Resolution of Novel Pancreatic Ductal Adenocarcinoma Subtypes by Global Phosphotyrosine Profiling

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Comprehensive characterization of signaling in pancreatic ductal adenocarcinoma (PDAC) promises to enhance our understanding of the molecular aberrations driving this devastating disease, and may identify novel therapeutic targets as well as biomarkers that enable stratification of patients for optimal therapy. Here, we use immunoaffinity-coupled high-resolution mass spectrometry to characterize global tyrosine phosphorylation patterns across two large panels of human PDAC cell lines: the ATCC series (19 cell lines) and TKCC series (17 cell lines). This resulted in the identification and quantification of over 1 tyrosine phospho-

Pancreatic ductal adenocarcinoma (PDAC) remains one of the most deadly solid cancers, characterized by extremely high mortality rates and poor treatment outcomes. Effective management of PDAC remains challenging, and there is a critical need for improved diagnostic and therapeutic strategies. The study of PDAC signaling networks is crucial for understanding the molecular aberrations driving this disease and for developing targeted therapies.

The characterization of signaling in PDAC cell lines was performed using immunoaffinity-coupled high-resolution mass spectrometry. This approach allowed for the identification and quantification of over 1800 class 1 tyrosine phosphorylation sites across two large panels of human PDAC cell lines: the ATCC series (19 cell lines) and the TKCC series (17 cell lines). The resulting data provided comprehensive insights into the signaling networks associated with PDAC.

Specifically, the three subtypes exhibited enhanced tyrosine phosphorylation of multiple RTKs including the EGFR, ERBB3, and MET. Interestingly, a similar RTK-enriched subtype was identified in the TKCC series, and ‘classifier’ sites for each series identified using Random Forest models were able to predict the subtypes of the alternate series with high accuracy, highlighting the conservation of three subtypes across the two series. Finally, RTK-enriched cell lines from both series exhibited enhanced sensitivity to the small molecule EGFR inhibitor erlotinib, indicating that their phosphosignature may provide a predictive biomarker for response to this targeted therapy.

These studies highlight how resolution of subtype-selective signaling networks can provide a novel taxonomy for particular cancers, and provide insights into PDAC biology that can be exploited for improved patient management.
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Poor survival rates and limited therapeutic options (1). Post-operative treatment of patients is largely limited to chemotherapeutics, such as gemcitabine, nab-paclitaxel or Folfirinox, although the addition of the EGFR-directed kinase inhibitor erlotinib to gemcitabine results in a modest improvement in patient survival (2). In addition, patient stratification for therapy remains in its infancy. These factors highlight an urgent need to better understand the molecular mechanisms that contribute to PDAC development, progression and heterogeneity.

Over the last two decades, a deeper understanding of the genetic and molecular basis of cancer has led to the development of targeted therapeutics and personalized treatment strategies that combine such approaches with companion biomarkers. This paradigm has yet to be successfully applied to PDAC, which likely explains its poor overall response to adjuvant therapy. Although almost all PDACs harbor activating mutations in KRAS, and inactivating mutations in TP53, SMAD4 and CDKN2A occur at a frequency of >30% (3, 4), mutations in these genes are not associated with clinically ‘actionable’ phenotypes. Evidence for the existence of different molecular phenotypes of PDAC has been found through exploration of the genomic landscape of this disease. Exome sequencing identified an unsuspected role for axon guidance pathway genes in ~20% of PDAC patients (3), whereas whole genome sequencing of PDAC specimens provided the basis for classification into four subtypes based upon patterns of genomic structural variation (4). In the latter study, a subtype of patients characterized by unstable genomes and/or a BRCA mutational signature was demonstrated to have increased sensitivity to platinum-based therapy. In addition, transcript profiling subclassifies PDAC into subtypes exhibiting contrast histopathological characteristics, mutation patterns and patient outcome (5) as well as differential sensitivity to erlotinib and gemcitabine (6), and a gene expression signature corresponding to the EGFR pathway correlates with erlotinib responsiveness in primary PDAC xenografts (7).

These genomic and transcriptomic studies strongly suggest that a greater understanding of the phenotypic heterogeneity of PDAC may lead to improved patient stratification for therapy.

A variety of proteomics approaches have been applied in the search for diagnostic or prognostic biomarkers for PDAC, although this work has yet to achieve clinical impact (8). In a recent study, proteomic and phosphoproteomic interrogation of PDAC versus normal tissue via a LC-MS/MS workflow identified several protein kinases exhibiting increased expression in tumor tissue, including HIPK1 and MLCK (9). This suggests that a comprehensive survey of deregulated signaling in PDAC might represent a fruitful strategy for identification of therapeutic targets and disease subclassification. Furthermore, it is well-established that tyrosine kinase signaling pathways can be deregulated at several different levels in PDAC. In addition to the near-universal presence of activating KRAS mutations (3), these include: overexpression of particular RTKs, including EGFR, ERBB3 and AXIN (10–13); increased activation of nonreceptor tyrosine kinases, such as Src (14); and enhanced activation of the phosphatidylinositol 3-kinase (PI3K) pathway because of AKT2 amplification (15) or PTEN loss (16). However, to date, interrogation of tyrosine kinase signaling in PDAC has been limited to candidate-based studies, and an unbiased, global survey of tyrosine phosphorylation patterns in this malignancy has yet to be reported. In the current study, we have addressed this knowledge gap by undertaking global MS-based phosphotyrosine profiling (17, 18) across two large, independent PDAC cell line cohorts, revealing novel PDAC subgroups characterized by contrasting tyrosine phosphorylation patterns, signaling networks, and sensitivity to erlotinib. This new molecular taxonomy for PDAC provides important insights into disease mechanisms and highlights potential biomarkers to help guide patient stratification for therapy.

EXPERIMENTAL PROCEDURES

Cell Culture—Two cohorts of pancreatic cancer cell lines were used. Cohort 1 cell lines (termed the ATCC Cohort) were purchased from and authenticated by the American Type Culture Collection (ATCC) (AsPC-1, BxPC-3, CFPAC-1, Capan-1, Capan-2, HPAC, HPAF-II, Hs700T, Hs766T, Panc 02.03, Panc 03.27, Panc 04.03, Panc05.04, Panc 08.13, Panc 10.05, Panc-1, PL45, MiaPaCa-2, SU.86.86, SW1990) and were used and cultured according to ATCC protocols. Cohort 2 cell lines (termed the TKCC Cohort) were isolated from primary patient-derived pancreatic ductal adenocarcinoma xenografts in house (TKCC 2.1, TKCC 04, TKCC 05, TKCC 06, TKCC 07, TKCC 09, TKCC 10, TKCC 12, TKCC 14, TKCC 15, TKCC 16, TKCC 17, TKCC 18, TKCC 19, TKCC 22, TKCC 26, TKCC 27) (4). Animal experimentation for cell line generation was approved by the Garvan Institute/St Vincent’s Hospital Animal Ethics Committee (approval number: ARA 12/21). Culture conditions used are summarized in supplemental Table S1.

Phosphoproteomic Profiling—

Generation of Protein Lysates—Cells were plated, allowed to reach ~70% confluence, and placed into base culture medium, without the addition of serum or additional growth factors, for 6 h prior to lysate collection. This was undertaken in order to minimize differences in signaling because of culture conditions and enable comparison of the inherent signaling network properties of the different cell lines. For each cell line, two independent biological replicates of ~2 × 10^6 cells were generated. ATCC cells were homogenized in 8 M Urea lysis buffer containing 20 mM HEPEs (pH 8), 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM sodium orthovanadate and the reducing agent 1 mM tris (2-carboxymethyl) phosphine (TCEP). The procedure for TKCC cells was similar except the lysis buffer was based on 6 M Guanidine hydrochloride containing 50 mM Tris-HCL (pH 8). Samples were sonicated on ice, cleared by centrifugation, then alkylated for 45 min with 4 mM iodoacetamide (in the dark) at room temperature (RT). These lysates were subsequently used to conduct pTyr profiling, as well as Western blotting.

Preparation of Lysates for Mass Spectrometry Analysis—20 mg (TKCC panel) or 30 mg (ATCC panel) of cell line lysate was diluted to 1 mg guanidine hydrochloride or urea respectively with 50 mM ammonium bicarbonate, and digested at 1:200 (w/w) with Lysyl Endopeptidase (WAKO Chemicals, Cape Charles, VA, USA) for 4 h (RT) and then at 1:100 (w/w) overnight (RT) with TPCK treated trypsin (Worthington, Lakewood, NJ, USA, ratio of 1:100). Two “spike-in” heavy
peptides with alanine (a) C3N15 modifications were added prior to 
1C-18 clean-up (Waters, Milford, MA, USA). These peptides were 
EEF1A1 (EHA(ala)LLAPYTLGVK) (500 fM) and MAPK14 (HTDDEMTG-
tC-18 clean-up (Waters, Milford, MA, USA). These peptides were 
buffer (50 mM MOPS (pH 7.4) with 10 mM sodium phosphate dibasic, 
ofhilitized overnight and dissolved in immunoaffinity purification (IAP) 
buffer (50 mM MOPS (pH 7.4) with 10 mM sodium phosphate dibasic, 
thon the ATCC panel, Sepharose 4B beads pre-conjugated to 100 
mu (ATCC panel) or 1% (v/v) n-octyl-b-D-glucopyranoside (TKCC panel)). For the 
the TKCC panel, the flowthrough from the PY100 IP was collected and immunoprecipitated again with 150 
mu of Purified Mouse Anti-Phosphotyrosine (pY20) antibody (BD 
Transduction Labs, San Jose, CA, USA). Peptides exhibiting nonspe-
cific binding were removed by three washes with IAP buffer and three 
the phosphate mass spectra were acquired on Q Exactive Plus for the 
TC cohort samples or Orbitrap Velos for the ATCC Cohort sam-
les were desalted by loading onto a C18 pre-column (100 
mu m, 2 cm column; particle size 5 m; pore size 100 Å; Acclaim 
PepMap RSLC, Thermo Scientific), and separated on a 12 cm 75 
LC analytical column pulled to an internal diameter of 5 
u by a 
P-2000 laser puller (Sutter Instruments Co, Novato, CA, USA) packed 
C18 Magic reverse phase material, using a Dionex Ultimate 3000 
for the Q Exactive, peptides were eluted at 250 NL/min 
gradient of acetonitrile in 0.1% (v/v) aqueous formic acid as 
2–10% in 1 min, 10–26% in 27 min, 26–34% in 2 min, and 
34–80% in 5 min. The eluent was directed into a nano-electrospray 
ion source (Nanospray Flex, Thermo Scientific) with a spray voltage of 
1.7 kV. Survey scans in the mass range of 375–1600 m/z were 
acquired with a resolution of 70,000 at m/z 200 and an AGC target of 
366 ions (max IIT 120 ms). The top 12 most intense ions (ion selection 
threshold > 1000 ions, charge state 2–5, preferred peptide match, 
exclude isotopes) were sequentially isolated (isolation window 1.8 
/120 ms). The top 12 most intense ions (ion selection threshold > 1000 ions, charge state 2–5, preferred peptide match, 
exclude isotopes) were sequentially isolated (isolation window 1.8 
m/z at a target of 1e6 ions (max IIT 100 ms), and fragmented in the 
HCD cell (normalized collision energy 27). MS/MS spectra were 
acquired in the Orbitrap mass analyzer at a resolution of 17,500 at 
m/z 200 and ions were excluded from selection for a further 15 s. For the 
Orbitrap Velos, peptides were electrospayed directly into the MS 
using a spray voltage of 1.8 kV, survey scans in the mass range of 
350–1750 m/z were acquired with a resolution of 60,000 at m/z 400 
and an AGC target of 1e6 ions. The top 10 most intense ions (ion selection 
threshold > 500 ions, charge state >2) were sequentially isolated 
(isolation window 2.5 m/z at a target of 1e5 ions and frag-
mented in the linear ion trap (normalized collision energy 30), and ions 
were excluded from selection for a further 30 s.

MS Data Analysis—The mass spectrometry proteomics data have been deposited in the ProteomeXchange Consortium (19) via 
the PRIDE partner repository with the data set identifier PXD003198 
(http://www.ebi.ac.uk/pride/archive/login Username: reviewer 
72200@ebi.ac.uk, Password: Mta7c1N4). Raw MS data were ana-
lyzed using MaxQuant software version 1.1.2.25 (20). Database 
searching was performed using the Andromeda search engine inte-
grated into the MaxQuant environment against the human UniProt 
database release 2010_10 (35,073 entries), concatenated with known 
contaminants as well as the reversed sequences of all entries (21). Protein, peptide, and site FDRs were controlled at a maximum of 1%, 
individualized precursor mass tolerances were applied as described 
(20), fragment mass tolerance was 20 ppm (default). The MaxQuant 
search included the fixed modification of cysteine carboxamidomethyl 
and methionine oxidation, acetyl (Protein N-term) and phosphoryla-
(tion (STY), and was limited to a maximum of 2 missed cleavages. For 
completeness, label-free data underwent a post MaxQuant process-
ing step to extract all MS quantitated peaks from MaxQuant results 
files independent of MS/MS identification. All data were analyzed as log2 
values.

Normalization and Imputation of Label-free Data with “Spike-in” 
Standard Heavy Peptides—The label-free intensity values for the 
EEF1A1 and MK14 heavy peptides in each cell line were averaged 
and a subsequent normalization factor generated. Label-free MS 
intensity values for each pTyr site quantified in the data set were 
divided by the appropriate normalization factor. Imputation of missing 
values, which is necessary prior to more advanced bioinformatic 
analyses, was performed using K-Nearest Neighbor (22) on normally 
distributed data.

Bioinformatics and Statistical Analysis—Data analyses were per-
formed using Microsoft Office Excel, R statistical programming lan-
guage (Bioconductor packages) and the bioinformatics platform Per-
seus (Max Planck Institute of Biochemistry) version 1.2.0.19.

Stepwise Hierarchical Clustering and Classifier List Prediction—A 
stepwise approach was developed to identify stable clustering topol-
y. Firstly, hierarchical clustering was performed multiple times with 
an incremental inclusion of phosphorylation sites (ordered by variance 
across cell lines) each time. It started from using the top 10 most 
variable and continued until the top 412 variable phosphorylation 
sites were included for ATCC data set and the top 433 sites for TKCC. 
The sites were included on the basis of each phosphorylation site 
having greater variance than the average variance of all sites. By 
comparison of individual stepwise clustering results, the most fre-
quent clustering topology was then identified. The random forest 
classifier (19) for subtypes of each cell line cohort was built respec-
tively with a two-step approach by the R package randomForest 
4.6–10. In the first step, a classifier was built based on the pTyr 
sites that were used in the stepwise clustering analysis. In the second step, 
only the most predictive pTyr sites with a mean decrease accuracy 
greater than 1.5 in the first classifier were selected, and used to build a 
refined classifier.

Subtype-Specific Site Identification, Pathway Enrichment and Pro-
tein-Protein Interaction Network Analyses—To identify subtype-spe-
cific sites, a multiple-sample ANOVA test was performed with multiple 
test correction. This identifies pTyr sites that have statistically signif-
icant (p < 0.05) specificity to one of the subtypes. This was followed 
by pairwise t-tests (p < 0.05) to identify subtype-specific high and 
subtype-specific low pY sites, representing those sites showing high 
and low relative tyrosine phosphorylation in a subtype-specific man-
ner. KOBAS was used to perform pathway enrichment analysis (23). 
The hypergeometric test was selected to test statistical enrichment of 
KEGG and Reactome pathways, and the p values were corrected for 
multiple comparisons. The protein-protein interactions among pro-
teins of interest were retrieved from the Protein Interaction Network 
Analysis (PINA) platform (24), and kinase-substrate relationships were 
downloaded from the PhosphoSitePlus database. The networks were 
generated and visualized using PINA4MS, a Cytoscape plugin for 
PINA.

Cell Viability Assay—Cell viability was assayed in biological triplic-
ate (n = 3) directly using a CellTiter-Glo Luminescence Cell Viability 
Assay (Promega, Madison, WI, USA). Cells were plated in full medium 
to a 96-well plate at 1.5 × 10^3 cells/well. Erlotinib (Symansis, 
Timaru, NZ), or vehicle (DMSO) were added the next day (Day 0) and 
the assay was undertaken at Day 5. All drugs were dissolved in DMSO 
and control cells received vehicle only. Significance was assessed by 
one-way ANOVA across treatments (p < 0.05 considered as signifi-
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Dose-response relationships were analyzed with GraphPad Prism.

**Antibodies and Western Blot Analysis**—Immunoblotting and densitometry analyses were performed in biological duplicate (n = 2) as previously described (17). Protein extracts were separated using 4–12% Bis-Tris precast gels (Invitrogen, Carlsbad, CA, USA) and were processed according to the manufacturer’s protocol. Primary antibodies used in this study were pY1189 ErbB3, ErbB3, pY1148-EGFR, pY1173-EGFR, pY1192-EGFR, Met (Cell Signaling Technology); EGFR, pY1234-MET, hnRNPs C1/C2, U and K and RMBX (Santa Cruz Biotechnology, Dallas, TX, USA); FAK (BD Transduction Laboratories); beta-Actin (Sigma-Aldrich, St Louis, MO, USA) and hnRNPs A2/B1, H, and R (Abcam, Cambridge, UK).

**DNA Extraction and Somatic Mutation Mapping**—DNA was extracted using the Qiagen Allprep Kit in accordance with the manufacturer’s instructions. Mutation screening of OncoCarta Panels v1.0, v2.0 and v3.0 (Sequenom, San Diego, CA, USA) was conducted by the manufacturer to detect and quantify mutation frequencies in the cell line panel.

**RESULTS**

**Determination of the PDAC Tyrosine Phosphoproteome**—Tyrosine phosphorylation profiling was undertaken across 19 PDAC cell lines sourced from the American Type Culture Collection (ATCC) (supplemental Table S1), as well as normal pancreatic duct epithelial (HPDE) cells and HPDE cells expressing oncogenic KRAS (25). Tryptic peptides underwent immunaffinity enrichment coupled with a high-resolution mass spectrometry (HR MS using LC-MS/MS) (Fig. 1A). A comprehensive list of 1622 pTyr sites in peptide sequences derived from 797 nonredundant proteins were quantified (Fig. 1B) (supplemental Table S2). Based on the MaxQuant derived phosphorylation site localization probability score, 89% of sites (1,435) were identified as having a high confidence localization probability of > 0.75 (class 1) (Fig. 1C). MS experiments were performed in duplicate with high biological replicative technical reproducibility (Fig. 1D). The phosphopeptide intensities showed a normal distribution over ~6 orders of magnitude (Fig. 1E). Approximately 50% of the phosphoproteins identified exhibited a gene ontology cellular compartment classification localizing them at the plasma membrane (supplemental Fig. S1A). In terms of molecular function, prominent categories were protein kinase, phosphatase and receptor activity, consistent with signaling functions (Supplemental Fig. S1B). However, a novel observation was that ~15% of the proteins identified exhibited RNA binding activity (supplemental Fig. S1B).

**Subclassification of PDAC Cell Lines Based on Tyrosine Phosphorylation Patterns**—The tyrosine phosphorylation profiles of the ATCC PDAC cell lines were compared using a normalization (using standard heavy peptides) and bioinformatics pipeline, the latter consisting of four major focal points (supplemental Fig. S1C and S1D). We chose to utilize only the most accurate 958 class I pTyr sites quantified by total ion current (TIC) in at least 75% of cell lines (Fig. 1B), with imputation of missing values based on k-nearest neighbor performed on the normally distributed and normalized data prior to further bioinformatics analysis. First, to identify whether the tyrosine phosphoproteome can group the cell lines into reproducible subtypes, we conducted a stepwise resampling of the phosphorylation sites to isolate the 10 most frequent hierarchical clustering topologies. The most frequently occurring topology (Fig. 2A) was identified to occur 58% of the time. This topology includes two broad clusters, with the second broad cluster splitting into two subclusters containing greater than two lines. For the purpose of downstream bioinformatic analysis, the three main cell line clusters were defined as three subtypes. The ATCC subtype 1 (green bar) contains lines MiaPaca-2, Panc-1, Hs700T, Hs766T, Panc 02.03 and SW1990. ATCC subtype 2 (pink bar) contains lines Capan-1, Capan-2, Panc 03.27, Panc 04.03 and Panc 08.13 and ATTC subtype 3 (blue bar) contains lines PL45, Panc 05.04, HPAC, AsPC-1, HPAF-II, BxPC-3, CFPC-1 and SU86.86.

To determine whether some somatic mutations in PDAC were relevant to the identified subtypes, all cell lines were assessed using the high-throughput Oncomap mutation profiling panels (Sequenom). Thirty-nine cancer genes with known mutation profiles in cancer, totalling 495 individual somatic mutations, were screened (supplemental Table S3). The predominant mutations identified were located in KRAS (Gly12Cys, Gly12Asp, Gly12Val, Gly61His) and TP53 (Arg248Trp, Arg248Gln, Gly245Ser, Arg273His) (supplemental Fig. S2A, supplemental Table S3). Additional mutations in CTNNB1 (Ser37Phe) (in Hs700T cells) and FBXW7 (Arg465Cys) (in AsPc-1 cells) were also detected. These analyses also confirmed that BxPC-3 and Hs700T do not contain a KRAS mutation. Because the Oncomap panel did not screen for all known TP53 mutations, we complemented this analysis by a survey of the literature and appropriate databases (http://p53.free.fr/Database/Cancer_cell_lines/Pancreatic_cancer.html, http://p53.iarc.fr/CellLines.aspx) (supplemental Table S3). This revealed that the majority of the ATCC cell lines exhibit mutant TP53. Overall, there does not appear to be a correlation between the somatic mutation profiles of the cell lines and their allocated subtypes.

**Identification of Segregation-driving Phosphosites**—To identify subtype-specific pTyr sites, an ANOVA-based approach was implemented and applied. It identified 144 pTyr sites that had statistically significant specificity (p < 0.05) to one of the three subtypes (Fig. 2A, supplemental Fig. S1C). All subtype-specific phosphoproteins then underwent combined bioinformatic analyses to identify whether particular signaling pathways or protein-protein interaction networks were characteristic of the different ATCC subtypes.

ATCC subtype 1 was characterized by hypophosphorylation of 65 pTyr sites exhibiting higher phosphorylation across ATCC subtype 2 and 3 cell lines (Fig. 2B). When the function of proteins harboring these sites was considered, there was a significant enrichment for proteins involved in formation and regulation of cell-cell adherent junctions (AJs) and tight junctions (TJs) (p = 0.002 and 0.06 respectively) (Fig. 2C). Isola-
Fig. 1. Phosphoproteomics workflow and overall results. A, The shot-gun phosphoproteomics workflow. B, Tyrosine phosphorylation sites quantified in the ATCC cell line cohort phosphotyrosine (pTyr) proteomics screen according to localization probability (LP) and valid values, including class 1 (> 0.75 LP), class 2 (> 0.5, ≤ 0.75 LP) and class 3 (> 0.25, ≤ 0.5 LP). The term “valid values” refers to the presence of a MS quantitated peak in the MaxQuant results files for a given phosphorylation site. Six sites have a localization probability of less than 0.25 and are not listed within the figure. C, Visualization of the localization probability distribution and mass error for pTyr sites quantified in the ATCC data set. D, ATCC cell line biological replicate technical reproducibility according to Pearsons correlation (R). E, Phosphopeptides ranked by respective intensities (log10), with overlaid histogram visualization. The bar heights of the histogram estimate the density of phosphorylation site intensities at the indicated intensity intervals (23 bins in total) that are shown on the y axis.
Identification of ATCC cell line cohort subtypes and function-based analysis. A, The 19 ATCC cell lines reproducibly form 3 subtypes (See colored bars, subtype 1 is green, 2 is pink and 3 is blue) using stepwise unsupervised hierarchical clustering. The heatmap shows the clustering topology of the subtype specific sites identified in a supervised manner by ANOVA. B, Unsupervised hierarchical clustering of ATCC subtype 1 (green bar) specific phosphosites with low relative phosphorylation compared with subtype 2 (pink bar) and 3 (blue bar). C, Isolated protein-protein interaction network of ATCC subgroup 1 specific phosphosites with low relative phosphorylation. Protein-protein (blue lines) and kinase-substrate (green lines) interactions are displayed. Blue circles indicate phosphoproteins enriched in this subtype that are involved in the adherens junction pathway, and purple circles represent proteins characterized by the GOBP term ‘cell adhesion’ (GO:0007155). D, E-cadherin (CADH1) tyrosine phosphorylation at pTyr sites 753/754/755 from a single peptide for each cell line, and then normalized by median centering (z score).
and polarity, KRAS expression is characteristic of PDAC cells expressing and spliceosome pathways, with corrected a striking enrichment for proteins involved in the mRNA proc-
respective). The sites enriched in ATCC subtype 2 exhibited than seen in subtype 1 and 3 (1 and 7 up-regulated pTyr sites [Fig. 3
PDAC cells exhibit characteristics of epithelial to mesen-
cells respectively, these data indicate that ATCC subtype 1
expression are associated with epithelial and mesenchymal
structures as important for maintaining epithelial cell architecture
and polarity, KRAS expression is characteristic of PDAC cells
with an epithelial phenotype (26), and E-cadherin and vimentin
expression are associated with epithelial and mesenchymal
cells respectively, these data indicate that ATCC subtype 1
PDAC cells exhibit characteristics of epithelial to mesen-
chyme transition (EMT).

ATCC subtype 2 was found to contain 54 up-regulated subtype-specific pTyr sites, (Fig. 3A), a far greater number than seen in subtype 1 and 3 (1 and 7 up-regulated pTyr sites respectively). The sites enriched in ATCC subtype 2 exhibited a striking enrichment for proteins involved in the mRNA processing and spliceosome pathways, with corrected p values of 0.000003 and 0.000030 respectively. Mapping protein–protein interactions among these phosphoproteins identified the presence of interaction ‘hubs’ that centered on specific members of these pathways (Fig. 3B). Differences in the abundance of a particular phosphopeptide might reflect altered relative phosphorylation of the corresponding protein, and/or changes in total protein levels. Western blotting revealed that expression of hnRNPs C, R, H, and K, as well as RBMX, was similar across the entire panel (supplemental Fig. S2B), indicating that the increased tyrosine phosphorylation of these proteins in Subtype 2 represents enhanced relative phosphorylation rather than an alteration in protein abundance. However, in the case of hnRNPA2B1, increased expression in subtype 2 may contribute to increased yield of hnRNPA2B1-derived phosphopeptides.

Splicing factor 3B subunit 1 (SF3B1), a splicing component of the U2 snRNP complex, is mutated in 4–5% of PDAC patients (3). By quantification of 2 pTyr sites on SF3B1, we identified an ATCC subtype 2-specific pattern of elevated SF3B1 tyrosine phosphorylation (Fig. 3C). Although the role of these phosphorylation events are unclear at present, their increased abundance in ATCC subtype 2 provides further evidence that dysregulated mRNA stability or processing characterizes this PDAC phenotypic subgroup.

RTK Expression and Activation in the ATCC Subgroups—Given the importance of oncogenic RTK signaling in carcinogenesis, we determined the association between all RTKs quantified in the ATCC data set and the identified subtypes. We first normalized the HPDE KRAS and ATCC cell line panel data to matched site intensities from HPDEs, which we utilized as a ‘normal’ cell line control. We then isolated the relative TIC for all RTK pTyr sites. This revealed that expression of active KRAS in HPDEs results in markedly elevated site-selective tyrosine phosphorylation of several RTKs, including RON, EPHA2, MET and ERBB2 (Fig. 4A). In addition, this identified that ATCC subtype 3 shows in-

**Fig. 3. Identification of a subtype of ATCC cell line cohort enriched for RNA processing function.** A, Unsupervised hierarchical clustering of ATCC subtype 2 (pink bar) specific sites with high relative phosphorylation. B, The ATCC subtypes 2 specific sites with high phosphorylation were analyzed for protein-protein (blue lines) and kinase-substrate (green lines) interactions. Blue circles indicate phospho-
proteins enriched in this subtype that are involved in mRNA processing, and purple circles those in the splicosome pathway. C, SF3B1 tyrosine phosphorylation at pTyr sites 70 and 84 were summed for each cell line, and then normalized by median centering (z score).
increased pTyr abundance of key oncogenic RTKs (Fig. 4A). Using a two-sample t test, 15 RTK pTyr sites exhibited significantly enhanced phosphorylation in subtype 3 versus the other subgroups. These included sites from the RTKs EGFR, MET, RON, EPHA4, EPHB2/3/4, and DDR2. Western blotting using phosphospecific and total antibodies against EGFR, MET, RON, EPHA4, EPHB2/3/4, and DDR2. Western blotting using EGFR, ErbB3 and MET phosphotyrosine specific antibodies. ATCC cell lines are divided into subtype 1 (green), 2 (pink) and 3 (blue). Gray bar: unclassified cell line.

Subtype Validation Using a Second Cell Line Cohort—PDAC is a highly heterogeneous cancer (3). Therefore, it was important to determine whether the three subtypes identified in the ATCC cell line panel could be detected in an independent PDAC cell line panel generated by our group from patient-derived xenograft models. Application of the phosphotyrosine profiling methodology to this panel of 17 PDAC cell lines, termed “The Kinghorn Cancer Centre” (TKCC) series, in biological duplicate achieved a depth of 1220 quantified pTyr sites, 90% of which were Class 1 sites with strong biological replicate technical reproducibility (Fig. 5A, supplemental Fig. S3A and supplemental Table S2). In total, 1833 class 1 phosphosites were identified in the ATCC and TKCC data sets. A nonredundant list of the 25 most abundant tyrosine phosphopeptides in each PDAC cell line cohort was generated, identifying phosphosites of possible significance to PDAC pathogenicity (supplemental Fig. S3B). Our study also identified over 400 novel phosphorylation sites (supplemental Fig. S3C, supplemental Table S3).

Using our bioinformatics pipeline (supplemental Fig. S1C), stepwise unsupervised hierarchical clustering of class 1 TKCC pTyr sites was conducted. A more relaxed data filtering approach was applied to this data set, to allow for the hypothesis of 3 subtypes being present. Thus we required greater than 1/3rd (6 of the 17) of the cell lines to be quantified for a given site. Interestingly, as with the ATCC cohort, unsupervised hierarchical clustering of the TKCC cohort also revealed 3 subgroups including 2 broad clusters, one of which contained 2 subclusters (Fig. 5B). We therefore defined the TKCC cell line cohort subtypes as follows; TKCC subtype 1 (purple bar) with cell lines TKCC 14, 26, 19, 16, 09, 10, and 15. TKCC subtype 2 (orange bar)
**Fig. 5.** Subclassification of an independent PDAC cell line cohort based on tyrosine phosphorylation patterns. A, Tyrosine phosphorylation sites identified and quantified in the TKCC cell line cohort phosphotyrosine (pTyr) proteomics screen according to localization probability (LP) and valid values, including class 1 (> 0.75 LP), class 2 (> 0.5, < 0.75 LP) and class 3 (≥ 0.25, ≤ 0.5 LP). B, The 17 TKCC cell lines reproducibility form 3 subtypes (See colored bars, subtype 1 is purple, 2 is orange and 3 is blue) using stepwise unsupervised hierarchical clustering. The heatmap shows the identified topology using the subtype specific sites identified in a supervised manner using ANOVA. C, A protein-protein interaction network formed by TKCC subtype 3 cell lines (blue bar), with phosphoproteins (purple nodes) connected via protein-protein (blue lines) or by kinase-substrate (green lines) relationships. Phosphoproteins enriched in this subtype for axon guidance pathways and ErbB signaling are highlighted with blue and green shading respectively. D, LC-MS/MS quantified RTK tyrosine phosphorylation across the TKCC cell lines according to the identified subtypes (See colored bars, subtype 1 is purple, 2 is orange and 3 is blue). Activation sites identified are highlighted by red text.
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contains TKCC lines 05, 27, and 22 and TKCC subtype 3 (blue bar) contains lines 06, 12, 07, 04, 18, 2.1, and 17. As for the ATCC cohort, there was no obvious relationship between these subtypes and the mutation status of KRAS or TP53 (supplemental Table S3).

Next we identified subtype-specific pTyr sites present in the TKCC cohort, identifying 383 pTyr sites using the ANOVA-based approaches. TKCC subtypes 1 and 2 were identified to only have down-regulated subtype-specific pTyr sites (101 and 73 sites respectively), whereas TKCC subtype 3 was identified to only have up-regulated subtype-specific pTyr sites (209 sites). We then utilized pathway analysis to characterize these subtype-specific sites, and identified enrichment for Ephrin (p = 0.04) and EGFR (p = 0.05) signaling in TKCC subtype 3. However, unlike the ATCC cohort, no enrichment for mRNA regulatory function was found within the 3 TKCC subtypes. Within the up-regulated subtype 3 sites we identified a dense network of protein-protein interactions, with strong representation from the axon guidance and EGFR signaling pathways (Fig. 5C). Isolation and mapping of the relative TIC of all RTK pTyr sites within the TKCC data set identified enhanced phosphorylation of many RTKs in TKCC subtype-3 (Fig. 5D). Tyrosine residues exhibiting enhanced phosphorylation included activation sites on EGFR, EPHA2, DDR1, FGFR1, INSR, MERTK, MET, and RON, indicating that increased activity of these RTKs characterizes this subtype.

Prediction of Subtypes Using a Random Forest-derived Signature—Next, we set out to identify, in a supervised manner, phosphosites that distinguish the three subtypes present in both the ATCC and TKCC series. We first isolated “classifier sites” for each cohort. These are sites identified using Random Forest (RF) models, based on having a high power to predict the defined topological subgroups (Mean Decrease Accuracy score of ≥ 1.5). Testing of these sites (ATCC, 69 sites and TKCC, 78 sites) using the left-out samples during the construction process of each RF model (out-of-bag approach) estimated that they predict the subtype topology of the corresponding cohort without error. We then tested the predictive power of the “classifier sites” in the alternate data set to where they were identified. The 33 ATCC classifier sites found in the TKCC cell lines data set could predict the TKCC topology with an accuracy of 94%. The 59 TKCC classifier sites quantified in the ATCC data set could predict the ATCC topology with an accuracy of 84%. Therefore, the 33 ATCC “classifier sites” represent the most robust signature to predict PDAC subgroups. The “classifier sites” from each cohort were then compared to identify the “common classifier sites” (Fig. 6A and 6B). These 8 sites predict the TKCC topology...
with an accuracy of 94%, and the ATCC topology with an accuracy of 89%. Therefore, we have identified a signature of 8 sites with a high predictive power across two PDAC cell line cohorts. Interestingly, this classifier does not include RTK phosphorylation sites characteristic of the “RTK-enriched” subgroup in the two cell line cohorts, indicating that the 3 subgroups identified in the independent cell line panels share additional conserved features. Specifically, subtypes 1, 2 and 3 in each panel clearly exhibit low, medium and high phosphorylation, respectively, of BAIAP2 Y337, PKP3 Y84, PKP2 Y166, CTNN1 Y174, and CTNN1 Y904. We note that although specific sites in RIPK2, TJPP2 and PLEC are present in the classifier, their subtype-selective relative tyrosine phosphorylation differs between the two panels. A summary of phosphorylation sites that exhibit subtype-selective patterns of phosphorylation that are conserved across the two different cohorts is provided in Fig. 6C.

**Sensitivity of the identified PDAC subgroups to the targeted therapeutic, erlotinib**—Given the increased expression and/or activation of EGFR and ErbB3 in ATCC and TKCC Subtype 3 (referred to as the RTK-enriched subtypes) and the clinical relevance of the small molecule EGFR inhibitor erlotinib in PDAC (27), we tested selected cell lines from each cohort for sensitivity to this EGFR kinase inhibitor. Indeed, both the ATCC and TKCC RTK-enriched subgroups exhibited enhanced sensitivity to erlotinib (Fig. 7A–7C) and therefore an increased dependence on EGFR kinase activity for cell proliferation. Importantly, this was not mirrored in the sensitivity profile of the TKCC cohort for the PDAC therapeutic gemcitabine (Fig. 7D), indicating a drug-selective effect. These data indicate that a phosphosignature characteristic of the RTK-enriched subtype could be used as a predictive biomarker for erlotinib sensitivity, and highlight a potential clinical application for this novel PDAC subclassification.

**DISCUSSION**

In this paper we interrogate, for the first time, the heterogeneity of PDAC at the level of protein tyrosine phosphorylation. Our study identifies ~1800 tyrosine phosphorylation sites across two large, independent PDAC cell line cohorts, provides new insights into dysregulated pathways and processes in PDAC, greatly extends our knowledge regarding the suite of RTKs activated in this disease, and also enables a novel subclassification based on tyrosine phosphorylation profiles. In addition, it identifies a PDAC subgroup characterized by high-level phosphorylation of multiple RTKs that exhibits increased sensitivity to erlotinib, providing ‘proof-of-principle’ that this approach could be exploited for development of predictive biomarkers.

A previous study based on gene expression profiling subclassified PDAC into three subtypes: classical, quasimesenchymal (QM), and exocrine-like, exhibiting differences in patient outcomes and therapeutic responses (6). Only the former two subtypes could be detected among human PDAC cell lines. In contrast, *in silico* microdissection of gene expression profiles resolved only two tumor subtypes, “classical” (equivalent to the namesake subtype identified by Collisson et al.) and “basal” (28). These workers suggested that the exocrine-like phenotype identified by Collisson *et al.* may reflect contamination from normal tissue, and that the “mesenchymal” component of the QM subtype might be derived from tumor stroma (28). However, a recent subclassification based on transcriptomics has identified four subtypes: squamous (corresponding to QM), aberrantly differentiated endocrine e
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crine (ADEX) (corresponding to exocrine), pancreatic progenitor (classical) and immunogenic (5). Comparing our study to that of Collisson et al., there is not a simple relationship between any of the three subtypes we identified and the gene expression-defined classical or QM subtypes. For example, although some of our ATCC subtype 1 lines in the QM subtype, this association is not perfect, because Panc 03–27, a Subtype 2 line, is also categorized as QM. In addition, the classical subtype contains examples of both subtype 2 and 3 cell lines from our study. Moreover, because all the PDAC cell lines characterized in the Moffitt et al. study (the majority of which are included in our ATCC panel) were assigned to a single subtype, basal, our approach is clearly resolving subtypes not detected by these workers. Also of note, for the ATCC panel, the tyrosine phosphorylation-defined subtypes do not exhibit an obvious relationship to patterns of mutation of known cancer-associated genes, including KRAS and TP53. Similarly, the different TKCC subtypes were not related to KRAS and TP53 status. Consequently, tyrosine phosphorylation profiling is providing a novel subclassification that is not mirrored in those obtained via transcriptomics or consideration of mutation patterns.

The identification of phosphosite classifiers enabled us to determine that the three subgroups resolved in each cell line cohort share distinctive characteristics in terms of tyrosine phosphorylation patterns. For example, subtypes 1, 2, and 3 in each panel clearly exhibit low, medium and high phosphorylation, respectively, of BAIAP2 Y337, PKP3 Y84, PKP2 Y166, CTNND1 Y174, and CTNND1 Y904. Interestingly, BAIAP2 regulates cytoskeletal organization, whereas PKP3, PKP2, and CTNND1 localize to or regulate cell-cell adhesions. The functional role of the latter three proteins is of particular interest given that a Sleeping Beauty mutagenesis screen for genes associated with PDAC development demonstrated an important role for the adherens and tight junction signaling pathways, and implicated CTNND1 as a tumor suppressor gene in this malignancy (29). Indeed, low expression of CTNND1 in a PDAC patient cohort was associated with poor survival (29). An interesting possibility is that the differential phosphorylation of proteins associated with cell-cell adhesions reflects, at least in part, modulation of the axon guidance pathway, a key regulator of cell-cell junction integrity and signaling that is strongly implicated in PDAC (3). Supporting this hypothesis, the phosphorylation of MET and several Eph receptors, which represent upstream regulators of this pathway, is markedly elevated in subtype 3.

One pathway commonly altered in PDAC is mRNA splicing, with mutations in splicing factors SF3B1 or RBM10 occurring in 4–5% of patients (3, 30). Interestingly, our analysis identified an alternative mechanism for dysregulation of RNA metabolism in PDAC, with markedly increased tyrosine phosphorylation of SF3B1, as well as many other proteins involved in regulation of RNA metabolism, including hnRNPC, H and K, and RBMX, being detected in ATCC subtype 2. Because aberrant expression of particular hnRNPs is associated with a switch in mRNA splicing of particular oncogenes or tumor suppressors toward pro-tumorigenic isoforms (31, 32), an attractive hypothesis is that the enhanced tyrosine phosphorylation of these regulators of RNA metabolism also positively impacts on oncogenic signaling pathways. In support of this, tyrosine phosphorylation of hnRNPs K and A2B1 is known to affect either their binding to, or the translation of, target mRNAs (33–35). The cell lines that characterize ATCC subtype 2 therefore present powerful models to test this hypothesis, as well as potential therapeutic opportunities. With regard to the latter, the splicing factors themselves could be targeted with small molecule inhibitors such as spliceostatin A (36). Alternative strategies include inhibition of upstream tyrosine kinases or downstream oncogenic pathways. However, subtype-selective phosphorylation of proteins associated with RNA metabolism was not detected in the TKCC cohort, and the reason for this is not clear at present. A potential explanation is the heterogeneity of PDAC, such that cell lines with this signature are under-represented in the TKCC cohort. Another possibility, which is not mutually exclusive, is that in the corresponding TKCC subgroup, alternative mechanisms act to dysregulate mRNA processing and/or the pathways that lie downstream.

An additional conserved feature of the molecular taxonomy provided by our phosphotyrosine profiling was the increased tyrosine phosphorylation of numerous RTKs, including EGFR, ERBB2, ERBB3, MET, FGFR1, and AXL, as well as several EphA and EphB receptors, in subtype 3 of both cell line panels. Additional lines of evidence support important roles for many of these receptors in PDAC pathogenesis. For example, signaling by the EGFR is required early in PDAC development (37), focal amplifications of MET, FGFR1, and ERBB2 occur in PDAC genomes (4) and contrasting levels of AXL activation can occur at different metastatic sites from the same PDAC patient (38). In addition, high expression of EphA7 or EphB2 is associated with poor prognosis in PDAC (39, 40). However, our study provides the first demonstration that a PDAC subtype exists that exhibits co-activation of many of these RTKs. This finding has important implications for the rational design of targeted therapies. In other cancers, co-activation of multiple RTKs is associated with resistance to treatment with individual TKIs, necessitating use of combination approaches using several TKIs or multikinase inhibitors (17, 41). Consequently, although subtype 3 cell lines exhibit a greater inhibition of proliferation upon treatment with the EGFR TKI erlotinib than those from other subtypes, and the subtype 3 cell lines HPAC and PL45 are sensitive to knockdown of MET and AXL, respectively (42), it remains possible that combined targeting of subtype 3-enriched RTKs will achieve a greater efficacy. This possibility is supported by a recent study demonstrating that the efficacy of MEK1 inhibition in PDAC is limited by compensatory signaling by multiple RTKs to the PI3K pathway, and that drug resistance can be
overcome by combined targeting of EGFR, ERBB2, PDGFRα, and AXL (12). Importantly, the detailed information provided by our study will assist the design of therapeutic approaches directed at effective RTK blockade, and also fuel further studies that characterize crosstalk between particular RTKs in PDAC.

In the clinical trial that demonstrated the efficacy of erlotinib and gemcitabine co-treatment in PDAC, improved response to erlotinib was not associated with EGFR expression levels (27). This is consistent with our data indicating that EGFR levels are not consistently higher in subtype 3 cell lines, versus those from the other subgroups. Instead, it appears to be the tyrosine phosphorylation pattern associated with subtype 3, rather than EGFR expression, that is associated with increased sensitivity to this drug. Of note, although loss of p53 function is associated with a decreased dependence on the EGFR in PDAC (43), the increased sensitivity of ATCC and TKCC subtype 3 (the RTK subtypes) to erlotinib cannot be explained by TP53 mutational status. In addition, it is unrelated to K-Ras ‘addiction’ (26), because two out of three K-Ras-addicted cell lines in the ATCC cohort (Panc-08–13 and Capan-1) fall outside of the erlotinib-sensitive subtype. One component of the erlotinib sensitivity “signature” could be ERBB3 phosphorylation, because expression of this receptor is positively correlated with the erlotinib response of PDAC cell lines (44, 45). Although our study was directed toward identification of novel PDAC phenotypic subgroups and phosphorylation-based classifiers for these, rather than drug response signatures, our findings suggest that phospho-signatures of response to targeted therapies in PDAC could be formally derived and applied to patient stratification for therapy. This could be implemented via use of RPPAs or targeted MS approaches. Ultimately this could lead to improved management of patients suffering from this devastating disease.

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