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Phosphorylation of NEUROG3 Links Endocrine Differentiation to the Cell Cycle in Pancreatic Progenitors

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SUMMARY

During pancreatic development, proliferating pancreatic progenitors activate the proendocrine transcription factor neurogenin 3 (NEUROG3), exit the cell cycle, and differentiate into islet cells. The mechanisms that direct robust NEUROG3 expression within a subset of progenitor cells control the size of the endocrine population. Here we demonstrate that NEUROG3 is phosphorylated within the nucleus on serine 183, which catalyzes its hyperphosphorylation and proteosomal degradation. During progression through the progenitor cell cycle, NEUROG3 phosphorylation is driven by the actions of cyclin-dependent kinases 2 and 4/6 at G_1 /S cell-cycle checkpoint. Using models of mouse and human pancreas development, we show that lengthening of the G_1 phase of the pancreatic progenitor cell cycle is essential for proper induction of NEUROG3 and initiation of endocrine cell differentiation. In sum, these studies demonstrate that progenitor cell-cycle G_1 lengthening, through its actions on stabilization of NEUROG3, is an essential variable in normal endocrine cell genesis.

Graphical Abstract

AUTHOR CONTRIBUTIONS

SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures and can be found with this article online at http://dx.doi.org/10.1016/j.devcel. 2017.02.006.

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INTRODUCTION

Diabetes mellitus is characterized by chronic hyperglycemia resulting from the loss or dysfunction of the insulin-producing β cells located in the pancreatic islets. A current treatment for diabetes is to replace these damaged cells through islet transplantation (Shapiro et al., 2000), which is limited by donor tissue availability. Production of large numbers of functional β cells from human embryonic stem cells (hESCs) could address this unmet need. Over the past decade, efforts to generate these cells have culminated in " β -like cells," which resemble β cells yet remain functionally immature (Johnson, 2016; Kieffer, 2016; Pagliuca and Melton, 2013). However, the number of β -like cells that are formed varies between biological replicates and laboratories (Rezania et al., 2014), making consistent endocrine cell formation difficult and expensive (Rostovskaya et al., 2015). Understanding the mechanisms that control endocrine cell differentiation during pancreas development will uncover ways to more uniformly generate mature β -like cells that could be used to treat those with diabetes (McKnight et al., 2010).

Pancreas formation is marked by the appearance of Pdx1-expressing pancreatic progenitor cells (Gu et al., 2002) that rapidly differentiate into two populations by approximately embryonic day 12 (E12): the tip progenitors that are competent to form all pancreatic cell types and the trunk cells that are lineage-restricted to endocrine and ductal fates (Zhou et al., 2007). Expression of Neurog3 induces trunk progenitor cell commitment to the endocrine lineage in a cell-autonomous manner (Apelqvist et al., 1999) and is required for the formation of endocrine cells during both mouse (Gradwohl et al., 2000) and human development (McGrath et al., 2015). High induction of Neurog3 is critical for proper commitment to the endocrine lineage (Wang et al., 2010) with glucagon (α) cells forming earliest in development, followed by insulin (β), pancreatic polypeptide (PP), and somatostatin (δ) cells (Johansson et al., 2007). Upon activation of Neurog3, pancreatic progenitors exit the cell cycle and differentiate, a process that is partially driven by Neurog3-

dependent upregulation of *Cdkn1a/p21* (Desgraz and Herrera, 2009; Gu et al., 2002; Miyatsuka et al., 2011).

The decision either to exit the cell cycle and differentiate or to undergo cell division occurs during the G_1 phase of the cell cycle. Progression through the cell cycle is controlled by cyclins and cyclin-dependent kinases (CDKs). During late G_1 , the cyclin D/CDK4/6 and cyclin E/CDK2 complexes phosphorylate the retinoblastoma protein (Rb), resulting in the commitment to cell division with progression through the G_1 -S phase transition. During the development of some tissues, G_1 lengthening is positively correlated with progenitor differentiation (Lange and Calegari, 2010). This correlation suggests that the cell cycle itself may directly regulate differentiation by altering the stability of obligatory, lineage-establishing transcription factors. For example, the CDK inhibitor P27^{Xic1} promotes neurogenesis by stabilizing *Xenopus* (Vernon, 2003) and mouse neurogenic transcription factors (Nguyen et al., 2006) through reductions in their ubiquitin-mediated proteasomal degradation (Vosper et al., 2007, 2009; Roark et al., 2012).

While cell-cycle proteins, such as P21, have been implicated in endocrine differentiation downstream of Neurog3, cell-cycle changes that might underlie induction of Neurog3 itself have not been investigated. As such, the aim of this work was to determine whether cell cycling itself regulates endocrine pancreas differentiation through fine-tuning the stability of Neurog3. This work demonstrates that lengthening of the G_1 cell-cycle phase is necessary for NEUROG3 stabilization and its transcriptional activity. Furthermore, hyperphosphorylation by CDK2 and CDK4/6 in rapidly cycling cells leads to NEUROG3 degradation and maintenance of the progenitor fate. Herein, a mechanistic link between progenitor cell-cycle length and endocrine pancreas genesis has been defined, explaining why only a subset of cycling progenitors robustly express NEUROG3 and differentiate to endocrine islet cells.

RESULTS

Cell-Cycle Length Increases during Early Pancreatic Development

As cell-cycle lengthening has been correlated with differentiation of embryonic, neural, and hematopoietic stem cells (Lange and Calegari, 2010), we first set out to understand whether a similar paradigm exists in mouse pancreas development. To accomplish this, we used cumulative 5-ethynyl-2'-deoxyuridine (EdU) labeling to experimentally determine the length of cell-cycle phases in mouse pancreatic progenitors between E11.5 and E13.5 (Arai et al., 2011). This approach requires serial injections of EdU to label all S-phase cells in vivo. The time required for a cell that started in S phase to complete the rest of the cell cycle (G₂-M-G₁) and re-enter S phase (Figure 1A) is measured by the time to reach maximal EdU labeling, i.e., the growth fraction (GF; Figure 1B), and equals the cell-cycle length (T_C; Figure 1B) less the S-phase length (T_S) (X₀ = T_C-T_S; Figure 1B). Segmental linear regression establishes the y intercept (y-int = T_S/T_C * GF; Figure 1B) and slope (m = GF/T_C). Using these data, T_S = y-int/m and T_C = y-int/m + X₀. To measure the length of G₂-M (T_{G2-M}), we experimentally determine the time for all EdU-labeled cells to transit through G₂ and enter mitosis (pHH3⁺) following a single bolus of EdU (Figure 1E). The length of G₁ (T_{G1}) can then be calculated by T_C - T_S - T_{G2-M}.

Using this approach, the time for Pdx1 immunoreactive (+) pancreatic progenitors to reach the GF was 6.50 \pm 0.38, 7.98 \pm 0.51, and 9.26 \pm 0.60 hr at E11.5, E12.5, and E13.5, respectively (Figures 1B–1D). T_C increased from 8.20 \pm 0.68 to 11.70 \pm 0.93 hr while both T_S and T_{G2-M} remained similar for each developmental day (Figures 1B–1E and Table 1). Therefore, the increases in total cell-cycle length were solely due to G_1 phase lengthening from 4.54 \pm 0.40 to 7.19 \pm 0.83 hr (Figure 1I and Table 1).

Close examination of the EdU labeling revealed that the EdU⁺ cells appeared in clusters near the tips prior to the trunks of the epithelium (Figure S1), suggesting a difference in cell-cycle length between the Pdx1⁺Cpa1⁺ tip and the Pdx1⁺Cpa1⁻ trunk progenitors. The time to reach the GF increased from E11.5 to E13.5 in both tip $(5.70 \pm 0.33 \text{ to } 7.69 \pm 0.34 \text{ hr})$ and trunk cells $(6.38 \pm 0.42 \text{ to } 9.43 \pm 0.64 \text{ hr})$ (Figures 1F–1H). In addition, on each developmental day the time to reach the GF differed between the tip and trunk cells, resulting in a G₁ phase length that is ~25% longer in the more differentiated trunk progenitors $(4.41 \pm 0.43 \text{ to } 7.36 \pm 0.64 \text{ hr})$ than in tip progenitors $(3.74 \pm 0.35 \text{ to } 5.62 \pm 0.34 \text{ hr})$ (Figure 1J and Table 1). Despite the increased G₁ length in trunk cells, their T_C was similar due to a reduction in the T_S (Figure 1J and Table 1). Taken together, these findings demonstrate that G₁ lengthening occurs within both progenitor pools during pancreas development.

Cell-Cycle Lengthening in Trunk Cells Allows Endocrine Differentiation during Pancreatic Development

To understand whether G₁ lengthening has functional consequences for endocrine pancreas development, we utilized two mouse models to manipulate pancreatic progenitor G_1 phase length. We previously noted (Chamberlain et al., 2014), and confirm here, that heterozygous Kras loss-of-function (Kras^{LSL}) pancreata have significantly more Neurog3 cells at E13.5 (Figures S2A and S2B). Since Kras activates cell cycling by driving G₁/S progression (Wennerberg et al., 2005), we hypothesized that the 1.2-fold increase in Neurog3⁺ cells in the Kras^{LSL} embryos resulted from G₁ lengthening. To determine whether Kras expression levels affected cell-cycle length, we carried out EdU labeling for 3.5 hr before harvest on E12.5 (Figure S2C). This EdU schedule targets the steep, linear phase of the labeling curve at E12.5 (Figure 1) and therefore enables inference of small changes in cell-cycle length. The proportion of Pdx1⁺EdU⁺ pancreatic progenitors was significantly reduced in Kras^{LSL} embryos at E12.5 (Figure S2D). While not significant, the Kras gain-of-function (Kras^{G12D}) embryos had slightly more EdU-labeled progenitor cells (Figure S2D), suggesting that they cycle more quickly. As the majority of Neurog3⁺ cells derive from the trunk cells, we next investigated whether changes in Kras signaling differentially affect the cell cycle of tip and trunk cells. EdU⁺ tip and trunk cells decreased in the Kras^{LSL} model and increased in the Kras^{G12D} model (Figures S2E and S2F) without affecting total cell numbers (Figure S2G). Notably, Kras^{LSL} tip cell EdU incorporation rates were similar to those of wild-type (WT) trunk cells, suggesting that their cell-cycle parameters and developmental competencies are similar (Figures S2D and S2E). In addition, altering Kras signaling during embryonic development does not significantly change the proportion of tip and trunk progenitor cells (Figure S2H). Taken together, these results suggest that altering cell-cycle length during mouse pancreatic development affects the generation of Neurog3⁺ endocrine progenitors.

To more directly test the consequence of altering G_1 length, we ectopically expressed Cdkn1b—a cyclin-dependent kinase inhibitor not normally expressed in Sox9-expressing pancreatic progenitor cells that blocks the G_1/S transition (Miyatsuka et al., 2011)—in a doxycycline-inducible manner. Control embryos were either single transgenic (ST) Sox9rtTA or tetO-Cdkn1b littermates of Cdkn1b-expressing Sox9-rtTA;tetO-Cdkn1b double transgenic (DT) embryos. Pregnant dams were injected with doxycycline on the morning of E12.5 and embryos were collected for analysis 9.5 hr later (Figure 2A). Both Cdkn1b⁺ and Neurog3⁺ cells increased significantly (Figures 2A and 2B), suggesting that Cdkn1b expression is sufficient to increase the number of Neurog3⁺ cells without altering total cell number (Figure 2C). To determine whether ectopic Cdkn1b expression slowed S-phase transition, doxycycline was given from E10.5 to E12.5, and EdU-labeling was carried out for 3.5 hr before harvest on E12.5 (Figure 2D) as described above. As expected, the proportion of EdU⁺ pancreatic progenitor cells significantly decreased in DT embryos (Figure 2E). To ensure that ectopic Cdkn1b expression did not cause cell-cycle exit, we quantified the GF using 11 hr of EdU labeling (GF; Figure 1C); no differences were observed between control ST and DT mice (Figure 2F).

Notably, 48 hr of Cdkn1b expression resulted in a 2.5-fold increase in both chromogranin A $(Chga)^+$ cells (Figure 2G) and glucagon (Gcg; $a)^+$ cells at E12.5 (Figure 2H) but did not significantly affect overall cell numbers (Figure 2I). These results suggest that the increase in Neurog3⁺ endocrine progenitors caused an increase in endocrine cell formation that was predominantly of the α -cell lineage. As competency of Neurog⁺ cells is predominantly down the α -cell lineage during early pancreas development (Johansson et al., 2007), we next asked whether Cdkn1b expression promotes the formation of other endocrine cell types later in pancreas development by administering doxycycline from E12.5 to E14.5 (Figure 2J). This doxycycline dosing regimen resulted in a significant decrease in the number of EdU⁺ cells at E14.5 after 3.5 hr of EdU labeling (Figure 2K) and a 3.3-fold increase in Chga⁺ endocrine cells at E14.5 (Figure 2L). In addition, the numbers of Gcg⁺ (2.8-fold; Figure 2M) and Ins⁺ (1.4-fold; Figure 2N) cells increased without affecting pancreatic size (Figure 2O). Consistent with their formation later in pancreas development, we did not observe any changes in the formation of δ or PP cells (data not shown). Combined with studies reported in Figure 1, these in vivo studies suggest that G_1 lengthening permits efficient Neurog3 expression and endocrine differentiation. Furthermore, ectopic expression of Cdkn1b, a G1-S CDK inhibitor, suggests that the activities of G_1 -S CDKs in the progenitor cells prevents efficient endocrine differentiation and maintains progenitor cell state.

Serine 174 and 183 Are Critical NEUROG3 Phosphorylation Sites

As it has previously been demonstrated that *Neurog3* message expression is more prevalent than its protein expression in rapidly cycling pancreatic progenitors (Bechard et al., 2016; Villasenor et al., 2008), we next sought to understand how NEUROG3 stabilization in pancreatic progenitors results in endocrine differentiation. Since CDKs regulate protein phosphorylation in a cell-cycle-dependent manner, we hypothesized that NEUROG3 phosphorylation in rapidly cycling progenitors results in its degradation. At its peak of expression in the mouse embryonic pancreas (Villasenor et al., 2008) and as previously noted (Wang et al., 2009), Neurog3 displays multiple distinct electrophoretic mobilities,

consistent with phosphorylation (Figure 3A). To determine which regions of NEUROG3 are phosphorylated, we generated three N-terminally FLAG-tagged human NEUROG3 truncation mutants. In addition, a phospho-mutant (T120A) was generated that is analogous to the T118 phosphorylation site of *Xenopus* Neurog2. Threonine 118 is required for the interaction and complex formation of Neurog2 with its heterodimerization partner, BETA-2/E12/E47, and acts to counter ubiquitination, prolonging its half-life (Vosper et al., 2007). The slower migrating forms of NEUROG3 were sensitive to phosphatase treatment, but 145–187 showed only a single, phosphatase-insensitive band (Figure 3B), implying the presence of key phosphorylation sites between the helix-loop-helix (HLH) and activation domains of NEUROG3. We were unable to visualize 6–78 on western blot despite significant effort (data not shown), possibly, as outlined below, because it lacks nuclear localization sequences important for stabilization.

As CDKs are serine/threonine kinases, we mutated each of the nine serine residues in the 145–187 region to alanine. Remarkably, mutating S183 alone prevented all phosphorylation, suggesting that S183 phosphorylation seeds subsequent multi-site phosphorylation (Figure 3C). A series of multi-serine mutations revealed that only S174 and S183 were phosphorylated when all the other serine residues were mutated (arrow and arrowhead in Figure 3D), indicating that in addition to S183, S174 can act as an initial phosphorylation target. A third series of mutants that leave S183 and a second serine intact revealed a third band in the presence of S161, S165, S174, S177, and S181 (arrowhead in Figure 3E). Combined, these data show that NEUROG3 is subject to sequential phosphorylation initiated by phosphorylation of S183.

NEUROG3 Phosphorylation Regulates Transcriptional Activity, but Not Nuclear Localization

Phosphorylation commonly serves to change transcription factor activity. Using a NEUROG3-responsive *PAX4* luciferase reporter (Smith et al., 2003), the functional consequences of NEUROG3 phosphorylation were assessed. As previously reported for T118 of *Xenopus* Neurog2, T120A showed reduced transcriptional activity relative to WT NEUROG3 (Figure 4A), as did 6–78, R93L (Wang et al., 2006), and removal of the HLH domain. S183A and a number of other serine mutations increased trans-activation using this reporter (Figure 4A). Furthermore, induction of *Pax4*, *NeuroD1*, and *Sst* expression in mPAC cells (Gasa et al., 2004) were significantly increased following transfection of the NEUROG3 phosphomutants (S183A, 8A/S183; Figures 4B–4D). Therefore, phosphorylation of the C-terminal region of NEUROG3 reduces transcriptional activity.

Phosphorylation of NEUROG3 could influence transcriptional activity by altering NEUROG3 subcellular localization. WT NEUROG3 accumulated exclusively in the nucleus (Figure S3A) but 6–78 showed impaired nuclear targeting (Figure S3B), suggesting that the N-terminal region harbors a nuclear localization sequence (NLS). Mutations of the other domains did not affect nuclear localization (Figure S3 and S4; data not shown), although

145–187 was enriched in the nucleoli marked by fibrillarin staining (Figure S3E). Transfected mPAC cells showed similar nuclear distributions of NEUROG3, S183A, and 8A/S183 (data not shown); thus, altered nuclear import does not explain the differences in transcriptional activity of the C-terminal NEUROG3 phosphomutants.

NEUROG3 Hyperphosphorylation Requires Nuclear Import

As C-terminal phosphorylation did not alter NEUROG3 subcellular localization, we next asked whether subcellular location affects NEUROG3 phosphorylation. Alanines were substituted for arginine and lysine in the predicted NLSs at 58–64, 79–82, and 84–87. None of the three individual region mutants reduced nuclear localization (Figures S3F–S3I), but mutating two out of the three NLSs impaired nuclear import (Figures S3J–S3M), even when nuclear export was blocked with leptomycin B (Figures S4A–S4J). As expected, mutating two out of the three NLSs also impaired NEUROG3 transcriptional activity (Figure S4P). Thus, NEUROG3 contains two redundant NLSs (58–64; 84–87), one of which overlaps with the basic DNA-binding domain (Figure 3B), as is common in other transcription factors (Cokol et al., 2000).

NLS mutation also reduced phosphorylation (Figures S4Q and S4R). However, the presence of a slower migrating form, most obvious for the cytoplasmically localized NLS58–64/84–87A (Figure S4Q, arrow), shows that some phosphorylation can occur outside the nucleus. Overexpression of E47, the basic HLH (bHLH) heterodimer partner of NEUROG3, rescued nuclear localization of the NLS mutants, but not the NLS58–64/84–87A/T120A mutant (Figures S4K–S4O), suggesting that E47 complexes with NEUROG3 outside the nucleus in a T120-phosphorylation-dependent manner and then tows NEUROG3 into the nucleus, as previously described for Neurog2 (Vosper et al., 2007). In sum, these studies show that C-terminal hyperphosphorylation of NEUROG3 occurs in the nucleus where the cell-cycle kinases typically function (Fiaschi-Taesch et al., 2013).

Phosphorylated NEUROG3 Is Degraded during G₂-M

As NEUROG3 phosphorylation reduced transcriptional activity, we next asked whether phosphomutant forms of NEUROG3 were able to elicit one of its main biological roles in the pancreas: to regulate G_1 cell-cycle arrest (Miyatsuka et al., 2011). Transfected HeLa cells were stained for DNA content and pHH3 to infer cell-cycle stage using flow cytometry. Overexpression of WT or phosphomutant NEUROG3 reduced cells in G_2 -M by 2-fold and increased G_1 -phase cells by 15% (Figure 5A), confirming that phosphomutant NEUROG3 remains able to arrest cells in G_1 . Only deletion of the HLH domain, which is essential for NEUROG3 dimerization with E47 and transcriptional activation of their targets (Figure 4A), abolished this property (Figure 5A). This observation demonstrates that NEUROG3 requires its bHLH binding partner, E47, and subsequent transcriptional activation (e.g., P21), but not phosphorylation, to arrest cells in G_1 .

To determine whether NEUROG3 is phosphorylated in a cell-cycle-dependent manner, we isolated G_1 , S, and G_2 -M populations by fluorescence-activated cell sorting (FACS), and used western blots to quantify the fastest, intermediate, and slowest migrating NEUROG3 bands. For each phosphoform of WT NEUROG3, expression was notably lowest in G_2 -M (Figure 5B). Within every cell-cycle stage, the unphosphorylated (fast) and maximally phosphorylated (slow) species of WT NEUROG3 were present at similar levels, while the

level of intermediately phosphorylated forms was significantly lower (Figure 5B, red bars). NEUROG3 _{HLH}, which does not cause differentiation and G₁ arrest (Figure 5A), had similar levels of the unphosphorylated and intermediately phosphorylated species at each stage, but the level of the maximally phosphorylated form was significantly lower (Figure 5B, blue bars). Furthermore, the relative amounts of total WT NEUROG3 were highest at S phase and lowest at G₂-M phase, whereas NEUROG3 _{HLH} levels were equal at G₀-G₁ and S phase while only slightly lower at G₂-M phase (cf. total red and blue bar heights in Figure 5B). Combined with the findings that WT NEUROG3 arrests the cells during S phase while NEUROG3 _{HLH} does not (Figure 5A), these results suggest that maximally phosphorylated WT NEUROG3 levels decline beyond S phase. Therefore, when NEUROG3 cannot heterodimerize to drive target gene transcription—which causes cell-cycle exit and differentiation—it is rapidly degraded.

To monitor the synthesis and degradation of NEUROG3, we generated fusion proteins with a fluorescent timer (FT) that changes its emission spectrum from green to red over several hours (Miyatsuka et al., 2009; Subach et al., 2009). Significantly fewer WT-NEUROG3 transfected cells expressed red protein than controls (FT only or FT-NEUROD1; Figure 5C), demonstrating that NEUROG3 has a short half-life and suggesting its presence promotes degradation of fused FT. Mutation of a putative ubiquitin/SUMO target residue (K121A) or the phosphorylation initiation site (S183A) stabilized fused FT, while mutating the analogous SUMOylation (K139A) site in NEUROD1 had no effect (Figure 5C). Blocking nuclear export (leptomycin B; Figure S4) or inhibiting the proteasome (MG132) significantly increased red WT-NEUROG3-expressing cells (Figure 5D), demonstrating that NEUROG3 is targeted for proteasomal degradation outside the nucleus. Coexpression of NEUROG3 heterodimerization partner E47 significantly increased the red population of WT-NEUROG3-expressing cells but not that of cells expressing T120A (Figure 5E), showing that E47 interaction stabilizes NEUROG3. Notably, S183A significantly increased NEUROG3 half-life (Figures 5C and 5E), confirming that S183 destabilizes NEUROG3 by allowing phosphorylation and degradation. These results indicate that phosphorylated NEUROG3 is targeted for cytoplasmic proteosomal degradation during G₂-M and that the S183 is the phosphorylation seed region that initiates this degradation.

NEUROG3 Is Phosphorylated by G₁ CDKs

The residues in Neurog1 and Neurog2 analogous to S183 in NEUROG3 are reportedly phosphorylated by ERK5 (Cundiff et al., 2009) and GSK3 (Ma et al., 2008), respectively. In addition, we found that CDK, p38 MAPK, and JNK family members are predicted candidates for the phosphorylation of NEUROG3 (Linding et al., 2007). Inhibiting ERK5/ MAPK7 (BIX02189), MAPK (LY2228820), JNK (SP600125), or GSK3 (SB216763) did not alter phosphorylation of NEUROG3 in HeLa cells (Figure 6A), suggesting that other kinases, such as CDKs, are responsible for phosphorylation of NEUROG3.

Blocking the HeLa cell cycle in M phase with nocodazole increased the hyperphosphorylated form of NEUROG3, supporting the model that NEUROG3 is progressively phosphorylated as cells transition through the cell cycle by CDKs (Figure 6B).

In addition, the CDK2 inhibitor roscovitine reverses nocodazole-stimulated NEUROG3 hyperphosphorylation (Bach et al., 2005; Bain et al., 2007).

As overexpression of Cdkn1b, an inhibitor of G_1 -S CDKs (i.e., CDK2, CDK4, CDK6), increased the number of NEUROG3⁺ cells (Figure 2B), the effect of inhibition of Cdk2, Cdk4, and Cdk6 using specific small-molecule inhibitors was next investigated in mouse embryonic pancreas explants. Embryonic pancreata were isolated at E11.5 and treated ex vivo for 24 hr with Cdk4/6 inhibitor PD-03329911 and Cdk2 inhibitors ii and iii (CDKi). This resulted in a 3-fold increase in the number of Neurog3⁺ cells without altering the total number of Sox9⁺ progenitor cells (Figure 6C), as assessed by immunofluorescence. Furthermore, treating human pancreatic progenitors, derived from human induced pluripotent stem cells (iPSCs) using a modified version of the Rezania et al. (2014) protocol, with CDKi for 24 hr also increased the expression of NEUROG3 protein detected by western blot (Figure 6D). These findings further bolster the role of CDKs in the destabilization of NEUROG3 during development.

To investigate the consequences of CDK inhibition on endocrine differentiation, we generated *NEUROG3-2A-eGFP* knock-in reporter CyT49 hESC lines using CRISPR/Cas9 (Figure S5). The three clonal lines that were tested (N2-2, N4-7, and N5-5) all formed definitive endoderm with similar efficiency to the parental CyT49 hESC line (Figure S5A). However, upon further differentiation to endocrine progenitors (stage 6), N5-5 had higher efficiency of forming NEUROG3⁺ cells (Figure S5B). To confirm the fidelity of the GFP fluorescence to the endocrine lineages, we differentiated *NEUROG3* reporter lines to stage 6 and GFP⁺ and GFP⁻ cells were FACS sorted prior to gene expression analyses. As expected, GFP⁺ cells had significantly increased expression of *NEUROG3* and *INSULIN* (Figure S5C) compared with GFP⁻ and CyT49 cells. Conversely, *SOX9* expression was enriched in GFP⁻ cells compared with GFP⁺ (Figure S5C). For further studies, N5-5 was used to identify NEUROG3⁺ lineage cells based on GFP fluorescence.

CDKi treatment of human endocrine progenitors derived from N5-5 significantly increased the number of GFP⁺ cells by 1.7-fold as determined by flow cytometry (Figure 6E). Consistent with our findings that phosphorylation of NEUROG3 reduces its transcriptional activity (Figure 4), the expression of *NEUROG3* and *NEUROD1*, which are direct targets of NEUROG3, and *PDX1*, which is a β/δ -cell marker, were all increased in NEUROG3-lineage cells following CDKi treatment (Figures 6F–6H).

To investigate the role of the cell cycle and CDK inhibition on human endocrine cell differentiation, we generated a transgenic CyT49 hESC cell-cycle reporter line using fluorescence ubiquitination-based cell-cycle indicator (FUCCI) (Sakaue-Sawano et al., 2008). This line (FUCCI-3) allows for the facile isolation of cells in G_1 (red) versus S-G₂-M (green) based on fluorescence. Treating stage-6 cells derived from the FUCCI-3 line with CDKi increased the proportion of cells in G_1 from 85% to 93%, suggesting the CDKi treatment slows the G_1 phase (Figure 6I). FACS was next used to isolate G_1 and S-G₂-M cells following CDKi and gene expression analyses were performed, which indicated that expression of *NEUROG3*, *NEUROD1*, and *PDX1* are significantly upregulated in G_1 compared with S-G₂-M cells (Figures 6J–6L). In addition, CDKi treatment significantly

increased the expression of both *NEUROG3* and *PDX1* in G_1 -phase cells compared with control vehicle-treated cells (Figures 6J and 6L).

Taken together, these findings demonstrate that G_1 lengthening occurs during mouse and human pancreatic endocrine genesis. This lengthening allows for NEUROG3 stabilization due to reduced activity of CDKs that are not active until the G_1 /S transition, resulting in the NEUROG3 transcriptional regulation that drives cell-cycle exit and endocrine differentiation. Conversely, in rapidly dividing progenitor cells with a short G_1 length, the combined actions of CDK2 and CDK4/6 lead to phosphorylation of NEUROG3 at the G_1 /S transition and rapid proteosomal degradation in the cytosol as cells progress through S- G_2 -M.

DISCUSSION

Pancreatic endocrine cell development is initiated by Neurog3 activation within bipotent trunk progenitor cells. Following the initiation of endocrine differentiation, silencing of further Neurog3 expression ensures normal endocrine differentiation and function (Miyatsuka et al., 2011). Prior to this work, it was not clear how Neurog3 was stabilized within a subset of trunk progenitors, nor was it understood how Neurog3 was rapidly degraded after initiating differentiation. This study aimed to address these two open questions.

Building on previous work (Bankaitis et al., 2015; Kim et al., 2015) we demonstrate that the mouse pancreatic progenitor cell cycle lengthens as a result of G_1 phase lengthening from 4.5 to 7.2 hr between E11.5 and E13.5. Furthermore, we show that this progenitor cell-cycle lengthening is required for efficient induction of Neurog3 and that CDKs active in cycling cells target Neurog3 for proteosomal degradation. This work demonstrates that cycling of pancreatic and endocrine progenitors may be essential for the dynamic regulation of Neurog3.

While our study identifies changes in G_1 length during pancreas development, there are several studies that have noted this phenomenon in other cell types. For instance, during neural development the length of G_1 increases from 3.2 to 12.4 hr over the course of 5 days in the pseudostratified ventricular epithelium, the region of the brain where the majority of neocortical neurons originate (Takahashi and Nowakowski, 1995). In addition, hESCs have a short cell-cycle length due to a reduced length of the G_1 phase (Becker et al., 2006), and differentiation is accompanied by an increase in G_1 length (Becker et al., 2010). Also, hESCs isolated in the G_1 phase have a greater propensity to differentiate than cells isolated in S and G_2 phases (Sela et al., 2012).

The data presented in this work support and extend the "cell-cycle length hypothesis" put forward for neural development (Salomoni and Calegari, 2010). This hypothesis posits that a particular threshold for a cell-fate determinant (e.g., NEUROG3) is required for differentiation to occur. By lengthening the G₁ phase of the cell cycle, more time is provided for the accumulation of Neurog3, which then surpasses its threshold to transcriptionally activate CDK inhibitors, such as *Cdkn1a/p21*, resulting in cell-cycle arrest and endocrine

differentiation. Consistent with this model, we find that transcription of *NEUROG3* is upregulated in the G_1 phase of the cell cycle during human development and that G_1 CDKs, whose actions result in the G_1 /S phase transition, act to destabilize NEUROG3. Notwithstanding the evidence, it remains possible that the G_1 CDKs do not directly phosphorylate NEUROG3 but initiate a phosphorylation cascade that results in NEUROG3 destabilization. We hope that our future work will uncover whether additional kinases phosphorylate NEUROG3 and determine whether these kinases regulate NEUROG3 in vivo.

In mice, the existence of a population of pancreatic progenitor cells with low expression of *Neurog3* mRNA but no detectable Neurog3 protein has been reported (Bechard et al., 2016; Villasenor et al., 2008). This could be explained by the rapid phosphorylation and degradation of Neurog3 due to high CDK expression. While Neurog3 can be detected for a few days in the developing pancreas, it lasts for only a few hours in individual cells (Miyatsuka et al., 2009), consistent with the rapid turnover of NEUROG3 protein (Figure 5). This interplay between cell-cycle machinery and differentiation has been demonstrated during mammalian neural development whereby inhibition of Cdk2 (Calegari and Huttner, 2003) or Cdk4 drives differentiation (Calegari et al., 2005). We have identified other transcription factors important for progenitor cell maintenance that we believe are stabilized during the G_1/S transition by cell-cycle kinases and ensure rapid transit through this checkpoint to maintain progenitor fate (F.C.L. and N.A.J.K., unpublished data). Future studies will be targeted at elucidating how cell-cycle kinases and transcription factor stability are inter-related and interdependent.

An interesting question that arises from this work, and the cell-cycle length hypothesis, is how does G_1 lengthening occur? Although we did not set out to answer this question, it likely results from both cell-autonomous and/or non-cell-autonomous mechanisms (Kim et al., 2015). For instance, this study suggests there is a complex relationship between cellcycle length and differentiation that is regulated, cell-autonomously, by relative levels of transcription factors. It is also possible that factors secreted by the mesenchyme could play a non-cell-autonomous role in regulating G_1 length. For example, mesenchymal Fgf10 is required for the proliferation of Pdx1⁺ pancreatic progenitors (Bhushan et al., 2001). The shorter G_1 length of the tip progenitors, which are in apposition with surrounding mesenchyme, supports a potential role for a mesenchymal secreted factor. Analysis of cellcycle length in models of impaired endocrine differentiation, using approaches developed herein, could elucidate mechanisms that endogenously control G_1 lengthening during pancreas development.

In addition to the CDK-dependent phosphorylation of NEUROG3, the half-life of Neurog3 is controlled by other post-translational modifications (PTMs) (Barrow et al., 2005; Miyatsuka et al., 2009). Mouse Neurog3 can be ubiquitinated on lysines as well as non-canonical residues (Roark et al., 2012). Mutating the lysine residues doubles the half-life of Neurog3 (K121A; Figure 5). In addition, the slow-migrating, phosphatase-insensitive form of NEUROG3 observed in the 8A/S183 lane of Figure 3D (double arrow) is likely additional evidence of ubiquitination or SUMOylation. It remains unclear whether there is a direct link between phosphorylation and other PTMs that target NEUROG3 for degradation, and future studies will aim to address this question. Furthermore, how NEUROG3 PTMs affect its

interactions with other proteins in the nucleus and its target gene repertoire are yet undescribed.

Studying the role of cell-cycle proteins during pancreas development has largely been through the use of knockout mice. Interpreting these studies is difficult, as altering expression of cell-cycle proteins could reduce overall numbers of pancreatic progenitors, which dictates the final size of the adult organ and its physiological function (Stanger et al., 2007). For instance, loss of *Cdk4* leads to reduced proliferation of $Pdx1^+$ pancreatic progenitors at E12.5, possibly due to a defect in pancreatic mesenchyme, resulting in a reduced number of Neurog3⁺ cells (Kim and Rane, 2011). However, our studies suggest that temporary reductions in Cdk4 may result in stabilization of Neurog3. It would be interesting to revisit this study using an inducible system that allows for spatiotemporal control of Cdk4 knockout in vivo.

In summary, this work shows that cell-cycle length regulates the balance between proliferation and differentiation during pancreatic development. In particular, G_1 lengthening, NEUROG3 stabilization by minimizing phosphorylation, and subsequent NEUROG3-driven cell-cycle exit combine to ensure that endocrine differentiation occurs in non-dividing cells. This research uncovers a complex and interacting set of protein modifications that govern NEUROG3 activity and, in turn, endocrine cell generation. Given the critical importance of pancreatic endocrine cell generation and the potential clinical utility of methods for controlling it, a better understanding of this complex system and the signals that influence it should be the subject of future investigation.

STAR * METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Donkey polyclonal Anti-mouse IgG	Jackson ImmunoResearch	Cat#715-001-003; RRID: AB_2307338
Goat polyclonal anti-Cpa1	R&D systems	Cat#AF2765; RRID: AB_2085841
Guinea Pig anti-NEUROG3	Schwitzgebel et al., 2000	German Lab
Guinea Pig polyclonal anti-Insulin	DAKO	Cat#A056401-2; RRID: AB_2617169
Mouse monoclonal anti-FLAG	Sigma-Aldrich	Cat#F3165; RRID: AB_259529
Mouse monoclonal anti-Glucagon	Sigma-Aldrich	Cat#G2654; RRID: AB_259852
Mouse monoclonal anti-Neurog3	Developmental Studies Hybridoma Bank	Cat#F25A1B3; RRID: AB_528401
Mouse monoclonal anti-Pdx1	Developmental Studies Hybridoma Bank	Cat#F109-D12; RRID: AB_1157903
Rabbit polyclonal anti-ChromograninA	Thermo Scientific	Cat#RB-9003-P1; RRID: AB_149730
Rabbit polyclonal anti-P27	Cell Signaling Technology	Cat#2552S; RRID: AB_10693314
Rabbit polyclonal anti-pHH3	Cell Signaling Technology	Cat#9701
Rabbit polyclonal anti-Sox9	Millipore	Cat#AB5535; RRID: AB_2239761
Rabbit polyclonal anti-β-actin	Abcam	Cat#AB8227; RRID: AB_2305186

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Sheep polyclonal anti-NEUROG3	R&D systems	Cat#AF3444; RRID: AB_2149527
Biological Samples		
CyT49 human embryonic stem cell (passages 20-p31)	Viacyte	CyT49; NIH hESC-10-0041
Human Peripheral Blood Mononuclear Cells	STEMCELL Technologies	Cat#70025.1
Chemicals, Peptides, and Recombinant Proteins		
5'-Ethynyl-2'-deoxyuridine (EdU)	Sigma-Aldrich	Cat#T511285
Alexa Fluor 594 Carboxamido-(6-Azidohexanyl), Triethylammonium Salt	ThermoFisher Scientific	Cat#A10270
Ascorbate	Sigma-Aldrich	Cat#A4544
Accutase	STEMCELL Technologies	Cat#07920
Activin A	E-biosciences	Cat#34-8993-85
Alexa-Fluor 594 azide	ThermoFisher Scientific	Cat#A10270
BIX02189	Tocris Biosciences	Cat#4842
Bovine Serum Albumin	Sigma-Aldrich	Cat#A3803
Cdk2 inhibitor II	EMD Millipore	Cat#219445
Cdk2 inhibitor III	EMD Millipore	Cat#238803
CHIR-99021	Sigma-Aldrich	Cat#SML1046
Copper Sulfate	Sigma-Aldrich	Cat#451657
DAPI	Sigma-Aldrich	Cat#D9542
DMEM/F12	Corning	Cat#15-090-CV
Doxycycline	Sigma-Aldrich	Cat#D9891
FBS (GIBCO)	ThermoFisher Scientific	Cat#12483020
Gamma secretase inhibitor XX	EMD Millipore	Cat#565789
Geltrex	ThermoFisher Scientific	Cat#A1433302
Glutamax	ThermoFisher Scientific	Cat#35050061
Herugulin	PeproTech	Cat#100-03
Hoescht 33342	ThermoFisher Scientific	Cat#H1399
Horse Serum (GIBCO)	ThermoFisher Scientific	Cat#16050122
Hyclone D-PBS	GE Healthcare Bio-Science	Cat#SH30028.01
Hyclone DMEM	GE Healthcare Bio-Science	Cat#SH30081
Hyclone Penicillin/Streptomycin (P/S)	GE Healthcare Bio-Science	Cat#SV30010
Insulin-Transferrin-Selenum (ITS-X)	ThermoFisher Scientific	Cat#51500056
KGF	R&D Systems	Cat#251-KG
LDN193189	EMD Millipore	Cat#509882
Leptomycin B	Sigma-Aldrich	Cat#L2913
LY2228820	SelleckChem	Cat#S1491
MCDB 131	USBiological Life Sciences	Cat#E3000-01B
MG132	Cayman	Cat#1211877-36-9
Nocodazole	Sigma-Aldrich	Cat#31430-18-9
Paraformaldehyde	Sigma-Aldrich	Cat#P6148

REAGENT or RESOURCE	SOURCE	IDENTIFIER
PD-0332991	Sigma-Aldrich	Cat#PZ0199
Primestar DNA polymerase	Takara Clontech	Cat#R010A
Puromycin	Sigma-Aldrich	Cat#P8833
QuickExtract	Epicentre	Cat#QE09050
ReLeSR	STEMCELL Technologies	Cat#05872
ReproTESR	STEMCELL Technologies	Cat#05920
Repsox	Sigma-Aldrich	Cat#R0158
Restriction Enzymes	New England Biolabs	N/A
Retinoic Acid	Sigma-Aldrich	Cat#R2625
Roscovitine	Sigma-Aldrich	Cat#R7772
SANT-1	Tocris Biosciences	Cat#1974
SB216763	SelleckChem	Cat#S1075
Sodium Bicarbonate	Sigma-Aldrich	Cat#S6297
SP600125	Sigma-Aldrich	Cat#S5567
Т3	Sigma-Aldrich	Cat#6397
TeSR-E8 Kit for hESC/hiPSC Matinenance	STEMCELL Technologies	Cat#05940
To-PRO-3 Iodide	ThermoFisher Scientific	Cat#T3505
Tris Base	Fisher Scientific	Cat#BP152
Tris HCl	Fisher Scientific	Cat#BP153
Triton X-100	Sigma-Adrich	Cat#T8787
Trizol	ThermoFisher Scientific	Cat#15596018
Vibrant DyeCycle Violet	ThermoFisher Scientific	Cat#V35003
XenoFree KOSR	ThermoFisher Scientific	Cat# A1099202
Zinc Sulfate	Sigma-Aldrich	Cat#Z0251
a-Amyloid Precursor Protein Modulator	EMD Millipore	Cat#565740
λ-phosphatase	New England Biolabs	Cat#P0753
Critical Commercial Assays		
CytoTune-iPS 2.0 Sendai Reprogramming Kit	ThermoFisher Scientific	Cat#A16517
Dual Luciferase Assay	Promega	Cat#E1960
Erythroid Progenitor Reprogramming Kit	STEMCELL Technologies	Cat#05924
Lipofectamine 2000	ThermoFisher Scientific	Cat#11668027
Lipofectamine 3000	ThermoFisher Scientific	Cat#L3000015
QuickExtract DNA Extraction Solution	Epicentre	Cat#QE09050
Experimental Models: Cell Lines		
Human: FUCCI-3: CyT49 FUCCI-3 transgenic hESC line	This paper	N/A
Human: N5-5: CyT49 Neurog3-2A-eGFP hESC line	This paper	N/A
Experimental Models: Organisms/Strains		
Mouse: CD-1	Charles River	Crl:022

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Mouse: Pdx1-Cre: B6.Cg-Tg(Pdx1-cre)89.1Dam/Mmuco	Mutant Mouse Regional Resource Centers	MMRRC:015970-UCD
Mouse: Kras ^{LSL-G12D} : B6.129S4-Krastm4Tyj/J	The Jackson Laboratory	JAX:008179
Mouse: TetO-Cdkn1b: C57BL/6(tetO-Cdkn1b)1Scpr/J	The Jackson Laboratory	JAX:017613
Mouse: Sox9-rtTA: C57BL/6-BAC-Sox9-rTTA	This paper	N/A
Recombinant DNA		
Hnf1a expression plasmid	Smith et al., 2003	N/A
pCAGGS-FUCCI	This paper	N/A
pCCC	Krentz et al., 2014	N/A
pCCC-LL502/3	This paper	N/A
px458	Ran et al., 2013	Addgene #48138
Sequence-Based Reagents		
Mouse Gusb (Mm00446953_m1)	ThermoFisher Scientific	Cat#4331182
Mouse Neurod1 (Mm01280117_m1)	ThermoFisher Scientific	Cat#4331182
Mouse Pax4 (Mm01159036_m1)	ThermoFisher Scientific	Cat#4331182
Mouse Sst (Mm00436671_m1)	ThermoFisher Scientific	Cat#4331182
Human NEUROD1 probe (FAM/ZEN/IaBKFQ): CGCCAGTTTCACCATTTCCGGG	IDT	N/A
Human NEUROD1 primer 1: TCCTTCGATAGCCATTCACATC	IDT	N/A
Human NEUROD1 primer 2: GCTGCCTTTTGTAAACACGAC	IDT	N/A
Human NEUROG3 gRNA: GGGTCGCTCCTCCAGCGACG	IDT	N/A
Human NEUROG3 probe (FAM/ZEN/IaBKFQ): AGCGGTCCTTCCCCAGAGC	IDT	N/A
Human NEUROG3 primer 1: TCTATTCTTTTGCGCCCGGTAG,	IDT	N/A
Human NEUROG3 primer 2: GCAGGTCACTTCGTCTTCC	IDT	N/A
Human PDX1 probe (FAM/ZEN/IaBKFQ): CGC TTGTTCTCCTCCGGCTCC	IDT	N/A
Human PDX1 primer 1: TGAAGTCTACCAAAGCTCACG	IDT	N/A
Human PDX1 primer 2: GGAACTCCTTCT CCAGCTCTA	IDT	N/A
Neurog3-2A-eGFP PGK-Puro 3' arm genotyping Forward primer: ATCAGCAGCCTCTGTTCCAC	IDT	N/A
Neurog3-2A-eGFP PGK-Puro 3' arm genotyping Reverse primer: GGAGTCATCTTGCCAAGGCT	IDT	N/A
Neurog3-2A-eGFP PGK-Puro 5' arm genotyping Forward primer: GGTAGAAAGGTAATATTTGGAGGCCT	IDT	N/A
Neurog3-2A-eGFP PGK-Puro 5' arm genotyping Reverse primer: CTGAACTTGTGGCCGTTTACG	IDT	N/A
Software and Algorithms		
CellProfiler	Carpenter et al., 2006	http://cellprofiler.org

REAGENT or RESOURCE	SOURCE	IDENTIFIER
NoD	Scott et al., 2010	http://www.compbio.dundee.ac.uk/www-nod/
NucPred	Brameier et al., 2007	http://www.sbc.su.se/~maccallr/nucpred/cgi-bin/single.cg
SUMOsp	Xue et al., 2006	http://sumosp.biocuckoo.org/
WoLF PSORT	Horton et al., 2007	http://wolfpsort.org/

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for reagents may be direct to, and will be fulfilled by, the Lead Contact, Dr. Francis C. Lynn (francis.lynn@ubc.ca).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse Models—The UBC and UCSF Animal Care Committees approved animal experiments. Mice were housed on laboratory chow diet on a 12-hour light-dark cycle. Noon on the morning of the discovery of a vaginal plug was considered embryonic day (E)0.5. CD-1 mice were obtained from Charles River. Frozen embryos of Pdx1-Cre (Gu et al., 2002) mice were obtained from the MMRRC and were rederived by the BC Preclinical Research Consortium (Vancouver BC, Canada). Kras^{LSL-G12D} mice (Jackson et al., 2001) and C57BL/6(tetO-Cdkn1b)1Scpr/J mice (Pruitt et al., 2013) were purchased from Jackson Laboratory. Sox9-rtTA transgenic mice were made by recombining the reverse tetracycline transactivator (Clontech; pTet-On) into the Sox9 coding region of a 202kb BAC (RP23-36D5); transgenic mice were generated by the UCSF transgenic core as described (Shimajiri et al., 2011). Transgene expression was induced in pregnant Sox9-rtTA; tetO-Cdkn1b with daily 1 mg intraperitoneal (IP) injections of doxycycline on indicated gestational days.

Generation of NEUROG3-2A-eGFP and FUCCI CyT49 hESC Lines—To generate *NEUROG3-2A-eGFP* knock-in hESC line, the CRISPR/Cas9 system was used as previously described (Krentz et al., 2014). pCCC contains a full-length CAGGS promoter to replace the Cbh promoter of px458, improving expression in hESC CyT49 cells. The gRNA (GGGTCGCTCCTCCAGCGACG; score 0.73) was designed using the algorithm reported by Doench *et al* (Doench et al., 2014) and was cloned into the BbsI sites of pCCC to generate pCCC-LL502/3 as described by Ran *et al* (Ran et al., 2013). The targeting vector was based on Addgene #31938, contained ~800bp *NEUROG3* homology arms and is available upon request. Electroporation, selection, and picking of clones was performed as previously described using 40 µg of donor and 15 µg of CRISPR/Cas vectors (Krentz et al., 2014). Genomic DNA was extracted using QuickExtract.

To generate FUCCI transgenic CyT49, 10^6 cells were plated into 6-well plate and 4 µg of FUCCI construct was transfected using lipofectamine 3000. Cells were selected for integration using 0.25 ug/mL puromycin and clones were picked based on fluorescence expression. mKO-hCDt-2A-mAG-hGem-2A-Puro was cloned into a custom pCAGGS expression vector; further details and vector are available upon request.

Induced Pluripotent Stem Cell Generation—Human iPSCs were generated in house from male human PBMCs by infecting erythroid progenitors (Erythroid Progenitor Reprogramming Kit) with Sendai virus as outlined in the manufacturer's protocol (Cytotune 2.0). Cells were plated on Geltrex after 48 hours and subsequently transitioned to ReproTeSR from days 4–7 post-infection (StemCell Technologies). At 3 weeks postinfection, iPSC clones were picked into and maintained in mTeSR-E8 (StemCell Technologies) in Geltrex-coated 96-well plates. Clones were passaged using ReLeSR (Stem Cell Technologies) every 4–6 days until passage 15. Sendai virus transgene expression was then analysed, and found to be absent, using Taqman (Life Technologies) and pluripotency assessed by immunostaining and qPCR.

Human Pluripotent Stem Cell Differentiations-CyT49 or hIPSCs were plated onto Geltrex (1:100)-coated 12 well plates at a density of 0.5×10^6 in 10/10 media [DMEM/F12, 10% Xenofree-KOSR, Glutamax, P/S, 10 ng/mL Activin A and 10 ng/mL Herugulin-1β]. Differentiations began 48 hours post-seeding using a modified version of Rezania et al (Rezania et al., 2014). Briefly, cells were rinsed with $1 \times DPBS$ and then basal culture media (MCDB 131 medium, 1.5 g/L sodium bicarbonate, $1 \times \text{Glutamax}$, $1 \times \text{P/S}$) with 10 mM final glucose, 0.5% BSA, 100 ng/mL Activin A, and 3 µM of CHIR-99021 was added for 1 day only. For the following two days, cells were treated with the same media without CHIR-99021 compound to generate definitive endoderm (Stage 1). On day four, cells were cultured in basal media with 0.5% BSA, 10 mM glucose, 0.25 mM ascorbic acid and 50 ng/mL of KGF for 2 days to generate primitive gut tube (Stage 2). To produce posterior foregut (Stage 3), cells were treated for three days with basal media with 10 mM final glucose concentration, 2% BSA, 0.25 mM ascorbic acid, 50 ng/mL of KGF, 0.25 µM SANT-1, 1 µM retinoic acid, 100 nM LDN193189, 1:200 ITS-X, and 200 nM a-Amyloid Precursor Protein Modulator (APPM). For stage 4, cells were treated with basal media with 10 mM glucose, 2% BSA, 0.25 mM ascorbic acid, 2 ng/mL of KGF, 0.25 µM SANT-1, 0.1 µM retinoic acid, 200 nM LDN193189, 1:200 ITS-X, and 100 nM APPM for 3 days to generate pancreatic progenitors. Cells were maintained as planar cultures and media was changed to basal media with 20 mM glucose, 2% BSA, 0.25 µM SANT-1, 0.05 µM retinoic acid, 100 nM LDN193189, 1:200 ITS-X, 1 µM T3, 10 µM Repsox, and 10 µM zinc sulfate for 3 days to generate pancreatic endocrine precursors (Stage 5). Finally, cells were treated for two days in stage 6 media: basal media with 20 mM final glucose concentration, 2% BSA, 100 nM LDN193189, 1:200 ITS-X, 1 μM T3, 10 μM Repsox, 10 μM zinc sulfate, 100 nM gamma secretase inhibitor XX. CDKi treatment (2.5 µM CDK4/6 inhibitor PD-0332991, 1 µm CDK2 inhibitor ii and 1 µm CDK2 inhibitor iii) was carried out for 24 hours in Stage 5 or 6. For flow cytometric analysis, cells were treated with Accutase to generate single cells and fixed in 4% PFA for 15 min before analysis for GFP expression on BD Canto or for FUCCI expression on BD Fortessa. For FACS, cells were treated with Accutase to generate single cells and sorted based on GFP or FUCCI proteins directly into Trizol for downstream RNA analysis.

METHOD DETAILS

Analysis of EdU Cumulative Labeling—EdU cumulative labeling studies were carried out as previously described (Takahashi et al., 1993). At 9AM, pregnant dams were given 1

mg 5-Ethynyl-2'-deoxyuridine (EdU; IP), followed by subsequent 0.25 mg IP doses at 1.5 hr intervals. Embryos were collected 0.5 hours after the last injection. Pancreata were fixed in 4% paraformaldehyde, dehydrated and paraffin embedded as described (Xu et al., 2015). At least 8 sections spaced across the pancreas of 4–13 embryos from 2 dams were analyzed at each time point. In total, over 100 embryos from 62 pregnant dams were used to quantify over 50,000 pancreatic progenitors. Custom CellProfiler pipelines were used for image analyses (Carpenter et al., 2006). To determine the lengths of the G1-, S-, G2-M-phases of the cell cycle, the percent of EdU+ progenitor cells was plotted against the labeling time and the data was fit using least squares linear regression. To measure the length of G2-M (T_{G2-M}), the time for all EdU-labeled cells to exit S, transit through G2 and enter mitosis (pHH3+ [1:1000]) after a single pulse of EdU was determined using least squares linear regression.

Immunofluorescence (IF) and Flow Cytometry-For fluorescent staining and EdU detection of fixed tissue, paraffin slides were deparaffinized and rehydrated followed by antigen retrieval for 20 minutes in a 95° C pH 6 citrate buffer. Slides were washed in PBS and permeabilized with 0.5% Triton-X before EdU detection was performed by incubating slides for 1h with: 100 mM Tris pH 8.5, 1mM CuSO₄, 30 uM Alexa Fluor 594 azide triethylammonium, 100 mM ascorbate in H₂0 (Salic and Mitchison, 2008). After three washes in PBS, slides were blocked for 30 minutes each in blocking buffer (5% horse serum) and 1:50 anti-mouse IgG in blocking buffer. Primary antibodies were incubated at 4° overnight at the following dilutions: mouse anti-Pdx1 (1:100), goat anti-Cpa1 (1:250), mouse anti-Neurog3 (1:100), rabbit anti-P27 (1:500), rabbit anti-Sox9 (1:500), rabbit anti-Chga (1:400), mouse anti-Glucagon (1:2000), rabbit anti-Insulin (1:1000). Secondary antibodies from Jackson ImmunoResearch were incubated at room temperature for 1-2hours at the following dilutions: anti-488 (1:250) and anti-Cy3 (1:450). Slides were mounted using SlowFade Antifade Reagents (Thermo Fisher Scientific) and were imaged with a $20 \times /$ 0.75 objective using a Leica TCS SP8 confocal system. Custom CellProfiler pipelines were used to quantified immunopositive cells.

For quantitative flow analyses, cells were dissociated with 0.05% trypsin, fixed and stained as described previously (Van Hoof et al., 2011), or stained for 1 h at 37°C in medium containing Vibrant DyCycle Violet. Flow cytometry was conducted with an LSR-II flow cytometer (BD Biosciences); the data were analyzed with FACSDiva software (BD Biosciences). To determine the percent of cells in each cell cycle phase, cells in G0/G1 (one set of chromatids) were identified from G2-M (one set of sister chromatids and pHH3+) based on intensity of DNA stain using Hoescht 33342 (Miyatsuka et al., 2011). Cells in S phase were identified as the population of cells between the G0/G1 and G2-M peaks based on intensity.

Cloning and Mutagenesis—Cloning and mutagenesis of NEUROG3, E47 and FT-NEUROG3 were done as previously described (Van Hoof et al., 2009). The FLAG-tagged wt *NEUROG3* ORF was generated by amplifying human *NEUROG3* cDNA with PrimeStar DNA polymerase (Takara) using the forward primer 5'ctcgagccaccATGgattacaaggatgacgacgataagacgcctcaaccctcgggtgcgc (NEUROG3-F) to

introduce a *Xho*I restriction site (underlined), a Kozak sequence (italics; start codon in capitals), and the N-terminal flag tag (bold) in combination with the reverse primer 5'-<u>gcggccgcgTTA</u>cagaaaatctgagaaagccagact (NEUROG3-R) to introduce a *Not*I restriction site (underlined) and a stop codon (bold capitals). The amplified product was ligated into pCR2.1 TOPO (Life Technologies). This pCR2.1-flag-hNEUROG3 construct was used to generate the single and multiple point mutants by site-directed mutagenesis as described previously (Van Hoof et al., 2009). The flag-tagged *NEUROG3* ₆₋₇ ORF was generated using the forward primer 5'-

<u>ctcgag*ccacc*</u>**ATGgattacaaggatgacgacgataag**cagcagcaggagtcggcgaaaga to introduce a *Xho*I restriction site (underlined), a Kozak sequence (italics; start codon in capitals), and the N-terminal flag tag (bold) in combination with the NEUROG3-R reverse primer. The flag-tagged *NEUROG3* ₁₄₅₋₁₈₇ ORF was generated using the NEUROG3-F forward primer in combination with the reverse primer 5'-

<u>gcggccgcg</u>**TTA**cagaaaatctgagaaagccagactgcctgggctcaagcaggc

ggaaaaggtggcccccagcagcccgggtcgctcctccaggtggtccgctatgc gcagcgtttg to introduce a Not restriction site (underlined) and a stop codon (bold capitals). The flag-tagged NEUROG3 HILH ORF was generated using the forward primer NEUROG3-F in combination with the reverse primer 5'-aggttggatcctcgattgcgct cgcggtcgttggc to introduce a BamHI restriction site (underlined), and the forward primer 5'-tgactggatccctgcgcatagcggaccacagcttg to introduce a BamHI restriction site (underlined) in combination with the NEUROG3-R reverse primer. These N-terminal and C-terminal amplification products were digested with *XhoI* in combination with *Bam*HI, and with *Bam*HI in combination with *NofI*, respectively, after which they were ligated into the pCR2.1-flag-hNEUROG3, the flag-hNEUROG3 ORF of which was removed using XhoI and NofI. The glycine and serine codons that were introduced as a result of the *Bam*HI restriction sequence were mutated to alanines codons by site-directed mutagenesis as described previously(Van Hoof et al., 2009). The 6 × Histagged *E47*ORF was generated by amplifying Syrian hamster *Pan1* cDNA (German et al., 1991) with PrimeStar DNA polymerase (Takara) using the forward primer 5'gtggcgctcgaggccaccATGgctctttctacgtggcagatggcagcct to introduce a XhoI restriction site (underlined) and a Kozak sequence (italics; start codon in capitals) in combination with the introduce a *Not* restriction site (underlined), a C-terminal $6 \times$ His tag (bold) and a stop codon (capitals). All constructs were sequenced prior to subcloning into the CMV promoterdriven mammalian expression vector pBAT12 (Smith et al., 1999) using XhoI and NotI.

Cell Culture, Transfection, Lysis and Phosphatase Treatment—HeLa cells and mPAC cells were grown in DMEM with 10% fetal bovine serum and penicillin/streptomycin (P/S) and transfected with Lipofectamine 2000. As indicated, cells were incubated with 50 ng/mL nocodazole, 1.0 nM leptomycin B, 10 μ M MG132, 50 mM roscovitine, 6.0 μ M BIX02189, 10 μ M LY2228820, 1.0 μ M SP600125, 10 μ M SB216763 for 24 hours before collection. Cells were washed and either fixed with 2.0% paraformaldehyde or collected in ice-cold PBS with protease inhibitors prior to lysis by sonication. Lysates were cleared by centrifugation and protein concentration determined by BCA assay (Van Hoof et al., 2006). λ -phosphatase treatment was carried out using the manufacturer's instructions.

SDS-PAGE and Western Blotting—Embryonic pancreata pooled from five E15.5 CD1 embryos were lysed in NRSB with protease inhibitors and analysed by western blot as described (Speckmann et al., 2016). 6.5–50 μg of HeLa cell protein lysate was separated by SDS-PAGE and blotted as described previously (Lynn et al., 2007). Antibodies included: 1:1,000 mouse-anti-FLAG; 1:1,000 rabbit-anti-β-Actin; 1:15,000 donkey-anti-mouse IRDye 800CW and 1:15,000 donkey-anti-rabbit IRDye 680RD (Li-Cor); 1:500 guinea-pig-anti-NEUROG3; 1:1000 sheep-anti-NEUROG3. Immunoreactivity was imaged using near-infrared immunofluorescent detection (Li-Cor Odyssey) or by chemiluminescence and film (Speckmann et al., 2016).

Functional Domain Prediction Software—Putative NLSs in WT NEUROG3 (NCBI accession number NP_066279) were identified with WoLF PSORT http://wolfpsort.org/ (Horton et al., 2007) and NucPred http://www.sbc.su.se/maccallr/nucpred/cgi-bin/single.cgi (Brameier et al., 2007), nucleolar localization signals with NoD http:// www.compbio.dundee.ac.uk/www-nod/ (Scott et al., 2010), and ubiquitination/ SUMOylation sites with SUMOsp http://sumosp.biocuckoo.org/ (Xue et al., 2006).

Luciferase Assay—Luciferase assays (Promega) were performed according to the manufacturer's instructions. Transcription factor-containing expression vectors were transfected in a 1:1:1 ratio with the previously described Hnf1a and Luciferase expression constructs (Smith et al., 2003).

Mouse Embryonic Pancreas Explants Treatment and Quantification—The dorsal pancreas of CD1 E11.5 embryos was dissected as previously described (Petzold and Spagnoli, 2012) and transferred to Millicell EZ slides (EMD Millipore) covered in Geltrex diluted 1:1 in DMEM/F12. Explants were allowed to recover and attach overnight in culture medium: DMEM/F12, 10% FBS, Glutamax, P/S, and insulin-transferrin-selenium. The following morning, media was topped up with CDKi for 24 hours: 2.5 µM CDK4/6 inhibitor PD-0332991, 1 µm CDK2 inhibitor ii and 1 µm CDK2 inhibitor iii. Explants were then washed with PBS and fixed in 4% PFA for 30 minutes. After removal of PFA, explants were washed three times with PBS and blocking solution (PBS +0.1% Triton-X and 3% horse serum) was added for a minimum of 1 hr. Primary antibodies (mouse anti-Pdx1 [1:100] and rabbit anti-Sox9 [1:500]) in blocking solution was added to explants and incubated overnight at 4°. The following morning, after three PBS washes explants were incubated with secondary antibodies in blocking solution for 1 hr: anti-mouse 488 (1:250), anti-rabbit 594 (1:450) and nuclei dye To-Pro-3 iodide (1:10,000). Explants were imaged on Leica SP8 confocal using z-stacks to image whole explant and images were quantified using custom CellProfiler pipelines.

qPCR—Cells were lysed in Trizol and standard phenol-chloroform extraction was used to isolate RNA. Following RNA extraction, Superscript III was used for reverse transcription followed by Taqman on ViiA7. Primer sequences are included in STAR Key Resources Table.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analyses were performed using Prism 5 (GraphPad Software). Segmental linear regression using 1000 iterations was used to mathematically model EdU cumulative labelling. Statistically significant differences were assessed using ANOVA followed by either a Dunnett or Tukey post hoc test, t-test or non-parametric tests as appropriate, with p value of < 0.05 deemed significant. Errors bars represent standard error of the mean.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

• The progenitor cell cycle lengthens during embryonic pancreas development

- Cell-cycle lengthening is essential for induction of NEUROG3
- In trunk progenitors, NEUROG3 is phosphorylated by G1-S CDKs and then degraded
- G1 lengthening drives endocrine differentiation by slowing NEUROG3 phosphorylation

Page 27



Figure 1. The Length of the G_1 Phase of the Cell Cycle Increases during Mouse Pancreatic Development

(A) Schematic describing experimental approach used to measure the lengths of the G_1 (T_{G1}), S (T_S), and G_2 -M (T_{G2-M}) cell-cycle phases with cumulative EdU labeling of cells in S phase (light blue) and the M-phase marker pHH3 (gray). The lengths of the G_1 , S, and G_2 -M phases of the cell cycle were determined from measurements of $T_C - T_S$ (green), T_S (blue), and T_{G2-M} (red).

(B) Cumulative EdU labeling of Pdx1⁺ cells at E11.5. Green arrow ($T_C - T_S$) indicates the length of time it takes to reach plateau (dotted black line = growth fraction [GF]). y intercept represents (T_S/T_C) * GF. Embryos (n = 4–13) from two dams were analyzed at each time point.

(C) Cumulative EdU labeling of Pdx1⁺ cells at E12.5. n = 4-13 embryos from two dams at each time point.

(D) Cumulative EdU labeling of Pdx1⁺ cells at E13.5. n = 4-13 embryos from two dams at each time point.

(E) The proportion of pHH3⁺ cells labeled with EdU after 1.0, 1.5, 2.0, and 2.5 hr at E11.5 (black), E12.5 (blue), and E13.5 (green). Red arrow indicates the length of G_2 -M (T_{G2-M}). n = 3 embryos from two dams at each time point.

(F–H) Cumulative EdU labeling of Pdx1⁺Cpa1⁺ tip (blue) and Pdx1⁺Cpa1⁻ trunk

progenitors (red) at (F) E11.5, (G) E12.5, and (H) E13.5. Arrows indicate time to reach GF. n = 4-13 embryos from two dams at each time point.

(I) The length in hours of the G_1 (black), S (gray), and G_2 -M (white) phases of the cell cycle at E11.5, E12.5, and E13.5 in the Pdx1⁺ progenitor population.

(J) Calculated cell-cycle parameters for tip and trunk progenitors at E11.5, E12.5, and E13.5: G_1 (red or blue), S (gray), and G_2 -M (white).

Data are presented as mean \pm SEM. See also Figure S1 and Table 1.



Figure 2. Ectopic Expression of Cdkn1b (P27) in Sox9⁺ Progenitors Causes Endocrine Cell Differentiation

(A) One injection of doxycycline (Dox) at E12.3 increased Cdkn1b (green) expression in the pancreatic epithelium (Pdx1⁺; red) of double transgenic (DT) but not single transgenic (ST) mice at E12.7.

(B) 9.5 hr of Cdkn1b induction at E12.5 significantly increased the number of Neurog3⁺ cells in DT embryos. n = 3. ***p < 0.0001 by unpaired t test.

(C) Ectopic expression of Cdkn1b (9.5 hr) did not alter the total number of pancreatic cells. n = 6.

(D) At E12.5, Cdkn1b-expressing (red) pancreatic epithelial cells (Pdx1⁺; blue) from Doxtreated (E10.5–12.5) DT embryos incorporated less EdU (green) at 3.5 hr.

(E) Significantly fewer $Pdx1^+EdU^+$ cells were counted following 3.5 hr of labeling of DT embryos. n = 7. *p < 0.05 by unpaired t test.

(F) Ectopic Cdkn1b expression did not alter the fraction of EdU⁺ progenitor cells (the GF) after 11 hr of labeling, indicating that cells remained in the cell cycle. n = 3.

(G) Chga⁺ cells were significantly increased at E12.5 DT embryos following 48 hr of Dox. n = 5. **p < 0.01 by Mann-Whitney U test.

(H) Gcg^+ cells were significantly increased at E12.5 in DT embryos following 48 hr of Dox. n = 6. **p < 0.01 by unpaired t test.

(I) Ectopic expression of Cdkn1b (E10.5–12.5) did not reduce total pancreatic cells at E12.5. n = 7.

(J) Dox treatment (E12.5–14.5) of DT embryos increased Cdkn1b (red) expression in subset of $Pdx1^+$ (blue) trunk cells.

(K) Dox-treated (E12.5–14.5) DT embryos showed significantly reduced EdU⁺ labeling after 3.5 hr at E14.5. n = 4. ***p < 0.0001 by unpaired t test.

(L) More Chga⁺ cells were present in Dox-treated (E12.5–14.5) DT embryos at E14.5. n = 5. **p < 0.001 by Mann-Whitney U test.

(M) More Gcg⁺ cells were present in Dox-treated (E12.5–14.5) DT embryos at E14.5. n = 5. **p < 0.01 by unpaired t test.

(N) Ins^+ cells were not significantly changed in Dox-treated (E12.5–14.5) DT embryos. n = 9. p = 0.09 by unpaired t test.

(O) The number of pancreatic cells per slide was unchanged, suggesting that growth was not affected by 48 hr of Cdkn1b expression at E14.5. n = 5.

Data are presented as mean \pm SEM. Scale bars, 50 µm. See also Figure S2.

Krentz et al.



Figure 3. Wild-Type Mouse and Human Neurog3 Can Be Hyperphosphorylated

(A) Western blot from E15.5 mouse pancreas reveals multiple Neurog3 bands.

(B) Schematic of FLAG-tagged (f) human NEUROG3 mutants: wild-type (WT), T \rightarrow A mutation predicted to prevent dimerization with E47 (T120A), basic (b), helix-loop-helix (HLH), activation (act) domains. Anti-FLAG western blot of lysates from HeLa cells expressing indicated mutants with (+) or without (-) phosphatase treatment.

(C) Anti-FLAG western blots of HeLa cell lysates expressing WT-Neurog3 (lane 1) and mutants that had one of the nine serine residues in the 138–190 region mutated to alanine as indicated.

(D) Anti-FLAG western blots of HeLa cell lysates expressing NEUROG3 variants with all nine serine residues in the 138–190 region mutated to alanine (9A) and variants with eight (8A) serine residues mutated to alanine, with indicated residue(s) left untouched.

(E) Anti-FLAG western blots of HeLa cell lysates expressing NEUROG3 variants with all nine serine residues in the 138–190 region mutated to alanine (9A) and variants with seven (7A) or eight (8A) serine residues mutated to alanine, with indicated residue(s) left untouched.

Asterisks, arrows, arrowheads, and double arrows represent the first, second, third, and fourth band, respectively. See also Figure S4.



Figure 4. Hyperphosphorylation Reduces the Transcriptional Activity of NEUROG3 (A) *PAX4* minienhancer-driven luciferase transcription is reduced by NEUROG3 phosphorylation. HeLa cells were cotransfected with the mini-enhancer, an Hnf1a overexpression construct, and FLAG-tag control (FLAG) or WT or mutant forms of NEUROG3.

(B) mPAC cells were transfected with FLAG-tag control, WT-NEUROG3, or the indicated mutants and analyzed for expression of endogenous *Pax4* by qPCR.

(C) mPAC cells were transfected with FLAG-tag control, WT-NEUROG3, or the indicated mutants and analyzed for expression of endogenous *NeuroD1* by qPCR.

(D) mPAC cells were transfected with FLAG-tag control, WT-NEUROG3, or the indicated mutants and analyzed for expression of endogenous *Somatostatin* (*Sst*) by qPCR.

Data are presented as mean \pm SEM. n = 3. *p < 0.05 by one-way ANOVA and Dunnett's post hoc test. See also Figures S3 and S4.



Figure 5. Phosphorylation of S183 Occurs during $\rm G_2\text{-}M$ Phase and Is Not Required for $\rm G_1\text{-}S$ Cell-Cycle Arrest

(A) Untransfected HeLa cells and those expressing FLAG-tagged NEUROG3 variants were trypsinized, fixed, and stained with anti-FLAG, anti-pHH3 antibodies, and DAPI, and flow cytometry utilized to determine the number of transfected cells in G_0 - G_1 , S, and G_2 -M. ** or $\dagger p < 0.005$ compared with untransfected by one-way ANOVA and Dunnett's post hoc test. (B) Live HeLa cells expressing FLAG-tagged WT NEUROG3 (red) or NEUROG3 _{HLH} (blue) were trypsinized and sorted by FACS into G_0 - G_1 , S, and G_2 -M populations. For each population, the fast-, medium-, and slow-migrating NEUROG3 species (i.e., unphosphorylated, intermediately phosphorylated, and maximally phosphorylated forms, respectively) were quantified by western blotting with anti-FLAG antibody and normalized to β -actin levels. Relative protein levels within each cell-cycle phase that are significantly different from others are indicated by asterisks above that phosphorylation form; *p < 0.05, **p < 0.005 by one-way ANOVA. n = 3.

(C–E) HeLa cells expressing the indicated fluorescence timer (pTimer/FT)-fusion proteins were untreated (C), incubated with leptomycin B or MG132 (D), or cotransfected with E47 (E), prior to flow cytometric analyses to determine the percentage of FT red-containing cells. Significant differences from FT only (C and E) or vehicle (D) are indicated by *p < 0.05, **p < 0.005, and ***p < 0.0005 by one-way ANOVA and Dunnett's post hoc test. n = 3.

Data are presented as mean \pm SEM.

Krentz et al.



Figure 6. NEUROG3 Is Phosphorylated by G₁-S Cyclin-Dependent Kinases

(A) Anti-FLAG western blot of HeLa cell lysates expressing indicated NEUROG3 variants were incubated with nocodazole alone or in combination with BIX02189, LY2228820, SP600125, or SB216763.

(B) Anti-FLAG western blot on HeLa cell lysates expressing FLAG-tagged WT-NEUROG3 were either untreated, incubated with nocodazole alone, or incubated with both nocodazole and roscovitine.

(C) Mouse pancreatic explants were cultured for 24 hr starting at E11.5 with Cdk2 and 4/6 inhibitors (CDKi) and the number of Neurog3⁺ cells/Sox9⁺ cells and the total Sox9⁺ cells were quantified by immunofluorescence staining. n = 5. **p < 0.01 by unpaired t test.

(D) Stage-6 endocrine progenitor cells derived from a human iPSCs were treated for 24 hr with CDKi and NEUROG3 protein was quantified by western blot. n = 3 wells. **p < 0.01 by unpaired t test.

(E) GFP⁺ cells were quantified using a *NEUROG3-2A-eGFP* hESCs (N5-5) differentiated to stage 6 prior to 24-hr CDKi treatment. n = 9 wells from two differentiations. *p < 0.05 by Mann-Whitney U test.

(F) *NEUROG3* expression in FACS purified GFP⁺ N5-5 cells following CDKi treatment at stage 6. n = 3 wells from two differentiations.

(G) CDKi treatment increased expression of *NEUROD1* in FACS purified GFP⁺ cells. n = 3 wells from two differentiations. *p < 0.05 by one-way ANOVA and Tukey post hoc test. (H) CDKi treatment increased expression of *PDX1* in FACS purified GFP⁺ cells. n = 3 wells from two differentiations. *p < 0.05, **p < 0.01, ***p < 0.001 by one-way ANOVA and Tukey post hoc test.

(I) CDKi treatment of stage-6 endocrine progenitors, derived from the FUCCI-3 hESC line, increased the proportion of cells in G₁ (red) and decreased the proportion of cells in S-G₂-M (green). n = 5 wells from two differentiations. ****p < 0.0001 by one-way ANOVA and Tukey post hoc test.

(J) CDKi treatment of stage-6 endocrine progenitors derived from FUCCI-3 hESC line significantly increased expression of *NEUROG3* in G₁ (red) cells. n = 4 wells from two differentiations. *p < 0.05, **p < 0.01 by one-way ANOVA and Tukey post hoc test. (K) Expression of *NEUROD1* was upregulated in G₁ (red) cells compared with S-G₂-M (green). n = 5 wells from two differentiations.

(L) CDKi treatment of stage-6 endocrine progenitors derived from FUCCI-3 hESC line significantly increased expression of *PDX1* in G₁ (red) cells. n = 5 wells from two differentiations. **p < 0.01 by one-way ANOVA and Tukey post hoc test. Data are presented as mean \pm SEM. See also Figure S5.

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Table 1

Cell-Cycle Phase Lengths of Pdx1⁺ Pancreatic Epithelium, Pdx1⁺Cpa1⁺ Tip, and Pdx1⁺Cpa1⁻ Trunk Progenitor Cells during Mouse Embryonic Development

		$\mathbf{T}_{\mathbf{C}}$	T_{G1}	T_S	T_{G2-M}	GF
$Pdx1^+$	E11.5	8.20 ± 0.68	4.54 ± 0.40	1.70 ± 0.38	1.96 ± 0.10	0.75 ± 0.02
	E12.5	9.73 ± 0.86	5.91 ± 0.51	1.75 ± 0.46	2.07 ± 0.07	0.73 ± 0.01
	E13.5	11.79 ± 0.93	7.19 ± 0.48	2.53 ± 0.48	2.07 ± 0.05	0.78 ± 0.01
Pdx1+Cpa1+	E11.5	8.41 ± 0.74	3.74 ± 0.35	2.71 ± 0.49	1.96 ± 0.10	0.94 ± 0.02
	E12.5	10.54 ± 0.66	5.29 ± 0.34	3.18 ± 0.41	2.07 ± 0.07	0.94 ± 0.02
	E13.5	11.57 ± 0.66	5.62 ± 0.34	3.89 ± 0.40	2.07 ± 0.05	0.95 ± 0.02
Pdx1+Cpa1-	E11.5	7.57 ± 0.74	4.41 ± 0.43	1.20 ± 0.41	1.96 ± 0.10	0.73 ± 0.03
	E12.5	9.49 ± 0.67	6.24 ± 0.44	1.17 ± 0.37	2.07 ± 0.07	0.70 ± 0.02
	E13.5	11.47 ± 0.95	7.36 ± 0.64	2.04 ± 0.47	2.07 ± 0.05	0.74 ± 0.01
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 $Mean \ values \pm SEM \ of \ total \ cell-cycle \ length \ (TC), \ G1 \ phase \ length \ (TG1), \ S \ phase \ length \ (TS), \ and \ G2-M \ phase \ length \ (TG2-M) \ red \ ($