Comprehensive molecular profiling to predict clinical outcomes in pancreatic cancer

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Abstract
Background: Pancreatic ductal adenocarcinoma (PDAC) has the worst prognosis among common cancers. The genomic landscape of PDAC is defined by four mutational pathways: kirsten rat sarcoma virus (KRAS), cellular tumor antigen p53 (TP53), cyclin dependent kinase inhibitor 2A (CDKN2A), and SMAD family member 4 (SMAD4). However, there is a paucity of data on the molecular features associated with clinical outcomes after surgery or chemotherapy.

Methods: We performed comprehensive molecular characterization of tumor specimens from 83 patients with PDAC who received surgery, using whole-exome sequencing and ribonucleic acid sequencing on tumor and matched normal tissues derived from patients. We also systematically performed integrative analysis, combining genomic, transcriptomic, and clinical features to identify biomarkers and possible therapeutic targets.

Results: KRAS (75%), TP53 (67%), CDKN2A (12%), SMAD4 (20%), and ring finger protein 43 (RNF43) (13%) were identified as significantly mutated genes. The tumor-specific transcriptome was classified into two clusters (tumor S1 and tumor S2), which resembled the Moffitt tumor classification. Tumor S1 displayed two distinct subclusters (S1-1 and S1-2). The transcriptome of tumor S1-1 overlapped with the exocrine-like (Collisson)/ADEX (Bailey) subtype, while tumor S1-2 mostly consisted of the classical (Collisson)/progenitor (Bailey) subtype. In the analysis of combinatorial gene alterations, concomitant mutations of KRAS with low-density lipoprotein receptor related protein 1B (LRP1B) were associated with significantly worse disease-free survival after surgery (p = 0.034). One patient (1.2%) was an ultrahypermutant with microsatellite instability. We also identified high protein kinase C lota (PRKCI) expression as an overlapping, poor prognostic marker between our dataset and the TCGA dataset.

Conclusion: We identified potential prognostic biomarkers and therapeutic targets of patients with PDAC. Understanding these molecular aberrations that determine patient outcomes after surgery and chemotherapy has the potential to improve the treatment outcomes of PDAC patients.

Keywords: KRAS, landscape, LRP1B, pancreatic cancer, PRKCI

Introduction
Pancreatic ductal adenocarcinoma (PDAC) has the worst prognosis among common cancers. The 5-year survival rate of PDAC is approximately 9%, despite the recent success with combination chemotherapies and targeted therapy. The incidence, mortality, and disease burden of PDAC are all rapidly increasing worldwide. PDAC is predicted to become the second leading cause of cancer-related death in the United States by 2030.

The Australian Pancreatic Cancer Genome Initiative (APGI) and the Cancer Genome Atlas
(TCGA) Research Network have provided the mutational landscape of PDAC.\textsuperscript{9,10} The genomic landscape of PDAC is defined by four mutational pathways that are essential drivers of carcinogenesis; \textit{kirsten rat sarcoma virus} (\textit{KRAS}), \textit{cellular tumor antigen p53} (\textit{TP53}), \textit{cyclin dependent kinase inhibitor 2A} (\textit{CDKN2A}), and \textit{SMAD family member 4} (\textit{SMAD4}). Along with these four major drivers, the additional frequently mutated genes including ring finger protein 43 (\textit{RNF43}) and AT-rich interactive domain-containing protein 1A (\textit{ARID1A}) have been identified in the previous systematic genomic profiling of PDAC. The key mutation in PDAC is activated \textit{KRAS}, which occurs early in carcinogenesis, whereas mutations in \textit{SMAD4} occur later and are thought to facilitate the progression of PDAC.\textsuperscript{9,10} Germline and somatic mutations in the DNA damage repair (DDR) genes, \textit{BRCA2}, \textit{partner and localizer of BRCA2} (\textit{PALB2}), and \textit{ATM}, were observed in a minority of cases, representing a class of patients for whom platinum-based chemotherapy and/or poly(ADP-ribose) polymerase (PARP) inhibition may have therapeutic benefit.\textsuperscript{9,10} Several groups have described transcriptomic subtypes in PDAC, with mostly consistent findings concerning two major lineages that separate PDAC into the squamous (alternatively named basal-like and quasi-mesenchymal) and classical subtypes.\textsuperscript{11,12} However, there is a paucity of data on the molecular features associated with clinical outcomes after curative surgery and previous studies do not provide any correlative information with respect to chemotherapy response. In the current study, we performed integrative genomics analyses of PDAC patients who received surgery and/or chemotherapy to facilitate identification of biomarkers and possible therapeutic targets.

Materials and methods

Patient enrollment

Among the patients with PDAC who received curative surgery at Samsung Medical Center, Seoul, South Korea, between September 2008 and June 2017, 83 patients with available samples were included in this study. The molecular analysis included whole-exome sequencing and ribonucleic acid (RNA) sequencing on tumor and matched normal tissues derived from patients. Clinical information including age, sex, primary tumor site, histologic type, stage, and treatment outcomes was extracted from hospital records. We also evaluated the outcomes of palliative chemotherapy. Computed tomography scan and/or magnetic resonance imaging were performed every two cycles to evaluate the chemotherapy response according to RECIST 1.1 criteria.\textsuperscript{13}

The collection of specimens and associated clinical data used in this study was approved by the Institutional Review Board of Samsung Medical Center (IRB #2015-10-062 and #2018-03-162). All patients who participated in this study provided written informed consent prior to enrollment and specimen collection. This study was performed in accordance with the principles of the Helsinki Declaration and the Korean Good Clinical Practice guidelines.

Tumor sample collection

Tumor tissues were obtained during surgery. If the tumor content was \(\geq 40\%\) after pathological assessment, tumor deoxyribonucleic acid (DNA), and RNA were extracted from freshly acquired tissues using a QIAamp Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions.\textsuperscript{9,10} Several groups have described transcriptomic subtypes in PDAC, with mostly consistent findings concerning two major lineages that separate PDAC into the squamous (alternatively named basal-like and quasi-mesenchymal) and classical subtypes.\textsuperscript{11,12} However, there is a paucity of data on the molecular features associated with clinical outcomes after curative surgery and previous studies do not provide any correlative information with respect to chemotherapy response. In the current study, we performed integrative genomics analyses of PDAC patients who received surgery and/or chemotherapy to facilitate identification of biomarkers and possible therapeutic targets.

Whole-exome sequencing for tumor tissue

For the generation of standard exome capture libraries, we used the Agilent SureSelect Target Enrichment protocol for Illumina paired-end sequencing library, version B.3, June 2015 (Agilent Technologies, Inc. Santa Clara, CA, USA) with 200 ng of input formaldehyde-fixed, paraffin-embedded DNA. In all cases, the SureSelect Human All Exon V5 probe set was employed. DNA quantity and quality were evaluated using PicoGreen\textsuperscript{14} and NanoDrop.\textsuperscript{15} Fragmentation of 1 ng, MA, USA with 200 ng of input formaldehyde-fixed, paraffin-embedded DNA. In all cases, the SureSelect Human All Exon V5 probe set was employed. DNA quantity and quality were evaluated using PicoGreen\textsuperscript{14} and NanoDrop.\textsuperscript{15} Fragmentation of 1 ng of Inc., MA, USA\textsuperscript{16} end, and Agilent adapters were ligated to the fragments. Once ligation had been assessed, the adapter-ligated product was amplified by polymerase chain reaction (PCR). Subsequently, the final purified product was quantified using quantitative PCR (qPCR) under the qPCR Quantification Protocol Guide and assessed using the Caliper High Sensitivity DNA LabChip Kit.

We also evaluated the outcomes of palliative chemotherapy. Computed tomography scan and/or magnetic resonance imaging were performed every two cycles to evaluate the chemotherapy response according to RECIST 1.1 criteria.\textsuperscript{13}
(PerkinElmer Inc., MA, USA). For exome capture, 250 ng of DNA library was mixed with hybridization buffers, blocking mixes, RNase block, and 5 µl of SureSelect All Exon capture library, according to the standard Agilent SureSelect Target Enrichment protocol. Hybridization to capture baits was performed at 65°C using the heated lid option of a thermocycler at 105°C for 24 h on a PCR machine. The captured DNA was amplified. The final purified product was quantified using qPCR according to the qPCR Quantification Protocol Guide and assessed using the TapeStation RNA ScreenTape (Agilent). Finally, we performed sequencing using the HiSeq™ 2500 platform (Illumina, San Diego, CA, USA).

Whole-exome sequencing data analysis
The resulting reads from whole-exome sequencing were mapped to human genome version 19 (hg19) using the Burrows-Wheeler Aligner (version 0.7.12-r1039) with the BWA-MEM algorithm. SAMtools sorted the aligned sequences by genomic coordinates (v0.1.19). The sorted reads were subjected to the Genome Analysis Toolkit (GATK, v3.6 and v.4.13) for duplicate marking, indel realignment, and base recalibration. We performed MuTect2 from GATKv4.13 to detect tumor somatic mutations using the recalibrated BAM files from tumor and matched normal samples. After removing possible germline events (population allele fraction < 2.5e−6 in gnomAD), we annotated the called mutations using variant effect predictors (VEP). The resulting variant events were converted to MAF format, and then the variants with altered reads < 4 were eliminated for further analysis in R. Using MSIsensor, we calculated the microsatellite instability (MSI) score of each tumor by comparing the BAM file of the tumor sample with that of the matched normal sample. We called the tumor ‘MSI’ if the MSI score of the tumor was > 3.5. Otherwise, the tumors were classified as ‘microsatellite stable (MSS)’. The COSMIC v3 single base substitution (SBS) mutational signatures of each tumor sample were determined by deconstructSigs (v1.8.0, R package). Significantly mutated genes were identified by MutSigCV (v1.41) on MATLAB. To estimate copy number alterations, CNVkit (v0.9.7.b1) was used for copy segmentation, followed by GISTIC2.0 to calculate copy number alterations for each gene with log2 threshold = 1.

RNA sequencing
Total RNA concentration was estimated using Quant-IT RiboGreen (Invitrogen, Waltham, MA, USA). To determine the DV200 (% of RNA fragments >200 bp) value, samples were run on the TapeStation RNA ScreenTape (Agilent). Overall, 100 ng of total RNA was subjected to sequencing library construction using a TruSeq RNA Access Library Prep Kit (Illumina) according to the manufacturer’s protocol. Briefly, total RNA was first fragmented into small pieces using divalent cations under elevated temperature. The cleaved RNA fragments were copied into first-strand complementary DNA (cDNA) using SuperScript II reverse transcriptase (Invitrogen, #18064014) and random primers. This was followed by second-strand cDNA synthesis using DNA polymerase I, RNase H, and deoxyuridine 5-triphosphate. The cDNA fragments were subjected to an end-repair process, addition of a single ‘A’ base, and subsequently, ligation of the adapters. Thereafter, the products were purified and enriched with PCR to create the cDNA library. All libraries were normalized, and six were pooled into a single hybridization/capture reaction. Pooled libraries were incubated with a cocktail of biotinylated oligos, corresponding to the coding regions of the genome. Targeted library molecules were captured via hybridized biotinylated oligo probes using streptavidin-conjugated beads. After two rounds of hybridization/capture reactions, the enriched library molecules were subjected to a second round of PCR amplification. The captured libraries were quantified using a KAPA Library Quantification Kit for Illumina Sequencing platforms according to the qPCR Quantification Protocol Guide (KAPA BIOSYSTEMS, #KK4854) and assessed using the TapeStation D1000 ScreenTape (Agilent Technologies, # 5067-5582). Indexed libraries were subsequently submitted to an Illumina HiSeq 2500 platform (Illumina), and paired-end (2 × 100 bp) sequencing was performed by Macrogen Inc. (Seoul, South Korea).

Gene expression calling and analysis
RNA sequence reads were mapped on hg19 by STAR (v2.6.1d) and sorted according to genomic coordinates. Cufflinks (v2.2.1) was used to calculate gene expression levels in fragments per kilobase million (FPKM) using Ensembl gene annotation. The FPKM values were log2-transformed for further analysis. Based on gene
expression profiling of tumor samples, to assign predefined tumor subtypes, nearest template prediction was applied using gene markers of each subtype in R. To identify the differentially expressed genes in tumor samples compared with their matched normal samples, we performed a paired t test and selected genes with adjusted p-values < 0.05 and log2 fold-change > 1. As a result, 5012 genes remained that we defined as ‘tumor-specific’ genes. Among these, highly variable genes were selected with a standard deviation (SD) > 1 for unsupervised hierarchical clustering (HC). To assume the expression scores of tumor microenvironment-associated cell markers, we used MCPcounter in R. Tumor purities based on gene expression profiles were calculated by R package ESTIMATE. Expression scores of gene sets were estimated using a single sample gene set enrichment analysis (ssGSEA) algorithm in GSVA (R package). GSEA was performed on GSEA-P. Gene ontology analysis was performed on DAVID (https://david.ncifcrf.gov/). Differentially expressed genes (log2 fold change > 1 and false discovery rate < 0.05) among HC-based tumor groups were identified through DESeq2 in R.

Results

Patient characteristics
The baseline characteristics of the study participants are summarized in Table 1. All 83 patients were pathologically confirmed as PDAC and received curative surgery. Fifty-two patients (62.7%) were male and the median age was 65 years (range, 37–82). There were 26 patients (31.3%) with stage I, 36 (43.3%) with stage II, and 17 (20.5%) with stage III/IV at diagnosis. The most common tumor location was pancreas head (49.4%). CA19-9 levels were higher than the upper limit of normal (ULN, 37 U/ml) in 65 patients (78.3%) at diagnosis.

We evaluated the clinicopathological factors that are associated DFS and OS. As expected, poorer differentiation and higher stage of cancer resulted in worse prognosis. In addition, a CA19-9 level higher than the ULN was significantly associated with short DFS in univariate analysis (Supplemental Tables 1 and 2).

Genomic landscape of 83 PDAC patients
To characterize genomic and transcriptomic profiles of the 83 PDAC patients, we obtained tumor and matched normal pancreas tissues with blood samples from each PDAC patient. The collected samples were subjected to whole-exome sequencing and RNA sequencing. Whole-exome sequencing data derived from tumor and blood samples were used to estimate somatic mutations and copy number variations.

In the PDAC cohort, the median number of somatic nonsilent coding mutations was 45 (7–10028), which is comparable with previous findings [Supplemental Figure 1(a) and (b)]. In addition, including four major drivers in PDAC, five significantly mutated genes (SMGs) were identified; KRAS (75%, $q = 4.19\times10^{-12}$), TP53 (67%, $q = 1.05\times10^{-11}$), CDKN2A (12%, $q = 9.07\times10^{-12}$), SMAD4 (20%, $q = 9.07\times10^{-12}$), and RNF43 (13%, $q = 1.01\times10^{-6}$), as was also previously reported (Figure 1).
Activating KRAS mutations are illustrated in Supplemental Figure 2(a). Multiple oncogenic KRAS alleles were identified, including G12D (n = 23, 27.7%), followed by G12V (n = 22, 26.5%) and G12R (n = 10, 12.0%), as well as another hotspot at codon 12, and 61 mutant alleles at a lower prevalence. KRAS mutations affecting two separate codons were found in one patient (KRASG12D and KRASQ92V). Patients who had KRAS mutations showed no significant difference in clinical outcomes, in terms of DFS and response to palliative chemotherapy, compared with patients with KRAS wild-type tumors. However, in the analysis of combinatorial gene alterations, concomitant mutations of KRAS with LRP1B, which is known as a functional tumor suppressor gene, were associated with significantly worse DFS (p = 0.034) [Supplemental Figure 2(b)].

Germline and/or somatic mutations in the DDR system were found in nine patients (10.8%), showing mutations in serine/threonine kinase 11 (STK11) (4.8%), ATM (3.6%), MutL homolog 1 (MLH1) (3.6%), BRCA2 (2.4%), BRCA1 (1.2%), and PALB2 (1.2%). It should be noted that one tumor (#74) was ultrahypermutant with MSI. The mutational signature of this sample showed that 72% of mutations were attributed to a DNA mismatch repair (dMMR)–MSI mechanism with a high frequency of C–T transition [Figure 1 and Supplemental Figure 3(a)]. When we examined its gene expression profiles using RNA sequencing data, this MSI tumor showed high expression scores for immune cell markers, particularly in cytotoxic lymphocytes, natural killer (NK) cells, and monocytic lineage cells. Overexpression of PDCD1 and CD274 was also detected [Supplemental Figure 3(b) and (c)].

Table 1. Baseline characteristics.

| Characteristics              | Total (N=83) |
|------------------------------|--------------|
| Age, years                   |              |
| Median                       | 65           |
| Range                        | 37–82        |
| Sex, no. (%)                 |              |
| Male                         | 52 (62.7)    |
| Female                       | 31 (37.3)    |
| Pancreas tumor location, no. (%) |            |
| Head                         | 41 (49.4)    |
| Body                         | 24 (28.9)    |
| Tail                         | 18 (21.7)    |
| Differentiation, no. (%)     |              |
| Well                         | 6 (7.2)      |
| Well to moderately           | 1 (1.2)      |
| Moderately                   | 49 (59.3)    |
| Moderately to poorly         | 5 (6.0)      |
| Poorly                       | 22 (26.5)    |
| Stage at diagnosis, no. (%)  |              |
| IA                           | 8 (9.6)      |
| IB                           | 18 (21.7)    |
| IIA                          | 5 (6.0)      |
| IIB                          | 31 (37.3)    |
| III                          | 15 (18.1)    |
| IV                           | 2 (2.4)      |
| Level of CA19-9 at diagnosis, no. (%) |          |
| Normal                       | 18 (21.7)    |
| ULN to <59 × ULN             | 60 (72.3)    |
| >59 × ULN                    | 5 (6.0)      |
| Adjuvant treatment (N=42), no. (%) |            |
| Chemoradiation               | 31 (73.8)    |
| Chemotherapy                 | 10 (12.0)    |
| Radiation                    | 1 (2.4)      |
| Palliative first-line chemotheraphy (N=29), no. (%) |       |
| Gemcitabine monotherapy      | 20 (70.0)    |
| Gemcitabine-based combination| 5 (17.2)     |
| FOLFIRINOX                   | 3 (10.3)     |
| TS-1/cisplatin               | 1 (1.2)      |

CA19-9, carbohydrate antigen 19-9; FOLFIRINOX, folinic acid/fluorouracil/irinotecan/oxaliplatin; TS-1, tegafur/gimeracil/oteracil; ULN, upper limit of normal.

Tumor-specific and normal tissue transcriptomic classification of PDAC

Because molecular characterization of PDAC has been limited by the high level of stromal cell involvement, we conducted tumor-specific and normal tissue transcriptomic analysis. Through differentially expressed gene (DEG) analysis between tumors and their matched normal tissues, we identified 5012 and 413 tumor-specific and normal genes, respectively (absolute log2-fold change > 1, adjusted paired t test p-value < 0.05). According to previous studies, PDACs can be classified into several transcriptomic subtypes.11,28,29 To identify tumor-specific and
normal subgroups in this cohort and compare them with the previously identified subtypes, we performed unsupervised HC using highly variable tumor-specific and normal genes (standard deviation > 1) [Figure 2(a) and Supplemental Figure 4(a)]. Tumor-specific gene expressing profiling was classified into two clusters (tumor S1 and tumor S2), which resembled the Moffitt tumor classification, and were also generated by tumor-specific genes [Figure 2(a) and (b)]. Tumor S1 displayed two distinct subclusters (S1-1 and S1-2). The transcriptome of tumor S1-1 overlapped with the exocrine-like (Collisson)/ADEX (Bailey) subtype while tumor S1-2 mostly consisted of the classical (Collisson)/progenitor (Bailey) subtype. On the other hand, 88% of basal-like, 96% of quasi-mesenchymal, and 92% of squamous subtypes were clustered as tumor S2 [Figure 2(b)]. We could not find any survival differences between the tumor-specific HC-based PDAC groups, although tumors with poor differentiation were significantly frequent in tumor S2 ($p=0.36$, OR=2.99, 19/27 poorly differentiated versus 23/56 well-differentiated and moderately
differentiated) [Figure 2(a) and Supplemental Figure 4(c)]. In addition, we investigated the association between genomic alterations and tumor groups. We found that TP53 mutations occurred with significantly greater frequency in tumor S1-1 than in other groups \((p = 0.0018, \text{OR} = 13.4)\) [Figure 2(c)].

To dissect the cellular composition of the tumor microenvironment (TME) according to HC-based tumor group, we applied ESTIMATE and MCPcounter algorithms [Figure 3(d) and (e)]. With lower tumor purities, higher scores for immune cell and endothelial markers were found in tumor S1-1 than in other types. By comparison, markers for fibroblasts were upregulated in tumor S2, which showed high mesenchymal traits. ESTIMATE and MCPcounter results demonstrated the low infiltration levels of TME in tumor S1-2. Collectively, we identified three clusters using tumor-specific genes: tumor S1-1 was characterized by high stromal/immune cell infiltration and high frequency of TP53 mutations; tumor S1-2 displayed high tumor purities, resembling the
conventional classical subtype, and; tumor S2, which is similar to the basal-like/quasi-mesenchymal/squamous subtype, showed increased expression levels for fibroblast markers and enrichment of tumors with poor differentiation.

Unsupervised HC with normal genes also revealed two clusters depending on the expression levels of overall normal genes [Supplemental Figure 4(a)]. The first cluster (normal S1) showed an absence of expression of normal genes, but tumors belonging to the other cluster (normal S2) expressed normal genes. Gene ontology analysis revealed that expressed normal genes in normal S2 were involved in proteolysis and digestion [Supplemental Figure 4(b)]. In addition, 13 of 16 tumor S1-1 samples were classified as normal S2, consistent with TME analysis. Tumor S1-1 showed transcriptomic similarity with the exocrine-like or ADEX subtype; it is controversial whether the exocrine-like/ADEX subtype is a result of normal tissue contamination or is a bona fide tumor subtype. Although tumor S1-1 had low tumor cellularity, there were PDAC tumor-specific genes present such as PROM1 (a well-known cancer stem cell marker40), the expression levels of which were significantly upregulated in tumor S1-1 (Supplemental Table 3). This suggests that tumor S1-1 is one of the PDAC tumor types rather than the result of simple contamination by normal tissue.

Figure 3. Identification of PRKCI as a poor survival predictor in PDAC. (a) Workflow for identifying poor prognostic markers in both SMC (Samsung Medical Center, this study cohort) and the TCGA PAAD cohort. (b–d) Kaplan–Meier curves showing the overall (b) and disease-free (c) survival differences between patients with relatively high and relatively low PRKCI expression levels in the SMC (b, c) and the TCGA (d) dataset. (e, f) PRKCI expression levels in tumor and normal tissues in the SMC (e) (p-value, paired t test) and the TCGA (f) dataset (p-value, unpaired t test).

DEG, differentially expressed genes; FC, log2-fold change; FDR, false discovery rate; HZ, hazard ratio; OS, overall survival; PDAC, pancreatic ductal adenocarcinoma; PRKCI, protein kinase C iota type.
Identification of poor prognosis-related genes

As PDAC remains one of the most fatal malignancies and displays low tumor purity, the identification of tumor-specific prognostic markers and therapeutic targets is important for PDAC patient treatment. To discover tumor-specific genes that predict prognosis of PDAC, we performed Cox regression survival analysis using gene expression levels [Figure 3(a)]. Univariate Cox regression analysis with OS was applied to the 5012 tumor-specific genes. This analysis revealed 164 genes that met the criteria (HR > 1, p-value < 0.05). Of these 164 genes, 82 were identified as worse prognosis-related genes via multivariate Cox regression analysis with the prognostic factors (tumor stage and differentiation). The same procedure was performed with TCGA PAAD data to identify the prognostic genes that can be validated in an independent dataset. As a result, only PRKCI overlapped with the TCGA PAAD dataset. PDAC patients with higher expression levels of PRKCI showed worse OS and worse PFS [Figure 3(b)–(d)]. A significant upregulation of PRKCI in tumors compared with normal controls was confirmed [Figure 3(e) and (f)].

Response to palliative first-line chemotherapy

Among 83 patients, 29 received palliative first-line chemotherapy due to disease progression [Figure 4(a)]. Twenty patients were treated with gemcitabine monotherapy and five were treated with gemcitabine-based combination therapy (2, gemcitabine/nab-paclitaxel; 2, gemcitabine/erlotinib; 1, gemcitabine/capecitabine). FOLFIRINOX treatment was performed for another three patients. As a result, one, 16, and 10 patients achieved partial response (PR), stable disease (SD), and progressive disease (PD), respectively, and two patients were not evaluable [Figure 4(b)]. The overall clinical benefit ratio (PR + SD) was 58.6% [95% confidence interval (CI), 3.3–30.7].

When we examined genomic correlates of clinical benefit of chemotherapy, none of the genomic alterations of core pathways were associated with clinical benefit of chemotherapy [Figure 4(b)]. Therefore, we conducted GSEA to explore whether there are distinct molecular gene sets among patients with clinical benefit of chemotherapy. We found ATP-binding cassette (ABC) transporter-associated genes were upregulated in patients who did not have clinical benefit of chemotherapy [Figure 4(c) and (d)].

Discussion

Using whole-exome and transcriptome sequencing, we performed integrative molecular characterization of 83 patients with PDAC receiving surgery and/or chemotherapy. The genomic landscape of PDAC, including four major driving mutations (KRAS, TP53, CDKN2A, and SMAD4), was comparable with the previous APGI and TCGA datasets9,10 and we also found that a substantial proportion of patients harbored germline and somatic mutations in their DDR systems. In addition, we identified factors that affect clinical outcomes, such as concomitant mutational status of KRAS and LRP1B, the expression level of PRKCI, and the overexpression of ABC transporter proteins.

KRAS, which plays an essential role in the signaling pathways that regulate cell growth and differentiation, is the most frequently mutated oncogene in PDAC. However, the association between KRAS mutational status and clinical outcome after surgery has not been clearly established. One recent study showed that KRAS G12D-mutant tumors were associated with worse DFS after surgery, compared with KRAS wild-type tumors.41 That study also showed that multiple alterations (≥3) among the four major driver mutations (KRAS, CDKN2A, SMAD4, and TP53) are associated with worse DFS. Our study did not find a significant association between KRAS mutational status and clinical outcomes after surgery. Our study population had a relatively lower proportion of head cancer (49% versus 72–79%) and a relatively higher proportion of T1-2 disease (81.9% versus 8–26%). The previous study also included a majority of white ethnicity and included only 21% of Asian populations. It should be noted that our study included a significantly higher proportion of patients who received adjuvant concurrent chemoradiotherapy according to the institutional protocol (73.8% versus 22–40%), even though the proportion of node-positive disease is similar or slightly lower (58.8% versus 67–77%). These differences in baseline tumor characteristics and adjuvant treatment strategy may contribute to the discordance of the association between KRAS mutational status and disease-free survival after surgery.

Interestingly, we found that concomitant mutation of KRAS and LRP1B is associated with worse PFS after surgery. LRP1B encodes an endocytic low-density lipoprotein (LDL)-family receptor
and the receptor binds to multiple extracellular ligands, including fibrinogen and apolipoprotein (apoE)-carrying lipoproteins.\textsuperscript{42} \textit{LRP1B} is known as a functional tumor suppressor gene; inactivating mutations of \textit{LRP1B} are frequently observed in diverse solid cancers, including melanoma, lung cancer, and gastric cancer.\textsuperscript{43–45} Owing to its large coding sequence (16 kbp), \textit{LRP1B} is often missed as significantly mutated on gene analysis, whereas its mutation could still have a functional consequence in tumorigenesis and heterogeneity.\textsuperscript{46} Our study identified nine patients (10.8\%) harboring \textit{LRP1B} mutations and found that \textit{LRP1B} was significantly downregulated in tumor tissues, compared with normal tissues. We also conducted GSEA and found that proliferation/cell cycle-associated gene sets were enriched in \textit{LRP1B} and \textit{KRAS} mutant samples, compared with tumors with only \textit{KRAS} mutations [Supplemental Figure 2(c)]. Interestingly, a recent report found that downregulation of \textit{LRP1B} in colon cancer promoted the growth and migration of cancer cells via aberrant activation of beta-catenin/T-cell factor signaling.\textsuperscript{47} These findings might indicate that

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**Figure 4.** Genomic and molecular correlates of clinical benefit of chemotherapy. (a) Swimmer plots illustrating overall survival from surgery for patients whose clinical response to palliative first-line chemotherapy was available (top panel) and patients who did not receive palliative first-line chemotherapy (bottom panel). (b) Genomic and molecular characteristics of 29 PDAC patients who received chemotherapy. From top to bottom, progression-free survival in months, tumor mutation burden (log10 scale), mutational signatures of the COSMIC V3 SBS, treated chemotherapy information and responses to the therapy, clinico-pathological features, and genomic alterations in major oncogenic pathways of PDAC. (c) GSEA plot showing the upregulation of the KEGG_ABC_TRANSPORTERS gene set in tumors derived from patients who achieved PD to chemotherapy compared with patients with clinical benefit of chemotherapy. (d) ssGSEA scores of the KEGG_ABC_TRANSPORTERS gene set were significantly higher in PD than in PR/SD patients ($p$-value, Wilcoxon rank sum test).
LRP1B loss of function may accelerates tumor cell proliferation in RAS-activated circumstances and promotes metastasis of tumor cells.

Our study also showed that ~10% of tumor samples harbored germline or somatic mutations in at least one of the DDR genes, including STK11 (4.8%), ATM (3.6%), MLH1 (3.6%), BRCA2 (2.4%), BRCA1 (1.2%), and PALB1 (1.2%), all of which are potential predictive biomarkers sensitive to PARP inhibition. Of note, one patient was ultrahypermutated (267 mutations/Mb) and showed high MSI. For PDAC, the frequency of ultrahypermutation and high MSI is not well known and varies considerably between studies. A recent systematic review reported that high MSI in PDAC is rare but exists in 1–2% of cases. That study also showed that high MSI in PDAC is strongly associated with medullary and mucinous/colloid histology and is usually KRAS-TP53 nonsense/colloid histology and is usually KRAS-TP53

In this study, we identified high PRKCI expression levels as an overlapping, poor prognostic marker between our dataset and the TCGA dataset. PRKCI (encoding for PKCι) is an oncogene that is frequently overexpressed and associated with poor outcomes in lung cancers. In KRAS-mutant lung adenocarcinoma, PKCι establishes and maintains an aggressive stem-like, tumor-initiating cell phenotype by activating a PKCι–ELF3–NOTCH3 signaling axis that drives transformed growth and tumor initiation. Together with previous lung adenocarcinoma studies, our study supports the notion that pharmacologic blockade of PKCι signaling can be a potential therapeutic option in KRAS-mutant PDAC.

Overexpression of ABC transporter proteins is responsible for drug efflux, which attenuates the efficacy of chemotherapy by protecting tumor cells against chemotherapeutic agents. In the precision oncology era, cytotoxic chemotherapy, such as gemcitabine, gemcitabine/nab-paclitaxel, and FOLFIRINOX, is still the mainstay of treatment for metastatic pancreatic cancer. As a result, our results highlight that ABC transporters could be important therapeutic targets for overcoming cytotoxic chemotherapy resistance in PDAC patients.

Conclusions
We performed integrative genomics analyses of PDAC patients who received surgery and/or chemotherapy, and reconfirmed genomic landscapes, major driver mutations, and DDR pathway mutations associated with PDAC. We also discovered potential prognostic biomarkers and therapeutic targets, such as concomitant mutations of KRAS and LRP1B, the expression level of PRKCI, and the overexpression of ABC transporter proteins. In the future, understanding the molecular aberrations that determine patient
outcomes after surgery and chemotherapy has the potential to improve treatment outcomes of PDAC patients.

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Conflict of interest statement

The authors declare that there is no conflict of interest.

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Supplemental material

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