

Enhancer, transcriptional, and cell fate plasticity precedes intestinal determination during endoderm development

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After acquiring competence for selected cell fates, embryonic primordia may remain plastic for variable periods before tissue identity is irrevocably determined (commitment). We investigated the chromatin basis for these developmental milestones in mouse endoderm, a tissue with recognizable rostro-caudal patterning and transcription factor (TF)-dependent interim plasticity. Foregut-specific enhancers are as accessible and active in early midgut as in foregut endoderm, and intestinal enhancers and identity are established only after ectopic *cis*-regulatory elements are decommissioned. Depletion of the intestinal TF CDX2 before this *cis* element transition stabilizes foregut enhancers, reinforces ectopic transcriptional programs, and hence imposes foregut identities on the midgut. Later in development, as the window of chromatin plasticity elapses, CDX2 depletion weakens intestinal, without strengthening foregut, enhancers. Thus, midgut endoderm is primed for heterologous cell fates, and TFs act on a background of shifting chromatin access to determine intestinal at the expense of foregut identity. Similar principles likely govern other fate commitments.

[*Keywords:* developmental competence; fate determination; tissue specification; lineage commitment; developmental plasticity; chromatin plasticity; homeodomain transcription factors]

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After embryonic cells acquire the competence to develop into a limited number of tissues, adult cell fates are determined in phases (Slack 1991). Possible cell identities are first specified, and developmental potential may remain flexible for some period until cells commit to unilineage differentiation. Each of these steps has a basis in transcription factor (TF) interactions with specific *cis*-regulatory elements (Shlyueva et al. 2014; Long et al. 2016). Enhancers recruited early in tissue ontogeny confer competence (Nord et al. 2013; Lara-Astiaso et al. 2014); cell identities emerge as organ primordia activate these and subsequently additional enhancers (Stergachis et al. 2013; Wang et al. 2015). For example, when human embryonic stem cells (ESCs) differentiate into pancreatic, liver, or lung lineages,

the active histone mark H3K4me1 is found on a large swath of enhancers, signifying broad tissue competence (Wang et al. 2015). Later, inductive signals drive different cell populations to stabilize a fraction of this early enhancer pool and thereby commit to a single fate.

We investigated the logic underlying enhancer recruitment and dynamic TF interactions with chromatin during development of the mouse alimentary canal, where distinctive epithelia with diverse morphologies and functions arise from undifferentiated gut endoderm. Because cell migration along the rostro-caudal axis is limited, the mucosa of each gut segment descends largely and directly from the corresponding regional endoderm

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(Fig. 1A). The molecular mechanisms are pertinent to human health because gastrointestinal atresias and malformations are among the commonest congenital disorders and because the largest risk for stomach and lower esophageal cancers occurs after native adult epithelia acquire heterologous (intestinal) character.

The pseudostratified epithelium lining the primitive mouse gut tube responds to regional cues to establish foregut (FG), midgut, and hindgut domains (Zorn and Wells 2009; Spence et al. 2011). Windows of developmental plas-

ticity are evident within these domains before cells adopt definitive squamous (esophagus [Eso] and forestomach [FS]) or columnar epithelial (glandular stomach and intestine) identities (Wells and Melton 1999; Grapin-Botton and Melton 2000). Some TFs are expressed broadly before approximately embryonic day 9 (E9)—when Wnt, bone morphogenetic protein (BMP), and other signals help establish an anterior–posterior pattern—and are later restricted to specific regions (Wells and Melton 2000). Moreover, endoderm transplanted before E13 adopts the

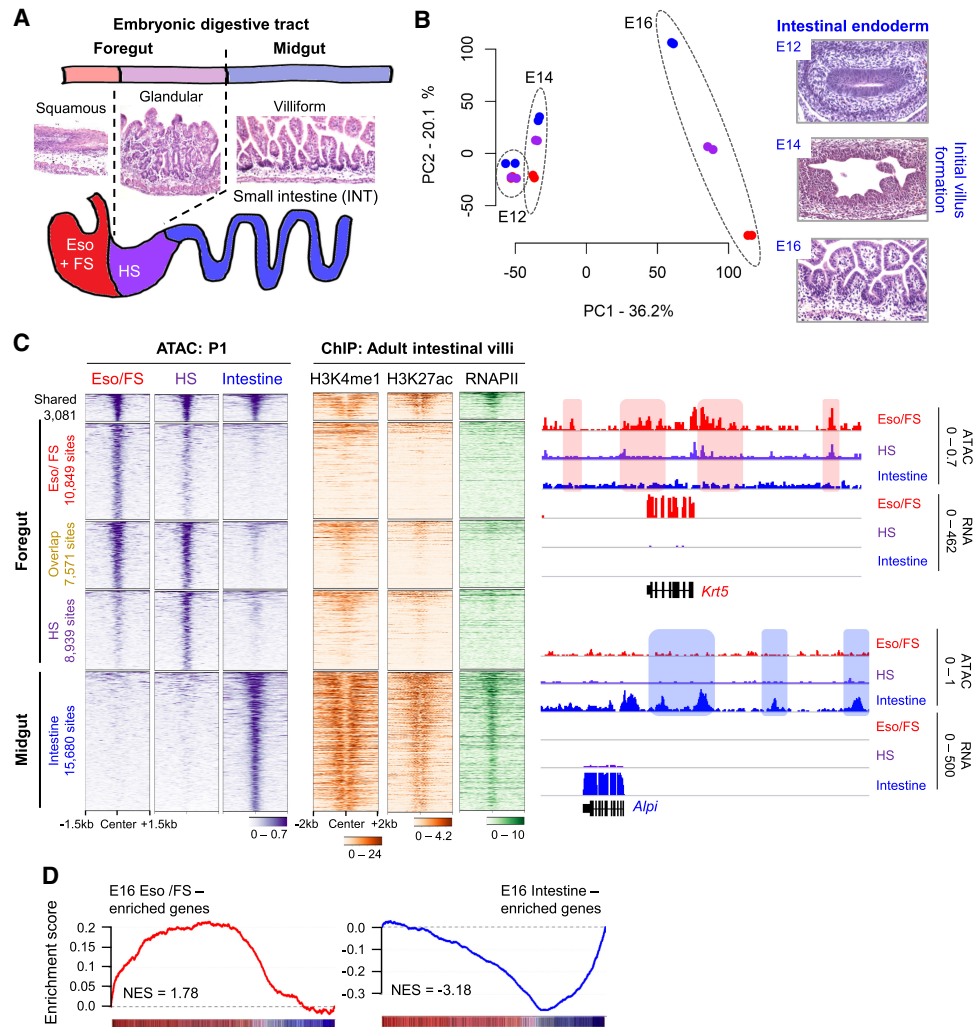


Figure 1. Transcriptional and chromatin landscapes in the developing and neonatal gut. (A) Gut epithelia with distinctive morphologies arise from contiguous endodermal regions. (Eso) Esophagus; (FS) forestomach; (HS) hindstomach; (Int) small intestine. (B) Principal component analysis (PCA) of Eso/FS (red), HS (purple), and Int (blue) transcriptomes at embryonic day 12 (E12), E14, and E16. $n = 2$ per group. The largest sources of variation reflect the emergence of tissue-specific mRNA profiles after E14. Photomicrographs of intestinal endoderm show pseudostratified epithelium at E12, early villus formation at E14, and mature villus structures at E16. (C) Differential chromatin access in postnatal day 1 (P1) endoderm from different regions of the tubal gut. Each cluster of ATAC-seq (assay for transposase-accessible chromatin [ATAC] using sequencing) summits ± 1.5 kb represents candidate enhancers selectively accessible in one region or common to the FS and HS (Overlap). In adult intestinal villi, the enhancers identified in P1 intestine carry H3K4me1, H3K27ac, and RNAPII, but these marks are largely missing from enhancers detected in P1 foregut tissues. Integrated Genome Viewer (IGV) tracks show ATAC-seq and RNA sequencing (RNA-seq) data at representative loci from Eso/FS and Int clusters. Numbers represent signal scales, and shaded boxes highlight region-specific enhancers. (D) Gene set enrichment analysis (GSEA) of genes located within 50 kb of sites selectively accessible in the Eso/FS or Int (derived using binding and expression target analysis [BETA]) (Wang et al. 2013), plotted with respect to differential mRNA expression in the E16 Eso/FS and Int. (NES) Normalized enrichment score.

regional identity of the host segment, whereas transplants of more advanced endoderm retain the identity of the donor region (Mizuno and Yasugi 1990; Duluc et al. 1994; Yasugi and Mizuno 2008). The chromatin and TF bases for initially labile midgut cell fate and subsequent commitment to intestinal differentiation are unclear.

In mid-gestation and beyond, the TF CDX2 is restricted to intestine-specific endoderm (Beck et al. 1995; Silberg et al. 2000) and adult intestinal epithelium (James et al. 1994). Forced expression in the developing mouse stomach (Mutoh et al. 2002; Silberg et al. 2002), but not in adult esophageal cells (Kong et al. 2011), induces enteric differentiation. Although germline CDX2 loss is lethal, heterozygote animals show focal stomach differentiation in the hindgut (Beck et al. 1999). Conditional loss at approximately E9 causes colonic atresia and replacement of the villous ileal mucosa by a stratified epithelium with Eso and FS features (Gao et al. 2009). In contrast, *Cdx2* deletion at E13 resulted in glandular stomach-like morphology and expression of gastric genes in the duodenum (Grainger et al. 2010), whereas deletion in adult intestines induced weak expression of few stomach genes (Verzi et al. 2010, 2011; Stringer et al. 2012) but precipitated lethal intestinal failure owing to collapse of CDX2-dependent enhancers (Verzi et al. 2013; Saxena et al. 2017). These studies implicate CDX2 in the highly contextual control of intestinal development and function. We postulated that investigation of CDX2–chromatin interactions during mouse development might illuminate the underpinnings of tissue competence, specification, and determination.

Results

Region-specific gene expression in the developing mouse gut is associated with distinct profiles of open enhancer chromatin

The gut endoderm generates a squamous lining in the Eso and FS and distinctive columnar epithelia in the hind-stomach (HS) and intestine (Zorn and Wells 2009). To study transcriptional and chromatin dynamics that underlie this rostro–caudal patterning, we purified EPCAM⁺ endodermal cells (Supplemental Fig. S1A; Sherwood et al. 2007) from discrete regions of the E12, E14, and E16 mouse gut (Fig. 1A): (1) the prospective FS and Eso, (2) the area between the FS and gastric pylorus (HS), and (3) the tube distal to the pylorus and proximal to the cecum (midgut or small intestine [Int]). RNA sequencing (RNA-seq) data from replicate samples (Supplemental Table S1) were highly concordant, and regional markers attested to the purity of cell isolates (Supplemental Fig. S1B). In principal component analysis (PCA) (Fig. 1B) and correlation analysis (Supplemental Fig. S1C), temporal changes accounted for the largest variation in gene expression, with mRNA profiles diverging by region after E12; these findings agree with observations that the intestinal lining is undifferentiated at E12 until villus primordia first appear at approximately E14 (Walton et al. 2012) and mature thereafter (Fig. 1B). *K*-means clustering of genes differentially expressed at E16 ($q < 0.05$, fold change ≥ 4 , reads

per kilobase per million mapped reads [RPKM] > 1) yielded groups with expression restricted to the Eso/FS, the HS, the Int, or two of these prospective epithelia (Supplemental Fig. S1D). These region-specific genes represent the purpose of digestive tract patterning and reflect the outcomes of spatio–temporal chromatin organization.

To determine the corresponding chromatin states, we first used the assay for transposase-accessible chromatin (ATAC) with sequencing (ATAC-seq) (Buenrostro et al. 2015) on Eso/FS, HS, and Int epithelia at postnatal day 1 (P1) (Supplemental Table S2). Replicate samples were highly concordant, regional differences in open chromatin were readily evident (Supplemental Fig. S2A,B), and difReps (Shen et al. 2013) identified genomic sites where chromatin access differed by region (Supplemental Fig. S2C). At sites located > 2 kb away from transcription start sites (TSSs), we thus detected candidate enhancers unique to each organ and sites shared among two or all three tissues (Fig. 1C). Areas selectively accessible in P1 intestine showed active histone marks and RNA polymerase II (Pol II) binding only in the adult intestine (Fig. 1C; Supplemental Fig. S2D), attesting that they are region-specific *cis* elements, and GREAT (Genomic Regions Enrichment of Annotations Tool) analysis (McLean et al. 2010) verified that genes < 50 kb from differential ATAC sites serve region-specific roles (Supplemental Fig. S2E). Tissue-restricted chromatin access of some *cis*-regulatory domains was readily apparent at known tissue-specific loci (Fig. 1C; Supplemental Fig. S2F,G), and binding and expression target analysis (BETA) (Wang et al. 2013) revealed association of the Eso/FS and Int site clusters with genes highly enriched in the E16 Eso/FS and E16 Int, respectively (Fig. 1D). Because enhancers can act at distances > 50 kb and on loci other than the nearest gene (Shlyueva et al. 2014; Long et al. 2016), such associations speak to likely activities of ATAC⁺ sites in aggregate and not the established function of any given site. Nevertheless, for simplicity, we refer to this collection of putative regulatory sites as “enhancers.”

Open chromatin dynamics identify a likely basis for cell fate plasticity in the developing intestine

To study trajectories of chromatin access that culminate in the neonatal tissues, we used ATAC-seq to map open chromatin in EPCAM⁺ cells purified from each regional endoderm at E12, E14, and E16. Figure 2A displays the normalized data from these series (quantified in Supplemental Fig. S3A,B) for every region-specific cluster. Although each P1 profile was strongest and restricted in the corresponding E16 epithelium, ATAC signatures in the FG were apparent in the respective regions by E14, whereas intestine-specific sites opened later (Fig. 2A). This differential trajectory is compatible with rostral-to-caudal progression of digestive tract differentiation, as judged by morphology in diverse vertebrate species (Wells and Melton 1999; Grapin-Botton 2005). Notably, FG-restricted profiles were evident at equal measure in all regions, including the Int, at E12 and E14; some FG sites even showed access in Int cells until E16. In contrast, a global view of Int

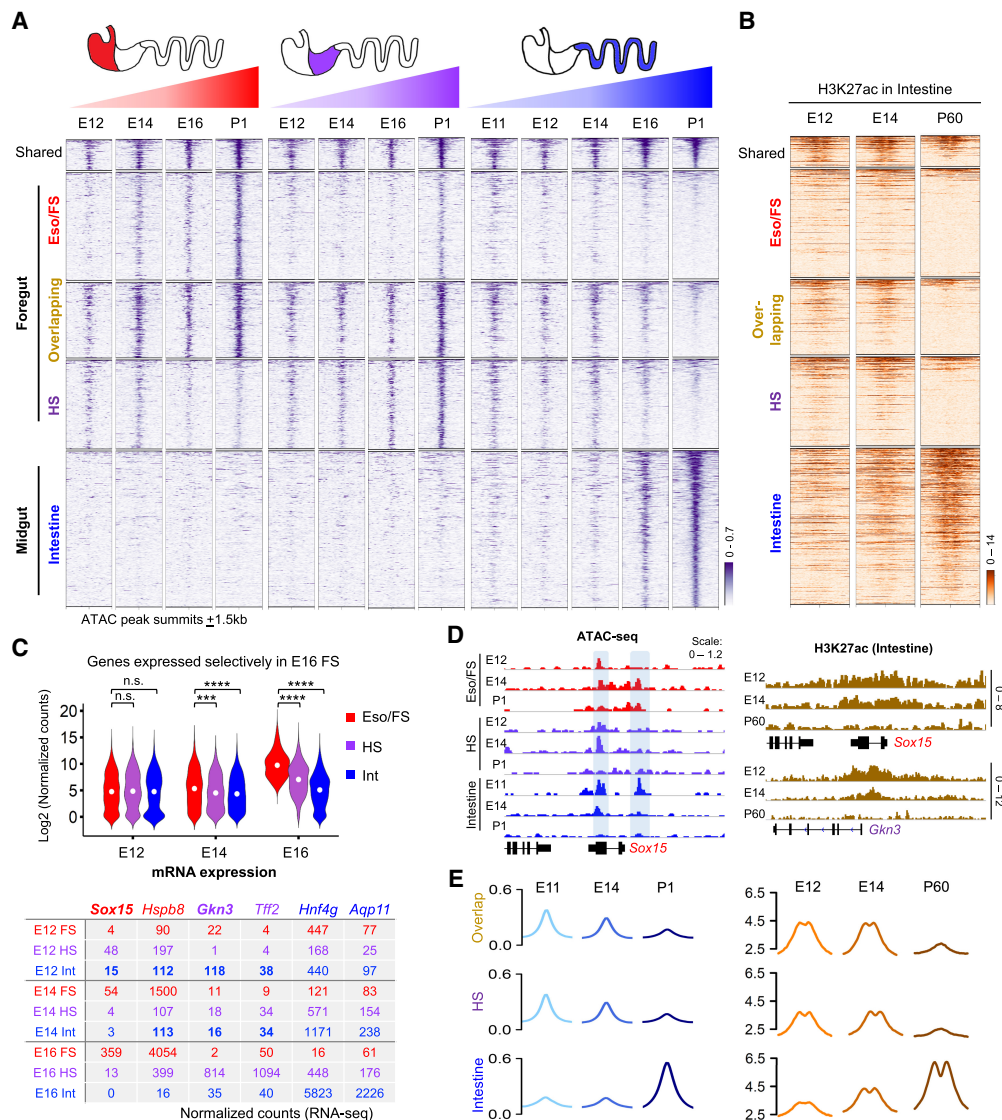


Figure 2. Extensive FG enhancer marking and activity in early Int endoderm. (A) ATAC-seq signals at region-specific P1 enhancers (from Fig. 1C) at successive stages in each regional endoderm: Eso/FS and HS (FG) and Int (midgut). FG endoderm lacked open chromatin at Int-specific enhancers at all times, whereas early Int endoderm showed extensive open chromatin at FG enhancers and loss of this chromatin access by P1. (B) H3K27ac ChIP-seq (chromatin immunoprecipitation [ChIP] combined with high-throughput sequencing) signals in E12, E14, and P60 Int endoderm (Kazakevych et al. 2017) at region-specific P1 enhancers. Early open chromatin at FG-specific enhancers is associated with H3K27ac, and both features are later lost. (C) Distributions of RNA levels among 824 Eso/FS-enriched genes (E16) (from Supplemental Fig. S1D) showing comparable global expression in the Eso/FS, HS, and Int at E12, before expression becomes selectively enriched in the Eso/FS. Dots in each violin plot represent median mRNA counts, and the significance of differences between groups was determined by the Mann-Whitney *U*-test. (n.s.) Not significant; (***) $P < 0.01$; (****) $P < 0.0001$. The table shows normalized RNA read counts for representative region-specific genes, illustrating FG-specific gene activity in early Int endoderm. (D) A representative Eso/FS locus, *Sox15*, carries open chromatin and H3K27ac in early (E11, E12, and E14) Int endoderm, later losing both features. Early, but not late, H3K27ac is also shown at a representative HS locus: *Gkn3*. (E) Aggregate intestinal ATAC-seq (blue; left) and H3K27ac ChIP-seq (brown; right) signals in the “Overlap,” HS, and Int enhancer clusters at different times. Chromatin access and H3K27ac at HS and “Overlap” enhancers decline during development, while both features arise at Int-specific enhancers.

enhancers revealed a lack of open chromatin at any time in FG-derived endoderm (Fig. 2A). These findings were evident in independent replicates of Int endoderm and do not reflect tissue contamination because the pylorus is an unmistakable landmark at the FG–midgut boundary. Nevertheless, we also examined Int endoderm at E11,

the earliest time that the Int and HS are visibly distinct and EPCAM is restricted to the endoderm. FG enhancers were as open in E11 Int endoderm as at E12 (Fig. 2A). As open enhancer chromatin reflects the potential to express nearby genes, these findings imply that before developing the ability to express Int genes, midgut endoderm may

harbor the potential to activate FG-specific genes and, by extension, for Eso/FS or HS differentiation. In contrast, FG endoderm appears to lack the reciprocal capacity for intestinal differentiation, at least by E12.

To ascertain whether open chromatin in Int bears other enhancer features, we analyzed data on the active histone mark H3K27ac in E12 and E14 EPCAM⁺ Int endoderm and adult epithelium (Kazakevych et al. 2017). Int-restricted enhancers carried H3K27ac in E14 and adult intestinal cells, as expected; so did many FG-specific sites in E12 and E14 intestinal endoderm (Fig. 2B). Both open chromatin and H3K27ac at FG enhancers were abrogated in P1 or adult Int cells (Fig. 2A,B), and to impute their *cis*-regulatory activity, we examined RNA levels of region-specific genes (Supplemental Fig. S1D), which the corresponding regional enhancers control (Fig. 1D). Median mRNA levels of FG genes were fourfold to >100-fold higher than those of Int genes in E16 FG endoderm but were comparable in E12 cells (Fig. 2C; Supplemental Fig. S3C). Eso/FS-specific *Sox15* (Sulahian et al. 2015), for example, shows selectively high mRNA in the rostral FG by E16 but even higher levels in Int than in Eso/FS at E12 (Fig. 2C). Like many such loci, *Sox15* and HS-restricted *Gkn3* are open and marked with H3K27ac in E12 Int endoderm and lose these features over time, even as the same sites gain enhancer features in the Eso/FS (Fig. 2D). These observations together reveal that thousands of FG enhancers are measurably active in undifferentiated Int endoderm and later decommissioned, concomitant with activation

of Int-specific enhancers and determination of Int cell fate (Fig. 2E).

The timing of CDX2 loss strongly influences homeotic intestinal transformation

Enhancer states represent the activities of sequence-specific TFs and recruited coregulators (Long et al. 2016). Indeed, motifs for known tissue-restricted TFs were the most enriched among the respective P1 tissue-specific enhancers (Fig. 3A). These TF transcripts were present at low levels in each E12 endoderm; over time, their levels increased in the cognate endoderm and declined reciprocally in heterologous epithelia (Fig. 3B). *Cdx2*, however, stands out because it is restricted to the intestine, as reported previously (Beck et al. 1995; Silberg et al. 2000), and peak levels were achieved by E12, much before regional endoderm specification. Conditional loss of *Cdx2* at different developmental stages has varied consequences (Gao et al. 2009). To determine its tissue requirements in relation to chromatin and TF dynamics, we deleted *Cdx2* using *Shh^{Cre}*, which acts in the luminal gut endoderm by approximately E10 (Echelard et al. 1993). Jejunal villi were stunted and, in lieu of crypt-villus units, the ileum had a stratified lining of the Eso/FS type (Fig. 3C, red arrows). Jejunal glands expressed the gastric parietal cell marker ATP4B at the base (Fig. 3C, white arrowheads), and, in place of scattered periodic acid Schiff (PAS)-stained goblet cells, duodenal and jejunal villi showed numerous gastric

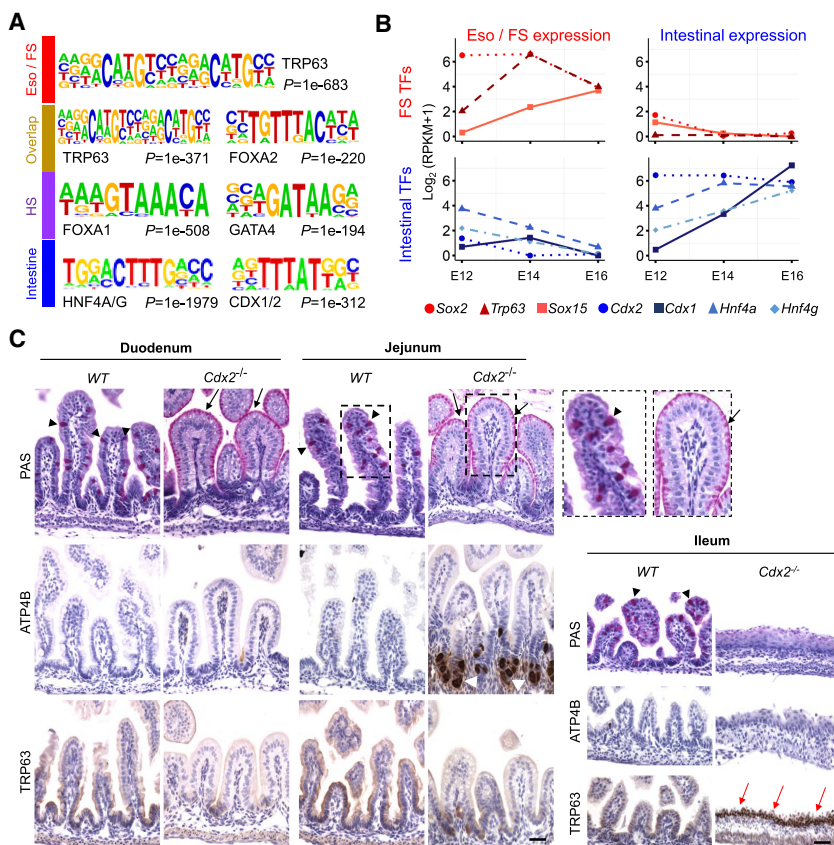


Figure 3. *Shh^{Cre}*-mediated *Cdx2* deletion results in anterior homeotic transformation. (A) The highest enriched sequences within region-specific enhancer clusters represent canonical TF motifs. (B) Trajectories of Eso/FS- and Int-specific TF transcripts in E12, E14, and E16 Eso/FS (left) and Int (right) endoderm. Log₂ transformed RPKM values, plotted over time, show increases and decreases in the different regions. (C) Intestinal regions (from the duodenum to the ileum) stained with PAS or with ATP4B or TRP63 antibody in E18 wild-type and *Shh^{Cre};Cdx2^{Fl/Fl}* embryos. In the mutant duodenum and jejunum, gastric foveolar cells (recognized by a mauve apical rim; black arrows) extensively replace intestinal goblet cells (recognized by a purple cytoplasmic blob; black arrowheads). These contrasting features of wild-type and mutant jejunal villi are highlighted in the dashed boxes and magnified at the right. In addition, ATP4B⁺ gastric parietal cells appear in the mutant jejunal (white arrowheads) but not duodenal crypts. The mutant ileum loses crypt-villus structure, including PAS⁺ goblet cells, and is lined instead by TRP63⁺ stratified squamous mucosa (red arrows). Bars, 25 μ m.

foveolar cells, where PAS classically stains the cellular apex (Fig. 3C, black arrowheads and arrows, respectively). Notably, these ectopic regional features were not intermingled: Squamous marker TRP63 was expressed in the *Cdx2*^{-/-} ileum but not the duodenum or jejunum, and, conversely, ATP4B was present in the jejunum but not the ileum (Fig. 3C). Thus, *Shh*^{Cre}-mediated CDX2 loss results in regionally confined alterations of endodermal fate determination, distinct from those reported when *Cdx2* is deleted earlier using *Foxa3*^{Cre} (Gao et al. 2009) or later using *Villin-Cre*^{ER-T2} (Grainger et al. 2010).

CDX2 requirements to establish intestinal fate shift during endoderm development

To study the transcriptional consequences of CDX2 depletion at different times, we isolated *Shh*^{Cre};*Cdx2*^{F1/F1} intestinal cells and treated pregnant *Villin-Cre*^{ER-T2};*Cdx2*^{F1/F1}

dams (Grainger et al. 2010) with tamoxifen to activate CRE on the 13th or 15th gestational day (Fig. 4A). When we activated CRE at or before E13, we isolated epithelial cells at E16 and (to give time for response to CDX2 loss after activating CRE at E15) E18. In every case, exon 2 mRNA (Supplemental Fig. S4A) and CDX2 protein (Fig. 4A) were depleted. *Shh*^{Cre}-driven loss of CDX2 yielded mRNA profiles that were the most distinct from wild-type E16 intestine and most similar to FG derivatives (Fig. 4B). Considering region-specific RNA clusters (Supplemental Fig. S1D), early CDX2 loss markedly attenuated Int-specific genes and produced substantial overlap with FG-restricted transcripts (Fig. 4C; Supplemental Fig. S4B). In contrast, deletion at E13 decreased Int transcripts modestly, and fewer FG genes were concomitantly activated, whereas deletion at E15 resulted in virtually no FG-specific gene expression (Fig. 4C), similar to adult *Cdx2*^{-/-} jejunum (Verzi et al. 2013). Moreover, after CRE activation

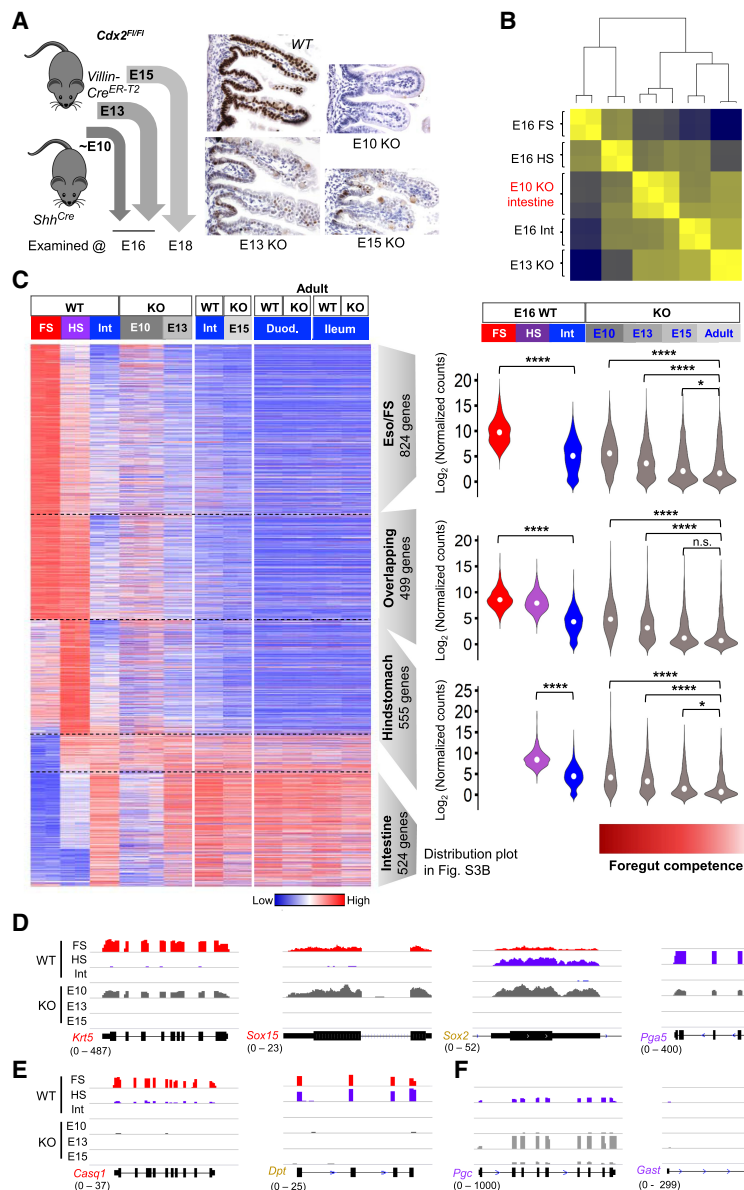


Figure 4. Time-sensitive activation of FG-specific transcriptional programs in the absence of CDX2. (A) Schema of *Cdx2* deletions mediated by either *Shh*^{Cre} (approximately E10) or treatment of *Villin-Cre*^{ER-T2} dams with tamoxifen at E13 or E15. Embryos deleted before E14 were assessed at E16, and those deleted at E15 were assessed at E18. CDX2 immunostains verify protein loss. (B) Unsupervised hierarchical clustering (Pearson coefficients) for the 2604 genes expressed differently (greater than fourfold, $q < 0.05$) in E16 wild-type tissues (Supplemental Fig. S1D) and E16 *Cdx2*^{-/-} Int after deletion at approximately E10 or E13. Transcriptomes in early, but not late, *Cdx2*-deleted Int endoderm resemble those in FG epithelia. (C) mRNA differences in *Cdx2*^{-/-} Int after deletion at approximately E10 ($n = 3$), E13, or E15 or in adult duodenum and ileum ($n = 2$ each). Clusters are those with differential expression in wild-type E16 EPCAM⁺ Eso/FS, HS, and Int cells (from Supplemental Fig. S1D). For each FG cluster, colored violin plots represent the distributions of mRNA levels (colors represent the region, and white dots represent the median) in wild-type tissues, and gray violins show the distributions after *Cdx2* deletion (knockout) at different times. Violin plots for Int genes are shown in Supplemental Figure S3B. The Mann-Whitney *U*-test was applied to determine the significance between *Cdx2* knockout at different times. (n.s.) Not significant; (*) $P < 0.05$; (****) $P < 0.0001$. (D–F) IGV tracks of RNA-seq data showing native (wild-type FS, HS, and Int) and ectopic FG gene expression in *Cdx2*^{-/-} intestines after deletion at E10, E13, or E15. Numbers represent the scale for normalized read counts.

in adult mice, duodenal and ileal *Cdx2*^{-/-} villus cells did not show significant levels of FG transcripts (Fig. 4C). Thus, CDX2 depletion markedly affects both FG and intestinal gene activity, with the largest effect on both programs occurring after early loss of CDX2 and progressively less disruption upon later losses.

Several features of FG gene activity in developing *Cdx2*^{-/-} intestines merit note. First, at any time of CDX2 loss, not all FG-specific genes were activated (Supplemental Fig. S4C). Expression of canonical Eso/FS and HS genes explains the FG morphology when *Cdx2* was deleted by E10 (Figs. 3C, 4D; Supplemental Fig. S4D), but other wild-type FG transcripts were never detected in the *Cdx2*^{-/-} midgut (Fig. 4E). Second, FG-specific mRNA levels in mutant intestines approached and often exceeded the native levels in the wild-type Eso or stomach (e.g., see Fig. 4D,F; Supplemental Fig. S4D), indicating robust ectopic gene activity. Third, transcripts expressed in both the wild-type Eso and stomach (“overlapping”) accounted for a substantial fraction of aberrant gene activity (Fig. 4C,D; Supplemental Fig. S4C), in line with the abundance of open chromatin at corresponding loci in the wild-type E11–E14 midgut (Fig. 2A). Fourth, whereas *Cdx2* deletion at E13 activated fewer FG genes and with less strength than deletion by *Shh*^{Cre}, some genes (especially overlapping and HS) were activated only after E13 deletion (Fig. 4F; Supplemental Figs. S4C,E). Finally, although global mRNA profiles after *Cdx2* deletion at E15 suggested almost no FG gene activity, ~20% of genes were activated greater than fourfold (Supplemental Fig. S4F).

These findings collectively demonstrate that midgut endoderm is initially biased toward FG differentiation and that CDX2 helps override this latent potential. Although midgut competence subsequently narrows, deletion of *Cdx2* as late as E13 increases intestinal mRNA levels of 53% of gastric genes by greater than twofold and 43% of gastric genes by greater than fourfold. By E15, however, the window for FG gene activity is closed and, except at a few residual enhancers, no longer requires CDX2 to prevent it. Of note, mRNA (Fig. 1B) and open chromatin (Fig. 2A) trajectories in wild-type embryos reveal a crucial transition at approximately E14, and *Cdx2* deletion through E13 reflects the same window of time, when endoderm is yet undetermined and FG genes can be stably activated in the midgut.

CDX2 loss at different times produces distinct tissue defects and open chromatin profiles

Foxa3^{Cre}-driven loss did not induce gastric morphology or ectopic stomach transcripts (Gao et al. 2009), whereas deletion at E13 induced HS cells only in part of the proximal duodenum and scant expression of gastric genes (Grainger et al. 2010). *Shh*^{Cre}-mediated *Cdx2* deletion likely occurs between these times and, in addition to ileal squamous differentiation, induces overt gastric features in the proximal and mid-intestine and robustly activates FG genes. The lower levels of FG-restricted transcripts when *Cdx2* was deleted at E13 or E15 predicted that anteriorizing homeotic transformation may be reduced or absent

at the later times. Indeed, following *Cdx2* deletion at E13 or E15, PAS staining throughout the intestine showed features of intestinal goblet—not gastric foveolar—cells, and the ileum did not carry a stratified epithelium but a typical columnar villous type (Fig. 5A). Thus, strikingly different phenotypes ensue when *Cdx2* is deleted at different times (Supplemental Fig. S4G), mirroring the inherent early accessibility and subsequent closure of FG enhancers.

The CDX2 sequence motif is enriched only at intestine-specific and not at FG-specific enhancers (Fig. 3A). Nevertheless, CDX2’s potent role in preventing FG gene activity could reflect repressive action from binding at FG enhancers or an indirect effect exerted by other means. To distinguish among these possibilities, we used CDX2 antibody for ChIP (chromatin immunoprecipitation) on EPCAM⁺ wild-type midgut endodermal cells, where it appeared at Int-specific enhancers by E14, and binding strengthened over time. In contrast, FG-specific enhancers globally showed negligible binding at E14 or E13, the earliest times that ChIP was feasible (Fig. 5B; Supplemental Fig. S5A), and inspection of individual FG sites failed to reveal appreciable CDX2 signals (Fig. 5C). This absence did not reflect ChIP insensitivity because binding at E13 and E14 was evident at other H3K27ac-marked enhancers with open chromatin (Supplemental Fig. S5B). Rather, the minimal overlap of E13 binding sites with regions of open FG-specific chromatin (Fig. 5D) implies that CDX2 precludes FG gene activation indirectly. In contrast, the sites and chronology of CDX2 binding at E14 and E17 (Fig. 5B) correlate with, and may underlie, intestine-specific gene activity.

To ascertain how FG- and Int-specific areas of open chromatin respond to loss of CDX2, we deleted *Cdx2* at various times and assessed Int endoderm by ATAC-seq. As with mRNA profiling, when *Cdx2* deletion occurred at E10 or E13, we studied EPCAM⁺ cells at E16, and, when *Cdx2* deletion occurred at E15, we examined epithelial cells 3 d later. We also examined adult *Cdx2*^{-/-} epithelium that we harvested from the ileum, the region of highest native expression (Silberg et al. 2000) and the most severely affected after embryonic (Fig. 3C) or adult (Verzi et al. 2010) CDX2 loss. *Shh*^{Cre}-driven loss of CDX2 and *Villin*^{Cre}-mediated loss at E13 caused a marked persistence of open chromatin at FG-specific enhancers, especially at “overlapping” and HS sites (Fig. 6A; Supplemental Fig. S5C,D). Among genes up-regulated after E10 deletion (Fig. 4C), 31% of TSSs lie <25 kb and 46% lie <50 kb from these areas of aberrantly open chromatin (Fig. 6B), and BETA analysis (Wang et al. 2013) confirmed strong statistical association (Supplemental Fig. S5E). Also in response to early *Cdx2* deletion, intestinal enhancers lacked open chromatin, indicating failure of intestinal gene activation. In contrast, deletion at E15 or in adults diminished intestinal ATAC signals to a lesser degree and failed to hold chromatin open at FG-specific enhancers (Fig. 6A; Supplemental Fig. S5C). These data identify a basis in chromatin for TF-dependent fate determination across thousands of region-specific *cis* elements (Fig. 6C). They also demonstrate CDX2’s dual role in de-commissioning FG enhancers without direct binding

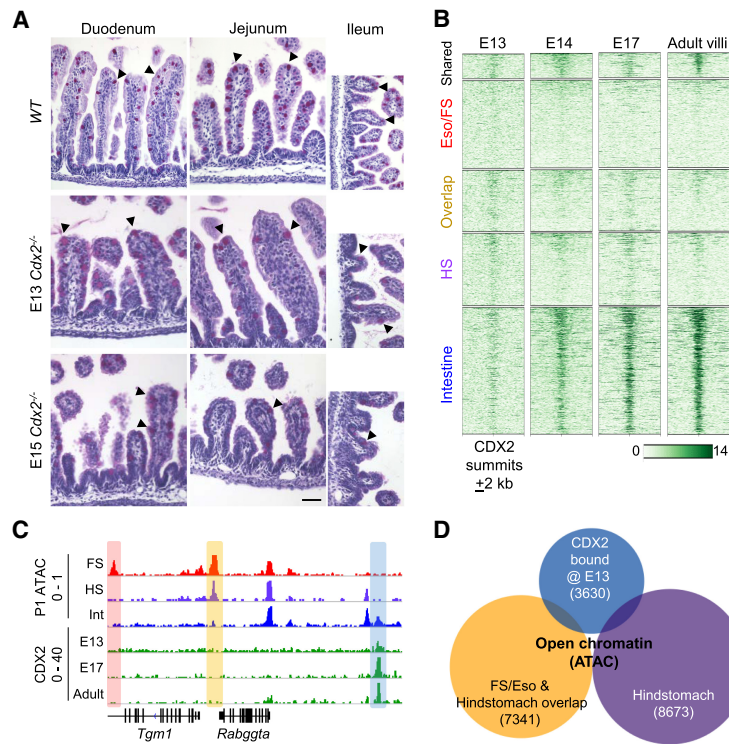


Figure 5. CDX2 binding in developing Int endoderm and the consequences of late depletion. (A) PAS staining of intestinal regions after *Cdx2* deletion at E13 or E15. Wild-type and mutant Ints were assessed at E18. Goblet cells (black arrowheads) were intact, and no region showed gastric foveolar differentiation (no cells were observed with apical rim staining). Bar, 25 μ m. (B) Normalized ChIP-seq data for CDX2 at different developmental stages. Binding at enhancers from any regional cluster is minimal at E13; thereafter, it increases progressively and selectively at Int enhancers. (C) Lack of CDX2 binding at P1 FG enhancers (red- and gold-shaded areas). The blue-shaded box demarcates a CDX2-bound intestinal enhancer. Numbers represent the scales of ChIP or ATAC signals. (D) E13-specific CDX2-bound sites overlap minimally with FG-specific enhancer regions, implying the absence of direct CDX2 control over FG genes.

and in establishing intestine-specific enhancers where CDX2 occupancy is pervasive.

Discussion

This study of mouse endoderm across developmental space and time reveals substantial chromatin, transcriptional, and cell fate plasticity foreshadowing the commitment to a unique epithelial identity. We report two core findings. First, before the midgut endoderm activates an intestine-specific transcriptional program and adopts villous morphology, enhancers that control FG-specific genes are broadly accessible to the same degree as in FG endoderm. At least after E11, the reverse is not true: Intestinal enhancers were inaccessible in multiple EPCAM⁺ cell isolates from the FS (future squamous) or HS (future glandular stomach). Beyond having accessible chromatin and the active histone mark H3K27ac, FG-specific genes near these *cis* elements are expressed in early midgut endoderm to levels at least the same as—and often higher than—the same transcripts in early FG endoderm. Together, these findings indicate that early midgut endoderm enhancers carry the potential to activate FG genes, and the tissue can therefore differentiate into stratified or glandular gastric epithelia. Thus, tissue determination reflects the imposition of intestinal—at the expense of FG—identity. Our observations mirror those showing broad priming of enhancers in human ESC-derived endoderm before differentiation of the pancreatic lineage (Wang et al. 2015). It is unclear why many FG genes express at even higher levels in early midgut than in FG endoderm (Fig. 2C). At a minimum, our findings imply that crucial FG TFs are present in

midgut endoderm and can engage accessible FG enhancers to drive transcription of FG genes.

A second core finding reveals the finite window in which CDX2 acts to suppress alternative tissue identity. Depletion of CDX2 at approximately E10 induces Eso/FS differentiation in the distal midgut, glandular HS differentiation in the proximal midgut, and failure to activate Int-specific genes. By E15 and in adult mice, acute CDX2 loss diminishes Int-specific transcripts but barely affects FG genes, whereas loss at E13 has an intermediate effect: activation of many FG-specific genes without induction of overt FG histology. Importantly, the window in which CDX2 is required to set chromatin marking and accessibility patterns (thus overcoming FG gene activity)—before but not after E14—coincides with that of initial epithelial divergence (Fig. 1B) and FG enhancer de-commissioning in the wild-type midgut (Fig. 2A). Thus, CDX2 loss before E14 stabilizes intrinsically accessible FG enhancers, which are subsequently unavailable to stabilize. These findings reveal the gene regulatory logic of rostro-caudal gut tube patterning and a chromatin basis for determination of one epithelial fate at the expense of others. Of note, our study examined only antero-posterior patterning of the luminal gut; development of other endoderm derivatives—the liver, pancreas, and lungs—may use the same or different regulatory logic.

The fact that FG enhancers open earlier than intestinal enhancers is consistent with observations that vertebrate gut endoderm differentiates in an anterior-to-posterior wave (Wells and Melton 1999; Grapin-Botton and Melton 2000). Moreover, judging by both profiles of open chromatin and the outcomes of *Cdx2* deletion at different times, anterior (FG) potential predominates throughout the early

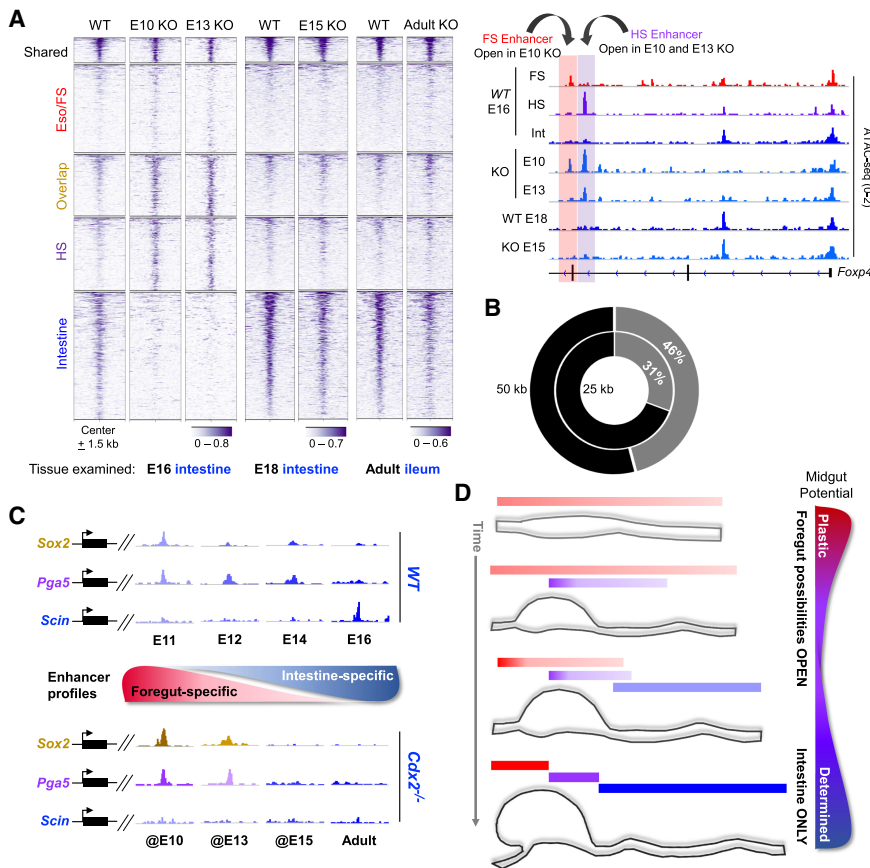


Figure 6. Persistence of open chromatin at FG enhancers after early, but not late, CDX2 loss. (A) Mice were treated to induce *Cdx2* deletion according to the schema in Figure 4A, and isolated Int epithelium was assessed for accessible chromatin by ATAC. “Overlap” and HS enhancers, which are ordinarily open in wild-type embryos until E14 (Fig. 2A), remained accessible after *Cdx2* deletion at approximately E10 or E13 but were closed after *Cdx2* deletion at E15 or in adult mice. Intestinal enhancers were significantly compromised after early, and less compromised after late, *Cdx2* deletion. The IGV track shows a representative locus: *Foxp4*. Shaded boxes denote Eso/FS- and HS-specific enhancers showing open chromatin after early *Cdx2* deletion. (B) The fractions of genes up-regulated in E10 *Cdx2*^{-/-} intestines (gray) located 25 or 50 kb away from “overlapping” or HS enhancers where ATAC signals persist in E10 *Cdx2*^{-/-} intestines. (C) Model for the temporal relation of enhancer accessibility to tissue plasticity in wild-type endoderm (top), as inferred from *Cdx2*^{-/-} intestines (bottom). IGV tracks show original ATAC-seq peaks at representative FG- and Int-specific enhancers located far from the respective TSSs. (D) Model depicting midgut developmental potential as a function of temporally regulated waves of enhancer accessibility and region-specific mRNA expression. Low-level expression

of FG genes is initially pervasive, while FG enhancers are accessible throughout the gut tube. Regional determination occurs after E14, when enhancer signatures become consolidated, gene expression domains become restricted, and regional anatomy approaches adult forms.

tubal endoderm and is superseded gradually by posterior (intestinal) fate in the midgut (Fig. 6D). Consequently, in the absence of factors that impose intestinal fate, distal regions may remain susceptible to anterior homeotic transformation for a longer time than proximal regions, which explains nicely why squamous differentiation is confined to the ileum when *Cdx2* is deleted at or before approximately E10. We note, however, that when *Cdx2* is deleted by *Shh*^{Cre}, prominent HS differentiation ensues in the jejunum (Fig. 3C), a homeotic transformation not seen with earlier deletion using *Foxa3-Cre* (Gao et al. 2009). To explain this difference, we postulate that Eso/FS and HS potential sweep through midgut endoderm in successive waves, each countered by CDX2-imposed intestinal identity. In terms of chromatin, Eso/FS-specific enhancers still dominate in the prospective ileum at approximately E10 (*Shh*^{Cre}), hence favoring squamous cell differentiation, while HS-specific enhancers have supplanted them in the future jejunum and promote gastric glandular differentiation. Meanwhile, Int-specific enhancers may replace both Eso/FS and HS enhancers early in the anterior midgut, so the prospective duodenum manifests intestinal potential when *Cdx2* is deleted using either *Foxa3-Cre* or *Shh*^{Cre}. Because our molecular analyses probed the whole

intestine, they lack the spatial resolution to confirm the details of this model (Fig. 6D), a limitation that also applies to other aspects of the study. For example, chromatin at thousands of “overlapping” and HS-specific enhancers was open in wild-type midgut endoderm until approximately E14, but we did not detect access at the global pool of FS-only enhancers. Moreover, depletion of CDX2 as early as E10 stabilized only “overlapping” and HS-unique enhancers and activated nearby genes but did not open FS-only sites, at least as detected in whole Int endoderm. Possibly, these enhancers were transiently open in the midgut before E11, or the signal was obscured because it was confined to small portions of the posterior wild-type and *Cdx2*^{-/-} midgut; alternatively, different patterning mechanisms may apply early and late in endoderm development. Distinguishing among these possibilities will require fine resolution of chromatin states in various regions of the E9–E11 midgut using gene-linked cell sorting or single-cell approaches to examine tiny cell populations before anatomic landmarks become clear.

Together with the tissue defects that result from CDX2 loss, the enhancer accessibility dynamics indicate that without intestinal TF activity, FG fates represent the

default outcome of mammalian endoderm differentiation. After FG potential is relinquished (approximately E14), however, loss of CDX2 alone is insufficient to restore it. Forced CDX2 expression also has different outcomes in the fetal mouse stomach, which develops with intestinal features (Mutoh et al. 2002; Silberg et al. 2002), and in adult gastric organoids, which, for the most part, retain stomach identity (Simmini et al. 2014). Thus, CDX2 can impose intestinal identity only during the brief window in development when enhancer dynamics permit choices among digestive endoderm fates. Such functions are not restricted to CDX2 and the intestine. SOX2 deficiency, in particular, results in a posterior homeotic transformation of the FS, with reduced keratinization and a columnar HS-type epithelium (Que et al. 2007), whereas continuous ectopic SOX2 expression in the midgut after E8.5 induces anterior homeosis with marked stomach differentiation (Raghoebir et al. 2012). SOX2 and CDX2 first appear at E7.75 in the most anterior and posterior endodermal cells, respectively, and their domains extend progressively toward the FG–midgut boundary, where the two TFs are coexpressed briefly before becoming mutually exclusive near the gastric pylorus by E9.5 (Sherwood et al. 2009). These temporal waves of expression—likely controlled by retinoic acid, BMP, and Wnt signaling posteriorly and by inhibition of these pathways anteriorly (Wells and Spence 2014; Davenport et al. 2016)—suggest how TF expression and chromatin dynamics together specify alternative endodermal fates. We propose further that initially accessible heterologous enhancers get decommissioned indirectly through mutual cross-repression by region-restricted TFs such as SOX2 and CDX2.

In conclusion, our findings offer a basis in chromatin for classical and recent observations on endoderm plasticity and determination. The repertoire of available enhancers, likely established by positional cues, dictates the limited spectrum of possible cell fates. As late as mid-gestation, the basis for cell plasticity is that different TFs can reinforce different sets of accessible tissue-specific enhancers and allow the remaining open sites to close. Irreversible commitment to one adult cell fate (determination) follows when specific TFs have reinforced the correct complement of *cis* elements, and the enhancers that enable alternative cell identities are no longer available. Similar enhancer logic and plasticity likely underlie the differentiation of other organs.

Materials and methods

Experimental mice and treatments

Wild-type CD1 mice were purchased from Charles River Laboratories. *Cdx2^{Fl/Fl}* (Verzi et al. 2010), *Shh^{Egfp-Cre} (Shh^{Cre})* (Harfe et al. 2004), and *Villin-Cre^{ER-T2}* (el Marjou et al. 2004) mice were reported previously. The morning of a vaginal plug was regarded as E0. Embryos from crosses between *Shh^{Cre};Cdx2^{Fl/+}* male and *Cdx2^{Fl/Fl}* female mice were used to isolate intestinal RNA or chromatin and subsequently genotyped to distinguish *Cdx2*-null from *Cdx2^{Fl/Fl};Cre⁻* and *Cdx2^{+/+};Cre⁺* controls. To delete *Cdx2* at later times, pregnant dams received 5 mg of tamoxifen (Sigma, T5648) dissolved in 0.5 mL of corn oil by oral gavage.

Adult *Cdx2^{Fl/Fl};Villin-Cre^{ER-T2}* mice received five daily intraperitoneal injections of tamoxifen (1 mg per 25 g of body weight or higher) prepared in corn oil and were euthanized on the third day after the last injection. *Cre⁻* mice served as littermate controls. All animal experiments were approved and monitored by Animal Care and Use Committees at Dana-Farber Cancer Institute and Rutgers University.

Tissue collection and cell isolation

Embryos were collected in ice-cold phosphate-buffered saline (PBS), and the digestive tract was collected in ice-cold Hank's balanced salt solution (HBSS; Thermo Fisher). The rostral half of E12 and E14 stomachs was considered the FS, and the caudal half was considered as the HS. In E16 embryos and neonates, the visible squamo-columnar junction was applied to separate the FS from the HS, discarding the interim zone. Intestines were harvested from 1 mm distal to the pylorus until (but excluding) the cecum. Tissues were digested with 0.25% Trypsin-EDTA (Thermo Fisher) for 30 min at 37°C. Released material was passed over 40- μ m filters (Corning) to obtain single cells. Neonatal intestines were washed with ice-cold PBS and incubated in 5 mM EDTA solution in PBS at 4°C with gentle shaking for 30 min to release villus epithelium. Suspensions were passed through 70- μ m filters (Corning) to retain villi, which were washed in cold PBS and dissociated into single cells by incubation in 4 \times TrypLE solution (Thermo Fisher) for 30 min at 37°C. After neutralization with 5% FBS, cells were strained over 40- μ m filters, pelleted at 1200g for 5 min, resuspended in FACS buffer (PBS, 5 mM EDTA, 2% FBS), and stained with APC-conjugated EPCAM antibody (1:100; Biolegend) for 1 h at 4°C in FACS buffer. EPCAM⁺ cells were sorted on an Aria II SORP flow cytometer (Becton-Dickinson). In some experiments, single cells were incubated with anti-CD326 (EPCAM) antibody conjugated to magnetic microbeads (Miltenyi Biotec, 130-105-958) for 40 min on ice and then collected over MS Columns (Miltenyi Biotec, 130-042-201) in a magnetic field. The purity of EPCAM⁺ cells isolated by magnetic beads or flow cytometry was comparable. All replicates in the study refer to biological replicates collected on different days. In rare experiments lacking a biological replicate, data from other samples or time points support the conclusions.

RNA-seq, ATAC-seq, and ChIP-seq (ChIP combined with high-throughput sequencing)

Total RNA was isolated from FACS-sorted epithelial cells using TRIzol (Thermo Fisher) and RNeasy microkit (Qiagen). Libraries were prepared using the SMARTer-Seq version 4 low-input mRNA library kit (Clontech). Five-thousand to 30,000 cells were used for ATAC-seq as described previously (Buenrostro et al. 2015), with minor modifications. Cells were lysed in ice-cold buffer (10 mM Tris at pH 7.4, 10 mM NaCl, 3 mM MgCl₂, 0.1% [v/v] Igepal CA-630), and nuclei were precipitated by centrifugation at 500g for 10 min at 4°C were treated with Nextera Tn5 transposase (Illumina, FC-121-1030) for 30 min at 37°C. Transposed chromatin was purified with QIAquick PCR purification kits (Qiagen) and amplified with High-Fidelity 2X PCR Master Mix (New England Biolabs) using a universal forward primer and unique reverse primers in a 50- μ L reaction for five cycles. Five microliters of the reaction mix was amplified by quantitative PCR for 20 cycles to determine the optimum number of additional cycles, defined as one-third the number needed to achieve maximum fluorescence intensity. Primer dimers were excluded from these PCR-amplified libraries using Agencourt AMPure XP beads (Beckman Coulter).

For fetal ChIP-seq, intestines were dissected from up to 36 embryos in ice-cold HBSS and disrupted by vigorous pipetting followed by cross-linking in 1% formaldehyde for 20 min at room temperature, incubation for 10 min on ice in chromatin preparation buffer (ChIP-IT high-sensitivity kit, Active Motif, 53040) with protease inhibitors (Roche, 11873580001) and phenyl methyl sulfonyl fluoride (PMSF), and centrifugation at 1250g for 5 min at 4°C. The pellet was resuspended in 0.5 mL of ChIP buffer (ChIP-IT high-sensitivity kit, Active Motif) and incubated for 10 min on ice. Lysates were sonicated in an E220 focused ultrasonicator (Covaris). CDX2 antibody-bound (Cell Signaling Technology) protein A and protein G magnetic Dynabeads (1:1 ratio; Thermo Fisher) were incubated overnight at 4°C with 2 µg of chromatin in 300 µL of ChIP buffer containing protease inhibitors and two carriers: 20 µL of recombinant human histone H2B solution (New England Biolabs, M2505S) and 1 µL of carrier RNA from the RNeasy microkit (Qiagen, 74004). Beads were washed five times with LiCl-RIPA buffer (10 mM Tris-HCl at pH 8, 500 mM LiCl, 1 mM EDTA, 1% NP-40) and once in 10 mM Tris-EDTA (pH 8). Beads resuspended in 100 µL of immunoprecipitation elution buffer containing 25 µg of RNase A were incubated for 30 min at 37°C followed by 1 µL of 20 mg/mL proteinase K (Active Motif), and cross-links were reversed overnight at 65°C. DNA was purified using MinElute PCR purification kits (Qiagen). ChIP-seq for adult villi with CDX2 (Cell Signaling Technology, 12306) or H3K4me1 (Diagenode, C15410194) antibody was as described previously (Saxena et al. 2017). Libraries were prepared using ThruPLEX kits (Rubicon, R400427).

All libraries were sequenced on a NextSeq 500 instrument (Illumina) to generate 75-base-pair (bp) single-end reads.

Histochemistry and immunohistochemistry (IHC)

Intestines were fixed overnight in 4% paraformaldehyde, washed in PBS, dehydrated in an ethanol series, and embedded in paraffin to prepare 5-µm-thick sections. Slides were incubated in 0.5% periodic acid and stained with Schiff's reagent (Alfa Aesar, J612171), and alkaline phosphatase (ALPI) activity was measured using AP staining kit II (Stemgent). For IHC, antigens were retrieved in 10 mM sodium citrate buffer (pH 6), and sections were then incubated overnight at 4°C with antibody against CDX2 (Cell Signaling Technology, 12306), ATP4B (MBL International Corp., D032-3), or TRP63 (Santa Cruz Biotechnology, sc-8343) followed by biotin-conjugated anti-rabbit or anti-mouse IgG (Jackson Laboratories); washed; and stained with Vectastain Elite ABC kit (Vector Laboratories) and 3,3' diaminobenzidine tetrahydrochloride (Amresco, 0430).

RNA-seq analysis

Reads were aligned to the mouse reference genome Mm9 (NCBI build 37) using STAR aligner version 2.5.3a (Dobin et al. 2013), and raw read counts from HTSeq (Anders et al. 2015) were used to estimate transcript levels. Unsupervised PCA was performed on log₂ transformed normalized counts using R (<https://www.r-project.org>). We then used DESeq2 (Love et al. 2014) to normalize data using size factors or rlog transformation from E16 wild-type FS, HS, and Int ($n = 2$ samples each) and identify significant ($q < 0.05$) differential expression. RPKM values were calculated using Cufflinks (Trapnell et al. 2012). Pearson correlation coefficients of normalized count values from biological replicates ($n \geq 2$ samples each) were calculated using the corrrgram package in R. Data tracks to visualize on the Integrative Genomics Viewer (IGV) version 2.3 (Robinson et al. 2011) were generated using bamCoverage modules in deepTools2 (Ramirez et al. 2016). We

determined optimal cluster numbers for K -means clustering of rlog transformed counts using the clusGap function in R, derived Gene Ontologies in DAVID (Huang et al. 2009), and generated violin plots using the ggplot2 package (Wickham 2009) in R studio. We used GENE-E software (Broad Institute) to plot log transformed values in heat maps.

To determine genes associated with P1 Eso/FS- and Int-specific enhancer clusters (Fig. 1) and up-regulated genes associated with retained FG enhancers (Supplemental Fig. S5E), we used BETA (Wang et al. 2013). Enriched expression of gene sets in the FS or Int was determined using gene set enrichment analysis (GSEA) (Subramanian et al. 2005), and normalized enrichment scores were derived using 1000 permutations of random gene sets of the same size. RNA-seq data from wild-type and *Cdx2*^{-/-} (deleted at E10) intestines were associated with a merged set of HS and “overlapping” ATAC⁺ sites persisting in E10 deleted *Cdx2*^{-/-} intestines.

ATAC-seq and ChIP-seq analyses

For ChIP-seq, 75-bp single-end reads were aligned to mouse reference genome Mm9 (NCBI build 37) using Bowtie2 (Langmead and Salzberg 2012). Following routine quality control, we reassessed all ATAC-seq data and aligned reads using the ENCODE pipeline (Supplemental Tables S1, S2). Nonunique and duplicate reads were removed using SAMtools version 1.3.1 (Li et al. 2009). We used individual or merged raw signals from replicate ChIP-seq and ATAC-seq files to visualize in IGV tracks and then created Bigwig files using the bamCoverage module in deepTools2. Data were quantile-normalized using Haystack version 0.4 (Pinello et al. 2014) followed by deepTools2 to plot heat maps. ATAC-seq signals were computed using SitePro in Cistrome tools (Liu et al. 2011), and intensity values were used to plot cumulative signals using R.

Differential ATAC-seq signals between tissue pairs were identified using DiffReps version 1.55.4 (Shen et al. 2013) with z -score cutoff of 5, window size of 500 bp, step size of 100 bp, and P -value of 0.05. Enhancers were defined by excluding promoters (± 1 kb from a TSS). We then used BEDTools (Quinlan and Hall 2010) to determine the overlap between the FS and HS in the DiffReps output. These overlapping regions were subtracted from the individual FS and HS sets and assigned to a distinct “Overlap” set, while others were considered specific to one region or the other. Similarly, sites common or unique to FS versus Int and HS versus Int pairs were determined by intersecting with the set of Int-specific regions. To identify the shared regions, peaks were called using MACS2 (Zhang et al. 2008) for each tissue and intersected using BEDTools, excluding promoters.

To determine correlations between samples, normalized read counts from replicate samples were collated over merged called peaks (generated by MACS2) from individual bam files using BEDTools. To determine cross-correlations, read counts were collated over peak files merged from each sample and plotted in R. Functions of genes within prescribed distances from ATAC sites were identified using GREAT version 3.0 (McLean et al. 2010), and motif enrichment was identified using HOMER version 4.7.2 (Heinz et al. 2010) based on cumulative binomial distributions against the genome background. ChIP-seq and ATAC-seq read counts ± 250 bp from peak summits were calculated from normalized data and are represented as log transformed values in violin plots. Differences among groups were determined using the Kolmogorov-Smirnov test. As P -values were generally significant (< 0.0001 to < 0.01) owing to large input sample sizes, we inferred similarities and differences using the D statistic, the maximum distance between two cumulative distribution functions (e.g., see Supplemental Figs. S3B, S5C).

Data availability

All data have been deposited in the Gene Expression Omnibus, series GSE115541.

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