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# Genomic and Epigenomic Landscaping Defines New Therapeutic Targets for Adenosquamous Carcinoma of the Pancreas

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#### Abstract

Adenosquamous cancer of the pancreas (ASCP) is a subtype of pancreatic cancer that has a worse prognosis and greater metastatic potential than the more common pancreatic ductal adenocarcinoma (PDAC) subtype. In order to distinguish the genomic landscape of ASCP and identify actionable targets for this lethal cancer we applied DNA content flow cytometry to a series of 15 tumor samples including five patient derived xenografts (PDXs). We interrogated purified sorted tumor fractions from these samples with whole genome copy number variant (CNV), whole exome sequencing (WES) and ATAC-seq analyses. These identified a variety of somatic genomic lesions targeting chromatin regulators in ASCP genomes that were superimposed on well characterized genomic lesions including mutations in TP53 (87%) and KRAS (73%), amplification of MYC(47%), and homozygous deletion of CDKN2A(40%) that are common in PDACs. Furthermore, a comparison of ATAC-seq profiles of three ASCP and three PDAC genomes using flow-sorted PDX models identified genes with accessible chromatin unique to the ASCP genomes, including the lysine methyltransferase SMYD2 and the pancreatic cancer stem cell regulator RORC in all three ASCPs, and a FGFR1-ERLIN2 fusion associated with focal CNVs in both genes in a single ASCP. Finally, we demonstrate significant activity of a pan FGFR inhibitor against organoids derived from the FGFR1-ERLIN2 fusion-positive ASCP PDX model. Our results suggest that the genomic and epigenomic landscape of ASCP provide new strategies for targeting this aggressive subtype of pancreatic cancer.

#### Keywords

adenosquamous carcinoma of the pancreas; flow sorting; somatic genomics; ATAC-seq; therapeutics

#### Introduction

It is estimated that in 2020, 57,600 Americans will be diagnosed with pancreatic cancer and 47,050 will die from the disease, making pancreatic cancer the third most common cause of cancer death (1). Adenosquamous cancer of the pancreas (ASCP) is a rare subtype of pancreatic cancer representing 1–4% of all pancreatic cancers (2). Strikingly, ASCP displays a higher metastatic potential and a worse clinical outcome than the more common pancreatic ductal adenocarcinoma (PDAC) (4). Yet, a large population-based analysis did not detect any differences in tumor stage at the time of diagnosis between PDAC and ASCP (5). There are various theories as to why this histological subtype exists, as normal pancreas tissue has no benign squamous epithelium. These include the induction of a squamous epithelium by inflammation, the collision of histologically distinct tumor lineages in the same tissue, and the aberrant differentiation and enrichment of tumor stem cells that acquire features of one or both subtypes (2). However, the rarity of ASCP, the scarcity of tissue samples suitable for high resolution genomic analyses, and the lack of validated preclinical models has limited the study of this deadly subtype of pancreatic cancer. Thus, little is known about the genomic landscape of ASCP, its relationship with PDAC and potential driver aberrations

We acquired both fresh frozen and formalin-fixed, paraffin-embedded (FFPE) ASCP samples, including primary and metastatic tissue from a rapid autopsy, from multiple institutions. We then applied DNA content flow cytometry, whole genome copy number and whole exome sequence analyses to the ASCP tumors to comprehensively profile the genomic landscape of this aggressive tumor. Notably, our integrated analyses of ASCP identified a variety of somatic genomic lesions targeting chromatin regulators in ASCP genomes that were superimposed on well characterized genomic lesions including mutations in KRAS and TP53, homozygous deletion of CDKN2A, and amplification of MYC, which are common to PDACs (6-8). We then compared the chromatin accessibility patterns of ASCP with PDACs using patient-derived xenograft (PDX) models with genomic lesions and mutations seen in patient samples. Of significant interest were genes with open chromatin regions unique to ASCP that included a FGFR1-ERLIN2 fusion, and the pancreas cancer stem cell regulator *RORC*. The latter is a nuclear hormone receptor known for its role in Th17 cell specification and regulation of inflammatory cytokine production and has been shown to be enriched during the emergence of chemorefractory tumors, a feature shared with ASCP, in the KPC mouse PDAC model (9). ASCP organoids were used to test the activity and functional significance of candidate therapeutic targets in ASCP. Specifically, organoids carrying the FGFR1-ERLIN2 fusion show a significant response to pharmacological FGFR inhibition. These results advance our understanding of this lethal subtype of pancreas cancer and provide new candidate targets for developing effective therapies for patients with ASCP and potentially refractory PDAC.

### Materials and Methods

#### **Tumor Samples:**

All patients gave written informed consent for collection and use of the samples. All tissue was collected for this study under Mayo Clinic IRB 14–003050. All tumor samples were histopathologically evaluated by a board certified GI pathologist at each institution prior to genomic analysis. As per the 2019 World Health Organization (WHO) Classification of Tumors, Digestive System Tumours, ASCP samples in this study both primary and metastatic, including patient matched samples from a rapid autopsy case, and patient derived xenograft (PDX) tissues (Supplemental Table 1), were defined by the presence of at least 30% squamous differentiation (Supplemental Figure 1). PDXs from resected ASCP tumors were obtained from the Mayo Clinic Institutional PDX program in Hepatobiliary and Pancreatic Tumors. All established tumors undergo extensive histologic comparison and DNA fingerprinting to confirm that each PDX is derived from each specific patient. The ASCP histology was confirmed for each PDX using WHO criteria. All research conformed to the Helsinki Declaration (http://www.wma.net/en/30publications/10policies/b3/).

#### Flow Cytometry:

Excess paraffin was removed from each FFPE sample with a scalpel from either side of  $40-60 \mu m$  scrolls then processed according to our published methods (10,11). We used one to

Page 4

three 50  $\mu$ m scroll(s) from each FFPE tissue block to obtain sufficient numbers of intact nuclei for sorting and molecular assays. Frozen tissue samples were minced in the presence of NST buffer and DAPI according to published protocols (10,12,13). Nuclei from each sample were disaggregated then filtered through a 40  $\mu$ m mesh prior to flow sorting with an Influx cytometer (Becton-Dickinson, San Jose, CA) with ultraviolet excitation and DAPI emission collected at >450 nm. DNA content and cell cycle were analyzed using the software program MultiCycle (Phoenix Flow Systems, San Diego, CA).

For sorting single nuclei into 96 well plates, organoid cultures were resuspended in DAPI (10  $\mu$ g/ml) then disaggregated using a 1mL 18g syringe followed by filtering through a 40  $\mu$ m mesh. Samples were examined under a microscope to confirm single nucleus suspensions. The forward scatter (FSC) and side scatter (SSC) signal and the fluorescence signals of the 488 nm Blue Laser and 355 nm UV laser were aligned and optimized using Ultra Rainbow fluorescent particles (3.0–3.4  $\mu$ m) containing a mixture of fluorophores that enable the Ultra Rainbow particles to be excited at any wavelength from 356 to 650 nm. The coefficient of variation (CV) was adjusted to a range of 1.9–2.9. A four- way sort was set up using a 100  $\mu$ l nozzle tip, 20 psi with a frequency 29.4 kHz. Nuclei were sorted in a one drop pure mode (high purity and high recovery of the sample) at 1500–2000 events per second to maintain purity and intactness of sorted material. In addition we deposited empty droplets in two wells and 50 nuclei and 100 nuclei in two additional wells as negative and positive controls for downstream amplification and sequencing.

#### Copy Number Analysis:

DNAs from frozen tissue and FFPE samples were treated with DNAse 1 prior to Klenowbased labeling. High molecular weight templates were digested for 30 minutes while DNAs from FFPE samples were digested for only 1 minute. In each case 1 µl of 10x DNase 1 reaction buffer and 2 µl of DNase 1 dilution buffer were added to 7 µl of DNA sample and incubated at room temperature then transferred to 70°C for 30 minutes to deactivate DNase 1. Sample and reference templates were then labeled with Cy-5 dUTP and Cy-3 dUTP respectively using a BioPrime labeling kit (Invitrogen, Carlsbad, CA) according to our published protocols (14). All labeling reactions were assessed using a Nanodrop assay (Nanodrop, Wilmington, DE) prior to mixing and hybridization to 400k CGH arrays (Agilent Technologies, Santa Clara, CA) for 40 hours in a rotating 65°C oven. All microarray slides were scanned using an Agilent 2565C DNA scanner and the images were analyzed with Agilent Feature Extraction version 11.0 using default settings. The aCGH data was assessed with a series of QC metrics then analyzed using an aberration detection algorithm (ADM2) (15). The latter identifies all aberrant intervals in a given sample with consistently high or low log ratios based on the statistical score derived from the average normalized log ratios of all probes in the genomic interval multiplied by the square root of the number of these probes. This score represents the deviation of the average of the normalized log ratios from its expected value of zero and is proportional to the height h (absolute average log ratio) of the genomic interval, and to the square root of the number of probes in the interval. All data have been deposited to GEO repository (accession GSE143256).

#### Whole Exome Sequencing:

DNAs from each sorted tumor population and a patient matched control sample were sequenced within the Mayo Clinic Medical Genome Facility (MGF) using established protocols for whole exome analysis. Briefly, whole exon capture was carried out with Agilent's SureSelect Human All Exon 71 MB v6 kit. 500 ng of the prepped library is incubated with whole exon biotinylated RNA capture baits supplied in the kit for 24 hours at 65 °C. The captured DNA:RNA hybrids are recovered using Dynabeads MyOne Streptavidin T1 (Invitrogen). The DNA was eluted from the beads and desalted using purified using Ampure XP beads (Beckman Coulter, Indianapolis IN). The purified capture products were then amplified using the SureSelect Post-Capture Indexing forward and Index PCR reverse primers (Agilent) for 12 cycles. Libraries were loaded onto paired end flow cells at concentrations of 4–5 pM to generate cluster densities of 600,000–800,000/mm<sup>2</sup> using the Illumina cBot and HiSeq Paired end cluster kit version 3. The flow cells are sequenced as 101 X 2 paired end reads on an Illumina HiSeq 2500 or 4000 using TruSeq SBS sequencing kit version 3 and HiSeq data collection version 1.4.8 software. Base-calling was performed using Illumina's RTA version 1.12.4.2.

#### Variant Calling and Annotation:

Aligned tumor and germline data (in bam format) were assembled for each patient. Tumorspecific variants were called using VarScan2 (version 2.3.9) (16) available on a high performance cluster computing environment. A minimum coverage of 10 reads in normal and tumor was used to call somatic variants, a minimum variant frequency of 0.08 to call a heterozygote, and a somatic p-value of 0.05 as a threshold to call a somatic site. The SNP calls were filtered to remove those near indel positions, and also removed likely false positives associated with common sequencing- and alignment-related artifacts (17). The variants were annotated functionally using Annovar (18) with hg19 reference genome.

#### Single cell sequencing:

We applied DNA content flow sorting to isolate single nuclei from PAX265-derived organoids and deposited them in individual wells of a 96 well plate. This approach includes fluorescence measures for each well that confirms the DNA content of each sorted nucleus and discriminates aggregates from single nuclei. DNA extraction and amplification of each sorted nucleus was performed using the REPLI-g Single Cell Kit (QIAGEN, Germantown, MD) according to protocols from the supplier. *KRAS* exon 2 was amplified using Platinum<sup>TM</sup> *Taq* DNA Polymerase (Thermo Fisher, Waltham, MA) according to the supplier's protocol, then sequenced by M13 primed Sanger sequencing. Sequencing was done through the Arizona State University Genomics Core. Sequences were visually inspected using Sequence Scanner V2 software (ABI, Foster City, CA).

#### **Omni-ATAC:**

Approximately 50,000 flow sorted nuclei were utilized for OMNI ATAC-seq following the protocol by Corces *et al.* (19). Nuclei were subjected to transposition reaction following cell lysis and washing steps. The transposition reaction was performed at 37 °C for 30 min in a thermomixer with shaking at 1,000 rpm. DNA was purified and indexed ATAC-seq libraries

amplified by PCR. The enrichment of accessible regions in each library was determined by real-time PCR targeting known highly accessible and closed chromatin sites and expressed as fold difference. Libraries passing this quality control step were sequenced to 51 base pairs from both ends on an Illumina HiSeq 4000 instrument at the Mayo Clinic Center for Individualized Medicine Medical Genomics Facility.

Omni-ATAC data were analyzed using the HiChIP pipeline developed by the Mayo Bioinformatics Core (20). Briefly, paired-end reads were mapped to the HG38 genome reference using Burrows-Wheeler Alignment tool (BWA) (21). Pairs of reads with one or both reads uniquely mapped were retained and duplicates removed using Picard MarkDuplicates command (http://broadinstitute.github.io/picard/). Peaks were identified using the Model-based Analysis of ChIP-Seq (MACS2) software package at FDR <=1% (22). For data visualization, BEDTools (23) in combination with in-house scripts were used to generate normalized tag density profiles at a window size of 200 bp and step size of 20 bp. Differential analysis of chromatin accessibility was performed using the DiffBind package (24). Raw read counts were normalized using the trimmed mean of M values (TMM) method, and sites with fold changes >=2 and FDR <=0.05 in the DiffBind test were extracted to represent differential chromatin accessibility sites. The ATAC-seq data are available in the GEO repository (accession GGSE143256).

#### Immunofluorescence:

Immunofluorescent staining was completed using FFPE sections. Slides were first deparaffinized using xylenes and a graded ethanol series, followed by permeabilization with Triton X-100. Antigen retrieval was completed using citrate buffer (pH 6) and steam heat. After blocking with 1 % BSA and 2% goat serum, slides were incubated with either anti-histone H3 acetyl K27 antibody (Abcam, Cambridge, UK) at 1:1000 dilution at 4 degrees overnight or anti-histone H3 mono methyl K4 antibody (Abcam), at 1:2000 dilution for 2 hours at room temperature. Slides were then incubated with goat anti-rabbit Alexa Fluor 594 secondary antibody (Life Technologies, Carlsbad CA) for one hour at room temperature and cover-slipped using ProlongGold antifade reagent with DAPI (Life Technologies) to preserve fluorescence and stain nuclei. Images were captured using a Zeiss Axio Oberserver Z1 microscope at 40x magnification.

#### Immunohistochemistry:

IHC staining was performed on a Leica BOND RX automated immunostainer using BOND primary antibody diluent and BOND Polymer Refine DAB Detection kit according to the manufacturer's instructions (Leica Biosystems, Buffalo Grove, IL). Pre-treatment was performed using citrate buffer at 100<sup>0</sup>C for 30 min, and tissue was stained using rabbit antihuman RORC(t) (polyclonal, PA5–23148, Thermo Fisher Scientific, Waltham MA) at a dilution of 1:4000. Stained slides were scanned (20x magnification) using a Pannoramic P250 digital slide scanner (3DHistech, Thermo Fisher Scientific).

#### Organoid cultures and treatments:

Organoids were seeded in 96-well plate coated with Matrigel (5,000 cells /well) in organoid media. Fresh organoid media and DMSO or drug (10 wells/condition) was added every 2–3

days. MTT assay were performed on Day 7 to measure proliferation. The absorbance listed for each drug is the average of the 10 wells per condition. Each experiment was performed in triplicate and the average  $IC_{50}$  reported.

## Results

#### Genomic landscape of ASCP identified alterations in chromatin regulators.

We screened 48 ASCP patient samples with DNA content flow cytometry. Tumor-specific fractions were detected in 15 samples. These included both fresh frozen and FFPE archival tissues. All tumor fractions were non-diploid by flow cytometry and genomically aberrant with both CNVs and somatic mutations. For example the sorted aneuploid fraction of sample A90–5 had multiple CNVs including gains of 18q11, 8q24, 7p12, 5p and 1p34.2 and multiple somatic mutations including in the lysine demethylase *KDM3A* (Figure 1). In contrast, the co-existing sorted diploid fraction was genomically normal at both the somatic CNV and mutation levels. The tumor fractions in the ASCP biopsies varied extensively (<5% to >50%) as a percentage of total cellular content (diploid, tetraploid and aneuploid fractions). However our sorting provided highly enriched (>90% tumor content) samples for analyses enabling the discrimination of homozygous deletions and mutations in these complex genomes. The samples that were diploid only by flow cytometry (n=20) were CNV neutral and lacked detectable somatic CNVs and mutations in their exomes (Supplemental Figure 2). In the remaining 13 cases there were insufficient intact nuclei for sorting due to extensive tissue degradation.

Biopsies from PDX models contain variable amounts of host tissue. Studies have used bioinformatic approaches to subtract mouse sequences and to extract and enrich human tumor profiles from these samples (25). In contrast we used our DNA content sorting to prepare PDX samples for analyses. Near-diploid peaks representing mouse host nuclei were present in each sample (Supplemental Figure 3). We selected peaks that corresponded to human tumor nuclei based on their ploidy and lack of overlap with DNA content and cell cycle profiles of the host tissue. The DNAs from each sorted population of interest were interrogated with CGH arrays to confirm the human tumor content and to provide a CNV profile of each tumor genome. These sorted human tumor samples were subsequently used for whole exome NGS with a patient matched normal sample.

We detected common recurring PDAC driver events in each of the ASCP genomes. These included *CDKN2A* and *SMAD4* homozygous deletions, *KRAS* and *TP53* mutations, and *MYC* amplifications (Figure 2). Our results support a model whereby ASCPs evolve from the same tumor lineage as the more common PDACs. However, superimposed on ASCP genomes are "hits" targeting key regulators of chromatin organization. These hits were heterogeneous and included homozygous deletions in *SMARCA2, ARID2*, and *ASXL2*, and somatic mutations in the DNA demethylase *TET1*, the histone ubiquitin E3 ligase *MSL2*, and the chromatin regulator *KANSL1* (Figure 2; Supplemental Figures 4–6). We also identified a series of high level (log<sub>2</sub> ratio >2) amplicons targeting genes associated with a variety of oncogenic signaling pathways. These included *TLK2, ASH2L, MPL*, and *FRS2* (Supplemental Figures 7 and 8). The latter, (Fibroblast Growth Factor Receptor Substrate 2) is an adapter protein that plays an important role in the activation of MAP kinases and in the

phosphorylation of PIK3R1, the regulatory subunit of phosphatidylinositol 3-kinase, in response to ligand-mediated activation of FGFR1 (26). In addition we detected a  $FGFR2^{A322G}$  mutation that has been reported as a pathogenic variant in esophageal cancer (27). Notably this mutation targets exon 8 encoding the second half of one of the three Ig domains and part of the IIIb isoform associated with epithelial tumors (28).

ASCP patient samples included a rapid autopsy case with a primary and multiple metastatic lesions. We detected small (<10% of total cell content) aneuploid tumor fractions in the primary tissue and each of three metastatic lesions (Figure 3). The CNV profiles of each flow-sorted aneuploid population overlapped and included gains of 8q21.3-qtel that contained the *MYC* locus, a focal amplicon at 15q23-q24.2 that included *PKM* and *CD276*, and a homozygous deletion at 5q21.3 targeting *EFNA5* and *FBXL17*. The primary and metastatic lesions also shared pathogenic mutations including *TP53*<sup>Y205S</sup> and *DCLK1*<sup>R699X</sup>. Strikingly none of these tissues had a *KRAS* mutation. However, we detected a somatic *ERBB4*<sup>\$183C</sup> mutation that was present in each sorted tumor population. Although not previously reported, this non-conserved variant is located adjacent to multiple pathogenic mutations in the Furin-like cysteine rich domain of ERBB4 (29).

The genomes of the five ASCP PDX models contain recurring pancreatic cancer lesions. These included pathogenic *KRAS* mutations in all 5 cases and homozygous *CDKN2A* deletions in 4/5 samples (Table 1). The one exception with an intact *CDKN2A* locus had a homozygous pathogenic *RB1* nonsense mutation. Four of these models had *TP53* mutations. The one *TP53*<sup>WT</sup> case (PAX208) had a homozygous deletion of *PTEN* a context associated with loss of TP53 protein and transcriptional activity (30). In addition, we detected a homozygous nonsense *SMAD4* mutation in PAX208 and loss of 18q21.1 in the remaining four cases. Thus our preliminary data highlight the presence of prevalent genomic drivers of pancreatic cancer in our PDX models. Notably the *KRAS* mutations included the common G12D variant in three of the PDXs and the less frequent G12S and Q61H variants in each of the two additional samples. Although limited to 5 PDXs for this rare cancer these diverse *KRAS* genotypes provide favorable models to study the biology of ASCP and to evaluate therapeutic strategies against this aggressive subtype of pancreatic cancer. Similar to the patient samples, the PDXs had homozygous deletions and mutations targeting epigenomic regulators including *ARID2*, *HMGB2* and *KANSL1* (Figure 2, Supplemental Figure 6).

#### ASCP shows a distinct chromatin accessibility pattern.

Given the presence of genomic lesions targeting key epigenomic regulators we hypothesized that ASCP and PDAC genomes would have shared and unique sites of active chromatin at key oncogenic drivers. We performed Omni-ATAC, an improved ATAC-seq protocol (19), on three of the ASCP PDXs and three PDAC PDXs that were also profiled with whole genome CNV analysis (Supplemental Figure 9). Active chromatin was present at the *KRAS* and the *MYC* loci in all six samples profiled (Supplemental Figure 10). Unique active chromatin sites included *SMYD2*, a lysine methyltransferase with specificity for the active chromatin mark H3K4me1, in all three ASCP samples (Figure 4A, B). Immunofluorescence analysis revealed higher H3K4me1 levels in ASCP PAX129 compared to PDAC PAX297 and normal pancreas tissue. In addition, all three ASCP PDX models had active chromatin at

the *CDK6* locus (Supplemental Figure 11). These included two, PAX139 and PAX217, with high level (log<sub>2</sub> ratio >2) *CDK6* amplicons. Given the presence of co-occurring *CDKN2A* deletions in these genomes we hypothesized that CDK4/6 inhibitors would have activity in these ASCP models. However we did not observe any response to either palbociclib or abemaciclib in organoid cultures derived from PAX139. The ASCP PDX genomes also had active chromatin at the *RORC* locus compared to PDACs (Figure 5A). Cytoplasmic and nuclear staining was observed in the ASCP and the PDAC PDX models, and in tissue from the rapid autopsy series (Figure 5B, Supplemental Figure 12). Notably there was an enrichment of nuclear staining in the metastatic lesions.

#### ASCP organoids carrying a FGFR1 fusion show sensitivity to pan FGFR inhibitor

We detected the CNV footprint of a *FGFR1-ERLIN2* fusion on 8p12 and its ATAC-seq profile in PAX265 (Figure 6A–C). These were consistent with the presence of an active fusion that included the extracellular domains and the kinase domains of FGFR1 fused to ERLIN2. This fusion has been reported in breast cancer and its' dimerization is believed to be sensitive to the FGF (31). Organoids developed from PAX265 with the *FGFR1-ERLIN2* fusion were sensitive to the pan FGFR inhibitor infigratinib with an IC<sub>50</sub> near 250nM in triplicate experiments (Figure 6D). To our knowledge this represents the first example of ASCP response to a targeted therapy. Notably this fusion co-occurred with a pathogenic *KRAS*<sup>G12V</sup> mutation (Figure 6E). The *KRAS*<sup>G12V</sup> variant was detected in 34/50 (68%) sorted single nuclei from the organoid model suggesting it did not affect the robust response to the pan FGFR inhibitor (Supplemental Figure 13). Given the presence of a pathogenic *FGFR2*<sup>A322G</sup> mutation and high level amplification of *FRS2* in patient samples our data suggests that FGFR inhibition may provide a therapeutic window for a subset of ASCP similar to that observed in cholangiocarcinoma (32,33).

## Discussion

To our knowledge this is the first study to apply DNA content sorting to the genomic analysis of ASCP. Our methods yielded purified (>95%) tumor samples for whole genome analyses from a variety of clinical samples. These include fresh frozen and FFPE tissues with low tumor content (<10-20%) and high amounts (>90%) of necrosis and debris. By profiling flow-sorted tumor fractions from each sample of interest we identified multiple heterogeneous deletions, mutations and amplicons targeting epigenomic and stem cell regulators superimposed on common PDAC driver lesions. Notably, many of these can be classified as either Tier 1 or Tier 2 variants based on ASCO/CAP guidelines (Supplemental Table 2) (34). Our flow assays exploit a cancer cell hallmark, an euploidy, present in >90%of solid tumors (35). The use of nuclei and DAPI provides discrimination of diploid, aneuploid, and tetraploid cells with C.V.s of 10% for DNA content (36,37). Thus, our assays can identify and sort tumor nuclei with DNA contents ranging from near diploid to hypertetraploid from clinical and preclinical samples of interest (10,14,38). In those relatively rare cases where DNA content overlaps with normal diploid content, the gating and collecting of proliferating fractions (e.g. 4N-G<sub>2</sub>/M) can enrich for tumor nuclei. Comparisons of ASCP with PDAC have been used to search for genomic lesions that account for the differences in appearance and clinical behaviors of these pancreas cancer

subtypes. These include reports of enrichment for *TP53* mutations and copy number losses at chromosome 3p in ASCPs relative to PDACs (39). However, we did not observe these differences in our small cohort of flow sorted samples. Additional studies have reported recurring mutations targeting *UPF1*, which encodes an RNA helicase essential for the highly conserved RNA degradation pathway, nonsense-mediated RNA decay (NMD), as a distinguishing feature of ASCP (40). The mutations were detected by targeted sequencing of archived tissues in 18/23 ASCPs and clustered in two regions spanning exon 10 through exon 11. Notably, we did not detect *UPF1* variants in any of our flow sorted patient and PDX samples.

Currently there are chemotherapy regimens with significant clinical activity in patients with PDAC (41). However, a major clinical challenge is the emergence of chemorefractory tumors even after significant clinical responses. In contrast, ASCPs typically fail to respond and progress rapidly similar to relapsed PDACs. Our combined CNV and exome results demonstrate that ASCP genomes contain the common lesions seen in PDACs including *KRAS* and *TP53* mutations, homozygous deletions of *CDKN2A* and *SMAD4* and amplification of *MYC*. This supports previous reports and a model whereby ASCP shares a common origin with PDAC. However, our results highlight that ASCP acquire additional heterogeneous lesions including homozygous deletions, mutations and focal amplicons targeting key regulators of the epigenome. In addition, our ATAC-seq analyses of the epigenome in preclinical models identified active chromatin at the *SMYD2* locus, a lysine methyltransferase with specificity for H3K4me1 as a distinguishing feature of ASCP compared to PDAC. This was associated with high expression of this active chromatin histone mark (Figure 4).

A genome-wide functional study of PDAC in the KPC mouse model identified a key role for immuno-regulatory genes in the self-renewal and maintenance of therapy resistant tumor stem cells in the pancreas (9). Notably, expression of *RORC*, a nuclear hormone receptor known for its role in Th17 cell specification and regulation of inflammatory cytokine production, increased with progression and its blockade via genetic or pharmacologic approaches depleted the pool of resistant cancer cells and profoundly inhibited tumor propagation in the KPC model. Our ATAC-seq data identified *RORC* as another distinguishing ASCP feature. Notably, IHC staining identified both cytoplasmic and nuclear *RORC* expression in all five PDX models and an enrichment of nuclear staining in metastatic lesions from the ASCP rapid autopsy (Figure 5, Supplemental Figure 12). This staining pattern compared to exclusive cytoplasmic expression, has been correlated with higher pathological tumor stages for PDACs at diagnosis (9).

The aggressive clinical behavior including lack of response to chemotherapy and an active *RORC* signature suggests that ASCPs are enriched for cell populations with cancer stem cell features. The presence of a clonal pathogenic *DLCK1* mutation in the primary and metastatic lesions of a rapid autopsy, and a homozygous deletion of *ASXL2* and a pathogenic *PROX1* mutation in separate primary tumors provide additional support for the putative role of active pancreas cancer stem cells in ASCP (Figures 2 and 3, Supplemental Figure 5) (42). Thus, our analyses suggest that the differences between these two subtypes of

pancreatic cancer may be driven through heterogeneous epigenomic lesions and cancer stem cell enrichment rather than selected ASCP-specific recurring genomic lesions.

Significantly, we also identified FGFR lesions including a focal FRS2 amplification, a FGFR2<sup>A322G</sup> activating mutation and a FGFR1-ERLIN2 fusion in ASCP samples (Figures 2 and 6, Supplemental Figure 7). Each of these lesions has been shown to activate FGFR/ FRS2 signaling pathway in cancer (31,43–45). To our knowledge these have never been reported in this aggressive subtype of pancreatic cancer. Strikingly these lesions co-occur in tumors with pathogenic KRAS mutations. The co-occurrence of KRAS and FGFR activation suggests a unique ASCP-specific molecular context. The presence of KRAS activating mutations typically result in resistance to therapies targeting upstream signaling such as the EGFR (46). Nevertheless, organoids from RORC positive ASCP PDX models with a FGFR1-ERLIN2 fusion and a KRAS<sup>G12V</sup> mutation were highly sensitive to single agent FGFR inhibition (Figure 6). KRAS mutations are early driver events in pancreatic cancer that can activate parallel signaling to bypass targeted inhibition of receptor tyrosine kinases. The *KRAS* mutant allele was present at  $\sim 60\%$  frequency in the sorted PAX265 tumor nuclei and organoid. This is typical for flow-sorted primary and metastatic pancreatic tumors we have profiled where low level 12p copy number gain of a pathogenic driver KRAS variant co-exists with wild type alleles in the same genome (38). Single-cell sequencing confirmed the presence of the pathogenic G12V variant in 64% of sorted nuclei from the organoid. Although selective amplification of one allele resulting in allelic drop out of the other may occur in single cell analyses our results support a model whereby the fusion and the *KRAS*<sup>G12V</sup> founder mutation co-existed in single cells. Future studies with these models, including DNA and RNA sequencing of sorted single cells and nuclei, will explore the basis of this response and investigate possible mechanisms of acquired resistance.

In summary, our data provide the basis for the development of novel approaches to treat ASCP. We propose that ASCP evolve from the same lineage as PDACs yet consist of enriched levels of *RORC* positive cancer stem cells, a feature that may drive other tumors with mixed adenosquamous and adenocarcinoma features. Furthermore, in addition to an epigenome that may promote stem cell features we report that a subset of ASCP has activated FGFR signaling that can be targeted with current inhibitors. Although currently limited in numbers, the availability of PDXs and organoids that recapitulate the genomic and epigenomic lesions found in patient samples provides initial preclinical models to interrogate therapeutic targets in this lethal chemoresistant cancer. Of significant interest will be clinical trials with FGFR and RORC inhibitors that include correlative studies of genomic and epigenetic lesions in both ASCP and PDAC.

#### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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# Significance:

These data provide a unique description of the ASCP genomic and epigenomic landscape and identify candidate therapeutic targets for this dismal cancer.

Lenkiewicz et al.



# **Figure 1: DNA content histogram and genomic profile of flow sorted ASCP biopsy.** A) Diploid (2N) and aneuploid peaks were identified and sorted from FFPE tissue sample A90. B–C) CNV and mutation profiles of the normal 2N and tumor aneuploid populations. The X and Y axes in the CGH plots represent chromosome and log<sub>2</sub> ratios for each genome.

Lenkiewicz et al.



Oncomap summary of somatic lesions in ASCP genomes.



**Figure 3: Genomic profiles of primary and metastatic biopsies from ASCP rapid autopsy 9R.** A) DNA content histograms of primary (top) and perigastric (bottom) biopsies. B) CNV profiles of flow sorted 2N and aneuploid populations. C) Homozygous deletion at 5q21.3 in sorted aneuploid population targeting *EFNA5* and *FBXL7*. The X and Y axes in the CGH plots represent chromosome and log<sub>2</sub> ratios for each genome. Red shaded areas denote homozygous deletions. D) IGV views of somatic mutations in *TP53, DCLK1*, and *ERRB4* detected in the primary (1°), and metastases in the liver (M1), perigastric (M2) and lung (M3). The sorted diploid fraction from the primary was used as the patient matched control.





**Figure 4: ATAC-seq profile of** *SMYD2* **and active chromatin mark in ASCP PDX models.** A) IGV views of chromatin accessibility in ASCP (top 3 rows) and PDAC (bottom 3 rows) PDX models. All samples were flow sorted prior to ATAC-seq analysis. B) Immunofluorescence analysis (40x magnification) of H3K4me expression in normal pancreas and tumor tissue (PAX 139).

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6-PAX295-ATA_sed_norm.stf		
7-PAX297-ATA_sed_nom.tdf		
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PAX208

#### Figure 5: ATAC-seq profile and expression of *RORC* in PDXs.

A) IGV views of chromatin accessibility in ASCP (top 3 rows) and PDAC (bottom 3 rows) PDX models. All samples were flow sorted prior to ATAC-seq analysis. B) IHC analysis (20x magnification) of *RORC* expression in ASCP (PAX139. PAX217, PAX265) and PDAC (PAX297, PAX295, PAX300) PDXs.



#### Figure 6: FGFR-ERLIN2 fusion in PAX265.

A) Whole genome and; B) locus specific analysis of CNV gains targeting *ERLIN2* and *FGFR1*. C) IGV views of ATAC-seq profiles of *ERLIN2* (left panel) and *FGFR1* (right panel). Red arrows denote open chromatin peaks in each gene. D) Dose response profile of PAX265 organoids to Infigratinib. Organoids were seeded in 96-well plate coated with Matrigel (5,000 cells /well) in organoid media. After 2–3 days of growth, fresh organoid media and DMSO or drug (10 wells/condition) was added. Fresh media and drug was added every 2–3 days. MTT assay were done on day 7 to measure proliferation. The absorbance listed for each drug is the average of the 10 wells per condition. Experiment was performed in triplicate with an average IC<sub>50</sub> near 250 nM. E) IGV view of *KRAS*<sup>G12V</sup> mutation in flow sorted PAX265. F) Single cell sequencing of KRAS exon 2 in flow sorted PAX265 derived organoids. Sanger sequencing of single nuclei sorted from 50 individual organoids identified 19 homozygous (top) and 15 heterozygous (middle) *KRAS*<sup>G12V</sup> variants. The other 16 were *KRAS*<sup>G12G</sup> wild type (bottom).

#### Table 1

#### ASCP PAX Genotypes

	KRAS	TP53	CDKN2A	SMAD4	Epigenome "Hits"
PAX139	G12D	E204_L206delEYL	_/_	_/+	HMGB2 <sup>49-50dupGC</sup>
PAX208	Q61H	wT <sup>1</sup>	_/_	W227X	ARID2-/-
PAX217	G12D	G223V	_/_	_/+	KANSL1Q396X
PAX232	G12S	Q165K	_/_	_/+	ASH2L <sup>+++</sup>
PAX265	G12D	P177_C182delPHHERC	+/+2	_/+	KANSL1 <sup>T221I</sup>

<sup>1</sup>PTEN<sup>-/-</sup>

<sup>2</sup><sub>RB1</sub>R787X