Lipoprotein receptors and a Disabled family cytoplasmic adaptor protein regulate EGL-17/FGF export in C. elegans

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Growth factors and morphogens need to be secreted to act on distant cells during development and in response to injury. Here, we report evidence that efficient export of a fibroblast growth factor (FGF), EGL-17, from the Caenorhabditis elegans developing vulva requires the lipoprotein receptor-related proteins Ce-LRP-1 and Ce-LRP-2 and a cytoplasmic adaptor protein, Ce-DAB-1 (Disabled). Lipoprotein receptors are transmembrane proteins best known for their roles in endocytosis. Ce-LRP-1 and Ce-LRP-2 possess a conserved intraluminal domain that can bind to EGL-17, as well as a cytosolic FXNPXY motif that can bind to Ce-DAB-1. Ce-DAB-1 contains signals that confer subcellular localization to Golgi-proximal vesicles. These results suggest a model in which Ce-DAB-1 coordinates selection of receptors and cargo, including EGL-17, for transport through the secretory pathway.

Keywords: Selective secretion; fibroblast growth factor; lipoprotein receptor traffic; Dab; cargo selection; endocytosis

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Many developmental morphogens and growth factors have specialized mechanisms of secretion. For example, Wingless [Wg] secretion requires posttranslational N-glycosylation by the transmembrane protein Porcupine [Porc] within the endoplasmic reticulum [van den Heuvel et al. 1993; Tanaka et al. 2002], whereas a Porc relative, Central missing [Cmn], is critical for Hedgehog [Hh] export [Amanai and Jiang 2001]. Spitz, a transmembrane epidermal growth factor [EGF], is held within the ER until Star chaperones it to the Golgi apparatus, where it then encounters Romboid-1, which triggers proteolytic cleavage and allows Spitz to move to the cell surface [Lee et al. 2001]. Most fibroblast growth factors [FGFs] act cell nonautonomously to coordinate cell migrations, proliferation, and fate specification during wound healing, angiogenesis, tumor metastasis, and development [Friesel and Maciag 1995; Goldfarb 1996; Sutherland et al. 1996; Borland et al. 2001]. The majority of FGFs contain standard signal sequences for secretion by the ER-Golgi pathway. However, FGFs 1, 2, and 11–14 lack signal sequences, and FGFs 9, 16, and 20 have a hydrophobic N terminus that may not be cleaved [Friesel and Maciag 1995; Ornitz and Itoh 2001]. Thus, certain FGFs may use distinct mechanisms for secretion.

A Caenorhabditis elegans fibroblast growth factor [FGF], EGL-17 [egg-laying defective], regulates the migrations of the sex myoblasts [SMs] during larval development [DeVore et al. 1995; Burdine et al. 1997, 1998]. The SMs are born at the posterior of L1 stage larvae and migrate anteriorly to the gonad center, where they generate the uterine and vulval musculature [Thomas et al. 1990]. EGL-17 is expressed in the developing vulva and gonad, and acts as an attractive cue for the SMs, which express an FRF receptor [FGFR], EGL-15. Mutations in genes for either the ligand [egl-17] or receptor [egl-15] prevent correct SM positioning around the gonad center, and egg-laying defects result [Stern and Horvitz 1991]. Although these data indicate that EGL-17 is secreted, the mechanism by which this occurs has not been investigated.

Here we report that efficient export of EGL-17 by the developing vulva involves the adaptor protein Disabled (Ce-DAB-1) and two lipoprotein receptor-related proteins, Ce-LRP-1 and Ce-LRP-2. Ce-DAB-1 and the receptors appear to function posttranscriptionally to promote secretion. Lipoprotein receptors cycle rapidly through the endocytic pathway, bringing bound ligands into the cell and releasing them in the acidic environment of endosomes before returning to the cell surface [Nykjaer and Willnow 2002]. Their extracellular domains contain variable numbers of repeated domains that allow them...
to interact with a diverse set of macromolecules bearing little structural similarity [Herz and Strickland 2001; Christensen and Birn 2002]. Although the primary roles of lipoprotein receptors appear to be in endocytosis, recent evidence suggests that they may participate in the transcytosis of at least some of their ligands [Dehouck et al. 1997; Marino et al. 2000, 2001]. However, there is no evidence regarding a possible role in export of proteins from the cell. Ce-DAB-1 is related to murine Dab1, which functions in a signaling pathway together with lipoprotein receptors VLDLR [very low density lipoprotein receptor] and ApoER2 [Apolipoprotein E receptor], and Dab2, which regulates endocytosis of lipoprotein receptors Megalin and LDLR [Howell and Herz 2001; Mishra et al. 2002a; Morris et al. 2002]. Dab1 and Dab2 contain a PTB [phosphotyrosine-binding] domain, also found in Ce-DAB-1, which mediates their association with an FXNPXY motif that is conserved in the intracellular domains of lipoprotein receptors [Howell et al. 1999; Trommsdorff et al. 1999; Morris and Cooper 2001]. The FXNPXY motif in the LDL [low-density lipoprotein] receptor is critical for endocytosis and also for trafficking to the basolateral surface of polarized epithelial cells [Nelson and Yeaman 2001]. The binding of Dab proteins to a conserved signal on lipoprotein receptors suggests that they may directly regulate receptor traffic.

Results

Roles of Ce-dab-1 during larval development

The C. elegans gene Ce-dab-1 encodes a protein related to Disabled [Dab] family members in mice and Drosophila [Fig. 1A]. The gene appears to be alternatively spliced, but all predicted proteins contain a conserved PTB domain and, like mammalian Dab2, have several sequences that may confer binding to clathrin adaptor proteins [Fig. 1A, red triangles] and Eps15 homology [EH] domain proteins [blue circles].

To identify a role for Ce-dab-1, gene function was disrupted by using RNA interference [RNAi; Fire et al. 1998]. Stage L1 larvae were soaked in Ce-dab-1 or control double-stranded RNA [dsRNA; Tabara et al. 1998; Maeda et al. 2001]. A number of phenotypes were reproducibly observed in the F₀ generation, including highly penetrant defects in cuticle molting [Mlt] and in egg laying [Egl; Table 1]. dsRNA prepared from alternate regions of Ce-dab-1 conferred the same phenotypes, indicating specificity [Table 1].

The Mlt defect of Ce-dab-1 RNAi-treated worms resembled that caused by mutation of Ce-lrp-1, which encodes a Megalin relative [Yochem et al. 1999]. The cuticle formed blisters, and some worms dragged cuticle remnants behind them. Many adults had a "wasp waist"
close to the vulva, apparently due to a girdle of unshed cuticle [Fig. 1C]. The same phenotype was induced by Ce-Irp-1 RNAi [Fig. 1D] or cholesterol starvation [Yochem et al. 1999]. This suggests that Ce-dab-1 and Ce-Irp-1 may regulate cholesterol availability for cuticle synthesis or molting.

Ce-dab-1 RNAi worms laid eggs at a slower rate and later stage than did control animals [Fig. 1B; Table 1]. Egl phenotypes can result from deficiencies in the development of the vulva, gonad, or egg-laying muscles [Sulston and Horvitz 1977; Newman et al. 1996]. The vulvae and gonads of Ce-dab-1 (RNAi) animals appeared to be normal, and the egg-laying rate did not correlate with the presence of cuticle fragments in or near the vulva, suggesting that the Egl phenotype was not caused by physical blockage. We therefore examined the egg-laying muscles, using worms expressing green fluorescent protein (GFP) in their SMs [Harfe et al. 1998]. Depletion of Ce-dab-1 caused significant posterior displacement of the SMs (P < 0.0001; Fig. 1E). These defects were similar to, but less severe than, those caused by egl-17 mutation [Burdine et al. 1997, 1998]. This suggests that Ce-dab-1 may be needed for proper functioning of the EGL-17/EGL-15 signaling pathway.

Ce-dab-1 is expressed in the developing vulva

To distinguish whether Ce-DAB-1 regulates SM migration through the EGL-17 producing cells or the EGL-17 responding SMs, we identified the sites of Ce-dab-1 gene expression. We used 5.5 kb of Ce-dab-1 upstream sequence to express either a DAB-1::GFP translational fusion [Ce-dab-1::DAB-1::GFP; Fig. 2A] or GFP [Ce-dab-1::GFP; data not shown] in transgenic worms. L2 stage worms showed GFP fluorescence in the ventral hypodermis [Fig. 2A]. In normal hermaphrodites, three hypodermal cells [the VPCs, P5.p, P6.p, and P7.p], give rise to the vulva [Fig. 2B; Sulston and Horvitz 1977]. Expression of Ce-dab-1 within the VPCs and their descendants continued through vulval development, and became restricted to the descendants of P5.p and P7.p by mid-L4 [Fig. 2]. Ce-dab-1 was also expressed in the anchor cell (AC), sheath cells surrounding the amphid neurons in the head, the gut, and several unidentified cells in the anus and uterus of L3-adult animals. However, no Ce-dab-1 expression was detectable in the SMs. The expression of Ce-dab-1 in the developing vulva and gonad partially overlaps that reported for egl-17 [Fig. 2H; Burdine et al. 1998; Branda and Stern 2000], suggesting that Ce-dab-1 may regulate SM migration through an interaction with egl-17.

Ce-DAB-1 prevents EGL-17 protein accumulation in the VPCs

We considered that Ce-dab-1 might regulate egl-17 gene expression or EGL-17 protein secretion. Because of weak expression of EGL-17 reporter constructs in the developing gonad [Branda and Stern 2000], we monitored egl-17 expression in the descendants of P6.p. To test whether Ce-dab-1 regulates egl-17 promoter activity, we used animals expressing GFP from the egl-17 promoter [Fig. 3A; Burdine et al. 1998]. Ce-dab-1 RNAi has no effect on the levels of egl-17::GFP expression during early [Fig. 3A] or late [data not shown] vulval development. Therefore, we tested whether Ce-dab-1 might regulate the export of EGL-17 from the developing vulva, by using the egl-17 promoter to express a translational fusion of full-length EGL-17 to GFP [Fig. 3C]. To determine whether the transgene is functional, it was injected into egl-17(n1377) mutant animals. Animals bearing the transgene showed a pronounced reduction in the characteristic egl-17(n1377) “bag of worms” Egl phenotype [Fig. 3B]. This suggests that the EGL-17::GFP fusion protein is exported and functional.

EGL-17::GFP protein traffic was monitored by both live GFP and indirect immunofluorescence in wild-type egl-17::EGL-17::GFP larvae. Long exposures revealed weak punctate fluorescence in the daughters or granddaughters of P6.p [Fig. 3C, top]. The rescue of the egl-17

Table 1. Ce-dab-1 RNAi phenotypes

| RNAIa | %eglβ | Laying rate [eggs/h]c | Staged | % mltε | Animals scoredδ |
|-------|-------|---------------------|--------|--------|-----------------|
| Control | 1.5 ± 1.4 | 4.8 ± 1.2 | -gastrulation | 0 | 720 |
| Ce-dab-1 | 74.9 ± 9.7 | 1.8 ± 0.9 | pretzel | 26.1 ± 6.3 | 249 |
| 1–546 | 77.7 ± 5.7 | ndb | pretzel | 25.6 ± 14.6 | 257 |
| 290–546 | 75.9 ± 5.2 | nd | pretzel | 31.8 ± 10.2 | 161 |

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- 1–546 77.7 ± 5.7 nd 25.6 ± 14.6
- 290–546 75.9 ± 5.2 nd 31.8 ± 10.2

*1 L1 animals were soaked overnight in solutions of dsRNA and scored as young adults. Control RNAi was prepared from an empty vector.

*2 Egg-laying defective animals were scored as young adults by using DIC microscopy and looking for postgastrulation embryos.

*3 The number of eggs laid by individual animals was counted over several hours and averaged (n = 10).

*4 The approximate stage of eggs upon laying were marked as either -gastrulation stage [30 to 150 cells] or -pretzel stage [400 to 550 cells].

*5 Molting-deficient animals were scored by DIC as predominantly young adults. Only animals with body constrictions due to unshed cuticle were scored as positive.

*6 Numbers represent a total of six independent experiments.

*7 Numbers represent residues in the corresponding protein sequence.

*8 nd indicates not determined.

Expression of Ce-dab-1 in the vulva and gonad regulated the levels of EGL-17 to GFP (Fig. 3C). To determine whether Ce-dab-1 might regulate the express of EGL-17::GFP, we used animals expressing GFP from the egl-17 promoter [Fig. 3A; Burdine et al. 1998]. Ce-dab-1 RNAi has no effect on the levels of egl-17::GFP expression during early [Fig. 3A] or late [data not shown] vulval development. Therefore, we tested whether Ce-dab-1 might regulate the export of EGL-17 from the developing vulva, by using the egl-17 promoter to express a translational fusion of full-length EGL-17 to GFP [Fig. 3C]. To determine whether the transgene is functional, it was injected into egl-17(n1377) mutant animals. Animals bearing the transgene showed a pronounced reduction in the characteristic egl-17(n1377) “bag of worms” Egl phenotype [Fig. 3B]. This suggests that the EGL-17::GFP fusion protein is exported and functional.

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Figure 2. Ce-dab-1 is expressed in the developing vulva. (A) Vulvae of qaEx4003[Ce-dab-1::Ce-DAB-1::GFP; unc-119(+); unc-119(e2498)] animals were observed under DIC and fluorescence and photographed as development progressed, increasing in time from top left to bottom right. Before the onset of vulval development (L2; top left panels), expression is seen in the cells of the ventral hypodermis, including the descendants of P5.p, P6.p, and P7.p. As vulval development progresses, expression continues and becomes restricted to the descendants of P5.p and P7.p by the time that vulval invagination is complete (L4; bottom right). Note that the P6.pxxx (great grand-daughters of P6.p) move out of the plane of focus as they adopt their terminal fate. The anchor cell is indicated by an inverted black triangle. Fluorescence above the anchor cell is gut fluorescence. Expression is identical in the independent integrated strain qaIs4000[Ce-dab-1::GFP; unc-119(+); unc-119(e2498)]. (B) Ce-dab-1 and egl-17 expression partially overlaps. Schematic illustration of the expression patterns of Ce-dab-1 (green) and egl-17 (red, as described; Burdine et al. 1998) during vulval development.
Figure 3. Ce-dab-1 RNAi induces Egl-17–GFP accumulation in the VPCs. (A) Ce-DAB-1 does not regulate egl-17 expression. The onset and level of GFP expression at various stages of vulval development was analyzed in ayIs4[egl-17::GFP, dpy-20(+)], dpy-20(e1282ts) animals (Burdine et al. 1998). Neither control nor Ce-dab-1 RNAi had an effect on the onset of GFP expression driven by the egl-17 promoter. Epifluorescence images were exposed equally. (B) An EGL-17-GFP fusion [egl-17::EGL-17::GFP] rescues the bag-of-worms phenotype of egl-17(n1377) animals. Progeny of animals expressing the injected pRF4[rol-6(su1006)] coinjection marker were observed for the bag-of-worms phenotype. Fifteen to 30 animals were scored from each set of injections. Empty GFP vector was used as a control. (C) Ce-DAB-1 regulates intracellular distribution of EGL-17/H11019GFP. qaEx4002[egl-17::EGL-17::GFP; unc-119(+)]; unc-119(e2498) transgenic animals were tested for changes in EGL-17::GFP distribution after treatment with control or Ce-dab-1 RNAi. Equal exposure times [long or short] were used to record GFP fluorescence. Ce-dab-1 [RNAi] resulted in accumulation of EGL-17–GFP in ∼70% of animals, compared with <5% in controls. A total of >14 independent experiments with >10 animals were performed. Similar results were obtained with the integrated strain qaIs4004[Ce-dab-1::Ce-DAB-1; H11019GFP; unc-119(+)]; unc-119(e2498). (D) LET-23 and AJM-1 localization is unaltered, whereas EGL-17::GFP accumulates in Ce-dab-1 [RNAi] animals. qaEx4002[egl-17::EGL-17::GFP; unc-119(+)]; unc-119(e2498) animals treated with control or Ce-dab-1 RNAi were analyzed by indirect immunofluorescence for LET-23 and AJM-1 (center panels) or LET-23 and EGL-17::GFP [anti-GFP, left and right panels]. Left panels show EGL-17::GFP in a single 0.2 µm confocal plane of P6.p, whereas right panels show a single confocal plane of P6.p after two divisions. Center panels show a single 0.2-µm confocal plane of P6.p after one division. Bar, 20 µm.
mutant phenotype and the pattern of fluorescence suggest that EGL-17::GFP is exported via vesicular structures from P6.p and its descendants and regulates SM migration. Ce-dab-1 RNAi introduced by either feeding [data not shown] or soaking caused a pronounced increase in fluorescence intensity in the descendants of P6.p [Fig. 3C, bottom]. Indirect immunofluorescence reveals an increased number of EGL-17::GFP-containing structures within P6.p and its descendants [Fig. 3D]. Combined with the Egl and SM migration effects of Ce-dab-1 RNAi, this suggests that Ce-dab-1 RNAi inhibits EGL-17 secretion and results in its accumulation in the secretory pathway. The effect was specific, because the cell surface protein LET-23 and the junctional protein AJM-1 were localized normally in these cells despite the retention of EGL-17::GFP [Fig. 3D].

To investigate whether accumulation of EGL-17::GFP can result from reduced secretion, we inhibited several genes implicated in vesicle traffic. The clathrin adaptor protein complex AP-1B is involved in basolateral secretion in mammalian cells [Folsch et al. 1999], whereas AP-3 is involved in Golgi/lysosome traffic and AP-2 in endocytosis [Lippincott-Schwartz et al. 2000]. RNAi for µ1/µ1B subunits of AP-1B (unc-101, apr-1, and apt-10; Shim et al. 2000) each caused a pronounced EGL-17::GFP accumulation in the P6.p descendants but did not cause a Mlt defect [Table 2]. RNAi for a α1 subunit of AP-1 (apt-2) caused both EGL-17::GFP accumulation and defective molting. Conversely, RNAi for potential β6 and β3 subunits of AP-3 (apt-5 and apt-6, respectively), had no phenotype, and RNAi for the α-adaptin subunit of AP-2 (apt-4) was lethal. These results support the hypothesis that retention of EGL-17::GFP can be caused by impaired secretion, and also show that EGL-17::GFP retention is independent of cuticle molting. In addition, Ce-dab-1 is unlikely to have an essential role in AP-2 function.

Lipoprotein receptors Ce-LRP-1 and Ce-LRP-2 participate in EGL-17 export from the P6.p descendants

The sequence of Ce-DAB-1 indicates that it is cytoplasmic. Ce-DAB-1 might therefore regulate secretion by binding to transmembrane receptors present in vesicles that carry EGL-17. Lipoprotein receptors span the membrane and associate with Dab family proteins in vertebrates and, thus, might connect Ce-DAB-1 with EGL-17 in the C. elegans developing vulva.

C. elegans has at least three genes related to mammalian lipoprotein receptors: Ce-lrp-1, rme-2 [receptor-mediated endocytosis; Grant and Hirsh 1999], and a gene related to mammalian Lrp [named here Ce-lrp-2] contained in the cosmids F48B3 and T21E3 [Springer 1998]. Ce-lrp-1 and Ce-lrp-2 expression has been detected in the developing vulva by in situ hybridization [Shin-i and Kohara 2001]. Therefore, we used RNAi to test whether these genes are involved in SM migration. Although inhibition of any of the genes individually had no effect, combined RNAi of Ce-lrp-1 and Ce-lrp-2 caused a significant reduction in SM migration [P < 0.0002, Fig. 4A]. Ce-dab-1 RNAi in addition to Ce-lrp-1 and Ce-lrp-2 had no additional effect on SM migration, suggesting that these genes may work in a common pathway. Rme-2 RNAi had no effect on SM migration when tested alone or in combination with Ce-lrp-1 or Ce-lrp-2 [Fig. 4A], suggesting that the effects of Ce-lrp-1 and Ce-lrp-2 RNAi are specific.

To test the roles of Ce-LRP-1 and Ce-LRP-2 in EGL-17 export, we observed their effects on egl-17 expression and EGL-17::GFP accumulation in P6.p descendants. Combined RNAi for Ce-lrp-1 and Ce-lrp-2 had no effect on egl-17 promoter activity [Fig. 4B] but did cause accumulation of EGL-17::GFP in these cells [Fig. 4C]. As expected, RNAi for Ce-lrp-1 or Ce-lrp-2 individually had no effect in this assay [data not shown]. These results suggest that Ce-LRP-1 and Ce-LRP-2 function with Ce-DAB-1 to facilitate EGL-17 export from P6.p descendants.

Ce-DAB-1 interacts with Ce-LRP-1 and Ce-LRP-2 and localizes to post-Golgi vesicles

In mammals, the PTB domains of Dab1 and Dab2 bind to a consensus FXNPFXY in the intracellular domains of lipoprotein receptors [Howell et al. 1999; Morris and Cooper 2001]. Ce-LRP-1 and Ce-LRP-2 contain such motifs. Therefore, we tested for Ce-DAB-1 PTB domain binding

Table 2. Adapin complexes and EGL-17 secretion

| Genea | Adapin complexb | Subunitb | Functionb | EGL-17::GFP accumulationc | mlt4 |
|-------|-----------------|----------|-----------|---------------------------|------|
| apm-1 | AP-1            | µ1/µ1B   | secretion | +                         | –    |
| apt-10|                 |          |           |                           |      |
| unc-101|                |          |           |                           |      |
| apt-2 |                 |          |           |                           |      |
| apt-4 | AP-2            | α        | endocytosis| n/a5                      | n/a5 |
| apt-5 | AP-3            | δ        | golgi-lysosome traffic | –   | –   |
| apt-6 |                 | β3       |           |                           |      |

aL1 animals were soaked overnight in solutions of dsRNA and scored as L3 larvae. Control RNAi was prepared from an empty vector. 
bPresumptive adaptin complex, subunit, and function by homology with mammalian proteins.
cAccumulation of the EGL-17::GFP fusion protein was measured in the descendants of P6.p as described.
dMolting-deficient animals were scored by DIC as L3/L4 larvae.
eRNAi was lethal prior to larvae reaching L3/L4 stage.
to the intracellular domains of Ce-LRP-1 and Ce-LRP-2. Using the yeast two-hybrid system, both Ce-LRP-1 and Ce-LRP-2 interact with Ce-DAB-1 [Fig. 5A,B]. Ce-DAB-1 PTB domain binding to Ce-LRP-1 was reduced by mutation of the consensus FXNPXY motif from amino acids 4653–4658 but not by mutation of related motifs [Fig. 5C,D], suggesting evolutionary conservation of the Dab PTB binding site. Ce-DAB-1 also contains sequence motifs that may permit binding to clathrin adaptor complexes [Fig. 1A], suggesting that it might localize to coated vesicles. Because there are currently few markers for subcellular compartments in C. elegans VPCs, we observed the subcellular localization of Ce-DAB-1 in mammalian tissue culture cells. As shown in Figure 5E,

Figure 4. Lipoprotein receptors Ce-LRP-1 and Ce-LRP-2, but not RME-2, regulate SM migration. [A] Ce-LRP-1 and Ce-LRP-2 regulate SM migration. The positions of individual SMs were measured as in Figure 1. dsRNA was added as indicated: (−) RNAi present; (+) no RNAi. Numbers in the right column indicate the number of animals scored. [B] The lipoprotein receptors Ce-LRP-1 and Ce-LRP-2 do not regulate EGL-17 expression. egl-17::GFP animals were analyzed after treatment with control or Ce-lrp-1, Ce-lrp-2 RNAi. [C] Ce-LRP-1 and Ce-LRP-2 regulate EGL-17 export from the VPCs. qaEx4002[egl-17::EGL-17::GFP; unc-119(+)]; unc-119(e2498) transgenic animals were tested for changes in subcellular distribution of the EGL-17::GFP fusion after treatment with control or Ce-lrp-1, Ce-lrp-2 RNAi.
Lipoprotein receptors interact with EGL-17

During synthesis and processing, receptor extracellular domains are in the lumen of the ER, Golgi, and transport vesicles, where they might associate directly or indirectly with EGL-17. Because the enormous extracellular domains of Ce-LRP-1 and Ce-LRP-2 (∼4600 and ∼4730 residues, respectively) were difficult to express, we first measured binding of EGL-17 to the extracellular domains of mammalian ApoER2 and VLDLR. Like Ce-LRP-1 and Ce-LRP-2, these receptors contain blocks of complement-like LDLR type A repeats, EGF repeats, and six-bladed YWTD/H9252 propellers (Fig. 6A). The extracellular domains of ApoER2 and VLDLR were expressed as fusions to the immunoglobulin Fc region (Hiesberger et al. 1999), together with myc epitope-tagged EGL-17. The receptor–Fc fusion proteins were recovered by using Protein A and were tested for bound EGL-17-myc (Fig. 6B). Both ApoER2 and VLDLR associated with EGL-17-myc.
whereas the unrelated EphA5 did not (Fig. 6B). This suggests that EGL-17 can bind to the extracellular domains of mammalian lipoprotein receptors.

To identify the domains responsible for EGL-17 binding, we made deletions of the ApoER2–Fc fusion protein. Surprisingly, deletion of the LDLR type A repeats, which bind multiple ApoE- and ApoB-containing lipoproteins (Nykjaer and Willnow 2002), had no effect on EGL-17 binding (Fig. 6C; ApoER2$^{\text{H9004A}}$). However, further removal of three EGF repeats and the $\varphi$/H9252 propeller abolished binding (ApoER2$^{\text{H9004AEY}}$). This suggests that the $\varphi$/H9252 propeller and/or EGF repeats are required to bind to EGL-17.

To determine whether a C. elegans lipoprotein receptor can bind to EGL-17, we inserted various regions of Ce-LRP-2 into the minimal ApoER2–Fc$^{\text{H9004AEY}}$ construct. EGL-17-myc bound to a fusion containing the eighth membrane-proximal $\varphi$ propeller and 11 EGF repeats from Ce-LRP-2 (Ce-LRP-2 YE), but not to a similar construct containing only the EGF repeats (Ce-LRP-2 E; Fig. 6D). This suggests that the association between Ce-LRP-2 and EGL-17 may be mediated by one of the $\varphi$ propellers present in the extracellular region of Ce-LRP-2. This physical interaction may be important for the ability of Ce-LRP-2 to regulate EGL-17 transport in the developing vulva.

**Discussion**

We have found that two lipoprotein receptors, Ce-LRP-1 and Ce-LRP-2, and an adaptor protein, Ce-DAB-1, mediate secretion of EGL-17 from the C. elegans VPC P6.p and its descendants. Interfering with expression of either
Ce-dab-1 or both Ce-lrp-1 and Ce-lrp-2 reduced SM migration, slowed egg laying, and caused accumulation of EGL-17::GFP in P6.p descendants [Figs. 1, 3, 4]. A similar accumulation of EGL-17::GFP in P6.p descendants was observed by using RNAi for subunits of clathrin adaptor complex AP-1, but not AP-2 or AP-3. On the other hand, traffic of LET-23 to the apical surface of the VPCs, and its localization to the basolateral surface by the LIN-2/7/10 complex (Simske et al. 1996; Kaech et al. 1998; Whitfield et al. 1999), appears to be normal, and the protein AJM-1 [Koppen et al. 2001] also localizes normally to junctions [Fig. 3]. Moreover, the LET-23 signal sequence, fused directly to either GFP or an EGL-17::GFP fusion protein, supports secretion that is independent of Ce-DAB-1 [data not shown]. Thus, the effects of Ce-DAB-1 on EGL-17 traffic are specific and consistent with a requirement for secretion. The cytoplasmic tails of Ce-LRP-1 and Ce-LRP-2 can bind to the Ce-DAB-1 PTB domain [Fig. 5], whereas a conserved β propeller domain in the extracellular regions of lipoprotein receptors can bind to EGL-17 [Fig. 6A–D], suggesting that protein–protein interactions link Ce-DAB-1 via the receptors to EGL-17. Ce-DAB-1 also localizes to Golgi-proximal subcellular structures when expressed in tissue culture cells, and to vesicles in P6.p descendants. These results support a model shown in Figure 6E. In this model, Ce-DAB-1 is recruited to transport vesicles at some stage of the secretory pathway and aids in the sorting of Ce-LRP-1 and Ce-LRP-2 into nascent export vesicles. We propose that EGL-17 is cosorted via a direct or indirect interaction with the extracellular domains of the receptors [Fig. 6E]. Ce-DAB-1 may also interact with other proteins important for vesicle budding or targeting.

Although it seems likely that reduced EGL-17 secretion from the developing vulva is a primary cause of the observed defects in SM migration and Egg, Ce-LRP-1, Ce-LRP-2, and Ce-DAB-1 may play additional roles. SM migration is regulated by a balance between gonad-dependent attraction and repulsion (GDA and GDR, respectively, Chen and Stern 1998). EGL-17 expression in the early gonad acts as the major attractant for the SMs [Branda and Stern 2000]. The role of EGL-17 coming from the developing vulva is less clear, but is sufficient to precisely position the SMs in the absence of the gonad [Burdine et al. 1998]. Although we detect some expression of Ce-DAB-1::GFP in the anchor cell at later stages of development [Fig. 2A], expression in the early gonad may be below our current limit of detection. Interestingly, Ce-dab-1 [RNAi] effects on SM migration are weaker than those described for null mutants of egl-17 [Stern and Horvitz 1991]. This may reflect reduced, but not eliminated, EGL-17 export, disrupting the balance between attractive and repulsive cues. It is also possible that EGL-17 from the VPCs regulates SM function or differentiation. Thus, the Egl phenotype of Ce-dab-1 [RNAi] animals may reflect a reduction both in SM migration and in SM function.

In addition to a role in secretion, we also found that depletion of Ce-DAB-1 or Ce-LRP-1, but not Ce-LRP-2, causes cuticle Mlt deficiencies similar to those caused by cholesterol starvation or Ce-lrp-1 mutation [Yochem et al. 1999]. This suggests that Ce-DAB-1 and Ce-LRP-1 have an additional role in cholesterol homeostasis. However, impaired cholesterol homeostasis due to Ce-lrp-1 RNAi is not sufficient to reduce SM migration. Interestingly, a genome-wide RNAi screen [Kamath et al. 2003] found that Ce-dab-1 [RNAi] animals are Unc, Gro, Slu, and Emb. Although we only observed the F1 generation and thus did not observe any embryonic lethality [Emb], we did observe the other phenotypes. These phenotypes may be secondary to the cuticle Mlt deficiency, which could be missed in a large-scale screen or could reflect other cellular functions for Ce-DAB-1.

The model suggested by our results does not define the step in EGL-17 secretion that is facilitated by Ce-DAB-1 and the receptors. Each stage of vesicular traffic requires the coordination of coat protein assembly with the selection of target-specific receptors and the appropriate vesicle contents [Schmid 1997; Lippincott-Schwartz et al. 2000]. Ce-LRP-1 and Ce-LRP-2 might function as cargo selection receptors and use Ce-DAB-1 to coordinate coat protein assembly at any of several steps. For example, cargo selection could occur at the ER to Golgi stage. Previously, coassembly of transmembrane proteins and COPII coats has been shown to allow concentrative sorting of receptor-associated soluble proteins in the ER lumen [Malkus et al. 2002]. In yeast, Erv29p is a transmembrane protein that is responsible for sorting α-factor to the Golgi [Malkus et al. 2002]. In vertebrate cells, transmembrane proteins, including BAP31, ERC53, and p24-family proteins, appear to select soluble proteins for transport from ER to Golgi [Lippincott-Schwartz et al. 2000]. Similarly, if a Ce-DAB-1/receptor/ EGL-17 complex forms in the ER, then Ce-DAB-1 might link these components to COPII coat assembly to facilitate transport to the Golgi.

Alternatively, a Ce-DAB-1/receptor complex may be involved in traffic of EGL-17 from the trans-Golgi network [TGN] to the basolateral surface of P6.p and its descendants. This would be akin to the role of GGA proteins, which bind to a DXXXL motif in the tail of the mannose 6-phosphate receptor and traffic it, and its associated soluble cargo, from the TGN to endosomes [Pucetollano et al. 2001; Zhu et al. 2001]. In mammalian epithelial cells, basolateral targeting of the LDL receptor depends on µ1B-containing clathrin adaptor proteins and an FXNFXY signal, but it is not known whether µ1B binds to the signal directly or via another protein [Nelson and Yeaman 2001]. The PTB-containing protein ARH was recently shown to be capable of binding to AP-1 components and to the FXNFXY of the LDL receptor, although it does not localize to the TGN [Mishra et al. 2002b]. Our data suggest that Ce-DAB-1 may be involved in recruiting a clathrin adaptor to the tails of Ce-LRP-1 and Ce-LRP-2 at the trans-Golgi network in P6.p and its descendants in the C. elegans developing vulva.

A third possibility is that Ce-DAB-1 might be involved in binding of EGL-17 to recycling lipoprotein receptors in a common endosome, where ligands and receptors from the cell surface can associate and enter the secre-
tory pathway without passing to lysosomes [Mostov et al. 2000]. This hypothesis is consistent with the recent observations that lipoprotein receptors participate in transcytosis [Nelson and Yeaman 2001]. Apical to basolateral transcytosis of LDL and thyroglobulin and vitamin D binding protein involves the LDLR and Megalin, respectively [Dehouck et al. 1997, Marino et al. 2000, 2001]. During transcytosis, vesicles move between basolateral and apical common endosome compartments, where they may encounter newly synthesized proteins coming from the TGN [Mostov et al. 2000]. In this way, endocytosed lipoprotein receptors that are either recycled or transcytosed may be able to carry newly synthesized ligands to their target membranes. Because Ce-DAB-1 binds to the FXNpXY endocytosis signal, it may assist delivery of Ce-LRP-1 and Ce-LRP-2 to the common endosome in order to collect EGL-17 for delivery to the surface, or may help target them from the common endosome to the basolateral surface. Distinguishing these possibilities will require detailed subcellular analysis of protein traffic in P6.p and its descendants.

Evolutionarily conserved interactions between Disabled family proteins and lipoprotein receptors

Dab family adaptor proteins interact functionally with lipoprotein receptors in both nematodes and mammals, even though the biological processes they mediate vary greatly [Fig. 6F]. Ce-DAB-1 regulates secretion, Dab2 regulates endocytosis in the kidney [Mishra et al. 2002a, Morris et al. 2002], and Dab1 relays extracellular signals during brain development [Howell and Herz 2001], each via lipoprotein receptors. Although the role of Ce-DAB-1 in signaling is unclear, the high degree of functional conservation across species suggests that vertebrate Dab family members or other PTB-containing proteins may participate in regulated traffic of lipoprotein receptors and associated cargoes to the cell surface. Indeed, it is possible that an early embryonic requirement for Dab2 [Morris et al. 2002] might be a consequence of altered protein traffic in polarized epithelial cells of the embryo.

Materials and methods

Nematode culture and microscopy

Nematodes were cultured and manipulated at room temperature by using standard techniques. The standard wild-type strain is Bristol N2. Strains used were egl-17(n1377), unc-119(e2498), lin-1(ku156); gld-1(q266), PD4666[yIs6(X)]; CeTwist hlb-8::GFP [Harfe et al. 1998], and ayIs4[egl-17::GFP, dpy-20(+)]; dp-41(e1282ts) [Burdine et al. 1998]. From this work this is included: qas4000[Ce-dab-1::GFP, unc-119(+)]; unc-119(e2498), qaEx4002[egl-17::EGL-17::GFP, unc-119(+)]; unc-119(e2498), qaEx4003[Ce-dab-1::Ce-DAB-1::GFP, unc-119(+)]; unc-119(e2498), qaEx4004[ecl-17::EGL-17::GFP, unc-119(+)]; unc-119(e2498). For microscopy, worms were mounted on a 3% agarose pad in M9 (42 mM Na2HPO4, 2 mM KH2PO4, 22 mM KCl, 85 mM NaCl, 1 mM MgSO4) supplemented with 10 mM Na3VO4. All fluorescence and DIC (differential interference contrast) images were obtained by using a standard epi-fluorescence microscope equipped with a digital camera and processed with Adobe Photoshop 5.0 (Adobe Systems). For comparisons of image intensity, all images in a set were processed equivalently. Cells were identified on the basis of their positions as described [Sulston and Horvitz 1977, Sternberg and Horvitz 1986]. When visible in the plane of focus shown, the AC is indicated by an arrowhead. Anterior is to the left, dorsal, to the top.

Plasmids and injections

Specifics of plasmid construction are available upon request. Transgenic animals were generated by using standard injection techniques [Fire 1986, Mello et al. 1991]. Multiple independent strains carrying each transgene were screened for expression and phenotype. Representative strains were chosen for each experiment. All PCRs for cloning were performed by using Pfu Turbo (Stratagene), and open reading frames were sequenced to confirm polymerase fidelity. Expression plasmids for ApoER2-V5–Fc, VLDLR-V5–Fc [Hiesberger et al. 1999], and EphA5–Fc [Davis et al. 1994] fusion proteins were constructed as described.

dsRNA-mediated interference (RNAi) and phenotype scoring

Standard techniques were used to synthesize dsRNA from T7 promoter-tagged, PCR-amplified cDNA. Templates for the synthesis of dsRNA are as follows: pDKF8 [Ce-dab-1], yk58584 [ltp-1], yk8d2 [rme-2], yk597d6 [Ce-ltp-2], and BSKS+ (control). After synthesis, RNA was ethanol-precipitated and resuspended in distilled H2O. Eggs were collected from hypochlorite-treated gravid adults [Lewis and Fleming 1995], and after hatching, L1 larvae were soaked overnight at room temperature in 5 µL of solution comprised of equivalent amounts of each RNA strand in 3 mM spermidine. Soaked animals were then transferred to NGM plates seeded with E. coli OP-50 and allowed to develop until L2 to L4, when they were observed and photographed as described above. When multiple RNAi species were used, equivalent amounts of each RNA strand were used. Egl rate experiments were performed as described [Byerly et al. 1976]. For scoring EGL-17::GFP retention, photographic exposures were set to Ce-dab-1 [RNAi] animals. Equal exposure times were then used to measure retention in controls and other RNAi treated animals.

SM migration and egl-17(n1377) rescue

SM studies were performed in the stain ayIs6(X) [CeTwist hlb-8::GFP] [PD4666], which expresses GFP in the nucleus of the descendants of M, and specifically in the SMs and their descendants during L3 to L4 [Harfe et al. 1998]. The final positions of the worms were scored relative to the Pn.p cells as described [Thommas et al. 1990]. For egl-17(n1377) rescue, pDKF47[ecl-17::EGL-17::GFP, rol-6(su1006)] [10 ng/µL] or pDKF58[ecl-17::LET-233p::EGL-17::GFP] with BSKS+ [130 ng/µL] was injected as above. To ensure hypodermal expression [from which the vulva is derived], pRF4[rol-6(su1006)] [10 ng/µL, Mello et al. 1991] was used as an injection marker. F1 animals expressing the rol-6(su1006) marker were picked to individual NGM + OP-50 plates and observed at least once a day for 4 d once becoming adults. If worms developed a “bag of worms” phenotype within 2 d of becoming adults, they were scored as positive for the phenotype. All other animals were scored as negative.

Yeast two-hybrid interactions

The physical interactions of lipoprotein receptor intracellular domains and the Ce-DAB-1 PTH domain were tested as de-
secondary antibodies were used where appropriate. Anti-germ agglutinin (WGA; 12.5 µg/mL, Molecular Probes), and antibody P1F4-E5 culture supernatant (1:10), Texas Red wheat in PBS. Cells were stained with anti-Ce-DAB-1 monoclonal antibody rinsed twice in PBS, fixed in 3.7% formaldehyde for 15 min at calcium phosphate or Lipofectamine Plus (Invitrogen), respectively. For cell culture experiments, NIH3T3 or HeLa cells were transfected with mAb3E6, Q-Biogene) at 1:50. Confocal z-series were used to 1999). Fixed animals were also stained with anti-GFP antibodies LET-23 and AJM-1 were localized as described (Whitfield et al. Immunofluorescence Cells were maintained in DMEM + 10% FBS. One hundred-millimeter dishes of 293T cells were transfected by using calcium phosphate with various combinations of ApoER2-V5–Fc, EphA5–Fc, pDKF33[eqgl-17::myc], pDKF64[ApoER2 ΔAEY], pDKF65[ApoER2 ΔA], pDKF66[Ce-LRP-2 E], or pDKF67[Ce-LRP-2 YE] (6 µg each). The following day, the medium was replaced with DMEM + 10% FBS. Two days after transfection, the cells were rinsed one time in ice-cold PBS and lysed in 500 µL of lysis buffer [1% Triton X-100, 50 mM Tris at pH 7.4, 120 mM NaCl, 2 mM CaCl2]. After clarification by centrifugation at 15,000 RPM, supernatants were precipitated with Protein A Sepharose (Sigma). Precipitated proteins or 20 µL of cell lysate were resolved by 9% SDS-PAGE and transferred to nitrocellulose, and Western blots were performed by using 9E10 (anti-myc) or Protein A-HRP (to detect Fc).

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