



Purity Independent Subtyping of Tumors (PurIST), A Clinically Robust, Single-sample Classifier for Tumor Subtyping in Pancreatic Cancer

Naim U. Rashid^{1,2}, Xianlu L. Peng¹, Chong Jin^{1,2}, Richard A. Moffitt^{3,4}, Keith E. Volmar⁵, Brian A. Belt⁶, Roheena Z. Panni⁷, Timothy M. Nywening⁷, Silvia G. Herrera¹, Kristin J. Moore¹, Sarah G. Hennessey¹, Ashley B. Morrison¹, Ryan Kawalerski³, Apoorve Nayyar¹, Audrey E. Chang¹, Benjamin Schmidt⁷, Hong Jin Kim⁸, David C. Linehan⁶, and Jen Jen Yeh^{1,8,9}

ABSTRACT

Purpose: Molecular subtyping for pancreatic cancer has made substantial progress in recent years, facilitating the optimization of existing therapeutic approaches to improve clinical outcomes in pancreatic cancer. With advances in treatment combinations and choices, it is becoming increasingly important to determine ways to place patients on the best therapies upfront. Although various molecular subtyping systems for pancreatic cancer have been proposed, consensus regarding proposed subtypes, as well as their relative clinical utility, remains largely unknown and presents a natural barrier to wider clinical adoption.

Methods: We assess three major subtype classification schemas in the context of results from two clinical trials and by meta-analysis of publicly available expression data to assess statistical criteria of subtype robustness and overall clinical relevance. We then devel-

oped a single-sample classifier (SSC) using penalized logistic regression based on the most robust and replicable schema.

Results: We demonstrate that a tumor-intrinsic two-subtype schema is most robust, replicable, and clinically relevant. We developed Purity Independent Subtyping of Tumors (PurIST), a SSC with robust and highly replicable performance on a wide range of platforms and sample types. We show that PurIST subtypes have meaningful associations with patient prognosis and have significant implications for treatment response to FOLFIRINOX.

Conclusions: The flexibility and utility of PurIST on low-input samples such as tumor biopsies allows it to be used at the time of diagnosis to facilitate the choice of effective therapies for patients with pancreatic ductal adenocarcinoma and should be considered in the context of future clinical trials.

Introduction

Recent treatment advances, including FOLFIRINOX (1), gemcitabine plus nab-paclitaxel (2), and olaparib for *BRCA*-mutant patients (3), have provided patients and providers with better options. With the substantial progress in molecular subtyping for pancreatic cancer (4–9), there is now an opportunity to determine the optimal choice of therapy given a patient's molecular subtype and other

biomarker information, enabling “precision medicine” approaches in pancreatic cancer (10, 11).

Transcriptomic molecular subtyping in pancreatic cancer is currently an area of active development, where multiple subtyping schemas for pancreatic cancer have been proposed. For example, three molecular subtypes with potential clinical and therapeutic relevance were first described by Collisson and colleagues (5), leveraging a combination of cell line, bulk, and laser capture microdissected (LCM) patient samples: Collisson (i) quasi-mesenchymal (QM-PDA), (ii) classical, and (iii) exocrine-like. A subsequent study of patients with pancreatic cancer, based on more diverse pancreatic cancer histologies in addition to the most common pancreatic ductal adenocarcinoma (PDAC), found four molecular subtypes (4): Bailey (i) squamous, (ii) pancreatic progenitor, (iii) immunogenic, and (iv) aberrantly differentiated endocrine exocrine (ADEX). More recently, Puleo and colleagues describe five subtypes that are based on features specific to tumor cells and the local microenvironment (7). Maurer and colleagues performed LCM of both tumor and stroma and showed the contribution of each to the three schemas above (8). Finally, we have previously shown two tumor-intrinsic subtypes of PDAC (6), which we called Moffitt (i) basal-like, given the similarities with basal breast and basal bladder cancer, and (ii) classical, given the overlap with the Collisson classical subtype.

However, consensus regarding proposed subtypes for clinical decision making in PDAC has been elusive. In addition, each proposed schema utilized independent cohorts of patients to demonstrate clinical relevance. As a result, the generalizability, robustness, and relative clinical utility of each proposed subtyping schema remains unclear. Comparative evaluations of these proposed subtyping systems have been limited, partially due to the difficulty in curating and applying these diverse subtyping approaches in new datasets.

¹Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina. ²Department of Biostatistics, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina. ³Department of Biomedical Informatics and Pathology, Stony Brook University, Stony Brook, New York. ⁴Department of Pharmacological Sciences, Stony Brook Cancer Center, Stony Brook University, Stony Brook, New York. ⁵University of North Carolina-Rex Healthcare, Raleigh, North Carolina. ⁶Department of Surgery, University of Rochester, Rochester, New York. ⁷Department of Surgery, Washington University, Saint Louis, St. Louis, Missouri. ⁸Department of Surgery, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina. ⁹Department of Pharmacology, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina.

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N.U. Rashid and X.L. Peng are the co-first authors of this article.

Corresponding Authors: Jen Jen Yeh, University of North Carolina at Chapel Hill, 1150 POB, 170 Manning Drive, Chapel Hill, NC 27599-7213. Phone: 919-966-5221; Fax: 919-966-8806; E-mail: jjyeh@med.unc.edu; and Naim U. Rashid, University of North Carolina at Chapel Hill, 3104E McGavran-Greenberg Hall, 135 Daur Drive, Chapel Hill, NC 27599. Phone: 919-966-8150; naim@unc.edu

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Translational Relevance

Molecular subtyping for pancreatic cancer has made substantial progress in recent years, facilitating the optimization of existing therapeutic approaches to improve clinical outcomes in pancreatic cancer. We show that a tumor-intrinsic two-subtype schema is the most replicable and clinically robust across different subtype schemas, with basal-like subtype tumors showing resistance to FOLFIRINOX-based regimens in two independent clinical trials. Our results strongly support the need to evaluate molecular subtyping in treatment decision-making for patients with PDAC in the context of future clinical trials. We present PurIST, a clinically usable single-sample classifier that is robust and highly replicable across different gene expression platforms and sample collection types, and may be utilized in future clinical trials.

Toward this end, we perform, for the first time, a systematic interrogation of the aforementioned subtyping schemas, based on a meta-analysis of their clinical utility across a large number of independent cohorts in PDAC including two clinical trials with treatment response data. We demonstrate that a tumor-intrinsic two-subtype schema from Moffitt and colleagues (6) is robust and best explains overall survival (OS) and treatment response across multiple validation datasets. Given the clinically replicable performance of this tumor-intrinsic two-subtype schema, we have developed a single-sample classifier (SSC) that we call Purity Independent Subtyping of Tumors (PurIST) to perform subtype calling for clinical use. We show that PurIST performs well on multiple gene expression platforms including microarray, RNA sequencing (RNAseq), and NanoString. In addition, given the preponderance of nonsurgical biopsies in the neoadjuvant and metastatic settings, we demonstrate its clinical utility for small sample volumes using a matched cohort of patients with bulk, archival, and fine-needle aspirations (FNA) samples. Finally, we show the stability of PurIST-predicted subtypes before and after treatment, and that PurIST basal-like subtype tumors are associated with treatment resistance to FOLFIRINOX, strongly supporting the need to incorporate subtyping into clinical trials of patients with PDAC.

Materials and Methods

Public datasets

Archival data were obtained from public sources (Supplementary Table S1; Fig. 1). For the public datasets, expression was used “as-is” with respect to the original publication, that is, RNAseq data were not realigned and gene-level expression estimates were provided in terms of fragments per kilobase per million reads (FPKM) or transcripts per million (TPM), depending on the study.

Sample collection

Deidentified bulk and FNA samples (“Yeh_Seq” dataset; Supplementary Table S1) were collected from the Institutional Review Board (IRB)-approved University of North Carolina Lineberger Comprehensive Cancer Center Tissue Procurement Core Facility after IRB exemption in accordance with the U.S. Common Rule and were flash frozen in liquid nitrogen. FNA samples were collected *ex vivo* at the time of resection. The FNA technique used mirrors standard cytopathology procedures, where three passes were performed using a 22-gauge needle. Palpation was used to localize the tumor. Samples were

frozen in either PBS or RNAlater (Millipore Sigma). FFPE samples were prepared, hematoxylin and eosin stained, and reviewed by a single pathologist (K.E. Volmar) who was blinded to the results. See Supplementary Materials and Methods for data processing and analysis of Yeh_Seq samples. RNAseq (GSE131050) and NanoString (GSE131051) data generated from these samples are deposited in Gene Expression Omnibus (GEO).

Sample inclusion for consensus clustering analysis and PurIST training

Each collected public dataset was subjected to sample filtering to retain samples for consensus clustering (CC)-based subtype calling by the Collisson, Bailey, and Moffitt schemas, with criteria specified in Supplementary Table S1. We implemented the methods utilized in the original publication pertaining to each schema (Supplementary Materials and Methods). When prior subtype calls were available, the original published calls were used. Specifically, in PACA_AU_seq and PACA_AU_array, the original Bailey subtype calls were used; in TCGA_PAAD, the Collisson and Bailey calls were used.

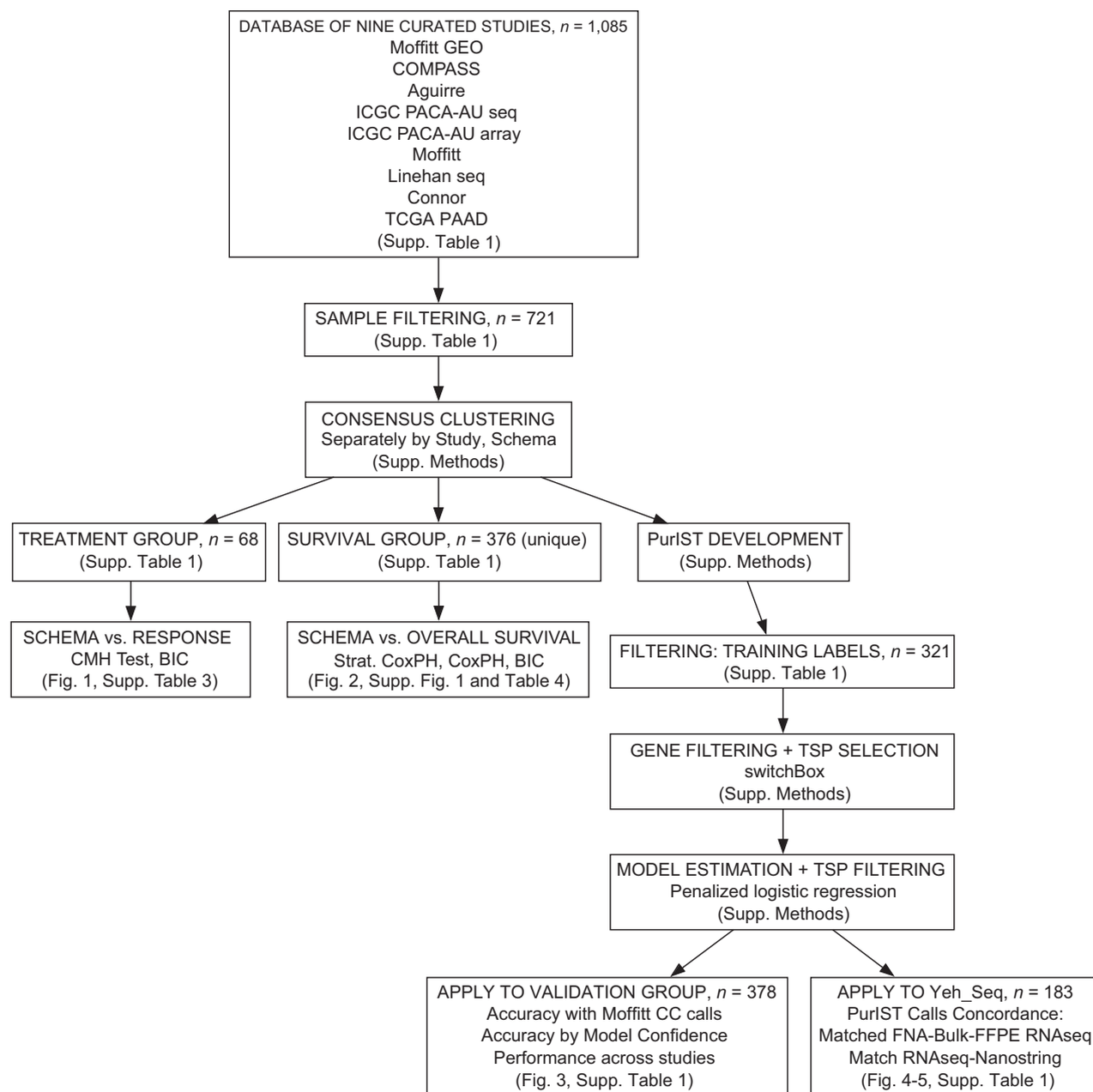
For treatment response and survival analysis, samples with available clinical and RNAseq data were used. Specifically, for the pooled survival analysis, samples from the following datasets with RNAseq data and CC calls were utilized: Linehan, Moffitt_GEO_array, PACA_AU_seq, PACA_AU_array, and TCGA_PAAD (survival group, Supplementary Table S1). Duplicated samples in PACA_AU_seq and PACA_AU_array datasets were only used once, with the subtypes called in PACA_AU_array used when mismatches of subtype calls were found between the two datasets. To train PurIST, Moffitt schema CC calls from the datasets in the training group (Aguirre, Moffitt_GEO_array, and TCGA_PAAD; Supplementary Table S1) were utilized. These samples were further filtered to provide final training labels for the PurIST algorithm (Supplementary Tables S1 and S2) by dropping poorly clustered samples on the clustered dendrogram in each dataset based on visual inspection. We considered these filtered calls as “training labels.” Model training for PurIST is described in the Supplementary Materials and Methods.

Results

The Moffitt tumor-intrinsic two-subtype schema has important implications for treatment response

To evaluate the potential impact of molecular subtypes on treatment response, we utilize transcriptomic and treatment response data from two independent clinical trials, and perform a systematic analysis of treatment response with respect to CC calls from each of the three different subtyping schemas (Supplementary Materials and Methods, Supplementary Fig. S1) for PDAC: Collisson, Bailey, and Moffitt (4–6). We first examined the association of the subtypes from each schema with treatment response using patient samples from a promising phase Ib trial by Nywening and colleagues (“Linehan,” Linehan_seq dataset; Supplementary Tables S1 and S2) of FOLFIRINOX in combination with a CCR2 inhibitor (PF-04136309) in patients with locally advanced PDAC, where an objective response was seen in 49% of patients (12). Enrolled patients had no prior treatment, and underwent core biopsies prior to the start of therapy. Twenty-eight patients with RNAseq and treatment data were available for analysis.

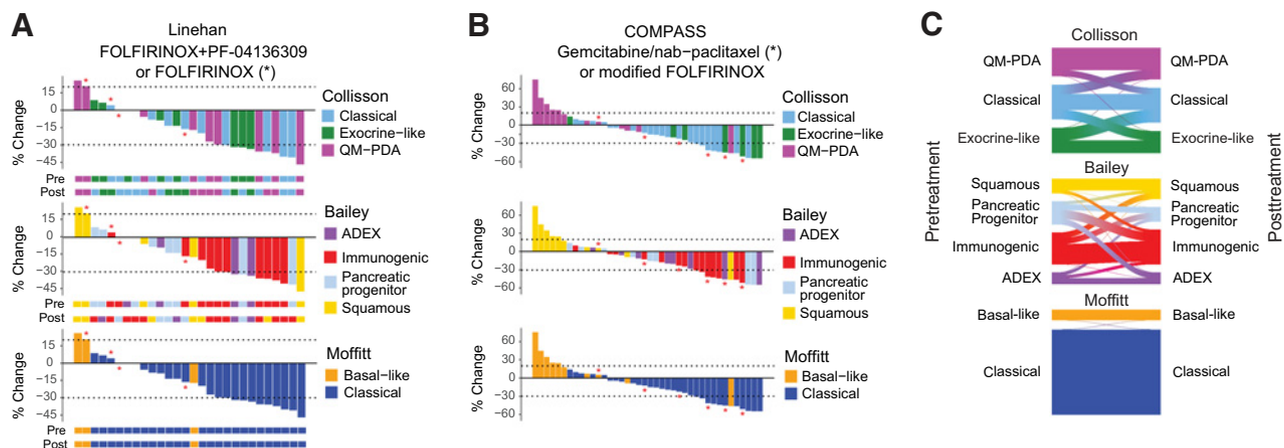
We found a significant overall association between categorical treatment response (based on RECIST 1.1 criteria) and pretreatment subtype classifications from the Moffitt schema ($P = 0.0117$; Supplementary Table S3), where basal-like tumors showed no response to

**Figure 1.**

Overall study workflow outlining the analyses performed, including the training and validation steps of the PuriST model. Supp., supplementary; CMH, Cochran-Mantel-Haenszel test; Strat. Cox PH, stratified Cox proportional hazards model.

FOLFIRINOX alone or FOLFIRINOX plus PF-04136309 after stratifying by arm [overall response rate (ORR) = 0%; disease control rate (DCR) = 33%; Supplementary Table S3, Fig. 2A, generalized Cochran-Mantel-Haenszel test], whereas classical tumors showed a much stronger response overall (ORR = 40%; DCR = 100%). In contrast, we were unable to identify a relationship between subtype and treatment response under the Collisson ($P = 0.428$) and Bailey ($P = 0.113$) schemas (Fig. 2A; Supplementary Table S3). As the sample size in this phase Ib trial ($n = 28$ patients) was small, we similarly reanalyzed the COMPASS trial results ($n = 40$ patients) in the context of the three subtyping schemas.

Patients enrolled in COMPASS underwent core-needle biopsies and were treated with one of two standard first-line therapies, modified-FOLFIRINOX or gemcitabine plus nab-paclitaxel. Collected patient samples in COMPASS underwent laser capture microdissection (LCM) followed by whole genome sequencing and RNAseq. Subtypes for each schema were determined as mentioned previously. Similar to our findings in the Linehan phase Ib trial, we found a significant association between the Moffitt two subtype schema with categorical treatment response stratifying by arm ($P = 0.00098$, generalized Cochran-Mantel-Haenszel test), where the basal-like subtype had much lower response to either treatment

**Figure 2.**

Subtype performance in discriminating treatment response. **A** and **B**, Waterfall plots showing the percent change (% change) in size of tumor target lesions from baseline in the context of the Collisson, Bailey, and Moffitt schemas in the Linehan (**A**) and COMPASS trials (**B**). +20% and -30% of size change are marked by dashed lines. Bar colors denote respective subtype calls of pretreatment samples. RECIST treatment response classification based on these values are given in Supplementary Table S3. **A**, Colored tracks below each plot denote the subtype calls in pre- and posttreatment. Patients marked with * were treated with FOLFIRINOX. The remainder of patients were treated with FOLFIRINOX + PF-04136309. **B**, Patients marked with * were treated with gemcitabine/nab-paclitaxel (GP)-based therapy. The remainder of patients were treated with modified FOLFIRINOX (m-FOLFIRINOX). **C**, Sankey plots showing transitions in subtype pre- and posttreatment in the Linehan trial in the context of the Collisson, Bailey, and Moffitt schemas.

(ORR = 10%; DCR = 50%) relative to the classical subtype (ORR = 36.7%; DCR = 100%). We also found significant associations between treatment response and the subtypes from the Collisson ($P = 0.0024$) and Bailey ($P = 0.0067$) schemas. However, we notably observe that the Bailey squamous subtype strongly overlaps with the Moffitt basal-like subtype, and the remaining nonsquamous Bailey subtypes appear to overlap strongly with the Moffitt classical subtype (Fig. 2B, Cohen Kappa = 1.0, $P = 2.54 \times 10^{-10}$). We similarly found that the Collisson QM-PDA and the remaining non-QM-PDA subtypes correspond strongly with the Moffitt basal-like and classical subtypes, respectively (Fig. 2B, Cohen Kappa = 0.875, $P = 2.44 \times 10^{-8}$), a fact also mirrored in the Linehan trial.

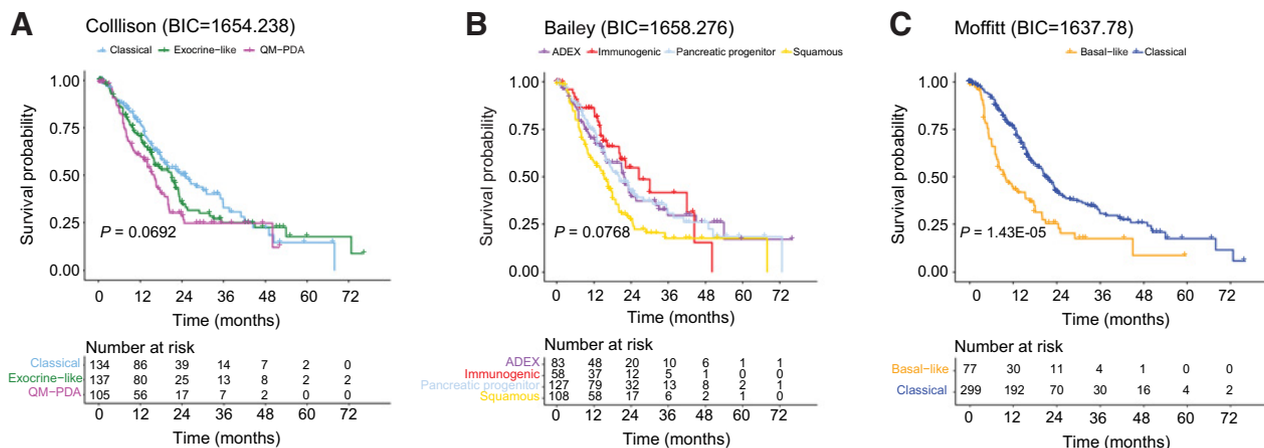
Given these observations, we formally evaluated the relative clinical utility of each subtyping system using nonnested model selection criteria such as Bayesian information criterion (BIC; ref. 13). Briefly, such criteria evaluate model fit relative to the complexity of the model, as models with more predictors (subtypes) may simply have better fit due to overfitting, and also may contain excess predictors (additional subtypes) that do not contribute meaningfully in differentiating clinical outcomes. The model with the lowest BIC in a series of competing candidate models is preferred in statistical applications, and is agnostic to the magnitude of the difference (14). Considering response as a continuous outcome (% change in tumor volume), we find that the Moffitt schema had the best (lowest) BIC score in both datasets (Linehan BIC = 247.37, COMPASS BIC = 378.75, two-way ANOVA model; Supplementary Table S3), compared with the Collisson (Linehan BIC = 254.63, COMPASS BIC = 382.8) and Bailey (Linehan BIC = 250.75, COMPASS BIC = 385.66) schemas. This result similarly held if we considered response as a categorical variable (ordinal regression model; Supplementary Table S3). This finding is also reflected among the non-QM-PDA and nonsquamous subtypes (Fig. 2A and B; Supplementary Table S3), where little difference in response can be seen between these subtypes. Our results using BIC suggest that the additional subtypes found in the Collisson and Bailey schemas do not demonstrate additional benefit in differentiating treatment response over the Moffitt two-subtype schema. Taken

together, these results suggest that the Moffitt basal-like and classical subtypes strongly and parsimoniously explains treatment response relative to other schemas in both clinical trials.

The Linehan phase Ib trial captured both pre- and posttreatment biopsies, providing a unique opportunity to evaluate the stability of molecular subtypes after treatment. As pre- and posttreatment biopsies are unlikely to be obtained from the same location, these samples may also provide an opportunity to evaluate intrapatient tumor heterogeneity. Interestingly, we found strong stability in the Moffitt schema subtypes in pre- and posttreatment biopsies (Fig. 2C; Cohen Kappa = 1.0; $P = 2.54 \times 10^{-10}$), suggesting that not only may there be less tumor-intrinsic subtype heterogeneity within a tumor, but also that the Moffitt schema subtypes are not affected by treatment, either with FOLFIRINOX or with the addition of the CCR2 inhibitor. In contrast, we found higher rates of switching in Collisson subtypes pre- to posttreatment (Fig. 2C; Supplementary Fig. S2), where changes in the exocrine-like and classical subtypes were more common. Similarly, the nonsquamous Bailey subtypes appeared to show the highest rate of subtype switching pre- and posttreatment, with the ADEX subtype demonstrating the highest rate of switching among these subtypes (Supplementary Fig. S2). It is unclear whether there is any clinical significance to such subtype transitions. Prior studies have suggested that the Bailey ADEX, Bailey immunogenic, and Collisson exocrine-like subtypes are confounded by tumor purity in contrast to the Moffitt subtypes (7–9), which may explain some of the increased heterogeneity in subtypes pre- and posttreatment in these schemas. In contrast, the Collisson QM-PDA and Bailey squamous subtypes, which were shown to overlap strongly with the Moffitt basal-like subtype, were observed to be much more stable between the two time points.

The tumor-intrinsic two-subtype schema strongly and replicably differentiates patient survival across multiple studies

Given the paucity of available genomic data in the context of treatment response in PDAC, we also perform a meta-analysis of five independent patient cohorts with OS data available: Linehan_seq, Moffitt GEO array (GSE71729), ICGC PACA_AU array, ICGC

**Figure 3.**

Subtype performance in predicting patient prognosis in pooled datasets from the survival group (Supplementary Table S1). Kaplan-Meier plots of OS in the context of the subtyping schemas of Collisson (A), Bailey (B), and Moffitt (C). Log-rank P values for overall association were determined from stratified Cox proportional hazards models, where dataset was used as a stratification factor to account for variation in baseline hazard across studies. BIC was calculated to compare the three subtyping schemas.

PACA_AU seq, and TCGA PAAD (survival group; Supplementary Tables S1 and S2). To determine the potential replicability of the different subtyping schemas (Collisson, Bailey, Moffitt) in differentiating clinical outcomes, we utilized CC subtype calls from each schema.

We find that the Moffitt tumor-intrinsic two-subtype schema reliably differentiates survival across individual datasets (Supplementary Fig. S1; Supplementary Table S4), showing significant associations with OS in the majority of individual studies in contrast to other schemas. After pooling datasets, we found that patients with Moffitt basal-like subtype tumors have significantly worse prognosis compared with the Moffitt classical subtype (Fig. 3C, stratified HR = 1.98, $P < 0.0001$, stratified Cox proportional hazards model). We also observed similar trends in the Bailey squamous and Collisson QM-PDA subtypes relative to other subtypes in the same schemas (Fig. 3A and B), mirroring our treatment response results from the previous section (Fig. 2A and B). However, overall subtype-specific survival differences were most pronounced within the two-subtype schema across studies (Supplementary Table S4), compared with the Collisson ($P = 0.069$) and Bailey ($P = 0.076$) schemas. Moreover, we find that nonsquamous subtypes in the Bailey schema have very similar OS to one another (Fig. 3B), where a direct overall comparison of these subtypes showed no statistically significant differences in survival in our pooled dataset (immunogenic vs. ADEX stratified HR = 1.07, pancreatic progenitor vs. ADEX HR = 1.01, overall $P = 0.82$). We find a similar result when comparing survival among patients from the non-QM-PDA subtypes in the Collisson schema in the pooled data (Fig. 3A; exocrine-like vs. classical stratified HR = 1.17; $P = 0.344$).

In our pooled dataset, strong correspondence was again found between the Bailey squamous, Collisson QM-PDA, and Moffitt basal-like subtypes, and between the Moffitt classical subtype and the remaining subtypes in the Bailey (Cohen Kappa = 0.56, $P = 0$; Supplementary Fig. S3A) and Collisson (Cohen Kappa = 0.4, $P = 0$; Supplementary Fig. S3B) schemas. In TCGA PAAD, where estimates of tumor purity were available, Moffitt classical patients that were also classified as QM-PDA in the Collisson schema had much lower tumor purity than other samples ($P = 0.0016$; Supplementary

Fig. S3C). The Bailey ADEX and immunogenic samples also had lower tumor purity, regardless of whether they were called Moffitt classical or basal-like (Supplementary Fig. S3D). These findings are similar to other studies (7–9), and suggest that the discordance in subtype assignment between schemas may be driven by tumor purity.

To determine the best fitting model for OS, we calculate BIC with respect to the stratified Cox proportional hazards model pertaining to each schema. Similar to our analysis of treatment response, we find that the Moffitt two-subtype schema has the best (lowest) BIC and therefore has the best and most parsimonious fit to the pooled survival data (Fig. 3; Supplementary Table S4). We also find this to be the case in the majority of individual studies, replicated across each of our validation datasets (Supplementary Table S4). These results reflect our finding that no difference in OS can be observed among the Collisson non-QM-PDA and Bailey nonsquamous subtypes in our pooled analysis. In total, these findings support the conclusion that the Moffitt two-subtype schema strongly and parsimoniously explains differences in OS, compared with alternate subtyping schemas. Our results further suggest that the additional subtypes found in the Collisson and Bailey schemas do not demonstrate additional clinical benefit in terms of predicting OS relative to the simpler Moffitt two-subtype schema, based on BIC and direct statistical comparison of the Collisson non-QM-PDA and Bailey nonsquamous subtypes. Given the robustness and highly replicable clinical utility of the Moffitt schema, we next developed a SSC based on this tumor-intrinsic two-subtype schema to avoid reliance on CC-based analysis.

PurIST SSC

The ability to resolve and assign subtypes via clustering is limited when applied to individual patients. Reclustering new samples with existing training samples may also change existing subtype assignments. Thus, we developed a robust SSC, PurIST, to predict subtype in individual patients, based on our three largest bulk gene expression datasets (TCGA PAAD, Aguirre Biopsies, and Moffitt GSE71729, training group; Supplementary Table S1). A key element of our method includes the utilization of tumor-intrinsic genes previously identified (6) to avoid the possible confounding of tumor gene expression with those from other tissue types. For model training,

we designated training labels (Supplementary Materials and Methods). We use rank-derived quantities as predictors in our final SSC model instead of the raw expression values, utilizing the k Top Scoring Pair (kTSP) approach to generate these predictors (Supplementary Materials and Methods). The motivation of this approach is that while the raw values of gene expression may be on different scales in different studies, their relative magnitudes can be preserved by ranks.

We find that this type of rank transformation of the raw expression data has several advantages. First, a single predictor (TSP) only depends on the ranks of raw gene expression of a gene pair in a sample. Hence, its value is robust to overall technical

shifts in raw expression values (i.e., due to variation in sequencing depth), and, as a result, is less sensitive to common between-sample normalization procedures of data preprocessing (15–17). Second, it simplifies data integration over different training studies as data are on the same scale. Finally, prediction in new patients is also simplified, as normalizing new patient data to the training set is no longer necessary, which may further affect the accuracy of model predictions (16).

Development and external validation of PurIST classifier

We apply a systematic procedure (Supplementary Materials and Methods) implementing the above approach to derive our PurIST

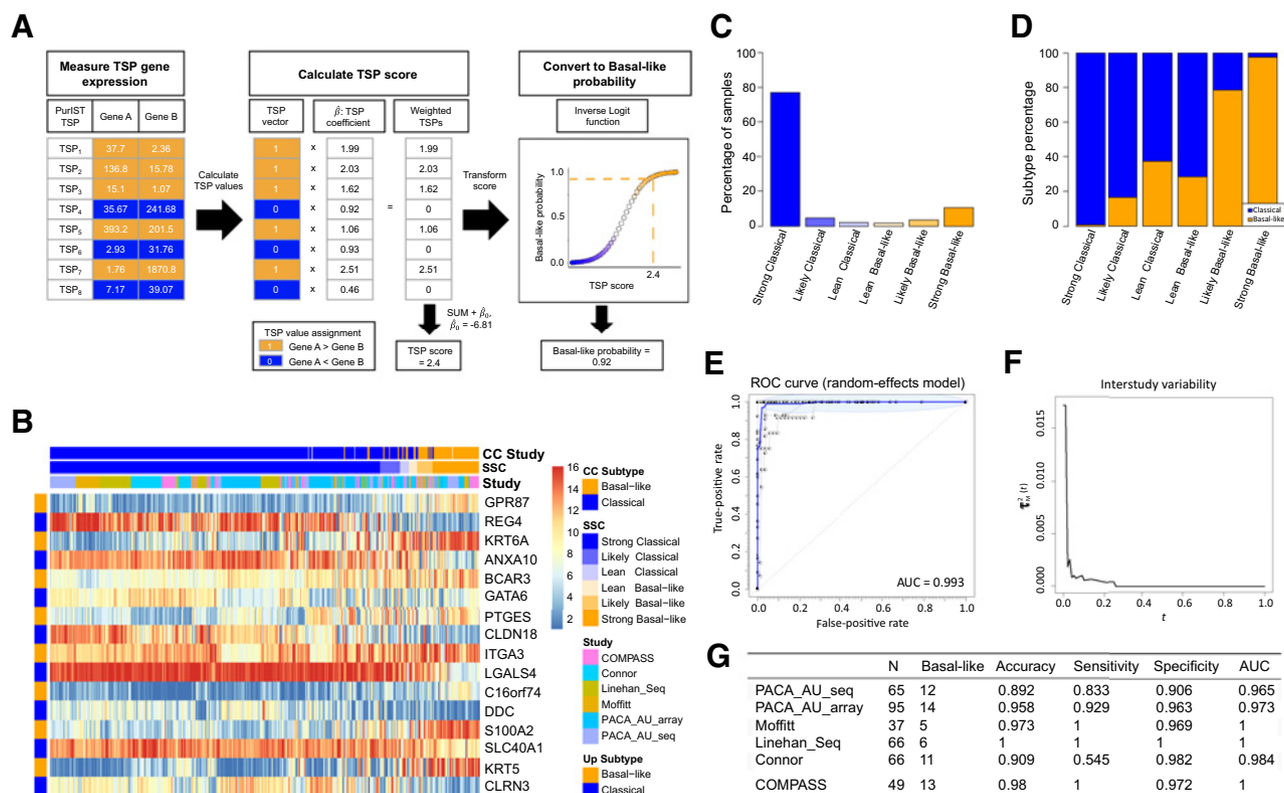


Figure 4.

Development and validation of the PurIST SSC classifier. **A**, Overview of PurIST prediction procedure. Gene expression for genes pertaining to each PurIST TSP is first measured in a new sample. Values are assigned for each TSP given the relative expression of each gene in the TSP (1 if gene A > gene B expression in the pair, 0 otherwise). Given the set of estimated PurIST TSP coefficients, a TSP score is calculated by summing the product of each TSP and its corresponding TSP coefficient, adjusting for the model intercept. This value is finally transformed into a predicted probability of belonging to the basal-like subtype for classification (inverse logit function). **B**, Heatmap of expression values pertaining to PurIST genes and patients from six validation studies belonging to the validation group (Group Membership, Supplementary Table S1). Columns are ordered by PurIST SSC-predicted basal-like probability. Genes pertaining to each TSP are presented in order along the rows, where genes with higher expression in the basal-like subtype in the pair are labeled with an orange bar on the left track and blue otherwise. CC subtype and training labels used for PurIST SSC training (white bars indicate not utilized for training) show strong correspondence with the SSC-predicted subtypes. Switching in relative gene expression within each gene pair can be observed with respect to subtype. Expression values across PurIST genes were rank transformed to equalize the expression scales across studies. SSC predicted basal-like probabilities were separated into subclasses to illustrate the level of model confidence in prediction (Materials and Methods). **C**, Barplot of SSC confidence levels categorizing the predicted basal-like class probabilities indicate that the majority of predictions are highly confident, with few in the “likely” ranges of each (predicted probabilities between 0.4 and 0.6). **D**, Misclassification rates among higher confidence predictions (strong classical/basal) in either subtype are very low. Shading of bars indicate the relative percentage of each CC subtype in a given prediction category. Lower confidence predictions (Likely/Lean categories), as expected, have higher misclassification rates with respect to CC subtypes but are less frequent overall. **E**, Consensus ROC curve derived across all six validation studies (blue) in addition to ROC curves pertaining to each individual study (gray) are presented. The consensus AUC is relatively high at 0.993, indicating excellent prediction performance utilizing the CC subtypes as ground truth. **F**, Interstudy variability curve indicates low variability (y-axis) across studies with respect to various basal-like predicted probability thresholds (x-axis) utilized to classify subjects at basal or classical. At the standard basal-like probability threshold of 0.5, the between study variability in ROC curves is very low, suggesting strong replicability in classification performance of the PurIST SSC across studies. **G**, Table showing individual metrics assessing PurIST performance in recapitulating CC subtypes in each validation dataset.

model for prediction in the tumor-intrinsic two-subtype schema given the training labels (Supplementary Materials and Methods) and rank transformed predictors for each training samples. The selected eight gene pairs (TSP), fitted model, and model coefficients are given in Supplementary Table S5. **Figure 4A** (Supplementary Materials and Methods) describes the validation that is performed in a hypothetical new patient by computing the values of each of the eight selected TSPs in that patient, where a value of 1 is assigned if the first gene in a TSP, gene A, has greater expression than the second gene, gene B, in that patient (and assigned 0 value otherwise). These values are then multiplied by the corresponding set of estimated TSP model coefficients, summing these values to get the patient "TSP Score" after correction for estimated baseline effects. This score is then converted to a predicted probability of belonging to the basal-like subtype, where values greater than 0.5 suggest basal-like subtype membership and the classical subtype otherwise.

To assess the quality of our prediction model, we evaluate the cross-validation error of the final model in our training group. We find that the internal leave-one-out cross-validation error for PurIST on the training group is low (3.1%). To validate this model, we apply it to the validation group datasets (Supplementary Table S1; **Fig. 4**) and determine whether PurIST predictions recapitulate the CC subtypes in each study. We find that pooled validation samples strongly segregate by CC subtype when sorted by their predicted basal-like probability, despite diverse studies of origin (**Fig. 4B**). These suggest that our methodology avoids potential study-level batch effects. The relative expression of classifier genes within each classifier TSP (paired rows, **Fig. 4B**) strongly discriminates between subtypes in each sample, forming the basis of our robust TSP-oriented approach for subtype prediction (Supplementary Fig. S4). We also find that, visually, predicted subtypes from PurIST have strong correspondence with independently determined CC subtypes. Overall, the PurIST classifier predicted subtypes with high levels of confidence (**Fig. 4C**), with most basal-like subtype predictions having predicted basal-like probabilities >0.9 (strong basal-like), and most classical subtype predictions with predicted basal probabilities of <0.1 (strong classical). Among these high confidence predictions, the majority of these calls corresponded with subtypes obtained independently via CC (**Fig. 4D**). Lower confidence calls (likely/lean basal-like/classical categories of prediction) had higher rates of misclassification, although these less confident calls were more rare in our validation datasets (**Fig. 4C**).

To evaluate the overall classification performance of PurIST across studies, we apply a nonparametric meta-analysis approach to obtain a consensus ROC curve based on the individual ROC curves from each validation study (18). We found that the overall consensus AUC is high, with a value of 0.993. ROC curves from individual studies were also consistent (**Fig. 4E**). In addition, we find that the estimated interstudy variability of these ROC curves with respect to predicted basal-like probability threshold t is low overall, with relatively higher variance at low thresholds and almost no variability at our standard threshold of 0.5 or greater (**Fig. 4F**). These reflect the similarity of individual ROC curves seen in **Fig. 4E**. We find that within our validation datasets, the prediction accuracy rates were in general 90% or higher, and individual study AUCs were 0.95 or greater (**Fig. 4G**). Furthermore, sensitivities and specificities were often high and in some cases equal to 1, reflecting near perfect classification accuracy. These results suggest that PurIST is robust across multiple datasets and platforms and reca-

pitulates the subtypes independently obtained via CC, which we have shown to have high clinical utility.

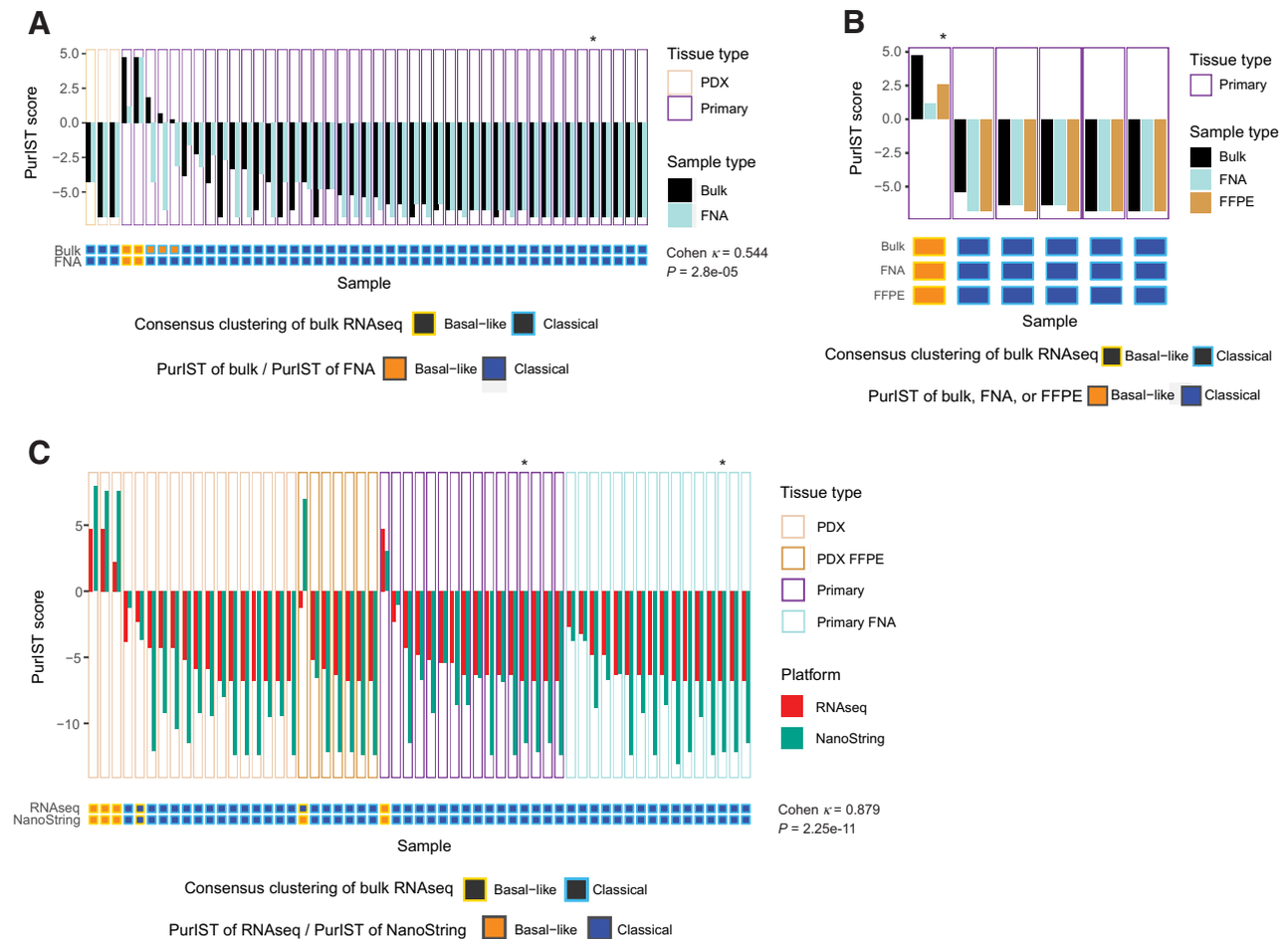
Replicability of PurIST in archival formalin-fixed and paraffin embedded and FNA samples

Because frozen bulk tumor samples are not commonly available in routine clinical practice, we next looked at the replicability of PurIST predictions across sample types that are more widely collected in clinical practice. Notably, nearly all preoperative and metastatic biopsies are obtained using either FNA or core biopsy techniques. Prior studies have shown the feasibility of performing RNAseq on core biopsies (11) and endoscopic ultrasound guided FNAs, both of which are commonly utilized in the diagnosis of pancreatic cancer (19). We therefore evaluated the performance of PurIST in both formalin-fixed and paraffin embedded (FFPE) and FNA samples.

Among 47 pairs of matched FNA and bulk samples that passed quality control (Yeh_Seq dataset; Supplementary Materials and Methods), we found significant agreement between the PurIST subtype calls of the matched FNA and bulk samples (Cohen Kappa = 0.544; $P = 2.8e-05$; **Fig. 5A**; Supplementary Table S1). Only three pairs of samples (6.4%) show disagreement in subtype calling results using PurIST. CC calls of the bulk samples are also shown as a comparison. We performed a similar evaluation with tumors that we had matched FFPE, FNA, and bulk samples available (**Fig. 5B**; Supplementary Table S1). We found complete agreement among PurIST subtype predictions among FFPE, FNA, and bulk samples in patients that had all three sample types available (five sets total), further supporting that PurIST is robust across different sample preparations. We also found that the genes pertaining to PurIST TSPs are comparatively less variable than genes not designated as tumor-intrinsic (Supplementary Fig. S5). For example, PurIST TSP genes, originally selected from our tumor-intrinsic gene list, have significantly higher Spearman correlation between sample types than Bailey immunogenic ($P = 0.0149$) or ADEX genes ($P = 0.0083$; Supplementary Fig. S5), using a permutation test (Supplementary Materials and Methods). The stability of TSP genes across sample types, support their robustness and their ability to identify tumor-intrinsic signals in samples that may be confounded by low-input or degradation.

Replicability of PurIST predictions on a NanoString platform

RNAseq assays in Clinical Laboratory Improvement Amendments (CLIA)-certified laboratories are still in their infancy. Thus, we evaluated the performance of PurIST on samples using NanoString nCounter, a gene expression quantification system that directly quantifies molecular barcodes. This platform has been widely used in cancer molecular subtyping (20), and is more widely available in CLIA-certified laboratories. In samples with both RNAseq and NanoString platform expression data available, we evaluated the consistency between subtype calls based on their RNAseq and NanoString expression data using PurIST-n (**Fig. 5C**; Supplementary Table S5; Supplementary Materials and Methods). This updated classifier is trained in a manner similar to PurIST, with the exception that genes were limited to those in common between the two platforms, as a more limited set of genes were available for our NanoString probeset. We found that there was strong agreement between PurIST-n calls in 51 patients with matched RNAseq/NanoString samples (Cohen Kappa = 0.879; $P = 2.25e-11$), where only one sample showed disagreement in its PurIST-n call. This discrepancy may be due to the relatively lower read count in the RNAseq sample for this patient. In addition, it is noteworthy that the PurIST-n call for this sample is a low confidence call ("lean classical"). These results

**Figure 5.**

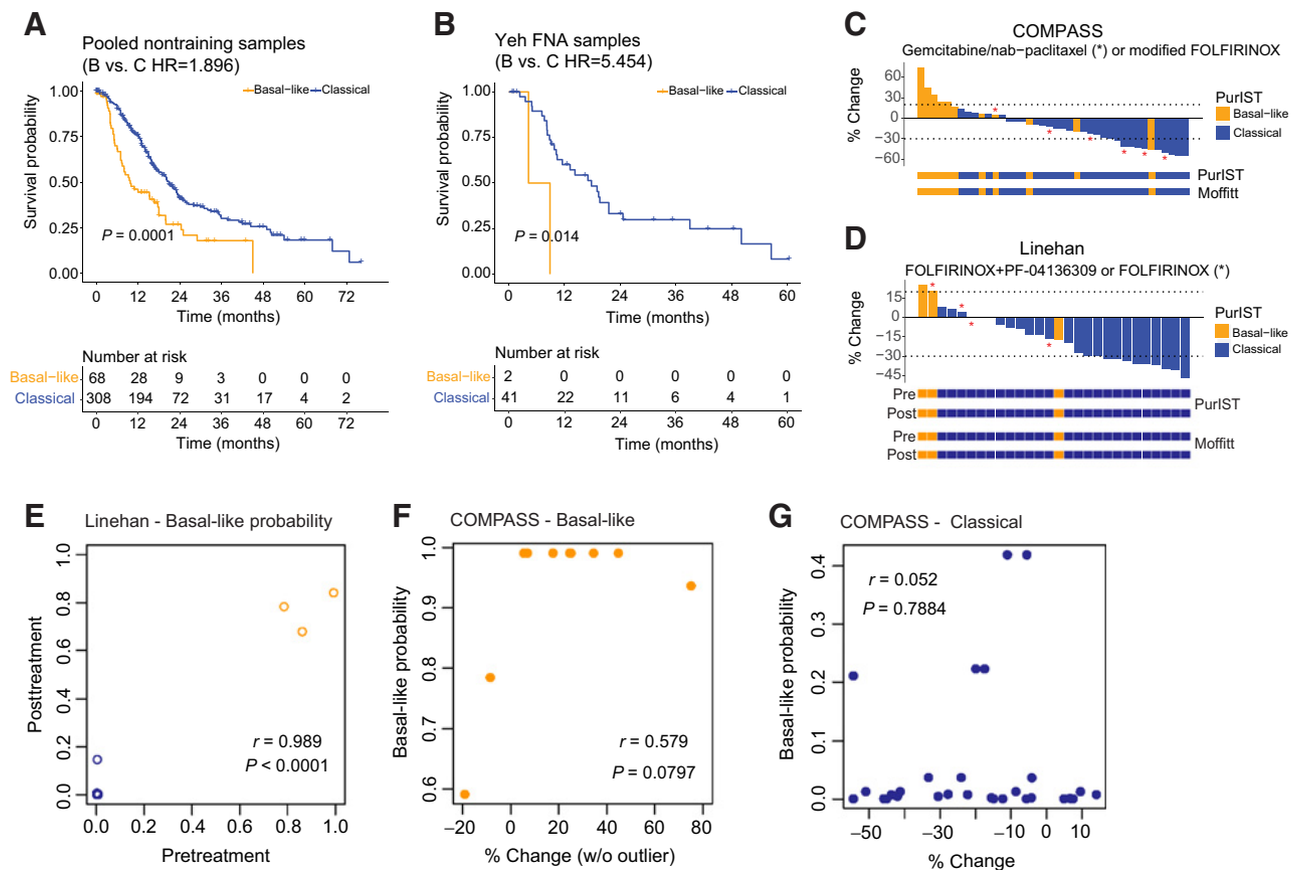
Comparison of CC and PurIST performance on different sample collections and gene expression platforms. **A**, Comparison of subtyping results using PurIST between matched FNA and bulk RNAseq samples. PurIST scores (estimated log-odds of a sample being basal-like vs. classical, Supplementary Materials and Methods) are in the upper waterfall plot (solid dark slate gray, bulk; solid light blue, FNA) and inferred subtypes are in the corresponding boxes below (solid blue, classical; solid orange, basal-like). Bulk RNAseq CC results are indicated by blue (classical) and orange (basal-like) square borders. *, ampullary carcinoma. **B**, Subtyping results using PurIST across matched FNA, FFPE, and bulk RNAseq samples. PurIST scores are in the top waterfall plot (solid dark slate gray, bulk; solid light blue, FNA; solid brown, FFPE) and inferred subtypes are in the corresponding boxes below (solid blue, classical; solid orange, basal-like). As a comparison, bulk RNAseq CC results are indicated by blue (classical) versus orange (basal-like) square borders. “*”, NanoString. **C**, Subtyping results using PurIST (and PurIST-n for NanoString) between matched RNAseq and NanoString samples. The PurIST(-n) scores (estimated log odds of a sample being basal-like vs. classical, Materials and Methods) are in the top waterfall plot (solid red, RNAseq; solid green, NanoString). Inferred subtypes are in the corresponding boxes below (solid blue, classical; solid orange, basal-like). Bulk RNAseq CC results are indicated by blue (classical) and orange (basal-like) square borders. **A-C**, Border colors of waterfall plots indicate sample origin. Cohen kappa coefficients and *P* values measure the agreement on subtype calls between bulk and FNA in **A**, and RNAseq and NanoString in **C**.

support the replicability of PurIST on the NanoString platform and suggest that NanoString may be more robust at overcoming the hurdles of low input or degraded samples.

Applicability of PurIST to treatment decision making

We next evaluated the potential utility of using PurIST for clinical decision making. In basal-like and classical samples that were classified by PurIST, we found significant survival differences in both the pooled public (with all training group samples removed) and the Yeh_Seq FNA datasets, with basal-like samples showing shorter OS (Fig. 6A and B; Supplementary Fig. S6; Supplementary Table S4). We then looked at the relevance of PurIST to treatment response in the COMPASS and Linehan trials (Fig. 6C and D). PurIST recapitulated

48 of 49 PDAC subtype calls compared with the previous CC-based calls in the COMPASS dataset, and 66 of 66 subtype calls in the Linehan dataset (Supplementary Tables S1 and S2). Only one patient with a CC classical tumor was called basal-like by PurIST and had stable disease (SD, % change >−30% and <20%) in the COMPASS trial. Notably, the only PR seen in a PurIST basal-like tumor was in a patient with an unstable DNA subtype (10). In agreement with our CC analysis (Fig. 2B), we find that PurIST-predicted subtype tumors had similar associations with treatment response (Fig. 6C and D; Supplementary Table S3). We also found no change in PurIST subtype or the confidence of the call after treatment, suggesting that PurIST tumor subtypes are unchanged after treatment with FOLFIRINOX ± PF-04136300 (Fig. 6D and E). Finally, after excluding the sample with an

**Figure 6.**

Clinical relevance of PurIST SSC in datasets belonging to the treatment group. **A** and **B**, Kaplan-Meier plots of OS in pooled datasets (**A**) belonging to the survival group minus datasets belonging to the training group and Yeh Seq FNA samples (**B**). *P* value and HRs for overall association were estimated by stratified Cox proportional hazards model in **A**, where dataset of origin was used as a stratification factor. **C** and **D**, Waterfall plots showing the percent change (% change) in size of tumor target lesions from baseline in the context of PurIST subtypes in the COMPASS (**C**) and Linehan trials (**D**). +20% and -30% of size change are marked by dashed lines. **C**, Bar colors denote PurIST subtype calls of the patient tumors. Patients marked with * were treated with gemcitabine/nab-paclitaxel (GP)-based therapy, and the rest were treated with modified FOLFIRINOX (m-FOLFIRINOX). **D**, Bar colors denote PurIST subtype calls of pretreatment samples. Colored tracks below to compare subtype calls for samples pre- and posttreatment of PurIST subtyping and the Moffitt schema. Patients marked with * were treated with FOLFIRINOX, and the rest were treated with FOLFIRINOX+PF-04136309. **E**, Correlation between the PurIST score (basal-like probability) for patient samples pre- and posttreatment in the Linehan trial. Basal-like samples were colored by orange and classical samples are colored by blue. **F** and **G**, Correlation between the percentage of change (% change) of tumors and the PurIST score (basal-like probability) derived from PurIST in basal-like (**F**) and classical samples (**G**), excluding the basal-like sample with an unstable DNA subtype.

unstable DNA subtype, we show a positive correlation between PurIST basal-like predicted class probabilities and worse treatment response in basal-like tumors (**Fig. 6F**). No association of PurIST classical confidence and treatment response was seen (**Fig. 6G**).

Discussion

Several subtyping systems for pancreatic cancer have now been proposed. Despite this, several limitations remain before they can be clinically usable. Here we leverage the wealth of transcriptomic studies that have been performed in pancreatic cancer to determine the molecular subtypes that may be most clinically useful and replicable across studies. Our results show that while multiple molecular subtypes may be used to characterize patient samples, the two tumor-intrinsic subtypes from the Moffitt schema: basal-like (overlaps with Bailey squamous/Collisson QM-PDA) and classical (overlaps with non-Bailey squamous/non-Collisson QM-

PDA) are the most concordant and clinically robust. The compelling findings of basal-like tumors showing resistance to FOLFIRINOX and the lack of objective studies comparing current first-line therapies FOLFIRINOX versus gemcitabine plus nab-paclitaxel strongly support the need to evaluate the role of molecular subtyping in treatment decision making for patients with PDAC. Therefore, we have developed a SSC based on the two tumor-intrinsic subtypes that avoids the instability associated with current strategies of clustering multiple samples and the low tumor purity issues in PDAC samples.

Prior studies have shown that merging samples from multiple studies (horizontal data integration) can improve the performance of prediction models, relative to training on individual studies (21). However, systematic differences in the scales of the expression values in each dataset are often observed, as some may have been separately normalized prior to their publication or were generated from a variety of expression platforms. Complicated cross-platform

normalizations are often employed in such situations prior to model training. Furthermore, new samples must be normalized to the training dataset prior to prediction to obtain relevant predicted values. This often results in a “test-set bias” (16), where predictions may change due to the samples in the test set or the normalization approach used. In addition, prediction models may change with the addition of new training samples, as renormalizations may be warranted among training samples. In all, this leads to potential complications for data merging, stability of prediction, and model accuracy (22, 23). We present PurIST, which is not dependent on cross-study normalization, and is robust to platform type and sample collection differences. We show that the sensitivity and specificity of PurIST calls are high across multiple independent studies, demonstrating that the PurIST classifier recapitulates the tumor-intrinsic subtype calling obtained initially by CC. Given the significant clinical relevance of the two tumor-intrinsic subtypes for both prognosis and treatment response, and the high accuracy of predicted subtype calls in our validation datasets, PurIST may have tremendous clinical value. Specifically, we show that PurIST works for gene expression data assayed across multiple platforms, including microarrays, RNAseq, and NanoString. Furthermore, the algorithm provides replicable classification for matched samples from snap-frozen bulk tissue as well as FNA, core biopsies, and archival tissues.

Thus, PurIST may be flexibly used on low input and more degraded samples and may be performed with targeted gene expression platforms such as NanoString, avoiding the need for a CLIA RNAseq assay. Our enduring findings that basal-like subtype tumors are significantly less likely to respond to FOLFIRINOX-based regimens strongly supports the need for the incorporation of molecular subtyping in future clinical trials to determine the association of molecular subtypes with this and other therapies. In addition, the stability of PurIST subtypes after treatment is a noteworthy finding and may point to fundamental biological differences in the tumor subtypes. However, larger clinical trials with pre- and posttreatment biopsies will be needed to determine whether this is a treatment-dependent observation. Our ability to subtype based on either core or FNA biopsies considerably increases the flexibility and practicality of integrating PDAC molecular subtypes into future clinical trials in the metastatic and neoadjuvant setting where bulk specimens are rarely available.

In summary, we present a clinically usable SSC that may be used on any type of gene expression data including RNAseq, microarray, and NanoString, and on diverse sample types including FFPE, core biopsies, FNAs, and bulk frozen tumors. Although results of the

association of FOLFIRINOX resistance in patients with basal-like subtype tumors is compelling, future prospective clinical trials in patients with PDAC will be needed to evaluate the utility of PurIST in treatment decision making, and in the context of different therapies.

Disclosure of Potential Conflicts of Interest

J.J. Yeh, N.U. Rashid, X.L. Peng, and R.A. Moffitt submitted US Provisional Application No.: 62/827473 on “Purity Independent Subtyping of Tumors (PurIST), A Platform and Sample Type Independent Single Sample Classifier for Treatment Decision Making in Pancreatic Cancer.” No potential conflicts of interest were disclosed by the other authors.

Authors' Contributions

Conception and design: J.J. Yeh, N.U. Rashid, R.A. Moffitt

Development of methodology: J.J. Yeh, N.U. Rashid, X.L. Peng, R.A. Moffitt

Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): J.J. Yeh, R.A. Moffitt, K.E. Volmar, B.A. Belt, R.Z. Panni, T.M. Nywening, S.G. Herrera, S.G. Hennessey, A.B. Morrison, A. Nayyar, B. Schmidt, H.J. Kim, D.C. Linehan

Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): J.J. Yeh, N.U. Rashid, X.L. Peng, C. Jin, R.A. Moffitt, R. Kawalerski, D.C. Linehan

Writing, review, and/or revision of the manuscript: J.J. Yeh, N.U. Rashid, X.L. Peng, C. Jin, R.A. Moffitt, K.E. Volmar, B.A. Belt, K.J. Moore, R. Kawalerski, A. Nayyar, B. Schmidt, H.J. Kim, D.C. Linehan

Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): J.J. Yeh, X.L. Peng, C. Jin, B.A. Belt, R.Z. Panni, T.M. Nywening, S.G. Herrera, K.J. Moore, R. Kawalerski, A. Nayyar, A.E. Chang, D.C. Linehan

Study supervision: J.J. Yeh, N.U. Rashid

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Purity Independent Subtyping of Tumors (PuriST), A Clinically Robust, Single-sample Classifier for Tumor Subtyping in Pancreatic Cancer

Naim U. Rashid, Xianlu L. Peng, Chong Jin, et al.

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