The homeobox factor PDX-1 is a key regulator of pancreatic morphogenesis and glucose homeostasis; targeted disruption of the PDX-1 gene leads to pancreatic agenesis in \( \text{pdx-1}^{-/-} \) homozygotes. PDX-1 heterozygotes develop normally, but they display glucose intolerance in adulthood. Like certain other homeobox proteins, PDX-1 contains a consensus FPWMK motif that promotes heterodimer formation with the ubiquitous homeodomain protein PBX. To evaluate the importance of PDX-1:PBX complexes in pancreatic morphogenesis and glucose homeostasis, we expressed either wild-type or PBX interaction defective PDX-1 transgenes under control of the PDX-1 promoter. Both wild-type and mutant PDX-1 transgenes corrected glucose intolerance in \( \text{pdx-1}^{-/-} \) heterozygotes. The wild-type PDX-1 transgene rescued the development of all pancreatic lineages in \( \text{pdx-1}^{-/-} \) animals, and these mice survived to adulthood. In contrast, pancreata from \( \text{pdx-1}^{-/-} \) mice expressing the mutant PDX-1 transgene were hypoplastic, and these mice died within 3 weeks of birth from pancreatic insufficiency. All pancreatic cell types were observed in \( \text{pdx-1}^{-/-} \) mice expressing the mutant PDX-1 transgene; but the islets were smaller, and increased numbers of islet hormone-positive cells were noted within the ductal epithelium. These results indicate that PDX-1:PBX complexes are dispensable for glucose homeostasis and for differentiation of stem cells into ductal, endocrine, and acinar lineages; but they are essential for expansion of these populations during development.

The homeobox protein PDX-1 functions critically in pancreatic development and glucose homeostasis; murine and human \( \text{pdx-1}^{-/-} \) homozygotes are apancreatic (1–3), and \( \text{pdx-1} \) heterozygotes develop glucose intolerance in adulthood, partly because of inadequate insulin gene expression (4, 5). Further underscoring the importance of this factor for glucose homeostasis, mutations in the PDX-1 gene have been identified in patients with maturity onset diabetes of the young-type 4 (6). Consistent with this phenotype, PDX-1 stimulates the expression of certain islet-specific genes such as somatostatin (7, 8), insulin (9–11), gluc-2 (12), and glucokinase (13), by binding as a monomer to promoter sites that contain a GATAAT consensus site.

Like certain homeobox factors, PDX-1 also binds as a heterodimer with the ubiquitous homeodomain protein PBX to target sites containing a consensus TGATTAAT motif. Detected in crude extracts from insulinoma cells (14), PDX-1:PBX complexes have also been observed in ductal and acinar cell lines, where they regulate the expression of exocrine target genes such as elastase (15). Compared with the PDX-1 monomer, the PDX-1:PBX heterodimer binds with 10- to 20-fold higher affinity to its sites (14), suggesting a potential mechanism by which PDX-1 target genes are hierarchically regulated during development.

To examine the requirement for PDX-1:PBX complexes in pancreatic morphogenesis, islet cell differentiation, and glucose homeostasis, we developed transgenic mice expressing either wild-type or PBX interaction defective PDX-1 transgenes under control of the PDX-1 promoter/enhancer. In the context of a \( \text{pdx-1}^{-/-} \) background, the wild-type PDX-1 transgene completely rescued pancreatic development and glucose homeostasis in adult mice. The mutant PDX-1 transgene also corrected glucose intolerance in \( \text{pdx-1}^{+/-} \) mice, but it was unable to fully rescue pancreatic development in \( \text{pdx-1}^{-/-} \) animals. All cell types within the pancreas of \( \text{pdx-1}^{-/-} \) mice expressing the mutant PDX-1 transgene were present; but proliferation and overall proliferation were greatly affected. These results show that PDX-1:PBX complexes are dispensable for the generation of the various cell types within the pancreas, but they are required for expansion of these cell populations during development. Taken together, these studies demonstrate that PDX monomers and PDX:PBX heterodimers perform distinct functions in the developing pancreas, reflecting in part their capacity to regulate discrete subsets of pancreatic target genes.

**Materials and Methods**

**Plasmids and Transgenic Mice.** Wild-type and mutant PDX-1 transgenes were constructed with the use of a 15-kb rat PDX-1 genomic clone (16) that, in addition to the 1-kb coding region, contains 6.5 kb of 5′ flanking sequence, a 4-kb intron, and 3 kb of 3′ flanking sequence. The PBX interaction defective PDX-1 clone contained mutations in sequences encoding aa 119–123 of the rat PDX-1 gene (FPWMK/AAGGO), which were generated by PCR-based mutagenesis (14). Plasmids were injected into C57BL6 mouse oocytes. Three independent lines of mice expressing the wild-type PDX-1 transgene and two independent lines expressing the mutant PDX-1 transgene were generated. Glucose tolerance tests were performed as previously described (4).

**RNase Protection and Western Blot Assays.** RNase protection assays were performed as previously reported (16), with a 450-base antisense PDX-1 riboprobe, extending from +130 to +530 of the rat PDX-1 cDNA and containing 50 nucleotides of plasmid sequence. Western blot assays were performed with the use of PDX-1-specific antisemir as reported (17).

**Immunolabeling.** Double immunofluorescence labeling was performed sequentially with primary antibodies made in different species: guinea pig anti-human insulin (Linco Research Immunodiagnostic, St. Charles, MO), rabbit anti-bovine glucagon (kindly donated by M. Appel, University of Massachusetts Medical School, Worcester, MA), and rabbit anti-synthetic somatostatin, a mixture of the three non-β cell hormone antibodies (antiglucagon, -somatostatin and -pancreatic polypeptide). Antibodies were used on paraffin sections of 4% buffered formaldehyde-
fixed pancreas, with the exception of PBX-1 antiserum (Santa Cruz Biotechnology), which was used on freshly frozen sections of 3-day-old wild-type mouse pancreas. The secondary antibodies used for immunofluorescence were Texas Red-conjugated donkey anti-guinea pig IgG, FITC-conjugated donkey anti-rabbit IgG, and streptavidin-conjugated FITC (all 1:100 dilution; Jackson ImmunoResearch). After extensive rinsing, slides were mounted with DABCO glycerol anti-fading mounting media. Immunoperoxidase labeling was carried out with the Histomouse kit (Zymed) for insulin (1:1000) or the Vectastain ABC kit (Vector Laboratories) for glucagon (1:1000) and somatostatin (1:1000). Detection of β-galactosidase from the pdx-1lacZKO allele was performed as previously described.

**Results**

Mutagenesis of the PBX interaction motif in PDX-1 (aa 119–123; FPWMK/AAGGO) has been shown to block complex formation with PBX on a heterodimer recognition site (14). Disruption of this motif in HoxA9 does not eliminate cooperativity with PBX (18), however, prompting us to evaluate functional interactions between a transcriptionally activated PBX (E2A-PBX) and wild-type or PBX interaction defective PDX proteins. Consistent with the selectivity of heterodimer binding sites for PBX:HOX complexes, wild-type and mutant PDX-1 proteins alone had marginal activity on a TSEII reporter that contains two PDX:PBX heterodimer recognition sites (Fig. 1A). After cotransfection with E2A-PBX, the wild-type PDX-1 construct induced reporter activity by 4- to 5-fold (Fig. 1A). In contrast, E2A-PBX had no effect on reporter activity in cells cotransfected with the mutant PDX-1 construct, demonstrating that disruption of the PBX interaction motif in PDX-1 is indeed sufficient to block functional cooperativity with PBX.

To evaluate the role of PDX-1:PBX complexes in promoting pancreatic morphogenesis and glucose homeostasis, we prepared transgenic mice expressing wild-type or PBX interaction defective PDX-1 proteins under the control of the PDX-1 promoter/enhancer (Fig. 1B). In previous experiments using lacZ reporter transgenes, both rat and mouse PDX-1 gene promoter/enhancers have been found to recapitulate the spatiotemporal expression pattern of the endogenous PDX-1 gene in transgenic mice (16, 19).

When fused to the rat PDX-1 promoter/enhancer, mutant and wild-type PDX-1 transgenes were comparably expressed in pancreatic tissue from founder animals by RNase protection and Western blot assays (Fig. 1C). The size and appearance of the pancreas in animals carrying mutant (mTg) or wild-type (wtTg) transgene mice were indistinguishable from nontransgenic littermates (not shown), indicating that neither transgene interferes with normal pancreatic development in this context.

To determine whether PDX-1:PBX complexes are required for glucose homeostasis, we crossed mTg and wtTg mice with pdx-1(+/−) mice (1). PDX-1 protein levels in pancreatic ex-
Fig. 2. Islet hypoplasia in pdx-1(−/−);mTg neonates. (A) Comparison of pdx-1(−/−);mTg (smaller) and wild-type (larger) littermates 16 days after birth. (B) Whole mount of pdx-1(−/−); wtTg and pdx-1(−/−); mTg mice. Ventral (Left) and dorsal (Right) views are included. p, pancreas; d, duodenum; sp, spleen; s, stomach. (C) Immunocytochemical analysis of insulin (red, arrows) and glucagon + somatostatin (green, asterisks) cells in sections from pdx-1(−/−); wtTg (Upper) and pdx-1(−/−); mTg (Lower). (D) Whole mount showing smaller developing islets (dark blue; arrowheads) in pdx-1(−/−) mTg compared with pdx-1(+/-) mTg mice. Blue color indicates expression of β-galactosidase from the endogenous pdx-1 allele pdx-1lacZKO (1).
tracts from *pdx-1*(+/-) mice were about half that of wild-type animals by Western blot analysis; and expression of either wild-type or mutant transgenes restored PDX-1 protein to wild-type levels in *pdx-1* heterozygotes (Fig. 1D). After i.p. glucose injection, blood glucose levels in wild-type mice rose to 175 mg/dl within 15 min, returning to baseline after 2 h. As expected, blood glucose levels in *pdx-1*(+/-) mice rose to 350 mg/dl after 20 min and remained abnormally elevated for the duration of the assay (ref. 4; Fig. 1D). Consistent with the notion that monomeric PDX-1 is sufficient for the activation of genes involved in glucose homeostasis, expression of either wild-type or mutant PDX-1 transgenes corrected the impairment in glucose homeostasis in *pdx-1* heterozygotes (Fig. 1D). These results suggest that PDX-1:PBX complexes are dispensable for islet cell function in the adult.

To determine the importance of PDX-1:PBX complexes in pancreatic development, we analyzed the activities of wild-type and mutant PDX-1 transgenes in the context of *pdx-1*(-/-) homozygous mice. *pdx-1*(-/-) mice were apancreatic and died 2–3 days after birth (1, 2); but *pdx-1*(-/-);wtTg mice developed normally and survived to adulthood. Mice expressing the mutant PDX-1 transgene (*pdx-1*(-/-);mTg) were comparable in size to nontransgenic mice at birth, but they were noticeably smaller after 2 weeks and failed to survive past 3 weeks (Fig. 2A). *pdx-1*(-/-);wtTg mice had a normal pancreas with a normal proportion of islet and exocrine cells (Fig. 2B and C). Compared with *pdx-1*(-/-);wtTg mice, the overall size of the pancreas in *pdx-1*(-/-);mTg mice at birth was 30–40% of normal, and the islets were markedly smaller (Fig. 2B–D).

Each of the cell types evaluated immunocytochemically (insulin, somatostatin, and glucagon) was present in...
pdx-1(-/-);mTg mice; but the distribution of these cells within individual islets was abnormal (Fig. 3). In particular, glucagon- and somatostatin-expressing cells were located throughout the small endocrine clusters and not exclusively at the periphery, as in wild-type mice (Fig. 3, compare C, D, G, and H).

The reduced size of the pancreas in pdx-1(-/-);mTg mice prompted us to attempt to determine whether the proliferation of pancreatic cells was correspondingly lower in these animals. By using the marker K-67 (20) as a reliable index of ongoing cellular proliferation, we observed intense staining in about 20–30% of pancreatic cells from wild-type newborn animals (Fig. 4A, Left). Consistent with their relative abundance in the neonatal pancreas, acinar cells accounted for the highest percentage of proliferating cells, followed by ductal and islet cells. K-67 staining was severely reduced in pdx-1(-/-);mTg mice; both the number and intensity of immunopositive cells were markedly lower, and all cell types within the pancreas were equally affected (Fig. 4A, Right). These results suggest that PDX-1:PBX complexes promote the expansion of various lineages within the pancreas by enhancing cellular proliferation.

To identify particular cell types where PDX-1:PBX complexes are likely to form, we conducted immunocytochemical studies with PBX-1 antiserum. PBX-1 was localized in the nucleus in all cells of the neonatal pancreas; the highest levels of PBX-1 were detected in ductal cells, with modest to low staining in islets and acinar cells, respectively (Fig. 4B). These results support the notion that PDX-1 interacts with PBX primarily in ductal epithelial cells, the site of presumed stem cells and the origin of islet endocrine cells. Remarkably, pancreatic ducts of pdx-1(-/-);mTg neonates contained numerous hormone-positive cells (Fig. 4C), whereas wild-type neonates did not (not shown). Indeed, all major hormone (insulin, glucagon, somatostatin) cell types were detected, suggesting a general defect in islet cell migration in these mice (Fig. 4C). Taken together, these results indicate that the interaction of PDX-1 with PBX is critical for early events in pancreatic development, including migration of differentiating endocrine cells or precursors out of the duct to form proper islets and proliferation of pancreatic cell types.

Discussion

PDX-1 has been shown to stimulate pancreatic gene expression in vitro, in part, as a heterodimer with PBX (14, 15), although the functional importance of this complex relative to the PDX-1 monomer has not been appreciated. Our results demonstrate a differential requirement in vivo for PDX-1:PBX heterodimers with respect to the initial specification of exocrine/endocrine cell types and the formation of the normal architecture of the mature organ.

In the absence of PDX-1, pancreatic development is arrested at the early bud stage, with complete absence of the major cell types (proper endocrine, exocrine, or ductal cells; refs. 9–11). Each of these pancreatic cell types is present, however, in pdx-1(-/-);mTg neonates, suggesting that monomeric (non-PBX-associated) PDX-1 activity is sufficient to promote initiation of a complete pancreatic genetic program. PDX-1:PBX complexes were also
dispensable for glucose homeostasis, inasmuch as overexpression of the mutant PDX-1 transgene rescued glucose intolerance in pdx-1 heterozygotes. Rather, the PBX interaction motif in PDX-1, and therefore the PDX-1:PBX complex, was essential for the expansion of each cell type within the developing pancreas. Pancreata from pdx-1 (−/−) mTg mice were markedly smaller than those of control littermates, owing in large part to a reduction in cellular proliferation among all endocrine/exocrine compartments.

Our immunocytochemical analyses indicate that PDX-1:PBX complexes are likely to form most abundantly in ducts of the developing pancreas, the sites of presumed islet cell precursors. Although cellular levels of the PDX-1:PBX heterodimer may be regulated in part by nuclear translocation of PBX via Meis proteins during development (21), PBX-1 protein was exclusively nuclear, at least in cells of the neonatal pancreas. Rather, PDX-1:PBX complexes appear to be limited primarily by the nuclear, at least in cells of the neonatal pancreas. Rather, PDX-1:PBX complexes are likely to form most abundantly in ducts of the developing pancreas, the sites of presumed islet cell precursors. During development, PDX-1 protein is, except for a few cells (22), undetectable in ducts of the mature pancreas (23). Remarkably, ductal expression of PDX-1 in adult cells is induced (22), undetectable in ducts of the mature pancreas (23). Alternatively, PDX-1:PBX may induce the expression of certain genes that are required for the subsequent migration of hormone positive cells from ducts. In either case, failure of these cells to migrate from the duct may contribute to the abnormal islet morphology in pdx-1 (−/−); mTg mice. In this regard, it will be of interest to identify PDX-1:PBX target genes that function in this developmental process.

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