Plant VAP27 proteins: domain characterization, intracellular localization, and role in plant development.

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Summary (200 words)

• The endoplasmic reticulum (ER) is connected to the plasma membrane (PM) through the plant specific NETWORKED protein, NET3C, and phylogenetically conserved Vesicle-Associated Membrane Protein-Associated Proteins (VAPs).

• Ten VAP homologues (VAP27-1 to 10) can be identified in the Arabidopsis genome and can be divided into three clades. Representative members from each clade have been tagged with fluorescent protein and expressed in Nicotiana benthamiana.

• Proteins from clades one and three localised to the ER as well as to ER/PM contact sites (EPCS), whereas proteins from clade two are found only at the PM. Some of the VAP27 labelled EPCS localised to plasmodesmata, and we show that the mobility of VAP27 at the EPCS is influenced by the cell wall. EPCS closely associate with the cytoskeleton, but their structure is unaffected when the cytoskeleton is removed.

• VAP27 labelled EPCS are found in most cell types in Arabidopsis with the exception of cells in early trichome development. Arabidopsis expressing VAP27-GFP fusions exhibit pleiotropic phenotypes including defects in root hair morphogenesis. A similar effect is also observed in plants expressing VAP27 RNAi.

• Taken together these data indicate that VAP27 proteins used at the EPCS are essential for normal ER-cytoskeleton interaction and for plant development.

Key words: ER/PM contact sites, endoplasmic reticulum, VAP27, Scs2, NET super-family
Introduction

Proteins and other cargos synthesised in the ER are transported to various destinations through the conventional vesicular trafficking pathway. In higher plants, the cortical endoplasmic reticulum (ER) network is a highly dynamic structure and its movements are regulated by the actin cytoskeleton in most cell types (Boevink et al., 1998; Sparkes et al., 2009b). However, direct association between the ER and other membrane compartments also exists (Stefano et al., 2014; Hawes et al., 2014), which may provide alternative transport routes, a so-called non-vesicular pathway. These may include the ER-Golgi interface (daSilva et al., 2004; Hawes et al., 2008; Sparkes et al., 2009a), an ER-chloroplast (Mehrshahi et al., 2014) connection and ER/PM contact sites (Sparkes et al., 2009b, Manford et al., 2012; Wang et al., 2014).

Various ER/PM contact sites (EPCS) have been reported in different species, and are regulated by various proteins. For example, the STIM1/Orai1 complex is found in animals and is required for intercellular calcium transport (Varnai et al., 2007; Carrasco and Meyer 2011). In yeast, the Scs2/Osh/Sac complex is found at the ER and is used for lipid transfer to the PM (Stefan et al., 2011; Loewen et al., 2005). This complex also regulates ER morphology during budding (Loewen et al., 2007). A few other candidates such as Ist2 and Tlb (known as synaptotagmins in animals and plants) have also been identified as candidates for regulating the formation of the EPCS (Manford et al., 2012; Perez-Sancho et al., 2015). In plants, early electron microscopy studies described the structure of the EPCS (Hepler et al., 1990), and persistency mapping identified these structures in living cells (Sparkes et al., 2009b). However, their protein composition has not been fully elucidated and recent studies have begun to redress this situation (VAP27/NET3C complex, Wang et al., 2014).

The ER/PM contact site is linked to the cytoskeleton and this is mediated by a VAP27/NET3C complex through its interactions with microtubules and F-actin (Wang et al., 2014). A similar association between the cytoskeleton and EPCS has also been reported in migrating cancer cells, where the ER/PM junction is formed at the leading edge and is associated with actin markers (Dingsdale et al., 2013). Therefore, it is likely that the close association between EPCS and the cytoskeleton is important for cell polarity and development.
Vesicle-Associated Membrane Protein (VAMP) - Associated Proteins (VAP) are conserved amongst phylogenetically distinct organisms, and were first identified in the SNARE protein complex that is involved in vesicle docking and fusion (Skehel et al., 1995). Their plant homologues are named VAP27 because the first member identified had a molecular weight of 27 kDa (Laurent et al., 2000). As the name suggests, animal VAPs bind to a wide range of SNARE proteins that are required for vesicle trafficking from the ER (Weir et al., 1998; Weir et al., 2001; Soussan et al., 1999). Their functions in lipid transfer have been well studied in yeast, and similar functions are likely to exist in plants due to the identification of their interaction with oxystereol-binding proteins (ORPs) and sphingolipid transfer proteins (Saravanan et al., 2009; Petersen et al., 2009). In addition, recent studies have shown that VAPs are also required in the virus infection pathway, which is unique to plants (Barajas et al., 2014).

Plant EPCS can be defined as persistent ER nodes/punctae that are static whilst the ER remodels (Sparkes et al., 2009b), as well as sites where ER membrane attach to the PM as observed in ultrastructural studies (Hepler et al., 1990). In this study, we use either VAP27-1-YFP or GFP-NET3C as markers for EPCS in plants (Wang et al., 2014, Perez-Sancho et al., 2015; Levy et al., 2015). We identify ten VAP homologues in the Arabidopsis genome, all of which contain a highly conserved major sperm domain (Laurent et al., 2000). We have chosen members of each of the three phylogenetic clades to study their intracellular localization, functional domains and effects on plant development.

Materials and Methods

Bioinformatic analysis

Multiple alignments were assembled in ClustalX (Larkin et al. 2007) and exported as graphics using Jalview. Domains were identified with the Simple Modular Architecture Research Tool, SMART (Schultz et al. 1998), Interpro (Hunter et al., 2011) Coils (Lupas et al., 1991) and TMHMM (Sonnhammer et al., 1998). The Maximum likelihood method was chosen for the VAP family phylogenetic tree as this method has been identified as one of the most robust optimality criterion. Maximum Likelihood trees were calculated in the MetaPIGA software.
package (Helaers and Milinkovitch, 2010), using stochastic heuristics for large phylogeny inference with the Metapopulation Genetic Algorithm (metaGA) (Lemmon and Milinkovitch, 2002). MetaGA is an evolutionary computation heuristic in which several populations of trees exchange topological information which is used to guide the Genetic Algorithm (GA) operators for much faster convergence. MetaPIGA calculations were stopped when the mean relative error of 10 consecutive consensus trees stayed below 5% using trees sampled every 5 generations or the Likelihood stopped increasing after 200 iterations. Trees were drawn and exported as graphical files from FigTree (Andrew Rambout, University of Edinburgh). Transcription profiles of VAP isoforms were generated with Gene Investigator (Zimmermann et al. 2004, NEBION / ETH Zurich) from publicly available DNA microarray data.

**Molecular biology**

Primers and plasmids used in vector constructions are listed in Supplementary table 1. The VAP27 full-length cDNAs were amplified by RT-PCR (Invitrogen, UK) with gene specific primers (Table S1). Fluorescent protein fusions to VAPs were made using Gateway recombination (Invitrogen) into various destination vectors as shown in Table S2. Full length VAP27s, as well as the major sperm, coiled-coil and transmembrane domain deletion mutants of VAP27-1 and 3, were generated using PCR with appropriate primers. The VAP27-3 Arabidopsis RNAi line was obtained from AGRIKOLA (Hilson et al., 2004) and the RNAi insertion was confirmed using AGRIKOLA specific primers (Table S1). The VAP27-1 RNAi construct obtained from AGRIKOLA was sub-cloned into the pHELLSGATE RNAi vector (Wesley et al., 2001) and transformed into Col-0 Arabidopsis.

**Plant transformation and GUS study**

*Arabidopsis* (Col-0) was grown on compost in a growth chamber with a 16hr light (22 °C) and 8hr dark (18 °C) regime. *N. benthamiana* were maintained in a growth room with a 16hr light (25°C) and 8hr dark (18°C) regime. Transient expression was performed by leaf infiltration using *N. benthamiana* with Agrobacterium (Sparkes et al., 2006). Stable transformed *Arabidopsis* lines were generated using floral-dipping (Zhang et al., 2006). The VAP27-1 and 3 genomic sequences (including promoter and open reading frame) were fused
in frame (without the termination codon) to the 5’ end of the GUS reporter sequence. Stable GUS plants were obtained by selecting floral-dipped seeds on half MS medium containing kanamycin. GUS staining and histological studies were performed as described in Deeks et al., 2012.

**Antibodies and Immunofluorescence study**

VAP27-1 cDNA corresponding to amino-acid residues 164-230 was cloned into pET28a plasmid (Novagen) which incorporates an N-terminal His tag into the expressed protein (see supplementary 1 for primers used). Recombinant proteins were generated in E.coli (Rosseta 2, Novagen) and purified using nickel agarose beads (Qiagen). Polyclonal antibodies were raised in mice as described (Ketelaar et al., 2004). The specificity of the antiserum was tested on a one dimensional gel western blot of a total protein extract from 14 day old Arabidopsis seedlings. For detection, the membrane was incubated in TBST buffer with 5% milk prior to primary antibody incubation (1:500-1000) and HRP-conjugated secondary ant-mouse IgG (1:3000) and developed using the ECL reagent (GE Heathcare). PageRuler pre-stained protein ladder (Life technologies) was included on the western blot. Immunofluorescence with freeze shattering was performed as described (Zhang et al., 2013). Antibodies were diluted and used at 1:300 for VAP27 and 1:500 for BIP2 (Agrisera), followed by secondary antibody incubation with TRITC-conjugated anti-mouse IgG and FICT-conjugated anti-rabbit (Jackson ImmunoResearch).

**Confocal microscopy and live cell imaging**

All the microscopy images in this paper are representative of more than three independent infiltrations or stable transformations.

Samples were imaged using laser scanning confocal microscopes (LSCM, Leica SP5). Images were taken in multi-track mode with line switching when multi fluorescence was used. FRAP experiments and data analyses were performed as described (Wang et al., 2011), using a minimum number of 15 areas of interest which were bleached from different cells. During the photobleaching step, full output from the laser line was used and low laser intensities (1% 514nm for YFP) were used for data collection. The difference in maximum recovery was analyzed using the Student’s t-test to confirm the statistical significance. FRET-
FLIM analysis was performed as described (Wang et al., 2014), 12 repetitions were performed for each sample. Protoplasts were prepared using infiltrated leaves of *N. benthamiana*. Leaves were cut with a blade every 2mm and placed in a petri dish containing enzyme mix (macerozyme 0.2%, cellulose 0.4%) with K3 buffer (B5 basal medium 3.78g/l; CaCl$_2$ 750mg/l; NH$_4$NO$_3$ 250mg/l; sucrose 136.2/l; xylose 250mg/l; 6-benzylaminopurine 1mg/l; Naphtalenacetic acid 1mg/l). Digestion was carried out over-night at room temperature. The enzyme mix was removed the next day, and cells were suspended in K3 buffer for microscopy studies. Cell wall as stained with calcofluor as described (Martiniere et al., 2011). Plasma membrane staining was performed by immersing small leaf segments into water solution containing FM4-64 (7.5µm, Sigma) for 10min. Plasmodesmata were labeled using aniline blue as described (Deeks et al., 2012). Cytoskeleton depolymerization drug treatment in this study was performed by incubating small leaf segments (3x3mm) in a solution containing latrulin b (25µM for 30-45min), Oryzalin (20 µM for 30-45min) or Amiprophos-methyl (APM; 50µM for 60-90min).

**Transmission electron microscopy and immuno-gold labelling**

Plant tissue was fixed by high-pressure freezing and freeze-substitution as described (Deeks et al., 2012). The VAP27 anti-serum was used at 1:100 dilutions and detected by 5nm gold-conjugated anti-mouse IgG.

**Gene Accession numbers**

The Arabidopsis genome initiative locus tags for VAP27 genes are: At3g60600 (VAP27-1), At1g08820 (VAP27-2), At2g45140 (VAP27-3), At5g47180 (VAP27-4), At2g23820 (VAP27-5), At4g00170 (VAP27-6), At1g51270 (VAP27-7), At4g21450 (VAP27-8), At4g05060 (VAP27-9), At5g54110 (VAP27-10).

**Results and Discussion**

**Phylogenetic analysis of Arabidopsis VAP27 proteins**

Ten Arabidopsis VAP homologues (Fig. 1a) have been identified from a BLAST search using the Major Sperm Domain (MSD, Fig. 1b). Analysis based on full length protein sequences
show that VAP27 isoforms fall into three distinct clades. Although members within a clade can have different organizations of domains, all members of group two in particular lack a transmembrane domain and have the MSD located centrally rather than at the amino terminus. The majority of VAP27 isoforms show expression across a variety of tissues as shown by the genevestigator analysis (Fig. 1c). In addition to many other tissues, VAP27-1 and VAP27-4 show a peak of transcription in pollen. The tissues in which we find transcription of VAP27-5 & 7 are much more limited. VAP27-5 is confined to shoot, pollen and the stele. VAP27-7 is only found in the leaf although transcription is also seen in mesophyll and root primary cell culture.

**Intracellular localization of VAP27 proteins in *N. benthamiana* leaf epidermal cells**

Five of the ten VAP27 proteins have been used to make chimeric constructs with yellow fluorescent protein (YFP) at the C-terminus. These constructs were used for Argobacterium mediated transient transfection of *N. benthamiana* leaf epidermal cells in order to study their intracellular localization and behaviour. Our previous study showed that VAP27-1 is an ER integral membrane protein that also localised to EPCS (Wang et al., 2014; Fig. 2a). A similar localisation pattern was observed for VAP27-3 (also known as PVA12, Saravanan et al., 2009) and VAP27-4, representative members of clade 1 and clade 3 respectively. Both proteins localise to the ER network, confirmed by their co-localisation with CFP-HDEL, in addition to immobile punctate structures that we previously identified as EPCS (Fig. 2b-c; Movie S1). At the cell periphery, VAP27-1 puncta appeared to co-localise with the PM (stained with FM4-64, Fig. 2a, inset). Signals from the rest of the ER are very distinct from the PM (FM4-64 labelled, red). In contrast, two members from Clade 2 of the VAP27 family namely, YFP fusions of VAP27-8 and VAP27-10 (also known as AtMAMI, Galaud et al., 1997), localise to the plasma membrane (Fig. 2d-f; Movie S2). They are very likely to be membrane peripheral proteins (as no transmembrane domain has been identified) recruited to the PM from a cytoplasmic pool. VAP27-8-YFP also labelled some immobile puncta (Fig. 2d) and is also found concentrated in the nucleolus (Fig. 2d, inset).

NET3C belongs to a plant specific family of actin binding proteins, the NET Family (Deeks et al., 2012); it locates to the EPCS (Wang et al 2014). We have previously shown that VAP27-1 co-localises and interacts with NET3C at these sites (Wang et al., 2014). In this study, we
show that VAP27-3-YFP also co-localises with GFP-NET3C at EPCS when co-expressed in the transient expression system (Fig. 3a). We have confirmed the physical interaction between RFP-VAP27-3 and GFP-NET3C in vivo using FRET-FLIM microscopy (Fig. 3b-d). The fluorescent life time (LT) of GFP-NET3C (donor complex) was found to be 2.61 ± 0.05ns, which reduces significantly in the presence of RFP-VAP27-3 (LT=2.41 ± 0.05ns; p=7.63E-8), indicating that they interact in a complex. It should be noted that the life time of GFP-NET3C in the nucleus does not change as no VAP27-3 is present and this also acts as an internal control for this FRET-FLIM study (Fig. 3d). A second negative control was carried out using GFP-HDEL and RFP-HDEL; the life time of GFP in cells expressing both constructs was measured at 2.66 ± 0.03ns and this indicates that these two proteins which do localise to the same compartment do not interact and therefore do not undergo FRET (Fig. 3c).

**Localisation of VAP27-1 in Arabidopsis**

In order to assess the level of translational expression of VAP27-1 and VAP27-3, each gene was ligated in frame with GUS at the 3’ end of each open reading frame. Expression in Arabidopsis revealed that both proteins are expressed ubiquitously (Fig. 4a-b), similar to their predicted transcriptional expression profiles (Fig. 1 c). Arabidopsis leaf epidermal cells were then used for further immuno-labelling studies.

A polyclonal antibody raised against VAP27-1 in mice detects a single band on a western blot of a total protein extract from Arabidopsis seedlings at a molecular weight similar to that for VAP27-1 (Fig. 4c). This antibody is specific to VAP27-1 when compared to its cross reactivity to VAP27-3 proteins, which have high overall sequence similarity (Fig. 4d). Cotyledons from a stable Arabidopsis line expressing VAP27-1-YFP were high pressure frozen and freeze-substituted for TEM and immuno-gold studies. The area of association (marked in red) between the ER and PM appears much enhanced by the expression of VAP27-1. Gaps between the ER and PM are almost undetectable (Fig. 4e-f). Gold labelled VAP27-1 is found throughout the ER network as well as at the EPCS (arrow, Fig. 4g).

Immunofluorescence studies using Arabidopsis leaf epidermal cells identifies the endogenous VAP27-1 on the ER network (which is stained by a BIP2 antibody) as well as some ER associated puncta that are distinct from the BIP2 labelled ER (Fig. 4h). The EPCS labelling of endogenous VAP27-1 in Arabidopsis is not as pronounced as the VAP27-1-YFP in
the tobacco transient expression system and this is likely to be because of the amount of protein that is present with more VAP-27-1-YFP being available in the transiently expressing cells. VAP27-1 signal is found at the same position as the BIP2 signal on the ER membrane. In contrast, VAP27-1 and BIP2 are only partially co-localized at the putative EPCS (Fig. S1). This makes sense because VAP27s (like yeast Scs2) are actively recruited to the EPCS, while other ER localised proteins are not.

Two strips of the same western blot of a 1D gel loaded with *N. benthamiana* leaf extract expressing VAP27-1-YFP were probed with 1. VAP27-1 antibody; 2. VAP27-1 antibody co-incubated with VAP-27 peptide immunogen. Incubating the VAP27-1 peptide immunogen with the VAP27-1 antibody abolished the ability of the antibody to detect a band on the western blot equivalent to VAP27-1-YFP indicating the antibody's specificity for VAP27-1 (Fig. 4j). Co-incubating the VAP27-1 peptide immunogen with the VAP27-1 antibody and using this mixture to stain Arabidopsis cells revealed no staining of the ER and EPCS in planta (Fig. 4i) further supporting the specificity of the VAP27-1 antibody used in this study. In summary, results from the immunocytochemistry are consistent with the live cell imaging data, confirming VAP27-1 as an ER network and EPCS localised protein in plants. We suggest that VAP proteins from clades 1 and 3, specifically VAP27-3 and VAP27-4, have a similar cellular location in Arabidopsis as their sequences are very similar to VAP27-1 (VAP27-3, 84%; VAP27-4, 57%) and their localisation in *N. benthamiana* is the same (Fig. 2a-c).

Stably transformed Arabidopsis expressing VAP27-1-GFP driven by its native promoter exhibit a similar subcellular localisation to that observed when using the same construct in the *N. benthamiana* transient expression system (Fig. 5a-c). Numerous ER-associated puncta are identified, reminiscent of the EPCS seen in leaf epidermal studies. However, these puncta are not seen in all cells. For example, in trichome development, EPCS labelling was only seen in mature trichomes (stage 6) and not found in the earlier developmental stages (stages 1-4) (Fig. 5 d-e). This is either because EPCS may not exist in this type of cell, or other proteins may be involved in EPCS formation. VAP27-1 labelled EPCS are found in close association with both the microtubule and actin cytoskeletons (Fig. 5g-i) in trichomes of the Arabidopsis stably transformed lines and in transiently expressing *N. benthamiana* leaf epidermal cells (Fig. S2a-f). In Arabidopsis trichomes for example, the percentage of EPCS that are associated with the actin cytoskeleton or with microtubules was found to be 81.2 ±
4.3% and 70.4 ± 14.3% respectively. A random association assessed by rotating the red channel (e.g. RFP-Lifeact) by 180° with respect to the green channel (VAP27-1-GFP) gave a percentage association of 40.1 ± 7.75%, which is significantly lower than the percentage association between EPCS and actin or microtubules indicating that their association with the cytoskeleton is a valid result.

VAP27-1 labelled EPCS are often located at the cross overs between F-actin and microtubules (Fig. S2d-f). This observation supports the observations that part of the ER sub-domain interacts with microtubules, forming so called C-MERs (cortical microtubule associated ER sites; Pena and Heinlein, 2013). However, these contact sites do not appear to be maintained by the cytoskeleton, as they still exist when either F-actin or microtubules are removed by drug treatments (Fig. S2g-i).

**The dynamics of VAP27 at the ER/PM contact site (EPCS) is influenced by the cell wall.**

A population of the VAP27-1-YFP labelled EPCS also associated with plasmodesmata as revealed by co-localisation of aniline blue staining of callose (Fig. 6a), suggesting that PDs may perform a similar function to the ER/PM contact site at the cell-cell border in terms of anchoring the peripheral ER. In this context, it is known that the desmotubule of plasmodesmata is comprised of highly constricted ER membrane (Wright *et al.*, 2007; Fitzgibbon *et al.*, 2010; Knox *et al.*, 2015) and as such a role of VAPs in anchoring the peripheral ER to the plasmodesmal channel is an attractive hypothesis.

Leaf epidermal cells expressing VAP27-1 were treated with mannitol to induce plasmolysis, designed to separate the plasma membrane from the cell wall (the PM in Fig. 6b-c is labelled with a fluorescence marker, PIP2-CFP). As a consequence, hechtian strands are found in the plasmolyzed cells that link the cell wall and plasma membrane (Lang-Pauluzzi 2000; Fig. 6b-c). Surprisingly, most of the VAP27-1 labelled EPCS were found within or at the tips of hechtian strands (Fig. 6c-d). Thus, we suggest that the plant ER/PM complex must interact indirectly with the cell wall through some PM localised proteins, which holds them together during plasmolysis while the rest of the ER network is separated from the cell periphery (Fig. 6d).
Subsequently, protoplasts were isolated from VAP27-1-YFP transformed leaves to study the influence of the cell wall on the EPCS (Movies S3). After photobleaching, the recovery of VAP27-1 at the ER/PM contact site is calculated as 74.45 ± 15.9% (Fig. 6f-g). It is known that cell wall can be re-generated around protoplasts (Martiniere et al., 2011). No cell wall staining is seen at 0 hours when the protoplasts were freshly prepared, whereas staining was seen at ca. 24 hours after isolation (Fig. 6e). When the cell wall reformed, the maximum recovery of VAP27-1 reduced to 57.48 ± 10.4%, significantly different from its recovery at 0 hours (p < 0.001). These differences in the percentage recoveries indicate that the immobile fraction of VAP27-1 within the photo-bleached region is greater when the cell wall has re-generated. This also indicates that the association between VAP27-1 labelled EPCS and the cell wall makes VAP27-1 largely immobile. However, the half time of recovery at both time points (0 and 24 hrs) does not change significantly (p>0.2) which indicates that the dynamics of VAP27-1 in the photobleached regions at 0 hours and ca. 24 hours are similar.

In conclusion, these results indicate that the cell wall affects the percentage recovery of the EPCS associated protein, VAP27-1. We suggest that this phenomenon is due to VAP27-1 interacting with a protein that both spans the plasma membrane and interacts with the cell wall and the EPCS at either terminus, or that VAP27-1 associates with a PM sub-domain whose mobility is constricted by the cell wall. Recently, the physical association of the cell wall with the plasma membrane has been implicated in the anchoring of many different proteins in the plasma membrane (Martiniere et al., 2012). Both scenarios could affect protein dynamics at the plasma membrane and subsequently the EPCS (Fig. 6h).

Expression of VAP27-1 and NET3C induces PM associated ER cisternae

High level expression of constructs in the *N. benthamiana* transient system can be obtained by increasing the optical density of the Agrobacteria used for infiltration (Batoko et al., 2000). When highly expressed, VAP27-1 interacts with NET3C and induces the formation of membrane cisternae, which are labelled by CFP-HDEL, suggesting that these cisternae are ER derived (Fig. 7a). EPCS labelling may still be seen in some parts of the cell (Fig. 7a, arrow), but the significant deformation of ER membrane makes these sites difficult to resolve. These ER derived membrane cisternae are closely attached to the PM, as seen by the close
association with FM4-64 fluorescence at the cell cortex (Fig. 7b-c; also compare with Fig. 1a, inset). Some filament-like structures in negative contrast can be observed within the membrane cisternae (Fig. 7d, arrow), and these co-localise with microtubules (labelled with the Kinesin Motor Domain fused to RFP; KMD-RFP). This microtubule related pattern in membranes has been previously described in several studies of plasma membrane integral proteins. This is likely due to the corralling of PM proteins by cortical microtubules (Martiniere and Runions, 2013). Not surprisingly, these negative images of microtubules disappear when microtubules are depolymerized by amiprophos-methyl (APM) treatment (Fig. 7e).

The enhanced association between the ER and PM membrane is only seen when both VAP27-1 and NET3C are present, and expression of VAP27-1 alone does not induce this phenomenon (Fig. 2a). This result suggests that only a small fraction of ER membrane can interact with PM associated NET3C to form the EPCS under native conditions where both VAP27 and NET3C expression are limiting. However, an excess level of both proteins appears to ‘glue’ the entire cortical ER system to the PM and induce this abnormal cellular phenotype (Fig. 7f).

**Domain characterization of VAP27 using deletion mutants and live cell imaging**

Three distinct functional domains are found in VAP27-1, namely, an N-terminal major sperm domain (MSD), a C-terminal transmembrane domain (TMD) and a coiled-coil domain (CCD). Domain deletion mutants of VAP27-1 were made and fused to YFP (Fig. 8a). VAP27-1ΔTMD was found to be cytosolic (Fig. 8e); VAP27-1ΔCCD-YFP is still ER localised but less puncta are observed than for full length VAP27 (Fig. 8c). The number of puncta was found to be 7.2 ± 3.1 per 30x30µm for VAP27-1ΔCCD compared to 31.4 ± 5.8 per 30x30µm for full length VAP27-1. VAP27-1ΔMSD-YFP forms numerous ER derived puncta, most of which are much more mobile than the full-length VAP27-1 puncta at ER/PM contact sites (Fig. 8d, Movie S4). Similar results were also obtained from a deletion study of another clade 1 VAP27, VAP27-3 (Fig. S3a-c). Little alteration of the ER network is seen in either the VAP27 full length (Fig. 2) or deletion mutant expressing cells (Fig. S3d-f). Therefore, the anchoring of the ER to the PM in plants may be complex and is likely to involve multiple proteins which includes the
association with NET3C (Wang et al., 2014), and possibly synaptotagmins (Perez-Sancho et al., 2015).

Protein dynamics within these puncta were studied using fluorescence recovery after photobleaching (FRAP; Fig. 8b). The maximum recovery of VAP27-1ΔCCD was found to be reduced (Rmax = 38.86 ± 18.5%) compared to full-length VAP27-1 (Rmax = 54.22 ± 18.2%; p < 0.05). VAP27-1ΔMSD showed almost no recovery over the same period (Fig. 8b). These results indicate that both MSD and CCD are important for the localization of VAP27 at the ER/PM contact site and most likely protein dynamics within the membrane, whereas, the TM domain determines the ER localisation of VAP27-1. Previously, we identified a functional motif on the major sperm domain required for the interaction between VAP27-1 and NET3C (Wang et al., 2014). This result is consistent with an observation from the co-expression of NET3C with VAP27-1 deletion mutants in this study. That is, co-localisation of GFP-NET3C is only seen with VAP27-1ΔTMD which contains the intact major sperm domain (Fig. 8g), whereas, no co-localisation is seen between NET3C and VAP27-1ΔMSD (Fig. 8f).

**Aberrant VAP27 expression effects plant development**

We stably transformed Arabidopsis producing lines expressing VAP27-1-GFP or VAP27-3-GFP driven by their native promoters. As we have shown that the GFP constructs localise to the EPCS in a pattern also observed using anti-VAP27-1 in planta, and that from yeast studies chimeric Scs2-reporter proteins (homologue of VAP27) are functional (Loewen et al., 2007), then these constructs are likely to be functional in plants. These plants exhibit defects in pollen, seed and root development. The most notable defect is in root hair development where hairs appear branched (compare Fig. 9a with 9b, arrow). High magnification images of these abnormal root hairs are shown in Fig. 9i. This phenotype is also observed in VAP27-1 RNAi lines (Fig. 9c, arrow), which show a significant knock-down of endogenous VAP27-1 protein expression (Fig. 9f-g). A root hair phenotype is also seen in plants expressing VAP27-3-GFP as well as in VAP27-3 RNAi lines (Fig. 9d-e). The percentage of branched root hairs was calculated for each line and there were ca. 40% abnormal root hairs in the VAP27-GFP expressing lines, and ca. 15-20% in VAP27 RNAi lines (Fig. 9h). These observations are consistent in more than three independent transformed lines. It is interesting that a similar phenomenon is seen in either VAP27 gain of function and loss of
function studies. These data suggest that tight control of VAP27 expression is essential. A similar phenomenon is observed when the actin regulating protein, Actin Depolymerising Factor, is both over-expressed and knocked down (Dong et al., 2001).

In the VAP27-1 expressing lines, both the ER and actin networks in the branched root hair are significantly different from those in the wild type (Fig. 9j-k). VAP27-1 labelled ER membrane aggregates formed at the point where the root hairs branch and the actin network appears disorganised in this zone (Fig. 9k). Root hair phenotypes have been observed previously when certain ER or F-actin regulating proteins are disrupted (Deeks et al., 2007; Guimil and Dunand, 2007).

In conclusion, the VAP27 protein family has been identified and representative candidates selected for further functional characterization. Proteins from clade 1 and 3 localised to the ER network as well as the ER/PM contact sites, whereas members of clade 2 are found at the PM. The function of the different domains have been characterised using VAP27-1 and 3 as examples. We have demonstrated that the major sperm domain and coiled coil domain are required for protein-protein interaction and that the transmembrane domain is required for intracellular localization. Pleiotropic defects were seen in plants expressing VAP27-GFP and also in VAP27 RNAi lines, suggesting that they are essential for normal plant development. Our results also suggest an indirect association between ER/PM contact sites and the cell wall, likely to be mediated through interaction with PM associated proteins.

Acknowledgments This work was supported by a BBSRC grant (BB/G006334/1) to P.J.H and a Leverhulme Trust grant (F/00382/G) to CH.

Author Contribution: P.W. and P.J.H. planned and designed the research. P.W. performed the research, and together with P.J.H and C.H wrote the manuscript. C.R., T.J.H. and I.S. contribute to data analysis.
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(a) Root hairs found within the differentiation zone of wild type Arabidopsis. (b) The root hairs from VAP27-1 expressing Arabidopsis lines exhibit an abnormal phenotype. They are much shorter and swollen compared to the wild type, and most are branched. (c) Brached root hairs were also seen in VAP27-1 RNAi lines (arrow), suggesting that either over- or under-expression of VAP27 affects root hair development. (d-e) The branched root hair phenotype is also observed in VAP27-3 expressing and VAP27-3 RNAi plants. (f) Western blot of VAP27-1 RNAi Arabidopsis (1-3) and wild type, the knock-down of VAP27-1 protein was confirmed in these RNAi lines. (g) Amido black staining suggested equal amount of proteins were loaded in all lanes (lower panel). (h) Statistical analysis of branched root hairs in VAP27-1 and 3 over-expression or knock-down lines. The percentage for each line is shown in the table. (i) Branched root hairs at high magnification, two root hairs were often seen bulged from one trichoblast cell. (j) The actin cytoskeleton (labelled by GFP-Lifact) in a wild type root hair cell, with fine filaments in the apical part and thick bundles at the base (3D maximum projection). (k) In the VAP27-1 expressing root hair cells, the ER and F-actin form aggregates, which affect its polarised growth. Instead of growing directionally, the root hair cell branched at the point where the membrane aggregates assemble (3D maximum projection; scale bar = 10µm).
Supplementary figure legends

**Figure S1.** Fluorescence signal distribution of VAP27-1-YFP on ER membrane.

**Figure S2.** The ER/PM contact sites in relation to the cytoskeleton.

**Figure S3.** VAP27 deletion mutants exhibit little effect on ER morphology

**Table S1.** List of primers used in this study.

**Table S2.** List of plasmids used in this study.

Supplementary movies

**Movies S1,** Z-stack images of leaf cells transiently expressing VAP27-4-YFP, which localises to the ER network as well as the EPCS.

**Movie S2,** Z-stack images of leaf cells transiently expressing VAP27-8-YFP, which localises to the PM.

**Movie S3,** Protoplast expressing VAP27-1-YFP.

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