ATRX, DAXX or MEN1 mutant pancreatic neuroendocrine tumors are a distinct alpha-cell signature subgroup

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The commonly mutated genes in pancreatic neuroendocrine tumors (PanNETs) are ATRX, DAXX, and MEN1. We genotyped 64 PanNETs and found 58% carry ATRX, DAXX, and MEN1 mutations (A-D-M mutant PanNETs) and this correlates with a worse clinical outcome than tumors carrying the wild-type alleles of all three genes (A-D-M WT PanNETs). We performed RNA sequencing and DNA-methylation analysis to reveal two distinct subgroups with one consisting entirely of A-D-M mutant PanNETs. Two genes differentiating A-D-M mutant from A-D-M WT PanNETs were high ARX and low PDX1 gene expression with PDX1 promoter hyper-methylation in the A-D-M mutant PanNETs. Moreover, A-D-M mutant PanNETs had a gene expression signature related to that of alpha-cells (FDR q-value < 0.009) of pancreatic islets including increased expression of HNF1A and its transcriptional target genes. This gene expression profile suggests that A-D-M mutant PanNETs originate from or transdifferentiate into a distinct cell type similar to alpha cells.

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Pancreatic neuroendocrine tumors (PanNETs) or islet cell tumors are a relatively rare neuroendocrine malignancy with an annual incidence of <1 per 100,000 per year (about 1000 new cases per year in the United States) but currently represent the second most common epithelial neoplasm after ductal adenocarcinoma of the pancreas and account for 1–2% of pancreatic tumors. PanNETs were erroneously considered a benign group of neoplasm because they were initially mostly comprised of benign symptomatic insulin-producing tumors (insulinomas). However, in the past three decades, it has become apparent that at least half of all PanNETs are nonfunctional, and they are a heterogeneous group of tumors with often unpredictable and varying degrees of malignancy. As many as 50–80% of PanNETs are associated with synchronous or metachronous metastatic disease. Knowledge of functional PanNETs has evolved from insulinoma to almost a dozen other diverse hormone-secreting tumors. These individual lesions may have specific clinical, pathological, and genetic associations, including multiple endocrine neoplasia type 1 (MEN-1), tuberous sclerosis, and von Hippel-Lindau (VHL) syndromes. Thus, the entity of PanNET represents a diverse group of heterogeneous neoplasms where combined clinical and pathologic assessment is required to further identify their genetic basis for neoplasia and to define their specific clinical behavior. The nonfunctional tumors require further elucidation to characterize their diverse pathogenesis and to predict outcome with potential biomarkers and molecular signatures. Current classification scheme for PanNETs include grade and stage. The World Health Organization (WHO) classification, which assesses the proliferative index of neoplastic cells, divides PanNETs into low grade (G1), intermediate grade (G2), and high grade (G3). The higher grade PanNETs are generally associated clinically with more aggressive behavior. Poorly differentiated neuroendocrine tumor of the pancreas is extremely rare and clinically aggressive, which represents a different pathogenesis from the well-differentiated counterpart. While well-differentiated PanNETs can be successfully treated with surgery, there are few treatments for metastatic PanNETs, and they do not respond to conventional chemotherapy. A greater understanding of PanNET pathogenesis may guide the development of novel therapeutic options.

Molecular studies have identified mutations in MEN1, ATRX, and DAXX to be the most commonly found in PanNETs (found in approximately 40, 10, and 20% of tumors, respectively). All three genes play a role in chromatin remodeling, MEN1 is a component of a histone methyltransferase complex that specifically methylate Lysine 4 of histone H3 and functions as a transcriptional regulator. ATRX and DAXX interact to deposit histone H3.3-containing nucleosomes in centromeric and telomeric regions of the genome. Additional mutations in mTOR pathway genes including TSC2, PTEN, and PIK3CA are found in one in six well-differentiated PanNETs. Other reported rare mutations in PanNETs include DNA damage repair genes (MUTYH, CHEK2, BRCA2) and chromatin remodeling gene SETD2.

The neuroendocrine cells in the pancreas include alpha, beta, delta, pancreatic polypeptide (pp)-producing and vasoactive intestinal peptide (VIP)-producing cells. The cell of origin for nonfunctional PanNETs is not well established. Here, we genotyped 64 well differentiated PanNETs for mutations in ATRX, DAXX, and MEN1 and performed RNA sequencing (n = 33) and DNA methylation (n = 32) analysis to identify distinct molecular phenotypes of A-D-M mutant PanNETs which potentially reveals their distinct cell of origin or transdifferentiated state.

Results

Clinical annotation and genotyping for ATRX, DAXX, and MEN1. We initially performed Sanger sequencing to genotype the ATRX, DAXX, and MEN1 genes in 64 individual PanNETs. All cases were histologically confirmed to be well-differentiated PanNETs of WHO G1/G2 grade, and cases of poorly differentiated neuroendocrine carcinoma were excluded. The mean patient age was 52 ± 1.5 years (mean ± standard error, ranging from 26–73) with a 59% male population. The locations of the tumors were 38% proximal/mid body, and 62% distal pancreas. Eighty-one percent of the cases were clinically non-functional and the remaining cases included insulinomas, glucagonomas, gastrinomas, and VIPomas. The median size of tumor was 3.6 ± 0.4 cm (ranging from 1.0–14.5 cm). Sixty-eight percent of patients had localized disease without distant metastasis at the time of initial diagnosis (Supplementary data 1).

An A-D-M mutant genotype was identified in 58% (37/64) of cases with ATRX, DAXX, MEN1, MEN1/ATRX, and MEN1/DAXX mutations in 8, 16, 20, 3, and 11% cases, respectively (Fig. 1b). The majority of mutations in ATRX, DAXX, and MEN1 were truncation mutations (stopgain or frameshift) and loss of function consistent with their role as tumor suppressors (Supplementary Data 2). Similar to the observations in our previously published data, the 5-year disease specific survival was associated with tumor stage (p-value < 0.04, log-rank tests), tumor grade (G1 vs G2 p-value < 0.02, log-rank tests), and distant metastasis (p-value < 0.002, log-rank tests), respectively. Among 44 patients who initially presented with localized disease without distant metastasis, those with the A-D-M mutant genotype had a worse recurrence free survival than those of A-D-M WT genotype (Fig. 1b). Furthermore, in comparison to A-D-M WT PanNETs, the A-D-M mutant PanNETs were associated with larger tumor size (3.6 ± 0.6 cm vs. 5.6 ± 0.7 cm, p-value < 0.03, log-rank tests) and higher tumor stage (T1 and T2 vs. T3, p-value < 0.04, log-rank tests). Other demographic and clinical characteristics (including gender, age, tumor functionality, and lymph node metastasis) revealed no statistically significant differences between the two genotypes of PanNETs.

Gene expression and DNA methylation reveal two subtypes of PanNETs. We performed RNA sequencing on 33 randomly selected tumors (19 A-D-M mutant, and 14 A-D-M WT). Unsupervised hierarchical clustering of the top 3000 variable genes across the PanNETs revealed two distinct clusters where almost all A-D-M mutant PanNETs were found in one cluster (Fig. 2a). The grouping of A-D-M mutant PanNETs into one distinct cluster by gene expression was robust to the number of most variable genes used for clustering (Supplementary Figure 1). Principal component analysis (PCA) separated the A-D-M mutant PanNETs from the A-D-M WT PanNETs along the first principal component (corresponding to the component comprising the largest variation in gene expression) (Fig. 2b). The separation of A-D-M mutant PanNETs from A-D-M WT PanNETs by PCA was robust to the number of top variable genes used (Supplementary Figure 2). These data show that A-D-M mutant tumors have a distinct gene expression pattern from that of A-D-M WT PanNETs. Neither hierarchical clustering nor PCA from gene expression revealed further subgrouping of the tumors with single mutations in ATRX, DAXX, or MEN1 or double mutations in ATRX/MEN1 or DAXX/MEN1.

In hierarchical clustering, the A-D-M mutant PanNETs formed a tighter cluster than the A-D-M WT PanNETs. In PCA, the A-D-M mutant PanNETs had smaller variance along PC1 than A-D-M WT PanNETs. Pair-wise correlation of gene expression between all PanNETs, showed a higher correlation among A-D-M mutant
PanNETs as compared to A-D-M WT PanNETs (Fig. 2c). Among A-D-M mutant PanNETs, mutants with the same genotype (mutations in ATRX/DAXX/MEN1) did not show greater gene expression correlation. These data suggest that A-D-M mutant PanNETs are a more homogeneous group compared to A-D-M WT PanNETs.

Within A-D-M mutant or A-D-M WT PanNETs groups, unsupervised clustering and PCA did not reveal differences between primary and metastatic tumors. Top 100 genes with highest variance across all samples separates mutant from A-D-M WT PanNETs and showed relatively high expression of “liver-specific” genes (APOH, ALDH1A1, FGB, APOC3 etc.) as well as complement and coagulation pathway genes (SERPINA1, FGA, F10, CP, MT3 etc.) in A-D-M mutant PanNETs (Fig. 2d; Supplementary Figure 3), both in primary (collected in absence of liver tissue) and metastatic tumors. Moreover, the pathological estimate of tumor purity was over 80% for all samples of PanNETs consistent with inference from ESTIMATE10 (median tumor purity of 90%, Supplementary data 3) showing high tumor purity characteristic of well differentiated PanNETs. In addition, seven A-D-M mutants and one A-D-M WT PanNETs were from the tissue of liver metastases and they had gene expression profile most similar to the genotype group of their primary PanNET counterpart (Fig. 2d). We confirmed the distinct gene expression signature of A-D-M mutant PanNETs in a larger tumor set (47 PanNETs including the 33 PanNETs where RNA sequencing was performed) using gene expression microarray technology. The 14 additional samples are comprised of 3 A-D-M WT PanNETs and 11 A-D-M mutant PanNETs (7 MEN1 mutant, 2 DAXX mutant, and 2 DAXX/MEN1 mutant)(Supplementary Figure 4).

To investigate epigenetic differences between PanNETs, we used the Illumina 450K chip to assay the DNA methylation at 411,549 CpG sites in 32 PanNETs. Unsupervised hierarchical clustering of the top 2000 variable DNA methylation sites across the PanNETs revealed two distinct clusters where almost all A-D-M mutant PanNETs were found in one cluster (Fig. 3a). Principal component analysis (PCA) separated the A-D-M mutant PanNETs from the A-D-M WT PanNETs along the first principal component (corresponding to the component comprising the largest variation in DNA methylation) (Fig. 3b). The separation of A-D-M mutant PanNETs from A-D-M WT PanNETs by PCA was robust to the number of top variable DNA methylation sites used (Supplementary Figure 5). These data reveal that A-D-M mutant PanNETs have a distinct DNA methylation pattern from that of A-D-M WT PanNETs. Neither hierarchical clustering nor PCA revealed differences in DNA methylation sites between the different combinations of genes mutated among the A-D-M mutant PanNETs. Within A-D-M mutant or A-D-M WT PanNETs groups, unsupervised clustering and PCA of DNA methylation did not reveal differences between primary and metastatic tumors.

To investigate the global histone methylation level in PanNETs with and without A-D-M mutations, we performed immunohistochemistry on H3K4me3, H3K9me3, H3K27me3, and H3K36me3 on 36 PanNETs. There was a general trend of lower histone methylation level for MEN1 mutant PanNETs when compared to WT PanNETs (Supplementary Figure 6 and Supplementary Table 1). A-D-M mutant PanNET gene expression resembles that of alpha cells. There are multiple neuroendocrine cell types in the pancreas including alpha, beta, gamma, delta, and epsilon. We used gene expression data for these various pancreatic neuroendocrine and exocrine cell types from a single cell sequencing study11 (Supplementary Table 2) to identify gene-set signatures representing highly expressed cell-type-specific genes (Supplementary data 4). The A-D-M mutant PanNETs uniformly exhibited a gene expression signature that was very similar to that of alpha cells (Fig. 4a). The A-D-M WT PanNETs were more heterogeneous in their expression of the genes among the gene set signatures for the different pancreatic neuroendocrine cell types. Greater heterogeneity of gene expression signature in A-D-M WT PanNETs was consistent with the greater heterogeneity found in global gene expression.

To further investigate the gene expression signature of A-D-M mutant PanNETs we performed gene set enrichment analysis12 (GSEA) on the 13 manually curated gene sets for pancreatic endocrine and exocrine cells from a previous study. This study assessed gene expression of individual pancreatic cell
types (alpha, beta, delta, PP, acinar, ductal, mesenchyme, and endothelial) enriched by flow cytometry and using single cell RNAseq (Supplementary Table 2). Our analysis indicates that only the alpha cell gene signature was significantly enriched in A-D-M mutant PanNETs (FDR q-value < 0.009) (Fig. 4b, c) (Supplementary Table 3).

Alpha and beta cell lineage specific genes were examined for the A-D-M mutant and WT PanNETs. ARX, IRX2, and TMSF4 were all highly expressed in A-D-M mutant PanNETs compared to A-D-M WT PanNETs (Supplementary Figure 7). Surprisingly, GCG (glucagon) expression was lower in A-D-M mutants as compared A-D-M WT PanNETs. For beta cell specific genes,
PDX1, MAFA, INS, and DLK1, all had lower expression in A-D-M mutant PanNETs than A-D-M WT PanNETs (Supplementary Figure 7). However, these genes had much greater expression heterogeneity in A-D-M WT PanNETs suggesting that some A-D-M WT PanNETs resemble beta cells and others did not (Supplementary Figure 7).

Validation of subtype and alpha cell signature in A-D-M mutant PanNETs. We derived an A-D-M mutant gene expression signature from significant differentially expressed genes between the A-D-M mutant and WT PanNETs from our data set \((n = 33)\). We used two independent panNET\(^6,13\) data sets to validate the gene expression signature of A-D-M mutant panNETs. We obtained a gene expression dataset from International Cancer Genome Consortium Pancreatic Endocrine Neoplasm (ICGC PAEN) \((n = 29)\)\(^6\) and Sadanandam et al. \((n = 75)\)\(^13\). The A-D-M mutant and WT PanNETs from both data sets have significant positive and negative correlations with our A-D-M mutant PanNET signature respectively (Fig. 5a) (Supplementary data 5). Additionally, we found alpha cell signatures to be significantly enriched \((FDR q < 0.001)\) only in the A-D-M mutant PanNETs from the two validation data sets using GSEA (Fig. 5b, c).

**HNF1A pathway is up-regulated in A-D-M mutant PanNETs and alpha cells.** HNF1A is one of the most significantly differentially expressed genes between A-D-M mutant and WT PanNETs. HNF1A is a homeobox family transcription factor that is highly expressed in the liver and is involved in the regulation of several liver-specific genes. The expression of HNF1A was 2.93 fold higher in A-D-M mutant PanNETs than A-D-M WT PanNETs (corrected \(p\)-value < 0.004, Benjamini–Hochberg) (Fig. 6a). Differentially expressed genes (DEgenes) between the A-D-M mutant and A-D-M WT PanNETs were found in 1478 genes (with greater than 3 fold change and corrected \(p\)-value < 0.05 Benjamini–Hochberg, see Methods section) (Supplementary data 6). Functional pathway enrichment for DEgenes using pre-ranked GSEA revealed the complement and coagulation cascades, retinol metabolism, and drug metabolism to be up-regulated in A-D-M mutant PanNETs (see Methods section) (Fig. 6b; Supplementary data 6). The differentially expressed genes were also enriched for HNF1A transcription factor motifs in their promoters \((FDR < 0.001, \text{Fig. 6c})\). The complete list of significant TF motifs is presented in Supplementary data 6. Taken together, the A-D-M mutant PanNETs had higher expression of HNF1A along with many of its transcriptional target genes associated with liver function. In addition, the transcriptional regulator of HNF1A, HNF4A\(^14\) was expressed 3.02 fold higher in A-D-M mutant

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**Fig. 4** A-D-M mutant PanNETs with alpha-cell signature. **a** Heatmap of gene expression for top 20 alpha and beta cell-specific genes from Muraro et al.\(^11\) revealed alpha cell specific genes are highly expressed in A-D-M mutant panNETs. A-D-M WT panNETs are more heterogeneous in gene expression but some show high beta cell specific gene expression. Red color represents higher correlation with alpha cell specific genes. **b** Gene set enrichment analysis show A-D-M mutant PanNETs to be enriched for expression of alpha cell specific genes. Pancreas cell type (alpha, beta, delta, PP, acinar, ductal) gene signatures were obtained from three different published dataset to access enrichment of cell type signatures in A-D-M mutant vs A-D-M WT PanNETs. Table represents GSEA results where size is the number of genes in gene set. All alpha cell gene sets (from three different sources) are significantly enriched in A-D-M mutant panNETs (highlighted in red). No other cell types were in A-D-M mutant or A-D-M WT panNETs. **c** GSEA plots of significant alpha cell signatures from Brasmig et al.\(^15\), Wang et al.\(^32\), and Muraro et al.\(^11\).
PanNETs ($p$-value $< 0.009$, DeSeq2). We used gene expression data from Bramswig et al.\textsuperscript{15} to show $HNF1A$ expression was increased in alpha cells compared to beta cells ($p$-value $< 0.008$), and the 465 alpha cell specific genes in the pancreas were enriched for transcriptional targets of $HNF1A$ and for having $HNF1A$ TF motif in their promoters (See Methods section; Supplementary Table 4).

Many of the most differentially expressed genes and highly expressed in A-D-M mutant PanNETs are targets of $HNF1A$ and are involved in protein secretion, transport and metabolism ($APOH$, $ALB$, $AFM$, $HAO1$, $UGT1A3$, $UGT1A1$, $GC$, $G6PC$, $TM4SF4$, $PKLR$ etc). $APOH$ is expressed 8.46 fold higher in A-D-M mutant PanNETs ($p$-value $< 10^{-5}$) and is potentially a good diagnostic biomarker for A-D-M mutant PanNETs. Moreover, $APOH$ has been shown to have high expression in only the alpha cells of the pancreatic islet in a single cell RNA sequencing study\textsuperscript{16}. We perform IHC staining for $APOH$ and show positive staining in 70 ± 2.5% of A-D-M mutant and only 18 ± 2.0% of A-D-M WT PanNETs (Supplementary Figure 8).

**PDX1 gene is hypermethylated with low expression in A-D-M mutant PanNETs.** There is no genome wide hypo or hypermethylation of DNA in A-D-M mutant or WT panNETs. DNA methylation differences between the A-D-M mutant and A-D-M WT PanNETs were found at 378 CpG sites (corrected $p$-value $< 0.05$ and difference in beta value $> 0.2$, Benjamini–Hochberg), 287 of which were found in genes and 91 in intergenic regions (Supplementary data 7). Of the 287 differentially methylated genic CpG sites, 70 (associated with 59 genes) were found at promoter (transcriptional start site, TSS1500 and TSS200) or within first exon, a region where DNA methylation is associated with transcriptional repression\textsuperscript{17}. Thirteen of the 59 genes were also found to be differentially expressed (with fold change greater than 3 and corrected $p$-value $< 0.05$, Benjamini–Hochberg) and seven genes that were hypomethylated in A-D-M mutant and over-expressed are $APOH$, $CCL15$, $EMID2$, $PDZK1$, $HAO1$, $BAAP21$, and $NPC11$. One gene, $TACR3$, was hypomethylated in A-D-M WT and over-expressed (Supplementary data 7). Four of the 70 CpG sites were found in the gene $PDX1$ (pancreatic and duodenal homeobox 1), a transcription factor necessary for pancreatic development and beta cell maturation. $PDX1$ functions in the cell fating of endocrine cells, favoring the production of insulin positive beta cells and somatostatin positive delta cells while repressing glucagon positive alpha cells\textsuperscript{18}. These four CpG sites were all hypermethylated in A-D-M mutant PanNETs (Fig. 7a) and the expression of $PDX1$ was 2.92 fold higher in A-D-M WT PanNETs ($p$-value $< 0.005$, DeSeq2) (Fig. 7a, b). In contrast, while ARX was highly expressed in A-D-M mutant PanNETs compared to A-D-M WT PanNETs, the promoter and first exon of ARX are not differentially methylated.

**Discussion**

Similar to a number of recent studies\textsuperscript{19,20}, we have demonstrated in this cohort of PanNETs that, in addition to pathologic stage and grade of the tumor, mutations in $DAXX$, $ATRX$, and $MEN1$ are associated with adverse clinical outcome in comparison to those without these mutations. Our results seem to be in contradiction to the findings initially reported by Jiao et al.\textsuperscript{5}, in that 15 patients with PanNETs carrying mutations in $DAXX$ or $ATRX$ genes had better survival than did 12 patients with wild-type PanNET. This discrepancy between our data and their data could be attributed to a different composition of the tumors. Indeed, all
the tumors analyzed in Jiao’s study were liver metastases from PanNETs as opposed to only 19% (12 out of 64) in our study. Other factors including sample size and length of follow up time may also contribute to discrepancies between different studies.

Here, we found that A-D-M mutant PanNETs form a distinct subgroup on the basis of their gene expression profile and DNA methylation pattern. Moreover, this subgroup is more homogeneous based on gene expression profile than the A-D-M WT PanNETs. The gene signature of the A-D-M mutant PanNETs strongly corresponds to the genes that are specifically expressed in alpha cells including genes known to define alpha cells such as ARX and “liver-specific” genes such as HNF1A and its transcriptional targets. Conversely, PDX1, a gene critical to the beta cell lineage is transcriptionally repressed in A-D-M mutant PanNETs and the PDX1 promoter is hypermethylated. On the other hand, WT PanNETs have heterogeneous gene expression profiles and their gene mutational landscape is less understood.

The pancreas is comprised of many different cell types including acinar, ductal, and at least five neuroendocrine cell types including alpha, beta, gamma, delta, and epsilon cells. There are two plausible explanations for the “alpha cell-like” expression pattern of A-D-M mutant PanNETs. Either an alpha cell or an uncharacterized cell type with an alpha-cell like gene expression profile is the cell-of-origin for PanNETs with mutations in ATRX, DAXX, and MEN, or loss of ATRX, DAXX, or MEN genes may promote pancreatic neuroendocrine (or progenitor) cell types to
MEG3 has been shown to upregulate expression of long noncoding RNA oncogenic pathways activated in A-D-M mutant PanNETs. The restricted deletion of MEG3 or ATRX/DAXX mutations occur alone or in a combined pattern suggest that they have independent oncogenic activities in A-D-M mutant PanNETs, making the idea of reprogramming to a homogeneous alpha-like cell state less probable. Some of the A-D-M WT PanNETs have a strong beta cell signature and these may have arisen from beta cells (Fig. 4a). However, other A-D-M WT PanNETs have neither alpha nor beta cell signatures, which may arise from other cell types in the pancreas.

Conditional knockouts of MEN1 in mice support the model of an alpha cell origin for A-D-M PanNETs.21 The restricted deletion of MEN1 to alpha cells surprisingly led to the development of insulinoma.22,23 Most of our PanNETs were nonfunctional (26 of 33 PanNETs) but the functional tumors were insulinoma and VIPoma, even though their gene expressions have alpha cell signature. Moreover, some PanNETs express combinations of neuroendocrine hormones (GGC, INS, SST, PPY, GHRL, VIP, and GAST), suggesting that regulation of cell type specific hormone may be disrupted. To create robust gene signatures that are not sensitive to changes in expression of a few genes, we use a large number of genes to create the A-D-M mutant and alpha cell signatures. In other mouse models of PanNETs21,24,25, MEN1 deletion using the insulin or PDX1 promoter driven Cre construct, insulinomas, glucagon-expressing tumors and well differentiated PanNETs were also observed. However, Cre expression may be leaky in these models and further study is needed to understand the heterogeneity of the cells in the tumors that develop and trace the cell of origin or transdifferentiated state of the cancer cells.

In our gene expression analysis, we have not identified the oncogenic pathways activated in A-D-M mutant PanNETs. MEN1 has been shown to upregulate expression of long noncoding RNA MEG3 in MIN6 mouse insulinoma cell line26. In the same study, they show MEG3 represses expression of the oncogene MET leading to delayed cell cycle progression and reduced cell proliferation. In a different study, MEN1 and DAXX were shown to repress the expression of the membrane metalloendopeptidase (MME) and mutations in MEN1 or DAXX result in loss of this repression leading to neuroendocrine tumor proliferation27. Our data is consistent with these studies when comparing A-D-M mutant to WT PanNETs, showing that A-D-M mutant PanNETs have lower expression of MEG3 (7.3 fold lower, p-value < 4.3E-07, DeSeq2), higher expression of MET (3 fold higher, p-value < 0.003, DeSeq2), and higher expression of MME (4 fold higher, p-value < 0.001, DeSeq2). Among A-D-M mutant PanNETs, we do not see expression differences of MEG3, MET, and MME depending on mutation status of ATRX, DAXX, and MEN1.

While PanNETs may seemingly represent as a single clinical disease, they can be further characterized into different subtypes based on their cell lineage and the associated molecular genotype. Understanding the epigenetic and transcriptional dysregulation of PanNETs will require comparison to their proper cells of origin which may explain the unpredictable outcome of the disease and facilitate the development of unique and targeted therapeutic strategies.

Methods

Patient’s information. Retrospective and prospective reviews of well-differentiated pancreatic neuroendocrine neoplasms were performed using the pathology files and pancreatic database at MSKCC with IRB approval. All patients were evaluated clinically at our institution with confirmed pathologic diagnoses, appropriate radiological and laboratory studies, and surgical or oncological management. Follow-up information was obtained for all cases.

Tissue acquisition and nucleotide extraction. Briefly, cases of pancreatic neuroendocrine neoplasms were identified. Fresh-frozen tumor and paired normal tissues were obtained from MSKCC’s tissue bank under an Institutional Review Board protocol. Histopathology of all tissues was evaluated on hematoxylin and eosin stained sections by an experienced gastrointestinal-hepato-pancreatobiliary pathologist to insure the nature of the tissue, greater than 80% tumorcellularity and absence of necrosis. The relevant tissues were then macro-dissected (20–25 mm), and DNA extractions were performed using the Qiagen’s DNeasy Blood and Tissue Kit and RNeasy Mini Kit, respectively was carried out according to the manufacturer’s protocols (Qiagen, Valencia, CA).

Sanger sequencing for gene mutation. All exons of the DAXX, ATRX, and MEN1 genes were amplified by PCR and then sequenced using Sanger sequencing. Every mutation detected was validated by bidirectional Sanger sequencing on the tumor DNA. Sequence data were deposited in the NCBI database (Supplementary Table 1).

PanNETs transcriptome sequencing and data analysis. RNA Library preparation and RNA sequencing was done by MSKCC Genomics Core Laboratory using Illumina HiSeq with (2 × 75 bp paired end reads) to a minimum depth of ~50 million reads were generated for each sample. Raw fastq files were probe for sequencing quality control using FastQC [http://www.bioinformatics.babraham.ac.uk/projects/fastqc]. Sequencing reads were mapped to human transcripts corresponding to Genepattern28 genome (hg19 version) GTF annotations using RSEM with default parameters. RSEM package29 was used to prepare the reference genome with given GTF and calculated expression from mapped BAM files. STAR30 aligner was used to map reads in RSEM algorithm. Transcripts mapped data were normalized to TPM (Transcript Per Million) from RSEM and log2 transformed (Supplementary data 8). This log2TPM values were used for all downstream analysis. Unsupervised clustering and Principal Component Analysis was conducted to elucidate subtypes structure using top 3000 variant genes as input. To query robustness of this subtyping, multiple variant gene sets were used and repeated the same process of unsupervised clustering. Top 100 variable genes were used to find genes, which were highly expressed in each subtype. Subset of these genes is selected to show in Fig. 2d for liver and complement system genes. To find differentially expressed genes (DEgenes) between A-D-M mutant PanNETs and A-D-M WT PanNETs, we used DeSeq2 R package31 on raw count (values from RSEM). We use significance cutoff with greater than 3 fold change and corrected p-value < 0.05 (Benjamini–Hochberg) to call a gene as DEgenes. GSEA Preranked32 method was used on DEgenes to find significant KEGG pathways, motif and biological process.

Clustering and principal component analysis. For unsupervised clustering on log2TPM, we used Pearson distance metric and ward.D2 hclust method (unless stated otherwise). PCA analysis was done using procarm in R. R (http://www.r-project.org/) was used for all the analysis and visualization of data.

PENGset from published dataset. The neuroendocrine cells in the pancreas include alpha, beta, delta, pancreatic polypeptide (pp)-producing and vasoactive intestinal peptide (VIP)-producing cells. Gene sets representing different endocrine cells (GSE97038) and pancreatic exocrine and endocrine cells (PENGset) were obtained from three meta-data1,13,32 (Supplementary Table 2). We created 13 PENGset representing all major cell from endocrine and exocrine pancreases. Supplementary Table 2 shows these gene sets with major cell types and number of genes in each set. These gene set were used as prior defined gene set for GSEA analysis.

Gene set enrichment analysis on major inlets cell types. Gene Set Enrichment Analysis (GSEA) was performed on the log2TPM expression values of all samples using downloaded version of GSEA software (Broad Institute, Cambridge, MA, USA) to identify the statistically enriched gene sets between A-D-M mutant and A-D-M WT PanNETs. Published pancreatic islet endocrine and exocrine cells signatures were used as prior defined sets as an input. We used all default parameters to perform GSEA on this gene sets to determine the enrichment of specific cell signatures between in the PanNET subtypes. We ran GSEA on 1000 permutation mode on phenotypic label to generate FDR and enrichment score (ES) for each gene set. Significant gene set was filtered based on FDR q-values (cutoff of 0.05).
Bramswig et al. FACs sorted alpha and beta cells gene expression. We extensively used Bramswig et al. FACs sorted RNAseq data to understand normal alpha and beta cells and correlated their gene signature sets with our A-D-M mutant and A-D-M WT PanNETs. We downloaded supplement file for total RNA seq normalized expression data for alpha (3 replicate) and beta (3 replicate) and exocrine cells (2 replicate). Bramswig et al. provide strong genes associated with alpha, beta and exocrine cells as supplement file. We used this strong cell specific genes and created a new set for alpha, beta and exocrine and named as Bramswig et al. gene set. HNF1A gene expression values were fetched to check whether HNF1A is over expression in normal alpha as compared to beta and exocrine. We applied Student t-test’s between three alpha and three beta samples to calculate p-value for HNF1A gene expression. Bramswig et al. strong alpha cell genes (n = 463) were queried to check for HNF1A transcription factor motif enriched using online GSEA version (C3 TFs motif database).

Microarray gene expression analysis. RNA extracted from PanNETs samples were submitted for Affymetrix microarray profiling using chip hgu333a2 array. All following analysis was done in R. Briefly, the raw Affymetrix CEL files were loaded in R (simplely7). Normalized expression values were calculated by the GC Robust Multi-array Average (GCRMA) algorithm and subjected to mean transformation in order to collapse all probes to respective genes in R using collapseRow8. We then followed the same clustering and PCA procedure that were done on RNAseq expression data. Collapse averaged gene expression values were imported in GSEA to run against 13 PEEGset cell types to find enrichment.

450 K DNA methylation array analysis. DNA extracted from PanNETs samples and interrogated using the Illumina 450 K platform (Illumina Inc. San Diego, CA) to access the DNA methylation profiles. All the analysis was performed using ChAMP version 2.6.0 open source software implemented in R, Briefly, IDAT raw data were imported in R and filtered to exclude samples with detection p-value < 0.01 (n = 32) and keeping 75% of samples and normalized using FunctionNormalization9. This normalization method correct for background; remove dye bias followed by Quantile normalization. Unsupervised clustering and PCA were done on top variants 2000 probes (Var2000) across all samples to find classes of PanNETs. We repeated this clustering using different number (Var10000, Var5000, Var3000, Var1000, and Var500) of probes to check robustness of this subtyping. Differentially methylated CpG sites (DMP) between the A-D-M mutant and A-D-M WT PanNETs were identified using chipMVP using all the default parameter method (Bonferroni-Hochberg) to adjust the p-value(<0.01, DeSeq2). Significant DMP sites from respective genes were compared to DEGesets to find overlapping deregulated genes in each subtype.

A-D-M mutant PanNET Signature and validation. Significant differentially expressed genes (fold change ≥3 and Corrected Pval <0.05, Benjamini–Hochberg) between A-D-M mutant and WT panNETs were used to log2 transformation of fold changes to create an A-D-M mutant PanNETs signature for validation. We downloaded gene expression and genotype data from two independent PanNETs cohort: (i) Pancreatic Cancer Endocrine Tumors and ICGC site and (b) Sadanandam et al. IGCC PAEN processed A-D-M mutation status and RNAseq gene expression dataset (FPKM normalized expression) were downloaded from ICGC website (http://icgc.org/icgc/cpp/68/304/1003406) for 29 samples (16 A-D-M Mutant and 13 A-D-M WT). Sadanandam et al. performed targeted sequencing of ATRX, DAXX, MEN1, PTEN, TSC2 and ATM on 75 PanNETs. Genotype information was downloaded from supplementary data of PMID: 28446169 and matched microarray gene expression data for 75 PanNETs (28 A-D-M and 47 A-D-M WT) were downloaded from NCBI GEO (Accession Number GSE73338). Normalized gene expression values obtained from GSE73338 were used for the enrichment analysis. Pearson correlation was calculated between our A-D-M mutant gene signature and the mean-variance normalized gene expression from IGCC PAEN and Sadanandam et al. and pval was calculated using Wilcoxon test. We performed GSEA analysis on A-D-M mutant and WT panNETs from Sadanandam et al.18

Immunohistochemistry (IHC). A representative, formalin-fixed, paraffin-embedded tissue section (4 μm thick) of each case was submitted to our institution’s core facility to perform immunohistochemistry using antibodies recognizing the APOH proteins. Briefly, sections were de-paraffinized and pre-treated in Cell Conditioning 1 (CCL mid; Ventana Medical Systems, AZ, USA) using an automated staining system (Ventana Discovery XT Autostainer; Ventana Medical Systems Inc., Tucson, AZ). Primary antibodies were applied for 60 min at a dilution of 1:100 for APOH (anti-APOH, polyclonal antibody; Proteintech). The sections were then incubated for 60 min with secondary antibody (1:200) followed by DAB Map detection (DAB visualization; Ventana Medical Systems). Cytoplasmic (APOH) labeling in at least 50% of the tumor cells was considered positive. In the case of APOH, normal liver tissue was used as a positive control in each experiment.

Histone marks IHC. Serial unstained slides (4 μm) were prepared from each block for subsequent immunohistochemistry with the following Histone 3 lysine antibodies conves: H3K4me3 clone C42D8 (Cell Signaling Technologies Catalog number 9751 1:1000 dilution), H3K9me3 clone EPR66101 (Abcam, Catalog number 9733 1:100 dilution) and H3K36me3 clone 036B11 (Cell Signaling, Catalog number 9733 1:100 dilution) and H3K36me3m, clone 333 (Active motif, Catalog number 61021, 1:500 dilution). Staining for all clones was performed on the Leica Bond immunohistochemistry platform according to the manufacturer’s protocol with the Leica DAB IHC detection kit. All slides were pretreated with epitope retrieval 2 solution (Leica Biosystems) for 30 min. Primary antibodies were incubated for 30 min. Multi-tissue normal positive control was used. The PanNET cases (n = 36, 14 A-D-M mutant and 22 A-D-M WT) were read and interpreted by an independent observer blinded to the clinicopathologic information. Scoring of all histomarks was performed using previously validated scoring systems H3K4me3, H3K9me3, H3K27me3, H3K36me3. The tumor was considered positive for the histone mark if there was histological evidence of nuclear staining. Every tumor was scored on a scale of 0–3 according to the percentage of cells with nuclear staining: 0, 0–5% positive cells; 1, 5–25% positive cells; 2, 25–75% positive cells; 3, 75–100% positive cells). Scores of 0–1 were estimated as low expression and scores of 2–3 indicated high expression. Student t-test was used to test significance for histone methylation across MEN1 panNETs as compared to MEN1 WT PanNETs.

Statistical analysis. Data are represented as mean ± standard deviation. GraphPad Prism 6 (GraphPad Software Inc, La Jolla, CA) was used for statistical and survival analyses. Survival analysis p-values (2-sided) were based on log-rank tests. Significance was defined as P<0.05.

Data availability. The authors declare that all data supporting the findings of this study are available within the article and its Supplementary data and figures. Data generated in this study were deposited to NCBI under GEO SuperSeries GSE117853 (GSE118014 for RNAseq, GSE117851 for Microarray and GSE117852 for 450K methylation). We do not impose any restrictions on any data availability.

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References
1. Hallforsdorsson, T. R., Rabe, K. G., Rubin, J. & Petersen, G. M. Pancreatic neuroendocrine tumors (PNETs): incidence, prognosis and recent trend toward improved survival. Ann. Oncol. 19, 1727–1733 (2008).
2. Tang, L. H. & Klimstra, D. S. Conundrums and caveats in neuroendocrine tumors of the pancreas. Surg. Pathol. Clin. 4, 589–624 (2011).
3. Tang, L. H., Basturk, O., Sue, J. J. & Klimstra, D. S. A practical approach to the classification of WHO Grade 3 (G3) Well-differentiated Neuroendocrine Tumor (WD-NET) and Poorly Differentiated Neuroendocrine Carcinoma (PD-NEC) of the pancreas. Am. J. Surg. Pathol. 40, 1192–1202 (2016).
4. Tang, L. H. et al. Well-differentiated neuroendocrine tumors with a morphologically apparent high-grade component: a pathway distinct from poorly differentiated neuroendocrine carcinomas. Clin. Cancer Res. 22, 1011–1017 (2016).
5. Jiao, Y. et al. DAXX/ATRX, MEN1, and mTOR pathway genes are frequently mutated in typical and atypical carcinoid tumors. Am. J. Surg. Pathol. 41, 1346–1353 (2017).
6. Hughes, C. M. et al. Menin associates with a trithorax family histone methyltransferase complex and with the hox/c locus. Mol. Cell 13, 587–597 (2004).
7. Lewis, P. W., Elsasser, S. J., Noh, K. M., Stadler, S. C. & Allis, C. D. Daxx is an H3-3-specific histone chaperone and cooperates with ATRX in replication-independent chromatin assembly at telomeres. Proc. Natl Acad. Sci. USA 107, 14075–14080 (2010).
8. Ferrone, C. R. et al. Determining prognosis in patients with pancreatic endocrine neoplasms: can the WHO classification system be simplified?. J. Clin. Oncol. 25, 5609–5615 (2007).
9. Yoshihara, K. et al. Infering tumour purity and stromal and immune cell admixture from expression data. Nat. Commun. 4, 2612 (2013).
10. Muraro, M. J. et al. Single-cell transcriptome atlas of the human pancreas. Cell 173, 335–347 (2018).
11. Subramanian, A. et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc. Natl Acad. Sci. USA 102, 15545–15550 (2005).
12. Sadanandam, A. et al. A cross-species analysis in pancreatic neuroendocrine tumors reveals molecular subtypes with distinctive clinical, metastatic, developmental, and metabolic characteristics. Cancer Discov. 5, 1296–1313 (2015).
14. Li, J., Ning, G. & Duncan, S. A. Mammalian hepatocyte differentiation requires the transcription factor HNF-4alpha. *Genes Dev.* **14**, 464–474 (2000).
15. Bramswig, N. C. et al. Epigenomic plasticity enables human pancreatic alpha to beta cell reprogramming. *J. Clin. Invest.* **123**, 1275–1284 (2013).
16. Baron, M. et al. A single-cell transcriptomic map of the human and mouse pancreas reveals inter- and intra-cell population structure. *Cell Syst.* **3**, 346–360.e4 (2016).
17. Bresc, F. et al. DNA methylation of the first exon is tightly linked to transcriptional silencing. *PLos One* **6**, e14524 (2011).
18. Mansouri, A. Development and regeneration in the endocrine pancreas. *ISRN Endocrinol.* **2012**, 640956 (2012).
19. Marini, I. et al. Loss of DAXX and ATRX are associated with chromosome instability and reduced survival of patients with pancreatic neuroendocrine tumors. *Gastroenterology* **146**, 453–460.e5 (2014).
20. Park, J. K. et al. DAXX/ATRX and MEN1 genes are strong prognostic markers in pancreatic neuroendocrine tumors. *Oncotarget* **8**, 49796–49806 (2017).
21. Shen, H. C. et al. Recapitulation of pancreatic neuroendocrine tumors in human multiple endocrine neoplasia type 1 syndrome via Pdx1-directed inactivation of Men1. *Cancer Res.* **69**, 1858–1866 (2009).
22. Shen, H. C. et al. Multiple endocrine neoplasia type 1 deletion in pancreatic alpha-cells leads to development of insulinomas in mice. *Endocrinology* **151**, 4024–4030 (2010).
23. Lu, J. et al. Alpha cell-specific Men1 ablation triggers the transdifferentiation of glucagon-expressing cells and insulinoma development. *Gastroenterology* **138**, 1954–1965 (2010).
24. Bertolino, P. et al. Pancreatic beta-cell-specific ablation of the multiple endocrine neoplasia type 1 (MEN1) gene causes full penetrance of insulinoma development in mice. *Cancer Res.* **63**, 4836–4841 (2003).
25. Li, F. et al. Conditional deletion of Men1 in the pancreatic beta-cell leads to glucagon-expressing tumor development. *Endocrinology* **156**, 48–57 (2015).
26. Modali, S. D., Parekh, V. I., Kebebew, E. & Agarwal, S. K. Epigenetic regulation of the IncRNA MEG3 and its target c-MET in pancreatic neuroendocrine tumors. *Mol. Endocrinol.* **29**, 224–237 (2015).
27. Feng, Z. et al. Menin and Daxx interact to suppress neuroendocrine tumors through epigenetic control of the membrane metallo-endopeptidase. *Cancer Res.* **77**, 401–411 (2017).
28. Reich, M. et al. GenePattern 2.0. *Nat. Genet.* **38**, 500–501 (2006).
29. Li, B. & Dewey, C. N. RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. *BMC Bioinformatics* **12**, 323 (2011).
30. Debin, A. et al. STAR: ultrafast universal RNA-seq aligner. *Bioinformatics* **29**, 15–21 (2013).
31. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol.* **15**, 550 (2014).
32. Wang, Y. J. et al. Single-cell transcriptomics of the human endocrine pancreas. *Diabetes* **65**, 3028–3038 (2016).
33. Wilson, C. L. & Miller, C. J. Simpleaffy: a BioConductor package for Affymetrix Quality Control and data analysis. *Bioinformatics* **21**, 3683–3685 (2005).
34. Gharabihi, R. Z., Fodor, A. A. & Gibas, C. J. Background correction using dinucleotide affinities improves the performance of GCRA. *BMC Bioinformatics* **9**, 452 (2008).
35. Miller, J. A. et al. Strategies for aggregating gene expression data: the collapseRows R function. *BMC Bioinform.* **12**, 322 (2011).
36. Morris, T. J. et al. ChAMP: 450k chip analysis methylation pipeline. *Bioinformatics* **30**, 428–430 (2014).
37. Fortin, J. P. et al. Functional normalization of 450k methylation array data improves replication in large cancer studies. *Genome Biol.* **15**, 503 (2014).
38. Wang, Z. J. et al. Decreased histone H2B monoubiquitination in malignant gastric carcinoma. *World J. Gastroenterol.* **19**, 8099–8107 (2013).
39. Noguchi, A. et al. SIRT1 expression is associated with good prognosis for head and neck squamous cell carcinoma patients. *Oncol. Surg. Oral. Med. Oral. Pathol. Oral. Radiol.* **115**, 385–392 (2013).
40. Wei, Y. et al. Loss of trimethylation at lysine 27 of histone H3 is a predictor of poor outcome in breast, ovarian, and pancreatic cancers. *Mol. Carcinog.* **47**, 701–706 (2008).
41. Ho, T. H. et al. Loss of histone H3 lysine 36 trimethylation is associated with an increased risk of renal cell carcinoma specific death. *Mod. Pathol.* **29**, 34–42 (2016).

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**Additional information**

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