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Published in: Cell Reports

DOI:

10.1016/j.celrep.2018.11.011

Publication date: 2018

Document version
Publisher's PDF, also known as Version of record

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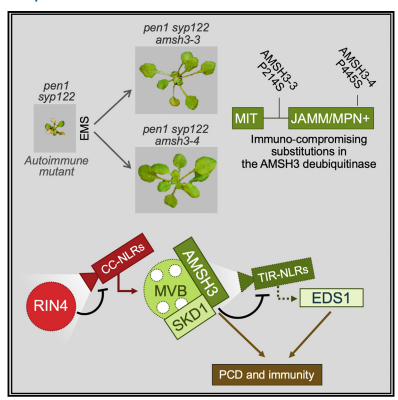
Citation for published version (APA): Schultz-Larsen, T., Lenk, A., Kalinowska, K., Vestergaard, L. K., Pedersen, C., Isono, E., & Thordal-Christensen, H. (2018). The AMSH3 ESCRT-III-Associated Deubiquitinase Is Essential for Plant Immunity. *Cell Reports*, 25(9), 2329-2338.e5. https://doi.org/10.1016/j.celrep.2018.11.011

Download date: 09. apr.. 2020

Cell Reports

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Graphical Abstract



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In Brief

NLR receptors are important for plant immunity as they monitor plant processes that are targeted by pathogens. Schultz-Larsen et al. find that the AMSH3 deubiquitinase enzyme is important for immunity activated by specific NLRs. Furthermore, their results suggest that AMSH3 itself is monitored by other NLRs.

Highlights

- The ESCRT-associated deubiquitinase AMSH3 is required for autoimmunity of pen1 syp122
- AMSH3 activity is required for immunity mediated by the CC-NLRs, RPM1 and RPS2
- Activity of SKD1 is required for activation of PCD downstream of RPM1
- The seedling lethality of the AMSH3 knockout is EDS1 dependent









The AMSH3 ESCRT-III-Associated Deubiquitinase Is Essential for Plant Immunity

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https://doi.org/10.1016/j.celrep.2018.11.011

SUMMARY

Plant "nucleotide-binding leucine-rich repeat" receptor proteins (NLRs) detect alterations in host targets of pathogen effectors and trigger immune responses. The Arabidopsis thaliana mutant pen1 syp122 displays autoimmunity, and a mutant screen identified the deubiquitinase "associated molecule with the SH3 domain of STAM3" (AMSH3) to be required for this phenotype. AMSH3 has previously been implicated in ESCRT-mediated vacuolar targeting. Pathology experiments show that AMSH3 activity is required for immunity mediated by the CC-NLRs, RPS2 and RPM1. Co-expressing the autoactive RPM1 $^{\mathrm{D505V}}$ and the catalytically inactive ESCRT-III protein SKD1^{E232Q} in *Nicotiana benthami*ana supports the requirement of ESCRT-associated functions for this CC-NLR-activated immunity. Meanwhile, loss of ESCRT function in A. thaliana is lethal, and we find that AMSH3 knockout-triggered seedling lethality is "enhanced disease susceptibility 1" (EDS1) dependent. Future studies may reveal whether AMSH3 is monitored by a TIR-NLR immunity receptor.

INTRODUCTION

Plants recognize pathogen invasion using plasma membrane-bound and intracellular receptors to trigger effective immune systems (Couto and Zipfel, 2016; Jones et al., 2016). To promote disease, pathogens transfer effectors into plant cells, where they target key immunity and susceptibility components. Plants monitor intracellular effector targets using "nucleotide-binding leucine-rich repeat" receptors (NLRs) (Jones et al., 2016). When effectors modify host proteins that are monitored by NLRs, effector-triggered immunity (ETI) is activated, and a complex signaling network translates it into an immune response involving the activation of hundreds of genes and often pro-

grammed cell death (PCD) of one or more cells (Jones et al., 2016; Spoel and Dong, 2012).

ETI activation follows different genetic pathways that are dependent on specific NLR N-terminal domains. In Arabidopsis thaliana, ETI activated by Toll-like/interleukin-1 receptor (TIR)-NLRs (TNLs) is dependent on "enhanced disease susceptibility 1" (EDS1) and PAD4, while ETI activated by coiled-coil (CC)-NLRs (CNLs) can require "non-race specific disease resistance 1" (NDR1) (Aarts et al., 1998; Wiermer et al., 2005; Day et al., 2006). Two CNLs, RPM1 and RPS2, mediate NDR1-dependent ETI when RIN4 is phosphorylated and cleaved by the bacterial effectors AvrRpm1 and AvrRpt2, respectively. A third CNL, RPS5, mediates NDR1dependent but RIN4-independent ETI (Jones et al., 2016). The TNL RPS4 activates EDS1-dependent ETI when RRS1-R is either acetylated by PopP2 or bound to AvrRps4 (Sarris et al., 2015). Screening for A. thaliana mutants with enhanced disease susceptibility has led to the identification of important immunity signaling components. Examples are EDS1, PAD4, and NDR1, which are required for NLR signaling. Meanwhile, SID2, EDS5, NPR1, and FMO1 are associated with salicylic acid (SA) and pipecolic acid (Pip) signaling, contributing to NLR activation of ETI (Wiermer et al., 2005; Zeier, 2013; Jones et al., 2016).

An alternative approach, based on lesion mimic mutants (LMMs) that exhibit constitutive autoimmune responses, has been highly successful for identifying novel immune components. Re-mutagenesis screens of LMMs have identified several defense components, such as 14 MOS genes (Johnson et al., 2012). Other examples are the LAZ genes identified in rescuing screens of acd11 (Rodriguez et al., 2016). It has been proposed that LMM phenotypes can be caused by the activation of NLRs, when guarded proteins, possibly pathogen effector targets, are lost due to mutations (Lolle et al., 2017; Rodriguez et al., 2016). Thus, rin4 is a severe LMM, rescued by rps2 CNL gene mutations (Mackey et al., 2003; Jones et al., 2016), and dependent on NDR1. Similarly, the LMM Isd1 is rescued by mutations in ADR1 CNL genes (Bonardi et al., 2011; Roberts et al., 2013). Other LMMs such as acd11, mpk4, smk7, and camta3 are partially or fully rescued by mutations in EDS1 and by mutations in TNL genes (Gloggnitzer et al., 2014; Lolle et al., 2017; Rodriguez et al., 2016; Zhang et al., 2012). The fact that EDS1 and NDR1 are required for NLR-mediated ETI (Aarts et al., 1998;



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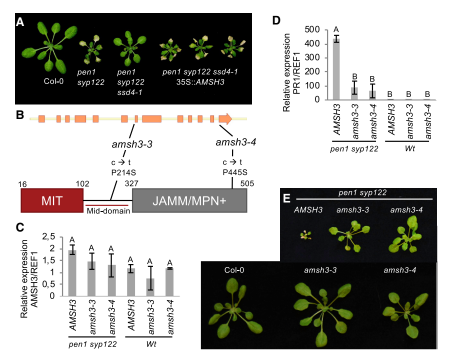
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Wiermer et al., 2005) can therefore be used to suggest that EDS1- and NDR1-dependent LMM autoimmunity are activated by monitoring NLRs.

PEN1, also referred to as SYP121, is an A. thaliana plasma membrane syntaxin that is required for penetration resistance against powdery mildew fungi (Collins et al., 2003). The closest homolog of PEN1 is SYP122, and pen1-1 syp122-1, hereafter referred to as pen1 syp122, is a severe LMM that requires EDS1, PAD4, NDR1, SID2, EDS5, FMO1, ALD1, and NPR1 (Zhang et al., 2007, 2008). Mutations in two of these genes can provide close-to-complete rescue of pen1 syp122 (Zhang et al., 2008). This shows that the LMM phenotype is genuinely due to autoimmunity and not developmental deficiencies directly caused by the loss of the two syntaxins. The EDS1, PAD4, and NDR1 dependency of this autoimmunity suggests that it is mediated by both TNL(s) and CNL(s), according to the reasoning above. The identities of those NLRs are unknown.

Here, we describe suppressor of syntaxin-related death 4 (SSD4), which is required for pen1 syp122 autoimmunity. We found that SSD4 is AMSH3, encoding a deubiquitinase associated with the "endosomal sorting complexes required for transport III" (ESCRT-III) machinery of multivesicular bodies (MVBs) (Isono et al., 2010), and that "associated molecule with the SH3 domain of STAM3" (AMSH3) is required for ETI activated by CNLs, RPM1, and RPS2. We present data suggesting that RPM1-mediated PCD triggered by the autoactive RPM1^{D505V} also requires SKD1, another ESCRT-III-associated component. We found that seedling lethality caused by knock out of AMSH3 (Isono et al., 2010) is EDS1 dependent, which makes us speculate that AMSH3 may not be a vital component of the cellular machinery per se. It may instead be guarded by one or more immunity-activating TNL.

Figure 1. The Full Lesion Mimic Phenotype of pen1 syp122 Requires AMSH3

(A) Representative pictures of 4-week-old plants: the 35S::AMSH3 complemented lines are independent T1 transformants.

(B) Schematic overview of the AMSH3 gene and

(C and D) AMSH3 (C) and PR1 (D) transcript profiling performed on 4-week-old plants. Error bars, SE (n = 3). Letters denote significant differences determined by ANOVA followed by a Tukey's honestly significant difference (HSD) post hoc test. (E) Five-week-old pen1 syp122 rescued by the two amsh3 mutant alleles, as well as amsh3-3 and amsh3-4 single mutants. These experiments have been repeated three times, with a similar result.

RESULTS

The Autoimmune Phenotype of pen1 syp122 Requires AMSH3

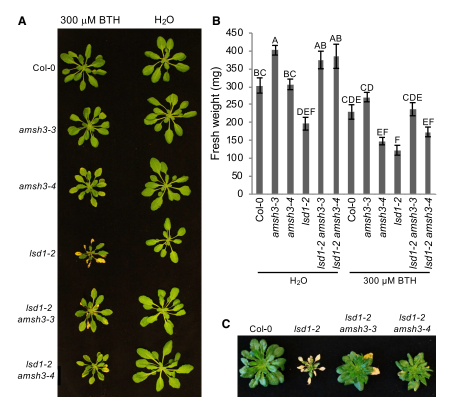
To identify components required for pen1 syp122 autoimmunity, an ethyl methanesulfonate (EMS) mutagenesis suppressor screen identified approximately 240 partially rescued pen1 syp122 ssd triple

mutants (Zhang et al., 2008). Here, we describe the pen1 syp122 ssd4-1 mutant (Figures 1A, 1E, S1, and S2). Note that the LMM phenotype of pen syp122, and thus the rescuing effect of ssd4-1, become more pronounced as the plants become older. F₁ plants of pen1 syp122 ssd4-1 × pen1 syp122 resembled pen1 syp122, and F2 plants segregated 3:1, which is consistent with a single recessive mutation causing the partial rescue.

Map-based cloning identified AMSH3 (At4g16144) as a candidate for SSD4 (see STAR Methods). AMSH3 is a deubiquitinase, and the ssd4-1 allele caused the P214S substitution (Figure 1B). qPCR showed that the AMSH3 was upregulated 2-fold in the pen1 syp122 mutant (Figure 1C). As transfer DNA (T-DNA) insertion alleles in AMSH3 cause seedling lethality (Isono et al., 2010), transgene complementation was used to confirm SSD4 being AMSH3. Here, pen1 syp122 ssd4-1 p35S::AMSH3 T₁ plants were pen1 syp122-like (Figure 1A). Complementation analysis within our collection of partially rescued pen1 syp122 ssd triple mutants identified another mutation at the AMSH3 locus, which caused the P445S substitution (Figures 1B and 1E). We designated our mutant alleles amsh3-3 (ssd4-1) and amsh3-4, respectively, following Isono et al. (2010). Thus, the autoimmune phenotype of pen1 syp122 requires AMSH3.

AMSH3 Function in the pen1 syp122 Autoimmune Phenotype Is Largely Independent of SA and Pip Signaling

The autoimmune phenotype of pen1 syp122 requires functional SA and Pip signaling (Zhang et al., 2008). To determine whether AMSH3 is required for SA and Pip signaling, we made use of the mutant alleles sid2-1, eds1-2, eds5-3, fmo1-1, pad4-19, and npr1-1 (Zhang et al., 2008). We introduced these six alleles



into the pen1 syp122 amsh3-3 triple mutant by crossing with triple mutants, in which they were already combined with pen1 and syp122 (all of these six triple mutants were partially rescued [Zhang et al., 2008; Figures S1 and S2]). The resulting quadruple mutants (e.g., pen1 syp122 pad4-19 amsh3-3) were larger than their triple mutant parents, suggesting that the AMSH3 function in pen1 svp122-conditioned autoimmunity is independent of SID2, EDS1, EDS5, FMO1, PAD4, and NPR1 (Figure S1). The pen1 syp122 npr1-1 triple mutant shows bleaching of the rosette, caused by SA hyperaccumulation (Zhang et al., 2007, 2008) (Figure S2A); and the pen1 syp122 npr1-1 amsh3-3 mutant also had bleached rosettes and stems (Figures S2A and S2B). As this phenotype is absent (e.g., after introducing sid2-1, which prevents SA accumulation, into pen1 syp122 npr1-1 [Zhang et al., 2008]), this suggested that SA accumulation still occurred in the presence of amsh3-3. Thus, the function of AMSH3 in autoimmunity did not appear to influence SA biosynthesis and signaling. This was corroborated by the fact that the expression of the NPR1-dependent SA marker gene PR1 (Cao et al., 1997) was only reduced by 80%-85% when pen1 syp122 was partially rescued by the amsh3-3 and amsh3-4 mutant alleles (Figure 1D); this suggests that SA signaling was largely independent of AMSH3. FMO1 plays a central role in Pip signaling (Zeier, 2013), and pen1 syp122 fmo1-1 amsh3-3 was also larger than the respective triple mutants (Figure S1). In summary, the genetic epistasis analysis suggests that the AMSH3 function in the pen1 syp122 autoimmunity is largely independent of SA- and Pipsignaling sectors.

Figure 2. AMSH3 Is Required for Isd1-Dependent rcd

(A and B) Phenotype (A) and rosette fresh weight (B) of 3-week-old plants, sprayed with 300 mM BTH or H₂O 7 days earlier. Error bars, SE (n = 9). For statistics, see Figure 1. The experiment was repeated three times, with a similar result.

(C) Thirteen-week-old short-day-grown plants, in which 3 weeks were at 20°C and then 10 weeks at 4°C. The experiments were repeated twice, with a similar result.

AMSH3 Is Required for ADR1-L1 and ADR1-L2-Dependent PCD

To obtain further insights into the biological relevance of AMSH3, the amsh3-3 and amsh3-4 single mutants were isolated after crossing to Col-0. They were developmentally similar to wild-type, but amsh3-3 plants were slightly larger and amsh3-4 plants slightly smaller than Col-0 (Figure 1E). In these 5-week-old plants, an age-related senescence phenotype appeared earlier on amsh3-4 plants than on Col-0. AMSH3 expression in amsh3-3 and amsh3-4 did not deviate from wild-type Col-0 (Figure 1C).

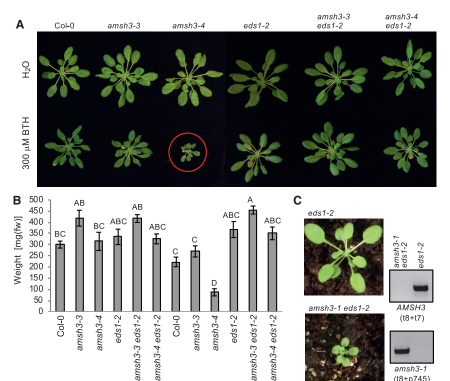
We found that AMSH3 was required for pen1 syp122 autoimmunity, but it was not

directly involved in SA and Pip signaling. To determine whether AMSH3 is required for cell death in other autoimmune mutants. we introduced amsh3-3 and amsh3-4 into the Isd1-2 and acd11-2 mutants (Figures 2C and S3). Isd1-2 has runaway cell death (rcd) governed by the CNLs ADR1-L1 and ADR1-L2 (Bonardi et al., 2011; Roberts et al., 2013), while acd11 has an autoimmune cell death phenotype that is dependent on the TNL LAZ5 (Rodriguez et al., 2016). The Isd1 autoimmune reaction can be enhanced by treatment with the SA analog bentothiadiazole (BTH). Seven days after BTH treatment, Isd1-2 plants showed typical rcd, whereas Col-0 wild-type amsh3-3 and amsh3-4 plants did not (Figure 2A). The Isd1-2 amsh3-3 and Isd1-2 amsh3-4 plants showed reduced rcd (Figures 2A and 2B), suggesting that AMSH3 is also required for the autoimmunity of Isd1-2 (see below regarding the intermediate effect of amsh3-4). Low temperature treatment also triggers rcd in Isd1-2 (Huang et al., 2010), which was strongly suppressed in the Isd1-2 amsh3-3 and Isd1-2 amsh3-4 (Figure 2C). However, introducing the amsh3-3 or amsh3-4 allele into acd11 failed to rescue its autoimmunity (Figure S3).

EDS1 Is Required for amsh3-4 Petiteness

We have noted that amsh3-4 plants, but not amsh3-3 plants, were significantly smaller than Col-0 wild-type after BTH application (Figures 2A and 3A). We therefore speculated that amsh3-4 displays a weak autoimmune phenotype. For this reason, we introduced the eds1-2 allele into amsh3-3 and amsh3-4 plants. The growth of amsh3-4 was recovered by the eds1-2 mutation, and the fresh weight of amsh3-4 eds1-2 was





300 µM BTH

Figure 3. EDS1 Is Required for amsh3-1 Lethality and amsh3-4 Petiteness

(A and B) Phenotype (A) and rosette fresh weight (B) of 3-week-old plants, sprayed with 300 mM BTH or H₂O 18 days earlier. Error bars, SE (n = 6-8). For statistics, see Figure 1. The experiment was repeated three times, with a similar result.

(C) Genotyped 4-week-old individuals from a segregating population from an amsh3-1 (+/-) eds1-2 parent. The AMSH3 wild-type and T-DNA insertion alleles were distinguished by PCR using AMSH3 primers (t8 and t7) and a T-DNA primer (p745).

indistinguishable from eds1-2 single mutants (Figures 3A and 3B). Furthermore, we noted that the amsh3-3 plants were slightly larger than Col-0, and likewise, the amsh3-3 eds1-2 plants were slightly larger than eds1-2. The amsh3-3 effect on plant weight was independent of BTH treatment (Figures 3A and 3B). amsh3-3 and amsh3-4 had opposing effects on the BTH response, despite that both alleles rescued the autoimmune phenotypes of pen1 svp122.

H₂O

The T-DNA insertion null mutations amsh3-1 and amsh3-2 and the ectopic overexpression of catalytically inactive AMSH3AXA cause developmental arrest and seedling lethality (Isono et al., 2010). Because this could be due to the aberrant activation of defense, similar to what the phenotype of amsh3-4 eds1-2 above suggests, we generated an amsh3-1 (+/-) eds1-2 (-/-) plant and analyzed the segregation of its progeny. We found that 39 plants resembled wild-type and 9 were slowly growing and dark green without necrosis (Figure 3C). This corresponds to a 3:1 segregation ratio. Genotyping confirmed that the slowly growing dark green plants were double homozygous amsh3-1 eds1-2 mutants (Figure 3C). Thus, the seedling lethality of AMSH3 knockout plants appears to reflect EDS1-dependent PCD.

The AMSH3 DUB Activity Appears Required for pen1 syp122 PCD

P445, substituted in AMSH3-4, is highly conserved in JAB1/ MPR1, PAD1 N-terminal+ (MPN+)/Mov34 metalloenzyme (JAMM)/MPN+-type deubiquitinases, and homology modeling and structure threading revealed that P445 is located in the catalytic cleft of the JAMM/MPN+ DUB domain; it is sterically juxtaposed to two catalytically important histidines, Ex(n)HxH(7)S(2)D

(Maytal-Kivity et al., 2002) (Figure 4A). Therefore, we analyzed whether AMSH3-4 and AMSH3-3 had altered substrate specificity or activity. We found that AMSH3-3 had catalytic activity comparable to that of wild-type AMSH3 protein (Figures 4B, 4C, and S5A). In contrast, AMSH3-4 had, as envisaged above, reduced catalytic activity and was unable to efficiently hydrolyze K48- and K63-linked ubiquitin oligomers in vitro (Figures 4B-4D). Yeast two-hybrid and subcellular localization revealed that neither AMSH3-3 nor AMSH3-4 interaction with the ESCRT com-

ponents VPS2.1 and ALIX (Figures S4A and S4B) and localization to SKD1^{E232Q}-induced aggregates deviated from those of AMSH3 (Figure S4C). Thus, the LMM phenotype of pen1 syp122 appears to require deubiquitinating enzyme (DUB) activity affected in AMSH3-4, as well as an unknown function affected in AMSH3-3.

amsh3-1 (t8+p745)

T-DNA knock out of AMSH3 or expression of AMSH3^{AXA} result in a dramatic accumulation of ubiquitin-conjugated proteins and developmental arrest (Isono et al., 2010). As AMSH3-4 had reduced catalytic activity, we examined the ubiquitination pattern in the amsh3-3 and amsh3-4 mutant plants. However, we found no significant alteration in the amount of ubiquitinconjugated proteins between wild-type Col-0, amsh3-3, and amsh3-4 plants (Figure 4E). This indicates that the altered catalytic activity of AMSH3-4 does not result in a dramatic global reorganization of the plant ubiquitinome. However, this can be observed during the induction of defense (Marino et al., 2012). Thus, we assayed global ubiquitin patterns in BTH-treated Col-0, amsh3-3, amsh3-4, eds1-2, amsh3-3 eds1-2, and amsh3-4 eds1-2 plants. Here, BTH-treated amsh3-4 plants experienced a dramatic increase in ubiquitin conjugates, which were significantly reduced, almost to wild-type levels, in amsh3-4 eds1-2 (Figure 4F). Thus, the increased accumulation of ubiquitin conjugates appeared to occur in plants as a result of EDS1-dependent PCD stimulation, not as a direct result of the reduced deubiquitinase activity found for AMSH3-4.

Resistance Conferred by RPM1 and RPS2, but Not RPS4 and RPS5, Depends on AMSH3

We speculated above that guarding of a PEN1/SYP122-mediated process by NLRs causes the autoimmunity and PCD of

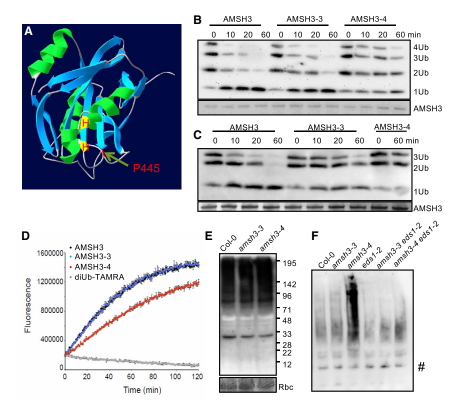


Figure 4. The P445S Substitution in AMSH3-4 Results in Reduced K48- and K63-Deubiquitinase Activity

(A) Homology model suggests that the substituted P445 in AMSH3-4 is juxtaposed to the catalytic site of the JAMM/MPN⁺ domain. Green, α-helices. Blue, β-sheets. Yellow, catalytic H411 and H413. Red, P445 substituted in AMSH3-4.

(B and C) K48-ubiquitin tetramers (B) and K63ubiquitin trimers (C) were incubated with AMSH3 variants in vitro for the indicated periods. The catalytic DUB activity was determined by the disappearance of ubiquitin oligomers visualized in the immunoblots by probing with antiubiquitin (top), and AMSH3, AMSH3-3, or AMSH3-4 protein levels were determined by anti-glutathione S-transferase (GST) (bottom). The experiments were repeated twice using independent AMSH3 protein purifications.

(D) In vitro fluorometric kinetics of AMSH3, AMSH3-3, or AMSH3-4 deubiquitinase activity using a TAMRA-based K63-linked diubiquitin. Error bars, SD (n = 4, each with three technical replicates). The experiment was repeated twice, with a similar result. See Figure S5A for gel with AMSH3 proteins. (E) Ubiquitin conjugation in total protein extracts from 4-week-old plants visualized on an immunoblot after probing with anti-ubiquitin (top). Rubisco (Rbc) loading control (bottom). The experiment was repeated four times, with a similar result.

(F) Ubiquitin conjugation in total protein extracts from 4-week-old plants, sprayed 18 days earlier with 300 μM BTH, visualized on an immunoblot after probing with anti-ubiquitin. #, non-specific crossreacting band serves as internal loading control.

pen1 syp122, and we have previously seen that proteins required for this autoimmunity are also required of immunity to pathogens (Zhang et al., 2007, 2008). To test whether AMSH3 is required for ETI, we spray inoculated amsh3-3. amsh3-4, and relevant control lines with Pseudomonas syringae pv. tomato (Pst) DC3000 expressing either AvrRpm1, AvrRpt2, AvrRps4, or AvrPphB. Col-0 expressed the NLRs guarding the function of these effectors (Jones et al., 2016). Increased bacterial titers of Pst DC3000 expressing AvrRpm1 and AvrRpt2 indicated that amsh3-3 and amsh3-4 were compromised in RPM1- and RPS2-conditioned resistance (Figures 5A and 5B). In contrast, bacterial titers in amsh3-3 and amsh3-4 mutant plants were indistinguishable from Col-0 wild-type when inoculated with Pst DC3000 expressing AvrPphB or AvrRps4 (Figures 5C and 5D). This suggested that AMSH3 is required for resistance conditioned by RPM1 and RPS2, but not by RPS4 and RPS5.

RPM1- and RPS2-Dependent HR-PCD Requires AMSH3

ETI is associated with PCD at the site of infection, and consequent vacuolar rupture and cellular ion leakage can be measured as changes in media conductivity (Hofius et al., 2009). To analyze whether the AMSH3 requirement in resistance was reflected in the PCD, we challenged Col-0, amsh3-3, amsh3-4, and ndr1-1 with Pst DC3000 (AvrRpm1) and tested for ion leakage. In Col-0, an increase in conductivity was observed by 3 hr post-inoculation (hpi). This increase was delayed in amsh3-3, amsh3-4, and ndr1-1 (Figure 5E). To corroborate these findings, we performed this experiment using Pst DC3000 (AvrRpt2) as well. The experiment revealed that Col-0 and amsh3-3 plants had similar conductivity kinetics, with a marked increase at ~5 hpi (Figure 5F). However, the ion leakage was considerably reduced in amsh3-4 plants, although not to the extent seen in the ndr1-1 control plants (Figure 5F). These results suggest that normal activation of PCD by RPM1 and RPS2 depends on AMSH3 and its catalytic deubiquitinase activity.

The Pseudomonas effectors AvrRpm1 and AvrRpt2 both target RIN4 to promote virulence. The CNLs RPM1 and RPS2 trigger PCD when RIN4 is altered: RPM1 is triggered by RIN4 phosphorylation and RPS2 is triggered by the loss of RIN4 interaction (Jones et al., 2016). However, overexpression of RIN4 has been shown to inhibit RPS2-dependent, although not RPM1-dependent, hypersensitive response (HR) (Mackey et al., 2003). Therefore, to test whether the effect of our amsh3 mutations on resistance involved changes in the amount of RIN4, we made an immunoblot of the RIN4 protein. However, no difference between Col-0 wild-type, amsh3-3, and amsh3-4 was found (Figure S5B). Thus, the reduced resistance to Pst DC3000 expressing AvrRpm1 or AvrRpt2 is not likely due to an increased steady-state level of RIN4.



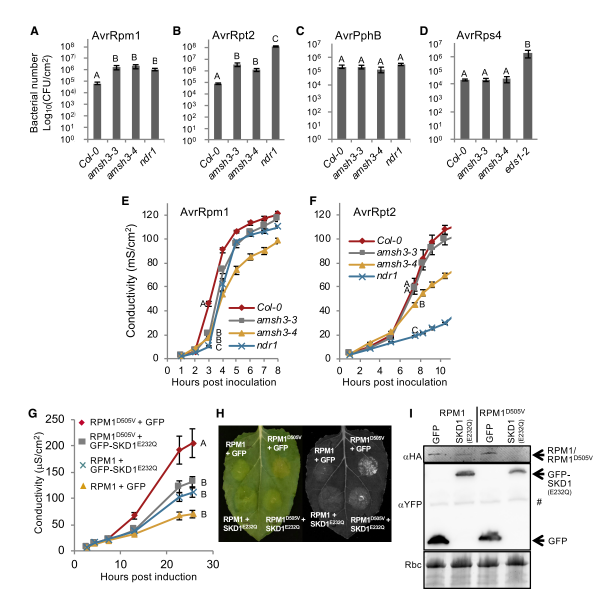


Figure 5. Involvement of AMSH3 and SKD1 in CNL-Mediated Immunity and Cell Death

(H) Leaf 3 days post-β-estradiol treatment. Left, visible light. Right, UV trans-illuminated.

(A-D) Four- to 5-week-old A. thaliana Col-0 plants were spray inoculated with Pst DC3000 expressing four different effectors: (A) AvrRpm1, (B) AvrRpt2, (C) AvrPphB, and (D) AvrRps4. The bacterial populations in the plants were determined 3 days post-inoculation (dpi). Error bars, SE (n = 12). (E and F) For ion leakage analyses, 4-week-old A. thaliana Col-0 leaves were infiltrated with Pst DC3000 AvrRpm1 (E) and Pst DC3000 AvrRpt2 (F). Error bars, SE

(G-I) Ion leakage and HR on N. benthamiana leaves 2 days after Agrobacterium-mediated transient co-expression of the indicated proteins. (G) GFP-SKD1^{E232Q} and GFP were 35S promoter regulated, while the expression of RPM1^{D505V} or RPM1 was induced by β-estradiol. Error bars, SE (n = 4).

(I) Immunoblot of total protein 5 hr after β-estradiol treatment. #, non-specific cross-reacting band. Rubisco (Rbc) loading control (bottom).

Letters denote significant differences at the indicated time points determined by ANOVA followed by a Tukey's HSD post hoc test. The experiments were repeated three times with a similar result.

Normal ESCRT-III Function Is Required for Activation of PCD Downstream of RPM1

As the AMSH3 deubiquitinase activity appeared important for immunity-related PCD, we wanted to test whether this could be due to a requirement for an intact ESCRT system. As discussed above, overexpression of SKD1 E232Q interferes with normal ESCRT function and stimulates MVBs to aggregate

(Haas et al., 2007). Therefore, we used SKD1 E232Q to block ESCRT function and test for RPM1-triggered PCD. To this end, we used the autoactive RPM1 D505V that mimics normal RPM1 activation both in A. thaliana and Nicotiana benthamiana in the absence of RIN4^{T166} phosphorylation (Gao et al., 2011). We transiently transformed N. benthamiana to co-express GFP-SKD1^{E232Q} or GFP with RPM1 or RPM1^{D505V}. RPM1

co-expressed with GFP-SKD1^{E232Q} displayed elevated levels of ion leakage relative to co-expression with GFP (Figure 5G). This was consistent with a previous report showing that SKD1^{E232Q} caused cell death (Haas et al., 2007). Expression of RPM1^{D505V} triggered a rapid increase in ion leakage, which was suppressed by co-expression with GFP-SKD1^{E232Q}, despite its own activation of PCD. The level of lesion formation in *N. benthamiana* after 3 days was consistent with the conductivity assays (Figure 5H), which suggest that cell death triggered by RPM1^{D505V} requires SKD1.

The stability of GFP and GFP-SKD1^{E232Q} was apparently not affected by the co-expression with RPM1 or RPM1^{D505V} (Figure 5I). However, SKD1^{E232Q} influenced RPM1 stability. Here, the autoactive RPM1^{D505V} was not detectable when co-expressed with SKD1^{E232Q} (Figure 5I). This may suggest that activated signaling-competent RPM1 is stabilized by SKD1.

DISCUSSION

The DUB Activity of AMSH3, Conserved in Eukaryotes, Is Required for PCD

To date, the ESCRT-I components VPS28-1 and VPS37-1, as well as the SKD1/VPS4-associated protein LIP5, have been identified as positive regulators of pathogen-associated molecular pattern (PAMP)-triggered immunity (Spallek et al., 2013; Wang et al., 2014). Here, we identified the ESCRT-III-associated deubiquitinase AMSH3 as a regulator of pen1 syp122 autoimmunity. In addition, our amsh3 mutations were able to suppress rcd conditioned by ADR1-L1 and ADR1-L2, as well as RPS2- and RPM1-conditioned defense responses seen as higher bacterial growth and reduced PCD. These data suggest that AMSH3 is required for PCD mediated by RPM1, RPS2, and ADR1s, all of which are CNLs. While RPM1 and RPS2 are interactors of RIN4, PCD triggered by the CNL RPS5, which is not an interactor of RIN4, did not require AMSH3. The same was true for PCD triggered by the TNL RPS4.

There are approximately 50 deubiquitinases in A. thaliana, and AMSH3 is 1 of 8 A. thaliana members of the JAMM subfamily found broadly in eukaryotes (Isono and Nagel, 2014). They are zinc metalloproteases that coordinate two zinc ions in their MPN+ domain, which activates a water molecule to hydrolyze ubiquitin isopeptide bonds to lysines on target proteins. AMSH3 can hydrolyze both K48- and K63-linked ubiquitin chains (Isono et al., 2010), and the P445S substitution in the MPN+ domain reduced this catalytic activity, whereas the P215S substitution did not. Proline is conformationally rigid, and it strongly affects the secondary structure of the amino acid chain in its vicinity. Therefore, a model could be that P445S affects the catalytically active site, whereas P215S affects another AMSH3 feature, which we have not been able to unravel. In summary, our observations point toward AMSH3 catalytic activity being required for autoimmunity and PCD being mediated by a subset of CNLs.

We found that AMSH3 was required for PCD triggered by AvrRpm1 and AvrRpt2 and mediated by the CNLs RPM1 and RPS2, which guard the status of RIN4. AMSH3 being an ESCRT-III component hinted that ESCRT-III in general is required for such PCD. To address this, we activated PCD in *N. benthamiana* by expressing the autoactive RPM1^{D505V} (Gao

et al., 2011) and co-expressed a dominant-negative version (SKD1^{E232Q}) of the AAA-ATPase, which is essential for ESCRT-III (Haas et al., 2007). This experiment supported the AMSH3 results and suggested a general function of this pathway in PCD triggered by RPM1. These data also imply that ESCRT-III functions downstream of RIN4. Furthermore, our data suggest that the stability of the PCD signaling active but not inactive RPM1 may be influenced by the SKD1^{E232Q} mutation. This result may explain why SKD1 inactivation resulted in reduced RPM1-dependent PCD. This line of reasoning also suggests a simple model explaining why AMSH3 is required for *pen1 syp122*-dependent PCD: loss of *pen1 syp122* function triggers PCD through an NLR, the stability of which is dependent on AMSH3.

Loss of AMSH3 Triggers PCD Dependent on EDS1

AMSH3 and its closest paralog, AMSH1, share domain organization in which both have an N-terminal microtubule interacting and transport (MIT) domain, a middle domain, and a C-terminal JAMM/MPN⁺ domain (Isono et al., 2010). AMSH3 makes a larger contribution to development, which is evident by the fact that amsh3-1 null mutant is seedling lethal, while the amsh1-1 knockdown mutant displays early senescence (Isono et al., 2010; Katsiarimpa et al., 2013). Because expression of AMSH3^{AXA} also results in seedling lethality, it is the loss of AMSH3 catalytic deubiguitinase activity and not the absence of the protein that triggers PCD (Isono et al., 2010). This is consistent with the observations above, as AMSH3-4 has reduced catalytic activity and amsh3-4 plants have reduced weight and rosette size, stimulated by BTH. It is noteworthy that both the amsh3-1 seedling lethality and the amsh3-4 size reduction require EDS1. Therefore, we suggest that these phenotypes reflect PCD governed by a TNL, as they are normally dependent on EDS1. The amsh1 knockdown mutant displays spontaneous PCD upon pathogen challenge (Katsiarimpa et al., 2013), which is consistent with the hypothesis that an NLR response is activated.

We speculate that a TNL is triggered by the state or reduced turnover of one or more AMSH1/AMSH3-DUB substrates. This putative substrate is presumably sorted through the ESCRT system, as mutations of core ESCRT-III components (e.g., SUCROSE NONFERMENTING 7.1 [SNF7.1], VPS2.1, VPS20.1, and VPS24.1, which are affected in vacuolar trafficking) also cause seedling lethality (Katsiarimpa et al., 2013; Cai et al., 2014). Likewise, overexpression of the catalytically inactive A. thaliana SKD1 E232Q and rice SKD1 E315Q caused lethality in tobacco BY-2 cells and rice leaves, respectively (Haas et al., 2007; Zhu et al., 2016). Similarly, knock down of positive regulator of SKD1 (PROS) or knock out of LIP5, both of which activate SKD1, has a negative effect on plant growth (Reyes et al., 2014). LIP5 interacts with ISTL1, whose yeast homolog regulates SKD1 activity (Tan et al., 2015), and the istl1 lip5 double mutant resembles an LMM by having stunted and necrotic leaves with constitutive PR1 expression (Buono et al., 2016). These findings agree with a model in which reduced turnover of an ESCRT-III cargo activates a TNL to trigger PCD. In addition, mutations in the ESCRT-II component gene VPS36 resemble an amsh3-1 null mutant (Wang et al., 2017). Similarly, the ESCRT-I double mutants vps28-1 vps28-2 and vps37-1 vps37-2 are unable to germinate (Chikkaputtaiah, 2008).



Future studies may reveal whether these ESCRT proteins are essential, although their inhibition causes severe alterations in MVBs. They may alternatively be monitored by one or more immunity-activating NLRs. Such a model is supported by evidence put forward by Sutipatanasomboon et al. (2017), who observed eds1-mediated rescue of the autoimmune phenotype of the ESCRT-I component mutant cfs1. Thus, it is intriguing that the ESCRT pathway is dispensable in yeast (Hurley, 2015). Having observed that AMSH3 and SKD1 play positive roles in immunity, it may be biologically meaningful that NLRs guard the ESCRT-III pathway in plants.

In summary, our analyses lead us to suggest a model for a role of the ESCRT-III-associated AMSH3 and SKD1 in the execution of CNL-triggered PCD in plant immunity, and a likely guarding of these components by TNLs.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures and one table and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.11.011.

ACKNOWLEDGMENTS

We would like to thank Mette Sylvan for excellent technical assistance and Prof. Jeffery L. Dangl (University of North Carolina, USA) for sharing the pEst-T7-RPM1-YFP-HA and pEst-T7-RPM1(D505V)-YFP-HA constructs. This work was funded by the Villum Foundation, Denmark (VKR023502), and the German Science Foundation (DFG) (IS221/4-1).

AUTHOR CONTRIBUTIONS

Conceptualization, T.S.-L., A.L., and H.T.-C.; Methodology, T.S.-L., E.I., and H.T.-C.; Investigation, T.S.-L., A.L., K.K., L.K.V., and C.P.; Resources, E.I.

and H.T.-C.; Writing – Original Draft, T.S.-L. and H.T.-C.; Writing – Review & Editing, T.S.-L., E.I., and H.T.-C.; Visualization, T.S.-L., E.I., and A.L.; Supervision, E.I. and H.T.-C.; Project Administration, H.T.-C.; Funding Acquisition, H.T.-C.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: December 30, 2017 Revised: March 4, 2018 Accepted: October 31, 2018 Published: November 27, 2018

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-AMSH3 Rabbit polyclonal	Isono et al. (2010)	N/A
Anti-ubiquitin Rabbit polyclonal	Sigma-Aldrich	Cat# Sigma-U5379
Anti-GFP Mouse monoclonal	Santa Cruz Biotechnology	Cat# sc-9996
Anti-RIN4 Goat polyclonal	Santa Cruz Biotechnology	Cat# sc-27369
Anti-PEN1 Rabbit polyclonal	Zhang et al. (2007)	N/A
Anti-HA Rat monoclonal	Chromotek	Cat# 7C9
Anti-GST Mouse monoclonal	Genscript	Cat# A00865-40
Anti-Gal4-BD (GBD) Mouse monoclonal	Clontech	Cat# 630403
Anti-Rat AP-conjugated	Sigma-Aldrich	Cat# A3687
Anti-Rabbit HRP-conjugated	Santa Cruz Biotechnology	Cat# sc-2004
Anti-Mouse HRP-conjugated	Sigma-Aldrich	Cat# A4416
Anti-Goat AP-conjugated	Santa Cruz Biotechnology	Cat# sc-2771
Bacterial and Virus Strains		
Pst DC3000 AvrRpm1	Grant et al. (1995)	N/A
Pst DC3000 AvrRpt2	Kunkel et al. (1993)	N/A
Pst DC3000 AvrRps4	Hinsch and Staskawicz (1996)	N/A
Pst DC3000 AvrRphB	Simonich and Innes (1995)	N/A
Agrobacterium tumefaciens GV3101 pMP90	Koncz and Schell (1986)	N/A
One Shot TOP10	Invitrogen	Cat# K2400-20
Escherichia coli Rosetta 2	Merck	Cat# 71405
Chemicals, Peptides, and Recombinant Proteins		
Gateway BP Clonase II Enzyme Mix	ThermoFisher Scientific	Cat# 11789100
Gateway LR Clonase II Enzyme Mix	ThermoFisher Scientific	Cat# 11791100
Beta-estradiol	Sigma-Aldrich	Cat# E8875
Trypan blue	Sigma-Aldrich	Cat# T-0776
c0mplete protease inhibitor cocktail	Sigma-Aldrich	Cat# 11697498001
Protino Glutathione Agarose 4B	Machery-Nagel	Cat# 745500.10
K48-ubiquitin tetramer	Boston Biochem	Cat# UC-210B
K63-ubiquitin trimer	Boston Biochem	Cat# UC-315B
ΓΑΜRA-labeled K63-ubiquitin dimer	R&D Systems	Cat# P62988
PTG	Sigma-Aldrich	Cat# I6758
BASTA	Bayer	Cat# L4872
ЗТН	Syngenta	Cat# A9180A
Critical Commercial Assays		
QuikChange II XL Site-Directed Mutagenesis kit	Agilent	Cat# 200521
SMART MMLV RT kit	Clontech	Cat# 634901
5x HOT FIREPol EvaGreen qPCR Mix Plus	Solis BioDyne	Cat# 08-24-00001
Experimental Models: Organisms/Strains		
Arabidopsis thaliana Col-0 wild-type	N/A	N/A
A. thaliana Col-0 amsh3-3	This study	N/A
A. thaliana Col-0 amsh3-4	This study	N/A
A. thaliana Col-0 amsh3-1	Isono et al. (2010)	N/A
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(Continued on next page)



Continued	0011005	IDENTIFIED
REAGENT or RESOURCE	SOURCE	IDENTIFIER
A. thaliana Col-0 eds1-2	Bartsch et al. (2006)	N/A
A. thaliana Col-0 acd11-2	Brodersen et al. (2005)	N/A
A. thaliana Col-0 lsd1-2	Torres et al. (2005)	N/A
A. thaliana Col-0 pen1-1 syp122-1	Zhang et al. (2007)	N/A
A. thaliana Ler pen1-1 syp122-1	Zhang et al. (2007)	N/A
A. thaliana Col-0 pen1-1 syp122-1 35S::AMSH3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 eds1-2	Zhang et al. (2007)	N/A
A. thaliana Col-0 pen1-1 syp122-1 fmo1-1	Zhang et al. (2008)	N/A
A. thaliana Col-0 pen1-1 syp122-1 npr1-1	Zhang et al. (2007)	N/A
A. thaliana Col-0 pen1-1 syp122-1 pad4-19	Zhang et al. (2008)	N/A
A. thaliana Col-0 pen1-1 syp122-1 sid2-1	Zhang et al. (2007)	N/A
A. thaliana Col-0 pen1-1 syp122-1 amsh3-3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 amsh3-4	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 eds1-2 amsh3-3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 fmo1-1 amsh3-3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 npr1-1 amsh3-3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 pad4-19 amsh3-3	This study	N/A
A. thaliana Col-0 pen1-1 syp122-1 sid2-1 amsh3-3	This study	N/A
A. thaliana Col-0 amsh3-1 eds1-2	This study	N/A
Oligonucleotides		
Primers used in this study, see Table S1	This study	N/A
Recombinant DNA		
DENTR-D-TOPO	ThermoFisher Scientific	Cat# K2400-20
pK7WGF2	Karimi et al. (2007)	N/A
pB2GW7: 35S-AMSH3	This study	N/A
pK7WGR2: p35S-RFP	Karimi et al. (2007)	N/A
pK7WGR2: p35S-RFP-AMSH3 (WT)	This study	N/A
pK7WGR2: p35S-RFP-AMSH3 (P214S)	This study	N/A
pK7WGR2: p35S-RFP-AMSH3 (P445S)	This study	N/A
pK7WGF2: p35S-GFP	This study	N/A
pK7WGF2: p35S-GFP-SKD1 (E232Q)	This study	N/A
pEst-T7-RPM1-YFP-HA	Gao et al. (2011)	N/A
pEst-T7-RPM1 (D505V)-YFP-HA	Gao et al. (2011)	N/A
pDEST-ACT2	Robertson (2004)	N/A
pDEST-ACT2: HA-GAD-AMSH3 (WT)	This study	N/A
pDEST-ACT2: HA-GAD-AMSH3 (P214S)	This study	N/A
pDEST-ACT2: HA-GAD-AMSH3 (P445S)	This study This study	N/A
pGBKT7: GBD	Kalinowska et al. (2015)	N/A
pGBKT7: GBD-ALIX	Kalinowska et al. (2015)	N/A
ogbkt7: gbd-Alix	Katsiarimpa et al. (2013)	N/A
	· · · · · ·	
pT7-GST-AMSH3	This study This study	N/A
TT CCT AMCUA (DOLAC)	inie etilav	N/A
pT7-GST-AMSH4 (P214S)	•	
pT7-GST-AMSH4 (P214S) pT7-GST-AMSH3 (P445S) Software and Algorithms	This study This study	N/A



CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Hans Thordal-Christensen (htc@plen.ku.dk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Arabidopsis

All A. thaliana (L.) Heynh. mutants and transgenic lines listed above were sown in soil, then incubated at 4°C for 3 days, and subsequently grown in growth chamber at 21°C and 70% relative humidity under short-day conditions with 8-hr light at 125 μ Einstein s⁻¹m⁻².

Nicotiana benthamiana

N. benthamiana plants were greenhouse-grown in soil under long-day conditions (16-hr light) at 24°C.

Bacterial strains

Pseudomonas syringae pv. tomato (Pst) DC3000 expressing either AvrRpm1, AvrRpt2, AvrRps4, or AvrPphB were grown on LB plates with 100 μ g/ml rifampicin and 50 μ g/ml kanamycin for 2 days for inoculation.

METHOD DETAILS

Map-based cloning

To identify the ssd4-1 mutation that suppressed the pen1 syp122 phenotype, an F₂ population from a cross between pen1 syp122 ssd4-1 in ecotype Columbia (Col-0) and pen1-1 syp122-1 backcrossed into ecotype Landsberg erecta (Ler) (Zhang et al., 2008) was used. Initial mapping positioned SSD4 on chromosome 4 in an interval between the markers, ciw6 (At4q13570) and FCA8 (At4g17200). Subsequent fine-mapping placed SSD4 in a region of 17 genes spanning from At4g16130 to At4g16215. The ssd4-1 mutation was identified after PCR-amplification and sequencing of all coding sequences in the interval based on genomic DNA.

BTH-treatment

Plants were sprayed with 300 µM BTH or water until the leaves were fully wet. The rosette fresh weight was determined after cutting the rosette from the stem.

Plasmid constructs

All cloning was done using the Gateway system (Thermo Fisher Scientific). The coding sequence for AMSH3 was cloned from cDNA into pENTR-D-TOPO, and those for AMSH3-3 and AMSH3-4 were generated by site-directed mutagenesis using the QuikChange II Clonase kit. All were transferred to pDEST15 (GST-tag), using the LR-II kit (Invitrogen), and transformed into the E. coli Rosetta 2 strain (Merck) for protein expression. Coding sequences for N-terminally GFP-tagged SKD1^{E232Q} (Katsiarimpa et al., 2011) and the AMSH3 versions were generated by recombining the pENTR clones with pK7WGF2 and pK7WGR2, respectively, using the LR-II Clonase kit. These constructs were then transformed into Agrobacterium tumefaciens GV3101 pMP90. For yeast two-hybrid assays, the AMSH3 alleles were introduced into the pDEST-ACT2 plasmid (Robertson, 2004) using the LR-II Clonase kit. Primers were synthesized at TAG Copenhagen (TAGC.dk).

Stable transformation of A. thaliana and transient protein expression in N. benthamiana

The A. tumefaciens strain GV3101 harboring binary protein expression vectors was grown overnight on LB plates (10 g tryptone, 5 g yeast extract, 5 g NaCl, 15 g Agar per liter) containing 100 μg/ml rifampicin, 100 μg/ml spectinomycin, and 25 μg/ml gentamycin. The cells were harvested and resuspended in 10 mM MgCl₂. Stable transformants of A. thaliana were obtained after floral dipping and subsequent selection of plants germinating from the harvested seeds on MS plates containing Basta (60 μ g/ml) (Clough and Bent, 1998). Co-expression in N. benthamiana was obtained by mixing the A. tumefaciens strains (each with an $OD_{600} = 0.3$) with the indicated plasmids, which then were infiltrated with a needleless syringe into leaves of 4-5-week-old plants.

Bacterial growth assays

Pst DC3000 expressing either AvrRpm1, AvrRpt2, AvrRps4, or AvrPphB were harvested and resuspended in 10 mM MgCl₂ and 0.01% v/v Silwet L-77 to 108 CFU/ml and spray-inoculated onto plants. To maintain high humidity the plants were covered with a translucent lid for one day. Bacterial populations within the leaves were determined 3 dpi. For each repeat, four leaf discs size 3 were sampled from the same rosette and ground in 200 µl 10 mM MgCl₂, using a pistle mounted on an electric drill, while adding 800 μl of 10 mM MgCl₂. A 6-times serial dilution was performed. Twenty microliters of each of the six dilutions were plated on LB plates supplemented with appropriate antibiotics. The colonies were counted 2-3 days after plating.



Ion leakage experiments

A. thaliana plants were infiltrated with 108 CFU/ml bacteria and five leaf discs harvested for each repeat using a cork-borer size 4. The discs were floated on water for 60 min. and then transferred to a 12 mL culture tube containing 5 mL water (5 discs per tube = one repeat). Experiments in N. benthamiana were performed in the same way 2 dpi with A. tumefaciens harboring the indicated plasmids. The expression of RPM1 or RPM1 D505V was induced with 40 μM β-Estradiol 2 hours prior to the leaf discs being cored and floated as described above. The change in conductivity of the solution was measured with a conductivity meter (CRM210, Radiometer) at the indicated time points.

Yeast two-hybrid assay

The interaction between AMSH3 versions and other ESCRT components were analyzed by co-transforming Saccharomyces cerevisiae Y190 with pDEST-ACT2:AMSH3, pDEST-ACT2:AMHS3-3 or pDEST-ACT2:AMSH3-4 combined with pGBKT7:VSP2.1 (Katsiarimpa et al., 2013) or pGBKT7:Alix (Kalinowska et al., 2015). The yeast two-hybrid assay was performed on synthetic complete medium without leucine and tryptophan (-LW) as well as histidine (-LWH). Higher stringency condition was obtained by adding 3-amino-1,2,4-triazole (3AT).

Total protein extraction

Total protein from N. benthamiana was extracted by grinding three leaf discs, made using cork-borer size 3, in liquid nitrogen with metal beads in 1.5 mL Eppendorf tubes using a Retsch mill. Two hundred and fifty microliters of 2 x modified Laemmli buffer (0.24 M Tris-Cl, pH 6.8, 6% SDS, 30% glycerol, 16% β-mercaptoethanol, 0.006% bromophenol blue, 10 M urea) were added to the samples, which then were heated for 5 min. at 95°C. The samples were then subjected to SDS-PAGE and western blot analysis as described below.

Total protein was extracted from A. thaliana by grinding 4-5-week-old plant leaf tissue in liquid nitrogen using a mortar and pestle. The proteins were extracted by incubation in cold buffer (100 mM Tris-HCl pH 7.5, 150 mM NaCl, 5 mM EDTA, 5% glycerol, 0.5% Triton X-100, 10 mM DTT and 1x c0mplete protease inhibitor cocktail (Roche)) for 30 min. at 4°C. The protein concentration of supernatants from a 12,000 x g centrifugation, for 20 min. at 4°C, was determined using BioRad's Bradford reagent.

For western blots, samples were heated for 5 min. at 95°C with 4x RunBlue LDS sample buffer (40% Glycerol, 4% LDS, 0.8 M Triethanolamine-Cl pH 7.6, 4% Ficoll-400, 0.025% Phenol Red, 0.025% Coomassie Brilliant Blue G250, 2 mM EDTA-2Na). The proteins were separated on Any kD Mini-PROTEAN® TGX Precast Protein Gels (BioRad) or 10% SDS-PAGE gels (Expedeon) and subsequently transferred to PVDF membranes (BioRad) using Trans-Blot Turbo Transfer System (BioRad). Immuno-blots were performed using 1:1000 anti-PEN1 (Zhang et al., 2007), 1:5000 anti-GFP (Santa Cruz Biotech.), 1:1000 anti-ubiquitin (Sigma-Aldrich), 1:1500 anti-AMSH3 (Isono et al., 2010), 1:1000 anti-HA (Chromotek), 1:1000 anti-RIN4 (Santa Cruz Biotech.), 1:1000 anti-Gal4 BD (GBD) (Santa Cruz Biotech.), 1:1000 anti-HA-HRP (Roche) and 1:2000 anti-YFP (Agrisera). Blots were reacted with AP- or HRP-conjugated secondary antibodies and revealed with BCIP/NBT (Promega) or Clarity Western ECL (Biorad), respectively.

RNA isolation and quantitative PCR

Total RNA was isolated from A. thaliana tissue using TRIzol (Ambion) and RQ1 RNase-free DNase (Promega)-treated. cDNA was synthesized using the SMARTTM MMLV RT kit (Clontech) and an oligo(dT) primer.

Transcript quantification was performed using the 5x HOT FIREPol EvaGreen qPCR Mix Plus (SOLIS BIODYNE) on a Stratagene MX3000P real-time PCR detection system. Gene specific primers are described in Table S1. We used the At4g26410 transcript as internal control (Czechowski et al., 2005).

DUB assavs

Over-night cultures (OD₆₀₀ = 0.6) of E. coli Rosetta 2 expressing the three versions of AMSH3 were supplied with IPTG to 500 µM and incubated at 18°C for another 18 h. The cells were harvested and resuspended in Buffer A (50 mM Tris-HCI, 100 mM NaCI, 10% glycerol) supplied to 1x c0mplete protease inhibitor cocktail (Roche) and 0.5% Triton X-100. Cells were sonicated for 10 × 15 s. at setting 5 of 10, and the cellular debris pelleted at 12.500 x g for 10 min. Hundred microliters Protino® Glutathione Agarose 4B (Macherey-Nagel) were added to the supernatant, and the slurry was incubated by rolling for 2 h. at 4°C. The beads were pelleted by a 500 x g centrifugation for 5 min. and washed three times in Buffer A. The bound proteins were eluted in 200 mM glutathione, pH 8.0. Gels with the purified proteins are shown in Figures 4B and 4C and S5.

Deubiquitination experiments were performed using K48- or K63-linked ubiquitin tetramer and trimer chains, respectively (Boston Biochem.). Two hundred fifty nanogram of ubiquitin chains were mixed with 2 pmol of E. coli-expressed AMSH proteins and incubated for the indicated time in a DUB Assay Buffer (50 mMTris-HCl pH 7.2, 25 mM KCl, 5 mM MgCl₂, 1 mM DTT) at 30°C. The reaction mixture was subjected to immunoblotting using an anti-Ubiquitin antibody. In a second deubiquitination assay 400 nM of K63-linked di-ubiquitin FRET TAMRA (position 3) (Boston Biochem) was mixed with 50 nM of E. coli-expressed AMSH proteins in a TAMRA DUB Buffer (50 mM Tris-HCl pH 7.5, 100 mM NaCl, 0.1% Pluronic F-127, 1 mM Tris (2-carboxyethyl) phosphine (TCEP)) and the fluorescence intensity was measured every minute for 120 minutes.



Protein localization in N. benthamiana

RFP:AMSH3 wild-type and mutant versions were co-localized with GFP:SKD1^{E232Q} in leaves of 4-5-week-old N. benthamiana, infiltrated with A. tumefaciens harboring the desired constructs. Two days later, leaf discs were punched using cork-borer size 4, and after water mounting, imaged on the abaxial side with an inverted Leica SP-5 confocal laser-scanning microscope. GFP was excited with the 488 nm argon-laser and fluorescence emission collected between 500 and 550 nm. RFP was excited at 543 nm and fluorescence emission collected between 640 and 645 nm. To visualize GFP and RFP simultaneously, Sequential Scan Mode was used. The confocal images were edited in Leica AF Lite (Version 2.6.3 build 8173) and Adobe Photoshop CS5.1 (64 Bit).

Structural homology modeling

The AMSH3 protein sequence was submitted to the online server SWISS-MODEL (Biasini et al., 2014). The best model with a QMEAN score of -2.00 and 51.83% identity was based on the crystal structure of Schizosaccharomyces pombe AMSH-like protease sst2 (4nql.1.A). The model encompassing AMSH3³¹⁸⁻⁵⁰⁷ was visualized in Swiss-PDBViewer V4.1.0 (https://www.expasy.org/spdv/).

QUANTIFICATION AND STATISTICAL ANALYSIS

The numbers of biological repeats (n) are provided in each figure legend. For transcript analyses, each biological repeat is based on an RNA sample extracted from four plants. For plant weight, repeats are individual plants. For bacterial growth, each repeat is based on four leaf discs from one plant. For ion leakage, each repeat is based on five leaf discs. Data are presented as means ± standard error (SE). We performed one-way analysis of variance (ANOVA) with a Tukey HSD post hoc test using R (http://www.R-project.org/). We employed an alpha of 0.05 as cut-off for significance.