

# Global Epigenomic Analysis of Primary Human Pancreatic Islets Provides Insights into Type 2 Diabetes Susceptibility Loci

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## SUMMARY

Identifying *cis*-regulatory elements is important to understanding how human pancreatic islets modulate gene expression in physiologic or pathophysiologic (e.g., diabetic) conditions. We conducted genome-wide analysis of DNase I hypersensitive sites, histone H3 lysine methylation modifications (K4me1, K4me3, K79me2), and CCCTC factor (CTCF) binding in human islets. This identified ~18,000 putative promoters (several hundred unannotated and islet-active). Surprisingly, active promoter modifications were absent at genes encoding islet-specific hormones, suggesting a distinct regulatory mechanism. Of 34,039 distal (nonpromoter) regulatory elements, 47% are islet unique and 22% are CTCF bound. In the 18 type 2 diabetes (T2D)-associated loci, we identified 118 putative regulatory elements and confirmed enhancer activity for 12 of 33 tested. Among six regulatory elements harboring T2D-associated variants, two exhibit significant allele-specific differences in activity. These findings present a global snapshot of the human islet epigenome and should provide functional context for noncoding variants emerging from genetic studies of T2D and other islet disorders.

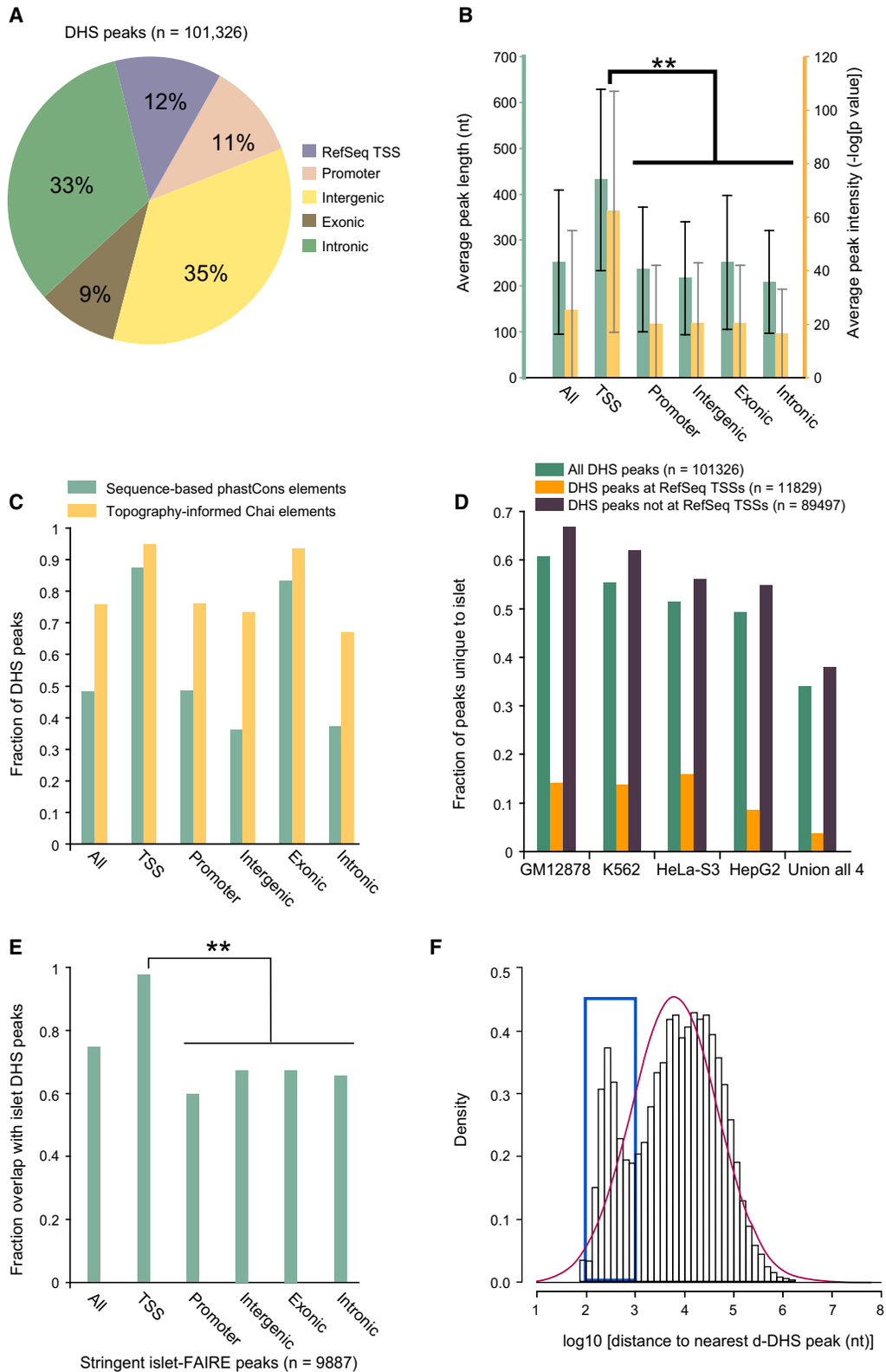
## INTRODUCTION

Type 2 diabetes (T2D) is a complex metabolic disorder that accounts for 85%–95% of all cases of diabetes and afflicts hundreds of millions of people worldwide (<http://www.diabetesatlas.org/content/diabetes>). It is a leading cause of substantial morbidity and is characterized by defects in insulin sensitivity and secretion resulting from the progressive dysfunction

and loss of  $\beta$  cells in the pancreatic islets of Langerhans (Butler et al., 2007; Muoio and Newgard, 2008). Both genetic predisposition and environmental factors contribute to these islet defects. Islets constitute 1%–2% of human pancreatic mass (Joslin and Kahn, 2005) and are composed of five endocrine cell types that secrete different hormones:  $\alpha$  cells (glucagon),  $\beta$  cells (insulin),  $\delta$  cells (somatostatin), PP cells (pancreatic polypeptide Y), and  $\epsilon$  cells (ghrelin). These cells sense changes in blood glucose concentration and respond by modulating the activity of multiple pathways, including insulin and glucagon secretion, to maintain glucose homeostasis (Joslin and Kahn, 2005). Several key transcription factors (TFs) that regulate these responses are known (Oliver-Krasinski and Stoffers, 2008). However, efforts to identify *cis*-regulatory elements upon which these and other factors act have been restricted primarily to promoter regions at specific loci (e.g., *INS*, *PDX1*) (Brink, 2003; Ohneda et al., 2000).

Results from genome-wide association studies (GWAS) of type 1 diabetes (T1D) (Barrett et al., 2009), T2D (reviewed in Prokopenko et al., 2008), and related metabolic traits (Dupuis et al., 2010; Ingelsson et al., 2010; Prokopenko et al., 2009) suggest that genetic variation in *cis*-regulatory elements may play an important role in  $\beta$  cell (dys)function and diabetes susceptibility (De Silva and Frayling, 2010). Of the 18 most strongly associated single-nucleotide polymorphisms (SNPs) in each of the T2D-associated loci, only 3 are missense variants; the remaining are noncoding (Prokopenko et al., 2008). Furthermore, there is evidence for allele-specific effects of two T2D-associated SNPs on the islet expression level of nearby genes (*TCF7L2* [Lyssenko et al., 2007] and *MTNR1B* [Lyssenko et al., 2009]). However, the dearth of annotation of functional regulatory elements has limited the capacity to investigate the role of regulatory variation in complex diseases such as T2D.

Recent characterization of histone modifications and DNase hypersensitivity in cultured cells has identified chromatin signatures predictive of regulatory elements and actively transcribed regions (Boyle et al., 2008; Guenther et al., 2007; Heintzman et al., 2007). The data generated so far suggest that regulatory



**Figure 1. Analysis of DNase I Hypersensitive Sites in the Islet Genome**

(A) Distribution of DNase I-hypersensitive (DHS) peaks across five genomic annotation sets. “Promoter” denotes proximal regions 5 kb upstream of RefSeq transcription start sites (TSSs) that do not overlap the TSS. “Exonic” represents regions that overlap at least one base with an exon.

element location and usage vary substantially among cell types (Heintzman et al., 2009; Xi et al., 2007). Also, extensive chromatin profiling has been conducted in very few human primary tissues to date (Bhandare et al., 2010). In this study, we describe a comprehensive genome-wide epigenomic map of unstimulated human pancreatic islets. Using DNase- and ChIP-seq approaches, we identified DNase I-hypersensitive sites that mark regions of open chromatin, loci enriched for active histone H3 lysine methylation modifications (H3K4me1, H3K4me3, and H3K79me2), and binding sites for the insulator CCCTC-binding factor (CTCF). These profiles provide a detailed chromatin snapshot of regulatory elements and actively transcribed units in the islet. Moreover, they identify regulatory elements harboring T2D-associated variants in 6/18 loci. These data provide a valuable resource for understanding and investigating *cis*-regulation in the human islet and for discovering regulatory elements that may play an important role in diabetes susceptibility.

## RESULTS

### Genome-wide Characterization of Open Chromatin in the Human Pancreatic Islet

Active regulatory elements reside in open chromatin regions hypersensitive to DNase I digestion (ENCODE Project Consortium, 2007; Boyle et al., 2008; Crawford et al., 2004; Hesselberth et al., 2009; Sabo et al., 2004). To identify all DNase-hypersensitive sites (DHS) in the human pancreatic islet, we performed DNase-seq (Boyle et al., 2008) and identified regions of the genome with significant enrichment of sequence reads using the MACS algorithm (Zhang et al., 2008) (Experimental Procedures). This approach identified 101,326 human islet DHS peaks (Table S1) covering ~27 million bases (~1% of the human genome). Consistent with observations in CD4+ T cells (Boyle et al., 2008), a substantive fraction of islet DHS peaks (23%,  $n = 23,408$ ) span annotated RefSeq transcription start sites (TSS) or are within regions 5 kb upstream (Promoter), but the majority reside within currently unannotated genomic regions that may harbor functional distal regulatory elements (Figure 1A). Peaks at TSSs are significantly longer and more intense than those at all other loci (Figure 1B). This observation supports the view that regions around TSSs are generally more susceptible to DNase I digestion than putative non-TSS regulatory elements (Boyle et al., 2008).

Approximately 48% ( $n = 48,777$ ) of all DHS peaks overlap phastCons vertebrate conserved elements (Siepel et al., 2005) (Figure 1C). Notably, ~87% (10,348/11,829) of peaks at TSSs overlap phastCons elements, compared to ~43% (38,429/89,497) at non-TSS loci (Figure 1C). This difference remains even after accounting for the longer peaks at TSSs (data not shown), supporting the model that TSS-proximal regions evolve under stronger sequence constraint than distal regulatory elements (Boyle et al., 2008). A recent study developed an algorithm (Chai) for topography-informed conservation analysis, which identified ~2-fold more bases in the human genome under evolutionary constraint compared to sequence-based methods (Parker et al., 2009). Accordingly, ~1.5 times as many (~76%) islet DHS peaks overlap these structurally constrained regions (Figure 1C).

To determine the extent of cell-type specificity of our islet DHS peaks, we obtained DNase-seq data generated for four different human cell lines: GM12878, K562, HeLa-S3, and HepG2 (Duke DNase, ENCODE Project Consortium, 2007). We identified DHS peaks for these cell lines (Experimental Procedures) and found that roughly half the islet peaks are shared with each individual nonislet cell type. Notably, ~35% ( $n = 34,273$ ) are completely unique to the islet (Figure 1D). Almost all (~99%) of these islet-unique peaks do not overlap RefSeq TSSs, which is consistent with the model that tissue-specific gene expression patterns are governed largely by distal *cis*-regulatory elements (Heintzman et al., 2009).

An independent method to map open chromatin is formaldehyde-assisted isolation of regulatory elements (FAIRE) (Giresi et al., 2007). Recently, this approach was used for human islets to identify three sets of candidate peaks, including “stringent” ( $n = 9887$ ) and “liberal” ( $n = 100,715$ ) peaks (Gaulton et al., 2010). Approximately 75% of the “stringent” islet FAIRE peaks overlap DHS peaks. However, this corresponds to only 7360 peaks, which is far fewer than the predicted number of functional regulatory elements genome-wide (ENCODE Project Consortium, 2007). The overlap is significantly greater at TSSs compared to non-TSSs (97% versus 65%) (Figure 1E). Comparing DHS peaks to the set of “liberal” islet FAIRE peaks, the overlap drops to ~29%. Therefore, the two approaches seem to identify distinct sets of non-TSS regulatory elements. Because it is difficult to assess the extent to which the dissimilarity between DHS and FAIRE data is explained by differences in islet sample purity, preparation methods, false positive signals, or population

(B) Average length (teal) and intensity (yellow) of DHS peaks across five genomic annotation sets. Peaks at RefSeq transcription start sites (TSSs) are significantly longer and more intense than those elsewhere (\*\*, two-tailed paired Student's *t* test,  $p$  value  $< 10^{-100}$ ). Error bars represent SD (SD measurements were often greater than the sample average due to highly skewed distributions, but error bars were cut off at zero for visualization).

(C) Sequence and structure constraint at DHS. DHS peaks at RefSeq TSSs are under substantially greater sequence constraint (assessed by phastCons vertebrate conservation scores) than intronic and intergenic DHS peaks. A large majority of DHS peaks within all genomic annotation sets are under strong structural constraint (assessed by the Chai algorithm) (Parker et al., 2009).

(D) Comparison of islet DHS peaks with peaks from four different human cell lines. Each data point represents the fraction of total peaks ( $n = 101,326$ ) unique to the human islet relative to each of the other four human cell types or all of them combined (Union of all 4). Roughly 35% are unique to the islet, and 99% of these are not located at RefSeq TSSs. Varying levels of similarity across cell types may be at least partially explained by differences in the stage of cellular differentiation and/or sequencing depth.

(E) Overlap between DHS peaks and formaldehyde-assisted isolation of regulatory elements (FAIRE) peaks. The overlap is significantly greater at RefSeq TSSs than elsewhere (\*\*, Fisher's exact test  $< 10^{-100}$ ).

(F) Logarithm-based distribution of the distance to the nearest distal DHS (d-DHS) peak among all d-DHS peaks. The blue box indicates an increased representation of peaks in the ~100–1000 bp range (clustered) relative to Gaussian expectation (red curve). This range is significantly enriched for islet-unique peaks (Fisher's exact test,  $p = 2.7 \times 10^{-9}$ ). Comparison of d-DHS, FAIRE, and GLTR locations is found in Figure S1.

diversity (McDaniell et al., 2010), more controlled comparisons of these techniques will be necessary to elucidate inherent preferences of each for specific classes of open chromatin.

Though many of the mechanistic details are not clear, it is widely accepted that distal and promoter regulatory elements can exert coordinated control of gene transcription via physical interactions (Dekker, 2003; Miele and Dekker, 2008). Therefore, it has been hypothesized that distal *cis*-regulatory elements may cluster together to form functional modules (Blanchette et al., 2006). To assess the clustering of putative islet-active distal *cis*-regulatory elements, we filtered from the islet DHS peaks ( $n = 101,326$ ) the regions that may represent promoters to identify a set of high-confidence distal peaks (d-DHS,  $n = 34,039$ ) (Table S2 and Figure S1 and Experimental Procedures). For each d-DHS peak, we computed the distance to the nearest d-DHS peak and observed an increased representation in the  $\sim 100$ – $1000$  bp range ( $n = 7652$ ) relative to the expectation from a normal distribution (Figure 1F). Furthermore, this set is significantly enriched for islet-unique peaks ( $p = 2.7 \times 10^{-9}$ ).

### Genome-wide Characterization of TSSs in the Islet Genome via H3K4me3 ChIP-Seq

To characterize human islet TSSs, we conducted ChIP-seq analysis of histone 3 lysine 4 trimethylation (H3K4me3) in four different human islet samples. H3K4me3 is enriched at CpG islands (Bernstein et al., 2007), TSSs (Li et al., 2007), and sites of active transcription (Kouzarides, 2007). Enriched regions present in all four islet samples, but absent from three mock-IP (anti-GFP) experiments, were designated as “H3K4me3 peaks.” This method identified 18,163 human islet H3K4me3 peaks (Table S3) covering  $\sim 1\%$  of the genome.

As expected, approximately two-thirds ( $n = 11,973$ ) of H3K4me3 peaks overlap RefSeq TSSs (Figure 2A). Greater than 70% of the remaining, unannotated peaks ( $n = 6190$ ) overlap computationally predicted TSSs and/or CpG islands. However, the significantly lower average length and intensity of unannotated H3K4me3 peaks compared to those at RefSeq TSSs (Figure 2B) suggests that at least some of these peaks may indicate weakly active TSSs, inactive but poised TSSs (Barski et al., 2007; Guenther et al., 2007; Mikkelsen et al., 2007), remnants of transcriptional activity from the developmental past or prior environmental stimulation (Barski et al., 2009), or chromatin looping with distal regulatory regions. While a subset of peaks could be false-positive signals, this is unlikely, as it would require a technical artifact that is consistent across all four islet samples.

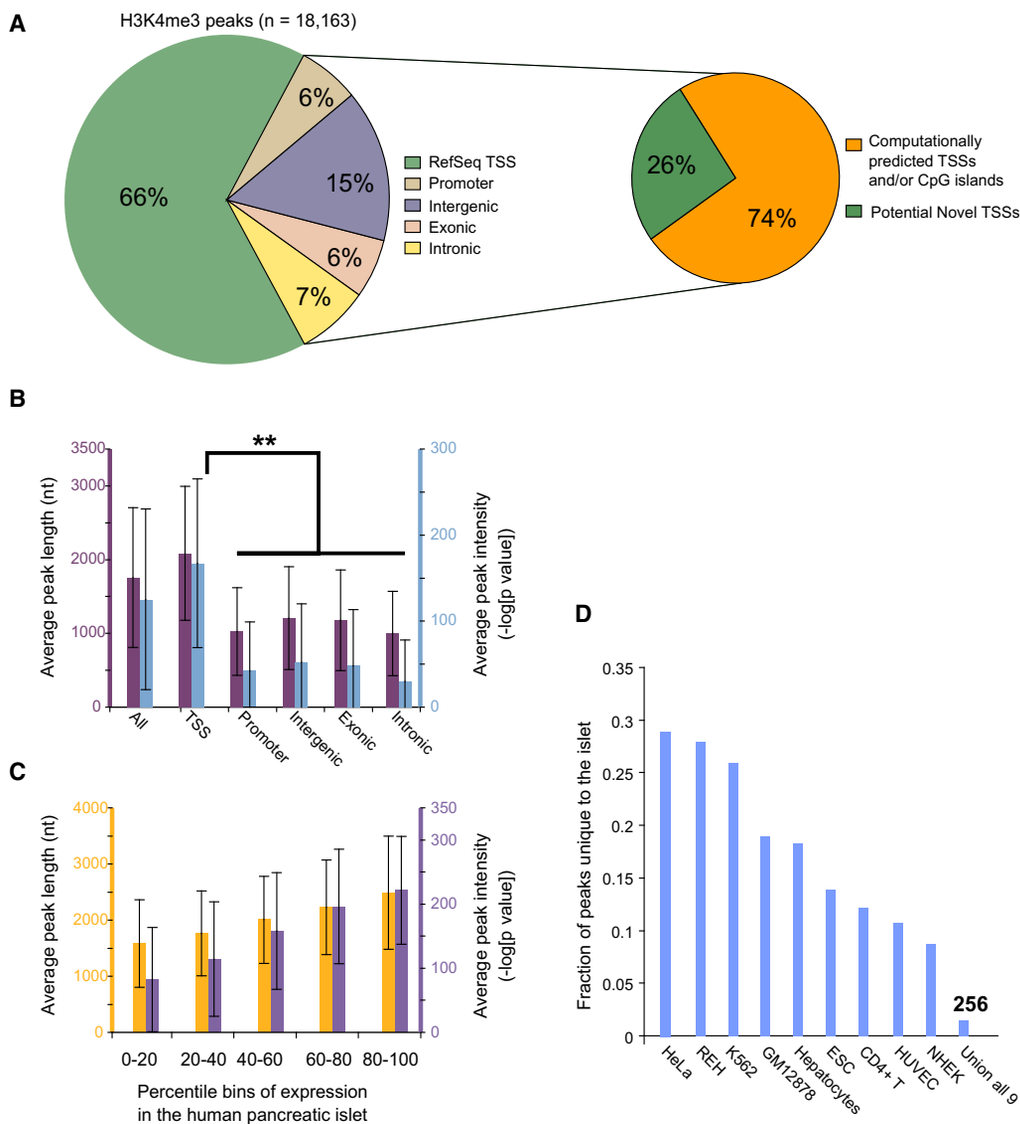
Previous genome-wide profiling studies have reported a positive correlation between the intensity of H3K4me3 signal and gene expression level (Barski et al., 2007; Guenther et al., 2007). To test this observation in islets, we downloaded human islet gene expression data from <http://T1Dbase.org> (Kutlu et al., 2009), partitioned gene expression into quintiles, and computed the average H3K4me3 signal length and intensity at the TSSs of genes within each bin. Although the average H3K4me3 peak length and intensity monotonically increases with gene expression, there is great variability within each expression bin (Figure 2C). Surprisingly, of the 245 most highly islet-expressed genes in this data set, 18% ( $n = 45$ ) have either no or extremely low associated H3K4me3 signal. Notably, 71%

(32/45) also lacked a DHS peak (data not shown). Gene ontology (GO) analysis revealed that these 45 genes are most significantly enriched for the molecular function of hormone activity ( $p = 0.029$  after Bonferroni correction for multiple testing) (Experimental Procedures). These genes include insulin (*INS*), glucagon (*GCG*), islet amyloid polypeptide (*IAPP*), pancreatic polypeptide preprotein (*PPY*), somatostatin (*SST*), and transthyretin (*TTR*). We confirmed by quantitative RT-PCR that *INS*, *GCG*, and *SST* are robustly expressed (Figure S2), so it is unlikely that low H3K4me3 at these TSSs is due to technical artifacts or adverse effects of the islet shipment or handling process. Because these genes are  $<10$  kb in length, we considered the possibility that weak H3K4me3 signal is simply associated with short genes. However, the proportion of short genes ( $<10$  kb in length) within the set of “most highly expressed with no/low H3K4me3 signal” (66.7%, 30/45) is not statistically different from the proportion of short genes within the entire set of most highly expressed (69.8%, 171/245). This result suggests that the transcriptional regulation of islet hormones and other related, highly islet-expressed genes occurs through a distinct mechanism as compared to most other genes.

H3K4me3 ChIP-chip (human embryonic stem cells, hepatocytes, REH cells [Guenther et al., 2007]) or ChIP-seq (human CD4+ T cells [Barski et al., 2007] and GM12878, HUVEC, NHEK, K562, and HeLa cell lines [Broad Institute ChIP-seq, Bernstein lab, ENCODE Project Consortium, 2007]) data are available for nine different human cell types. Comparisons between islet and each other cell type indicated that, on average, 10%–30% of the islet peaks are unique (Figure 2D). Not surprisingly, this value drops to  $\sim 1.5\%$  ( $n = 256$ ) when compared with all nine cell types together. Only 34 of the 256 islet-unique peaks correspond to TSSs of annotated RefSeq genes, and these are enriched for known pancreatic  $\beta$  cell functions such as secretion ( $p = 9.3 \times 10^{-3}$ ) and  $\text{Ca}^{2+}$ -dependent exocytosis ( $p = 6.6 \times 10^{-3}$ ) (Table 1). Furthermore, several of the genes (*SLC30A8*, *GCK*) harbor genetic variants that confer significant risk for T2D and elevated plasma fasting glucose levels (Dupuis et al., 2010; Ingelsson et al., 2010; Prokopenko et al., 2008, 2009). The remaining 222 islet-unique peaks may represent alternative TSSs of genes with function in developing and/or mature islets or TSSs of unannotated coding or noncoding transcription units.

### Identification of Unannotated Islet-Active TSSs

H3K4me3 peaks in unannotated genomic space ( $n = 6190$ ) are TSS candidates. Because H3K4me3 may also be enriched at inactive TSSs (Guenther et al., 2007), we adopted a two-step approach to identify the subset of these 6190 peaks that are likely to be active in the human islet (Figure S3A). First, we developed an algorithm that uses DHS peaks to assign directionality to H3K4me3 peaks (Experimental Procedures). DHS peaks tend to be sharply focused around the TSS, while H3K4me3 peaks are broader and extend well into the body of the transcription unit. We hypothesized that the location of the DHS peak relative to the H3K4me3 peak could predict the directionality of the underlying gene. Using the strongest DHS peak within an H3K4me3 peak, this simple algorithm performed at  $\sim 90\%$  accuracy on annotated RefSeq genes known to be expressed in the human islet (Experimental Procedures). Interestingly, the majority ( $\sim 80\%$ ) of the incorrectly assigned TSSs (based on



**Figure 2. Analysis of Histone 3 Lysine 4 Trimethylation Loci in the Islet Genome**

(A) Distribution of H3K4me3 peaks across five genomic annotation sets as described in Figure 1A. Two-thirds of the peaks span RefSeq transcription start sites (TSSs, left pie chart). Non-RefSeq H3K4me3 peaks are enriched for computationally predicted TSS and/or CpG islands (right pie chart). Additional information is provided in Figure S3.

(B) Average length (purple) and intensity (blue) of H3K4me3 peaks across five genomic annotation sets as described in Figure 1B. The average length and intensity of peaks is significantly higher at TSSs (\*\*, two-tailed paired Student's *t* test,  $p$  value  $< 10^{-100}$ ). Error bars represent SD.

(C) Relationship between average H3K4me3 peak length (yellow)/intensity (purple) and average gene expression level. Error bars represent SD.

(D) Comparison of islet H3K4me3 peaks with peaks from nine different human cell types. Each data point represents the fraction of total peaks ( $n = 18,163$ ) unique to the human islet relative to each of the other nine human cell types or all of them combined (Union of all 9). ~1.5% of the peaks are unique to the islet. Varying levels of similarity across cell types may be at least partially explained by differences in the stage of cellular differentiation and/or sequencing depth.

current annotation) harbored multiple DHS peaks, positioned on either end of the H3K4me3 peak. These H3K4me3 peaks are slightly (~200 nt) longer than those for which the orientation was correctly assigned, increasing the likelihood of overlapping non-TSS-related DHS peaks, which can confound the prediction algorithm. Many of these non-TSS DHS peaks may correspond to CTCF-binding sites that are located on the opposite side of the DHS with respect to the TSS (Boyle et al., 2008) and RNA polymerase (Pol) III-bound loci found in chromatin domains

occupied by Pol II and associated with enhancer-binding factors (Oler et al., 2010). We observe examples of each case in our data set (Figure S4).

Second, we performed ChIP-seq to profile genome-wide histone 3 lysine 79 dimethylation (H3K79me2), which is enriched in actively transcribed regions (Guenther et al., 2007). If the relative density of H3K79me2 reads on either side of an H3K4me3 peak was consistent with its predicted directionality (as determined from the pattern of the DHS and H3K4me3 signal), then

**Table 1. Examples of Islet-Unique H3K4me3 Peaks**

Gene Symbol	Relevance to Islet Biology
GCK	Involved in glucose metabolism
	T2D GWAS locus (Dupuis et al., 2010)
	Harbors an islet-specific promoter (Magnuson, 1990)
SLC30A8	Involved in cation (Zn <sup>2+</sup> ) transport important for insulin secretion (Chimienti et al., 2004)
	T2D GWAS locus (Prokopenko et al., 2008)
	Exhibits islet-specific expression (Chimienti et al., 2004)
REG1A	Derived from regenerating islets (Terazono et al., 1988)
FFAR1	Exhibits islet-specific expression (Bartoov-Shifman et al., 2007)
	Regulates insulin secretion (Itoh et al., 2003)
SYT4	Involved in Ca <sup>2+</sup> -dependent trafficking and exocytosis of secretory vesicles (Tsuboi and Rutter, 2003)
KCNK16	Exhibits pancreas-specific expression (Girard et al., 2001)
ELAVL4	Regulates cell proliferation (Joseph et al., 1998)
UCN3	Regulates glucose-stimulated insulin secretion (Li et al., 2007)
PRSS1	Harbors mutations that underlie hereditary pancreatitis and pancreatic cancer (Teich et al., 1998)

Nine examples among the 34 islet-unique peaks that are at RefSeq transcription start sites (TSSs). The corresponding genes have known pancreatic islet function (such as insulin secretion), and some harbor genetic variants that confer significant risk for type 2 diabetes (*SLC30A8* and *GCK*).

the underlying TSS was classified as islet active. Intragenic TSSs are difficult to assess using this method, because the H3K79me2 signal may be due to transcription from an upstream TSS. Restricting the analysis to intergenic space, we identified 263 candidates for unannotated, islet-active TSSs (Table S4), of which 75% (n = 196) overlap CpG islands and/or computationally predicted TSSs (Figure S3A). These candidates include islet-active TSSs for noncoding RNAs such as the let-7a-1 cluster of microRNAs (Figure 3A) and the miR-1179/miR-7-2 cluster (Figure S3B). We also identified putative alternative TSSs for genes with important islet function such as pancreatic peptidyl-glycine  $\alpha$ -amidating mono-oxygenase (*PAM*), which encodes for an islet secretory granule membrane protein (Figure 3B). Finally, we identified an active promoter locus that is contained within a recently reported T1D-associated region on chromosome 12 (index SNP rs1701704). This promoter could underlie an unannotated transcript or could be an alternative promoter for the downstream gene Ikaros family zinc finger 4 (*IKZF4*) (Figure S3C), which is considered a strong functional candidate for T1D (Hakonarson et al., 2008).

#### Identification of Distal *cis*-Regulatory Elements

Sites bound by the CTCF are an important class of *cis*-regulatory elements that can mediate insulator or other regulatory activities (Phillips and Corces, 2009). To generate a genome-wide CTCF-binding site profile in the human islet, we performed ChIP-seq and designated enriched regions as “CTCF peaks” (n = 21,304) (Table S5 and Experimental Procedures). We assessed the genomic distribution of peaks (Figure 4A), computed the

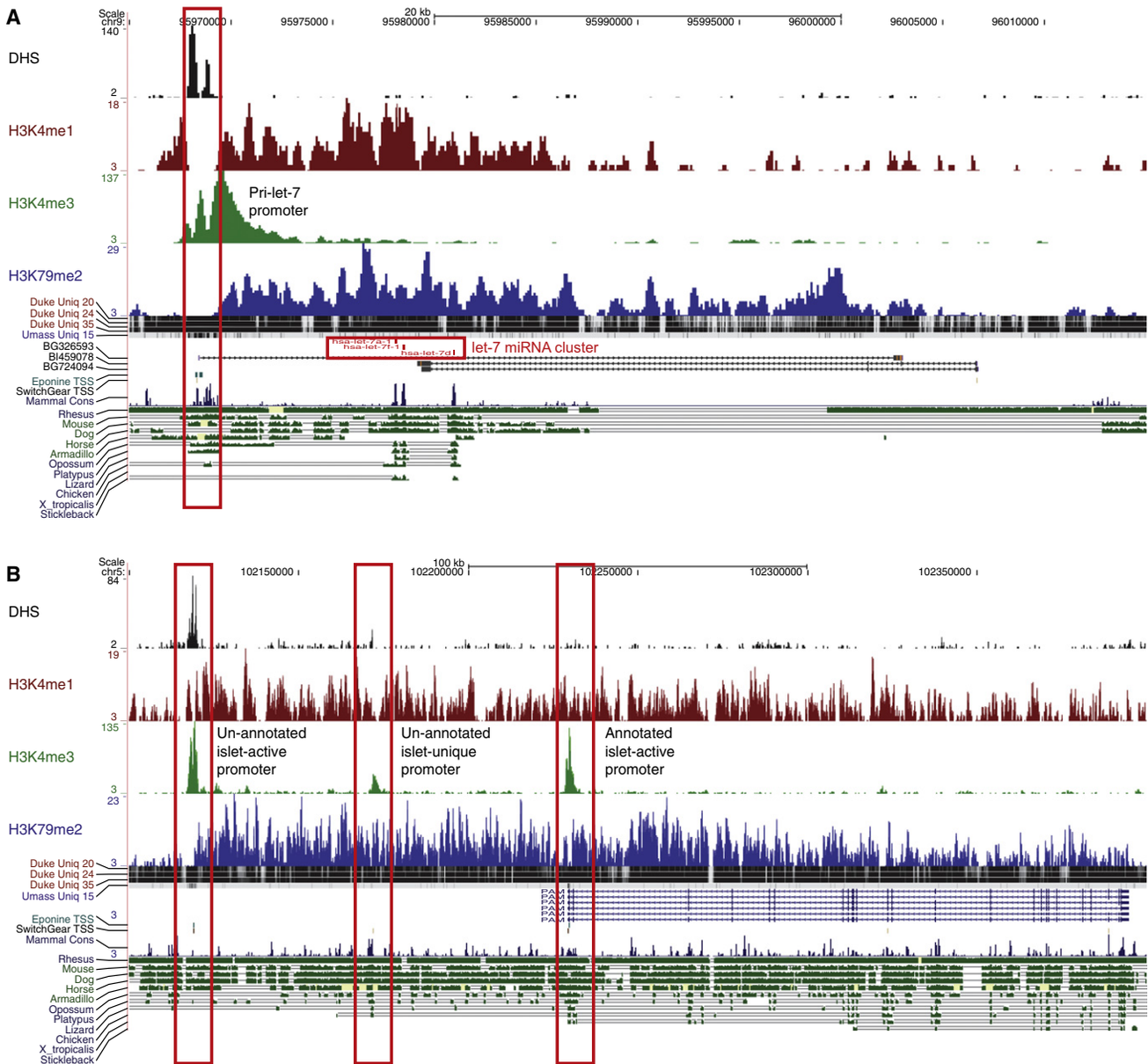
average peak intensity/length across various genomic categories (Figure 4B), and identified the most significantly overrepresented motif within the peaks using MEME (Figure 4C and Supplemental Experimental Procedures). The results corroborate those from previously described studies in other cell types (Kim et al., 2007; Jothi et al., 2008; Cuddapah et al., 2009). Further, only 0.6% (n = 123) of CTCF peaks were islet unique (Figure 4D). Finally, we observed that among the 77% of CTCF peaks that overlap 22% of DHS peaks, the CTCF peaks are positioned near the center of the DHS peak with a slight 5' shift (Figure 4E).

Previous studies have observed depletion of monomethylated histone 3 lysine 4 (H3K4me1) at TSSs and enrichment at putative enhancers such as distal STAT1 and EP300 sites (ENCODE Project Consortium, 2007; Heintzman et al., 2007, 2009; Robertson et al., 2008) and nonpromoter DHS (Barski et al., 2007; Robertson et al., 2008; Wang et al., 2008). To profile H3K4me1 across the human islet genome, we repeated the ChIP-seq strategy described above for three islet samples. We computed the average ratio of the density of extended H3K4me1 sequence reads in DHS peaks at RefSeq TSSs (t-DHS, n = 11,829) and d-DHS peaks (n = 34,039) (Experimental Procedures) to the density in flanking control regions that do not harbor DHS signal (Experimental Procedures). t-DHS peaks are significantly depleted for H3K4me1, whereas d-DHS peaks are significantly enriched (Figure 5). Further, there was no significant difference in H3K4me1 enrichment between CTCF-positive and CTCF-negative d-DHS. Although we detected depletion of H3K4me1 at t-FAIRE peaks, there was no enrichment at d-FAIRE peaks (Figure 5).

We did not detect dramatically different H3K4me1 enrichment levels between intergenic and intragenic d-DHS peaks (Figure S5). Interestingly, although the average H3K4me3 read density in d-DHS peaks was ~3-fold less than that of H3K4me1, d-DHS peaks were still enriched for H3K4me3 signal relative to flanking control regions (Figure S5). These observations are consistent with the previous finding that although H3K4me1 often marks distal regulatory regions, a substantial portion is also associated with H3K4me3 signal (Robertson et al., 2008). Overall, the enrichment of active histone modifications suggests that islet d-DHS peaks are strong candidates for putative regulatory elements. Fifty published index SNPs (<http://www.genome.gov/gwastudies/>) and their linkage disequilibrium partners ( $r^2 > 0.6$ ) for diabetes (T1D, T2D) and related quantitative traits (fasting glucose, fasting insulin) are found within 500 bp of nonpromoter d-DHS peaks (Table S9 and Experimental Procedures), suggesting that these SNPs may contribute to diabetes or altered islet physiology by modulating regulatory element activity.

#### Application of Chromatin Profiles to T2D Susceptibility Loci

To identify regulatory elements and transcripts that may underlie molecular mechanisms of T2D, we analyzed the chromatin profiles in the 18 GWAS-derived genomic loci conferring risk for T2D (Prokopenko et al., 2008). The genomic boundaries of each association signal (Table S6) were defined by the Spotter algorithm (Experimental Procedures). The chromatin profiles do not predict any alternative promoters or unannotated/noncoding



**Figure 3. Identifying Unannotated Islet-Active Transcription Start Sites**

(A) Candidate islet-active TSS for the primary transcript of the ubiquitous let-7a-1/7d/7f-1 microRNA cluster. The TSS (red box; DHS+, H3K4me3+, H3K4me1+) is ~10 kb upstream of the 5'-most microRNA in the cluster, and the full-length primary transcript (H3K79me2+) of ~35 kb matches a known EST (BSG326593). This EST likely represents a noncoding RNA primary transcript from which the let-7 cluster of miRNAs is processed (Marson et al., 2008). The strategy for predicting TSSs is shown in Figure S3A.

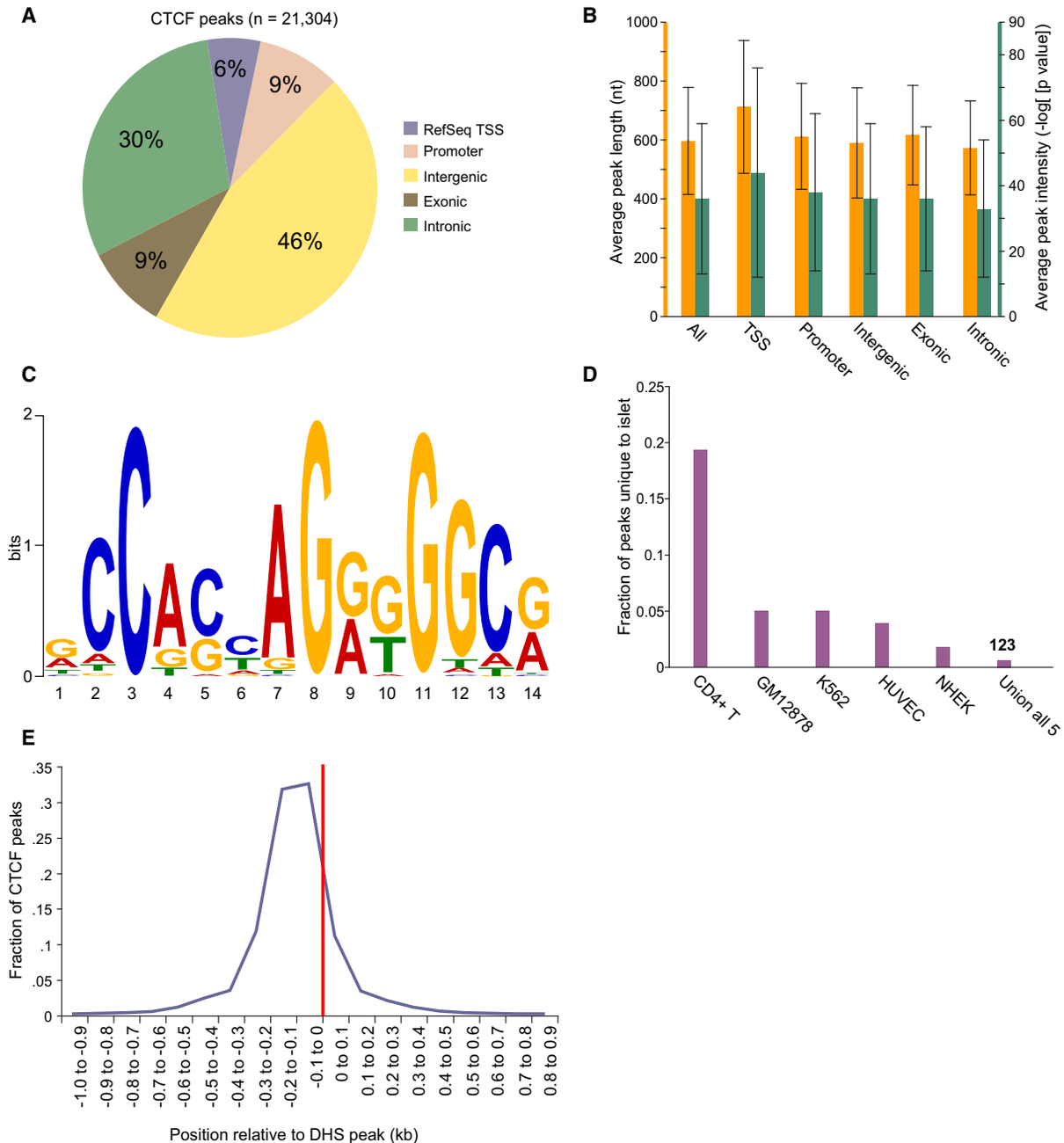
(B) Two candidate islet-active alternative TSSs (red boxes) for the gene *PAM*, which encodes an islet secretory granule membrane protein. One of the candidate TSSs is also islet unique and occurs between the annotated TSS and an unannotated islet-active TSS. Examples of confounding factors for predicting islet-active TSSs are shown in Figure S4.

transcripts in these regions. However, they do identify 118 d-DHS peaks, which represent putative distal regulatory elements (Table S7 and Experimental Procedures). About one-quarter of these elements (n = 28) are bound by CTCF in the islet. Six of the 118 elements contain one or more T2D-associated SNPs (index SNP or SNP with  $r^2 > 0.6$ ) (Table S8). These six include a previously identified element containing the index SNP rs7903146 in the *TCF7L2* locus (Gaulton et al., 2010). The

remaining five map to the *IGF2BP2*, *KCNQ1*, *WFS1*, *FTO*, and *CDC123/CAMK1D* loci. Only the *CDC123/CAMK1D* element is bound by CTCF in the islet.

**Validation of Putative Islet Regulatory Elements in T2D Loci**

To determine whether predicted regulatory elements in the islet can function as enhancers, we cloned two classes of elements



**Figure 4. Profiling of Binding Sites for the CCCTC-Binding Factor**

(A) Distribution of CTCF peaks across five genomic annotation sets as described in Figure 1A.

(B) Average length (orange) and intensity (green) of CTCF peaks across five genomic annotation sets is fairly uniform. Error bars represent SD.

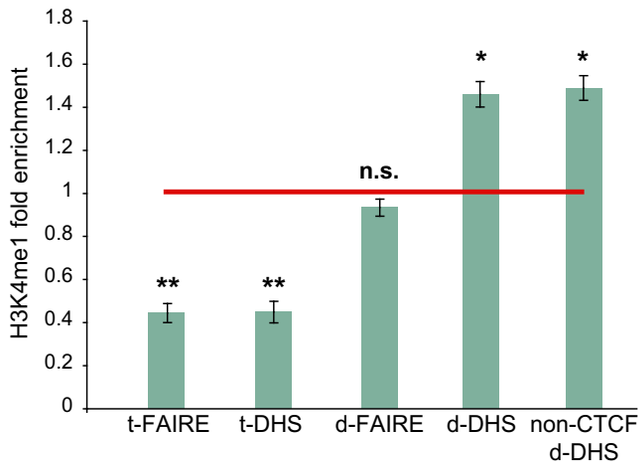
(C) Motif determined by MEME (Bailey and Elkan, 1994) using the top 10% of CTCF peaks.

(D) Comparison of islet CTCF peaks with peaks from five different cell types. Each data point represents the fraction of total peaks (n = 21,304) unique to the human islet relative to each of the other five human cell types or all of them combined (Union of all 5). Less than 1% of the peaks are unique to the islet (n = 123). Varying levels of similarity across cell types may be at least partially explained by differences in the stage of cellular differentiation and/or sequencing depth.

(E) Positioning of CTCF peaks relative to the center of overlapping DHS peaks (red line). Almost all CTCF peaks that overlap DHS peaks are within 200 bp of the DHS peak center.

containing d-DHS peaks into luciferase reporter vectors (Figure 6): those bound by CTCF (“C,” n = 11) and those that are not (“P,” n = 33). We also cloned a number of non-DHS, non-CTCF controls (“N,” n = 15). Because human islet cell lines

are not available, we tested these elements for enhancer activity in murine pancreatic MIN6 (Figure 6A) and HeLa (Figure 6B) cell lines. Only ~15% (4/26) of the negative controls exhibited enhancer activity in any orientation or cell type (~9% [1/11] of



**Figure 5. Representation Analysis of Histone H3 Lysine 4 Monomethylation in Candidate Regulatory Regions**

DNase I-hypersensitive site (DHS) and formaldehyde-assisted isolation of regulatory elements (FAIRE) peaks at RefSeq TSSs (t-DHS and t-FAIRE, respectively) are significantly depleted for H3K4me1 signal (\*\*, two-tailed paired Student's t test,  $p < 0.005$ ), and DHS peaks at distal candidate regulatory elements (d-DHS) are enriched for H3K4me1 signal (\*, two-tailed paired Student's t test,  $p < 0.01$ ). Error bars represent SD among three islet samples. FAIRE data were obtained from Gaulton et al. (2010). Representation analysis of additional histone modifications is shown in Figure S5.

"C" elements and 20% [3/15] of "N" elements) (Figures 6A and 6B). In contrast,  $\sim 2.5$ -fold more "P" elements demonstrated enhancer activity (12/33). This positive rate (36.4%) is comparable to that of predicted HeLa enhancers (Heintzman et al., 2009) that exhibited increased luciferase activity in our HeLa reporter assays (38.5%, 5/13).

Four of 12 "P" elements exhibiting enhancer activity (P4, *KCNJ11/ABCC8*; P12, *TCF7L2*; P17, *WFS1*; P20, *HHEX/IDE*) are unique to the islet; one of these (P17, *WFS1*) is also undetected by at least three other methods for the prediction of regulatory element potential: PReMod (Ferretti et al., 2007), phastCons (Siepel et al., 2005), and islet-FAIRE (Gaulton et al., 2010). The average H3K4me1 enrichment among the 12 d-DHS peaks in the elements exhibiting enhancer activity was similar to that computed for all d-DHS ( $\sim 1.3$ -fold) (Figure 6C). However, there was large variation in H3K4me1 enrichment among individual elements (0.6- to 3.4-fold), with only 3/12 enriched above baseline (1.0) (Figure 6C).

#### Allele-Specific Analysis of Five Regulatory Elements Containing T2D-Associated SNPs

Five "P" elements tested contain T2D-associated SNPs (P9, *IGF2BP2*; P12, *TCF7L2*; P17, *WFS1*; P21, *KCNQ1*; P23, *FTO*) (Figures 6A and 6B). Notably, four out of the five elements (all except P9) exhibited enhancer activity in at least one orientation and cell type tested. To assess allele- or haplotype-specific effect(s) of T2D-associated variants on enhancer activity, we cloned these four regions from the genomic DNA of individuals with risk and nonrisk genotypes/haplotypes and compared luciferase reporter activity (Figures 6D and S6A). We confirmed significantly stronger enhancer activity for the *TCF7L2* element

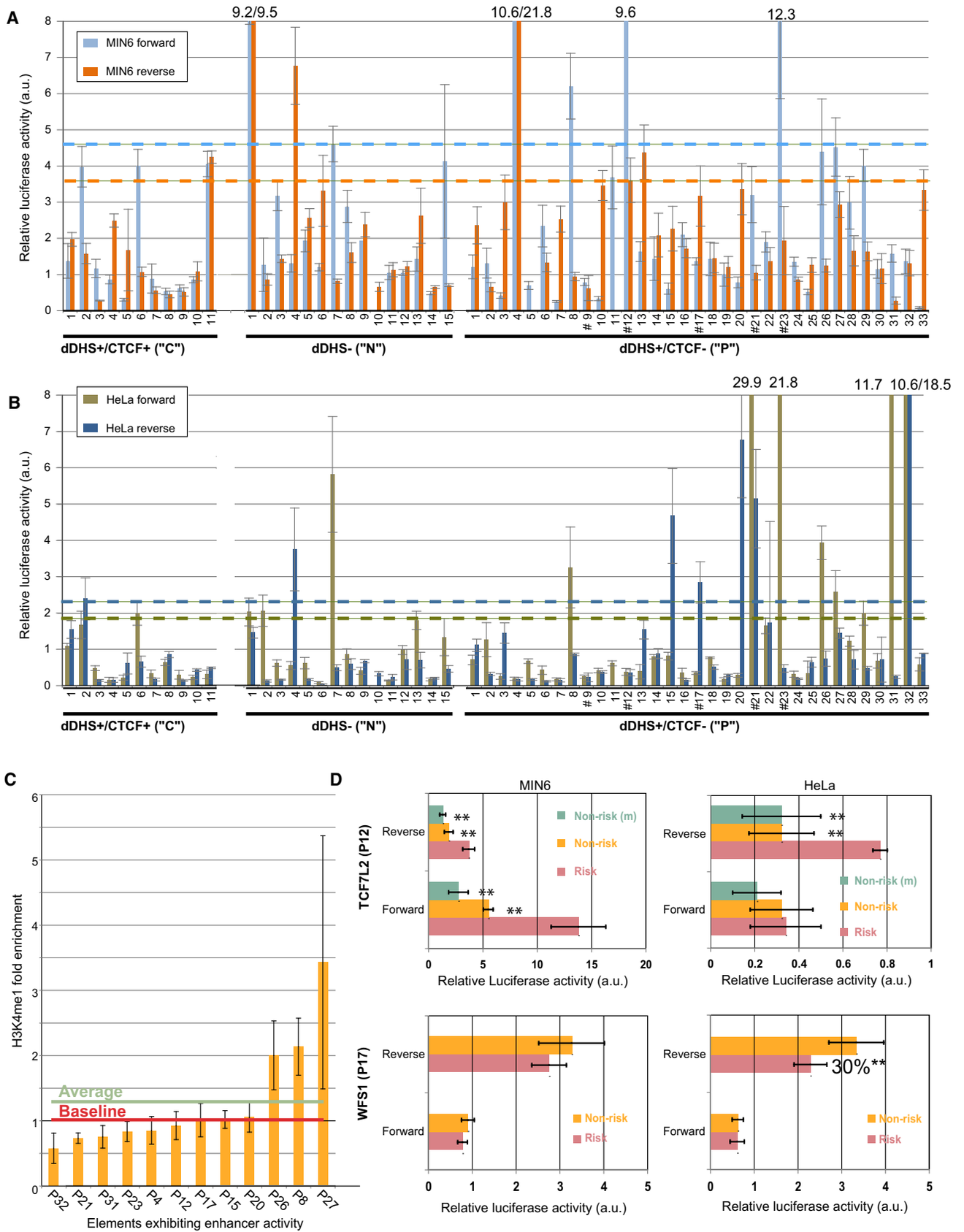
(P12) containing the rs7903146 risk allele relative to the nonrisk allele ( $\sim 3$ -fold) (Figure 6D) (Gaulton et al., 2010). *TCF7L2* allelic enhancer effects were specific to the MIN6 cell line (Figure 6D, compare MIN6 and HeLa). Sequencing of the *TCF7L2* inserts from each haplotype revealed two variant bases, a novel variant (C/G at Chr10:114,747,977; hg18) and rs7903146; only rs7903146 mediated allele-specific effects on enhancer activity (Figure 6D, compare Risk to Nonrisk and Nonrisk(m)) (Figure S6B). We also identified a haplotypic effect on enhancer activity for the *WFS1* element (P17), which contains four SNPs (rs4689397, rs6823148, rs881796, and rs4234731). The risk haplotype exhibited  $\sim 30\%$  lower activity than nonrisk in HeLa cells (Figure 6D).

#### DISCUSSION

In this study, we describe the most comprehensive characterization to date of the epigenomic profile of unstimulated human pancreatic islets. Using DNase- and ChIP-seq techniques, we profiled open chromatin, CTCF-binding sites, H3K4me3, H3K4me1, and H3K79me2 across the entire genome in human islets. Integrated analysis of these large-scale data sets identified  $\sim 18,000$  putative TSSs,  $\sim 30\%$  of which were previously unannotated by RefSeq. Further computational genomic analyses revealed that at least several hundred of these are islet-active TSSs, including those for major islet miRNAs previously implicated in the control of glucose homeostasis (Lynn, 2009). Interestingly, active chromatin marks (H3K4me3, DHS, H3K79me2) were absent from a subset of highly islet-expressed genes, including those encoding islet-specific hormones (*INS*, *GCG*, *SST*, *IAPP*, *PPY*, and *TTR*). This observation suggests that some genes critical for islet function have an unconventional promoter chromatin signature, indicative of a unique transcriptional control mechanism. Mutskov and Felsenfeld (2009) have proposed such a model based on detailed analysis of the *INS* locus in human islets.

We also identified  $\sim 34,000$  candidate distal regulatory elements in human islets. A substantial number of these putative elements were clustered ( $< 1000$  bp from each other). Comparisons with other cell types indicated that these clustered elements are significantly enriched for islet-unique sites and thus may represent islet-specific regulatory modules worthy of more extensive future investigation. Based on CTCF-binding profiles,  $\sim 22\%$  of the  $\sim 34,000$  candidate distal regulatory elements are predicted insulator sites. Previous studies have reported that the H3K4me1 signal is enriched in distal regulatory elements (Heintzman et al., 2007, 2009). Though our analyses confirm this finding in aggregate, we show that H3K4me1 enrichment may not be a reliable predictor of regulatory activity for individual elements.

Fifty SNPs associated with islet-related diseases and traits map to within 500 bp of a candidate nonpromoter regulatory element. Focusing on T2D, 4 of the 12 elements that function as enhancers in vitro (*FTO*, *KCNQ1*, *TCF7L2*, and *WFS1* loci) harbor T2D-associated SNPs, including two (*TCF7L2* and *WFS1* loci) that exhibit significant allele-specific differences in activity. These results suggest that altered enhancer activity plays a role in the molecular mechanism underlying at least a subset of T2D genetic association signals.



These data sets should provide functional context for noncoding variants identified through additional association, targeted resequencing, or whole-genome sequencing studies. Further analysis of the repertoire of regulatory elements in the human islet will enhance the understanding of gene regulation in the islet and should offer additional insight into the molecular mechanisms that underlie diabetes susceptibility.

## EXPERIMENTAL PROCEDURES

### Human Islets

Fresh human pancreatic islets were obtained from the ICR Basic Science Islet Distribution Program and National Disease Research Interchange (NDRI). Islet viability and purity were assessed by the distribution centers and are shown along with phenotypic/clinical information of each donor in Table S10. Islets were warmed to 37°C and washed with calcium- and magnesium-free Dulbecco's phosphate-buffered saline (Invitrogen; Carlsbad, CA) prior to crosslinking. For chromatin immunoprecipitation (ChIP) studies, cells were crosslinked for 20 min in 1% formaldehyde at room temperature, frozen in liquid nitrogen, and stored at -80°C.

### DNase-Seq and DHS Peak Identification

For DNase-seq experiments, fresh pancreatic islets were disaggregated to achieve single-cell suspension. Islets were washed with prewarmed 1X PBS once and resuspended with dissociation solution (1 ml of 1X PBS, 50  $\mu$ l of 0.05  $\mu$ l Dispase I stock solution [Roche; Indianapolis, IN]). Islet suspension was transferred to a 6-well culture dish, incubated at 37°C for 30 min, dissociated with a 2 ml sterile pipette, and incubated for another 30 min. This incubation-agitation cycle was repeated 4 or 5 times until >90% of islets were disaggregated into single cells. Cells were washed with prewarmed 1X PBS once and prepared for DNase-seq experiments as previously described (Song and Crawford, 2010). Libraries from three primary human islet samples (Table S10) were sequenced using the Illumina GAI platform. Peaks were identified using MACS (Supplemental Experimental Procedures) (Zhang et al., 2008).

### ChIP and Illumina GAI Sequencing

ChIP assays were carried out as previously described (Scacheri et al., 2006), with the following modifications. Intact nuclei were isolated and chromatin was sheared on ice using a Branson 450 Sonifier (constant duty cycle, output 4, 12–16 cycles of 20 s sonication with 1 min rest between cycles) to a size of 200–1000 bp. Antibodies used for ChIP were anti-H3K4me3 (ab8580, Abcam; Cambridge, MA), anti-H3K4me1 (ab8895, Abcam), anti-H3K79me2 (ab3594, Abcam), anti-CTCF (ab70303, Abcam; 07-729, Millipore; Danvers, MA), and anti-GFP (sc-8334, Santa Cruz Biotechnology; Santa Cruz, CA).

Islet ChIP-seq libraries were prepared and sequenced using the Illumina GAI protocol and platform. The number of sequencing lanes, clusters, aligned reads, repeat-filtered reads (no satellite reads), and unique starts is shown for each islet and ChIP experiment in Table S12. MACS (Zhang et al., 2008) was used to call H3K4me3 and CTCF peaks (Supplemental Experimental Procedures).

### Genome-wide Analysis of Chromatin Marks

Perl and R scripts were written to perform the genomic characterization and comparative analysis of DHS, H3K4me3, and CTCF peaks. Unless otherwise noted, functional annotation data sets (including RefSeq and UCSC known genes, predicted TSSs and bidirectional promoters, phastCons elements, CpG islands, and ChIP-seq data sets) were downloaded from the UCSC Table Browser on November 1, 2009 (<http://genome.ucsc.edu/cgi-bin/hgTables>).

For "computationally predicted TSSs," both the Eponine and the Switchgear data sets from the UCSC Table Browser were utilized. Human pancreatic islet gene expression data were downloaded from T1DBase (<http://T1DBase.org>), and expression data for other tissues were downloaded from BioGPS Human U133A/GNF1H Gene Atlas (<http://biogps.gnf.org/downloads/>). Islet-selective gene expression was defined as at least 3-fold greater expression in the islet relative to any other tissue represented. Genome-wide results of the Chai algorithm were determined according to the parameters in Parker et al. (2009), and islet-FAIRE data sets were obtained from Gaulton et al. (2010). GO analyses were performed using the web-based tool NIH DAVID 6.7 (<http://david.abcc.ncifcrf.gov/>). For the DHS peak clustering analysis (Figure 1F) and the histone modification enrichment/depletion analysis (Figures 5 and S5), we stringently defined d-DHS peaks as those that are not within H3K4me3 peaks and  $\geq 5$  kb away from RefSeq TSSs, UCSC Known Gene TSSs, Eponine or Switchgear computationally predicted TSSs, and CpG islands, yielding 34,039 d-DHS. To select regulatory elements to test for enhancer activity (Figure 6), the definition of d-DHS was slightly loosened ( $\geq 5$  kb upstream and  $\geq 1$  kb downstream from known and predicted TSSs and CpG islands). P values for statistical comparisons were computed using either the two-tailed paired Student's t test or the Fisher's exact test. Details of the remaining computational analyses are described in Supplemental Experimental Procedures.

### Molecular Cloning

Putative regulatory elements were amplified from human genomic DNA with primers designed using PrimerTile (<http://research.nhgri.nih.gov/tools/>). Element boundaries were determined by manual H3K4me1 profile inspection. Coordinates of amplified elements and primer sequences for amplification are found in Table S13. Putative regulatory elements were cloned using the Gateway system (Invitrogen). Generation of Gateway-compatible vectors is described in Supplemental Experimental Procedures. Variants of interest were introduced using QuikChange Lightning (Stratagene; La Jolla, CA). Mutagenesis primer sequences are available upon request. Mutagenesis was confirmed by direct sequencing.

### Transfection and Dual Luciferase Assays

Cells were seeded in 96-well plates (40,000 cells/well HeLa, 60,000 cells/well MIN6) and cotransfected with 0.072 pmol Gateway-modified firefly (pGL 4.23, Promega; Madison, WI) and 2 ng Renilla (pRL-TK, Promega) vectors using Lipofectamine 2000 (Invitrogen). Two vector preparations per insert orientation were tested. Transfections were performed in triplicate.

Cells were lysed in 1  $\times$  passive lysis buffer (Promega) 36–48 hr posttransfection, and dual luciferase assays were run on a Centro/Centro XS3 Microplate Luminometer LB 960 (Berthold; Bad Wildbad, Germany). Firefly values were normalized to Renilla to control for differences in cell number or transfection efficiency. Luciferase assays were performed in triplicate. For each element tested, at least two independent vector preparations were used. Activity was

## Figure 6. Luciferase Reporter Activity Validates Putative Enhancer Elements

(A) Relative luciferase activity of constructs in three element classes tested in MIN6 cells. Genomic locations of elements are found in Table S13. Blue and orange dashed lines indicate 2.33 standard deviations ( $p = 0.01$ ) (Heintzman et al., 2009) above the median activity of tested CTCF-bound regions for elements cloned in the forward or reverse orientations, respectively. Data represent the mean  $\pm$ SD of three replicates each for two separate clones (six total measurements). C, d-DHS+/CTCF+ element; N, d-DHS-/CTCF-; P, d-DHS+/CTCF- element. # marks elements containing T2D-associated SNPs. Numbers above the bars indicate the luciferase activity for elements beyond the scale of the y axis; a.u. denotes arbitrary units.

(B) Relative luciferase activity of constructs in three element classes tested in HeLa cells. Data are analyzed and annotated as in (A).

(C) H3K4me1 representation in the 12 elements exhibiting enhancer activity. Though the overall average enrichment of H3K4me1 is  $\sim 1.3$ -fold (green line), only 3/12 elements are above baseline (red line). Error bars represent SD among three islet samples.

(D) Relative luciferase activity of *TCF7L2* (P12) and *WFS1* (P17) elements in MIN6 (left panels) or HeLa (right panels) cells containing the risk or nonrisk alleles of T2D-associated SNPs. For *TCF7L2*, (m) denotes a mutation generated by site-directed mutagenesis from the risk to nonrisk allele. Data represent the mean  $\pm$ SD of three replicates each from at least two independent clones. \*\*, two-tailed unpaired Student's t test,  $p < 0.01$ . Additional allelic analysis is shown in Figure S6.

defined as 2.33 standard deviations (SD) ( $p = 0.01$ ) above the median activity of negative controls (Heintzman et al., 2009), defined as CTCF-bound elements in this study.

#### ACCESSION NUMBERS

The NCBI Gene Expression Omnibus (GEO) umbrella accession number, which links to the individual ChIP-seq and DNase-seq data sets, is GSE23784.

#### SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and 13 tables and can be found with this article online at doi: 10.1016/j.cmet.2010.09.012.

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