The V0-ATPase mediates apical secretion of exosomes containing Hedgehog-related proteins in Caenorhabditis elegans

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Polarized intracellular trafficking in epithelia is critical in development, immunity, and physiology to deliver morphogens, defensins, or ion pumps to the appropriate membrane domain. The mechanisms that control apical trafficking remain poorly defined. Using Caenorhabditis elegans, we characterize a novel apical secretion pathway involving multivesicular bodies and the release of exosomes at the apical plasma membrane. By means of two different genetic approaches, we show that the membrane-bound V0 sector of the vacuolar H+-ATPase (V-ATPase) acts in this pathway, independent of its contribution to the V-ATPase proton pump activity. Specifically, we identified mutations in the V0 “α” subunit VHA-5 that affect either the V0-specific function or the V0+V1 function of the V-ATPase. These mutations allowed us to establish that the V0 sector mediates secretion of Hedgehog-related proteins. Our data raise the possibility that the V0 sector mediates exosome and morphogen release in mammals.

Introduction

All developmental and physiological functions performed by epithelia depend on the polarized targeting of the plasma membrane and secreted proteins to either the apical or basolateral plasma membranes (Rodriguez-Boulan et al., 2005). Cargo proteins sorted in the Golgi apparatus and the endosomal system through sets of basolateral- and apical-specific sorting determinants are transported to the plasma membrane following partially different routes (Hoekstra et al., 2004; Rodriguez-Boulan et al., 2005). Although basolateral secretion has been fairly well characterized, the mechanisms involved in apical trafficking remain poorly defined (Rodriguez-Boulan et al., 2005).

Basolateral sorting signals usually correspond to tyrosine or dileucine residues found in the COOH terminus of proteins. They are recognized by basolateral-specific adaptor complexes (Bonifacino and Lippincott-Schwartz, 2003; Rodriguez-Boulan et al., 2005), such as AP-1B in epithelial cells (Folsch et al., 1999). Before membrane fusion and SNARE action, vesicles are thought to be tethered to the basolateral membrane by the exocyst complex (Whyte and Munro, 2002), which was initially identified in yeast (TerBush and Novick, 1995). In metazoans, the exocyst is required for basolateral delivery of the LDL receptor in MDCK cells (Grindstaff et al., 1998; Yeaman et al., 2001), of E-cadherin in the Drosophila melanogaster notum (Langevin et al., 2005), and for Rhodopsin1 transport in D. melanogaster photoreceptor cells (Beronja et al., 2005). Recent results suggest that AP-1B and the exocyst act primarily in recycling endosomes (Ang et al., 2004; Beronja et al., 2005; Langevin et al., 2005; Lock and Stow, 2005; Satoh et al., 2005), which underlines the central role of this organelle in sorting processes. Indeed, recycling endosomes may be compartmentalized into apical- and basolateral-related domains, or even divided into distinct organelles, suggesting that they could also play a critical role in apical trafficking (Hoekstra et al., 2004; Rodriguez-Boulan et al., 2005).

Aside from this possible role of recycling endosomes, all other aspects of sorting along the basolateral and apical routes seem to differ. Apical signals are more diverse and often correspond to posttranslational adducts, such as lipids or glycosyl ceramide (Schuck and Simons, 2004; Rodriguez-Boulan et al., 2005). For instance, the Hedgehog morphogen is secreted apically upon
cholesterol addition, but basolaterally otherwise (Gallet et al., 2003). No specific apical cytolsolic complex, akin to AP-1B or the exocyst, has been identified so far. Instead, protein clustering, possibly through lipid rafts, is thought to mediate the sorting and transport of apical cargoes (Schuck and Simons, 2004; Rodriguez-Boulan et al., 2005). In particular, glycosyl phosphatidylinositol–linked proteins appear to form oligomers that are directly targeted to the apical membrane (Paladino et al., 2004, 2006; Hua et al., 2006). Several proteins have been proposed to play an active role in apical protein clustering, raft formation, and/or apical delivery, such as caveolins (Kurzchalia et al., 1992), annexin 13b (Fiedler et al., 1995), and the tetraspan protein VIP17/MAL (Cheong et al., 1999; Puertollano et al., 1999). However, their mechanistic roles have not been fully elucidated, or their implication has been questioned (Manninen et al., 2005). In addition to the limited understanding of apical secretion at the molecular level, it is not clear whether the terminal fusion process involves small vesicles, such as those defined at synapses, or larger organelles, such as secretary lysosomes (Blott and Griffiths, 2002).

Hence, despite the many critical findings originating from tissue culture cells (Rodriguez-Boulan et al., 2005), investigations with other systems and other cargo proteins could help to elucidate the mechanisms involved in apical exocytosis. Caenorhabditis elegans, which has contributed to decipher the mechanisms controlling vesicular trafficking (Nurrish, 2002), provides such an in vivo model. We have chosen to analyze apical secretion of cuticle proteins by the epidermis. The cuticle includes glycosylated collagens, glycosyl phosphatidylinositol–linked cuticins, and lipid-modified Hedgehog-related peptides (McMahon et al., 2003; Sapiro et al., 2005; Zugasti et al., 2005). We previously suggested that the gene che-14 is required for cuticle secretion (Michaux et al., 2000). The CHE-14 protein is the C. elegans orthologue of Drosophila Dispatched, which participates in apical targeting of cholesterol-modified Hedgehog (Burke et al., 1999; Gallet et al., 2003).

While searching for che-14 alleles (Michaux et al., 2000), we uncovered several additional mutations inducing che-14–like phenotypes and reasoned that they might identify new components of the apical trafficking pathway. Two such mutations, mc37 and mc38, proved to be small deletions behaving as genetic-null alleles of the gene vha-5 (unpublished data). The gene vha-5 encodes one of the four C. elegans “a” subunits of the V0 sector of the vacuolar H⁺-ATPase (V-ATPase), and is required for development beyond the L2 larval stage (Oka et al., 2001; Pujol et al., 2001). The V-ATPase is a multisubunit protein complex consisting of two subcomplexes called the V0 and V1 sectors (Fig. 1 A). The cytosolic V1 sector hydrolyses ATP and provides the energy to pump protons through the transmembrane proteolipid pore formed by the V0 sector (Nishi and Forgac, 2002). The V-ATPase is the main proton pump establishing a pH gradient in the secretory and endocytic pathways. It generates a proton-motive force that is essential to load synaptic vesicles with neurotransmitters before secretion (Amara and Kuhar, 1993). The V-ATPase is also found at the apical plasma membrane of polarized cells, where it is essential for osmoregulation in animal excretory systems (Nishi and Forgac, 2002). More recently, biochemical and genetic data suggested that the V0 sector can play a role independently from the V1 sector. In Saccharomyces cerevisiae, vacuoles deficient for the “a” subunit Vph1p do not fuse efficiently (Peters et al., 2001; Bayer et al., 2003). In D. melanogaster, neurons lacking the “a” subunit Vha100 accumulate vesicles in synaptic terminals (Hiesinger et al., 2005). In both cases, the defects were independent of the proton gradient and placed downstream of SNARE function (Peters et al., 2001; Bayer et al., 2003; Hiesinger et al., 2005).

By further dissecting the role of vha-5 using targeted mutagenesis, and by comparing phenotypes resulting from the inactivation of V1 or V0 subunits, we uncover a specific role for the V0 sector in mediating secretion to the apical membrane. In particular, we show that the V0 sector is required for apical secretion of Hedgehog-related peptides through a multivesicular compartment able to release exosomes.

Results

The V0 “a” subunit VHA-5 is apical and required for cuticle formation

To determine the distribution and subcellular localization of VHA-5, we raised polyclonal antibodies against its cytoplasmic NH₂ terminus. In addition, we generated a COOH-terminal VHA-5::GFP fusion, which rescued the larval lethality caused by the vha-5 (mc38) deletion (Fig. 1 B). The VHA-5 antiserum recognized a 105-kD protein in wild-type extracts (Fig. 1 C, lane a). To prove its specificity, we examined extracts from vha-5 (mc38) homozygous animals carrying the rescuing VHA-5::GFP construct. The VHA-5 antiserum failed to detect the ~105-kD band in these extracts, but detected an ~135-kD band (Fig. 1 C, lane b). These results are consistent with vha-5 (mc38) being a small deletion associated with a frameshift (Fig. 1 B and not depicted) and with the presence of 257 additional residues in the GFP-fusion protein. We conclude that the VHA-5 antiserum is specific and that vha-5 (mc38) is a molecular null mutation.

In agreement with previously published observations (Oka et al., 2001; Pujol et al., 2001), we found that VHA-5 is expressed in the H-shaped excretory cell corresponding to the C. elegans kidney-like organ (Fig. 1, D and E). It is also expressed in the main epidermal syncytium (Fig. 1, D–F), which had previously been overlooked. The excretory cell extends long processes called excretory canals where osmoregulation takes place (Nelson and Riddle, 1984), whereas the epidermis controls body length and apical cuticle secretion (White, 1988). VHA-5 colocalized apically with the V1 subunit VHA-8 in both tissues (Fig. 1 E; note that VHA-5 is not expressed in the lateral epidermis). VHA-5 was localized at the level of apical membrane stacks by immunogold staining (Fig. 1 G). Consistent with VHA-5 distribution and a role of the V-ATPase in osmoregulation (Nishi and Forgac, 2002), vha-5 (mc38) larvae filled with fluid and died at the L1 stage (unpublished data), which corresponds to the phenotype observed after laser ablation of the excretory cell (Nelson and Riddle, 1984). In addition, vha-5 (mc38) L1 larvae had a severe malformation of the lateral cuticular specializations known as alae (Fig. 1 D and Fig. 2 A), which are primarily synthesized by the lateral seam cells.
Although VHA-5 is not expressed in these cells, the main epidermal syncytium also contributes to their morphogenesis (Sapio et al., 2005). Because VHA-5 is transmembrane and not cuticular, the simplest interpretation for this phenotype is that vha-5 mutations compromise the secretion of proteins needed for alae formation.

The V0 sector alone is required for cuticle formation

As outlined in the previous section, the V0 sector may fulfill two distinct functions; either working together with the V1 sector to mediate proton pumping or working alone, as in yeast and D. melanogaster neurons, to mediate membrane fusion. To determine which of these functions could account for the cuticle secretion defect observed in vha-5(mc38) larvae, we examined the role of other V-ATPase subunits in cuticle formation using the RNAi approach. If improper proton pumping is responsible for the aforementioned vha-5 cuticle defects, RNAi knockdown of either V0 or V1 subunits should result in similar cuticular defects. Conversely, if the loss of a V0-specific function accounts for the vha-5 cuticular phenotype, only RNAi knockdown of V0 subunits should phenocopy vha-5 cuticle defects. We chose two V1 subunits (VHA-8 and -13) and one V0 subunit (VHA-4) encoded by single-copy genes, which were, thus, expected to be ubiquitously expressed and to display RNAi phenotypes of comparable severity. In addition, we tested RNAi against the three genes encoding the V0 “c” subunit (vha-1, -2, and -3), which are >78% identical at the nucleotide level. We found that the RNAi phenotype of vha-1 was the strongest and was directly comparable to that of vha-4, -8, and -13 (Fig. 2 B’), presumably because it reflects knockdown of all three paralogs. RNAi against these V1 or V0 subunit genes led to 100% lethality in the progeny of treated animals (Fig. 2 B’, bottom bars). It is likely that most embryos died because of a defect in yolk endocytosis, which is known to be sensitive to proton pumping (Choi et al., 2003). In agreement, we found that yolk vitellogenin-GFP accumulated in the pseudocoelom of RNAi-treated animals rather than in oocytes and embryos (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1).

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although lethality was weaker because VHA-5 is not ubiquitously expressed like VHA-4. One trivial explanation for the persistence of normal alae after V1 subunit knockdown could be that RNAi was less efficient than for V0 subunits. It is unlikely, as the lethality rates and the larval osmoregulation defects observed after V1 and V0 subunit knockdown were comparable (Fig. 2, B' and B''), hinting that both RNAi were equally effective. To support this idea, we submitted a \textit{vha-8::gfp} transgenic strain to \textit{vha-8} RNAi and verified that it induced a drastic decrease of VHA-8::GFP fluorescence (Fig. 2 D). We conclude that the V0 sector is required independently from the V1 sector for apical secretion of some cuticle components.

The two functions of V0 are genetically separable

If the V0 sector has two distinct functions, it should be possible to recover \textit{vha-5} alleles that affect either its V0-specific secretion function or its V0+V1 proton-pump function. The \textit{vha-5} distribution and the aforementioned phenotypes indicate that reducing V0-specific function should affect cuticle secretion, whereas impairing proton pumping should affect the excretory canal responsible for osmoregulation. To identify such mutations, we used a plasmid rescue strategy, whereby we generated mutations by using PCR on a rescuing \textit{vha-5::gfp} construct, introducing them into \textit{vha-5(mc38)/+} animals, and recovering live homozygous \textit{vha-5(mc38)} animals whenever possible (Fig. 3 A). We modified charged or large hydrophobic residues, as well as residues previously mutated in the yeast Vph1p (Leng et al., 1996, 1998). We generated 56 mutations (Fig. 3, B and C; and Fig. S2, available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1); 42 had no obvious phenotype by differential interference contrast (DIC) microscopy (Fig. 3 B, stars), and eight failed to rescue, indicating that those residues are essential for VHA-5 function (Fig. 3 B, white boxes). More interestingly, six substitutions rescued the \textit{vha-5(mc38)}-induced lethality and affected the cuticle, the excretory canal, or both.

These six mutations defined three classes, which we will call “cuticle mutations” (L786S, E830Q, and V844F), “canal mutations” (W190A and R191A), or “mixed mutations” (W327A). First, animals carrying cuticle or mixed mutations were significantly shorter and dumpier than wild-type animals or animals carrying canal mutations (Fig. 4 A). Western blot analysis using the VHA-5 antiserum detected similar amounts of mutant VHA-5::GFP proteins (Fig. 4 B), implying that expression level...
differences do not explain phenotypic differences. Second, scanning electron microscopy (SEM) showed that adult alae were strongly affected in the former, but not the latter, animals (Fig. 4 C and Fig. S3 A, available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1). Third, confocal microscopy using the mutant VHA-5::GFP as a marker and transmission electron microscopy (TEM) revealed that the excretory canal of animals with canal or mixed mutations, but not with cuticle mutations, was abnormal (Fig. 4, D and E; and Fig. S3, A and B). Their excretory canals had an increased section, often with multiple lumens, and 3–10 abnormal whorls per canal (see mc38; Ex[W190A] in Fig. 4 [D and E]). Strikingly, we observed similar phenotypes by knocking down V1 or V0 subunits by RNAi from L3 larval stage until adulthood (Fig. 5). Thus, we infer that the defects induced by canal mutations reflect an impairment of V0 + V1 proton pumping, and that they are caused by loss-of-function rather than by gain-of-function mutations. Expansion of the excretory canal in these mutants might help

Figure 3. Mutations introduced in the V0 subunit VHA-5. (A) Strategy to generate vha-5 mutations based on complementation of the vha-5-null allele vha-5(mc38). (B) Predicted topology for VHA-5 based on yeast Vph1p (Nishi and Forgac, 2002) and positions of substitutions. Box, symbols for the most important phenotypes (see also Fig. S2). DcClustal alignment (http://bips.u-strasbg.fr/PipeAlign/jump_to.cgi?DcClustal+noi) of VHA-5 with the three other C. elegans “a” subunits, VHA-6 (intestinal), VHA-7 (epidermal), and UNC-32 (ubiquitous in the embryo, and then muscular and neuronal), the most closely related human and fly “a” subunits (human ATP6V0a1 and D. melanogaster V100), and the S. cerevisiae “a” subunit Vph1p. The positions of the mutations and the predicted positions of the transmembrane domains (numbered with roman letters) are indicated above VHA-5. Fig. S2 is available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1.
to compensate for the decrease in proton-pumping efficiency. Surprisingly, animals with canal mutations did not show any proton-pumping defect in the epidermis. Possibly, proton-pumping is preserved in this tissue because the "a" subunit VHA-7, which is expressed in the epidermis but not in the excretory cell (Oka et al., 2001; Pujol et al., 2001), compensates for the mutated VHA-5. In contrast, we infer that the cuticle defects induced by cuticle mutations reflect an impairment of the V0-specific secretion function, which would not be compensated by other "a" subunits, probably because they are not endowed with this specific function. The existence of various "a" subunits with possibly different functions in the epidermis is reminiscent of the difference observed in yeast between Vph1p and Stv1p (Kawasaki-Nishi et al., 2001). Lastly, we suggest that the mixed mutation W327A affects both V0+V1 and V0-specific functions. We note that cuticle mutations are located in the last

Figure 4. Genetic separation of the V0-specific and V0+V1 functions of the V0 subunit VHA-5. (A) Body length of adults at the same age (Error bars represent the SD; ***: significantly different from wild-type with \( P < 0.0001 \)). In this and subsequent figures, vha-5(mc38) animals with a mutant vha-5::gfp or vha-5::rfp transgenic are noted mc38; Ex[substitution]. (B) Western blot analysis with VHA-5 antiserum of extracts prepared from the three main mutants described in the text. The VHA-5::GFP protein levels are comparable, relative to an actin loading control. (C) Adult outer cuticle, alae (arrows), and annuli (arrowheads) observed by SEM [genotypes indicated above images]. Note the stunted alae and annuli defects induced by L786S and E830Q mutations. (D) GFP fluorescence of the VHA-5 construct in the excretory canal of similar adults [top row, single XY confocal section; bottom row, transverse XZ projection. Compare the normal lumen [arrow] in the control animal with the whorls [arrowheads] induced by the W190A mutation. (E) Excretory canal in similar adults observed by TEM, and quantification of the canal section area. Note the multiple lumens (black arrowheads) in mc38; Ex[W190A] animals (dotted lines outline the excretory canal). NS, not significantly different from control animals; SD, standard deviation.

Figure 5. Knockdown of V0 and V1 subunits induces whorls in the excretory canal. VHA-5::GFP fluorescence in the excretory canal of adults after RNAi against V0 (vha-1 and vha-4) or V1 (vha-8 and vha-13) subunits performed during larval development. Note the presence of whorls [arrowheads; the normal lumen is outlined with arrows], as in mc38; Ex[W190A] animals (Fig. 4 D).
transmembrane domain or in the COOH-terminal luminal tail, whereas canal mutations are in the NH$_2$-terminal cytoplasmic part, which is more likely to interact with the V1 sector (Nishi and Forgac, 2002). We conclude that the V0-specific and V0+V1 functions of VHA-5 are genetically separable.

**VO mediates secretion of exosomes through MVBs**

If indeed the V0 sector is involved in secretion, *vha-5* cuticle mutants should accumulate secretory organelles. At low magnification, TEM through the epidermis showed that animals carrying cuticle or mixed mutations contained significantly more and larger dense organelles than wild-type adults or animals carrying canal mutations (Fig. 6 A and Fig. S3 C). At higher magnification these organelles appeared as multivesicular bodies (MVBs; Fig. 6, B and C). MVBs are endosome-derived organelles containing 30–90 nm vesicles, which grow from early and recycling endosomes or from the trans-Golgi network and evolve into lysosomes or into secretory organelles (Raiborg et al., 2003). Hence, MVB accumulation may reflect either an endocytic/degradation pathway or a secretory defect. To distinguish between these two possibilities, we examined whether *vha-5* mutants had normal lysosomes. In addition, we compared *vha-5* defects to those induced by strong mutations in *vps-27, rme-8*, and *cup-5*, which are three essential genes acting at different steps along the endocytic route (Zhang et al., 2001; Treusch et al., 2004; Roudier et al., 2005). The rationale for this comparison is that if *vha-5* cuticle mutations affect endocytosis,
then vps-27, rme-8, or cup-5 mutations should induce cuticle phenotypes comparable to those of vha-5 mutants. As found in other systems (Luzio et al., 2001), we could observe in all vha-5 mutants intermediate late endosome–lysosome compartments corresponding to enlarged MVBs with multilamellar structures (called hybrid-MVBs in the next section) and normal lysosomes, suggesting that the endocytic/degradation pathway was not qualitatively affected (Fig. 6 C). In contrast, cup-5(ar465) mutants accumulated large MVBs in their epidermis that did not evolve into lamellar structures (Fig. 7 A). Furthermore, vps-27(ok579), rme-8(b1023), and cup-5(ar465) mutants had normal alae, unlike vha-5 cuticle mutants (Fig. 7, B and C). We conclude that cuticle mutations are unlikely to affect degradation, and, rather, they affect a secretory pathway.

To understand the relevance of MVB accumulation in vha-5 mutants, we reinvestigated secretion in the wild-type C. elegans epidermis. In hematopoietic cells, some MVBs release their vesicle content into the extracellular space and, thus, play a role in exocytosis, in addition to their well-defined role in the endosomal pathway (de Gassart et al., 2004). The vesicles released by fusion of MVBs with the plasma membrane were originally called exosomes in antigen-presenting cells. In support of the notion that secretion in the C. elegans epidermis involves exosomes, we observed small light MVBs containing 50–100-nm vesicles just beneath the apical plasma membrane. Moreover, we occasionally saw vesicles immediately above the plasma membrane in the inner cuticular layer, strongly suggesting that a MVB had released its intralumenal vesicles (Fig. 6 B). These MVBs were always found in the vicinity of epidermal apical membrane stacks, a structure whose role is so far unknown (White, 1988). In contrast, we rarely observed similar MVBs adjacent to the plasma membrane in cuticle mutants, or they were darker (Fig. 6 C). These data suggest that the MVB-limiting membrane can fuse with the apical membrane to release exosomes in the cuticle, and show that this process is impaired in vha-5 cuticle mutants. It raises the possibility that the V0 sector is critical for MVB fusion with the apical membrane during exosome release.

V0 mediates secretion of hedgehog-like peptides

An important expectation of the cuticle defects described so far is that we should be able to identify cuticular proteins whose secretion depends on VHA-5 activity. Cuticle proteins include collagens and Hedgehog-related peptides (McMahon et al., 2003; Zugasti et al., 2005). We found that the collagen DPY-7 was efficiently secreted in vha-5(mc38)–null animals, in animals carrying cuticle mutations, as well as in che-14(mc35) mutants (Fig. S4, available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1). We turned our attention onto Hedgehog-related peptides, which appeared as good candidates for three reasons. First, vha-5 alae defects partially resemble those observed in che-14 mutants. Second, CHE-14 is homologous to Dispatched, which is a protein required for Hedgehog release (Burke et al., 1999; Michaux et al., 2000). Third, despite the absence of a Hedgehog homologue in C. elegans, its genome contains several Hedgehog-related peptides required to generate a normal cuticle, although their precise roles remain unknown (Aspock et al., 1999; Zugasti et al., 2005).

We tagged with GFP the secreted domain of the Hedgehog-related peptides WRT-2 and -8 (Fig. 8 A and Fig. S3 E), which are expressed in the epidermis (Aspock et al., 1999). We found that animals bearing cuticle mutations, but not canal mutations, accumulated VHA-5::RFP and WRT-2::GFP or -8::GFP in discrete entities in their epidermis (Fig. 8, B and C; and Fig. S3, D and F). These entities most likely correspond to the dense and hybrid MVBs (Fig. 6, C and D) because VHA-5::RFP also colocalized (Fig. 8 C) with the MVB marker VPS-27::GFP (Roudier et al., 2005). Moreover, both VHA-5 antiserum and a GFP antiserum targeting WRT-2::GFP decorated the MVBs of cuticle mutants (Fig. 8 D and Fig. S5 B, available at http://www.jcb.org/cgi/content/full/jcb.200511072/DC1). Last, in wild-type nontransgenic animals, in addition to membrane stacks (Fig. 1 G), VHA-5 was found at the MVB-limiting membrane, in

![Figure 7](image-url). Fluid-phase endocytosis mutations do not affect cuticle formation. (A) Epidermis of a cup-5(ar465) adult visualized by TEM; note the enlarged electron-dense MVB (demarcated by dotted lines). (B) Adult alae visualized by DIC; vps-27(ok579) is an L2 lethal mutation and rme-8(b1023) is a temperature-sensitive lethal mutation. vps-27, rme-8, and cup-5 mutations did not affect alae formation, in contrast to vha-5 cuticle mutations.
intralumenal vesicles, and in the cuticle (Fig. 8 E and Fig. S5 A), suggesting that it could act at different steps in the secretion of vesicle (see Discussion). Importantly, the VHA-5 protein with the substitutions L786S (Fig. 8 B) or E830Q (not depicted) could reach the plasma membrane in heterozygous vha-5(mc38)/+ animals, which strongly suggests that their intracellular retention in homozygous mc38 animals is caused by the loss of a trafficking function rather than by misfolding. Consistently, the WRT-2/8 proteins were not retained intracellularly either in heterozygous vha-5(mc38)/+ animals, despite the presence of the L786S (Fig. 8 B), or in E830Q transgenes (not depicted). These results indicate that the V0 sector plays a key role in a specific apical secretion pathway that is taken on by Hedgehog-related proteins, but not by collagens.

Discussion
Whereas basolateral secretion is known to depend on the activity of specific complexes (AP-1B and the exocyst), no such complex has been implicated in apical secretion. In addition, although recycling endosomes appear to play a central role in basolateral secretion, their importance in apical secretion is still under active investigation. Our characterization of mutations affecting the V-ATPase “a” subunit VHA-5 sheds new light on the apical biosynthetic secretory pathway. We could observe the fusion of MVBs with the apical plasma membrane in wild-type animals, and the subsequent release of exosomes. In contrast, we found that some VHA-5 mutations accumulate MVBs in their epidermis and prevent the secretion of Hedgehog-related proteins. Thus, we propose a model whereby apical secretion of Hedgehog-related proteins involves their incorporation into the intralumenal vesicles of MVBs, and their subsequent release when MVBs fuse with the apical plasma membrane (Fig. 9). Furthermore, we suggest that the V0 sector of the V-ATPase plays a key role in this process.

We can envision two scenarios for the role of the V0 sector. First, vha-5 mutations affecting cuticle formation could decrease V-ATPase proton pumping along the biosynthetic secretory route to indirectly impair secretion. Consistent with this possibility, mutations in S. cerevisiae Vph1p (L746S, E789Q, and V803F) corresponding to the cuticle mutations (L786S, E830Q, and V844F, respectively) strongly reduce, but do not
membrane are compromised so that MVBs grow and become denser by accumulating their content, which can nevertheless be normally degraded.

vacuole fusion and at the plexes between both membranes (Fig. 9), as suggested in yeast the apical membrane via the formation of V0 sectors transcomplexes and at the MVB-limiting membrane (Fig. 8 E and Fig. S5 A), and by the accumulation of dense or hybrid MVBs in cuticle mutants (Fig. 8 E and Fig. S5 A). We expect the existence of two distinct V0 populations, some mediating fluid-phase endocytosis, others mediating proton pumping with the V1 sector. In cuticle defective vha-5 mutants (right), most fusion events between MVBs and the plasma membrane are overcome so that MVBs grow and become denser by accumulating their content, which can nevertheless be normally degraded.

Figure 9. **Model for apical secretion mediated by the V0 sector of the V-ATPase.** We propose that in a wild-type animal (left) the V0 sector mediates fusion between the limiting membrane of MVBs and the apical plasma membrane (Fig. 1, E–G) and the MVB-limiting membrane (Fig. 8 E and Fig. S5 A), and by the accumulation of dense or hybrid MVBs in cuticle mutants (Fig. 8 E and Fig. S5 A). We expect the existence of two distinct V0 populations, some mediating secretion, others mediating proton pumping with the V1 sector. In cuticle defective vha-5 mutants (right), most fusion events between MVBs and the plasma membrane are compromised so that MVBs grow and become denser by accumulating their content, which can nevertheless be normally degraded.

What could explain a common requirement for the V0 sector during the C. elegans apical exocytosis, yeast vacuole fusion, and D. melanogaster synaptic transmission? The prevailing view is that a SNAREpin complex initiates membrane fusion once a vesicle has been docked to a proper membrane (Chernomordik and Kozlov, 2003; Jahn et al., 2003). Although the V0 sector is thought to act downstream of SNAREs in yeast and D. melanogaster (Peters et al., 2001; Bayer et al., 2003; Hiesinger et al., 2005), we cannot exclude that it also acts in parallel to SNAREs, at least in C. elegans, to dock MVBs. Another possibility is that V0 transcomplexes initiate the formation of a protein pore, as initially suggested in yeast (Peters et al., 2001). On the other hand, expansion of the fusion pore is considered as the limiting step in membrane fusion, and might require additional catalysts in vivo (Chernomordik and Kozlov, 2003). Such a role could be fulfilled by the V0 sector, either to overcome constraints caused by the big size and/or the specific lipoprotein content of C. elegans epidermal MVBs and yeast vacuoles, or to allow rapid synaptic transmission in D. melanogaster neurons (Hiesinger et al., 2005).

Irrespective of the precise role of the V0 sector in membrane fusion, our findings bear potentially important implications. First, morphogens such as Wingless and Hedgehog in D. melanogaster, or Sonic-Hedgehog at the mouse node, might be secreted through a similar pathway because their secretion involves particles possibly related to exosomes (Greco et al., 2001; Gallet et al., 2003; Panakova et al., 2005; Tanaka et al., 2005). A major objective will be to determine whether CHE-14 and Dispatched act in the aforementioned secretory pathway, and, if so, at which step. Second, several other cell types, such as antigen-presenting cells, reticulocytes, and some epithelial cells, can release exosomes (de Gassart et al., 2004), which might thus also require the V0 sector for their secretion. In particular, the V0 sector might be directly associated with the transmission of some infectious diseases because viruses, such as HIV and the prion protein, can be disseminated through MVBs and the exosome-releasing machinery (de Gassart et al., 2004; Fevrier et al., 2005). Likewise, the aforementioned secretory pathway could be involved in innate immunity because expression of the Hedgehog-related peptide GRD-3 is induced in C. elegans upon bacterial infection (Couillault et al., 2004).
Third, our findings raise the issue of the origin of the MVBS. Interestingly, the apical recycling endosomes have been recognized to play an important role in biosynthetic secretory pathways (Hoekstra et al., 2004). Future studies should reveal whether the secretory MVBS that we described could originate from this compartment.

In conclusion, our work shows that trafficking to the apical membrane of at least some lipid-modified proteins involves specific protein complexes (the V-ATPase V0 sector), much as trafficking to the basolateral membrane, and predicts a key role for MVBS in apical exocytosis.

Materials and methods

Strain maintenance and RNA interference

Worms were grown at 20°C (unless noted otherwise; Brenner, 1974). The identification of mc37 and mc38 as vha-5/F53H10.4::null mutations will be described elsewhere (the gene affected by mc37 and mc38 was initially named rdy-1; see www.wormbase.org). Marker and alleles used are as follows: cup-5000(e546) (Treusch et al., 2004); mc38; Ex[vha-5(E830Q)::rfp] (Roudier et al., 2005). The L1 alae phenotype of mc38 was scored by allowing adults to lay eggs for 1 h at 15°C, and then transferring embryos to 25°C after egg laying. RNAi was performed using the following bacterial clones from the Wellcome- Medical Research Council library (Kamath et al., 2003); vha-1, il5420; vha-4, il5716; vha-5, IV-AO13; vha-8, IV-3I08; and vha-13, V9-O06 (www.wormbase.org). To score L1 larvae, RNAi was induced in larvae at the L2–L3 molt. Feeding plasmids were retransformed into fresh HT115 (DE3) bacteria, selecting for ampicillin resistance. To score adults, RNAi was induced in larvae at the L2–L3 molt. Feeding plasmids were retransformed into fresh HT115 (DE3) bacteria, selecting for ampicillin resistance.

Plasmids

Cloning of the vha-5 coding sequence with a 2.8-kb promoter upstream of the GFP coding sequence in the pPD95.75 vector (Fire kit) generated a recombinant plasmid. A vha-5::mrfp construct was obtained by replacing the GFP with the monomeric red fluorescent protein (mRFP) coding sequence in the vha-5:gfp construct. A vha-8::yfp construct was obtained by cloning the vha-8 coding sequence and a 3-kb promoter upstream of the YFP coding sequence in the pPD136.64 vector (Fire kit). To generate vha-5::gfp; rol-6(su1006) construct. A vha-5::gfp construct was mutated using the QuikChange Site-Directed Mutagenesis kit (Strategene). Each desired mutation, and the entire vha-5::gfp construct. A vha-5::mrfp construct was used to generate a strain with a vha-5 mc38; Ex[vha-5(E830Q)::rfp] construct. A vha-5::mrfp construct was obtained by replacing the GFP with the monomeric red fluorescent protein (mRFP) coding sequence in the vha-5:gfp construct. A vha-8::yfp construct was obtained by cloning the vha-8 coding sequence and a 3-kb promoter upstream of the YFP coding sequence in the pPD136.64 vector (Fire kit). To generate vha-5::gfp; rol-6(su1006) construct. A vha-5::gfp construct was mutated using the QuikChange Site-Directed Mutagenesis kit (Strategene). Each desired mutation, and the entire vha-5::gfp construct. A vha-5::mrfp construct was used to generate a strain with a vha-5 mc38; Ex[vha-5(E830Q)::rfp] construct. A vha-5::mrfp construct was obtained by replacing the GFP with the monomeric red fluorescent protein (mRFP) coding sequence in the vha-5:gfp construct.
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