The vacuolar ATPase is required for physiological as well as pathological activation of the Notch receptor

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SUMMARY

Evidence indicates that endosomal entry promotes signaling by the Notch receptor, but the mechanisms involved are not clear. In a search for factors that regulate Notch activation in endosomes, we isolated mutants in Drosophila genes that encode subunits of the vacuolar ATPase (V-ATPase) proton pump. Cells lacking V-ATPase function display impaired acidification of the endosomal compartment and a correlated failure to degrade endocytic cargoes. V-ATPase mutant cells internalize Notch and accumulate it in the lysosome, but surprisingly also show a substantial loss of both physiological and ectopic Notch activation in endosomes. V-ATPase activity is required in signal-receiving cells for Notch signaling downstream of ligand activation but upstream of γ-secretase-dependent S3 cleavage. These data indicate that V-ATPase, probably via acidification of early endosomes, promotes not only the degradation of Notch in the lysosome but also the activation of Notch signaling in endosomes. The results also suggest that the ionic properties of the endosomal lumen might regulate Notch cleavage, providing a rationale for physiological as well as pathological endocytic control of Notch activity.

KEY WORDS: Drosophila, V-ATPase, Notch signaling, Endocytosis, ESCRT

INTRODUCTION

Cell-cell signaling via the Notch receptor is used throughout development to regulate multiple cell behaviors, and inappropriate activation of Notch is emerging as a common hallmark of an increasing number of cancers (reviewed by Schweigsguth, 2004; Miele et al., 2006; Roy et al., 2007). Thus, resolving the mechanisms by which Notch signaling is regulated is of great importance and of widespread interest in order to understand human development as well as to devise effective anticancer therapies. In response to ligand engagement, the Notch receptor is activated by S3 cleavage, a γ-secretase-mediated intramembrane proteolysis that liberates the Notch intracellular domain from its transmembrane anchor, allowing the soluble form to travel to the nucleus, where it regulates the transcription of a variety of important target genes (Schweisguth, 2004).

Mounting evidence, particularly in Drosophila, has pointed to the unexpected involvement of the endosomal system in regulating activation of the Notch receptor in signal-receiving cells. Importantly, endosomal regulators can either increase or decrease Notch signaling depending on the specific site of the endocytic pathway at which they act. For example, factors that positively regulate traffic from the cell surface to endosomes, including Dynamin (Shibire – FlyBase), the ubiquitin ligase Deltex, the syntaxin Avalanche (Avl; Syntaxin 7), the GTPase Rab5, the Rab5 effector Rabenosyn-5, and the Sec1/Munc18 family protein Vps45, are required to promote signaling. By contrast, Endosomal sorting required for transport (ESCRT) complex components, the C2-domain protein Lethal (2) giant discs 1 (Lgd), and the ubiquitin ligase Su(dx), which subsequently sort cargo within the endosome towards lysosomal degradation, are required to prevent excess signaling (reviewed by Brou, 2009; Fortini and Bilder, 2009; Fürthauer and González-Gaitán, 2009; Tien et al., 2009). Much of this evidence points to early endosomes as important sites of signaling activation. However, the molecules and mechanisms that restrict activation of the Notch receptor to this endosomal compartment remain unknown.

MATERIALS AND METHODS

Genetics

Mosaic eyes were generated as described previously (Tapon et al., 2001). Completely mutant eye discs (referred to in the text as mutant discs) were generated as described (Newsome et al., 2000). Follicle cell clones were generated as described previously (Bilder et al., 2000). Other alleles utilized were: Vha53e9, VhaSFDEf00464 (Bloomington Stock Center #15758), vps22e9, VhaSFDEf00464 (Vaccari et al., 2009), HrsDEf00464 (Lloyd et al., 2002), EtsplmB-lacZ (Nellesen et al., 1999) and UAS GFP-Lamp-1 (a gift of T. E. Rusten, Norwegian Radium Hospital, Oslo, Norway). Recombined chromosomes were generated using standard genetic techniques. The lesion in Vha68-2 was identified by sequencing PCR products amplified from genomic DNA; at least two independent reactions were sequenced for each allele. Detailed genotypes for each experiment are available upon request; the primers used to sequence Vha68-2 are listed in Table S1 in the supplementary material.

Live disc culture assays

Notch internalization assays were performed as described (Vaccari et al., 2008). For Lysotracker incorporation, mosaic eye discs or ovaries were dissected in Drosophila cell medium (M3, Sigma) and incubated in medium containing 1 μM Lysotracker (DND-99, Molecular Probes) for 5 minutes at room temperature, washed in fresh medium, mounted and imaged. For Dextran-uptake assays, mosaic eye discs dissected in M3 medium were incubated for 10 or 40 minutes in Texas Red-Dextran MW 3000, lysine-fixable (Molecular Probes), washed, fixed and counterstained using an antibody to Notch ICD, mounted and imaged.

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Immunohistochemistry and optical microscopy
Immunohistochemical analyses were performed as described (Vaccari et al., 2009). Actin stains used TRITC-conjugated phalloidin (Sigma). Primary antibodies to the following antigens were used: Avl (Lu and Bilder, 2005), Hrs (Lloyd et al., 2002), ubiquitin (FK2; Biomol), Caspase-3 (Decay – FlyBase) (Cell Signaling Technologies), β-gal (Cappell) and Notch ECD, Notch ICD, Mmp1, Hnt, Cut, Elav and Dlg (DSHB). Alexa 488- or Alexa 647-conjugated secondary antibodies (Molecular Probes) were used. All images shown are single confocal sections taken with a TCS microscope (Leica) using 16×/NA 0.5, 40×/NA 1.25 or 63×/NA 1.4 oil-immersion lenses. Images were edited with Adobe Photoshop CS and were assembled with Adobe Illustrator. Quantification was performed using ImageJ (NIH). The data presented are the average of measurements from eight confocal sections for each genotype (Fig. 1H), of two discs for each genotype (Fig. 3L), and of four discs for each genotype (Fig. 4M).

Electron microscopy
Wild-type (WT), Vha55 and Vha68-2 completely mutant eye discs were placed in 0.1 M cacodylate buffer containing 2.5% glutaraldehyde for 3 hours at room temperature. Samples were post-fixed in osmium tetroxide (Electron Microscopy Science, Hatfield, PA, USA) for 2 hours and in uranyl acetate (Electron Microscopy Science) for 1 hour at room temperature. Samples were then dehydrated through a graded ethanol series with propylene oxide as a transition fluid (TAAB Laboratories Equipment, Aldermaston, UK) and embedded in epoxy resin (Poly-Bed; Polysciences, Warrington, PA, USA) overnight at 42°C then for 2 days at 60°C. Subsequent examination of the region of interest (the eye disc epithelium) was performed on semi-thin sections (500 nm) stained with Toluidine Blue. Ultra-thin sections (50 nm) were then cut and stained with uranyl acetate and lead citrate and images were taken with a CM10 (Philips, Eindhoven, The Netherlands) or Tecnai 12-G2 (FEI, Eindhoven, The Netherlands) electron microscope.

Quantification of the multivesicular body (MVB) compartment within ultra-thin sections was performed using iTEM software (FEI) (Fig. 1L). We determined the density of MVBs as the total number of MVBs in 187, 195 and 186 μm² of WT, Vha68-2 and Vha55 mutant tissue, respectively, and normalized each to 1000 μm². We measured MVB diameter as the length of the major axis, measuring only those completely enclosed in the micrograph.

RESULTS
V-ATPase function controls Notch protein levels, endosomal acidification and overall tissue size and morphology of Drosophila epithelial organs
To identify genes involved in endocytic control of the Notch receptor, we screened a collection of Drosophila mutants enriched for endocytic regulators (Menut et al., 2007) to identify those that display altered Notch localization. We generated eye imaginal discs consisting predominantly of mutant cells (see Materials and methods) and immunostained using Notch antibodies. We found that discs mutant for the single allele complementation group MENE2L-R6 accumulate high levels of Notch as compared with wild-type (WT) discs (Fig. 1A,B). Clones of MENE2L-R6 mutant cells generated in the context of a mosaic eye disc also displayed substantial accumulation of Notch, which was found in large intracellular puncta (Fig. 1C). MENE2L-R6 discs were consistently smaller than WT discs and also displayed aberrant morphology, compromised epithelial polarity, and upregulated Matrix metalloprotease 1 (Mmp1) production (Fig. 1A,B; see Fig. S1A-B’ in the supplementary material). Together, these data suggest that R6 disrupts a gene that regulates Notch trafficking as well as the proper growth and organization of imaginal disc tissue.

We mapped the R6 mutant by complementation to a deficiency that removes the chromosomal region 34A3. Tests with lethal transposon insertions within the deficiency region revealed that R6 fails to complement P[PZ]/h20150, which is inserted in the first intron of the predicted gene CG3762 (Fig. 1D). CG3762 is also known as Vha68-2, the most widely expressed of the three Drosophila genes that encode the A subunit of the multiprotein vacuolar ATPase (V-ATPase) proton pump (Allan et al., 2005) (Fig. 1E). The A and B subunits form a heteromer that binds nucleotide in the peripheral V1 sector; nucleotide binding and hydrolysis are essential for transmembrane proton conductance through the integral membrane V0 sector (reviewed by Nishi and Forcaga, 2002) (Fig. 1E). Sequestration of Vha68-2 in R6 mutants revealed a nonsense mutation in the region encoding the C-terminal domain. Based on the crystal structure of the related A-ATPase subunit A (Maegawa et al., 2006), the truncated C-terminal domain of the R6 mutant protein, even if expressed, is likely to lack part of the region that contacts subunits D and F, which are involved in torque generation (Fig. 1D,E). Since torque generation is essential for proton transport and thus endomembrane acidification, we tested the ability of Vha68-2 mutant cells to incorporate the vital dye Lysotracker, which accumulates in acidified lysosomes. Compared with WT cells or cells mutant for the endosomal component Hrs, we found that Vha68-2 mutant cells incorporate very low levels of Lysotracker, indicating impairment of the ability to acidify endocytotic organelles (Fig. 1F-H).

Failure of Lysotracker incorporation into Vha68-2 mutant cells is not due to absent endocytic structures, as staining with endosomal markers and electron microscopy revealed that both early and late endosomes, as well as lysosomes, are present in mutant cells (Fig. 1I,J,L; Fig. 2E-G). In addition, ultrastructural analysis of the morphology of the endocytic compartment revealed that Vha68-2 mutant cells contain, compared with WT, more multivesicular bodies (MVBs), a degradative endosomal organelle. These MVBs were also enlarged and contained a high number of internal luminal vesicles (ILVs) (Fig. 1I,J,L). Consistent with a role for acidification in activating low-pH lipid hydrolases (Soyombo et al., 2006), these data suggest that Vha68-2 is required for lysosomal degradation of ILVs. Overall, the Lysotracker incorporation and the ultrastructural analyses indicate that Vha68-2 causes a substantial loss of proton transport activity.

To confirm that the Notch trafficking, growth and architecture phenotypes observed in Vha68-2 imaginal disc cells are due to a lack of V-ATPase activity, we analyzed the phenotypes of mutations that disrupt other V-ATPase subunits. Vha55j2e9 is a transposon insertion that creates a strong mutant in the single Drosophila gene encoding the B subunit of V-ATPase (Davies et al., 1996) (Fig. 1E). We found that, similar to Vha68-2 discs, Vha55j2e9 mutant discs display Notch accumulation, Mmp1 expression, small size and aberrant morphology; they also showed ultrastructural defects similar to, although less strong than, those of Vha68-2 (Fig. 1K,L; see Fig. S1C and Fig. S2C,D in the supplementary material). In addition, we found that eye discs homozygous for EY04644, a transposon insertion in the 5’UTR of VhaSFD, the single Drosophila gene encoding V-ATPase subunit H (Allan et al., 2005), phosphoryl Vha68-2 and Vha55 mutant discs (see Fig. S2E,F in the supplementary material). Moreover, a portion of the cells mutant for any of the three genes is eliminated from the disc, especially when surrounded by WT cells (see Fig. S3 in the supplementary material). These data strongly suggest that the common phenotypes observed in Vha55, Vha68-2 and VhaSFD mutant cells are caused by impaired V-ATPase function. Taken together, they indicate that the V-ATPase controls endosomal acidification and is also a major regulator of Notch localization, epithelial organization and growth in imaginal disc cells.
Lysosomal degradation, but not endocytic trafficking to late endosomes/lysosomes, is impaired in cells lacking acidic compartments

The high levels of intracellular Notch seen in V-ATPase mutant cells could in theory result from impaired exocytosis or endocytosis. To test whether V-ATPase controls endocytic transport in imaginal cells, we first followed the uptake of Dextran over time in mosaic eye discs containing clones of Vha68-2 mutant cells that were cultured ex vivo. Compared with WT cells, internalization of Dextran was reduced at the 10-minute time point in Vha68-2 mutant cells, but after 40 minutes Vha68-2 cells accumulated similar amounts of Dextran to WT cells (Fig. 2A,B). These data indicate that although its rate may be reduced, endocytic trafficking is still functional upon impairment of Vha68-2 expression. To directly test endocytic trafficking of Notch, we used an ex vivo assay for Notch internalization and found that, compared with WT cells, Notch is internalized but fails to be degraded in Vha68-2 mutant cells (Fig. 2C). In mutant cells, Notch accumulated in large and distinct intracellular puncta (Fig. 2C/H11032). These intracellular puncta resemble those seen in discs treated with chloroquine, a compound that prevents lysosomal acidification (Vaccari and Bilder, 2005).

To determine the specific site of endocytic Notch accumulation in Vha68-2 mutant cells, we colabeled eye disc cells for Notch and Avl or Hrs, two early endosomal markers, or GFP-Lamp-1 (CG3305), a late endosomal/lysosomal marker. We found that in Vha68-2 mutant cells, Notch accumulated in large intracellular puncta, which are similar to those seen in discs treated with chloroquine. These data indicate that although endocytic trafficking is still functional upon impairment of Vha68-2 expression, the accumulation of Notch in intracellular puncta suggests that Notch trafficking is impaired in these cells.
V-ATPase activity is required for Notch activation in signal-receiving cells prior to \(\gamma\)-secretase-dependent S3 cleavage.

Since V-ATPase mutant cells possess high levels of undegraded Notch, we asked whether this phenotype affects Notch signaling. Because other mutants that trap Notch in late endosomes, such as \(f\text{ab}1\), do not alter Notch signaling (Rusten et al., 2006; Vaccari et al., 2008), we expected Notch activity to be unaffected in cells lacking V-ATPase activity. Surprisingly, we found that expression of the Notch signaling reporter \(\text{mb}l\)-\(\text{lacZ}\) is significantly reduced in \(\text{Vha}68-2\) mutant disc cells, as compared with the stereotyped levels and patterns seen in WT discs (Fig. 3A,B) (Nellesen et al., 1999). Interestingly, the degree of reduction in reporter expression paralleled the degree of Notch accumulation in mutant cells (Fig. 3B''). Similarly, compared with WT, \(\text{mb}l\)-\(\text{lacZ}\) expression was absent in most \(\text{Vha}68-2\) mutant eye disc cells (Fig. 3C). As also observed for Notch accumulation (Fig. 1C), this phenotype was variably penetrant, with some cells showing mild phenotypes. This could reflect a functional redundancy of \(\text{Vha}68-2\) with other \(\text{Vha}68\) genes or hypomorphy of the allele. Despite the phenotypic variability, these results suggest that V-ATPase activity is in fact positively required for Notch signaling.

Lack of Notch signaling in V-ATPase mutant tissue could be due to effects on Notch ligands or the Notch receptor. To discriminate between these possibilities, we utilized \textit{Drosophila} ovaries. We first asked whether V-ATPase mutant somatic follicle cells (FCs) show the acidification and Notch degradation defect observed in V-ATPase mutant eye disc cells. We made genetic mosaic ovaries in which some FCs were mutant for \(\text{Vha}68-2\). Similarly to mosaic eye discs, we found that, compared with neighboring WT cells, mutant FCs failed to incorporate Lysotracker and accumulated high amounts of Notch intracellularly (see Fig. S4A,B in the supplementary material).

In ovaries, the Notch ligand Delta is upregulated in the germ cells during stages 6-7 of oogenesis to activate Notch signaling in the adjacent FCs (Deng et al., 2001; Lopez-Schier and St Johnston, 2001), leading to downregulation of target genes such as \(\text{cut}\) and to activation of target genes such as \(\text{hind}5\text{ight}\) (\(\text{hht}\); \textit{pebbled} – FlyBase) (Fig. 3D,F) (Sun and Deng, 2005; Sun and Deng, 2007). Thus, to assay the extent of Notch signaling activation in \(\text{Vha}68-2\) mutant FCs, we immunostained mosaic ovaries in which some FCs were mutant for \(\text{Vha}68-2\). Similarly to mosaic eye discs, we found that, compared with neighboring WT cells, mutant FCs failed to incorporate Lysotracker and accumulated high amounts of Notch intracellularly (see Fig. S4A,B in the supplementary material).

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in the supplementary material). Consistent with an impairment in Notch signaling activation in V-ATPase mutant FCs, we also found a substantial reduction in Hnt expression in Vha55 mutant FCs (Fig. 3G; see Fig. S4D in the supplementary material). Taken together, these data indicate that V-ATPase activity is required cell-autonomously in signal-receiving cells for activation of the Notch receptor.

Activation of the Notch receptor following ligand binding requires a proteolytic cleavage called S3, which is mediated by γ-secretase (reviewed by Schweisguth, 2004). To test whether V-ATPase function is required for this cleavage event, we compared WT and mutant cells expressing either of two activated signaling forms of Notch. Notch extracellular truncation (NEXT) is a membrane-tethered form of Notch that is a constitutive substrate for S3 cleavage by γ-secretase in WT cells; Notch intracellular domain (NICD) represents the soluble cleavage product of this reaction (Fig. 3H). Clones of WT imaginal disc cells expressing NEXT overproliferate due to unrestrained Notch signaling activation (Fig. 3I). However, Vha55 mutant cells expressing NEXT did not overproliferate and instead created only small clones, of a size similar to Vha55 clones (Fig. 3J, compare with Fig. 4E), indicating that the production of an active signaling form of Notch is blocked in these cells. Finally, Vha55 mutant cells expressing NICD overproliferated similarly to WT clones expressing either transgene, revealing that the requirement for V-ATPase activity in Notch signaling can be bypassed by providing a cleaved form (Fig. 3K). These data show that V-ATPase activity is required downstream of ligand binding, but upstream of the S3 activating cleavage.

Acidification of the early endosomal lumen is a prerequisite for efficient activation of Notch signaling

Although the above data suggest a role for V-ATPase activity in Notch cleavage, they do not specify in which organelle this activity might be required, as V-ATPases acidify most compartments of the

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**Fig. 3. V-ATPase activity is required to activate the Notch receptor downstream of ligand binding in epithelial tissue.** (A-C) Drosophila eye discs stained to detect the Notch reporter mβ-lacZ and Notch (single mβ-lacZ channel is shown in A’-C’). Compared with WT (A), mβ-lacZ expression is reduced in predominantly mutant Vha68-2R6 discs (B) and in clones of Vha68-2R6 mutant cells located posterior to the morphogenetic furrow in mosaic discs (C). Mutant tissue is outlined in B’, C’. The regions of the tissue that accumulate Notch the most appear to express mβ-lacZ the least (compare B with B’). In C, mβ-lacZ is expressed at high level in WT developing photoreceptors but is absent in clones of mutant cells. (D-G) Egg chambers at stages 5-7 of oogenesis stained for Cut (D,E) and Hnt (F,G). In WT cells, Cut expression is subjected to Notch-dependent downregulation after stage 5 (D), whereas Hnt is expressed in a Notch-dependent manner from stage 6 onwards (F). In Vha68-2 mutant FCs, Cut expression is maintained after stage 5 (E; for GFP channel see Fig. S4C in the supplementary material), whereas in Vha55 mutant FCs, Hnt expression is reduced after stage 5 (G; for GFP channel see Fig. S4D in the supplementary material), in both cases indicating impairment in Notch signaling activation in V-ATPase mutant FCs. (H) Schematic of the Notch cleavage events preceding signaling activation. The full-length Notch is cleaved at the S2 site to generate the γ-secretase cleavage substrate NEXT. NEXT is subsequently cleaved by γ-secretase at the S3 site to generate the active cytoplasmic NICD fragment. (I-K) Mosaic eye discs generated using the MARCM system stained to detect NICD. GFP-positive cells expressing NEXT overproliferate to form very large clones (I). By contrast, GFP-positive cells expressing NEXT that lack V-ATPase activity underproliferate and form small clones (J). In GFP-positive cells that express NICD and lack V-ATPase activity, normal proliferation is restored (K). (L) Quantification of the experiment shown in I-K. Scale bars: 100 μm in A-B’, I-K; 10 μm in C-G.
endocytic pathway. Previous work has shown that efficient Notch activation following ligand binding requires entry of Notch into early endosomes, whereas a failure to sort and transport Notch out of the Hrs-positive early endosomal compartment, such as in ESCRT or lgl mutant cells, causes ectopic Notch activation (reviewed by Brou, 2009; Fortini and Bilder, 2009; Fürthauer and González-Gaitán, 2009; Tien et al., 2009). To determine whether V-ATPase activity is required in the early endosome for Notch activation, we generated chromosomes that were double mutant for Vha55 and the ESCRT-III component vps22 (larsen – FlyBase). Similar to vps22 mutant cells, vps22 Vha55 double-mutant cells accumulated ubiquitin and showed extensive colocalization of Notch with Hrs, indicating that V-ATPase activity is not required for Notch to access the early endosomal compartment (Fig. 4A–F). However, in contrast to vps22 mutant cells, expression of mβ-lacZ was significantly reduced in vps22 Vha55 double-mutant cells, to a level that was similar to that of Vha55 mutant cells (Fig. 4A–C,G–I). These data indicate that although Notch is trapped in early endosomes in vps22 Vha55 mutant cells, as in cells mutant for ESCRT components alone, it is nevertheless unable to signal. Consistent with a suppression of ectopic Notch signaling, in contrast to vps22 mutant cells, vps22 Vha55 mutant cells in mosaic eyes were unable to overproliferate (Fig. 4J–L). Thus, whereas ESCRT phenotypes are epistatic to V-ATPase with respect to Notch trafficking, V-ATPase phenotypes are epistatic to ESCRT with respect to Notch signaling. These results indicate that V-ATPase function is required for Notch activation in early endosomes.

**DISCUSSION**

Recent evidence suggests that the route and rate of Notch traffic through endocytic compartments can be regulated to either potentiate or diminish signaling activity. Our results identify a physiological mechanism to account for this effect on signaling. We identify a set of genes required for endosomal Notch signaling that encode subunits of the V-ATPase, the molecular machine that creates a proton motive force to acidify most compartments of the endosomal system. The data thus establish a novel role of V-ATPase that is relevant to both physiological and pathological Notch signaling. While this work was under review, an article describing a similar role for V-ATPase regulators and a different subunit of the V-ATPase in Notch signaling was published (Yan et al., 2009). Our independent results confirm those findings and expand them by reporting the ultrastructure of V-ATPase mutant cells, extending the role to tumor contexts that depend on excess Notch signaling activity and, importantly, providing evidence for the early endosome as the site of the requirement for V-ATPase activity. Interestingly, the aquaporin Big brain (Bib) was recently shown to promote Notch signaling at a similar step of Notch processing, and bib mutant clones, like V-ATPase mutant clones, show defects in endosomal acidification (Kanwar and Fortini, 2008). It is possible that some other, as yet uncharacterized, V-ATPase function unrelated to its well-established role in proton pumping might be required for Notch signaling activation. Alternatively, the common acidification and Notch signaling phenotypes of V-ATPase and bib mutant cells might indicate that Notch signaling normally requires endosomal acidification.

The best-understood requirement for V-ATPase-dependent acidification is in promoting lysosomal degradation by ensuring the proper targeting, release and activation of lysosomal proteases. Indeed, we find that Notch and other cargoes are trapped in a lysosome-like compartment in V-ATPase-deficient cells, where they accumulate rather than being degraded. The cargo-trapping phenotype resembles that seen in the presence of chemical inhibitors of lysosomal acidification, such as chloroquine, as well as in cells mutant for regulators of lysosomal genesis such as the HOPS components Dor and Car and the PIKfyve Fab1.
(Sevrioukov et al., 1999; Rusten et al., 2006; Akbar et al., 2009). However, in striking contrast to these latter mutants, in which Notch signaling is largely unaffected, a substantial loss of Notch signaling is seen in V-ATPase-deficient cells. This indicates that the requirement for V-ATPase in Notch signaling must precede lysosomal entry.

How could acidification of earlier endosomal compartments promote Notch signaling? One possibility is that the role of acidification reflects the requirement of endosomal transport for Notch activation. In mammalian cells, V-ATPases have been implicated in the recruitment of proteins that modulate traffic between early and late endosomes (Hurtado-Lorenzo et al., 2006). We find a reduced rate of endocytosis in V-ATPase mutant cells, and it is possible that a dampened flux of Notch through the endocytic pathway contributes to reduced Notch signaling. However, we believe this role is unlikely to account for the V-ATPase mutant phenotype for the following reasons. First, although its endocytic rate may be reduced, Notch can clearly reach both early and late endosomal compartments in mutant cells; this mild quantitative reduction in traffic contrasts with the more substantial and qualitative loss of Notch signaling. Second, in mammalian cells, loss of endosomal acidification reduces progression between early and late endosomes. Since the requirement for Notch signaling is in entry into early endosomes, Notch activation would seem to be less sensitive to this step. Moreover, because Notch reaches late endosomal compartments in V-ATPase mutant cells, in Drosophila, at least, redundant mechanisms to energize endocytic traffic must exist. Third, the striking failure of Notch trapped within Hrs-positive early endosomes in V-ATPase ESCRT double mutants to signal suggests that even when endocytic traffic is blocked at the endosomal sorting step, Notch activation still requires V-ATPase activity. The absence of Notch activation in vps22 Vha55 cells is particularly striking because even when most Notch cannot reach endosomes in cells double mutant for the ESCRT component TSG101 (erupted) and the endosomal syntaxin avl (Syx7), the minor fraction of Notch that does reach endosomes is efficiently cleaved and strongly activates ectopic Notch signaling (Vacari et al., 2008). This suggests that the role of V-ATPase in Notch activation goes well beyond trafficking. The data also reveal that it is not mere access to the early endosome that is required for Notch signaling, but rather that a specific physiological feature of that endosome is central to the signaling activation mechanism.

An alternative hypothesis is that V-ATPase-dependent acidification creates an endosomal environment that is conducive for productive Notch S3 cleavage and, therefore, signaling. In this scenario, Notch transits through the early endosome in V-ATPase mutant cells, but is not efficiently activated because of altered γ-secretase function. Such a role is consistent with our genetic experiments, which show that a membrane-tethered Notch truncation that is automatically processed to an active form by γ-secretase in WT cells cannot signal in cells that lack V-ATPase function. Although it is appealing to speculate that an acidic pH could promote overall cleavage by γ-secretase, which requires low luminal pH for optimal activity (Pasternak et al., 2003), experiments testing the acid dependence of other γ-secretase substrates have failed to show a reduction in overall S3-type processing (Vingtedoux et al., 2007). Although these assays are limited in their ability to reflect the in vivo situation, an interesting possibility suggested by mammalian studies is that pH might alter the precision of the S3 cleavage site, producing NICD forms that are quickly degraded and cannot effectively signal (Tagami et al., 2008). Additional possibilities exist, such as that rather than creating a cleavage-promoting environment per se, the V-ATPase could serve as a pH sensor that couples the generation of a low pH to the recruitment of γ-secretase regulators in order to restrict cleavage to an endosomal site. Finally, acidification could directly affect the correct maturation and localization of the γ-secretase enzyme. Future work will distinguish between these possibilities.

Regardless of the specific mechanism governing the role of V-ATPase in Notch cleavage, overall our data suggest that drugs that impair V-ATPase function and consequently reduce endosomal acidification might be used to curtail pathologic overactivation of Notch, such as that observed in cancers characterized by Notch overexpression (Roy et al., 2007). Considering that such a goal is presently being pursued by the use of γ-secretase inhibitors (Rasul et al., 2009; Tanaka et al., 2009), the use of already established V-ATPase inhibitors, such as Bafilomycin A1, either alone or in combination with γ-secretase inhibitors, could represent a promising therapeutic avenue.

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Competing interests statement
The authors declare no competing financial interests.

Supplementary material
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References


Table S1. Primers used to sequence Vha68-2

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