGB proteins with molecular mass of 51 to 63 kDa in hordei-like TGBs, consists of three domains, the N-terminal domain, internal domain and the C-terminal NTPase/helicase domain [10]. In *Poa semilatilis virus*, the N-terminal domain represents an unfolded protein region which includes two positively charged sequence motifs involved in virus long-distance movement via the phloem, whereas the internal domain is a small structured domain able to interact with both the N-terminal domain and the NTPase/helicase domain [10, 11]. The latter domain has the NTPase and helicase activities and is capable of self-interactions [10-14]. Additionally, all three domains of the hordeiviral TGBp1 bind RNA exhibiting either a cooperative (the internal and the NTPase/helicase domains) or non-cooperative (the N-terminal domain) binding [10, 11, 13]. These RNA binding activities are responsible for interaction with viral RNAs and the formation of vRNPs; such RNPs were isolated from BSMV-infected plants and were shown to contain BSMV positive-stranded RNAs and TGBp1 but not other viral or non-viral components [15]. The presence of multiple binding sites in TGBp1 and the ability of internal domain for interactions with other protein domains, as well as the proposed ability of the N-terminal domain to conditionally adopt a structured conformation, provide the basis for structural remodeling of vRNPs that would be necessary for different stages of virus movement in plant [10]. TGBp2 and TGBp3 are small integral membrane proteins required for delivery of hordei-like TGBp1 to plasmodesmata [7, 8]. While TGBp2, when expressed in the absence of TGBp3, is localized to the cortical endoplasmic reticulum (ER) or numerous ER-associated vesicular bodies, TGBp3 localizes to plasmodesmata or plasmodesmata-associated membrane compartments called peripheral membrane bodies (PMBs) and, moreover, is able to target TGBp2 from the ER structures to plasmodesmata and PMBs [7-9]. Furthermore, TGBp2 and TGBp3 direct TGBp1 to PMBs and plasmodesmata, and in the case of BSMV this targeting could involve direct interactions between TGBp1 and TGBp3 [16, 17]. Recent evidences suggest that TGBp3 contains signals of plasmodesmata targeting [18, 19] and that TGBp3 serves therefore as a ‘driving force’ for the targeting of TGBp2, TGBp1, and most probably TGBp1-containing vRNPs to plasmodesmata [8].

Previously we analyzed PMTV TGB functions using an experimental system in which intra- and intercellular transport of TGB proteins was reconstructed using individually cloned PMTV TGB genes and a particle bombardment-based delivery of recombinant plasmids into plant cells [16]. Specifically, this experimental system includes two expression vectors which carry, under the control of the cauliflower mosaic virus 35S promoter, either the PMTV TGBp1 gene fused to the green fluorescent protein (GFP) gene, or the cloned region of PMTV genome comprising overlapping genes of TGBp2 and TGBp3. The latter construct mimics the natural expression of TGBp2 and TGBp3 from a single subgenomic RNA [20]. After bombardment of plant leaves with metal particles carrying the GFP-TGBp1 expression vector, the fusion protein expressed in individual epidermal cells was distributed uniformly in the cytoplasm and also found in the nucleus. However, when leaves were bombarded by a mixture of GFP-TGBp1 and TGBp2/TGBp3 expression vectors, GFP-TGBp1 was directed to large PMBs and to cell wall-embedded punctate bodies supposed to correspond to plasmodesmata and, moreover, was transported into cells immediately adjacent to the bombarded cell showing that PMTV TGBp2 and TGBp3 can mediate transport of GFP-TGBp1 to and through plasmodesmata in the absence of other virus-encoded components [16]. Further experiments revealed that specific interactions between TGBp1 and TGBp2/TGBp3 are required for TGBp1 delivery to plasmodesmata and translocation to adjacent cells as well as allowed us to carry out a preliminary mapping of protein region involved in interaction between TGB proteins [16]. Therefore, the experimental system based on co-expression of TGB genes by use of particle bombardment proved to be a useful tool for studies of TGB-mediated transport.

In this paper we report elaboration of an experimental system based on agrobacteria-mediated transient expression of three PMTV TGB proteins from a single binary vector providing controlled synthesis levels for TGBp1, TGBp2 and TGBp3. Using this system we demonstrate a transient association of GFP-tagged TGBp1 with microtubules and microtubule-associated bodies that preceded TGBp1 targeting to cell wall-embedded structures which are proved to be plasmodesmata. We also show that disruption of cell microtubules with colchicine prevents TGB1 targeting to plasmodesmata, and TGB1 mutations which block interactions with microtubules also impede its plasmodesmata targeting.

**MATERIALS AND METHODOLOGY**

**Recombinant Clones**

Recombinant plasmids pRT-GFP-TGBp1 [16] and pCB-GFP-TAL [18] have been described earlier. To generate the YFP-TAL construct the GFP gene in pCB-GFP-TAL was replaced with the YFP gene using appropriate restriction sites. To obtain YFP-MA4 fusion gene, the microtubule-binding domain of mouse microtubule-associated protein 4 (MAP4) was amplified with specific primers as described [21] and cloned as a fusion with the YFP gene into pRT100 [22]. Expression cassettes including the 35S promoter, the fluorescent protein fusion gene and transcriptional terminator were cut out of the vectors rYFP-TGBp1, pCB-YFP-TAL and rRT-YFP-MA4 by digestion with HindIII and ligated into HindIII-digested binary vector pLH7000 [23] provided by Dr L. Hausmann (Federal Centre for Breeding Research on Cultivated Plants, Germany). pLH-35S:GFP-TGBp1-NOS:TGBp2/TGBp3 was constructed in three steps. Using pBin19 [24] as a template, NOS promoter was replaced with the YFP gene using appropriate restriction sites. The latter construct mimics the natural expression of TGBp2 and TGBp3 from a single subgenomic RNA [20]. After bombardment of plant leaves with metal particles carrying the GFP-TGBp1 expression vector, the fusion protein expressed in individual epidermal cells was distributed uniformly in the cytoplasm and also found in the nucleus. However, when leaves were bombarded by a mixture of GFP-TGBp1 and TGBp2/TGBp3 expression vectors, GFP-TGBp1 was directed to large PMBs and to cell wall-embedded punctate bodies supposed to correspond to plasmodesmata and, moreover, was transported into cells immediately adjacent to the bombarded cell showing that PMTV TGBp2 and TGBp3 can mediate transport of GFP-TGBp1 to and through plasmodesmata in the absence of other virus-encoded components [16]. Further experiments revealed that specific interactions between TGBp1 and TGBp2/TGBp3 are required for TGBp1 delivery to plasmodesmata and translocation to adjacent cells as well as allowed us to carry out a preliminary mapping of protein region involved in interaction between TGB proteins [16]. Therefore, the experimental system based on co-expression of TGB genes by use of particle bombardment proved to be a useful tool for studies of TGB-mediated transport.
promoter and terminator sequences, was digested with XhoI and XbaI and ligated with XhoI-XbaI-fragment of pRT-
TGBp2/TGBp3 [16] containing overlapping PMTV TGBp2 and TGBp3 genes. The expression cassette from the
resulting vector was cut out with SalI and subcloned into SalI-digested pH7000-GFP-TGBp1.

Particle Bombardment, Agroinfiltration and Fluorescent Microscopy

Particle bombardment of N. benthamiana leaves was performed using the flying disc method with a high-pressure
equipment-based PDS-1000 system (Bio-Rad) as described [25]. Agroinfiltration and imaging of cells with a Leica TCS SP2
system was carried out as described [26]. The software
package provided by the manufacturer was used for
projections of serial optical sections and image processing.

Leaf Treatments

Callose was visualized in leaf cells with 0.1% (w/v)
sirolfluor-containing aniline blue (Merck) in 1 M glycine, pH 9.5, as previously described [27]. To induce plasmolysis, leaf
pieces were incubated in 0.5 M mannitol solution for 5 - 10
min. The colchicine treatments were done by infiltrating 50
μM colchicine solution in an agroinfiltration buffer (10 mM
MES [pH 5.85], 10 mM MgCl2, 150 μM acetylsyringone) into N. benthamiana leaves.

RESULTS

TGBp1 Interaction with Microtubules in Particle
Bombardment Assay

In our previous particle bombardment experiments,
PMTV 35S:GFP-TGBp1 construct gave rise to diffuse
fluorescence observed in the nucleus and the cytoplasm of
bombarded cells. Importantly, the cytoplasmic TGBp1
fluorescence was not associated with the cell wall or plasmodesmata [16]. Additionally, the cytoplasmic
fluorescence was sometimes associated with numerous faint
thin strings resembling in their appearance microtubules
(MTs) (not published). In these earlier experiments,
microscopy of bombarded leaves was carried out at 15 - 18
hours post bombardment that would hinder detection of the
MT-like structures. Therefore in this study we used a longer
incubation of bombarded leaves before microscopy. When
GFP-TGBp1 was observed in epidermal cells of N. benthamiana leaves at 24 hours post bombardment or at later
time points, strong association of GFP fluorescence with the
MT-like structures was easily detected in most examined
cells (Figs. 1A, 3B). To identify the nature of the GFP-
TGBp1-containing string-like structures, co-localization
studies with markers of MTs and microfilaments were
conducted. To visualize actin cables the actin-binding
domain of mouse talin fused to yellow fluorescent protein
(YFP) (YFP-Tal) known to bind the actin cytoskeleton in
different types of living plant cells was used [28]. MTs were
visualized by use of marker protein YFP-MAP4 that
represented yellow fluorescent protein (YFP) fused to the
microtubule-binding domain of mouse microtubule-
associated protein 4 (MAP4) as described [21]. This fusion
has been previously shown to bind MTs in plant cells [21].
Co-expression of GFP-TGBp1 with YFP-Tal by particle
bombardment revealed that GFP and YFP fluorescent signals
were associated with different non-overlapping filamentous
structures (Fig. 1C). In contrast, in cells where GFP-TGBp1
and YFP-MAP4 were co-expressed the string-like structures
containing GFP-TGBp1 coincided with MTs labeled with
YFP-MAP4 (Fig. 1D). A similar co-localization with MAP4-
labeled MTs was recently reported for Tobacco mosaic virus (TMV) MP [29]. Thus our observations demonstrated
association of TGBp1 with MTs.

Expression of Three PMTV TGB Proteins from a Single
Binary Vector

Observations of GFP-TGBp1 in association with MTs and MT-associated structures made in the absence of two
other TGB proteins remained uncertain the significance of MT localization for the TGBp1 intracellular movement.
Previously, we analyzed TGBp2/TGBp3-dependent intra-
and intercellular transport of PMTV GFP-TGBp1 using the
experimental system based on particle bombardment with
two expression vectors [16]. This system had two major
disadvantages. First, the relative amounts of the expression
plasmids delivered in each particular cell can vary that may
result in variations of the TGBp1 to TGBp2/TGBp3 ratio in
different cells. Second, both the TGBp1 gene and the
TGBp2/TGBp3 bicistronic gene region were cloned in
expression cassettes under the control of the 35S promoter
that results, taking into account that translation of a BSMV
bicistronic template gives a TGBp2:TGBp3 ratio of 10:1
[20], in an approximate TGBp1:TGBp2:TGBp3 molar ratio
of 10:10:1 in co-bombarded cells instead of 100:10:1 ratio
expected for natural virus infection on the basis of indirect
data such as mRNA levels and in vitro translation efficiencies [9]. Therefore, both TGBp2 and TGBp3 were
present in co-bombardment experiments in approximately
ten-fold excess relative to TGBp1 amounts.

To improve the described system for further PMTV TGB
studies we constructed a binary vector harboring two expressing cassettes, one of which is driven by the 35S
promoter and contains GFP-fused PMTV TGBp1 gene, while
another contains the TGBp2/TGBp3-encoding region cloned under the control of the promoter and transcription
terminator of the A. tumefaciens nopaline synthase (NOS).
The NOS promoter is known to be considerably weaker
compared to the 35S promoter [30]. The two-cassette
construct was termed pLH-35S:GFP-TGBp1-NOS:TGBp2/
TGBp3 (Fig. 2).

Epidermal cells of N. benthamiana leaves agroinfiltrated
with the construct 35S:GFP-TGBp1-NOS: TGBp2/TGBp3 were
imaged using confocal laser scanning microscopy at
different time points. At one day post infiltration (dpi) no
visible GFP fluorescence was observed in infiltrated leaf
areas (data not shown). At 2 dpi the GFP fluorescence was
detected in the cytoplasm and within the nucleus as
previously observed [16]; the cytoplasmic fluorescence was
mainly associated with MTs similar to those found in
bombardment experiments (Fig. 3A). At 3 dpi GFP-TGBp1
was again found in the nucleus and association with MT.
Additionally, GFP-TGBp1 was detected in numerous granular bodies of 0.3 - 0.8 μm in size which were located
both along the MTs and close to the cell wall (Fig. 3B, C).
Using time-lapse microscopy we failed to detect movement
of GFP-TGBp1-containing MT-associated granular structures in time periods of up to 1 min (data not shown). At 4 dpi, GFP-TGBp1 was detected mostly in cell wall-associated bodies and also in residual MTs, the nucleoplasm and large amorphous inclusions comparable in size to the nucleus (Fig. 3D). Typically, one such inclusion was found in each transformed cell close to the nucleus (Fig. 3D). These structures resembled aggresomes which are formed in MT-dependent manner in response to excessive amounts of misfolded proteins in mammalian cells [31] and described recently also for plants [32]. It should be noted that aggresome-like structures were observed in some cells also at 3 dpi (data not shown). Therefore, similarly to previous bombardment experiments with two expression vectors [16], agrobacterium-mediated expression of PMTV TGBp1, TGBp2 and TGBp3 from a single binary vector resulted in targeting of GFP-TGBp1 to cell wall-associated structures. However, in the case of agroinfiltration, localization of GFP-TGBp1 to large membranous ‘intermediate’ bodies, or PMBs, described earlier [16], was not observed. These structures therefore could have arisen due to TGBp2 and/or TGBp3 overexpression and represented artificial perturbations of cellular membranes [8]. On the other hand, observations of infiltrated leaves at different time points revealed temporal localization of GFP-TGBp1 to different subcellular structures. Importantly, in these experiments association of GFP-TGBp1 with MT-like structures always preceded its detection in cell wall-associated sites.

Fig. (1). Localization of PMTV TGBp1 fused to GFP in bombarded N. benthamiana epidermal cells imaged 24 hours post bombardment. A and B, localization of GFP-TGBp1. B, single optical section in the cortical region of cell expressing GFP-TGBp1. C, co-expression of GFP-TGBp1 with YFP-Tal. D, co-expression of GFP-TGBp1 with YFP-MAP4. In C and D, GFP signal is shown in the left panel, YFP signal (the middle panels) was digitally pseudocolored with red to facilitate interpretation of merged images (the right panels). The images, except B, are reconstructed by superposition of series of confocal optical sections. Scale bars: A, 20 μm; B, 5 μm; C and D, 10 μm.
Co-Localization of GFP-TGBp1 with Plasmodesmata

To determine whether the GFP-TGBp1-containing cell wall-associated structures are related to plasmodesmata, plasmodesmata-associated callose was visualized by staining with sirofluor, a specific fluorescent dye [27]. Leaves infiltrated with 35S:GFP-TGBp1-NOS:TGBp2/TGBp3 construct were stained at 3 dpi and analyzed by fluorescent microscopy. Two-color imaging revealed stained callose depositions along cell walls, and some of them coincided with GFP-TGBp1-containing structures (Fig. 4A). These data confirmed directed targeting of GFP-fused PMTV TGBp1 to plasmodesmata in the presence of TGBp2 and TGBp3.

Additionally, we analyzed whether the TGBp1-contained structures are peripherally associated with plasmodesmata or localize within cell wall-crossing microchannels and/or plasmodesmata central cavity. To this end, the portions of N. benthamiana leaves infiltrated with 35S:GFP-TGBp1-NOS:TGBp2/TGBp3 construct were subjected to plasmolysis by mannitol treatment, and epidermal cells were studied microscopically. After the treatment, the plasma membranes were retracted from cell walls, and gaps between them were visible in the bright field (Fig. 4B, arrows). Fluorescent microscopy revealed that after plasmolysis many GFP-TGBp1-containing bodies retained in cell walls (Fig. 4B, arrowheads). Thus, TGBp1-containing structures reside within cell walls, most probably - within plasmodesmata central cavity.

Colchicine Treatment Abolishes TGBp1 Plasmodesmata Targeting

Upon leaf agroinfiltration with 35S:GFP-TGBp1-NOS:TGBp2/TGBp3 construct, GFP-TGBp1 association with MTs preceded to its localization to plasmodesmata. Therefore we verified whether interaction with MTs is required for TGBp1 intracellular transport to plasmodesmata. To this end we analyzed the transport of GFP-TGBp1 to plasmodesmata in cells where MTs were disrupted by colchicine. In this series of experiments, colchicine was added to an infiltration buffer prior to agroinfiltration of N. benthamiana leaves with 35S:GFP-TGBp1-NOS:TGBp2/TGBp3 construct. In parallel, control leaves were infiltrated with 35S:YFP-MAP4 and treated with colchicine. At 3 dpi, microscopy of YFP-MAP4-agroinfiltrated leaves treated with colchicine revealed diffused fluorescence in the cytoplasm and in the nucleus (Fig. 5B). Additionally, as in non-treated leaves, YFP-MAP4 was detected in the nucleolus (Fig. 5A, B). In leaves agroinfiltrated with 35S:GFP-TGBp1-NOS:TGBp2/TGBp3 construct MTs, MT-associated bodies, plasmodesmata and aggresome-like structures visible in non-treated leaves at 3 dpi (Fig. 5C) were not observed in colchicine-treated leaves at 3 dpi (Fig. 5D) and upon further observations at 4 dpi (data not shown). In these samples GFP-TGBp1 was found in the cytoplasm and in the nucleus. Surprisingly, GFP-TGBp1 was localized in the cytoplasm not diffusely
Fig. (3). Localization of PMTV TGBp1 fused to GFP in epidermal cells of *N. benthamiana* leaves agroinfiltrated with the construct 35S:GFP-TGBp1-NOS:TGBp2/TGBp3. Cells were imaged at 2 days post infiltration (dpi) (A), 3 dpi (B and C) and 4 dpi (D). The images, except C, are reconstructed by superposition of series of confocal optical sections. C represents a single optical section in the cortical cell region. Scale bars: A, B and D, 20 μm; C, 5 μm.

Fig. (4). Co-localization of GFP-TGBp1 with cell wall-embedded plasmodesmata structures in epidermal cells of *N. benthamiana* leaves agroinfiltrated with the construct 35S:GFP-TGBp1-NOS:TGBp2/TGBp3. A, GFP-TGBp1 co-localization with callose stained with sirofluor-containing aniline blue dye; GFP signal is shown in the left panel, callose signal (the middle panel) was digitally pseudocolored with red to facilitate interpretation of the merged image (the right panels). Arrows in the right panel point to overlapping signals. B, GFP-TGBp1 localization in cells subjected to plasmolysis by mannitol treatment. Left panel shows the GFP signal, the middle panel - a bright field image, the right panel - superposition of the images. Arrows show the plasma membranes retracted from cell walls. Arrowheads point to GFP-TGBp1-containing structures retained in the cell wall. Scale bars, 15 μm.
but in association with an irregular network located in cell periphery and closely resembling the cortical endoplasmic reticulum (ER) (Fig. 5D). Further experiments are being carried out to analyze the TGBp1 potential to interact with ER membranes in different experimental conditions. Whatever the nature of the observed ER-like structures, our data show that disassembly of cell MTs by colchicine prevented GFP-TGBp1 targeting to plasmodesmata and the MT-dependent aggresome formation.

Deletion Analysis Reveals Correlation Between Microtubule Association and Plasmodesmata Targeting

To identify protein regions involved in TGBp1 association with MTs, a series of TGBp1 mutants was constructed. They included four deletion mutants containing deletions in the TGBp1 N-terminal extension region, dN with the deletion of residues 1-34 including a stretch of positively charged residues, Mlu1 with deletion of residues 37-85 described previously [16], Mlu2 with deletion of residues 37-184, and HEL with deletion of residues 1-187 (Fig. 6). Additionally, three point mutants were constructed, GKN with T to N substitution in the conserved motif I of the TGBp1 NTPase/helicase domain, and mutants C2 and C3 with substitutions in the motif VI (Fig. 6). T to N mutations in NTPase motif I (Walker A nucleotide-binding site) are well documented to result in much higher affinity of NTPases for ADP/GDP than for ATP/GTP thus disabling energy conversion even in the presence of ATP and blocking therefore the protein enzymatic/regulatory functions [33, 34]. Mutation introduced in C2 and C3 fall into a region potentially involved in interactions with TGBp2 or TGBp3 which are necessary for the TGBp2/TGBp3-directed targeting of TGBp1 to plasmodesmata [16]. The mutant TGBp1 genes were cloned into pLH-35S:GFP-TGBp1-NOS:TGBp2/TGBp3 to replace the wild type sequence, the resulting vectors were transformed into A. tumefaciens, and the obtained agrobacterial cultures were used for infiltration of N. benthamiana leaves. Association of mutants with subcellular structures was analyzed by microscopy at 2, 3 and 4 dpi and compared to that of the wild-type protein. It was found that the mutations dN, Mlu1, GKN, C2 and C3 had, to different extent, a suppressive effect on the GFP-TGBp1 ability to interact with MTs, to localize to MT-associated aggresome and to be targeted to plasmodesmata; however this group of mutants retained the ability of wild-type protein to interact with these cell sub-structures (Table I). In contrast, the mutants Mlu2 and HEL completely lost the ability for association with MTs and plasmodesmata targeting (Table I;
Fig. (6). Schematic presentation of mutations introduced in PMTV TGBp1. The protein amino acid sequence is shown as the filled box. The dark grey region indicate the position of the NTPase/helicase sequence domain with the seven conserved motifs I - VI. The box labelled with ‘+’ indicates the positively charged stretch in the N-terminal protein region. Horizontal arrows indicate deletions; the positions of residues at the endpoints of deleted regions are indicated above the arrows. The original sequences and the mutated sequences resulting from point substitutions in mutants GKN, C1 and C2 are shown in boxes.

These data show the correlation between these two interactions characteristic of PMTV TGBp1 and support the colchicine inhibition data pointing to the dependence of TGBp1 plasmodesmata targeting on the integrity of cell MTs. Additionally, one can conclude that the TGBp1 region comprising residues 86 - 184 is involved in the protein association with MTs. This region includes a C-terminal portion of the TGBp1 N-terminal natively unfolded domain and most of the internal structured domain [10]. Amino acid sequence alignment of TGBp1s encoded by four pomoviruses revealed that this region exhibits considerable sequence conservation in its C-terminal portion corresponding to the protein internal domain, whereas the N-terminal domain appeared to be weakly conserved even among these closely related viruses (Fig. 8). This observation suggests that the PMTV TGBp1 internal domain can be involved in specific interactions with MTs.

Table 1. Association of GFP-TGBp1 with Sub-Cellular Structures in N. benthamiana Leaves Infiltrated with 35S-GFP-TGBp1-NOS:TGBp2/TGBp3 Construct and its Derivatives Carrying Mutations in the TGBp1 Gene

| TGBp1 | MTs, 2 dpi | MT-Associated Bodies, 3 dpi | Plasmodesmata, 4 dpi |
|-------|------------|-----------------------------|---------------------|
| wt    | +++        | +++                         | +++                 |
| dN    | +          | +                           | +                   |
| Mlu1  | +          | +                           | +                   |
| Mlu2  | -          | -                           | -                   |
| HEL   | -          | -                           | -                   |
| GKN   | +          | ++                          | +                   |
| C2    | ++         | +++                         | ++                  |
| C3    | +          | ++                          | +                   |

**DISCUSSION**

In this paper we report co-localization of GFP-fused PMTV TGBp1 with MTs. Thus, TGBp1 can be added to a list of plant virus MPs capable of interaction, direct or indirect, with MTs. The most extensively studied MP encoded by TMV is among these proteins. During infection, TMV MP interacts with endoplasmic reticulum (ER) membranes and MTs, as well as localizes to plasmodesmata and induces an increase of their size exclusion limit [3, 4, 35]. TMV MP also has the capacity to bind single-stranded nucleic acids *in vitro*, and the movement of viral RNA genome likely occurs in the form of vRNP [3]. Thus TMV MP shares with PMTV TGBp1 at least two activities involved in formation of vRNPs and interaction with cortical MTs [8, 35]. TMV MP sequences required for interaction with MTs are identified by mutation analyses [36, 37]. Moreover, a region within TMV MP is revealed that shares sequence similarity to tubulins, and single amino acid substitutions within this so-called ‘tubulin mimicry domain’ are sufficient to confer temperature-sensitive loss of MT association pointing to the possibility that TMV MP is capable of making binding contacts equivalent to tubulin-tubulin interaction [38]. These data, together with biochemical evidences obtained *in vitro* [39-41], indicate that TMV MP can interact with tubulin and MTs directly. Although we were unable to find a sequence similar to the ‘tubulin mimicry domain’ in PMTV TGBp1 (data not shown), deletion analysis suggest that the TGBp1 region encompassing residues 86 - 184 is involved in MT association. Assuming that the predicted subdomain organization of PMTV TGBp1 is very similar to that of hordeivirus TGBp1 [10], one can conclude that among three domains of the hordei-like TGBp1s the structured ID domain is responsible for the protein association with MTs. Taking into account our preliminary data indicating that PMTV TGBp1 can interact with tobacco tubulin A1 *in vitro*, we presume the direct physical TGBp1 interaction with MTs.
What is the possible significance of the targeting of PMTV TGBp1 to the MTs? Recent findings illuminated the role of MTs in the intracellular transport of TMV vRNPs. Using a TMV derivative encoding a temperature-sensitive TMV MP, small mobile MT-proximal particles representing MP-RNA complexes were revealed in cells at the leading front of infection \[41, 42\]. Importantly, the TMV MP-containing particles formed upon MP transient expression in the absence of viral infection are functionally related to the particles observed in virus-infected cells; these particles contain RNA and are thought to be assembled on the MT network; their anchorage to MT sites and release can contribute to the intracellular transport of these specific RNPs to plasmodesmata \[41\]. Similarly to TMV MP, PMTV TGBp1 was also found in granular bodies associated with MTs. However, in contrast to the TMV MP, we were unable to detect movement of TGBp1-containing bodies. Nevertheless, association of TGBp1 with MTs and its localization in MT-associated granular bodies can be necessary for the protein targeting to plasmodesmata in the presence of TGBp2 and TGBp3, since treatment with the microtubule-disrupting drug, colchicine, inhibits localization of TGBp1 to plasmodesmata. One possible function of TGBp1 association with MTs is that the MT network may provide a scaffold for assembly of TGBp1-containing granular bodies as an initial step in TGBp1 targeting to plasmodesmata by TGBp2 and TGBp3. It should be noted that the *Cauliflower mosaic virus* proteins P2, P3 and P6 also form bodies that localize to MTs and, moreover, require intact MTs for their efficient formation \[43, 44\], lending support to the possibility that a similar mechanism may be utilized for the formation of MT-associated granular structures containing PMTV TGBp1.

Additionally, localization to MTs could facilitate TGBp1 degradation. The degradation of *Potato leafroll virus* MP, for example, was found to require intact MTs, and a similar model has been proposed for the MT-associated degradation of TMV MP \[32, 45\]. It was suggested that individual MP molecules or their small aggregates could traffic along MTs to stationary sites of the formation of aggresome-like structures \[32\]. Indeed, a similar model has been proposed for the formation of aggresomes, perinuclear structures formed in animal cells in response to protein overproduction and misfolding \[46\]. Evidently, transient expression of PMTV TGBp1 alone or together with TGBp2 and TGBp3 induced formation of typical single perinuclear aggresome-like structure in transformed cell. Moreover, colchicine
blocked both TGBp1 localization to plasmodesmata and formation of perinuclear aggresomes.

Current view of TMV MP intracellular transport is based on the fact that ER membrane proteins can laterally diffuse within the membrane [47] and move from cell to cell [4, 46]. Since the ER is tightly associated with the actin network [49], microfilaments could play an important role in supporting the ER-mediated targeting of MP and/or vRNA to plasmodesmata. Assuming that in plant cells MTs and actin cables may co-align at cell periphery [50] it can be proposed that the MP or viral transport RNP, initially anchored to MT-proximal sites, may be able to detach and undergo movements by an ER-driven mechanism [41]. Our data indirectly support applicability of a similar model to the PMTV transport system. After the colchicine treatment GFP-TGBp1 was localized in the cytoplasm not diffusely but in association with an irregular network closely resembling the cortical ER, suggesting its potential ER binding ability. Although the nature of these ER-like structures remains to be established, one can speculate that TGBp1 association with the ER could be due to integral membrane proteins TGBp2 and/or TGBp3 associated with the ER or ER-derived membrane compartments [7].

CONCLUSION

To conclude, we suggest the model of PMTV TGBp1 functioning emphasizing the important role of MTs. Early in infection TGBp1 can laterally associate with MTs to form MT-associated granular bodies. After some amounts of TGBp2 integrated into ER membranes are accumulated in infected cell, TGBp1-containing granular bodies, due to the suggested TGBp1 ability to interact with TGBp2 [16], are directed to the sites of TGBp2 localization at close contacts between MTs and ER. It can be speculated that the TGBp1-containing granular bodies include also RNA because of RNA binding activities of TGBp1 and TGBp2 [8]. Increase of TGBp3 amount results in a rapid ER-driven movement of TGBp1/TGBp2/RNA complexes to plasmodesmata and through plasmodesmata to neighboring cells. Further increase of concentration of MT-bound TGBp1 results in its targeting to aggresomes for degradation that can be important at the later steps of infection cycle when virions are forming, and excessive amounts of RNA-binding protein TGBp1 can compete with the coat protein for viral genomic RNAs. Thus, we propose that TGBp1 interaction with MTs may be important for both the formation of vRNP bodies early in infection and degradation of the excessive TGBp1 at late infection stages.

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