

RESEARCH ARTICLE

ROCK-nmMyoII, Notch and *Neurog3* gene-dosage link epithelial morphogenesis with cell fate in the pancreatic endocrine-progenitor niche

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ABSTRACT

During mouse pancreas organogenesis, endocrine cells are born from progenitors residing in an epithelial plexus niche. After a period in a lineage-primed *Neurog3*^{LO} state, progenitors become endocrine committed via upregulation of *Neurog3*. We find that the *Neurog3*^{LO} to *Neurog3*^{HI} transition is associated with distinct stages of an epithelial egression process: narrowing the apical surface of the cell, basalward cell movement and eventual cell-rear detachment from the apical lumen surface to allow clustering as nascent islets under the basement membrane. Apical narrowing, basalward movement and *Neurog3* transcriptional upregulation still occur without *Neurog3* protein, suggesting that morphogenetic cues deployed within the plexus initiate endocrine commitment upstream or independently of *Neurog3*. *Neurog3* is required for cell-rear detachment and complete endocrine-cell birth. The ROCK-nmMyoII pathway coordinates epithelial-cell morphogenesis and the progression through *Neurog3*-expressing states. NmMyoII is necessary for apical narrowing, basalward cell displacement and *Neurog3* upregulation, but all three are limited by ROCK activity. We propose that ROCK-nmMyoII activity, *Neurog3* gene-dose and Notch signaling integrate endocrine fate allocation with epithelial plexus growth and morphogenesis, representing a feedback control circuit that coordinates morphogenesis with lineage diversification in the endocrine-birth niche.

KEY WORDS: Plexus, Organogenesis, Morphogenesis, Differentiation, Progenitor, Niche

INTRODUCTION

In mouse pancreatogenesis, tissue growth, morphogenesis and differentiation must be coordinated, although how remains largely unresolved. We have reported that bipotent duct and endocrine progenitors exist within an epithelial niche called the ‘plexus state’ (Bankaitis et al., 2015), which comprises a web-like epithelium arising around E12.5 within an anlage of multipotent progenitors (MPC) (Kesavan et al., 2009; Hick et al., 2009; Villasenor et al., 2010). From E12.5–18.5, the plexus undergoes rapid expansion and remodeling, very different from classical branching morphogenesis, concurrent with a prolonged wave of endocrine-cell birth (Pan and

Wright, 2011). The progressive replacement of plexus by an arbor-like ductal epithelium is associated with a great reduction in the production of new endocrine cells (Kopp et al., 2011; Bankaitis et al., 2015). Here, we investigate the coordination of plexus morphogenesis and endocrine cell-fate allocation.

Pancreatic endocrine cells derive from a Notch-responsive Sox9⁺ progenitor pool (Apelqvist et al., 1999; Kopp et al., 2011; Serup, 2012) via activation of the endocrine-lineage determinant neurogenin 3 (*Neurog3*; Gradwohl et al., 2000; Schwitzgebel et al., 2000; Gu et al., 2002). *Neurog3* is necessary and sufficient for endocrine-cell birth (Gradwohl et al., 2000; Johansson et al., 2007). Cells with high-level *Neurog3* (*Neurog3*^{HI}) exit the cell cycle (Miyatsuka et al., 2011), egress from the epithelium (Gouzzi et al., 2011) and cluster adjacent to the forming nascent islets of Langerhans. Lineage tracing in *Neurog3*-null and hypomorphic mice (Wang et al., 2010; Shih et al., 2012) showed that preventing acquisition of the *Neurog3*^{HI} state causes cells to remain intra-epithelial and to differentiate into mature acinar and ductal fates. The mechanisms regulating progression of epithelial progenitors through various *Neurog3* expression states are poorly defined.

Non-autonomous feedback from *Neurog3*-expressing cells regulates numerous processes in forming duct versus endocrine tissues. Inactivating *Neurog3* greatly expands low-level *Neurog3* (*Neurog3*^{LO}) transcription throughout the Sox9⁺ epithelium and causes the loss of *Neurog3*^{HI} cells (Wang et al., 2010; Bechard et al., 2016). *Neurog3* loss causes a widespread reduction in epithelial Notch-pathway activity (Magenheim et al., 2011; Shih et al., 2012; Qu et al., 2013). Thus, *Neurog3*^{HI} cells normally induce lateral inhibition of *Neurog3* expression in surrounding Sox9⁺ progenitors to balance endocrine differentiation with progenitor maintenance (Afelik et al. 2012; Qu et al., 2013; Apelqvist et al., 1999; Murtaugh et al., 2003). *Neurog3* deficiency causes a persistent reduction in Sox9⁺ cell replication, suggesting additional functions in supporting replicative growth of progenitor epithelium (Bankaitis et al., 2015). *Neurog3* loss results in a dysmorphic plexus that is precociously transformed into more-mature epithelial duct and branched states, again suggesting that egressing endocrine cells modulate normal morphogenesis by maintaining their parental ‘plexus niche’ (Magenheim et al., 2011; Bankaitis et al., 2015). Collectively, these studies position *Neurog3*-expressing cells as a coordination nexus for progenitor maintenance, cell-fate allocation and plexus morphogenesis, to regulate growth and assembly of the ductal network with formation of a dispersed endocrine pancreas (Rieck et al., 2012; Bankaitis et al., 2015).

In epithelia, the molecular motor nmMyoII acts on filamentous-actin (F-actin) substrates in apical constriction, tissue folding, cell delamination and migration, and generation of inter- and intra-cellular tension (Gorfinkiel and Blanchard, 2011; Heisenberg and

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Bellaïche, 2013; Martin and Goldstein, 2014). Rho-associated kinase (ROCK) positively regulates nmMyoII activity (Wang et al., 2009), but also functions as a modulator of actin remodeling (Watanabe et al., 1999; Maekawa et al., 1999; Reinto and Ridley, 2003), affecting tissue and cell-shape change, cell polarization and migration (Li et al., 2005; Nakayama et al., 2005, 2008). ROCK-nmMyoII activities influence many complex morphogenetic and intracellular signaling processes (Amano et al., 2010). Rho-GTPase signaling, an upstream ROCK/nmMyoII regulator, influences neural-crest specification in frog embryos (Kim et al., 2015), and specific cell-fate decisions in mesenchymal cells (Sordella et al., 2003; McBeath et al., 2004). These reports clearly point to integration between morphogenesis signaling and the intrinsic gene-regulatory networks that drive cell fate.

We report here on the mechanistic interconnections between epithelial morphogenesis and fate in the pancreatic plexus niche. We use three-dimensional tissue reconstructions, gene-reporter analyses and genetic and pharmacological interventions to study interdependencies between *Neurog3* gene activity, Notch signaling and ROCK nmMyoII-controlled epithelial-cell morphogenesis. We propose that sequential, dissociable steps in endocrine fate allocation are mediated by morphogenetic changes at an apical versus basal cell surface. *Neurog3*^{LO} expression initiates concurrently with apical surface contraction, and becomes upregulated as its apical surface becomes increasingly focalized and the cell begins basalward movement. Initial contraction, focalization and *Neurog3* upregulation occur in the absence of *Neurog3* protein, suggesting that endocrine specification and entry to commitment occur via epithelium-intrinsic inputs upstream or independent of *Neurog3*. nmMyoII and ROCK oppositely regulate apical narrowing, focalization and basalward cell movement, and thus acquisition of the *Neurog3*^{HI} state. We propose a model in which a circuit comprising ROCK-nmMyoII, *Neurog3* gene dosage and Notch signaling balances apportionment of endocrine cells from the plexus while enabling proper growth and morphogenesis of the pancreatic epithelium.

RESULTS

Morphological transitions of the F-actin⁺ apical cortex are associated with cell-fate determination

A prominent feature in polarized epithelial cells is a belt of filamentous actin (F-actin) circumscribing the sub-apical cell cortex (Martin and Goldstein, 2014). These belt-like structures (hereafter F-actin^{BELT}) are closely apposed to tight and adherens junctions, and are important in mediating remodeling processes such as apical constriction, tissue folding, cell intercalation and epithelial egress or extrusion, among others (Heisenberg and Bellaïche, 2013). To probe whether specific cell-shape changes are associated with duct versus endocrine cell-fate decisions, we compared the F-actin^{BELT} topologies in cells located within the endocrine progenitor-rich plexus, the endocrine progenitor-poor duct state and in cells expressing *Neurog3* using an EGFP knock-in null allele (Lee et al., 2001). For the plexus, confocal z-stack reconstructions showed an intense multicellular meshwork of cortical F-actin closely associated with the Muc1⁺ apical lumen surface (Fig. 1A). The F-actin^{BELT} was located nearby the apical ZO1⁺ tight junctions, near the apical-most extent of Ecad⁺ adherens junctions, and apical with respect to the nuclei of all epithelial cells (Fig. S1A-H' and data not shown), as expected for F-actin^{BELT} in most epithelia. Plotting the F-actin^{BELT} aspect ratio against F-actin^{BELT} perimeter length from many individual cells served as a metric of apical surface shape (Fig. S2A-B) and revealed a spectrum of apical surface shapes in

Neurog3^{+/+} and heterozygous-null *Neurog3*^{EGFP/+} plexus (Fig. 1D). This spectrum in the plexus was similar throughout mid-to-late gestation, indicating that the overall mix of topologies is maintained during the major wave of cellular differentiation (Fig. 1E). Also in the plexus, numerous intense F-actin puncta were located directly at the Muc1⁺aPKC⁺ luminal surface, in a highly focalized pattern (hereafter, F-actin^{FOCAL}), signifying tightly narrowed apical contacts or tethering points (Bechard et al., 2016) of egressing *Neurog3*-expressing cells (Fig. 1B,C; Fig. S3A-D). Essentially all Muc1⁺ cells expressing *Neurog3*^{EGFP} exhibited a narrowed F-actin^{BELT} or F-actin^{FOCAL} signal (Fig. 1F). In contrast, the F-actin^{BELT} of essentially all cells in the duct state exhibited a larger perimeter and decreased aspect ratio (Fig. 1G-J). Scoring E-cad⁺ cell boundaries near the apical surface of these cells confirmed a large apical domain (Fig. S4A-D) and cell-flattening phenotype consistent with ductal differentiation (Grapin-Botton, 2005). In duct-state epithelium, *Neurog3*-reporting cells and F-actin^{FOCAL} structures were reduced or absent compared with plexus, consistent with a large reduction in *Neurog3*⁺ cell numbers (Bankaitis et al., 2015). EGFP-expressing cells without Muc1⁺ apical lumen-contact were considered to have exited the epithelium as endocrine-committed cells, and were not investigated further. These analyses suggest that F-actin^{BELT} narrowing and F-actin^{FOCAL} formation are associated with activation of *Neurog3* expression, while the larger F-actin^{BELT} shapes are associated with ductal or non-endocrine cell fates.

Morphogenetic influences mediate endocrine commitment in the plexus upstream of *Neurog3* protein

In E14.5 *Neurog3*^{NULL} (*Neurog3*^{EGFP/EGFP}) epithelium, *Neurog3*^{HI} (EGFP^{HI}) cells are lost, *Neurog3*^{LO} (EGFP^{LO}) cell numbers are increased, and the epithelial plexus is dysmorphic (Wang et al., 2010; Magenheim et al., 2011; Bankaitis et al., 2015; Bechard et al., 2016). Accordingly, we observed a dramatic loss of Muc1⁺F-actin^{FOCAL} structures, and a shift in apical F-actin^{BELT} shapes toward a more uniform circular morphology relative to *Neurog3*^{+/+} and *Neurog3*^{EGFP/+} samples (Fig. 2A-D,I). In addition to the previously reported 'epithelial fattening' observed in the *Neurog3*^{NULL} state, we also detected increased numbers of Sox9⁺ nuclei per unit of Muc1⁺ epithelium (Fig. S5A,B), consistent with a relative cell-crowding effect caused by the lack of egress of endocrine-committed cells. Although not establishing cause and effect, these data show that multiple morphological features of the plexus state, including apical F-actin^{FOCAL} structures formed in *Neurog3*^{HI} cells, are altered or lost in the absence of *Neurog3* function.

We recently reported that the abnormal E14.5 *Neurog3*^{EGFP/EGFP}-null epithelium undergoes a delayed morphological compensation during late gestation (E16.5-18.5) to adopt an epithelial branched structure highly similar in gross morphology to controls (Bankaitis et al., 2015). This 'corrective remodeling' process provides a stringent model to assess whether *Neurog3* promoter activity, which is reduced and expanded across the abnormal E14.5 *Neurog3*^{EGFP/EGFP} plexus, can be stimulated by *Neurog3*-independent cues that regulate the late-stage remodeling of the plexus to a more regular epithelial tree. Numerous inspections of E17.5 and E18.5 thick-sectioned and Muc1-labeled *Neurog3*^{EGFP/EGFP} pancreata confirmed epithelial remodeling to the grossly wild-type relative proportions of plexus, duct and branched states (Fig. S6A-D). The number of Sox9⁺ nuclei per linear unit of Muc1⁺ epithelium was comparable in E17.5 control and *Neurog3*^{EGFP/EGFP} plexus (Fig. S5B), in contrast to the increased cell density in E14.5

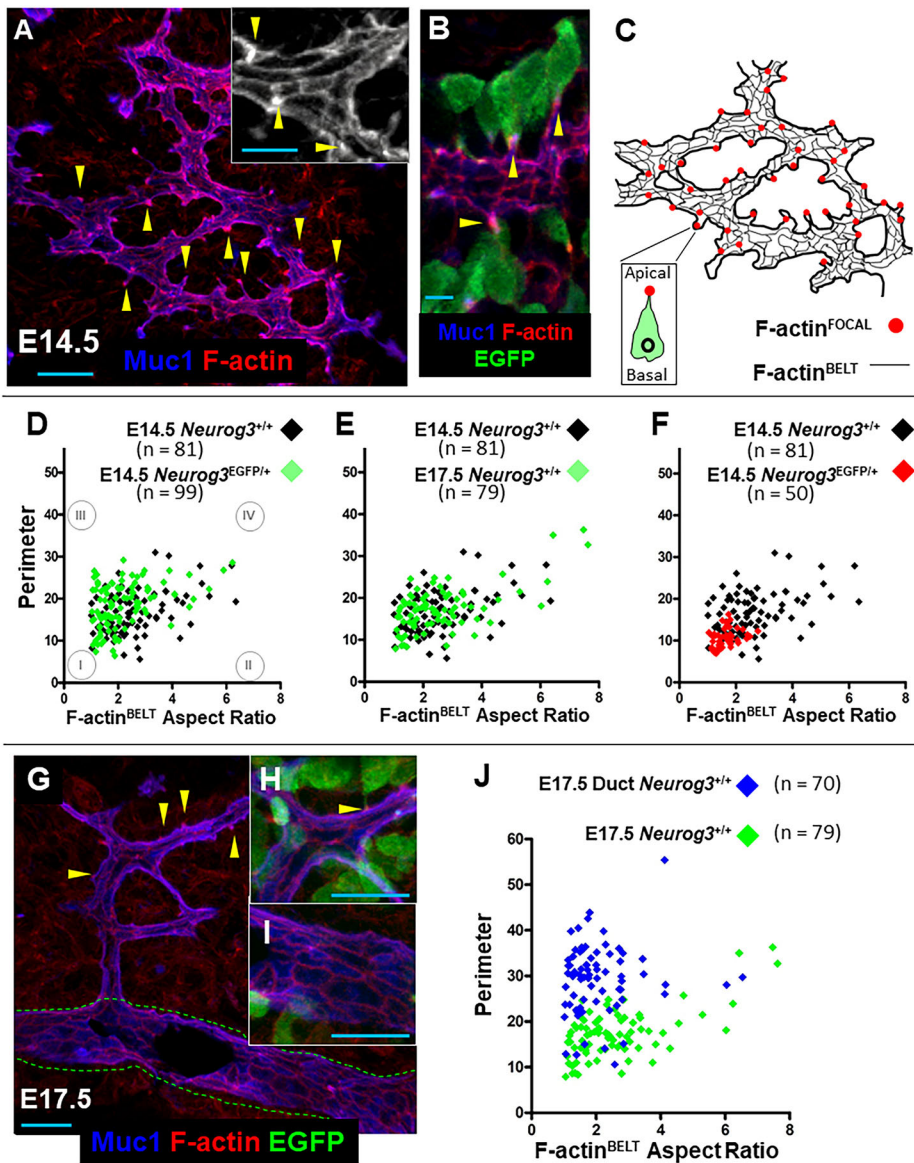


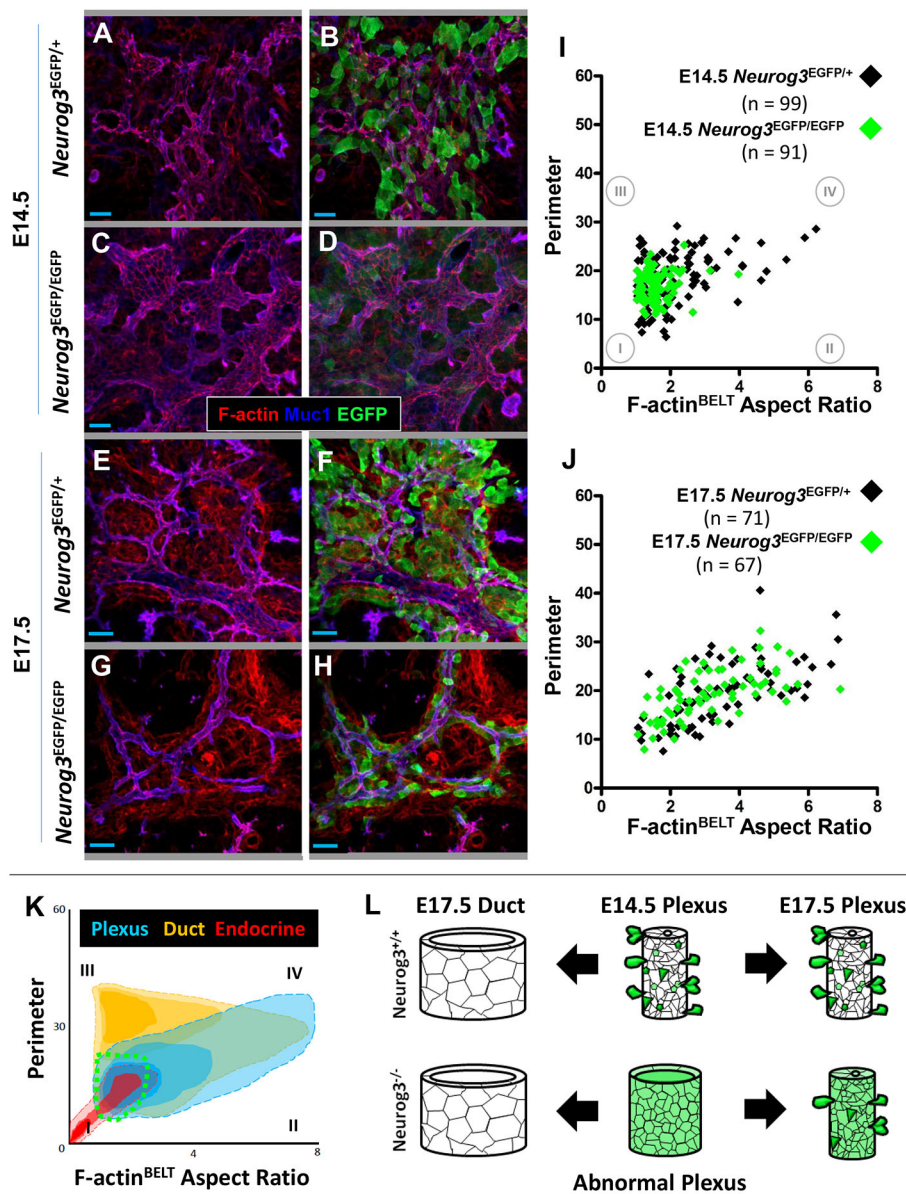
Fig. 1. Duct versus endocrine differentiation is associated with apical expansion or narrowing of the F-actin⁺ cell cortex.

(A) Confocal z-stack (30 μm depth) of E14.5 dorsal pancreas showing plexus labeled with Muc1 and F-actin. Inset shows cortical F-actin (F-actin^{BELT}) in individual epithelial cells. (B) F-actin^{FOCAL} marks the apical-lumen contacts of egressing *Neurog3*^{EGFP/+} EGFP⁺ cells. (C) Manual trace of F-actin^{BELT} meshwork (thin black lines) from A showing F-actin^{FOCAL} structures (red dots) and *Neurog3*-expressing cells (inset). (D) Cortical F-actin^{BELT} aspect ratio versus perimeter of n cells (at least three separate samples) for control or *Neurog3*^{EGFP/+} cells randomly selected from E14.5 plexus. I, narrowed F-actin^{BELT}; II, elongated F-actin^{BELT}; III, expanded F-actin^{BELT}; IV, expanded and elongated F-actin^{BELT}. These indicators apply to E, F and J. (E) F-actin^{BELT} distributions from E14.5 and E17.5 *Neurog3*^{+/+} plexus. (F) F-actin^{BELT} shape spectrum from E14.5 wild-type plexus and apically narrowed *Neurog3*-expressing cells from *Neurog3*^{EGFP/+} plexus. (G) Confocal z-stack (30 μm) of representative plexus and duct states at E17.5; duct state indicated by dashed green line. Yellow arrowheads indicate F-actin^{FOCAL} structures in A, B and G. (H,I) Enlargements of the plexus and duct state in G. F-actin^{BELT} is expanded and there is a lack of F-actin^{FOCAL} in the duct state. (J) F-actin^{BELT} shape spectrum from E17.5 plexus and duct states. Scale bars: 20 μm in A, G-I; 5 μm in B.

Neurog3^{EGFP/EGFP} plexus. Moreover, the spectrum of apical surface shapes within the E17.5 plexus of *Neurog3*^{EGFP/EGFP} samples was essentially indistinguishable from controls (Fig. 2E,G,J). Thus, multiple aspects of the abnormal morphogenesis in *Neurog3*^{NULL} tissue at E14.5 were corrected by late gestation. Surprisingly, such corrected plexus also displayed periodically spaced *Neurog3*^{EGFP-HI} cells (Fig. 2F,H), with narrowed F-actin^{BELT} structures at their apical surface (not shown). Although the EGFP^{HI} cell number in late-stage *Neurog3*^{EGFP/EGFP} plexus was reduced compared with control (quantified below), these data indicate that defects in *Neurog3* transcriptional upregulation are substantially bypassed, independently of *Neurog3* protein function, when the *Neurog3*^{EGFP/EGFP} plexus undergoes corrective remodeling. This effect was essentially specific for cells in the plexus state, because very few EGFP^{HI} cells were observed in non-plexus ductal or branched states, which is the case for normal ductal and branched state (Fig. 5C-D; Fig. S6A-D). These data suggest that morphogenetic cues deployed within a ‘normal’ remodeling plexus can profoundly influence *Neurog3* levels, and that some initial steps in endocrine-cell commitment are initiated in part through *Neurog3*-independent pathways (Fig. 2K,L).

Apical narrowing, focalization and basalward cell displacement mark the endocrine-committed state

To study how particular morphogenetic inputs could be linked to the progression of cells through *Neurog3*-expressing states, we focused on the morphological alterations occurring as epithelial (Muc1⁺) *Neurog3* expression is activated and upregulated during endocrine-cell birth. In E14.5 *Neurog3*^{EGFP/+} pancreata, we observed Muc1⁺EGFP^{LO} cells (6.5% of Muc1⁺EGFP⁺ cells, $n=95$; Fig. S7A,C) without a narrowed or focalized apical surface (Fig. 3A,A’). Another 19.3% of Muc1⁺EGFP⁺ cells showed a narrowed but not focalized apical surface, and all were EGFP^{HI} (Fig. 3B,B’). The majority (74.2%) of Muc1⁺EGFP⁺ cells exhibited clear apical focalization, and again all were EGFP^{HI}. The mean *Neurog3*^{EGFP} intensity of apically non-narrowed, narrowed and focalized Muc1⁺EGFP⁺ cells indicated a strong link between narrowing and focalization and the EGFP^{HI} state (Fig. 3E; Fig. S7C). The majority of Muc1⁺EGFP^{HI} cells with a focalized apical surface also showed a basalward displacement of the cell body – assessed as a displaced nuclear location relative to the Muc1⁺ epithelial plane – and indications of basal-surface ruffling and protrusive distortions shown in our previous publications to



represent filopodial migratory activity at the basal leading edge of the cell (Fig. 3C,C'; Fig. S7A) (Bechard et al., 2016; Löf-Öhlin et al., 2017). Progressive basalward movement was associated with elongation of the Muc1⁺F-actin^{FOCAL} structures, and an eventual detachment from the luminal surface (Fig. 3D,D'), signifying completion of cell egress from the epithelium. A *Neurog3*^{BAC} transgenic reporter line *Neurog3*^{RG1} (hereafter *Neurog3*^{RG}) (Bechard et al., 2016) independently confirmed the association of apical narrowing, focalization and basalward cell displacement with acquisition of the *Neurog3*^{HI} state (Fig. 3F-I'; Fig. S7B-D'). These results suggest that *Neurog3* upregulation within the plexus occurs concomitantly with a finely resolved sequence of events beginning with apical narrowing, then F-actin^{FOCAL} formation and basalward cell movement during endocrine-cell birth (Fig. 3E,J).

Neurog3-independent and Neurog3-dependent regulation of the endocrine-cell birth process

Although *Neurog3*^{EGFP/EGFP} (null) pancreata lack differentiated endocrine cells (Gradwohl et al., 2000; Shih et al., 2012), the late-stage correctively remodeling *Neurog3*^{EGFP/EGFP} plexus

contained many *Neurog3*^{EGFP-HI} cells. To define the stage in endocrine-cell birth when *Neurog3*-deficient cells fail, we compared the progression of *Neurog3*^{EGFP/+} and *Neurog3*^{EGFP/EGFP} Muc1⁺EGFP⁺ cells through apical narrowing, F-actin^{FOCAL} formation and basalward movement. In E14.5 *Neurog3*^{EGFP/EGFP} pancreata, cells exhibited a uniformly circular and narrowed apical surface compared with heterozygous controls, with essentially all being EGFP^{LO} (Fig. 3K,M). In the late-stage 'corrected' *Neurog3*^{EGFP/EGFP} plexus, however, 58.5% of all Muc1⁺EGFP^{HI} cells ($n=91$) were apically narrowed, versus 18.3% in the *Neurog3*^{EGFP/+} plexus ($n=82$) (Fig. 3K-R), suggesting that *Neurog3*^{NULL} cells do not efficiently progress past apical narrowing (Fig. S8A). The lengths of F-actin^{FOCAL} structures in *Neurog3*-null Muc1⁺EGFP^{HI} cells were reduced relative to control (Fig. S8C,D), suggesting reduced basalward movement and consistent with indications of diminished protrusive distortions at the basal surface (Fig. S8C-E). Finally, a small proportion of Muc1⁺EGFP^{HI} *Neurog3*^{EGFP/EGFP} cells were non-apically narrowed and not basally displaced (15.3% of all *Neurog3*^{EGFP/EGFP} Muc1⁺EGFP^{HI} cells versus 2.3% in *Neurog3*^{EGFP/+} controls;

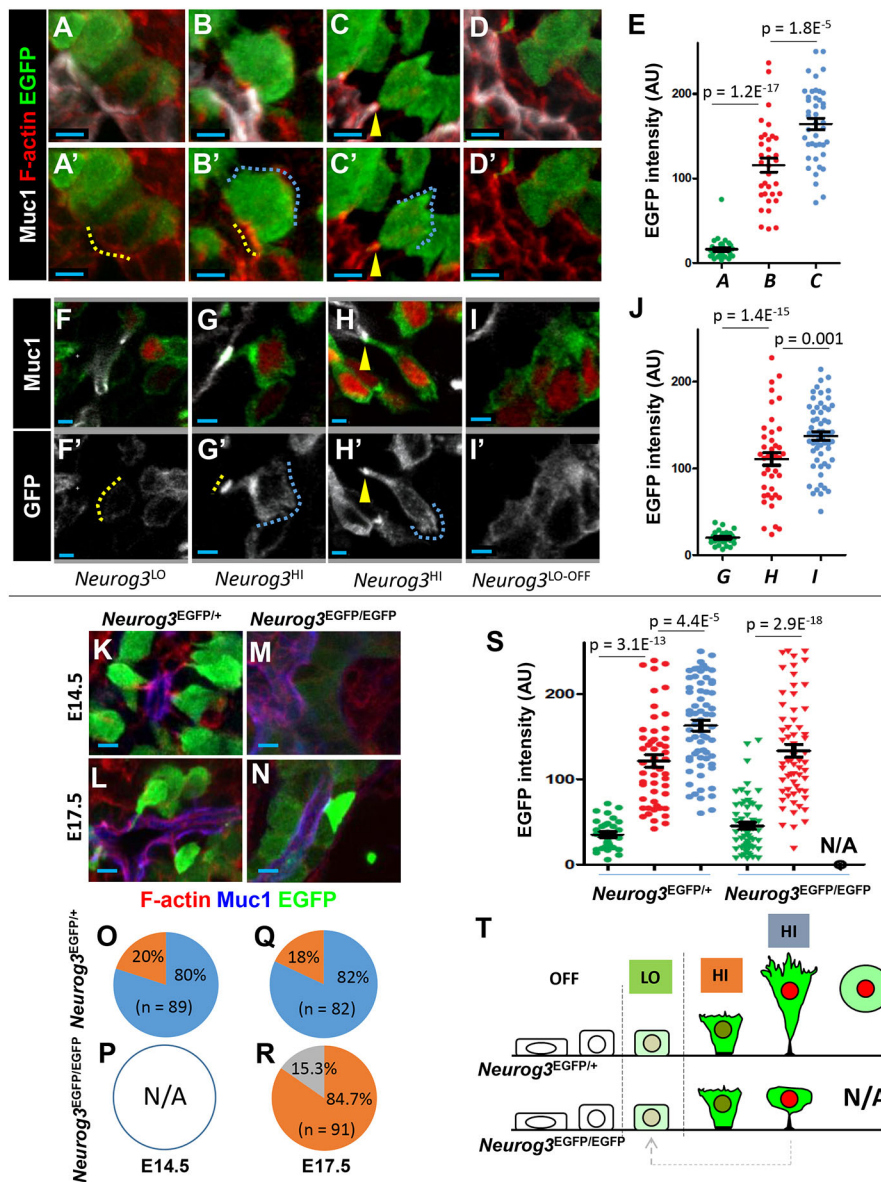


Fig. 3. Neurog3-independent apical narrowing and Neurog3-dependent basalward cell displacement guide endocrine-cell birth.

(A-D') *Neurog3*^{EGFP/+} EGFP⁺ cells showing stepwise progression from non-apically narrowed and non-egressing EGFP^{LO} to an apically narrowed (yellow dashed line) or F-actin^{FOCAL}-associated basally displaced (blue dashed line) EGFP^{HI} state, and finally to an apically detached state. (E) Mean fluorescence intensity measurements (AU, arbitrary units) in cells non-apically narrowed, apically narrowed or basally displaced. (F-I') *Neurog3*^{RG+} cells showing similar progression as in A-D'. (K) Mean fluorescence measurements as in E. (K,L) EGFP⁺ cells in *Neurog3*^{EGFP/+} plexus. (M,N) EGFP⁺ cells in *Neurog3*^{EGFP/EGFP} plexus. (O-R) Percentage of Muc1⁺EGFP^{HI} cells in non-apically narrowed (gray), apically narrowed (orange) and basally displaced (blue) states in each genetic condition. (S) Fluorescence intensities from non-apically narrowed (green), apically narrowed (red) and basally displaced (blue) EGFP⁺ cells in E17.5 *Neurog3*^{EGFP/+} and *Neurog3*^{EGFP/EGFP} epithelium. (T) Schematic showing *Neurog3*-independent and *Neurog3*-dependent steps during *Neurog3* upregulation and cell egression. Scale bars: 5 μm in A-D', K-N; 3 μm in F-I'.

Fig. 3R-T). We interpret this population as cells that had previously entered the initial stages of apical narrowing, focalization and *Neurog3* upregulation, but then failed in apical-surface detachment and returned to the epithelium. These data show that complete *Neurog3* deficiency compromises, but does not completely block, cells in the plexus from entering the sequence of apical narrowing, focalization and basalward movement normally defined by the endocrine-committed *Neurog3*^{HI} state. Moreover, these data suggest that a program of morphogenetic regulation of apical narrowing, focalization and basalward cell movement can be effected independent of *Neurog3* protein function (Fig. 3T).

NmMyoII-dependent pathways mediate plexus morphogenesis and endocrine-cell birth

The molecular motor nmMyoII, produced as IIa (Myh9), IIb (Myh10) or IIc (Myh14) isoforms, acts on F-actin substrates to drive processes such as apical constriction and cell delamination (Martin and Goldstein, 2014). The active phosphorylated form of nmMyoII light chain (p-nmMyoII) was detected at the apical aspect of pancreatic epithelial cells, in a pattern consistent with

close localization to the F-actin^{BELT} network (Fig. S9A,B). The p-nmMyoII signal was detected on most but not all Muc1⁺ F-actin^{FOCAL} structures (Fig. S9C,D). Immunodetection of nmMyoIIa/IIb isoforms showed a substantial signal in the blood vessels and a lower pan-mesenchymal signal (Fig. S9K,L), consistent with our detecting p-nmMyoII in both tissues (data not shown). The available reagents did not detect nmMyoIIa/IIb in the pancreatic epithelium or in *Neurog3*^{EGFP}-expressing cells. Analysis of nmMyoIIc using a GFP fusion knock-in mouse strain (Ebrahim et al., 2013), however, showed a pattern surprisingly selective for the epithelium at E14.5, which became largely restricted to non-acinar epithelium by E17.5 (Fig. S9E-J), and a low nmMyoIIc-GFP signal in differentiated endocrine cells (data not shown). These data are consistent with roles for nmMyoII activity in mediating epithelial morphogenesis and endocrine differentiation in the plexus.

NmMyoII activity maintains the plexus state and promotes endocrine differentiation

To assess potential function(s) of nmMyoII in apical-surface narrowing and focalization within the plexus, we explanted

plexus-rich segments of E15.5 dorsal pancreata (Fig. 4A), and cultured them with or without the nmMyoII inhibitor blebbistatin (BBS) (Straight et al., 2003), which directly interferes with intrinsic GTPase motor activity (Kovacs et al., 2004) to inhibit all nmMyoII-dependent processes. With vehicle treatment over 36 h of culture, the plexus maintained gross morphological features similar to those observed at equivalent *in vivo* stages. Conversely, BBS inhibition caused a broad, rapid and reproducible transformation of the plexus into an abnormal ‘duct-like’ state that exhibited increased lumen diameter and flattened epithelial cell morphologies (Fig. 4B-G). Within these transformed duct-like states, F-actin^{BELT} perimeters became enlarged compared with those in untreated explants (Fig. 4H-L). Much of any remaining plexus exhibited distended lumens that were prominent at the ‘nodes’ where epithelial segments of the plexus intersected, and acinar lumens were also distended (Fig. S10A-F). The BBS-mediated effects on epithelial morphology were dose dependent and reversible. An 18 h BBS washout led to restoration of normal F-actin^{BELT} morphologies (Fig. S11A-H), indicating a robust recovery in the 25-50 μ M dose-range of live tissue with differentiating endocrine cells (not shown). These BBS doses and time frames of exposure that caused minimal cell death in the explants (not shown) were consistent with the notion that the effects on epithelial morphology were likely the outcome of acute nmMyoII inhibition. The effects of BBS on plexus morphology were not caused by loss of epithelial integrity or apical polarity, because aPKC, the primary cilium marker γ -tubulin, and E-cad maintained their respective apical and basolateral localizations during the experimental timeframe (Fig. S12A-F). Concomitant with BBS-induced duct-like transformation, there was a marked decrease in the number of Neurog3⁺ cells (Fig. 4C,F, quantified

below). In rare regions where plexus remained, large numbers of both Neurog3⁺ cells still existed, suggesting that BBS had not accessed all areas of the explant volume equivalently, and that in such areas endocrine-cell birth could progress forward. Additionally, although the immunofluorescent signal for Sox9 was somewhat reduced in BBS-treated explants, there was no change in epithelial Pdx1 or Prox1 immunodetection intensity (Fig. S13A-H), suggesting that the reduced Neurog3⁺ cell numbers did not result from loss of pancreatic epithelial identity (Westmoreland et al., 2012). These experiments targeting nmMyoII activity, although not cell-type specific in nature, suggest functions for nmMyoII in limiting apical surface size, in maintaining the plexus morphology and in mediating the production of Neurog3-expressing populations within the plexus (Fig. 4M).

ROCK-nmMyoII activity mediates sequential steps in the delamination of Neurog3-expressing cells

ROCK-nmMyoII pathway activity has been shown to regulate apical narrowing, cell migration and cell-rear retraction (Riento and Ridley, 2003; Worthylake et al., 2001), and our data presented here and that of others (Gouzhi et al., 2011) suggest that these processes are essential for endocrine-cell birth. To test how separate parts of the ROCK-nmMyoII pathway influence epithelial morphogenesis and endocrine-cell birth, we tested whether pharmacological inhibitors of ROCK and nmMyoII block or augment apical narrowing, focalization and basalward cell displacement in *Neurog3*-expressing cells. Consistent with our results above, BBS-treated *Neurog3*^{RG} explants showed reduced numbers of *Neurog3*^{RG+} cells along the distended Muc1⁺ lumens (Fig. 5A-D; Fig. S14A-E; quantified below). Among the *Neurog3*^{RG+} cells

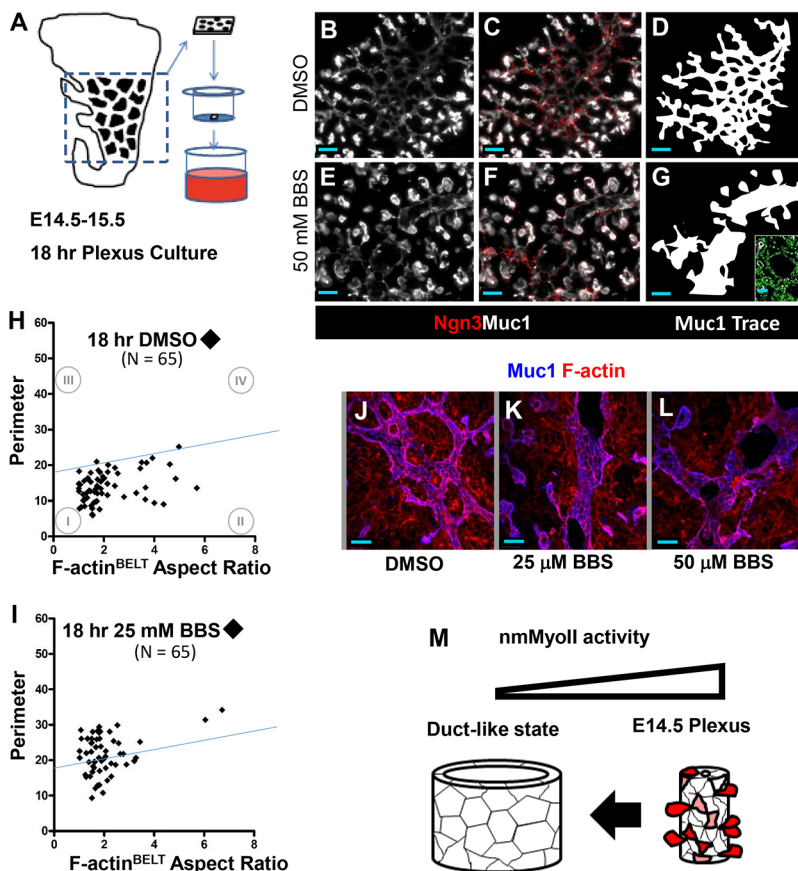


Fig. 4. NmMyoII activity limits apical expansion and promotes generation of Neurog3⁺ cells. (A) Diagram of the culture method. (B-G) Muc1 and Neurog3 immunodetection in explants cultured for 18 h under vehicle or BBS-treated conditions; manual epithelial tracings are shown in D and G. Inset in G shows a sectional plane at higher magnification, with DAPI to indicate position of nuclei related to the Muc1⁺ lumen surface. (H, I) F-actin^{BELT} shapes in explants in vehicle or BBS explants. Blue lines indicate the maximum y-axis F-actin^{BELT} dimension measured *in vivo*. I, narrowed F-actin^{BELT}; II, elongated F-actin^{BELT}; III, expanded F-actin^{BELT}; IV, expanded and elongated F-actin^{BELT}. (J-L) Muc1 and F-actin after treatment with increasing BBS doses. (M) Schematic showing the morphological transformation of the plexus into a duct-like state under nmMyoII inhibition. Scale bars: 50 μ m in B-G; 30 μ m in inset of G; 20 μ m in J-L. Data are from $n=3$ explants for each condition.

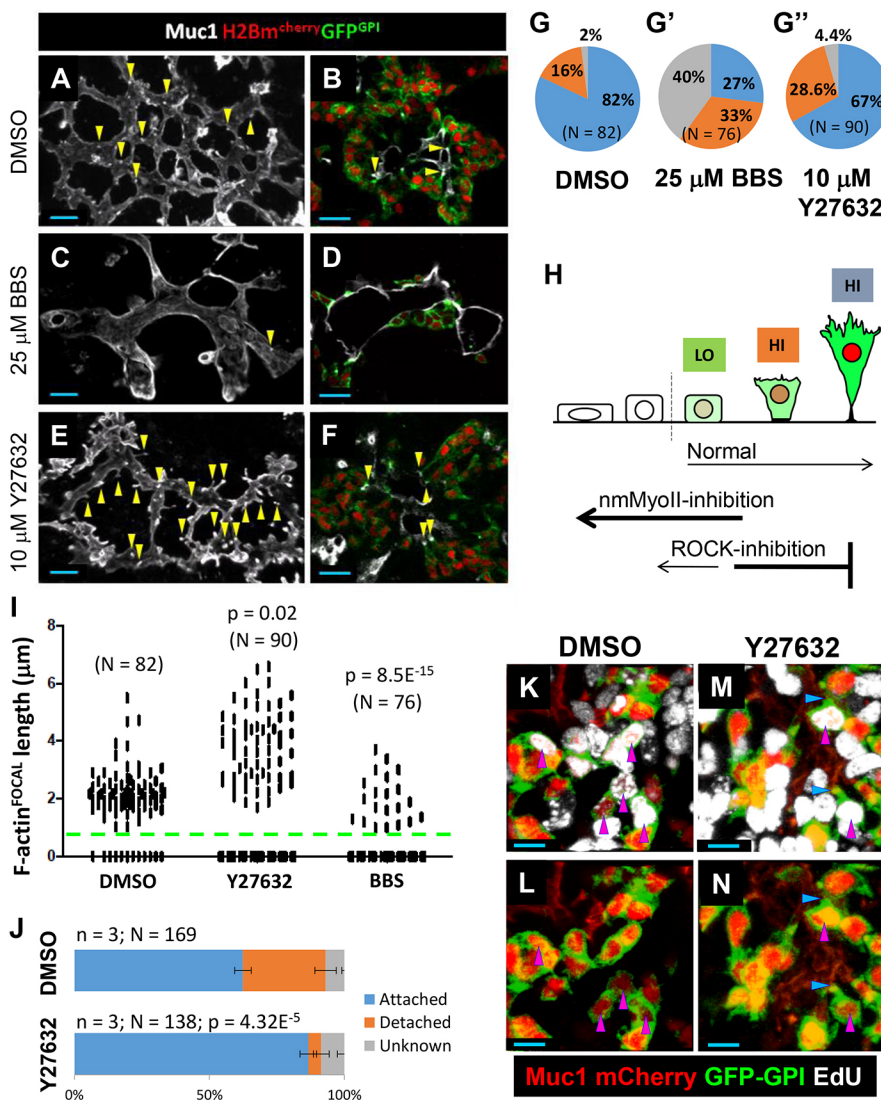


Fig. 5. ROCK-nmMyoII activities mediate morphological transitions during endocrine-cell birth. (A-F) *Muc1*⁺ and *Neurog3*^{RG+} cells in explants treated with vehicle alone, 25 μM BBS or 10 μM Y27632 for 18 h; *Muc1*⁺ F-actin^{FOCAL} structures are indicated by yellow arrowheads. There is a reduction in apically narrowed and basally displaced populations after BBS treatment. (G-G'') Proportions of non-narrowed (gray), apically narrowed (orange) and basally displaced (blue) *Neurog3*^{RG+} populations after vehicle, BBS or Y27632 treatment. (H) Schematic showing non-apically narrowed, apically narrowed and basally displaced *Neurog3*-expressing cells in G, and the effect of nmMyoII and ROCK inhibition. (I) Lengths of typical F-actin^{FOCAL} apical tether structures after vehicle, Y27632 or BBS treatment. Green dashed line indicates the distinction between an apically narrowed surface (no F-actin^{FOCAL} length or <1.0 μm) versus an elongated F-actin^{FOCAL} structure (>1.0 μm). (J) EdU pulse-chase quantification of defective cell-rear retraction in Y27632-treated explants. (K-N) EdU⁺ *Neurog3*^{RG+} cells exhibiting attached (blue arrowheads) or detached (pink arrowheads) cell rears. Scale bars: 20 μm in A-F; 5 μm in K-N. Data are from *n*=3 explants for each condition.

present in BBS-treated explants, quantification of non-apically narrowed, apically narrowed and basally displaced *Neurog3*^{RG+} cells indicated a distinct shift toward a non-apically narrowed and non-displaced positioned state (Fig. 5G,G'). The *Muc1*⁺ F-actin^{FOCAL} structures observed showed a decreased length compared with controls (Fig. 5H,I). These results fit the notion that nmMyoII activity is crucial for apical narrowing, focalization and basalward cell movement during endocrine-cell birth.

Blocking ROCK activity with Y27632 (Uehata et al., 1997) is known to alter cell migration and also cause defects in cell-rear retraction in some cell types, with such changes attributed, at least in part, to ROCK-mediated modulation of nmMyoII activity, actin-filament stabilization and disassembly of integrin-signaling complexes (Riento and Ridley, 2003; Worthylake et al., 2001). In pancreatic explants, ROCK inhibition led to several effects on tissue and cell morphology that were noticeably distinct from those under nmMyoII inhibition. Y27632-treated explants showed no evidence of plexus transformation into 'duct-like' states, and there was no decrease in the number of *Muc1*⁺ *Neurog3*^{RG+} cells exiting from the epithelium (Fig. 5E-G,G'). Upon close inspection, and consistent with known functions for ROCK in regulating cell migration, reporter-positive *Neurog3*^{RG+} cells (we did not specifically distinguish between reporter-high versus low states in these drug-

treatment conditions) scored in Y27632-treated explants displayed increased basalward movement relative to control *Neurog3*^{RG+} cells (Fig. S14A,F-I). Treatment with Y27632 increased the number of *Neurog3*^{RG+} cells with, as well as length of, F-actin^{FOCAL} structures at their apical surface (Fig. 5E,F,I) suggesting a block in detachment of the apical-cell rear, even with the exaggerated basalward movement of these cells. We confirmed a cell-rear detachment defect in *Neurog3*^{RG+} cells using EdU pulse-chase methods. Briefly, replicating progenitor cells incorporate EdU at S phase, and over a 17-24 h period a proportion of them differentiate and become epithelially detached (*Muc1*⁻) (Bankaitis et al., 2015). Defective rear detachment caused by Y27632 treatment should thus cause an accumulation of EdU⁺ *Muc1*⁺ *Neurog3*^{RG+} cells. One hour after EdU injection into pregnant dams, E13.5 pancreas was explanted and cultured for 18 h with or without Y27632. Confocal z-stack volumes of EdU pulse-labeled explants were scored for attached (*Muc1*⁺) or detached (*Muc1*⁻) EdU⁺ *Neurog3*^{RG+} states. In vehicle controls, 63.4±3.1% of EdU⁺ *Neurog3*^{RG+} cells were *Muc1*⁺, 30.6±3.9% were *Muc1*⁻ and 7.0±1.0% were non-categorizable (Fig. 5J-L). In Y27632-treated explants, 86.6±3.1% of EdU⁺ *Neurog3*^{RG+} cells were *Muc1*⁺, 4.8±2.9% *Muc1*⁻ and 8.6±2.7% were non-categorizable (Fig. 5J,M,N). These data are consistent with two functions for ROCK in the birth of endocrine

cells: first in limiting basalward movement, and then in directing cell-rear detachment to allow epithelial exit. Finally, short-term (6 h) time-lapse videos of pancreatic explants cultured on fibronectin, with or without BBS or Y27632, confirmed that nmMyoII inhibition reduced, whereas ROCK inhibition augmented, basalward movement of *Neurog3*^{RG+} cells (Fig. S15A-O). Thus, ROCK and nmMyoII coordinate different aspects of endocrine-cell delamination by regulating apical narrowing, focalization, basalward movement and detachment of the rear of the cell from its focalized apical contact.

nmMyoII activity is necessary for *Neurog3* activation in the absence of Notch-mediated repression

Because nmMyoII inhibition blocked behaviors associated with endocrine commitment and reduced the numbers of *Neurog3*⁺ cells, we hypothesized that nmMyoII activity is required for acquisition of the *Neurog3*-expressing state. We therefore quantified how BBS affected the number of *Muc1*⁺*Neurog3*⁺ cells formed in explants, as a metric for the BBS effect on all *Neurog3*-expressing states. Notably, because the processes of apical narrowing and focalization were perturbed under BBS-treated conditions, we lacked key criteria necessary for quantifying changes in the numbers of *Neurog3*^{LO} versus *Neurog3*^{HI} cells, and thus do not report on such changes here.

Comparison of the (*Neurog3*⁺*Muc1*⁺*DAPI*⁺)/(*Neurog3*-negative *Muc1*⁺*DAPI*⁺) cell fraction in E15.5 explants treated with BBS or vehicle showed a near 50% reduction in all *Neurog3*⁺ cells under nmMyoII inhibition (Fig. 6A-B',E). We next tested the effect of nmMyoII inhibition in the presence or absence of the small-molecule Notch inhibitor DBZ (Milano et al., 2004), which causes 'default' derepression of *Neurog3* and acquisition of the *Neurog3*⁺ state (Serup, 2012; Bankaitis et al., 2015). Consistent with this effect, DBZ-only explants showed ~2.5-fold increase in total (*Neurog3*⁺*Muc1*⁺*DAPI*⁺)/(*Neurog3*-negative *Muc1*⁺*DAPI*⁺) cells over vehicle alone (Fig. 6C,C',E). In BBS/DBZ-treated explants, *Neurog3*⁺ cell numbers increased slightly over vehicle alone, but were greatly reduced compared with DBZ alone (Fig. 6D-E). These results indicate that nmMyoII inhibition interferes with processes controlling *Neurog3* transcriptional activation or upregulation, or both, even when in the context of induced escape from Notch repression, suggesting that endocrine-cell birth requires permissive inputs supplied by nmMyoII activity.

ROCK activity suppresses the *Neurog3*^{LO-HI} transition

Apical focalization and initiation of basalward movement are associated with the *Neurog3*^{HI} state in normal and *Neurog3*-null epithelium (Fig. 3). Because ROCK inhibition augmented apical

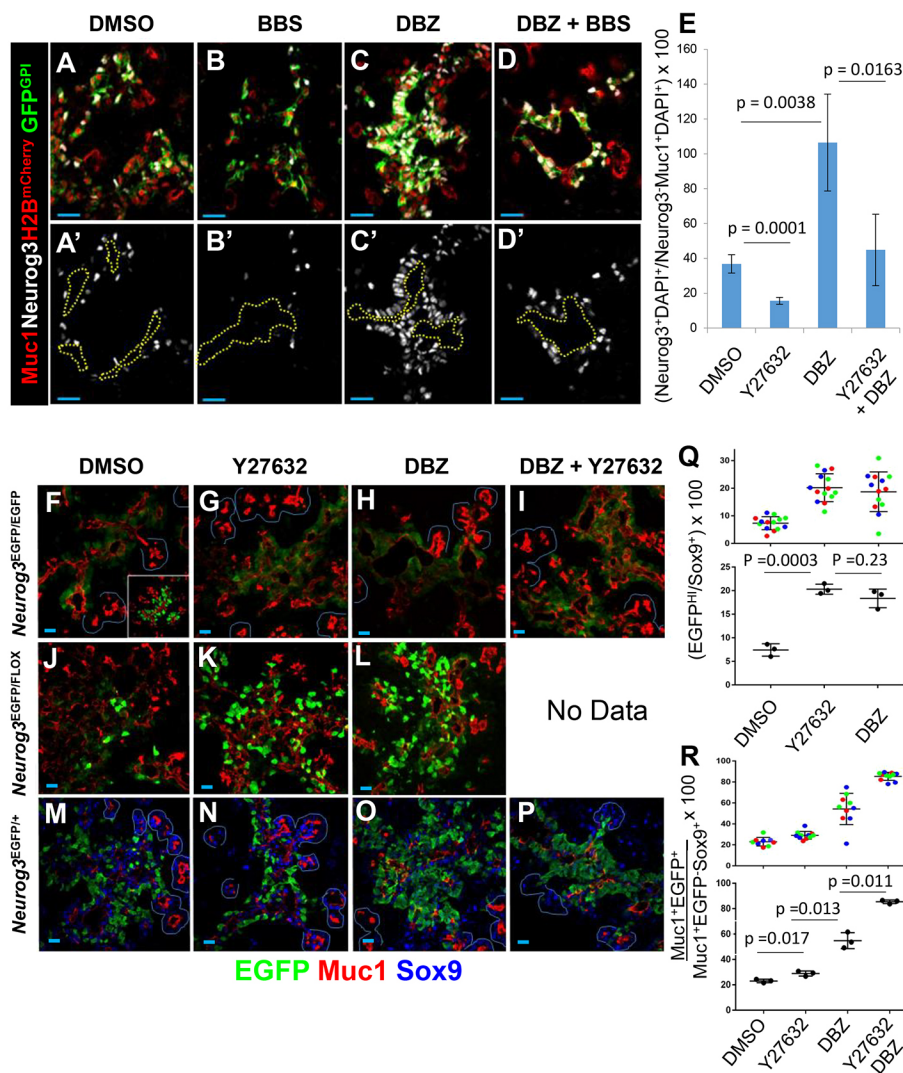


Fig. 6. ROCK-nmMyoII, Notch and *Neurog3* gene dosage regulate acquisition of *Neurog3*-expressing states. (A-D') *Muc1* and *Neurog3* protein in E15.5 *Neurog3*^{RG+} explants after treatment with vehicle alone, 25 μ M BBS, 20 nM DBZ, or BBS and DBZ. BBS was applied for 18 h with DBZ added 6 h into culture (DBZ exposure, 12 h). (E) Numbers of *Muc1*⁺ *DAPI*⁺ cells that are *Neurog3*⁺ in each condition (*DAPI*⁺ signal not shown). (F-I) *Muc1*, EGFP and nuclear Sox9 (latter not shown) in E13.5 *Neurog3*^{EGFP/EGFP} explants treated with 10 μ M Y27632, 20 nM DBZ or Y27632 plus DBZ for 18 h. (J-L) E13.5 *Neurog3*^{EGFP/Flox} explants treated and analyzed according to F-H. (M-P) *Muc1*, EGFP and Sox9 in E13.5 *Neurog3*^{EGFP/+} explants treated according to F-I. (Q) Quantification of endocrine-committed EGFP^{HI} Sox9⁻ cells compared with Sox9⁺ cells in 40 \times z-stacks (~20 μ m) *Neurog3*^{EGFP/Flox} hypomorph explants treated with vehicle, Y27632 or DBZ. Upper colored panel shows spread of raw data, each dot representing one z-stack and each color representing an individual biological replicate (mean \pm s.d.). (R) Quantification of total (*Muc1*⁺EGFP⁺)/(total *Muc1*⁺EGFP⁻Sox9⁺) cells in *Neurog3*^{EGFP/+} explants treated with vehicle, Y27632, DBZ, or DBZ and Y27632. Upper colored panel shows data represented as in Q (mean \pm s.d.). Scale bars: 20 μ m for A-D'; 10 μ m for F-P. Blue lines indicate acinar clusters. $n \geq 3$ pancreata for each condition. For E, Q and R, $n = 3$ pancreata for each condition; significance is determined using Student's *t*-test.

focalization and basalward movement in *Neurog3*-expressing cells (Fig. 5), we hypothesized that ROCK normally functions to limit endocrine-cell birth. We first tested whether Y27632 treatment can alleviate the defective *Neurog3* transcriptional upregulation observed in early-stage *Neurog3*^{EGFP/EGFP} (null) explants. In *Neurog3*^{EGFP/EGFP} explants, *Neurog3* is broadly activated at low levels, but Y27632 did not increase the number of *Neurog3*^{HI} cells (Fig. 6F,G). Thus, Y27632 treatment is not able to promote *Neurog3* upregulation in the absence of *Neurog3* protein function. We next tested whether ROCK inhibition could raise the level of *Neurog3* expression in cells already expressing low levels of functional *Neurog3* protein, by exposing explants from a strong hypomorphic condition, *Neurog3*^{EGFP/Flox} (Wang et al., 2010), to Y27632. *Neurog3* hypomorphs are much more prone to maintaining cells in the *Neurog3*^{LO} state than are wild-type or heterozygous-null controls in which most cells progress forward to the *Neurog3*^{HI} state (Wang et al., 2010; Becharad et al., 2016). Accordingly, vehicle-treated *Neurog3*^{EGFP/Flox} tissues had significantly increased *Muc1*⁺*Sox9*⁺*EGFP*^{LO} cell numbers over vehicle-treated *Neurog3*^{EGFP/+} explants, and much reduced numbers of endocrine-committed *Muc1*⁺*Sox9*⁺*EGFP*^{HI} cells (Fig. 6J versus M). The Y27632-treated *Neurog3*^{EGFP/Flox} explants, however, showed near-complete restoration of *Muc1*⁺*Sox9*⁺*EGFP*^{HI} numbers towards those seen in vehicle-treated *Neurog3*^{EGFP/+} and in 20 nM DBZ-treated *Neurog3*^{EGFP/Flox} explants (Fig. 6K-Q). Therefore, we conclude that ROCK activity normally gates the transition from *Neurog3*^{LO} to *Neurog3*^{HI} states, and ROCK inhibition can bypass the defects in *Neurog3* upregulation seen under *Neurog3* hypomorphism. Interestingly, Y27632-treated *Neurog3*^{EGFP/+} explants exhibited only a slight increase in the total *Muc1*⁺*EGFP*⁺/*Muc1*⁺*EGFP*⁻*Sox9*⁺ cell number relative to vehicle, indicating that Y27632 does not cause augmentation of *Neurog3* activation or upregulation when a normal number of *Neurog3*-expressing cells are present (Fig. 6M,N,R).

Notch and ROCK suppress endocrine differentiation through distinct mechanisms

That ROCK inhibition does not substantially increase *Neurog3*^{HI} cell numbers in *Neurog3*^{EGFP/+} explants, but does in *Neurog3* hypomorphs (Fig. 6Q,R), implies that ROCK inhibition cannot promote *Neurog3*^{HI} cell numbers if large numbers of *Neurog3*^{HI} cells are already present. We proposed that this inability resulted from Notch-mediated lateral inhibition from the *Neurog3*^{HI} cell population. To test this idea, we treated heterozygous *Neurog3*^{EGFP/+} explants with DBZ and Y27632 together, and compared total cell numbers (*Muc1*⁺*EGFP*⁺/*Muc1*⁺*EGFP*⁻*Sox9*⁺) with those after vehicle, Y27632 or DBZ treatment. DBZ/Y27632 co-treated *Neurog3*^{EGFP/+} explants showed a large increase in total (*Muc1*⁺*EGFP*⁺/*Muc1*⁺*EGFP*⁻*Sox9*⁺) cells over DBZ or Y27632 alone (Fig. 6M-P,R). These data suggest that the failure of Y27632 to promote robust *Neurog3* upregulation in *Neurog3*^{EGFP/+} explants is circumvented by inhibiting Notch activity, and thus that ROCK inhibition stimulates *Neurog3* activation and upregulation provided that cells escape Notch repression. Notably, the majority of *Muc1*⁺*EGFP*⁺ cells in DBZ-treated *Neurog3*^{EGFP/+} explants remained *Sox9*⁺ (Fig. S16B), suggesting that pharmacological Notch inhibition derepresses *Neurog3* expression but interferes with the normal process of *Sox9* downregulation normally associated with *Neurog3* upregulation (Shih et al., 2012). Because this effect is not seen in the context of Y27632 alone, we propose it likely that ROCK and Notch activity regulate endocrine differentiation through separate mechanisms or at different stages of *Neurog3*

activation and upregulation. Like Y27632 alone, DBZ alone had no apparent effect on the number of cells upregulating EGFP-reporter in homozygous null explants, nor did DBZ and Y27632 in combination (Fig. 6F-I), indicating that the ability of DBZ or Y27632, both individually and in combination, to cause *Neurog3* upregulation is dependent on the presence of at least low levels of *Neurog3* protein. These results suggest that, provided *Neurog3* protein is available, ROCK activity limits the normal transition of *Neurog3*^{LO} cells to *Neurog3*^{HI} endocrine-committed cells.

Large-scale compensatory synergy between morphogenetic and endocrine-birth programs under reduced *Neurog3* function

The results presented above, together with evidence from other reports, suggest that *Neurog3* gene dose, Notch and ROCK-nmMyoII work together in a feedback control circuit that coordinates endocrine fat allocation, progenitor growth and epithelial plexus morphogenesis (Wang et al., 2010; Magenheim et al., 2011; Bankaitis et al., 2015). We were interested in the degree of robustness of this circuit, and its ability to yield, under various degrees of challenge, compensatory functions in driving the linked processes of morphogenesis and endocrine differentiation. We reasoned that the late-stage corrective remodeling of an initially (E14.5) abnormal *Neurog3*^{EGFP/EGFP} plexus, occurring concurrent with a partial restoration of *Neurog3*^{HI} cell number (Fig. 2), reflects the presence but only partial success of a compensatory response. We proposed that the failure to fully restore *Neurog3*^{HI} cell numbers resulted from the absence of feedback signals normally derived from endocrine-committed *Neurog3* protein-producing cells during their departure from the plexus epithelium. We therefore tested whether a hypomorphic level of *Neurog3* represented a sufficient feedback stimulus to confer a full compensatory response linking conversion of the plexus into an epithelial structure with a full restoration of *Neurog3*^{HI} endocrine-committing cells. We quantified the number of apically narrowed *Muc1*⁺*EGFP*^{HI} cells in the plexus (*Muc1*⁺*EGFP*^{HI} cells as a proportion of *Sox9*⁺ cells) at E13.5, E16.5 and E18.5 to compare endocrine-cell birth as a function of plexus remodeling in *Neurog3*^{EGFP/+}, *Neurog3*^{EGFP/Flox} and *Neurog3*^{EGFP/EGFP} states. *Neurog3*^{EGFP/+} pancreata showed robust endocrine differentiation in all E13.5, E16.5 and E18.5 samples. The *Neurog3*^{EGFP/EGFP}-null pancreata again showed that the early and pervasive defects in *Neurog3* upregulation were progressively but only partially rescued by E16.5 and E18.5, confirming that *Neurog3*-dependent processes induce the normal rate of *Neurog3*^{HI} cell birth (Fig. 7A-F). In the *Neurog3*^{EGFP/Flox} hypomorph, remarkably, although very low numbers existed at early stages, there was a complete restoration of the number of *Neurog3*^{HI} cells by E18.5 (Fig. 7A-J). Thus, even a strongly reduced level of *Neurog3* is sufficient to engage, albeit late, an adaptive feedback control circuit that links endocrine differentiation and plexus morphogenesis to establish and maintain the balance between cell differentiation and epithelial remodeling.

DISCUSSION

Our results focus on a new ‘global’ understanding of the functional integration between programs of *Neurog3*-driven endocrine-fate determination, Notch-mediated progenitor maintenance, and ROCK/nmMyoII-mediated epithelial morphogenesis. We describe distinct ways in which the ROCK/nmMyoII pathway mediate a dissociable sequence of epithelial egression events linked to the acquisition of pre-endocrine-committed and endocrine-committed *Neurog3*-expressing states within the dynamically remodeling

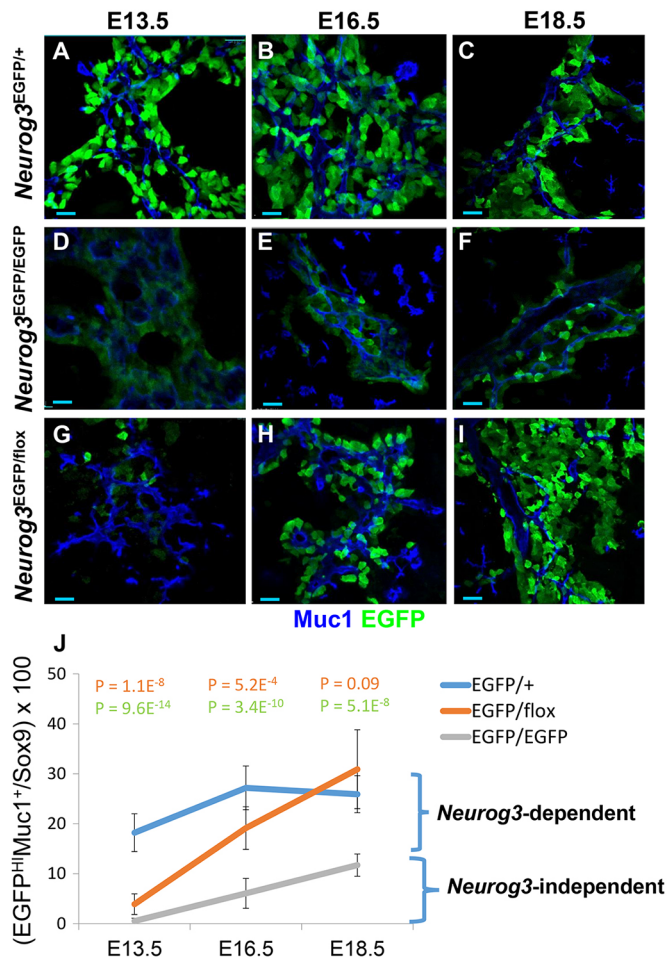


Fig. 7. Plexus morphogenesis intercommunicated with *Neurog3* gene dose drives robust endocrine differentiation. (A–I) Muc1 and EGFP shown as a time-course for plexus remodeling in sectioned (30 μ m) *Neurog3*^{EGFP/+} (A–C), *Neurog3*^{EGFP/EGFP} (D–F) and *Neurog3*^{EGFP/Flox} (G–I) pancreata. (J) Quantification of the amount of delaminating endocrine committed (EGFP^{HI}Muc1⁺Sox9⁻ cells)/(total Sox9⁺EGFP⁻ cells) at each stage. Scoring from images of plexus epithelium was averaged from $n=3$ pancreata (*Neurog3*^{EGFP/+}=11, 10 and 9 images for E13.5, E16.5 and E18.5, respectively; *Neurog3*^{EGFP/EGFP}=12, 10 and 9 images for E13.5, E16.5 and E18.5, respectively; *Neurog3*^{EGFP/Flox}=9, 11, and 14 images for E13.5, E16.5 and E18.5, respectively). Error bars are s.e.m. Scale bars: 15 μ m.

epithelium. The main findings of this paper are presented schematically in Fig. S17. We propose that a feedback circuit comprising ROCK/nmMyoII, Notch and *Neurog3*-dependent influences bestows self-organizing feedback properties that ensure the correct balance between progenitor growth, differentiation and morphogenesis.

Morphogenetic determinants mediate endocrine cell-fate acquisition in plexus progenitors

ROCK/nmMyoII-dependent processes mediate a step-wise cell-egression sequence for progression into and through *Neurog3*^{LO} and *Neurog3*^{HI} states. Inhibiting nmMyoII activity interferes with apical narrowing and focalization, and basalward cell movement, and limits formation of *Neurog3*⁺ cells. Similar deficits in nmMyoII-inhibited explants occurred even when endocrine cells were overproduced when lifting Notch-mediated repression. Because our experiments did not discern cell-autonomous effects of nmMyoII inhibition, future dissection of its function specifically

within *Neurog3*-expressing cells will likely be informative. ROCK inhibition augmented apical narrowing, focalization and basalward cell movement, and efficiently promoted progression of *Neurog3*^{LO} into *Neurog3*^{HI} cells in *Neurog3* hypomorphs, but not in heterozygous *Neurog3*^{EGFP/+} explants with much more robust Notch-mediated inhibition. A pro-endocrine modulation by ROCK inhibition occurred in heterozygous explants only with Notch co-inhibition, implying that, in normal tissue, ROCK de-repression promotes *Neurog3*^{LO-to-HI} transitions up to a ‘set point’ where an appropriate number of *Neurog3*^{HI} cells exert lateral inhibition as a dominant patterning determinant. Thus, we suggest that ROCK and nmMyoII are important factors in mediating timely and spatially localized cellular responses to signals that both drive (*Neurog3* as a cell-autonomous instructor) and inhibit (via Notch) endocrine-fate acquisition. Intimate connections between epithelial morphogenesis programs and intrinsic cell-fate gene-regulatory networks are supported by additional studies, such as when *Cdc42* inactivation led to failed progenitor-cell apicobasal polarization, block of epithelial tubulogenesis and severe endocrine-cell depletion (Kesavan et al., 2009). This phenotype resulted at least in part from improper positioning of cells within environments conducive for endocrine differentiation, as discussed below. Moreover, perturbation in *Pdx1*⁺ epithelial cells of CELSR proteins, components of the Wnt/PCP (planar cell polarity) signaling orthogonal to apico-basal polarity, caused large reductions in *Neurog3*⁺ cells (Cortijo et al., 2012). Other reports connect morphogenesis and differentiation via RhoA (Petzold et al., 2013; Azizoglu et al., 2017) and Rac signaling (Löf-Öhlin et al., 2017). An important link between Rac, apical domain size control and endocrine cell-fate acquisition occurs in mice and humans (Löf-Öhlin et al., 2017). It will be important in the future to dissect how morphogenetic pathways mediate recognition and processing of epithelial cues, possibly under precise local spatial control, that control progenitor growth and cell fate. Targeting such activities could alter the growth and differentiation properties of pancreas-differentiated iPES/ES cells *in vitro*.

Morphogenetic behavior of endocrine-biased progenitors in an asymmetric niche

Stem-cell niche microenvironments are commonly asymmetric, e.g. showing pro-maintenance or pro-differentiation cues at basal or apical aspects of a cell or tissue (Li and Xie, 2005). Theoretically, controlling the timing and exposure to such separated cues could regulate which cells undergo cell-fate allocation, and when. Particularly relevant, therefore, are the associations drawn here, in both *Neurog3*^{EGFP/+} and *Neurog3*^{EGFP/EGFP} pancreata, between apical surface size/shape, basalward cell movement within and away from the epithelium, and *Neurog3* expression. Cells in all genetic conditions studied here, including the *Neurog3* protein null, only underwent *Neurog3* upregulation upon narrowing and focalization of their apical surface (F-actin^{FOCAL}) and basalward cell displacement. We suggest, therefore, that apical-basal cell-body positioning within the epithelial plane controls, at least in part, acquisition of distinct *Neurog3*-expressing states upstream of *Neurog3* protein function. This finding seems conceptually aligned with zebrafish retinal cell behavior, wherein neurogenesis is coupled to interkinetic nuclear migration along an apical-to-basal Notch gradient (Del Bene et al., 2008). In addition, in the zebrafish retina, apical domain size has been linked to modulation of Notch signaling (Clark et al., 2012). In the fish and mouse pancreas, high, medium and low Notch signaling impart duct, bipotent progenitor and endocrine fates, respectively (Ninov et al., 2012; Shih et al.,

2012), associated in our studies with large, intermediate or narrowed/focalized apical surfaces. Because ROCK and nmMyoII inhibition showed opposite effects on the derivation of *Neurog3*⁺ cells, in the presence and absence of Notch inhibition, we suspect that regulation of apical surface size and basalward cell movement is a fundamental function by which ROCK and nmMyoII mediate cell fates.

Transcriptional determinants of cell fate regulate tissue and cellular morphogenesis

Collectively, our data support previous studies suggesting that differentiating endocrine-cell progeny perform a ‘feedback support role’ for the epithelial plexus niche, and extend the range of functions for *Neurog3* beyond its roles as an endocrine-lineage determinant and initiator of Notch-mediated lateral inhibition. The picture emerging is that *Neurog3* influences numerous developmental processes in the plexus niche, including driving robust progenitor replication (Bankaitis et al., 2015), non-cell-autonomously refining gene-expression patterning (Wang et al., 2010; Bechard et al., 2016), cell-autonomously determining endocrine-cell fate (Gradwohl et al., 2000; Gu et al., 2002) and regulating tissue morphogenesis (Magenheim et al., 2011; Bankaitis et al., 2015). We show the first evidence that *Neurog3*-dependent non-autonomous mechanisms stimulate the production of new endocrine cells from the plexus (Fig. 7). We propose that a feedback function for *Neurog3*⁺ cells drives, or is a strong supportive influence in, the systematic integrative sustenance of epithelial growth and morphogenesis that is balanced with the apportionment of duct versus endocrine cell types. Future analyses might lead to understanding how transcription-factor determinants feed into the morphogenetic programs to build precise micro-environments, and their coordination to ensure timely placement of the right types and numbers of lineages throughout organogenesis. Under *Pdx1* deficiency, early dorsal pancreatic bud evagination, cell polarization and microlumen formation all occur (Marty-Santos and Cleaver, 2016; Villasenor et al., 2010) but there is a complete failure in subsequent plexus assembly. Moreover, tissue-specific inactivation of transcription factors – such as *HNF1b* (De Vas et al., 2015), *HNF6* (Pierreux et al., 2006) or *Prox1* (Wang et al., 2005; Westmoreland et al., 2012) – all cause epithelial malformations concurrent with defective endocrine lineage allocation. Although these phenotypes were attributed largely to direct effects on genes such as *Neurog3*, equally consistent is the idea that an altered epithelial architecture precludes efficient spatial deployment of signals that drive endocrine differentiation and epithelial morphogenesis programs. Building systems-like network models linking epithelial morphogenesis, gene transcription and intracellular signaling as principal components for a larger ‘niche-framework’ will be important.

Adaptive self-organization in the plexus niche

We propose that cooperative engagement of adaptive morphogenetic regulation with *Neurog3*-dependent processes confers robustness on endocrine differentiation, affording resilience even when genetic insults cripple endocrine differentiation (Bankaitis et al., 2015; Fig. 7). In the case of *Neurog3*^{EGFP/EGFP}-null epithelium, that the morphologically abnormal plexus state is populated very broadly with *Neurog3*^{LO} cells indicates that signals triggering *Neurog3* transcriptional activation and specifying the endocrine-biased state are extensive over the epithelium. That adaptive restoration of *Neurog3*^{HI} cells occurs only during late-stage ‘corrective remodeling’ (in association with a specific sequence of cell-shape

changes) suggests that localized morphological alterations in the plexus epithelium exert central control over endocrine commitment via *Neurog3* upregulation. Importantly, the restoration of *Neurog3*^{HI} cells is only moderate in the absence of *Neurog3* protein (Fig. 7), which we interpret as indicating that plexus morphogenesis normally synergizes with *Neurog3*-dependent processes to instruct appropriate numbers of endocrine cells to begin exit from the growing, dynamically remodeling epithelium. For example, even with a low level of *Neurog3* protein function, the strongly hypomorphic epithelium could essentially completely restore *Neurog3*^{HI} cell numbers back from the massive mid-gestation reduction. We feel that such cross-regulatory feedback between non-autonomous morphogenetic determinants and *Neurog3* represents a large-scale ‘community effect’ (Gurdon, 1988) to modulate progenitor flux toward the various epithelially derived lineages. We propose that such a community effect is likely influenced by biomechanical (e.g. intra/intercellular tension) and biochemical factors (Cebola et al., 2015; Marinari et al., 2012), some derived directly from endocrine cells engaged in a prolonged process of epithelial egression. Because cell departure is associated with fate acquisition in developing systems across phyla, linking this process to the architecture and behavior of the parental epithelium could reflect a broad process of organ-system self-organization.

MATERIALS AND METHODS

Mice

Neurog3^{EGFP} [B6.129.S-*Neurog3*^{tm1(EGFP)Khk/Mmed}] knock-in (Lee et al., 2001) and *Neurog3*^{FLOX} [*Neurog3*^{tm1(cre/ERT)Ggu}] mice (Wang et al., 2008) were from Guoqiang Gu (Vanderbilt University, Nashville, TN, USA). *Neurog3*^{EGFP/+} and *Neurog3*^{EGFP/EGFP} pancreata were assigned definitively via EGFP-fluorescence level and lack of EGFP⁺ islet clusters, all readily observable under epifluorescence and agreeing with PCR genotyping and immunolabeling confirmation of genotype, as expected for a coding-region deletion allele. Tables S1 and S2 describe *Neurog3*^{EGFP/EGFP} and *Neurog3*^{Flox/+} genotyping, and *Neurog3*^{EGFP/Flox} PCR genotyping followed Wang et al. (2010). *Neurog3*^{RG1} mice were genotyped as described (Bechard et al., 2016). nmMyoIIC^{GFP} mice were a kind gift from Robert Adelstein (National Heart, Lung, and Blood Institute, Bethesda, MD, USA). Wild-type mice were of mixed genetic background. All protocols were approved by Vanderbilt University IACUC.

Immunofluorescence analysis

Embryonic pancreas was fixed in paraformaldehyde (4%, 4°C for 4–6 h, Fisher 04042-500) and for cryosectioning was sucrose equilibrated (30%, 4°C overnight, VWR 0335-12KG) and embedded in OCT (Tissue-Tek, Sakura). A Leica CM3050 S produced thick sections of 30–40 μm or 10 μm thin sections. Table S1 lists primary antibodies incubated overnight at 4°C in PBS containing 0.1% Triton X100, 0.5% BSA and 0.01% sodium deoxycholate. All secondary antibodies were from Jackson ImmunoResearch Laboratories, used at 1:500. Imaging was carried out using a Zeiss Apotome or confocal Zeiss LSM 510 META microscope. Confocal z-stack volumes were acquired at 40× magnification with 3D reconstructions using Imaris software. To measure apical F-actin cortices, 2D sections (1.5–2.0 μm sectional planes) from 3D volumes were assessed for whole F-actin^{BELTS} visible along the *Muc1*⁺ lumen surface, perimeter: aspect ratio was determined by line or polygon-trace functions in Imaris.

Quantification and statistical analyses

Distinguishing plexus and non-plexus states was carried out as described (Bankaitis et al., 2015). Fluorescence intensity measurements used mean pixel intensity in ImageJ. Sox9 and *Neurog3* numbers were counted manually. For *Neurog3*⁺ or EGFP⁺ explant analysis, cell numbers in each state were summed from individual plexus volumes (six or more for each sample, ~30% of total explant), averaged within each sample, and averaged between all samples. Measurements of morphological structures and nuclear

densities were made using line or polygon-trace functions in Imaris. Nuclei were defined by H2B-mCherry signal in *Neurog3^{RG}* samples or Sox9 immunopositivity in *Neurog3^{EGFP/+}* and *Neurog3^{EGFP/EGFP}* samples. Low-level Sox9 immunopositivity in cells initiating epithelial egression in *Neurog3^{EGFP/+}* samples was digitally enhanced (brightness and contrast) in Image J. All Student's *t*-tests were two-tailed.

Drug treatments

Explants were cultured at 37°C in 5% CO₂ at the air-liquid interface on trans-well supports (Costar 3422) in DMEM (no phenol red) with 10% FBS, penicillin/streptomycin (10 U/ml each) and 1% L-glutamine (Shih and Sander, 2014). Blebbistatin [Abcam, S(-) form, CAS, 856925-71-8], Y27632 (Selleckchem, S1049) and DBZ (Cayman Chemical, 209984-56-4) or vehicle (0.1% DMSO final) were added. Explants were fixed (for 2 h in 4% paraformaldehyde) for analysis.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: E.D.B., M.E.B., G.G., M.A.M., C.V.E.W.; Methodology: E.D.B., M.E.B.; Software: E.D.B.; Validation: E.D.B., C.V.E.W.; Formal analysis: E.D.B.; Investigation: E.D.B., M.E.B., C.V.E.W.; Resources: G.G., M.A.M., C.V.E.W.; Data curation: E.D.B.; Writing - original draft: E.D.B., C.V.E.W.; Writing - review & editing: E.D.B., M.E.B., C.V.E.W.; Visualization: E.D.B.; Supervision: G.G., C.V.E.W.; Project administration: C.V.E.W.; Funding acquisition: C.V.E.W.

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Supplementary information

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