Eukaryotic endoplasmic reticulum (ER)-plasma membrane (PM) contact sites are evolutionarily conserved microdomains that have important roles in specialized metabolic functions such as ER-PM communication, lipid homeostasis, and Ca$^{2+}$ influx. Despite recent advances in knowledge about ER-PM contact site components and functions in yeast (*Saccharomyces cerevisiae*) and mammals, relatively little is known about the functional significance of these structures in plants. In this report, we characterize the Arabidopsis (*Arabidopsis thaliana*) phospholipid binding Synaptotagmin1 (SYT1) as a plant ortholog of the mammal extended synaptotagmins and yeast tricalbins families of ER-PM anchors. We propose that SYT1 functions at ER-PM contact sites because it displays a dual ER-PM localization, it is enriched in microtubule-depleted regions at the cell cortex, and it colocalizes with Vesicle-Associated Protein27-1, a known ER-PM marker. Furthermore, biochemical and physiological analyses indicate that SYT1 might function as an electrostatic phospholipid anchor conferring mechanical stability in plant cells. Together, the subcellular localization and functional characterization of SYT1 highlights a putative role of plant ER-PM contact site components in the cellular adaptation to environmental stresses.
Under mechanical stresses, different cell types, including meristematic, expanding, and fully differentiated cells, undergo physiological changes based on the sensing and integration of various mechanical signals (Monshausen and Haswell, 2013). Although it is well established that plants sense and respond to mechanical cues, our understanding of the various molecular mechanisms by which this is accomplished is limited, and most of our knowledge relies on comparisons with mechanosensors and transduction pathways identified in *Escherichia coli* and mammalian cells (Arnadóttir and Chalfie, 2010). Currently, two nonmutually exclusive mechanosensing models, namely, the ion channel and the tensegrity models, coexist in the plant literature. In the ion channel model, plant homologs of the bacterial mechanosensitive channel of small conductance and putative stretch-activated Ca\(^{2+}\)-permeable channels are gated in response to mechanical forces and trigger a signaling cascade through the rapid influx of extracellular Ca\(^{2+}\) toward the cytosol (Arnadóttir and Chalfie, 2010; Jensen and Haswell, 2012; Sukharev and Sachs, 2012; Kurusu et al., 2013). In the tensegrity model, plant cells operate as self-supporting structures stabilized by a dynamic prestress state in which all elements are in isometric tension (Fuller, 1961). In such a structure, the mechanical disturbance of any individual element allows stress signals to propagate and be transduced at relatively distant locations (Ingber, 2008). Thus, the mechanically stable cell walls provide structural support, and the constant remodeling of the underlying cytoskeleton in response to mechanical disturbances acts as a tensegrity sensor (Komis et al., 2002; Berghöfer et al., 2009; Nick, 2013).

In this article, we characterize the Arabidopsis (*Arabidopsis thaliana*) type I anchor Synaptotagmin1 (SYT1/SYTA/NTMC2T1.1 [hereafter SYT1]; Craxton, 2010; Yamazaki et al., 2010; Lewis and Lazarowitz, 2010) as an important component required to withstand mechanical stress in plant cells. SYT1 belongs to a five-member family in Arabidopsis and shares a common modular structure with different members of the mammalian extended synaptotagmins (E-Syts) and yeast (*Saccharomyces cerevisiae*) tricalbins families of organelle tethers (Manford et al., 2012; Fig. 1A; Supplemental Fig. S1). These proteins act as molecular bridges between the ER and the PM at sites where both cellular membranes are in close proximity, called ER-PM contact sites. These specialized microdomains carry out important roles in organelle communication, lipid and Ca\(^{2+}\) homeostasis, and intracellular signaling in animal and yeast cells (Toulmay and Prinz, 2011; Helle et al., 2013; Prinz, 2014). In plants, these ER-PM contact sites have been morphologically described for decades (Staehelin, 1997), but their physiological roles have not been thoroughly characterized. Furthermore, the identity of molecular components at these sites has remained elusive until the recent characterization of the ER-PM localized Vesicle-Associated Protein27-1 (VAP27) and NETWORKED 3C (NET3C) markers in Arabidopsis (Wang et al., 2014).

In previous reports, SYT1 has been described as an essential component for PM integrity maintenance, especially under conditions of high potential for membrane disruption such as freezing or salt stresses (Schapire et al., 2008, 2009; Yamazaki et al., 2008). In these reports, SYT1 was proposed to act as a Ca\(^{2+}\)-dependent regulator of...
membrane fusion, in analogy to the classical animal SYTs that mediate Ca\textsuperscript{2+}-triggered vesicle fusion during neurotransmission (Carr and Munson, 2007). Another report highlighted a role for SYT1 in viral spreading from cell to cell (Lewis and Lazarowitz, 2010). The recent characterization of mammalian E-Syts in stress tolerance (Herdman et al., 2014) and the phylogenetic relationships of tricalbins and E-Syts with SYT1 (Craxton, 2010; Yamazaki et al., 2010) led us to hypothesize that SYT1 could be a functional ortholog of E-Syts and tricalbins in plants. Consistently, we found that it is localized in specific ER-PM subdomains in cortical cytoskeleton-depleted regions, it colocalizes with the VAP27 marker, and it anchors negatively charged phospholipids through its PM-targeted C2 domains. Additionally, we show that SYT1 loss of function causes mechanical instability at the tissue and cellular level without altering the gross ER morphology. Based on these findings, we conclude that SYT1 acts at ER-PM contact sites as part of structural platforms adjacent to the cortical cytoskeleton that are required for mechanical stress tolerance in Arabidopsis.

RESULTS
The Arabidopsis SYT1 Localizes at the ER and the PM and Accumulates at ER-PM Contact Sites

SYT1 contains a modular structure similar to ER-PM tether proteins such as E-Syts and tricalbins (Craxton, 2010; Yamazaki et al., 2010), proteins that are localized at specific ER-PM subdomains (Giordano et al., 2013; Helle et al., 2013; Prinz, 2014). Moreover, cell biological and biochemical analyses have reported SYT1 being at the ER, PM, and plasmodesmata (Schapire et al., 2008; Lewis and Lazarowitz, 2010; Yamazaki et al., 2010). To unequivocally establish the SYT1 subcellular localization, we developed new polyclonal anti-SYT1\textsubscript{244-541} antibodies (hereafter, anti-SYT1), and demonstrated their specificity by the absence of signal in the null syt1-2 mutant (hereafter, syt1) via western blot and immunolocalization (Supplemental Fig. S1). Next, we performed immunolocalization analyses in transgenic Arabidopsis lines expressing the GFP-tagged ER retention peptide HDEL (HDEL-GFP), and the PM markers PIN-FORMED2 (PIN2-GFP), BRASSINOSTEROID INSENSITIVE1 (BR1-GFP), and PLASMA MEMBRANE INTRINSIC PROTEIN 2A (PIP2A-GFP). As shown in Figure 1B, the anti-SYT1 perinuclear signal colocalized with HDEL-GFP, confirming the broad ER localization of SYT1. Interestingly, at the cell cortex, the anti-SYT1 signal was characterized by a punctate pattern that was closely associated with HDEL-GFP but did not colocalize (Supplemental Fig. S1). Remarkably, in those cortical regions, the anti-SYT1 signal colocalized with PM markers PIN2-GFP, BR1-GFP, and PIP2A-GFP (Fig. 1C; Supplemental Fig. S2), suggesting a dual ER-PM localization. To analyze the SYT1 subcellular localization in leaf epidermal cells, we generated transgenic Arabidopsis lines expressing a C-terminal translational fusion of GFP to SYT1 (SYT1\textsubscript{proSYT1-GFP}; hereafter, SYT1-GFP). At the cell cortex, the SYT1-GFP marker appeared as a reticulate network of beads and strings (Fig. 1D) that was clearly different from the classical network of sheets and interconnected tubules observed in the luminal ER marker HDEL-GFP and the GFP-tagged membrane-associated ER marker SECRETORY 12P-LIKE-GFP (Supplemental Fig. S3). At the equatorial plane, the SYT1-GFP marker appeared as a punctuate signal at the cell periphery (Fig. 1E), and signal movement through transvacuolar strands was observed (Supplemental Movie S1). Transient colocalization in Nicotiana benthamiana of a Red Fluorescent Protein (RFP)-tagged SYT1 fused to its endogenous promoter(SYT1\textsubscript{proSYT1-RFP}) with the ER-targeted yellow Cameleon 4.60 (Iwano et al., 2009) showed that the SYT1 signal was enriched at specific ER subdomains (Supplemental Fig. S3). A high-resolution subcellular localization for SYT1 was obtained by immunogold Transmission Electron Microscopy (immunoTEM) (immunoTEM) of high-pressure frozen, freeze-substituted wild-type roots probed with the anti-SYT1 antibody. Similar to previous immunoTEM results (Schapire et al., 2008), the anti-SYT1 signal was observed at both the ER and PM (Supplemental Fig. S4). However, the use of more specific and sensitive antibodies together with the better structural preservation of the high-pressure frozen samples enabled us to observe an enrichment of the anti-SYT1 signal in regions where the cortical ER was in close apposition to the PM (Fig. 2A),
and between cortical microtubule bundles (Fig. 2B). Since the SYT1 localization at the ER-PM interface strongly resembled that of the recently reported Arabidopsis VAP27 and NET3C ER-PM markers (Wang et al., 2014), we next performed transient colocalization studies using the RFP-tagged VAP27 fused to the Ubiquitin-10 promoter (UBQ10proVAP27-RFP) and SYT1proSYT1-GFP markers in N. benthamiana leaves. As shown in Figure 2, C and D, the markers colocalized to a great extent in both intracellular and cortical regions of the cell. Together, these data strongly suggest that SYT1 is enriched at ER-PM contact sites, a feature shared with the E-Syts and tricalbins in animals and yeast, respectively.

The SYT1-Labeled ER-PM Contact Sites Are Predominantly Stationary

Once we determined that SYT1 populations with different subcellular localizations coexist in plant cells, we analyzed the dynamics of SYT1-GFP using time-lapse and fluorescence recovery after photobleaching (FRAP) assays. The time-lapse experiments showed that the intracellular SYT1-GFP signal was highly dynamic, and different SYT1-GFP localizations within the transvacuolar strands were detected on frames separated by 15 s (Supplemental Movie S1). In contrast, the cortical SYT1-GFP signal was spatially restricted to largely stationary positions (Fig. 3A; Supplemental Fig. S5; Supplemental Movie S2). FRAP analysis of the cortical SYT1-GFP signals showed the recovery of 50% of the signal was achieved in less than 5 min (Fig. 3, B and C; Supplemental Movie S3). Notably, the recovered signal positions strongly resembled those marked by the prebleached SYT1-GFP, further supporting the nearly stationary nature of the SYT1-GFP cortical sites (Fig. 3D; Supplemental Fig. S5; Supplemental Movie S4).

The ER-PM Contact Sites Localize in Microtubule-Depleted Regions at the Cell Cortex

In actively dividing root epidermal cells, the SYT1 signal transitioned from the ER to the mitotic spindle at metaphase and was delivered to the forming cell plate during cytokinesis (Fig. 4A). This result suggests that the spatial localization of SYT1 might be partially controlled by the microtubule cytoskeleton. To further investigate this possible link, we performed colocalization studies of

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**Figure 3.** The SYT1-labeled ER-PM contact sites are predominantly stationary. A, SYT1-GFP signal comparison at the cortical regions of 8-d-old shoot epidermal cells during a 600-s time-lapse experiment. The initial (t = 0 s) and final (t = 600 s) frames are shown. Scale bars = 40 μm. B and C, Time course FRAP showing the recovery of the SYT1-GFP cortical signal at the equatorial plane (B) or cortical plane (C) in 8-d-old shoot epidermal cells. Scale bars = 40 μm. D, Pre- and post-FRAP SYT1-GFP signal comparison at the cortical regions of 8-d-old shoot epidermal cells. The initial (prebleach) and final (postbleach) frames of a representative 600-s FRAP experiment are shown. Scale bars = 40 μm.
SYT1 with different MT markers. In roots, the anti-SYT1 and the GFP-tagged mammalian MICROTUBULE-ASSOCIATED PROTEIN4 (MAP4-GFP) signals were mutually exclusive, indicating that the SYT1 puncta were mainly located within MT-depleted regions (Fig. 4B). The same result was obtained when the anti-SYT1 signal was compared with that generated by a monoclonal antibody that recognizes the tyrosinated form of the alpha-tubulin subunit (Fig. 4C). In shoots, we analyzed the SYT1 localization using a stable double transgenic line harboring the RFP-tagged TUBULIN6 (TUA6-RFP) and SYT1-GFP markers. As shown in Figure 4, D and E, both signals were largely separated with an estimated colocalization rate that ranged from 4% to 12%, depending on the sample and microscopy settings used (n = 24 cells). Finally, we treated lines expressing the Arabidopsis SYT1-GFP marker with cytoskeleton depolymerizing drugs. As shown in Figure 4, F to H (controls shown in Supplemental Fig. S6), F-actin depolymerization induced by latrunculin B abolished SYT1-GFP cytoplasmic streaming (Supplemental Movie S5) and caused the aggregation of the intracellular/ER SYT1-GFP pool, whereas the cortical pool remained mostly unaffected. Interestingly, oryzalin-mediated MT depolymerization changed neither the intracellular ER nor the cortical SYT1-GFP distributions in shoot epidermal cells, but changes in SYT1-GFP distributions were observed in mature guard cells (Fig. 4H).
results suggest that, once established, the ER-PM contact sites do not require the cytoskeleton scaffold to maintain their position in leaf epidermal cells, but this process might be differentially regulated in guard cells.

The SYT1 C2 Domains Are Targeted to the PM and Bind Negatively Charged Phosphatidylinositol Phosphates

Similar to most tricalbins and E-Syts (Toulmay and Prinz, 2012; Giordano et al., 2013), SYT1 contains a predicted TM domain and phospholipid binding C2 domains through which it could connect the ER to the PM in trans. To assess the accuracy of these predictions and to distinguish between the two possible SYT1 orientations (TM anchored at the ER and C2 domains tethering the PM, or TM anchored at the PM and C2 domains tethering the ER), we first analyzed where isolated SYT1 C2 domains were targeted. For this purpose, we generated stable transgenic lines harboring the SYT1244-541 peptide containing the GFP-tagged SYT1 C2 domains C2A and C2B (SYT1-C2AB; Schapire et al., 2008) fused to the 35S promoter. We also used a 35SproGFP and the SYT1-GFP lines as controls. As shown in Figure 5A, the 35SproGFP and the 35SproC2AB-GFP markers were both partially targeted to the nucleus, but whereas the 35SproGFP construct strongly labeled the cytosol, the 35SproC2AB-GFP construct was mainly targeted to the PM. This subcellular localization was further assessed by localization analysis of the different markers in plasmolyzed cells (Supplemental Fig. S7) and colocalization studies with the endocytic tracer N-(3-triethylammoniumpropyl)-4-{6-[4-(diethylamino) phenyl] hexatrienyl} pyridinium dibromide (FM4-64) that labels the PM in short-term treatments (Vida and Emr, 1995; Supplemental Fig. S8). In all cases, the subcellular localization of C2AB-GFP was different from the punctuated PM pattern observed for SYT1-GFP, suggesting that (1) the SYT1 TM domain is anchored to the ER and the C2 domains tether the PM, and (2) the SYT1 TM and SMP domains might restrict the positions where the SYT1 C2 domains bind to the PM; that is, to regions where the ER is in close contact with the PM.

In mammals, the E-Syts C2 domains have been reported to bind phosphatidylinositol phosphates (PtdInsPs) in both a Ca2+-dependent and Ca2+-independent manner (Giordano et al., 2013). Once we determined that the SYT1 C2 domains were targeted to the PM, we aimed to identify phospholipid species that might be involved in the C2-PM tethering process. Thus, we performed in vitro protein-lipid overlay assays using the purified SYT1-C2AB peptide in the presence or absence of Ca2+. In the presence of Ca2+ (10 μM Ca2+; Fig. 5B), the SYT1-C2AB peptide bound phosphatidylinerine (PS), phosphatidic acid (PA), lysosphatidic acid (LPA), and each of the negatively charged PtdInsPs, but not phosphatidylinositol (PtdIns). In the absence of Ca2+ (5 mM EGTA; Fig. 5B), the SYT1-C2AB strongly reduced the affinity for PS and slightly reduced the affinity for most of the PtdInsPs species. Together, these results suggest that the SYT1-C2AB binding to PM-localized PtdInsPs is largely Ca2+-independent and largely relies on electrostatic interactions.

SYT1 Confers Cellular Resistance to Mechanical Stress

Previous reports have shown that SYT1 loss of function causes hypersensitivity to conditions that

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**Figure 5.** The SYT1 C2 domains are targeted to the PM and bind negatively charged PtdInsPs. A, Subcellular localization of the 35SproGFP, 35SproC2AB244-541-GFP, and SYT1-GFP markers in 5-d-old shoot epidermal cells. The equatorial (left) and cortical (right) planes are shown. Scale bars = 40 μm. B, Protein-lipid overlay assay. PIP Micro Strips (Life Technologies) were incubated with 0.5 μg mL−1 of the SYT1244-541 C2AB soluble peptide in the presence (10 μM Ca2+) or absence (5 mM EGTA) of Ca2+. Binding was detected using the anti-SYT1 antibody. S(1)P, Sphingosine 1-phosphate; PE, phosphatidylethanolamine; PC, phosphatidylcholine.
generate changes in cellular turgor, such as osmotic, ionic, and freezing stresses (for review, see Schapire et al., 2009). Due to its subcellular localization and putative tethering function, we hypothesized that SYT1 might function as a structural component required to withstand external and internal physiomechanical forces. To gain further insights into this putative role, we determined whether the expression of SYT1 responds to mechanical damage using transgenic plants transformed with the reporter β-glucuronidase gene driven by the SYT1 promoter (SYT1proGUS). As shown in Figure 6A, the SYT1proGUS signal was strongly induced by mechanical damage. Next, we analyzed the behavior of the SYT1-GFP marker subjected to controlled pressure (6,750 Pa). As shown in Figure 6B, the organized SYT1-GFP reticulated structures developed into more defined SYT1-GFP aggregates after 20-s treatments. A similar behavior was observed when the HDEL-GFP marker was used, suggesting that changes in ER morphology might represent an early response to mechanical stress. Under similar conditions, the cytoskeleton MAP4-GFP marker partially depolymerized, indicating that different cortical structures were quickly responding to mechanical stress cues (Fig. 6B). To investigate whether SYT1 has a functional role in mechanical stress responses, we next analyzed the cellular stability of the syt1 mutant upon mechanical stress. First, we observed that syt1 protoplasts displayed reduced tolerance to mechanical stress compared with the wild type, suggesting that SYT1 has a role in mechanical stress adaptation that is, at least in part, cell wall independent (Supplemental Fig. S9). In leaves, the application of mechanical stress inhibited their expansion and caused more extensive tissue damage in syt1 compared with the wild type (Fig. 6C). The extent of the damage was further evaluated using trypan blue, a vital stain that cannot be excluded by dead cells. Although trypan blue stain was limited to a few individual cells within the pressed area in the wild type, the stain was observed in most cells either within or adjacent to the

Figure 6. SYT1 confers tissue and cellular mechanotolerance. A, Histochemical GUS staining of control and mechanically damaged 3-week-old SYT1proGUS plants. Plants were mechanically pressed with a 5-inch hemostate set to the first pressure level, and GUS staining was performed 1 h after the pressure treatment. B, Changes in the SYT1-GFP, HDEL-GFP, and MAP4-GFP localization upon mechanical stress load. Five-day-old seedlings were mounted on microscopy slides using water. Before imaging, a 330-g (6,750-Pa) weight was applied to the 22- × 22-mm coverslip for 20 s. Whereas the SYT1-GFP and HDEL-GFP markers form cortical aggregates when pressure approaches the cellular collapse limit, the MAP4-GFP marker shows MT depolymerization. Scale bars = 20 μm. C, Phenotype of mechanically wounded 6-week-old wild-type Columbia (Col) and syt1 rosette leaves. Pictures were taken 3 d after wounding. D, Trypan blue staining of 6-week-old wild-type and syt1 rosette leaves 1 h after wounding. E, Electrolyte leakage in 6-week-old wild-type and syt1 leaf discs. Data are the means ± SO from three replicates (n = 45 leaf discs). Student’s t test: P ≤ 0.01. F and G, Cellular collapse of syt1 epidermal cells upon mechanical load. Five-day-old Col and syt1 harboring the HDEL-GFP and MAP4-GFP markers were loaded with a 330-g (6,750-Pa) weight and imaged after 20 s. Collapsed cells are marked with asterisks. Scale bars = 50 μm.
pressed area in syt1 (Fig. 6D). This difference in damage was quantified by measuring electrolyte leakage of mechanically excised wild-type and syt1 leaf discs. Figure 6E shows that the normalized conductivity of the solution containing syt1 leaf discs was approximately 45% higher than that of the solution containing wild-type discs. Consistent with the observed mechanical stress hypersensitivity, the application of heavy mechanical loads (6,750 Pa) caused the collapse of 5% of the wild-type (n = 122) and 44% of the syt1 (n = 120 cells) epidermal cells harboring the HDEL-GFP marker, and 3% of the wild-type (n = 103) and 28% of the syt1 (n = 118 cells) epidermal cells harboring the MAP4-GFP marker (Fig. 6, F and G). Remarkably, the application of light mechanical loads (1,125 Pa) enhanced the depolymerization of the MAP4-GFP marker in syt1 (74%; n = 131) compared with the wild type (14%; n = 107), indicating that this early mechanical response was also altered in the syt1 background (Supplemental Fig. S10). Despite the multiple phenotypes associated with SYT1 depletion, no clear morphological differences at the cortical ER were observed between the wild type and syt1 (Supplemental Fig. S11). This result suggests that SYT1 is important for ER-PM function, but additional components might be required for ER-PM contact sites establishment.

DISCUSSION
SYT1 Is a Plant ER-PM Phospholipid Binding Anchor

In yeast, three conserved protein families have been identified as ER-PM tetherers: INCREASED SODIUM TOLERANCE2 (related to the mammalian anoctamin family of ion channels), the VAP proteins SUPPRESSOR OF CHOLINE SENSITIVITY2 (Scs2) and Scs22 and the tricalbin (Tcb) proteins Tcb1 to Tcb3 (orthologs of the extended synaptotagmin-like proteins E-Syt1 to E-Syt3 in mammals; Manford et al., 2012). Recently, a plant homolog of the yeast Scs2 protein, VAP27, was reported to localize at Arabidopsis ER-PM contact sites, identifying the first molecular components of such sites in plants (Wang et al., 2014). The Arabidopsis SYT1 contains the modular structure typical of tricalbins and E-Syts (Craxton, 2010; Yamazaki et al., 2010), is enriched at ER-PM subdomains, and similar to yeast where Scs2 and TcbS interact (Manford et al., 2012), it colocalizes with the ER-PM contact sites marker VAP27. Together, these results support the notion that an ER-PM tethering mechanism similar to that in yeast and mammals exists in plants.

To achieve their tethering function, most yeast tricalbins and mammalian E-Syts establish ER-PM contact sites in durable or transient states using a single amino-terminal ER anchor (Manford et al., 2012), the SMP domain that localizes them to specific ER-PM subdomains (Toulmay and Prinz, 2012), and a variable number of phospholipid binding domains (C2 domains) that connect the proteins to the PM in trans via electrostatic interactions with negatively charged phospholipids (Toulmay and Prinz, 2011; Manford et al., 2012; Giordano et al., 2013; Schauder et al., 2014). Our results suggest that the SYT1 localization at the nexus between the ER and the PM is restricted by its TM and SMP domains, and its docking to the PM mainly occurs through nonspecific electrostatic interaction with negatively charged PtdInsPs (Fig. 7A). These results highlight mechanistic similarities between the ER-PM contact sites establishment in plants and mammals, such as the Ca2+-independent binding of phosphatidylinositol-4,5-bisphosphate observed in SYT1 and the mammalian E-Syt2 and E-Syt3 (Giordano et al., 2013). There are also

![Figure 7. Putative roles of SYT1 in ER-PM contact sites. A. Docking mechanism: The SYT1 TM domain is anchored to the cortical ER (cER) and together with the SMP domain restricts the positions where the C2 domains dock to the PM. The C2 domains bind PM-localized PtdInsPs through largely Ca2+-independent electrostatic interactions. B. SYT1 responses to mechanical stress. Top left, In a wild-type (WT) background, SYT1 is enriched in MT-depleted regions, colocalizes with ER-PM marker VAP27, and contributes to the PM stabilization by mechanically reinforcing the ER-PM contact sites. Top right, In a syt1 background, SYT1 does not reinforce the ER-PM contact sites, but VAP27 and likely other ER-PM contact site components are sufficient to maintain the PM stability. Bottom left, In a mechanically challenged wild-type background, SYT1, the VAP27/NET3C complex, and the cortical MTs cooperate to distribute mechanical loads. Bottom right, In a mechanically challenged syt1 background, the VAP27/NET3C complex and the cortical MTs are not sufficient to properly distribute mechanical loads; as a result, the cortical MTs depolymerize and the PM collapses. The mechanism (black arrows) by which the ER-PM contact sites might exchange information with cortical MTs remains unclear.](https://www.plantphysiol.org/)# Figure 7. Putative roles of SYT1 in ER-PM contact sites. A. Docking mechanism: The SYT1 TM domain is anchored to the cortical ER (cER) and together with the SMP domain restricts the positions where the C2 domains dock to the PM. The C2 domains bind PM-localized PtdInsPs through largely Ca2+-independent electrostatic interactions. B. SYT1 responses to mechanical stress. Top left, In a wild-type (WT) background, SYT1 is enriched in MT-depleted regions, colocalizes with ER-PM marker VAP27, and contributes to the PM stabilization by mechanically reinforcing the ER-PM contact sites. Top right, In a syt1 background, SYT1 does not reinforce the ER-PM contact sites, but VAP27 and likely other ER-PM contact site components are sufficient to maintain the PM stability. Bottom left, In a mechanically challenged wild-type background, SYT1, the VAP27/NET3C complex, and the cortical MTs cooperate to distribute mechanical loads. Bottom right, In a mechanically challenged syt1 background, the VAP27/NET3C complex and the cortical MTs are not sufficient to properly distribute mechanical loads; as a result, the cortical MTs depolymerize and the PM collapses. The mechanism (black arrows) by which the ER-PM contact sites might exchange information with cortical MTs remains unclear.
key differences between yeast and plant contact sites, such as the Ca<sup>2+</sup>-dependent binding of PS observed in SYT1 that substitutes the Ca<sup>2+</sup>-dependent binding of phosphatidylinositol-4,5-bisphosphate in the mammalian E-Syt1 (Giordano et al., 2013). Still, important mechanistic information, such as the putative presence of SYT1 in multiprotein tethering complexes involving VAP27/NET3C and other components, and the putative enrichment of ER-PM contact sites in specific phospholipids, is not yet defined in plants.

**SYT1 Anchors the PM at MT-Depleted Regions**

Similar to the Arabidopsis ER-PM contact site markers VAP27 and NET3C, the SYT1-GFP signal appears as a highly punctuated structure reminiscent of ER-PM anchor sites in both *N. benthamiana* and Arabidopsis epidermal cells (Wang et al., 2014). FRAP and time-lapse experiments showed that these structures were remarkably stationary, providing a clear distinction between the pools of highly dynamic intracellular SYT1 and stable cortical SYT1. Interestingly, a previous study has shown that VAP27 and NET3C associate tightly to the cortical MTs and F-actin cables, directly connecting the ER to the cortical cytoskeleton at ER-PM contact sites (Wang et al., 2014). Our data support and expand this view and suggest that the cortical tubulin cytoskeleton also provides positional cues for the establishment of ER-PM contact sites. In our model (Fig. 7B), the association of SYT1 puncta with the PM is largely restricted to MT-depleted regions in both roots and shoots, possibly because the tight associations between thick MTs (25 nm wide on average; Goddard et al., 1994) and the PM could pose steric impediments to the establishment of the narrow 10- to 30-nm (West et al., 2011) ER-PM contact sites. Remarkably, the organization of the putative SYT1-GFP anchors in cortical regions is mostly unaffected by the MT-depolymerizing drug oryzalin, suggesting that once established, the ER-PM contact sites do not require the cortical MT scaffold to maintain their fixed positions. Similar to the results observed for NET3C and VAP27 (Wang et al., 2014), latrunculin B abolished the SYT1-GFP movement through transvacuolar strands, indicating that the F-actin cables are important for the proper intracellular SYT1 turnover and transport. Taken all together, we propose two different roles for the plant cytoskeleton in ER-PM contact site formation: (1) The internal F-actin network functions as a carrier of SYT1 (and probably NET3C and VAP27) toward specific contact sites in different cell types; and (2) The cortical MT array provides positional information that allows the close apposition of ER and PM membranes for the establishment of ER-PM contact sites.

**Putative Roles of SYT1 in Abiotic Stress Tolerance**

Proper thigmomostimuli perception is essential for plant growth and development, and two nonexclusive models of mechanoperception coexist in the plant literature. In the tensegrity model, mechanical deformations of the cell wall are finely modulated by the intrinsic instability of the MT cytoskeleton that senses the stress patterns. This process modifies the mechanical properties of plant cells through cellular endomembrane organization rearrangements (Brandizzi and Wasteneys, 2013) and targeted the deposition of rigidifying cell wall components (Landrein and Hamant, 2013; Sampaikumar et al., 2014). According to the tensegrity model, the SYT1-labeled ER-PM contact sites might function as deformable platforms that coordinate the cortical ER and cytoskeleton responses to minimize the mechanical loads at the PM. This model would not require the activation of transcriptional programs and can explain the instantaneous PM instability observed upon exposure to multiple stresses, including freezing, osmotic, and mechanical challenge (Yamazaki et al., 2008; Schapire et al., 2008; this study). This model could also explain the enhanced cytoskeleton depolymerization observed in the *syt1* mutant due to the limited ability of SYT1-depleted contact sites to properly transfer mechanical cues to the cortical cytoskeleton (Fig. 7B). An alternative tensegrity model for SYT1 is derived from mammalian cells where mechanical stimulation results in a reduced distance between the ER and the PM, thereby possibly enhancing transcompartment signaling and structural stabilization (Prinz, 2014). In this scenario, defects in transcompartment signaling could partially explain the general abiotic stress sensitivity observed in *syt1* that resembles the reduced survival under stress observed in mice lacking E-Syt2 and E-Syt3 tethers (Herdman et al., 2014).

Additionally, the ion-sensing model predicts that stress-inducible Ca<sup>2+</sup>-dependent sensors decode mechanically triggered Ca<sup>2+</sup> signatures (Kurusu et al., 2013). These putative sensors would transduce Ca<sup>2+</sup> signatures by selectively binding to negatively charged phospholipids localized in the PM, such as the well-known PtdInsPs signaling intermediates (Mosblech et al., 2008), or through direct interaction with putative stress-inducible Ca<sup>2+</sup>-binding partners (Hashimoto and Kudla, 2011). Our experimental results suggest that the SYT1 binding to PtdInsPs might be electrostatic and largely Ca<sup>2+</sup>-independent, which makes it an unlikely sensing mechanism; however, an intriguing possibility is that PS, and not PtdInsPs, might control downstream Ca<sup>2+</sup>-dependent responses to mechanical stress in plants. At this point, we consider that the implications of the Ca<sup>2+</sup>-dependent binding of PS during mechanical stress as well as the identification of putative stress-inducible and Ca<sup>2+</sup>-dependent SYT1 binding partners are important aspects that need to be addressed by future research.

**CONCLUSION**

Our study identifies the Arabidopsis SYT1 as an essential component required for cellular resistance to mechanical stress acting as a putative ER-PM anchor. These
findings represent progress toward the understanding of ER-PM contact site functions in plant cells. Based on our results, we propose that the SYT1-labeled ER-PM contact sites represent conserved eukaryotic platforms where the close apposition between membranes coordinates the responses at the cell cortex and provides structural support against multiple environmental stresses. Future analysis of the SYT and VAP families of ER-PM markers represents a starting point for more precise elucidation of the role of plant ER-PM contact sites in the maintenance of PM stability upon stress exposure.

MATERIALS AND METHODS

Plant Growth and Chemical Treatments

For microscopy assays, surface-sterilized and cold-stratified Arabidopsis (Arabidopsis thaliana) seeds were grown as described in Rosado et al. (2012). For drug treatments, 4-d-old seedlings grown on solid one-half-strength Murashige and Skoog salts supplemented with 1% (w/v) Suc were transferred to 12-well plates containing liquid basal medium supplemented with 1 μM latrunculin B, 25 μM oryzalin, or mock (0.1% [v/v] dimethyl sulfoxide) and incubated for 24 h. After treatments, seedlings were rinsed and mounted on microscopy slides using water.

Cloning

The SYT1 and VAP27 constructs were generated through multisite gateway cloning. A 970-bp SYT1 promoter fragment (Benhamdi et al., 2008) was cloned into pDONR221 (Invitrogen), and a 615-bp UBQ10 promoter fragment was cloned in the pENTR 5’-TOPO vector. The genomic fragments of the SYT1 and VAP27 coding sequence were amplified from Col-0 DNA and recombined into pDONR221 (Invitrogen). The ER-targeted yellow cameleon 4.60 was subcloned from pBII121 vector (Iwano et al., 2009) into the multicloning site of pGEMZf(+), using restriction fragments generated by XbaI and SacI, allowing for further subcloning into pENTR1A using the resulting Sall and EcoRI restriction fragment. For the C-terminal XF fusions (GFP, pEN-R2-F-L3; mRFP, pEN-R2-R-L3,0 [Karimi et al., 2007]; and TagRFP, pEN-R2-TagRFPSTOP-L3 [Myle et al., 2013]), the respective entry vectors were recombined together with the respective promoter and coding sequence entry vectors into pKm43GW or pBm43GW (Karimi et al., 2005). For the promoter GUS construct, the SYT1 promoter fragment was recombined with pEN-L1-S-L2 into pKm42GW (Karimi et al., 2005). The obtained constructs were transformed into Col-0 via floral dip (Clough and Bent, 1998) or were used for transient expression in Nicotiana benthamiana.

Transient Expression in N. benthamiana

Suspensions of Agrobacterium tumefaciens strain LBA4404 harboring the different constructs were grown to an optical density at 600 nm of 0.5 and infiltrated into 4-week-old N. benthamiana leaves as previously described in Goodin et al. (2002). Transiently transformed leaves were imaged 3 to 5 d after infiltration using a Pascal Excite laser scanning confocal microscope (Zeiss) equipped with a 488-nm argon laser for GFP and a 543-nm He-Ne laser for RFP. Sequential line scanning mode was used to separate signals. The SYT1-GFP images were obtained using a Zeiss Pascal Excite laser scanning confocal microscope equipped with a 488-nm argon laser for GFP and a 543-nm He-Ne laser for propidium iodide. For FRAP analysis, we adjusted the laser intensity and performed a single bleaching scan using a 405-nm diode laser. We collected images before and after bleaching using low laser intensities and monitored FRAP for 15 min with data acquisition points every 20 s. All microscopy images were processed using the Leica LAS AF Lite software. In brief, the FRIP strips were blocked for 1 h in 3% bovine serum albumin in Tris-buffered saline plus Tween 20 supplemented with either 10 μM CaCl2 or 5 mM EGTA and then incubated overnight at 4°C in blocking buffer containing 0.5 μg mL⁻¹ SYT1244-541 C2AB peptide. The strips were incubated with the primary anti-SYT1 antibody (1:1,000) for 2 h and with the secondary horseradish peroxidase-conjugated goat anti-rabbit antibody (IgG-HRP, Santa Cruz Biotechnology, 1:14,000) for 1 h. The chemiluminescent signal was detected using the Clarity western enhanced chemiluminescence substrate (Bio-Rad).

Immunostaining

Five-day-old root tips were used and all working solutions were prepared in microtubule stabilization buffer (50 mM PIPES, 5 mM EGTA, and 5 mM MgSO4 [pH 7]). Sample preparation followed the basic protocol described in Sauer (2000). For immunodetection, a polyclonal rabbit anti-SYT1 antibody (1:1,000) or a monoclonal anti-Tyr-tubulin mouse IgG (TUB-1A2; London Resin Company Ltd), sectioned, and poststained as described (McFarlane et al., 2008). Immunolabeling was performed according to McFarlane et al. (2008) with 1:100 primary anti-SYT1 antibody and 1:100 goat-anti-rabbit 10-nm gold conjugate (BBI Solutions). As negative controls, wild-type sections were treated without primary antibody, and syt1 mutant sections were treated with anti-SYT1 and the gold-conjugated secondary antibody. Samples were imaged at 0.8 kV using a Zeiss EM910 coupled to an Olympus Quemsa CCD camera.

Confocal Imaging

Immunolocalization and cell viability images were obtained using a Leica TCS SP5 II or a Zeiss 710 confocal microscope equipped with 405-nm diode lasers for 4,6-diamino-phenylindole, a 488-nm argon laser for GFP, Alexa488 and fluo-rescin diacetate, and a 561-nm He-Na laser for FM4-64, Cy3, and TRITC signals. For localization and molecular integrity analyses, sequential line scanning mode was used to separate signals. The SYT1-GFP images were obtained using a Zeiss-Pascal Excite laser scanning confocal microscope equipped with a 488-nm argon laser for GFP and a 543-nm He-Ne laser for propidium iodide. For FRAP analysis, we adjusted the laser intensity and performed a single bleaching scan using a 405-nm diode laser. We collected images before and after bleaching using low laser intensities and monitored FRAP for 15 min with data acquisition points every 20 s. All microscopy images were processed using the Leica LAS AF Lite software. In brief, the FRIP strips were blocked for 1 h in 3% bovine serum albumin in Tris-buffered saline plus Tween 20 supplemented with either 10 μM CaCl2 or 5 mM EGTA and then incubated overnight at 4°C in blocking buffer containing 0.5 μg mL⁻¹ SYT1244-541 C2AB peptide. The strips were incubated with the primary anti-SYT1 antibody (1:1,000) for 2 h and with the secondary horseradish peroxidase-conjugated goat anti-rabbit antibody (IgG-HRP, Santa Cruz Bio-technology, 1:14,000) for 1 h. The chemiluminescent signal was detected using the Clarity western enhanced chemiluminescence substrate (Bio-Rad).

Mechanical Treatments

Fully expanded 6-week-old rosette leaves from plants grown under short-day photoperiod (8 h of light/16 h of dark) were mechanically pressed perpendicularly to the midrib using the first pressure level of a surgical hemostatic clamp. After 1-h treatment, leaves were harvested and incubated for 6 min in boiling trypan blue staining solution (trypan-blue stock solution [10 g of phenol, 10 mL of lactic acid, 10 mL of glycerol, 10 mg of trypan blue] diluted 1:1 with ethanol) and changed to chloral hydrate destaining solution (2.5 g mL⁻¹ in water). The destaining solution was changed several times during the next 3-d period, and the destained leaves were mounted on microscopy slides using 50% glycerol. Images were acquired using a Nikon Eclipse E 800 differential interference contrast microscope. For mechanical treatments at the cellular level, 5-d-old seedlings were mounted on microscopy slides using water, and variable weights ranging from 50 to 350 g were applied to the 22 × 22-mm coverslip for 20 s. Pressed cotyledons were observed under a confocal microscope immediately after treatment.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number At2g20990 for SYT1. The syt1-2 allele used in the study corresponds to the transfer DNA insertion line SAIL_775_A08 (Arabidopsis Biological Resource Center stock no. CS634629).
SUPPLEMENTAL DATA

The following supplemental materials are available.

SUPPLEMENTAL FIGURE S1. Functional domains of SYT orthologs and anti-
SYT1 antibody specificity.

SUPPLEMENTAL FIGURE S2. SYT1 colocalization with different PM markers.

SUPPLEMENTAL FIGURE S3. Comparison between SYT1 and different ER
markers.

SUPPLEMENTAL FIGURE S4. ImmunoTEM controls.

SUPPLEMENTAL FIGURE S5. SYTI-GFP time lapse and FRAP experiments.

SUPPLEMENTAL FIGURE S6. Chemical treatment controls.

SUPPLEMENTAL FIGURE S7. Subcellular localization of the 35SproGFP,
35SproC2AB, and SYT1-GFP markers in plasmolyzed cells.

SUPPLEMENTAL FIGURE S8. Colocalization of the 35SproGFP, 35SproC2AB,
and SYT1-GFP markers with FM4-64.

SUPPLEMENTAL FIGURE S9. Protoplast viability assay in wild-type and syt1
backgrounds.

SUPPLEMENTAL FIGURE S10. Severeing activity upon mechanical stress in
wild-type and syt1 backgrounds.

SUPPLEMENTAL MOVIE S1. SYTI-GFP marker movement through transva-
cular strands.

SUPPLEMENTAL MOVIE S2. Time-lapse experiment using the SYT1-GFP
marker.

SUPPLEMENTAL MOVIE S3. FRAP analysis using the SYT1-GFP marker.

SUPPLEMENTAL MOVIE S4. FRAP analysis of the SYT1-GFP cortical signal.

SUPPLEMENTAL MOVIE S5. SYTI-GFP signal upon treatment with LatB.

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