HID-1 controls formation of large dense core vesicles by influencing cargo sorting and trans-Golgi network acidification

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HID-1 controls formation of large dense core vesicles by influencing cargo sorting and trans-Golgi network acidification


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ABSTRACT

Large dense core vesicles (LDCVs) mediate the regulated release of neuropeptides and peptide hormones. They form at the trans-Golgi network (TGN), where their soluble content aggregates to form a dense core, but the mechanisms controlling biogenesis are still not completely understood. Recent studies have implicated the peripheral membrane protein HID-1 in neuropeptide sorting and insulin secretion. Using CRISPR/Cas9, we generated HID-1 KO rat neuroendocrine cells, and we show that the absence of HID-1 results in specific defects in peptide hormone and monoamine storage and regulated secretion. Loss of HID-1 causes a reduction in the number of LDCVs and affects their morphology and biochemical properties, due to impaired cargo sorting and dense core formation. HID-1 KO cells also exhibit defects in TGN acidification together with mislocalization of the Golgi-enriched vacuolar H⁺-ATPase subunit isoform a2. We propose that HID-1 influences early steps in LDCV formation by controlling dense core formation at the TGN.

INTRODUCTION

The ability to control the secretion of proteins regulates many aspects of biology, including physiology, development, and behavior. The regulated secretion of peptide hormones, such as insulin and glucagon, is essential for glucose homeostasis, and defects in their secretion lead to serious metabolic deficits. The regulated release of many neuropeptides by the hypothalamus contributes to the modulation of brain function and controls a variety of physiological behaviors ranging from feeding to sleep and reproduction (Argiolas and Melis, 2013; Sohn et al., 2013; Richter et al., 2014).

In addition to the constitutive secretory pathway enabling the immediate release of newly synthesized proteins, specialized secretory cells such as neurons and endocrine cells also express a regulated secretory pathway (RSP). This pathway enables them to store a subset of secretory proteins (e.g., peptide hormones, neuropeptides) into vesicles that accumulate intracellularly. These vesicles are called secretory granules or large dense core vesicles (LDCVs), and their exocytosis can be triggered by an extracellular physiological stimulus, leading to an increase in cytosolic Ca²⁺ concentration. The mechanisms controlling this regulated exocytosis depend on the presence of specific membrane proteins both at the plasma membrane and on LDCVs, including synaptotagmins that act as Ca²⁺ sensors for vesicle fusion and release (Sudhof, 2012). Despite the significance of peptide hormones and neuropeptides for physiology and many human diseases, the cellular and molecular mechanisms controlling the biogenesis of LDCVs, which determines this molecular composition, still remain poorly understood.

Analysis of budding using metabolic labeling in rat neuroendocrine PC12 cells has demonstrated that LDCVs form at the trans-Golgi network (TGN), where sorting of soluble regulated secretory proteins from constitutively secreted proteins occurs (Tooze and Huttner, 1990). LDCVs contain large amounts of “granulogenic” proteins, such as members of the granin family, which aggregate to form a dense core under the specific pH and redox conditions of the TGN (Gerdes et al., 1989; Chanat and Huttner, 1991). It has been proposed that these luminal interactions represent a driving force in LDCV formation (Arvan and Halban, 2004), with proteins destined for other organelles being removed from nascent LDCVs after
budding. This process of maturation can last up to several hours after budding and depends on clathrin as well as on the adaptor proteins AP-1, GGAs, and PACS (Dittie et al., 1996, 1997; Kakhlon et al., 2006). However, even immature LDCVs can undergo regulated release in rat neuroendocrine PC12 cells (Toose et al., 1991), suggesting that LDCV maturation might not be a strict functional requirement for all secretory cells.

Recent studies have revealed the contribution of several other cytosolic factors controlling various aspects of LDCV formation and maturation to neuroendocrine secretion. For example, the EARP complex and its interactor EIPR-1 (Topalidou et al., 2016), as well as the adaptor protein AP-3 (Asensio et al., 2010; Sirks et al., 2013) and its putative coat VPS41 (Asensio et al., 2013), have been involved in cargo sorting to LDCVs, whereas the BAR (Bin-Amphi-physin-Rev) domain protein Arfaptin-1 has been proposed to influence the size of insulin granules by regulating the timing of Arf1-mediated scission at the TGN (Gehart et al., 2012). The BAR domain proteins (Pick1 and ICA69) are thought to bind to immature LDCVs to regulate their number and size (Cao et al., 2013; Holst et al., 2013). Finally, several Rab proteins (Rab-2, Rab-5, and Rab-10), as well as Rab-2 effectors (Edwards et al., 2009; Sumakovic et al., 2009; Hannemann et al., 2012; Sadsharan et al., 2012; Aillon et al., 2014; Pinheiro et al., 2014), are involved in LDCV maturation.

A forward genetic screen in the model organism Caenorhabditis elegans has previously identified HID-1 as a factor implicated in neuroendocrine secretion (Mesa et al., 2011). HID-1 null worms display reduced levels of LDCV soluble cargo and impaired neurosecretion (Mesa et al., 2011, Yu et al., 2011), and conditional knockout (KO) mice lacking HID-1 in beta-cells of the pancreas display a defect in insulin secretion (Du et al., 2016), suggesting that this factor might be significant for glucose homeostasis. Interestingly, HID-1 is a peripheral membrane protein associated with the Golgi/TGN and its expression seems restricted to specialized secretory cells (Wang et al., 2011), suggesting that it might contribute directly to LDCV biogenesis. Here we test this hypothesis and show that HID-1 promotes mammalian neuroendocrine secretion by influencing LDCV cargo sorting and dense core formation. We further demonstrate that HID-1 is required for TGN acidification and propose that it controls an early step in LDCV formation at the TGN.

RESULTS

HID-1 is required for large dense core vesicle–soluble cargo storage and secretion

To investigate a potential role for HID-1 in the mammalian RSP, we generated HID-1 knockout PC12 cells using genome editing. Specifically, we relied on homologous directed repair after cleavage by Cas9 to knock in a fluorescent protein (tdTomato) while effectively knocking out HID-1. Our repair template was designed in such a way that the fluorescent reporter should not be expressed without proper, in-frame recombination (Supplemental Figure S1, A and B). Although this approach does not guarantee biallelic homologous recombination, we reasoned that, as the efficiency of Cas9-mediated cleavage is much higher than that of homologous recombination repair, cells with monoallelic recombination would have a high probability of exhibiting indels on the other allele due to nonhomology end joining. After transfecting PC12 cells with Cas9, gRNA, and our repair template, we sorted cells positive for tdTomato by flow cytometry. After three consecutive rounds of cell sorting, the proportion of positive cells shifted from an initial 0.2% to more than 90%. We further validated that our repair template was integrated at the proper locus by performing PCR on genomic DNA isolated from our sorted cells (Supplemental Figure S1C). We also observed the existence of alleles without tdTomato, suggesting that the integration is monoallelic in some cases. However, sequencing of these nonrecombined alleles revealed the presence of various indels (Supplemental Figure S1D). The most common mutation that we identified (11 out of 24) is a 47 deletion directly downstream of the cleavage site. Supplemental Figure S1D lists all the identified mutations. Importantly, the presence of the wild-type (WT) allele was only found on three occasions. All the other mutations led to out-of-frame deletions or insertions (Supplemental Figure S1E). To confirm that HID-1 was indeed deleted from our cell population, we analyzed our cells by Western blot and immunofluorescence, testing commercial antibodies against HID-1. Although we were unable to find any antibodies suitable for immunoblotting, we identified a mouse monoclonal antibody (see Materials and Methods) displaying a specific signal by immunofluorescence (Figure 1A). Indeed, WT cells displayed a strong perinuclear staining that was absent from HID-1 KO cells. The same signal could be observed in HID-1 KO cells expressing HID-1-HA (Figure 1B). Consistent with our cell sorting enrichment data, we observed by immunofluorescence that HID-1 was absent from more than 90% of our cells, which is consistent with our genotyping results. This genome-editing approach offers the advantage of relying on transient expression of Cas9 and gRNAs, which should limit off-target effects.

Loss of HID-1 significantly reduced basal cellular levels of the LDCV-soluble marker secretogranin II (SgII) to ~40% of that observed in control cells by fluorescence Western blot and impaired the stimulation of SgII release by depolarization (Figure 1, C–E). However, we found no significant reduction after normalizing secreted SgII values to total SgII levels, suggesting that there is no impairment in regulated exocytosis per se and that the decrease in SgII content is the major contributor to the defect in release (Figure 1F). We observed a similar phenotype in a second HID-1 KO PC12 cell line generated using an independent gRNA (Supplemental Figure S2). Furthermore, we observed no change in SgII mRNA levels by qPCR (Supplemental Figure S3) and found a similar storage and secretion defect of a transfected, exogenous soluble LDCV marker (ANF-GFP), whose expression is under the control of a strong CMV promoter, suggesting that the decrease in cellular content is unlikely to be transcriptional (Supplemental Figure S4). Importantly, lentivirus-mediated expression of full-length HID-1 bearing a C-terminal HA-tag (HID-1-HA) in HID-1 KO cells restored SgII to WT levels and rescued the secretion phenotype (Figure 1, C–F), thus ruling out Cas9 off-target effects.

To further assess secretion of LDCV-soluble cargo, we transfected WT and HID-1 KO PC12 cells with NPY fused to the superecliptic pHuorin (NPY-pHuorin), which has been shown to undergo regulated exocytosis (Kogel et al., 2010; Sirks et al., 2013), and monitored individual exocytotic events by live imaging using spinning disk confocal microscopy (Figure 1G and Supplemental Movies S1 and S2). As expected, WT PC12 cells displayed very few events under basal conditions and responded to depolarization (~7-fold over basal). The number of basal events observed in HID-1 KO cells remained low and unchanged, but we observed a striking reduction in the amount of exocytotic events in response to stimulation in these cells (Figure 1G). Altogether, our results suggest that the absence of HID-1 reduces storage and secretion of LDCV cargoes.

HID-1 does not influence the endolysosomal or constitutive secretory pathway

To rule out the possibility that the impact of HID-1 KO on the RSP was due to indirect effects on constitutive secretion or endocytosis, we analyzed secreted fractions obtained from WT and HID-1 KO PC12 cells under nonstimulatory conditions by Coomassie (data not
HID-1 contributes to large dense core vesicle biogenesis by influencing trans-Golgi network acidification

Our observations suggest that HID-1 might be directly involved in LDCV biogenesis. We thus tested whether the loss of HID-1 influences the composition of LDCVs. Relying on equilibrium sedimentation through sucrose, we found a pronounced shift of SgII to lighter fractions (~1 M sucrose compared with ~1.5 M) in HID-1 KO cells (Figure S3A). Interestingly, the loss of HID-1 also redistributed the calcium sensor synaptotagmin 1 away from the WT LDCV peak (bottom of the gradient) toward lighter fractions (Figure S3B), but had no effect on the distribution of the synaptic-like microvesicle marker synaptophysin (Figure S3C). The change in the steady-state distribution of both soluble and transmembrane LDCV markers indicates that HID-1 influences LDCV biochemical properties and might

FIGURE 1: HID-1 is required for SgII storage and secretion from PC12 cells. (A) WT and HID-1 KO PC12 cells were stained with a mouse mAb to HID-1, followed by an anti-mouse antibody conjugated to Alexa Fluor 488 and mounted with Fluoromount containing DAPI. Representative confocal micrographs show the absence of HID-1 staining in HID-1 KO PC12 cells. (B) HID-1 KO PC12 cells transfected with HID-1-HA were stained with a mouse mAb to HID-1 and a rat mAb to HA, followed by an anti-mouse antibody conjugated to Alexa Fluor 488 and an anti-rat antibody conjugated to Alexa Fluor 647. Representative confocal micrographs show the presence of HID-1 staining only in cells expressing HID-1-HA. Scale bar indicates 5 μm. (C) PC12 cells (WT, HID-1 KO, or HID-1 KO transduced with HID-1-HA lentivirus) were washed and incubated for 30 min in Tyrode’s solution containing 2.5 mM K+ (basal) or 90 mM K+ (stimulated). Cellular and secreted SgII were measured by quantitative fluorescence immunoblotting (C), with the secreted SgII normalized to tubulin (D) or total SgII (F) and expressed as percentage of basal secretion in the control, and the basal cellular SgII was normalized to tubulin (E). Multiple comparison statistical analysis was performed by one-way analysis of variance (ANOVA) followed by a post hoc Tukey test; *p < 0.01 relative to KO (n = 4). No statistical difference was observed between WT and KO + rescue. The bar graphs indicate mean ± SEM. (G) WT and HID-1 KO PC12 cells were transfected with NPY-pHluorin and then imaged live by spinning disk confocal microscopy for 15 s in basal Tyrode’s solution. Regulated exocytosis was triggered by the addition of an equal volume of 90 mM K+ Tyrode’s solution (45 mM K+ final) and imaged for an additional 30 s. Images show representative maximum-intensity time projections of 150 basal and stimulated frames. At the end of the experiment, cells were imaged in Tyrode’s solution containing 50 mM NH4Cl, pH 7.4, to reveal total NPY-pHluorin fluorescence by alkalinization and identify transfected cells. Scale bar indicates 5 μm. Bar graph shows the number of exocytotic events per second normalized to cell surface area. Multiple comparison statistical analysis was performed by one-way ANOVA followed by post hoc Tukey test; *p < 0.01 relative to stimulated exocytosis from WT (n = 15 cells for WT and n = 11 cells for KO from two independent experiments).
such as the granins. The acidic environment of the TGN lumen is thought to drive this aggregation in PC12 cells (Chanat and Huttner, 1991). As HID-1 colocalizes with Golgi markers (Wang et al., 2011), we hypothesized that HID-1 might contribute to TGN acidification. To test this directly, we targeted pHluorin to the lumen of the TGN using the transmembrane domain of the TGN marker sialyltransferase (Wong et al., 1992). We confirmed that our reporter indeed localizes to the TGN by transfecting WT and HID-1 KO PC12 cells with TGN-pHluorin and comparing its steady state localization to TGN38 by immunofluorescence (Figure 5A). Next, we used live imaging to measure the fluorescence of our reporter expressed in WT and HID-1 KO PC12 cells under basal conditions. We incubated our cells with nigericin/monensin (Wu et al., 2000) to generate individual pH calibration curves for every cell that we imaged (Figure 5B). From the sigmoidal curves, we extrapolated absolute TGN pH values for WT and HID-1 KO cells. Strikingly, we found that the absence of HID-1 led to significant alkalinization of the TGN (Figure 5C). Importantly, this phenotype could be rescued by transient expression of HID-1-HA.

How does a peripheral membrane protein such as HID-1 influence TGN pH? The acidification of intracellular organelles depends primarily on the activity of the vacuolar H+-ATPase (V-ATPase), which consists of two multisubunit sectors (V1 and V0). The cytoplasmic V1 sector mediates ATP hydrolysis, whereas V0 assembles into a transmembrane pore involved in proton translocation (Forgac, 2007; Cipriano et al., 2008; Marshansky and Futai, 2008). PC12 cells express three isoforms (a1, a2, and a3) of the largest V0 subunit, and each isoform displays a particular subcellular localization with enrichment at specific organelles (Saw et al., 2011). As a2 accumulates at the Golgi in PC12 cells, this isoform is likely to contribute significantly to Golgi acidification. Thus, to test whether V-ATPases contribute to the phenotype caused by the lack of HID-1, we looked at the distribution of a2. For this, we relied on velocity sedimentation of PC12 membranes through sucrose to separate organelles based on their size, with the Golgi migrating at the bottom of the gradient under these conditions (Tooze and Huttner, 1990; Asensio et al., 2010). Using WT membranes, a2 displayed a bimodal distribution with a major peak at the bottom of the gradient, consistent with its reported Golgi localization (Figure 5D). Strikingly, the loss of HID-1 led to a dramatic redistribution of a2 to the top of the gradient, away from Golgi fractions (Figure 5D). We also monitored changes in pH following treatment with the V-ATPase inhibitor bafilomycin A1 (BafA1) (Yoshimori et al., 1991). Whereas BafA1 rapidly neutralized the TGN pH of WT cells, the drug had a more modest effect in HID-1 KO cells (Figure 5E). We thus conclude that HID-1 contributes to TGN acidification by controlling the proper accumulation and/or retention of a2 at the Golgi.

LDCV-soluble cargo aggregation is thought to be a key determinant for efficient sorting to the RSP. Indeed, treatment with drugs neutralizing intraluminal pH redirects adrenal corticotropin hormone or granins to the constitutive secretory pathway (Moore et al., 1983; Gerdes et al., 1989; Taupenot et al., 2005). Surprisingly, neither our biochemical secretion experiments nor our live-imaging experiments with NPY-pHluorin (Figure 1) revealed any increase in basal exocytosis, which would be expected from cargo being rerouted to the constitutive secretory pathway. Where does the soluble cargo go? Previous work has shown that reduced levels of LDCV-soluble cargo observed in hid-1 mutant worms could be partially rescued by genetically inhibiting lysosome biogenesis (Yu et al., 2011). To test whether increased lysosomal degradation is contributing to the depletion of SgII observed in HID-1 KO PC12 cells, we incubated our cells with protease inhibitors for 24 h to inhibit lysosomal hydrolases...
HID-1 influences the sorting of transmembrane large dense core vesicle cargo

Our data indicate that HID-1 influences sorting of soluble LDCV cargo by impairing its aggregation into a dense core, but does it contribute to sorting of transmembrane LDCV proteins as well? Similarly to chromaffin cells, PC12 cells store monoamines into LDCVs through the action of an integral membrane protein: the vesicular monoamine transporter (VMAT), which depends on a cytoplasmic dileucine-like motif for sorting to LDCVs (Liu et al., 1994; Erickson et al., 1995). Mutations within this motif cause an increase in VMAT cell surface delivery by diverting the transporter from the regulated to the constitutive secretory pathway (Li et al., 2005; Asensio et al., 2010). To test whether HID-1 influences the sorting of VMAT, we monitored its cell surface delivery using a flow cytometry assay that we previously developed (Asensio et al., 2010). For this, we transfected WT and HID-1 KO PC12 cells with HA-VMAT2-GFP, which contains a luminal HA tag to monitor surface level as well as a cytosolic GFP to determine total expression. After incubating the transfected cells with an anti-HA antibody conjugated to Alexa647, we measured GFP and Alexa647 fluorescence by flow cytometry. The cumulative frequency distribution of the ratio of surface (HA) to total (GFP) showed no difference between WT and HID-1 KO PC12 cells. As a positive control, we observed that WT cells transfected with EE/AA HA-VMAT2-GFP, a trafficking mutant that missorts to the plasma membrane (Li et al., 2005, Asensio et al., 2010), exhibited the expected shift in distribution (Figure 6A). VMAT2 is thus not being rerouted to the constitutive secretory pathway in the absence of HID-1.

This observation raises the intriguing possibility that VMAT2 might be able to sort onto LDCVs lacking a dense core. To test this directly, we transfected WT and HID-1 KO PC12 cells with HA-VMAT2-GFP, a trafficking mutant that missorts to the plasma membrane (Li et al., 2005, Asensio et al., 2010), and monitored individual exocytotic events using live-imaging (Figure 6B and Supplemental Movies S3 and S4). As expected, WT PC12 cells displayed very few events under basal conditions, but showed a massive response to depolarization. Consistent with our flow cytometry data, the number of non-stimulated exocytotic events in HID-1 KO cells remained unchanged, again suggesting that VMAT2 is not being rerouted to the constitutive secretory pathway by default in the absence of HID-1. However, we observed a dramatic reduction in the number of events triggered by depolarization, suggesting that VMAT2 does not sort into “empty” LDCVs in the absence of HID-1. As
a complementary approach, we developed an assay to assess monoamine secretion taking advantage of the false fluorescent neurotransmitter (FFN206), which is a VMAT substrate (Hu et al., 2013). After loading WT and HID-1 KO PC12 cells with FFN206, we measured the fluorescence of secreted and cellular fractions under basal and stimulated conditions using a plate reader. Similarly to the effect observed with SgII, we found that loss of HID-1 led to a decrease in basal storage as well as both basal and regulated secretion of FFN206. However, normalization of the secretion data to the cellular content showed no difference in regulated secretion per se (Figure 6, C–E). Altogether, these data indicate that the absence of HID-1 reduces the number of VMAT2-positive LDCVs available for regulated exocytosis, but also suggest that VMAT2 does not sort into “empty” LDCVs.

If the number of VMAT2-positive LDCVs available for release is reduced, but VMAT2 does not traffic to the plasma membrane constitutively, where does excess VMAT2 accumulate? To address this, we immunostained WT and HID-1 KO PC12 cells transfected with VMAT2-HA and determined the steady-state localization of the transporter. The images revealed VMAT2 enrichment in the perinuclear area, suggesting that the transporter might be retained in the Golgi area. Consistent with this, we observed an increase in the amount of VMAT2 overlapping with TGN38 (Figure 6F). Finally, we also determined the steady-state localization of an endogenous transmembrane LDCV cargo. Immunostaining for Syt1 showed that this marker similarly accumulates in the TGN area (Figure 6G).

**DISCUSSION**

These results establish that the peripheral membrane protein HID-1 is required for LDCV biogenesis from rat neuroendocrine PC12 cells. The absence of HID-1 leads to defects in both storage and secretion of SgII as well as monoamines. Our phenotype is consistent with earlier studies in worms showing a decrease in accumulation and release of neuropeptides (Mesa et al., 2011; Yu et al., 2011). More recently, it has been reported that conditional KO mice, in which HID-1 has been specifically inactivated from beta-cells of the pancreas, display impaired insulin secretion. This phenotype has been attributed to a defect in homotypic fusion of immature insulin granules (Du et al., 2016), a mechanism that is somewhat surprising considering that this fusion process had never previously been observed in this cell type. Although we cannot definitively rule out that this might contribute to our phenotype, we did not observe any obvious defects in LDCV homotypic fusion in our HID-1 KO PC12 cells by electron microscopy. It is also important to note that immature LDCVs can still undergo exocytosis in PC12 cells (Tooze et al., 1991). Thus, it is unlikely that our secretion defect is caused uniquely by a maturation defect. Instead, we suggest that HID-1 controls a step upstream of LDCV maturation, probably at the level of budding.
studies have demonstrated that pH values of 6.9 or above dramatically impair granin aggregation (Chanat and Huttner, 1991). Consistent with this, we observed a dramatic reduction in the total number of vesicles with a discernible dense core by electron microscopy. In addition, the few remaining LDCVs exhibited smaller cores and displayed abnormal biochemical properties with a shift of SgII toward lighter sucrose fractions after equilibrium sedimentation. Given the known significance of core formation for efficient sorting of soluble cargo to the RSP, we expected SgII to be missorted to the constitutive secretory pathway by default. Surprisingly, basal secretion of SgII remained unchanged in HID-1 KO cells, and instead, we found that a fraction of SgII was rerouted to the lysosome. We propose that lack of SgII aggregation leads to its dilution and diffusion throughout the endomembrane system (Figure 7). Alternatively, defective LDCVs might be targeted for degradation by crinophagy after budding from the TGN (Orci et al., 1984).

As the expression of HID-1 is restricted to specialized secretory cells (Wang et al., 2011), it raises the question of why HID-1 would be needed in some cell types but not others. Indeed, the existence of an acidic Golgi apparatus is not specific to secretory cells but rather is a general feature of eukaryotic cells. It is tempting to speculate measured using a plate reader. (E) Secreted FFN206 was expressed as a percentage of total FFN206 fluorescence. Statistical analysis was performed by one-way ANOVA followed by a post hoc Tukey test: **p < 0.01 relative to basal secretion from WT; ***p < 0.001 relative to stimulated secretion from WT (n = 8). The bar graphs indicate mean ± SEM. (F) WT and HID-1 KO PC12 cells were transfected with VMAT2-HA, fixed, and stained with HA and TGN38 antibodies followed by Alexa Fluor 488- and Alexa Fluor 647-conjugated secondary antibodies or (G) WT and HID-1 KO PC12 cells were transfected with TGN38-GFP, fixed, and stained with Syt1 antibody followed by Alexa Fluor 647-conjugated secondary antibody. The cells were imaged by spinning-disk confocal microscopy. The amount of HA (F) or Syt1 (G) immuno-reactivity overlapping with TGN38 was expressed as a percentage of total fluorescence: **p < 0.001 relative to WT (for Syt1, n = 20 cells for WT and n = 16 cells for KO; for VMAT2, n = 26 cells for WT and n = 38 cells for KO from two independent experiments). The scale bars indicate 5 μm. The data shown indicate mean ± SEM.

FIGURE 6: Loss of HID-1 leads to TGN enrichment of transmembrane LDCV cargoes. (A) WT and HID-1 KO PC12 cells were transfected with the indicated HA-VMAT2-GFP constructs, incubated with an HA antibody conjugated to Alexa Fluor 647 for 1 h, washed, and analyzed by flow cytometry to determine the fluorescence of individual cells. The ratios of surface (HA) to total (GFP) were computed and expressed as a cumulative frequency distribution. (B) WT and HID-1 KO PC12 cells were transfected with VMAT2-pHluorin and then imaged live by spinning disk confocal microscopy as described in Figure 1. Images show representative maximum-intensity time projections of 150 basal and stimulated frames. At the end of the experiment, cells were imaged in Tyrode’s solution containing 50 mM NH₄Cl, pH 7.4, to reveal total VMAT2-pHluorin fluorescence by alkalinization and identify transfected cells. Scale bar indicates 5 μm. Bar graph shows the number of exocytotic events per second normalized to cell surface area. Statistical analysis was performed by one-way ANOVA followed by a post hoc Tukey test: **p < 0.01 relative to stimulated secretion from WT (n = 20 cells for WT, n = 16 cells for KO from two independent experiments). The data shown indicate mean ± SEM. (C–E) PC12 cells were loaded with FFN206 (a vesicular monoamine transporter fluorescent substrate) and subjected to a secretion assay as described in Figure 1. Secreted (C) and cellular (D) fluorescence values were
that HID-1 acts as a gatekeeper to prevent a2 from leaking into the RSP (Figure 7). In PC12 cells, knockdown of a1, the isoform enriched on LDCVs, is not sufficient to impair LDCV acidification, but double knockdown of both a1 and a2 leads to alkalinized LDCVs, suggesting that a2 has the propensity to escape onto LDCVs (Saw et al., 2011). In the absence of an efficient retention mechanism, the continuous budding of LDCVs and other vesicles might effectively deplete a2 from the TGN. It will be interesting to test this in the future by assessing whether HID-1 is able to interact directly with a2 or whether it mediates its effect indirectly through another partner. Another explanation could be related to the biochemical properties of the secretory cargo that specialized secretory cells need to accommodate. Indeed, with a predicted isoelectric point of ~4.7, both SgII and chromogranin A are acidic proteins carrying a net negative charge at neutral pH. As these proteins are expressed at very high level in most specialized secretory cells, they might significantly increase the buffering capacity of the biosynthetic pathway of these cells, making it harder to build a pH gradient. These secretory cells might thus rely on some additional mechanisms to ensure efficient Golgi acidification, and HID-1 might play an important role by influencing a2 localization.

We have previously observed that knockdown of AP-3 or VPS41 leads to defects in SgII storage and regulated secretion, together with constitutive delivery of transmembrane LDCV cargoes to the cell surface, suggesting that these cytosolic factors recruit and concentrate membrane proteins onto LDCVs (Asensio et al., 2010, 2013). In striking contrast, transmembrane LDCV cargoes do not traffic constitutively to the plasma membrane in the absence of HID-1, but instead accumulate in a perinuclear compartment, where they partially overlap with TGN markers. In addition, although loss of HID-1 also results in defects in SgII storage, regulated secretion per se, when normalized to cellular stores, remains unchanged. Thus, the few LDCVs budding from the TGN of HID-1 KO cells must be fully competent for regulated exocytosis, and presumably have the proper membrane protein composition. This suggests that HID-1 does not influence the ability of the cytosolic sorting machinery to interact with transmembrane cargoes. So why do Syt1 and VMAT2 display TGN enrichment? We propose that inefficient core formation not only reduces the efficiency of soluble cargo sorting, but also influences the total number of budding vesicles. If transmembrane cargoes can still interact with the cytosolic sorting machinery, then reductions in the amount of budding would result in their accumulation at the TGN, with the cytosolic sorting machinery effectively preventing them from leaking into the constitutive secretory pathway.

**MATERIALS AND METHODS**

**Molecular biology**

The human codon-optimized Cas9 and chimeric guide RNA expression plasmid (pX330) developed by the Zhang lab (Cong et al., 2013) were obtained from Addgene. To generate gRNA plasmids, a pair of annealed oligos (20 base pairs) were ligated into the single guide RNA scaffold of pX330. The following gRNAs sequences were used: Forward1: 5′-CACCCTGACT-TCACGGGCTGCAG-3′; Reverse1: 5′-AAACCTGACCCCGCTGGAGGCCAAC-3′ to generate HID-1 KO#1; Forward2: 5′-CACCCTGACCCCGCTGGAGGCCAAC-3′; Reverse2: 5′-AAACAGAAAGCATTCGCTGGCCT-3′ to generate HID-1 KO#2. For the homologous directed repair template, tdTomato-polyA was inserted into pBSKII (+), and HID-1 homology arms were amplified by PCR and inserted 5′ and 3′ of tdTomato. The following primers were used to amplify the homologous arms: 5′ arm Forward: 5′-TCCCTGAAAA-GTGAATGGAGC-3′; 5′ arm Reverse: 5′-CTGAGGGGAATCGAGTAGAC-3′; 3′ arm Forward: 5′-CTTGTAGCACAGGGTGGCC-3′; 3′ arm Reverse: 5′-TGGTATCCCTCTTATTAAAGGATGC-3′. The human codon-optimized Cas9 and chimeric guide RNA expression plasmid (pX330) developed by the Zhang lab (Cong et al., 2013) were obtained from Addgene. To generate gRNA plasmids, a pair of annealed oligos (20 base pairs) were ligated into the single guide RNA scaffold of pX330. The following gRNAs sequences were used: Forward1: 5′-CACCCTGACT-TCACGGGCTGCAG-3′; Reverse1: 5′-AAACCTGACCCCGCTGGAGGCCAAC-3′ to generate HID-1 KO#1; Forward2: 5′-CACCCTGACCCCGCTGGAGGCCAAC-3′; Reverse2: 5′-AAACAGAAAGCATTCGCTGGCCT-3′ to generate HID-1 KO#2. For the homologous directed repair template, tdTomato-polyA was inserted into pBSKII (+), and HID-1 homology arms were amplified by PCR and inserted 5′ and 3′ of tdTomato. The following primers were used to amplify the homologous arms: 5′ arm Forward: 5′-TCCCTGAAAA-GTGAATGGAGC-3′; 5′ arm Reverse: 5′-CTGAGGGGAATCGAGTAGAC-3′; 3′ arm Forward: 5′-CTTGTAGCACAGGGTGGCC-3′; 3′ arm Reverse: 5′-TGGTATCCCTCTTATTAAAGGATGC-3′. The main isoform amplified from these cells corresponds to isoform #2, which has been reported for mouse (Uniprot: Q8R1F6-2) and for human (Uniprot: Q8IV36-2). The PCR products were then subcloned into FUGW. The following primers were used to genotype HID-1 KO cells: P1F: 5′-CGACAT-AACGGGGAGTCC-3′; P1R: 5′-GCCATTAGCAGCGATGGGACC-3′; P2F: 5′-GGCCATGCGAGCCTGACA-3′; P2R: 5′-CCAGTCCACTGGGATGC-3′; P3F: 5′-GTGAATCCCTGCTACTGATTCC-3′; P3R: 5′-CCCTGAGCACAGGGTGC-3′. To test for the presence of indels, primers P3F/P3R were used. The resulting PCR products were ligated into pBluescript II KS. Isolated plasmids from 24 random colonies were then analyzed for the presence of indels by sequencing.

**Cell culture and lentivirus production**

PC12 cells were maintained in DMEM supplemented with 10% horse serum and 5% calf serum under 5% CO2 at 37°C. Transfection of PC12 cells was performed using Lipofectamine 2000 (LifeTechnologies) or Fugene HD (Promega) according to the manufacturer’s instructions. Weekly, PC12 cells were maintained in DMEM supplemented with 10% horse serum and 5% calf serum under 5% CO2 at 37°C. Transfection of PC12 cells was performed using Lipofectamine 2000 (LifeTechnologies) or Fugene HD (Promega).
instructions. HEK293T cells were maintained in DMEM with 10% fetal bovine serum under 5% CO₂ at 37°C. Lentivirus was produced by transfecting HEK293T cells with FUGW, psPAX2, and pVSVG using Fugene HD according to the manufacturer’s instructions.

**Antibodies**

HA.11 mouse monoclonal antibody (mAb) was obtained from Covance (USA), HA (3F10) rat mAb from Roche, SgII rabbit antibody from Meridian Life Science (USA), TGN38 mouse mAb from BD Biosciences (USA), tubulin mouse mAb from Developmental Hybridoma Bank Studies (USA), synaptotagmin 1 mouse antibody from Synaptic Systems (Germany), synaptophysin (p38) mAb from Covance (USA), HID-1 mouse mAb from Novus Biologicals (USA), and goat anti-rabbit Alexa Fluor 647, anti-mouse Alexa Fluor 488, anti-mouse Alexa Fluor 647, and anti-rat Alexa Fluor 488 secondary antibodies from Molecular Probes (USA). a2 rabbit polyclonal antibody was a generous gift from Xiao-Song Xie (UT Southwestern). The following antibodies did not work by Western blot: HID-1 mouse mAb from Novus Biologicals (2F4), HID-1 mouse mAb from LifeSpan BioSciences (clone 2E9), HID-1 goat polyclonal from Santa Cruz (S-14).

**Secretion assays**

PC12 cells were plated on poly-l-lysine, washed, and incubated in Tyrode’s buffer containing 2.5 mM K⁺ (basal) or 90 mM K⁺ (stimulated) for 30 min at 37°C. The supernatant was then collected, cell lysates prepared as previously described (Asensio et al., 2010), and the samples analyzed by quantitative fluorescence immunoblotting. For FFN206 assays, PC12 cells were preloaded with 1 μM FFN206 in Tyrode’s buffer for 45 min. Following incubation, cells were washed twice with Tyrode’s buffer and subjected to a secretion assay as described above. Supernatant and lysates were loaded on 96-well plates and fluorescence was read using a plate reader (excitation: 369 nm; emission: 464 nm).

**Immunofluorescence and confocal microscopy**

PC12 cells were rinsed with PBS and fixed in 4% paraformaldehyde in PBS and incubated for 20 min at room temperature. Cells were permeabilized in PBS containing 0.1% Triton X-100 for 10 min at room temperature and blocked in PBS containing 2% BSA, 1% fish skin gelatin, and 0.02% saponin. Primary antibodies were diluted in blocking solution at 1:1000 (SgII), 1:1000 (TGN38), 1:1000 (Syt1), 1:500 (HA)m and 1:500 (HA.i1). The secondary goat anti-rabbit antibodies were diluted in blocking solution at 1:1000. Images were acquired using an Olympus Fluoview scanning confocal microscope and a 63× oil objective (NA 1.42) at a resolution of 512 × 512 pixels with a sampling speed of 12.5 μs/pixel with Kalman filter (integration count 5).

**Density gradient fractionation**

Equilibrium sedimentation through sucrose was performed as previously described (Asensio et al., 2010, 2013). Briefly, a postnuclear supernatant was prepared from PC12 cells by homogenization with a barbed bearing device (12 μm clearance), loaded onto a 0.6–1.6 M continuous sucrose gradient, and sedimented at 30,000 rpm in an SW41 rotor for 14–16 h at 4°C. Fractions (~750 μl each) were collected from the top and were analyzed by quantitative fluorescence immunoblotting using an FX Imager (BioRad) or a Fluochem R (ProteinSimple). Velocity sedimentation through sucrose was performed as described above, with postnuclear supernatant loaded onto a 0.3–1.2 M continuous sucrose gradient and sedimented at 25,000 rpm for 19 min. Fractions (1 ml) were collected from the top and analyzed as above.

**Electron microscopy**

PC12 cells were plated onto Aclar film disks coated with poly-l-lysine, fixed with 2.5% glutaraldehyde, 2% paraformaldehyde, and 2 mM calcium chloride in 0.15 M cacodylate buffer (pH 7.4). Coverslips were rinsed and fixed in a mixture of 2.5% glutaraldehyde and 2% paraformaldehyde in 0.15 M cacodylate buffer at pH 7.4 with 2 mM calcium chloride warmed to 37°C for 5 min in an incubator. Subsequently, samples were transferred to a refrigerator at 4°C in the same fixative solution until they were ready to be processed. Once ready to be processed, each coverslip was rinsed in 0.15 M cacodylate buffer three times for 10 min each and subjected to a secondary fixation step for 1 h in 1% osmium tetroxide/0.3% potassium ferrocyanide in cacodylate buffer on ice. Following this, samples were washed in ultrapure water three times for 10 min each and stained en bloc for 1 h with 2% aqueous uranyl acetate. After staining was complete, samples were briefly washed in ultrapure water, dehydrated in a graded acetone series (50%, 70%, 90%, 100%, 100%) for 10 min in each step, infiltrated with microwave assistance (Pelco BioWave Pro, Redding, CA) into LX112 resin, and flat embedded between two slides that had been coated with PTFE release agent (Miller-Stephenson MS-143XD; Danbury, CT). Samples were cured in an oven at 60°C for 48 h. Once the resin was cured, the slides were separated and the Aclar coverslips were peeled off. A small region was excised and glued onto a blank stub with epoxy. Thin sections (70 nm) were taken and imaged on an FE-SEM (Zeiss Crossbeam 540, Oberkochen, Germany) using the aSTEM detector. The SEM was operated at 28 KeV and probe current 0.9 nA, and the STEM detector was operated with the annular rings inverted for additional image contrast. Whole cells were imaged at 6114 × 4608 pixels with a resolution of 3.722 nm/pixel from random sections. Zoomed regions were imaged at 2048 × 1536 pixels with a resolution of 1.861 nm/pixel. Images were analyzed with ImageJ. Morphologically identifiable LDCVs were counted per cell section and performed blind to the conditions of the experiments.

**Lysosomal inhibition**

PC12 cells were incubated for 24 h in complete medium supplemented with vehicle or a cocktail of lysosomal protease inhibitors (Sigma) including 10 μM antipain, 10 μM leupeptin, and 5 μM pepstatin A. Cells were washed on ice with cold PBS and lysed by the addition of 50 mM Tris-HCl, pH 8.0, 150 mM NaCl, 1% Triton X-100, and SIGMAFAST protease inhibitor cocktail (Sigma-Aldrich) plus 10 mM EDTA and 1 mM PMSF. Samples were analyzed by quantitative fluorescence immunoblotting.

**EGF degradation assay**

WT and HID-1 KO PC12 cells were washed twice with PBS and starved of serum for 2 h in DMEM with 0.1% BSA (GoldBio). During starvation, EGF-biotin (GoldBio) streptavidin-647 conjugate was prepared. EGF-biotin (5 μg/ml) was incubated for 30 min at 4°C at a 5:1 ratio to streptavidin-Alexa647 (Life Technologies). Following starvation, cells were washed twice with ice-cold PBS on ice and incubated with EGF-A647 conjugate at a final concentration of 100 ng/ml for 1 h on ice. Excess unbound EGF was removed by washing with ice-cold PBS with 0.5% BSA. Cells were chased for indicated times before fixation with 4% PFA in PBS for 20 min at room temperature. Cells were analyzed by flow cytometry (CyAn ADP Analyzer, Beckman Coulter, USA) or by scanning confocal microscopy.

**Spinning disk confocal live imaging**

PC12 cells were co-transfected with NPY-phluorin or VMAT2-phluorin together with BDNF-mCherry. At 1 d after transfection,
cells were transferred to poly-l-lysine coated 22 mm glass coverslips. After an additional 2 d, cells were washed once with Tyrode’s buffer and coverslips were transferred to an open imaging chamber (Life Technologies). Cells were imaged close to the coverslips, focusing on the plasma membrane (determined by the presence of BDNF-mCherry positive plasma membrane docked-vesicles), using a custom-built Nikon spinning disk at a resolution of 512 x 512 pixels. Images were collected for 100 ms at 10 Hz at room temperature with a 63x objective (Oil Plan Apo NA 1.49) and an ImageEM X2 EM-CCD camera (Hamamatsu, Japan). Following baseline data collection (15 s), an equal volume of Tyrode’s buffer containing 90 mM KCl was added to stimulate secretion and cells were imaged for an additional 30 s. At the end of the experiment, cells were incubated with Tyrode’s solution containing 50 mM NH₄Cl, pH 7.4, to reveal total fluorescence and to confirm that the imaged cells were indeed transfected. Movies were acquired in MicroManager (UCSF) and exported as tiff files. To automatically detect newly appearing exocytic events within a cell, difference images were constructed between the averages of adjacent pairs of frames in a given movie, that is, mean (n + 2, n + 3) – mean (n, n + 1), where n is any frame between the first and last frame minus 3. Positive differences in intensity between the averaged frames were taken to be candidate events. The difference images were then passed through a Gaussian filter to reduce image noise. Transfected cells were cropped from each movie by hand and analyzed individually. Event detection was performed using a wavelet-based method as previously described (Olivo-Marin, 2002; Jaqaman et al., 2008). Noise events were filtered from the true events on the basis of intensity and the stipulation that events occur within the confines of the cell, and not overlap with previously counted events. A convex hull around the remaining events was generated to approximate cell area and to find the density of events. Cell activity could then be assessed by producing the cumulative sum of events produced by a cell over the course of the movie. All image analysis was performed using MATLAB and the MATLAB image processing toolbox.

pH imaging
PC12 cells were transfected with TGN-pHluorin. At 1 d after transfection, cells were transfected to poly-l-lysine-coated 22-mm glass coverslips. After an additional 2 d, cells were washed once with Tyrode’s buffer (pH 7.4) and imaged using a Zeiss Axiovert 5100 TV widefield microscope. Images were collected with 100 ms exposure at a resolution of 512 x 512 pixels at room temperature with a 40x objective (NA 1.30) and a CoolSNAP HQ2 camera (Photometrics, USA). Cells were then perfused with enriched KCl buffer supplemented with 5 μM nigericin (Sigma-Aldrich) and 5 nM monensin (Sigma-Aldrich) at pH 8.5 and incubated for 10 min before image acquisition. This process was repeated in pH increments of 0.5 down to pH 5.5. For the V-ATPase inhibition experiments, cells were perfused with Tyrode’s buffer (pH 7.4) supplemented with 200 nM BafA1 (Sigma Aldrich) and imaged for the indicated amount of time before the generation of the calibration curve as described above.

Quantitative PCR
RNA was isolated from HID-1 KO or WT PC12 cells with the E.Z.N.A total RNA isolation kit (Omega); then isolated RNA was DNase treated with Turbo DNase (Ambion). A quantity of 2 μg of total RNA was reverse transcribed (SuperScriptIV, Thermo) and subjected to triplicate qPCR for the last exon–exon junction of ActB and SgII transcripts. qPCR was performed using SYBR Green qPCR Master Mix (BioRad) and a BioRad iQ5 real-time PCR machine (BioRad) with gene-specific primers. The results were normalized to expression of the housekeeping gene ActB. SgII forward primer: CCTACTTGAGAAGGATTTTGCG. SgII Reverse primer: ACCAACCCTATTGGTTCTCT. ActB Forward primer: CCTAGACCAGTGAAGATCActB. ActB Reverse primer: GATAGAGCCACCAATCCAC.

Statistics
Unless indicated otherwise, all statistical analysis was performed using the two-tailed Student’s t test. Statistical analyses were conducted using Excel or Prism.

Figure preparation
Images were processed using ImageJ; any changes in brightness and contrast were identical between samples meant for comparison.

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REFERENCES


Figure S1. Generation of HID-1 KO PC12 cells using CRISPR/Cas9

(A) Illustration of the strategy used to generate HID-1 KO PC12 cells, relying on homologous recombination repair after cleavage by Cas9 to knock-in a fluorescent protein (tdTomato), while effectively knocking-out HID-1. The repair template was designed in such a way that the fluorescent reporter should not be expressed without proper, in-frame recombination. (B) Schematic illustrating the sequence of the homology arms flanking tdTomato in the repair construct. (C) Genotyping PCR was performed from genomic DNA isolated from WT and HID-1 KO cells using the indicated primers. The location of primers with regard to the homology arms is shown in panel A. (D) The subpopulation of PCR products corresponding to alleles without integrated tdTomato were ligated into pBlueScript II. Plasmids were isolated from twenty-four random colonies and analyzed for the presence of indels by sequencing. (E) Predicted protein sequences associated for the five types of identified indels.
Figure S2. HID-1 KO PC12 cells generated using an independent gRNA show similar defects in SgII storage and secretion

HID-1 KO PC12 cells generated with an independent gRNA (KO#2) were plated and subjected to a secretion assay as described in Figure 1. Cellular and secreted secretogranin II (SgII) were measured by quantitative fluorescent immunoblotting with the secreted SgII normalized to actin (B) or total SgII (C) and expressed as percent of basal secretion in the control. The cellular SgII content was normalized to actin (A). Multiple comparison statistical analysis was performed by one-way ANOVA followed by posthoc Tukey test. ***, p< 0.001 relative to stimulated secretion from WT (n=4).
Figure S3. Loss of HID-1 does not reduce SgII mRNA

Quantitative RT-PCR analysis of WT and HID-1 KO PC12 cells transcripts shows no change in SgII expression. n=3.
WT and HID-1 KO PC12 cells were transfected with ANF-GFP and subjected to a secretion assay as described in Figure 1. Cellular and secreted ANF-GFP were measured with a plate reader and expressed as percent of basal control values (A,B). Alternatively, secretion was normalized to total ANF-GFP (C). Multiple comparison statistical analysis was performed by one-way ANOVA followed by posthoc Tukey test. *, p < 0.05 relative to WT; **, p < 0.01 relative to stimulated secretion from WT, ***, p < 0.001 relative to stimulated secretion from WT (n=7).
Figure S5. Lysosomal inhibition partially restores the levels of SgII in HID-1 KO cells

WT and HID-1 KO PC12 cells were incubated where indicated with protease inhibitors for 24 hours. Cellular SgII was quantified by fluorescent western analysis and the values normalized to tubulin immunoreactivity. *, p< 0.05 relative to KO; (n=3). The data shown indicate mean ± s.e.m.