

Arabidopsis BTB/POZ protein-dependent PENETRATION3 trafficking and disease susceptibility

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The outermost cell layer of plant roots (epidermis) constantly encounters environmental challenges. The epidermal outer plasma membrane domain harbours the PENETRATION3 (PEN3)/ABCG36/PDR8 ATP-binding cassette transporter that confers non-host resistance to several pathogens. Here, we show that the Arabidopsis ENDOPLASMIC RETICULUM-ARRESTED PEN3 (EAP3) BTB/POZ-domain protein specifically mediates PEN3 exit from the endoplasmic reticulum and confers resistance to a root-penetrating fungus, providing prime evidence for BTB/POZ-domain protein-dependent membrane trafficking underlying disease resistance.

The PENETRATION3 (PEN3/ABCG36/PDR8) ATP-binding cassette transporter of *Arabidopsis thaliana* is a crucial component of preinvasive defence against some fungal and bacterial non-host pathogens entering by direct penetration^{1–4}. In above-ground organs, PEN3 is recruited to sites of pathogen attack at the cell surface^{3,4}. In seedling roots, PEN3 polarly localizes to the epidermal outer membrane domain in the absence of pathogens^{5,6}. Root epidermal cells display four major polar plasma membrane domains: the outer domain facing the environment, the inner domain oriented towards the cortical cell layer, the shootward-oriented, apical, and the root tip-oriented, basal, domain⁶. Proteins in the outer domain that function in regulating the transport of inorganic compounds include, for example, the NIP5;1 boric acid uptake channel⁷. Factors required for PEN3 and NIP5;1 trafficking from the trans-Golgi network to the outer domain have been identified^{8–10}, and exocyst complex components promote polar tethering of several outer domain proteins^{9,11}. However, factors that specifically mediate trafficking of polar outer membrane cargos involved in responses to root-penetrating pathogens remain to be discovered.

In a genetic screen for mislocalization of PEN3 fused to green-fluorescent protein (PEN3-GFP) in the root epidermis of seedlings⁹, we recovered one recessive mutant in which PEN3-GFP localized to a cytoplasmic structure resembling the endoplasmic reticulum (ER) (Fig. 1a–d). This *er-arrested pen3-1* (*eap3-1*) mutation indistinguishably affected localization of PEN3-GFP from that of PEN3-mCherry (Supplementary Fig. 1a,b), which colocalized with the ER-intrinsic chaperone BIP in the *eap3-1* mutant (Supplementary Fig. 1c,d), corroborating an ER arrest of PEN3.

We mapped the *eap3-1* mutation to a 275-kb interval on chromosome 3 by recombination mapping (Fig. 1e) combined

with mapping and mutation identification by deep sequencing. This interval contained one non-synonymous, single-nucleotide polymorphism that caused a G to A missense mutation resulting in the conversion of glycine 192 to glutamate (*eap3*^{G192E}) in the predicted open reading frame of gene model At3g09030.1 (Fig. 1f). Wild type and homozygous *eap3-1* mutants displayed similar *EAP3* transcript levels (Fig. 1g), whereas the SALK_101331 T-DNA insertion located in the single exon of At3g09030 (designated *eap3-2*) caused a lack of full-length *EAP3* messenger RNA (Fig. 1g). In contrast to wild type (Fig. 1h), PEN3-GFP colocalized with the ER marker ER-ECFP-HDEL in *eap3-2* seedlings (Fig. 1i), in which PEN3-mCherry colabelled with BIP immunofluorescence (Supplementary Fig. 1e,f). *trans*-heterozygous *eap3-1/eap3-2* seedlings also displayed ER-mislocalization of PEN3-GFP (Supplementary Fig. 1g,h); this non-complementation strongly supported that the *eap3* mutations caused the PEN3 ER-mislocalization. The *EAP3* gene model predicts a 460-amino-acid protein with a molecular weight of 49.3 kDa carrying two potential protein–protein interaction domains, an N-terminal BTB/POZ domain and a central six-bladed beta-propeller domain.

To localize the *EAP3* protein, we tagged the genomic *EAP3* sequence with the coding sequence for a C-terminal myc tag (*pEAP3:EAP3-myc*) or a GFP tag (*pEAP3:EAP3-GFP*). Expressing *EAP3-myc* in *eap3-2* mutants fully rescued their PEN3-mCherry ER-mislocalization phenotype (Fig. 1j–l), whereas *EAP3-GFP* provided partial rescue in *eap3-2* mutants (Fig. 1m), suggesting full functionality of *EAP3-myc* and partial functionality of *EAP3-GFP* (Fig. 1n).

EAP3-GFP was expressed at very low levels, displayed cytosolic localization *in vivo* (Fig. 1m, Supplementary Fig. 1i,j) and was detectable just above background levels by western blotting (Supplementary Fig. 2a,b), consistent with very low At3g09030 mRNA levels reported in publicly available expression datasets (Supplementary Fig. 3). Functional *EAP3-myc* could neither be detected above background by western blotting nor by immunofluorescence microscopy (Supplementary Fig. 2a,b). However, immunoprecipitation revealed full-length *EAP3-myc* and *EAP3-GFP* proteins (Supplementary Fig. 2a,b), suggesting expression at low levels with full and partial functionality, respectively. Therefore, visualization of full-length *EAP3-GFP in vivo* supported a cytosolic localization of *EAP3*.

Strikingly, PEN3-mCherry accumulated in the *eap3-2* mutant, when examined by western blotting (Supplementary Fig. 2c–f),

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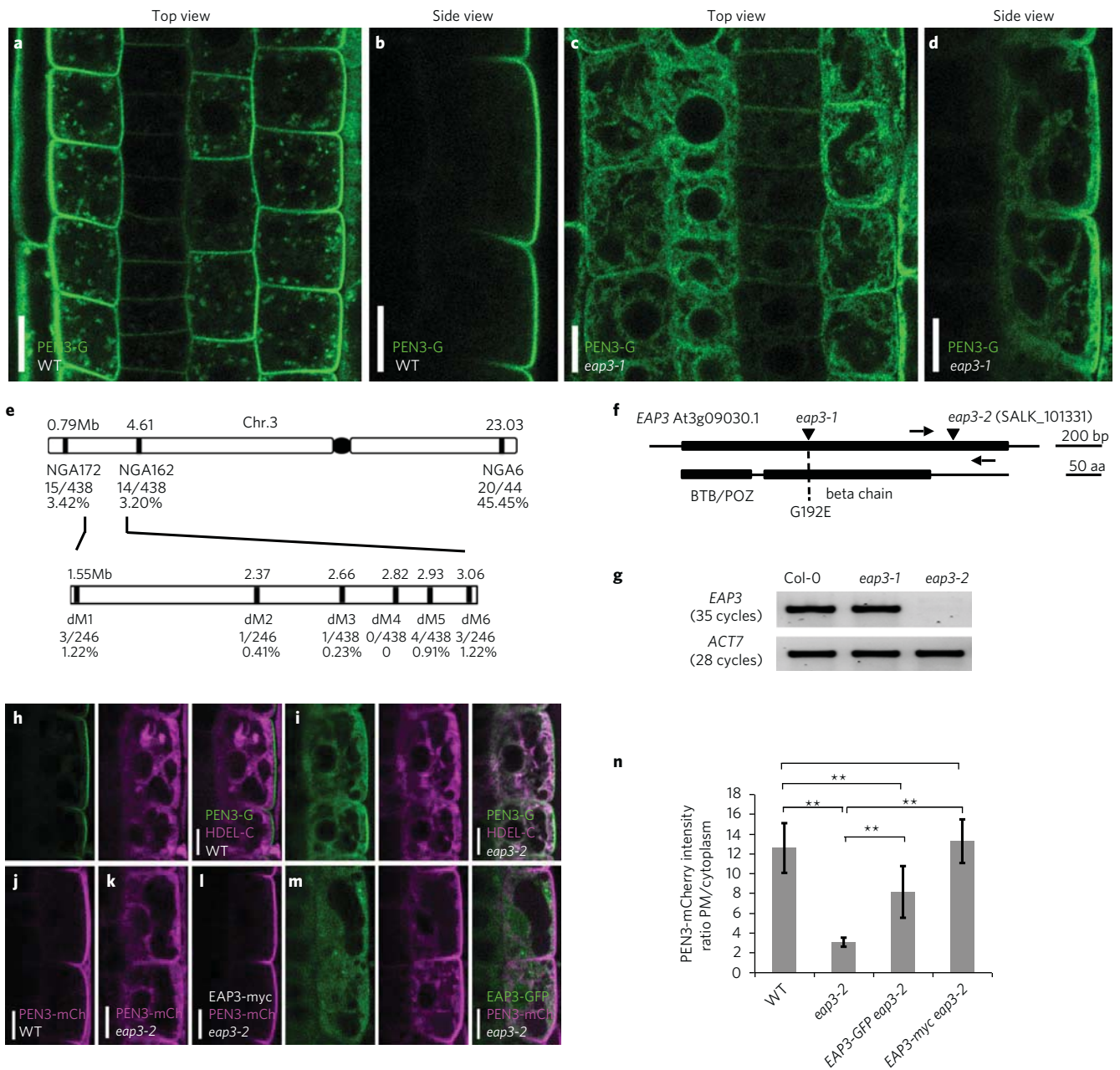


Fig. 1 | *eap3* mutations affect a cytosolic BTB/POZ-domain protein. a-d, PEN3-GFP (PEN3-G) in Col-0 wild type (**a,b**) and *eap3-1* (**c,d**). Root epidermal layer in top view on a tangential optical section (**a,c**) and from side view on a median, longitudinal, transversal optical section (**b,d**). **e**, Map-based identification of *eap3-1*. Simple sequence length polymorphism (SSLP; NGA) and derived cleaved amplified polymorphic sequences (dCAPS; dM) markers indicated with respective positions, recombinant/total chromosomes and recombination frequency (%). **f**, EAP3 At3g09030.1 genomic region (black line, top), exon (black bar). *eap3-1* point mutation and *eap3-2* T-DNA insertions indicated as well as amino acid sequence (black line, bottom) with BTB/POZ domain and six-bladed beta-propeller domain (beta chain). **g**, Semiquantitative PCR with reverse transcription analysis of Col-0, *eap3-1* and *eap3-2* 5-week-old leaves, using primers flanking the T-DNA insertion (arrows in **f**) and for *ACTIN7* (*ACT7*). PCR with reverse transcription cycle numbers indicated. Three independent experiments gave similar results. **h,i**, Colocalization analysis between PEN3-G (green) and ER-ECFP-HDEL (HDEL-C; magenta) in wild type (**h**) and *eap3-2* (**i**). **j-n**, Complementation analyses. **j,k**, PEN3-mCh at outer domain in Col-0 (**j**) and in ER-like structure in *eap3-2* (**k**). **l**, EAP3-myc restores PEN3-mCh localization in *eap3-2*. **m**, EAP3-GFP partially restores PEN3-mCh localization in *eap3-2*. **n**, Quantification of PEN3-mCherry outer PM domain/cytoplasm signal intensity ratio in respective backgrounds. ****P** <10⁻⁷ determined by Student's two-tailed *t*-test with equal variance, *n* = 30 cells. Scale bars, 10 μ m. Chr., chromosome; PM, plasma membrane; WT, wild-type Col-0.

whereas PEN3-mCherry displayed wild-type levels in lines expressing EAP3-myc, and an intermediate level between wild type and the mutant in the line expressing EAP3-GFP (Supplementary Fig. 2c-f). These results further supported PEN3 protein accumulation in *eap3* mutants.

We addressed whether PEN3 accumulation in the ER reflected the arrest of newly biosynthesized PEN3 protein by whole-cell fluorescence recovery after photobleaching (FRAP). In wild type, PEN3-mCherry displayed FRAP at the outer membrane domain after 90–180 min (Fig. 2a). Plasma membrane levels of PEN3-mCherry

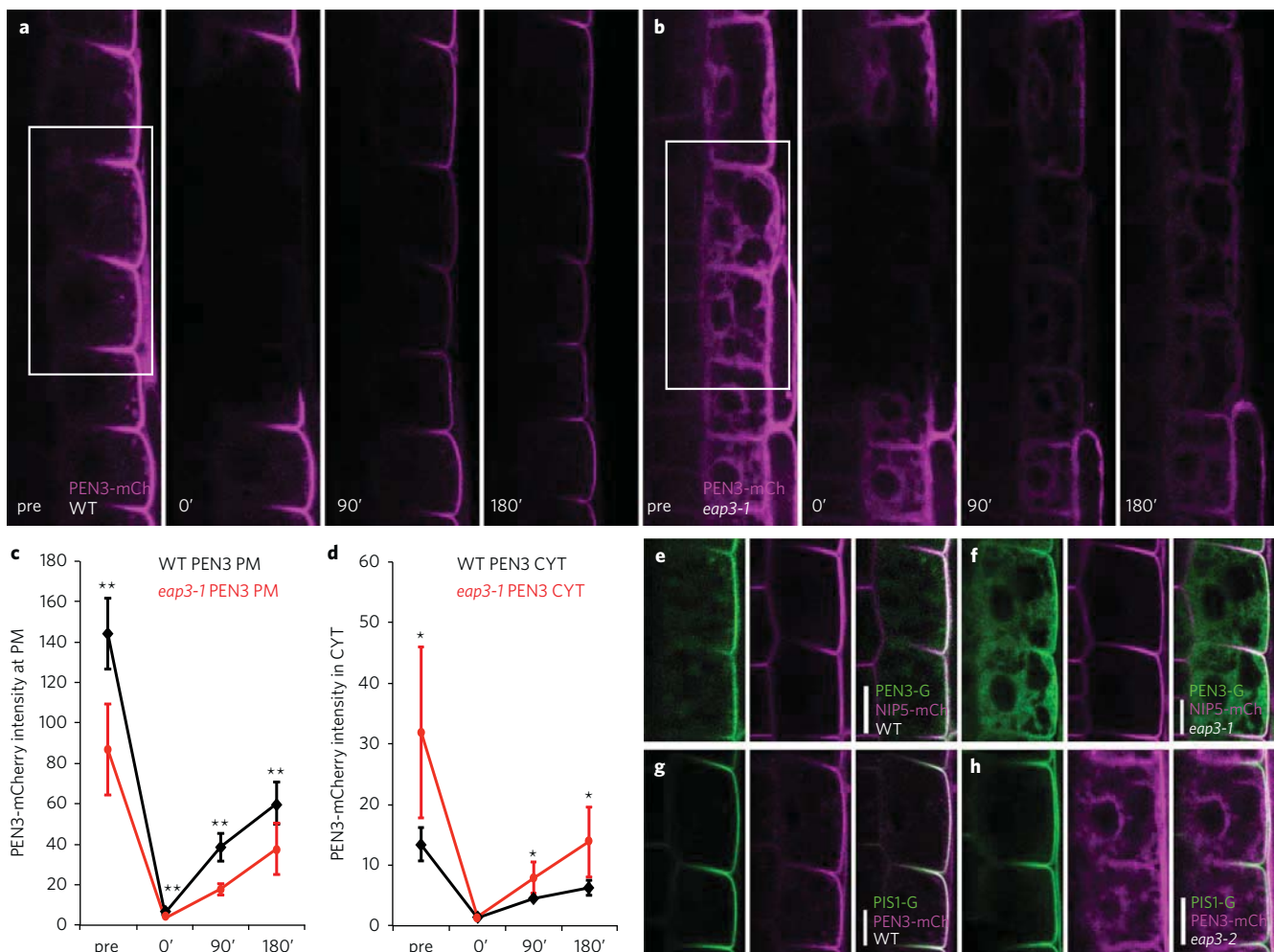


Fig. 2 | EAP3 mediates PEN3 secretory trafficking from the ER with high cargo specificity. **a,b**, Representative images of PEN3-mCherry (PEN3-mCh) whole-cell FRAP analyses with PEN3-mCh bleached at 561nm in wild type also expressing PEN3-GFP (WT) (**a**) and in *eap3-1* also expressing PEN3-GFP (**b**). Prebleach images (pre) with rectangular bleach ROIs are indicated. PEN3-GFP is not displayed in **a,b,c,d**. Quantitative analysis of FRAP experiments. Intensities of PEN3-mCh fluorescence at the outer plasma membrane domain (PM; **c**) region and in the cytoplasmic (CYT; **d**) region measured and compared between WT and *eap3-1*. Significances of differences in PEN3-mCh fluorescence intensities at different time points were determined employing Student's two-tailed *t*-test with unequal variance from $n=12$ WT cells and $n=10$ *eap3-1* cells. Results are displayed as averages with bars representing 95% confidence intervals. Significance level was $P < 0.05$. ** $P < 0.01$ and * $P < 0.05$. **e,f**, PEN3-GFP (PEN3-G; green) and NIP5;1-mCherry (NIP5-mCh; magenta) in wild type (**e**) and *eap3-1* (**f**). **g,h**, PIS1-GFP (PIS1-G; green) and PEN3-mCh (magenta) in wild type (**g**) and *eap3-2* (**h**). Scale bars, 10 μ m. ROI, region of interest.

were low in the *eap3-1* mutant prior to bleaching, and recovery remained low when compared to wild type (Fig. 2b,c). However, the *eap3-1* mutant showed stronger ER-like fluorescence of PEN3-mCherry, which displayed stronger FRAP than in wild type (Fig. 2b,d). In the wild type, PEN3-mCherry fluorescence recovered at the plasma membrane, colocalizing with non-bleached PEN3-GFP (Fig. 2a, Supplementary Fig. 4a), whereas localization of PEN3-mCherry during FRAP was indistinguishable from non-bleached PEN3-GFP in the ER of *eap3-1* (Fig. 2b, Supplementary Fig. 4b). Hence, EAP3 mediates trafficking of newly synthesized PEN3 through the early secretory pathway.

We next examined whether *eap3* mutations affected localization of other membrane proteins. Both the plasma membrane marker EGFP-LTI6a (Supplementary Fig. 4c,d) and the apical domain marker PIN2-EGFP (Supplementary Fig. 4e,f) were correctly localized in the *eap3-1* mutant (Supplementary Fig. 4d,f). Similarly, NIP5;1-mCherry that co-localized with PEN3-GFP at the outer domain in wild type (Fig. 2e) maintained its wild-type

outer-domain localization in *eap3-1* mutant cells with mislocalized PEN3-GFP (Fig. 2f). Strikingly, even the closest PEN3 homologue ABCG37/PIS1/PDR9 fused to GFP (PIS1-GFP), which colocalized with PEN3-mCherry at the outer domain in wild type (Fig. 2g), remained localized to this domain in *eap3-2* mutant cells (Fig. 2h). Thus, *eap3* mutations selectively disrupt PEN3 trafficking, when compared to other membrane proteins.

Consistent with EAP3 having a high cargo specificity for PEN3, we neither observed obvious developmental defects in *eap3* seedlings nor adult plants (Supplementary Fig. 5a-d). To address whether EAP3 and PEN3 act in a common pathway for pathogen resistance, we challenged *pen3* and *eap3* mutants by soil-inoculation with the root-penetrating pathogenic fungus *Fusarium oxysporum* *f. sp. conglutinans* (*F. oxysporum*)¹². Compared to non-treated controls (Fig. 3a-f) and to *F. oxysporum*-treated wild type (Fig. 3g), the *F. oxysporum*-treated *pen3-4* (Fig. 3h), *eap3-2* (Fig. 3i) and *eap3-2 pPEN3: PEN3-mCherry* mutants (Fig. 3j) displayed increased disease susceptibility indicated by many chlorotic and necrotic

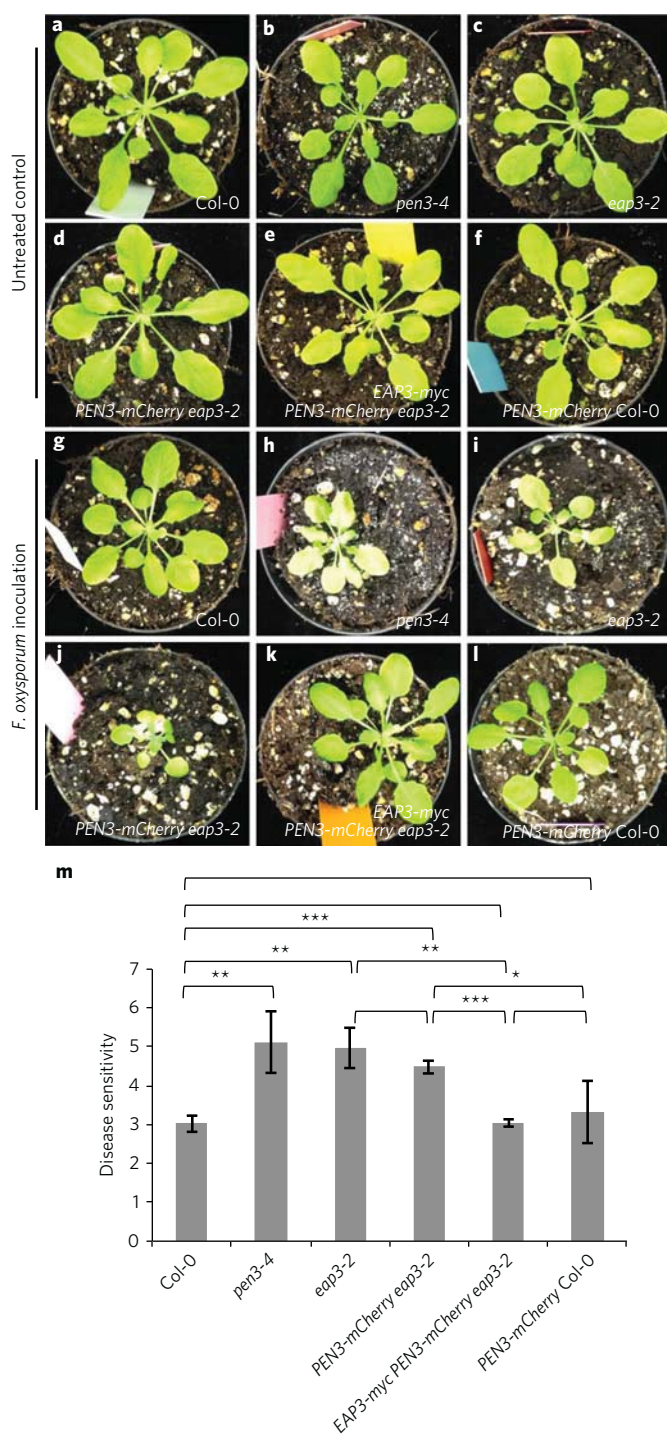


Fig. 3 | *pen3* and *eap3* mutations increase susceptibility to *F. oxysporum*. **a-f**, Five-week-old plants on soil without treatment. **g-l**, Five-week-old plants on soil 2 weeks post *F. oxysporum* inoculation. **h-j**, Note large number of chlorotic and necrotic leaves compared to **a-f** and **k-l**. **m**, Quantitative and statistical analysis of disease sensitivity. Averages from three independent experiments ($n=3$) are displayed, each employing 15–30 plants per genotype, with error bars indicating SD. Statistical differences with respect to increased disease susceptibility between genotypes were determined from the averages of three experiments ($n=3$) employing Student's one-tailed t-test with equal variance. Significance level was $P < 0.05$. *** indicates $P < 0.001$. ** indicates $P < 0.01$. * indicates $P < 0.05$. Exact P values are displayed in Supplementary Table 1.

leaves (Fig. 3h–j). As above, expressing EAP3-myc in the *eap3-2* *pPEN3:PEN3-mCherry* background rescued this mutant phenotype (Fig. 3k–m, Supplementary Table 1). The reduction of plant fresh weight after *F. oxysporum* inoculation showed a similar trend, revealing a significant reduction in the *pen3-4* and *eap3-2* mutants (Supplementary Fig. 6, Table 1). Together, these findings indicate a common function for EAP3 and PEN3 in disease resistance to a root-penetrating fungal pathogen.

Our study unravels a specific requirement of a BTB/POZ-domain protein for PEN3 membrane trafficking. The BTB/POZ-domain is an evolutionarily conserved motif and protein–protein interaction domain found in proteins involved in a wide range of cellular functions^{13–17}. The *Arabidopsis* genome encodes for around 80 BTB/POZ-domain proteins, of which a small subset act as substrate adaptors for E3 ubiquitin ligase complexes by interacting with CULLIN3 (CUL3)^{13–16}. Whereas no BTB/POZ-domain protein has previously been associated with ER membrane trafficking in plants, a precedent exists in mammals: Transport of procollagen I from the ER involves monoubiquitylation-dependent regulation of the COPII component SEC31, which requires the KLHL12 BTB/POZ-domain protein as a CUL3 adaptor^{13,17,18}. However, KLHL12 and EAP3 are molecularly unrelated except for harbouring a BTB/POZ domain. Moreover, the BTB/POZ domain of EAP3 displays poor conservation of the residues required for CUL3 binding and is not likely to function as an E3 ligase adaptor^{15,16}. Thus, we hypothesize that EAP3 mediates protein–protein interactions at the cytosolic face of the ER, where the ATP-binding cassette domains of PEN3 are located¹. Our findings pave the way for identification of cytosolic EAP3 interactors and for characterising the precise functions of the EAP3 protein domains during PEN3 exit from the ER. Interestingly, the ER accessory protein AUXIN RESISTANT4 and the PHOSPHATE TRANSPORTER TRAFFIC FACILITATOR1 protein, both unrelated to EAP3, mediate trafficking of specific cargos from the ER^{19,20}, although the underlying molecular mechanisms remain unknown. These examples highlight future opportunities in studying molecular mechanisms underlying ER exit of specific cargos involved in important physiological processes such as hormone or phosphate transport, and disease resistance in plants, to understand the physiological relevance of such specificity.

Methods

Plant materials and growth conditions. Plant materials and growth conditions are described in the Supplementary Methods.

Mapping and identification of the *eap3-1* mutation. The identification of the *eap3-1* mutation by recombination mapping and by mapping by genome sequencing is described in the Supplementary Methods.

Immunofluorescence labelling and confocal laser scanning microscopy analyses. Methods employed for immunofluorescence labelling and confocal laser scanning microscopy analyses are detailed in the Supplementary Methods.

FRAP and quantification of PEN3-mCherry localization. FRAP experiments were performed as described⁹ with the following minor modifications. In brief, FRAP experiments were conducted with the Zeiss LSM 780 system and objective (Zeiss, Oberkochen, Germany) described in the Supplementary Methods. Two to three consecutive cells per root were selected for full photobleaching. Within these cells, 25 bleaching iterations were performed at three to five different Z positions with 100% laser power of the 561 nm laser. Postbleach images were acquired at indicated time points using four-line averages in line-scanning mode with the same microscope settings. In order to quantify FRAP, PEN3-mCherry fluorescence intensities at the outer plasma membrane domain were measured using the segmented line function in Fiji software (<http://www.imagej.nih.gov/ij/>) in bleached cells for pre- and postbleach time points. For fluorescence intensities in the cytoplasmic area of bleached cells, the freehand selection function in Fiji was used to select the whole cytoplasmic area surrounded by plasma membrane for pre- and postbleach time points. Fluorescence intensities in non-bleached neighbouring cells at the outer plasma membrane were measured to normalize for the loss of fluorescence during FRAP caused by initial photobleaching and excitation during postbleach imaging. To correct for background fluorescence, background intensities were measured from root cells that did not express

PEN3-mCherry using the freehand selection function in Fiji software (<http://www.imagej.nih.gov/ij/>).

Quantification of PEN3-mCherry localization. The method for quantitative analysis of subcellular PEN3-mCherry fluorescence is detailed in the Supplementary Methods.

Immunoprecipitation and western blot analyses. Methods employed for immunoprecipitation and western blotting are described in the Supplementary Methods.

Quantification of PEN3-mCherry protein level. The procedure for quantification of PEN3-mCherry protein levels after western blotting is detailed in the Supplementary Methods.

RT-PCR analysis. The method employed for semiquantitative RT-PCR analysis is described in the Supplementary Methods. Primer sequences are listed in Supplementary Table 2.

Developmental map of transcript levels employing the eFP Browser tool. A description of the use of the eFP Browser tool for extraction and display of publicly available expression data shown in Supplementary Fig. 3 is available in the Supplementary Methods.

Plasmid construction and transformation. Descriptions of plasmid construction and transformation are available in the Supplementary Methods.

***Fusarium oxysporum* root infection assays in soil.** *Fusarium oxysporum* root infection assays in soil are described in the Supplementary Methods.

Accession numbers. Sequence data related to this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: *EAP3* (At3g09030), *PEN3* (At1g59870), *ACT7* (At5g09810), *PIN2* (At5g57090), *NIP5;1* (At4g10380), *PIS1* (At3g53480).

Data availability. The data that support the findings of this study are available from the corresponding author upon request.

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Author contributions

M.Gr. conceived the project. M.Gr., H.M., B.A. and M.Ge. designed experiments. H.M. performed all experiments except for *Fusarium oxysporum* work and for SHORE mapping. T.L. and J.H. performed SHORE mapping of *eap3-1*. B.A. and M.Ge. performed *Fusarium oxysporum* experiments. M.Gr. and H.M. wrote the manuscript. All authors read and edited the manuscript prior to publication.

Competing interests

The authors declare no competing financial interests.

Additional information

Supplementary information is available for this paper

Correspondence and requests for materials should be addressed to M.Gr.