Phragmoplastin Polymerizes into Spiral Coiled Structures via Intermolecular Interaction of Two Self-assembly Domains*

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Phragmoplastin, a high molecular weight GTPase belonging to the dynamin superfamily of proteins, becomes associated with the cell plate during cytokinesis in plants. Growth of the cell plate requires continuous fusion of vesicles, and phragmoplastin appears to play a role in the formation of vesicle-tubule-vesicle structures at the cell plate. In this study, we have demonstrated that two self-assembly domains (SA1 and SA2) are involved in polymerization of phragmoplastin. SA1 is about 42 amino acids long and is located near the N terminus overlapping with the GTP-binding region. SA2, containing at least 24 amino acids, is located in the middle of the molecule outside the GTP-binding domain. Peptides containing either SA1 or SA2 interact efficiently with the full-length phragmoplastin. The SA1 domain of one phragmoplastin molecule also binds to SA2 of another as confirmed in vitro by using radiolabeled peptides. This interaction leads to the formation of polymers with a staggered contoured spiral structure. Electron microscopy studies revealed that helical arrays of phragmoplastin can be induced by reducing salt concentration. Our results suggest that phragmoplastin may assemble into helical arrays that wrap around and squeeze vesicles into vesicle-tubule-vesicle structures observed on the forming cell plate.

Phragmoplastin is a GTPase with molecular mass of 68 kDa and shares significant overall amino acid sequence similarity, particularly in the N-terminal half, with the members of the dynamin superfamily. Dynamin-like proteins have been isolated from a variety of eukaryotes, ranging from yeast to plants to humans. These proteins are mostly membrane-bound with different subcellular locations and have been implicated in diverse biological processes, including vesicle formation, protein sorting, resistance to viral infection, mitochondrial and chloroplast biogenesis, endocytosis, and cell plate formation (1–5). Phragmoplastin was shown to be specifically associated with cell plate during cytokinesis in plants (1, 6). Three Arabidopsis cDNAs encoding dynamin-like proteins (aG68, ADL1, and ADL2) have been identified, of which ADL1 and ADL2 have been implicated in chloroplast biogenesis (3, 7, 8). Similar to yeast Vps1 and human Drp1 (4, 9, 10), phragmoplastin lacks a C-terminal proline-rich domain that interacts with Src homology 3 (SH3) proteins (11). Phragmoplastin also lacks the pleckstrin homology (PH) domain that mediates the binding of dynamin to membranes containing acidic phospholipids (12).

To self-assemble into helical arrays is one of the common features shared by dynamin family of proteins. One class of the dynamin homologues, Mx proteins, can polymerize into helical structures that are 11 nm thick and 100–150 nm long (13). Mx proteins contain a “self-assembly” domain that is conserved in the dynamin family of proteins (14), including phragmoplastin (1). Dynamins can form helical structures both in solution and in vivo (15, 16). This property enables dynamin to wrap around the base of clathrin-coated pits and constricts their neck upon addition of GTP (17, 18). Dynamin alone is sufficient for the formation of constricted necks of coated pits (17). By hydrolyzing GTP, it functions as a force-generating molecule responsible for the fission of membrane vesicles. The presence of lipid promotes the formation of polymeric complexes of dynamin (15). Self-assembly of dynamin does not require guanine nucleotides, but GTP binding promotes self-assembly, whereas GTP hydrolysis may trigger disassembly of dynamin polymers (17). Assembly of dynamin into spirals can be induced by the addition of GDP and γ-phosphate analogues under physiological conditions (19). Multiple peptide domains have been implicated in self-assembly of dynamin and its homolog proteins, including Dnm1p/Vps1p-like (DVLP). Whereas the C-terminal proline rich domain of dynamin is not required for self-assembly (19), DVLP contains a conserved domain DVH2 (for Dnm1p/Vps1 homology 2) at the C terminus that is involved in oligomerization of DVLP (20). The process of dynamin self-assembly may be accounted for by both intra- and intermolecular interactions between three different protein domains (21). Coiled-coils have also been demonstrated to be involved in dynamin self-assembly (22). However, they may not play a role in phragmoplastin self-assembly, because the probability of coiled-coil formation in phragmoplastin is very low (only 20%). These data suggest that members of the dynamin family may form helical structures using different domains that provide intra- and intermolecular interactions.

Assembly and disassembly of dynamin-like proteins on the surface of biological membranes may provide a mechanism to control vesicle budding, vesicle fusion, and the formation of new membrane systems, such as vesicle-tubule-vesicle structures, that occur during cell plate formation. Phragmoplastin appears to be a “simpler” molecule than dynamin and Mx proteins because it does not contain the proline-rich domain or leucine zipper at the C terminus and lacks the pleckstrin homology domain. Nevertheless, this protein retains the ability to form oligomers of its own and forms complexes with other proteins (6). Using a yeast two-hybrid system in combination with an in vitro protein-protein interaction assay, we identified two domains that are involved in phragmoplastin oligomeriza-

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Self-assembly of Phragmoplastin

EXPERIMENTAL PROCEDURES

Bacteria and Yeast Strains—Escherichia coli DH5α and Top10F were used for plasmid construction. HB101 was used to recover plasmids from yeast cells. Yeast (Saccharomyces cerevisiae) Y190 (MATa, ura3-52, his3-200, leu2-3, 112, gal4, α, cyr1/2, lys2–801, ade2–101, trp1–901, leu2-3, 112, gal4, α, cyr1/2, lys2–801, HIS3, URA3, GAL1, GAL1, GAL1, lacZ) was used as the host for the two-hybrid experiments.

Plasmid Construction—Full-length phragmoplastin cDNA (pSDL12α(1)) was partially digested with BgII and XhoI, and the 2-kb fragment was ligated into the pAS1-CYH2 and pACT2 vectors, generating pAS-Phr and pACT-Phr, respectively. The BglII-HindIII fragment was also inserted into the BamHI-Sall sites of pUC18. The resulting plasmid (pUC-Phr) was digested with SmaI-BglII fragment (0.57 kb) from pUC-Phr into the SnaBI-BamHI sites of pACT2. A HindII fragment was released from pUC-Phr, followed by self-ligation, generating pUC-HC2A. pACT-HC2 containing the SA2 domain was made by subcloning a SnaBI-HindIII fragment (0.9 kb) into pACT2. For construction of pACT-H3, a SnaBI-HindIII fragment (0.75 kb) was transferred from pUC-HC2A to pBlue-script II SK+ to give pBS-HC2A, from which a SnaBI-XhoI fragment was subcloned into the same sites of pACT2. pACT-DRI1 was constructed by cloning a Drai-BamHI fragment (0.62 kb) into the SnaBI-BamHI sites of pACT2.

Yeast Two-hybrid Assay—To assay protein-protein interaction, cDNAs encoding the respective peptides were subcloned into pAS2 in-frame with the DNA-binding domain of Gal4 and into pACT2 in-frame with the transcription activation domain. Yeast Y190 was transformed with the two plasmids by electroporation followed by selection on yeast medium containing complete medium SC-Trp-Leu-His. Colonies were transferred to interaction medium (SC-Trp-Leu-His containing 30 mg/liter 3-amino-1,2,4-triazole). Colonies growing on SC-Trp-Leu medium were assayed for transcription activation domains (pAS-Phr and pACT-Phr, respectively; pAS-Phr; 3, 100 munits were added to 1 ml of interaction buffer (20 mM Tris-Cl, pH 8.0, 100 mM KC1, 2 mM MgCl2, 0.5% lutarol, 0.5% bovine serum albumin) and incubated with 10 µl of purified GST fusion protein bound to glutathione-agarose beads for 1 h. The beads were washed five times with the interaction buffer without bovine serum albumin, and the bound proteins were eluted directly in SDS-polyacrylamide gel electrophoresis sample buffer. After electrophoresis, the gel was stained with Coomasie Blue R-250, dried, and autoradiographed.

RESULTS

Intermolecular Interaction of Phragmoplastin in the Yeast Two-hybrid System—Using a cross-linking approach, we previously reported that soybean phragmoplastin forms homologimers (6). This was explained by the presence in phragmoplastin of a self-assembly domain similar to that described in Mx proteins (28). During the course of screening for phragmoplastin-interacting proteins using the yeast two-hybrid system, we isolated from an Arabidopsis library several isoforms of phragmoplastin-like proteins that lacked the N-terminal half of the molecule, where the self-assembly domain is located. These results suggested that an additional domain(s) must exist in the second half of the protein that allows intermolecular interaction. We subcloned soybean phragmoplastin cDNA in both pAS2 and pACT2 and determined the domains involved in the assembly of phragmoplastin that can be assayed using the two-hybrid system. Data from Fig. 1 demonstrate that cells expressing phragmoplastin from both vectors grew well on His-; SC medium containing 30 mM SC-Trp-Leu-His and 30 mg/liter 3-amino-1,2,4-triazole, suggesting that an interaction occurs between phragmoplastin molecules. When phragmoplastin was fused only with the DNA-binding domain in pAS2, or only with the transcription activation domain in pACT2, or when two empty vectors were used, the yeast cells were not...
able to grow on His− SC medium. The self-interaction was very strong, as revealed by X-Gal reaction that took less than 1 h to develop blue color (data not shown). β-Galactosidase activity assays produced consistent results with His selection experiments and X-Gal reaction. This interaction was equally strong when the self-assembly domain was missing (see below), suggesting the presence of a new domain that has not yet been described in the dynamin family of proteins.

Two Self-assembly Domains Are Involved in the Polymerization of Phragmoplastin—A set of deletion constructs were made to delimit peptide domains required for self-interaction of phragmoplastin. The BgII fragment (amino acids 1–185) contains a peptide domain (amino acids 49–93) that was previously thought to be responsible for phragmoplastin self-interaction because this domain is homologous to the self-assembly domain (amino acids 51–96) of mouse Mx1 protein (13). The deletion fragments were subcloned in pACT2 vector in-frame with the transcription activation domain (ACT) of pACT2. Transformants from each strain were streaked on X-Gal plates.

Full-length phragmoplastin served as a positive control in this experiment. Fig. 4 shows that SA1 does not have binding affinity toward itself but interacts with SA2. SA2, on the other hand, may interact with itself and SA1. To exclude the possibility that SA2 region may activate transcription on its own, we used “empty” vector pACT2 as a negative control plasmid along with pAS-SA2 to transform the yeast cells. These cells were negative in X-Gal assay, suggesting that SA2 cannot act as a transcription activator and that the X-Gal staining is brought about through protein-protein interaction. Although the importance of SA2-SA2 interaction in phragmoplastin polymerization is not clear at this point, the interaction between SA1 and SA2 could be significant as it may provide a novel mechanism for the formation of phragmoplastin polymers in a staggered helical array (see under “Discussion”).

To test whether any other protein component(s) may be required for the interaction between two phragmoplastin fragments, we used radiolabeled peptide from in vitro translated SA2 to react with purified phragmoplastin deletion fragments. Full-length phragmoplastin and the BgII and HincII fragments were cloned in pGEX-KG vector in-frame with GST. The recombinant proteins were expressed in E. coli and purified as GST fusion proteins using glutathione-agarose beads. The 35S-labeled peptides were obtained by in vitro translation and were reacted with purified GST fusion proteins bound to glutathione-agarose beads. As shown in Fig. 5, GST alone did not react with 35S-labeled SA2. GST fusion with SA1, SA2, and the full-length phragmoplastin interacted with 35S-labeled SA2 product, suggesting that SA2 can interact with SA1, SA2, and the full-length phragmoplastin. These results are consistent with that obtained in vivo using the yeast two-hybrid system and provide further evidence for the intermolecular interaction between SA1 and SA2 domains. More importantly, this experiment demonstrated that such an interaction occurs through direct contact and that no additional protein is required.

Phragmoplastin Assembles into Helical Structures—Purified phragmoplastin under high salt conditions (HCB150) did not polymerize and appeared to exist predominantly as monomers and dimers, as shown by negative-stain electron micrographs (Fig. 6A). Polymerization could be induced by dialysis against a low salt buffer (HCB15; Fig. 6, B and D). The presence of 10 μM GTP reversed the polymerization process and disassembled the helical structures into oligomers (data not shown). Incubation with GTPγS, a nonhydrolyzable analog of GTP, maintained the helical structures, suggesting that hydrolysis of GTP by phragmoplastin GTPase results in disassembly of helical arrays. Comparison of the helical arrays formed with or without the presence of GTPγS revealed that addition of GTPγS makes the helical arrays into more compact structures (Fig. 6, B and C). These results may be improved by using native phragmoplastin peptides. These data demonstrate that assembly of phragmoplastin into higher order helical structures can be induced by reducing salt concentration in the buffer without the presence of GTP. Hydrolysis of GTP appears to alter the stereo-structure of helical arrays and disassembles the arrays into oligomers.

**DISCUSSION**

Identification of Two Self-assembly Domains Involved in Polymerization of Phragmoplastin—Members of the dynamin family including phragmoplastin have been shown to form polymers. These polymers are formed in a highly ordered manner, a process known as self-assembly. The process of self-assembly can be demonstrated in vitro. Native dynamin can be extracted in high salt (300 mM NaCl) buffer. When salt concentrations were reduced to 25–50 mM by dialysis, dynamin spontaneously self-assembled into a mixture of partial rings, intact rings, and small stacks of interconnected rings (15). The ring...
stacks can further assemble into striking tubules of dynamin that can be reversibly disassembled into simple structures. Similar helical arrays of dynamin have also been observed in vivo, being bound to microtubules (29, 30), or as coats surrounding tubular membrane invaginations in nerve terminals (16, 31). We dissected the structural components required for self-assembly of phragmoplastin and demonstrated here that two self-assembly domains (SA1 and SA2) are involved in this process. One of them, SA1, is homologous to the so-called self-assembly domain identified in Mx1 and other dynamin family of proteins (14). SA1 is located between consensus sequence I (GXXXXGK(S/T)) and consensus II (DXXG) of the tripartite GTP-binding motif. SA1 is 48 amino acids long in mouse Mx1 but is 6 amino acids shorter in phragmoplastin. It consists of two parts: region 1 is highly conserved among members of the dynamin family and contains the “dynamin signature” motif (LP(R/G)(S/T/N)(G/N)IVTR). Region 2 of SA1 is variable in length and has little homology among dynamin-like proteins (Fig. 3). Our data demonstrated that SA2 (24 amino acids long) is necessary for phragmoplastin self-interaction. However, it is not clear whether this fragment of 24 amino acids alone is sufficient for the interaction. This region could be part of a longer domain. SA2 shows limited homology among dynamin-like proteins. It is not known whether the homologous SA2 domain in dynamins is indeed involved in self-assembly.

**Phragmoplastin Forms Polymers through Staggered Interactions of Intermolecular SA1 and SA2 Domains**—Human MxA contains a C-terminal leucine zipper that can interact with an internal region of the MxA molecule. Intermolecular interaction between the leucine zipper and the internal region leads to the formation of MxA polymers (32). Human dynamin, yeast Vps1, and plant phragmoplastin do not contain a leucine zipper but may use the SA2 domain instead. We do not have evidence so far for an intramolecular interaction between SA1 and SA2, which would otherwise cause a significant conformational change (N-terminal forward folding). Our results also suggest that the SA2 domains of two phragmoplastin molecules can interact with each other. The significance of such an interaction, if taking place in planta, also remains to be elucidated. In this report, we have presented evidence obtained both from

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**FIG. 3.** Alignment of amino acid sequences of the two self-assembly domains of phragmoplastin and their homologous regions in the dynamin protein family. Amino acid sequences of soybean phragmoplastin (GmPhr), Arabidopsis dynamin-like protein 1 (AtADL1) (GenBank™ accession number L36939), AtADL2 (AF012833), human dynamin 1 (HsDyn1) (L07807), HsDyn2 (L36983), rat dynamin 3 (RnDyn3) (D14076), human MxA (HsMxA) (M30817), HsMxB (M30818), human dynamin-related protein 1 (HsDRP1) (AF000430), yeast Vps1 (ScVps1) (P21576), ScDnm1 (L40588), and ScMgm1 (X62834) were aligned using DNA Star software. The aligned regions corresponding to phragmoplastin amino acid residues 51–96 and 347–371 are presented in A (SA1 domain) and B (SA2 domain), respectively. In parentheses are numbers of amino acid residues that were omitted to maximize the alignment. The dynamin signature motif is marked by asterisks. Positions of the first amino acid residues of SA1 and SA2 in the proteins are indicated on the left.
radiographed (B). The gel was stained with Coomassie Blue R-250 (B), dried, and auto-
sample buffer and resolved by SDS-polyacrylamide gel electrophoresis. 

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Phragmoplastin cDNA frag-
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pression of phragmoplastin in plant cells could be regulated by many 
leads to the formation of a helical array (Fig. 6). Such helical arrays can be disassembled by the pres-
staggered polymeric helical structure (Fig. 6). The length of 
mechanism by which such structures are formed 
ological steps have been distinguished during cell plate for-
hydrolysis of GTP and in interaction with other proteins.

Assembly and Disassembly of Phragmoplastin into Helical Arrays Is Regulated by GTP Hydrolysis—Whereas the exact molecular shape of the purified native phragmoplastin remains to be determined through x-ray crystallography, GST-fused phragmoplastin molecules do not appear to be “rod-shaped” as shown on negative stained electron micrographs (Fig. 6A). Inter-
teractions between SA1 and SA2 of neighboring molecules may lead to the formation of a helical array (Fig. 6F). Purified phragmoplastin indeed forms helical structures when the salt concentrations were reduced slowly by dialysis to 15 mM NaCl (Fig. 6). Such helical arrays can be disassembled by the presence of GTP (data not shown) but not GTP-S. Similar helical structures have also been observed in dynamin and Mx pro-
protostomian nerve terminals or isolated endosome vesicles formed in the 
monolayer, B). The gel was stained with Coomassie Blue R-250 (B), dried, and auto-
and SA2 peptides that were adsorbed to glutathione-agarose beads. After extensive washing 
the buffer, peptides retained on the beads were eluted with SDS sample buffer and resolved by SDS-polyacrylamide gel electrophoresis. The gel was stained with Coomassie Blue R-250 (B), dried, and auto-
numbers in (from low to high) of 14, 21, 31, 45, 66, and 97 kDa. Lane numbers in B and C correspond to row numbers in A.

yeast two-hybrid system (in vivo) and in vitro experiments, to 
demonstrate that SA1 and SA2 interact intermolecularly with each other. Continuous interaction between SA1 of one mole-
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