

The Circadian Clock is Disrupted in Pancreatic Cancer

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43 **Abstract**

44 Disruption of the circadian clock is inextricably linked to cancer development and progression.
45 Establishing this connection has proven beneficial for understanding cancer pathogenesis, determining
46 prognosis, and uncovering novel therapeutic targets. However, barriers to characterizing the circadian
47 clock in human pancreas and human pancreatic cancer – one of the deadliest malignancies – have
48 hindered an appreciation of its role in this cancer. Here, we employed normalized coefficient of variation
49 (nCV) and clock correlation analysis in human population-level data to determine the functioning of the
50 circadian clock in pancreas cancer and adjacent normal tissue. We found a substantially attenuated clock
51 in the pancreatic cancer tissue. Then we exploited our existing mouse pancreatic transcriptome data to
52 perform an analysis of the human normal and pancreas cancer samples using a machine learning method,
53 cyclic ordering by periodic structure (CYCLOPS). Through CYCLOPS ordering, we confirmed the nCV
54 and clock correlation findings of an intact circadian clock in normal pancreas with robust cycling of
55 several core clock genes. However, in pancreas cancer, there was a loss of rhythmicity of many core clock
56 genes with an inability to effectively order the cancer samples, providing substantive evidence of a
57 dysregulated clock. The implications of clock disruption were further assessed with a *Bmal1* knock-out
58 pancreas cancer model, which revealed that an arrhythmic clock caused accelerated cancer growth and
59 earlier metastatic spread, accompanied by chemoresistance and enrichment of key cancer-related
60 pathways. These findings provide strong evidence for clock disruption in human pancreas cancer and
61 demonstrate a link between circadian disruption and pancreas cancer progression.

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67 **Introduction**

68 The circadian clock is a conserved molecular feedback loop that regulates many signaling pathways to
69 control metabolism, immunity, apoptosis, and other critical cellular functions in the body.¹ At its core, the
70 positive arm of the clock (i.e. CLOCK and BMAL1 [also known as ARNTL or MOP3]) drives
71 transcription of the negative arm, including PER1-3 and CRY1-2.^{2,3} The negative arm represses the
72 transcriptional activation of CLOCK and BMAL1, and a second interlocked loop involves the nuclear
73 receptors ROR $\alpha/\beta/\lambda$ and NR1D1-2 (also known as REV-ERB α/β), which activate and suppress BMAL1
74 expression, respectively. These, along with other core clock components, form a tightly regulated series
75 of transcriptional-translational feedback loops that ensure rhythmic expression over 24 hours and function
76 to maintain cellular and organ homeostasis.²⁻⁴

77 Environmental cues influence the synchronization of circadian rhythms in various organ systems, and
78 misalignment of external cues with the internal clock (e.g. shift work) can cause clock dysfunction with
79 consequent metabolic derangements and pathologic states.⁴⁻⁶ For instance, circadian dysregulation has
80 been strongly linked to obesity and diabetes, both risk factors for cancer.⁷⁻¹¹ Concordantly, landmark
81 studies have shown that disruption of the endogenous clock through mutations in or suppression of the
82 core clock genes is intricately linked to tumor growth in several cancers.¹²⁻¹⁴ For example, knockout of
83 *Bmal1* in *Kras*- and *p53*-mutant lung cancer causes marked tumor progression *in vivo*,¹³ while MYC-
84 induced repression of *BMAL1* in human neuroblastoma drives decreased overall survival in a BMAL1-
85 dependent manner.¹⁵ Importantly, targeting dysfunctional clock components in certain cancers has proven
86 an effective treatment strategy.^{15,16} Thus, identifying an aberrantly functioning circadian clock in cancer
87 can lead to key advancements such as understanding pathogenesis, prognosis, and uncovering novel
88 therapeutic targets.

89 Although indeterminate, there is some evidence that the clock may be dysregulated in pancreatic ductal
90 adenocarcinoma (PDAC), leading to a worse prognosis;^{17,18} this is alarming for a deadly malignancy
91 where only 11% of patients survive beyond 5 years.¹⁹ To advance our understanding of how clock

92 disruption impacts PDAC pathogenesis, and ultimately foster the identification of therapeutic targets, an
93 essential first step is to establish that clock dysfunction exists in human PDAC. Unfortunately, to date, the
94 cumulative data does not definitively support this assertion and is inconclusive. Prior studies have relied
95 on contrasting expression differences between the core clock genes in tumor compared to normal
96 pancreas as a basis for clock disruption.^{17,18,20,21} But differential expression alone is limited, and does not
97 provide insight into critical components of clock health such as relative amplitude, rhythmicity, or
98 correlation of expression amongst core clock genes.²² Moreover, in the pancreas, phase advancement as a
99 result of chronic jetlag causes differential expression of core clock genes while maintaining a robust and
100 healthy clock (i.e. strong relative amplitude, rhythmicity, and clock correlation).²³ Thus, much more
101 substantive data is required to affirm clock disruption in PDAC.

102 In pre-clinical studies (e.g. mouse or cell-line models), the determinants of clock health can be identified
103 by obtaining longitudinal data under controlled conditions.^{23,24} However, in human studies – particularly
104 human pancreas – the requisite periodic data by multiple sampling is not feasible (or ethical). Therefore,
105 alternative means of determining rhythmically expressed genes in humans are necessary. Following the
106 emergence of several bioinformatics tools, the principal aspects of clock health including relative
107 amplitude, core clock gene correlation, and statistical determination of rhythmicity can be resolved when
108 using the appropriate reference data.^{22,25–28} We recently generated a robust pancreas dataset demonstrating
109 diurnally expressed genes over 48 hours.²³ With this foundation, we were able to apply normalized
110 coefficient of variation (nCV), clock correlation analysis, and cyclic ordering by periodic structure
111 (CYCLOPS) to test the hypothesis that the circadian clock is disrupted in human PDAC.^{22,25,28} While on
112 the surface this may be construed as a simplistic hypothesis, several limitations have hindered the ability
113 to evaluate the circadian clock in human pancreas, including the lack of human periodic data and an
114 absence of a reference transcriptional dataset.

115 Here, we employed nCV, clock correlation analysis, and CYCLOPS on publicly available human
116 population-level expression data from The Cancer Genome Atlas (TCGA) and Clinical Proteomic Tumor

117 Analysis Consortium 3 (CPTAC-3) datasets to determine the health of the circadian clock in PDAC and
118 adjacent normal tissue.^{29,30} For the first time, we identified an intact circadian clock in human normal
119 pancreas with robust cycling of several core clock genes. We also found a markedly weakened clock in
120 the cancer tissue, providing substantive evidence of a dysregulated clock in PDAC. These findings
121 represent significant advancements in evaluating clock function in PDAC, and the potential clinical
122 implications of clock disruption were further assessed with a pre-clinical PDAC model. We used
123 CRISPR/Cas9 technology to selectively target *Bmal1* and examined the effects of clock dysfunction *in*
124 *vitro* and *in vivo*. This revealed that loss of clock function caused accelerated cancer growth, earlier
125 metastatic spread, enrichment of key cancer-related pathways, and resistance to commonly used cytotoxic
126 chemotherapies for PDAC. These findings provide strong evidence for circadian clock disruption in
127 human PDAC and demonstrate a link between circadian disruption and pancreas cancer progression.

128 **Materials and Methods**

129 *Mouse Care*

130 All animal studies were conducted according to an approved protocol (M005959) by the University of
131 Wisconsin School of Medicine and Public Health (UW SMPH) Institutional Animal Care and Use
132 Committee (IACUC). Male and female C57Bl/6J mice were housed in an Assessment and Accreditation
133 of Laboratory Animal Care (AALAC) accredited selective pathogen-free facility (UW Medical Sciences
134 Center) on corncob bedding with chow diet (Mouse diet 9F 5020; PMI Nutrition International) and water
135 ad libitum.

136 *Clock Correlation, nCV, and CYCLOPS Pipeline*

137 We assessed the overall clock gene correlation and robustness of the clock with the clock correlation
138 matrix and normalized coefficient of variation (nCV) – nCV is known to be correlated with the relative
139 amplitude (rAMP) of oscillating clock genes, indicating the clock robustness.^{22,25} The nCV was calculated
140 for the overall condition with the nCVnet and nCVgene functions.²⁵ Clock correlation matrices were

141 created using an available shiny app (<https://github.com/gangwug/CCMapp>) which compares the
142 correlation of clock components (17 individual clock genes) to a baseline correlation from the circadian
143 atlas using the Mantel test.^{22,26,27,31,32}

144 Cyclic ordering by periodic structure (CYCLOPS)²⁸ was validated for use in the pancreas utilizing our
145 existing murine normal circadian and chronic jetlag pancreas RNA sequencing (RNA-seq) data (Gene
146 Expression Omnibus (GEO) Accession number: GSE165198).²³ Specifically, the seed genes for use in
147 CYCLOPS were selected by cross-referencing genes found to be rhythmically expressed in our dataset
148 with those genes either rhythmically expressed in the liver (similarly metabolic organ) or those used by
149 Wu *et al.* when validating CYCLOPS in the skin (**Supplemental Data File 1**).^{23,26,31,32} The updated
150 CYCLOPS pipeline by Wu *et al.* was then used to reorder our murine pancreas datasets with known
151 sample collection times.²⁶ Eigengenes were selected with the Oscope package to sharpen CYCLOPS.³³
152 Clusters with a $p < 0.05$ and $\text{Met}^{\text{smooth}} < 1$ were considered to be significantly reordered.²⁸ Rhythmicity of
153 the reordered genes was determined on cosinor analysis with a $p < 0.01$, $\text{rAMP} > 0.1$, goodness of fit (rsq)
154 > 0.1 , and $\text{fitmean} > 16$.²⁶ Significantly rhythmic gene phase was then compared to the rhythmic gene
155 phase detected from the known sample time collection using the meta3d function of Metacyle.³⁴ Clock
156 genes were highlighted to demonstrate a correlation between the predicted and actual phase.

157 The clock was evaluated in human normal and human PDAC RNA-seq datasets from TCGA and
158 CPTAC-3.^{29,30} After batch correction with ComBat and filtering, 50 matched normal and 318 PDAC
159 samples were obtained for analysis.³⁵ The pipeline described above was then used to obtain the clock
160 correlation matrix, nCV, and CYCLOPS reordering. Cosinor analysis was performed to test for
161 rhythmicity. Given the additional biologic heterogeneity of the human data, a $p < 0.05$, $\text{rAMP} > 0.1$,
162 goodness of fit (rsq) > 0.1 , and $\text{fitmean} > 16$ were used as a rhythmicity cutoff. Rhythmic genes from
163 normal samples were assessed with phase set enrichment analysis (PSEA).³⁶ Rhythmic gene sets ordered
164 by significance were inputted with their calculated phase of expression. Default settings were used for
165 PSEA, including domain 0-24, min item 10, max sims/test 10,000. The gene set enrichment analysis

166 (GSEA) gene ontology (GO) (c5.go.bp.v7.5.1.symbols) set was leveraged as the pathway input. The top
167 15 significant pathways ($q < 0.05$) were selected for representation.

168 *KPC Cell Line Creation and Maintenance*

169 Pancreas cancer cells that harbor *Kras*^{G12D} and *Trp53*^{R172H} mutations (KPC cells) were acquired from
170 Ximbio (Catalog Number 153474; Westfield Stratford City, UK). Cells were cultured in DMEM
171 supplemented with 10% Fetal Bovine Serum (Cytiva, Marlborough, MA), 1% L-glutamate-Penicillin-
172 Streptomycin (Gibco, ThermoFisher Scientific, Waltham, MA), and 1% non-essential amino acids
173 (Gibco) at 37°C at 5.0% CO₂ to their appropriate confluence for use. CRISPR/Cas9 technology was used
174 to introduce *Bmiall* mutations into the well-established KPC cells, generating the *Bmiall* functional
175 knock-out line (KPC-BKO).³⁷ Synthetic tracrRNA and target-specific crRNAs (crRNA:tracrRNA
176 (ctRNAs)) were annealed as per manufacturer instructions (Integrated DNA Technologies (IDT),
177 Coralville, IA). Ribonucleoproteins were formed with Cas9 protein (V3, catalog # 1081059, IDT) and
178 ctRNAs individually (ctRNA 1:AATATGCAGAACACCAAGGA, ctRNA 2:
179 TTAGAATATGCAGAACACCA) (IDT). Nucleofection was used to introduce RNPs (1.95 μM Cas9, 2
180 μM ctRNA, 2 μM electroporation enhancer (IDT)) into 2 x 10⁵ KPC cells on a Nucleofector 4D (Lonza
181 Biosciences, Walkersville, MD) with an SF kit (Catalog Number V4CX-2032, Lonza Biosciences) as per
182 manufacturer instructions using pulse code CM-120. After 48 hours of recovery in DMEM growth media
183 (described above) at 37°C at 5.0% CO₂, single cells were deposited in 96 well plates using a BD
184 FACSaria III (BD Biosciences, Franklin Lakes, NJ). Outgrowing clones were condensed to a 96 well
185 plate in duplicate to propagate clones and generate a genomic DNA source. Genomic DNA was harvested
186 from KPC cells, nucleofected with ctRNA-1 RNPs and ctRNA-2 RNPs, and the targeted region of *Bmiall*
187 was PCR amplified using primers: Forward: acactttccctacacgacgcttccgatct NNNNNNN
188 CCAAGAACCTTGTGTCTG and Reverse: gtgactggagttcagacgtgtcttccgatct
189 AGAGGACTCCACAGACATGAAC (IDT). PCR products were dual-indexed with indexing PCR,
190 pooled, sequenced on an Illumina MiSeq instrument (San Diego, CA), and analyzed with CRISPResso2.³⁸

191 *Creation of Per2-dLuciferase reporter KPC cell line*

192 KPC cells were stably transfected with a mammalian gene expression vector harboring a destabilized
193 luciferase reporter driven by the *Per2* promoter fused to intron 2 of *Per2* and a puromycin resistance
194 cassette (VectorBuilder, Santa Clara, CA).³⁹ The vector was transfected with lipofectamine 2000
195 (ThermoFisher Scientific) and incubated for 2 days, and then exposed to media containing 2.5 µg/mL
196 puromycin for 3 days, and surviving clones were selected. Luciferase activity was measured in the
197 selected clones with the luciferase assay system (Promega, Madison, WI) on a BMG CLARIOstar (BMG
198 Labtech, Ortenberg, Germany) plate reader. The two selected clones were then subcloned using a BD
199 FACS Aria III after staining with DAPI. Luciferase activity was again measured in the subclones to
200 validate expression before use in downstream experiments.

201 *Clock Function Testing*

202 To evaluate for clock function, KPC and KPC-BKO cells were synchronized with 200 nM
203 dexamethasone for 2 hours in FBS-free DMEM media, followed by RNA isolation 24 hours after the
204 synchronization using the RNeasy protocol (Qiagen, Hilden, Germany) according to the manufacturer's
205 recommendations.⁴⁰ Samples were collected at 4-hour intervals for 24 hours. Quantitative real-time
206 polymerase chain reaction (RT-qPCR) was performed for the downstream core clock gene (CCG) *Per1*
207 (ID: Mm00501813_m1, Life Technologies, Carlsbad, CA) and the housekeeper gene *Hprt* (ID:
208 Mm03024075_m1) using GoTaq® Probe qPCR and RT-qPCR System (Promega, Madison, WI) and
209 Quantstudio 7 flex RT-PCR system (ThermoFisher Scientific). Expression was measured and the ΔCT
210 was calculated. The mean ΔCT values were then tested for rhythmicity using the meta2d function of
211 Metacycle.³⁴ For Metacycle settings, the min period and max period were set to 24, and “JTK”, “LS”, and
212 “ARS” were selected for cycMethod. An integrated FDR corrected *q* value < 0.05 and rAMP > 0.1 were
213 taken as rhythmic. To separately evaluate clock function in KPC cells, luciferase activity driven by the
214 *Per2* promoter was measured. Cells from two independent clones were synchronized with 200 nM
215 dexamethasone for 2 hours in FBS-free DMEM media, and 24 hours later luciferase activity was

216 measured at 4-hour intervals. To measure activity, 1x cell culture lysis reagent (Promega) was added and
217 cells were incubated for 5 minutes. Firefly luciferase assay reagent (Promega) was added and
218 luminescence was measured on a BMG CLARIOstar plate reader.

219 *Western Blotting*

220 Western blotting was performed to determine BMAL1 expression in the KPC cell lines. After
221 synchronization, protein samples were isolated at 6-hour intervals for 24 hours using CellLytic M lysis
222 reagent (MilliporeSigma, Burlington, MA) and Halt Protease and Phosphatase Inhibitor Cocktail
223 (ThermoFisher Scientific). A total of 30 μ g of each sample was loaded onto a Mini-PROTEAN TGX
224 7.5% precast mini-gel (Bio-RAD Laboratories, Hercules, CA). The gel was then transferred using the
225 semi-dry transfer technique to an Immobilon-P PVDF membrane (MilliporeSigma). The membrane was
226 then blocked with 5% skim milk and incubated with 1 μ g/ μ L rabbit BMAL1 antibody (NB100-2288,
227 Novus Biologicals, Littleton, CO) and 1:2000 rabbit β -ACTIN antibody (4967, Cell Signaling
228 Technologies, Danvers, MA). Finally, the membranes were incubated with alkaline phosphatase-
229 conjugated goat anti-rabbit IgG (1:10000) (Jackson ImmunoResearch West Grove, PA) and stained using
230 1-Step NBT/BCIP solution (ThermoFisher).

231 *RNA Isolation, Sequencing, Differential Gene Expression Analysis*

232 To evaluate for transcriptomic differences between wild-type and BKO KPC cells, bulk RNA-seq was
233 performed on 6 independent samples collected from each condition. Isolation was carried out as above
234 and quality was tested for an RNA integrity number (RIN) > 7.5 on the Agilent 2100 bioanalyzer (Agilent
235 Technologies, Santa Clara, CA). A total of 300 ng of mRNA was enriched with poly-A selection and
236 sequencing on the Illumina HiSeq2500 platform by the University of Wisconsin Biotechnology
237 Sequencing Core. FASTq files were processed with Skewer and genes were filtered to remove those with
238 low expression.⁴¹ Samples were normalized by the method of trimmed mean of M-values.⁴² Contrasts
239 were drawn with the edgeR package, with differential expression taken when the FDR $q < 0.05$.⁴³

240 Pathway testing was performed with the KEGG database (Kyoto Encyclopedia of Genes and Genomes)
241 using previously described methods.⁴⁴ The top 500 significant genes were inputted, ordered by *q* value,
242 and the top 9 significant pathways (where *p* < 0.05) were plotted for visualization. Pathway dot size is
243 indicative of the number of genes in each pathway. The RNA-seq data is publicly available through GEO
244 (Accession number: GSE213680).

245 *Heterotopic Tumor Modeling*

246 To create flank tumors, 1 x 10⁵ KPC or KPC-BKO cells were injected into the right flanks of
247 immunocompetent C57BL/6J mice obtained from Jackson Laboratory (Bar Harbor, ME). Cells were
248 mixed in a 1:1 50 µL solution of DMEM media and Matrigel (Corning Inc, Corning, NY). In the first
249 experiment, a single dose of KPC or KPC-BKO cells was injected into C57BL/6J mice (male: n = 10,
250 female: n = 10, each group) and tumors were measured twice weekly for four weeks starting on day 6
251 with the caliper method as previously described.⁴⁵ Tumor length and width were measured and tumor
252 volume was calculated using the formula: tumor volume = length x width² x 1/2. Tumor weight was also
253 measured (in mg) at the conclusion of the study period. Two independent replicates were performed for
254 each condition. In the second experiment, the same dose of KPC or KPC-BKO cells was injected into
255 C57BL/6J mice (n = 7 in each group) and tumors were measured weekly until the mice became moribund
256 or died. Kaplan Meier log-rank analysis was then performed to compare survival differences between
257 conditions with the survival package.⁴⁶

258 *Analysis*

259 All analyses were performed in R version 4.2.0 or Julia version 0.3.12 unless otherwise indicated.

260 **Results**

261 *The Clock in Human Pancreatic Ductal Adenocarcinoma is Less Robust Than in Normal Pancreatic*
262 *Tissue*

263 To test the hypothesis that the circadian clock is disrupted in human pancreatic ductal adenocarcinoma
264 (PDAC), we used clock correlation, nCV, and CYCLOPS analysis on publicly available TCGA and
265 CPTAC-3 pancreatic datasets (**Figure 1**, **Figure 2**, and **Figure 3**).^{29,30} However, we first needed to ensure
266 the validity of the pipeline in the pancreas since these had not been previously applied to pancreatic
267 tissue. We assessed the clock in our existing murine normal circadian and chronic jetlag pancreas RNA-
268 seq datasets by examining the clock correlation matrix and nCV.²³ These datasets contained pancreas
269 samples acquired every 4 hours for 48 hours in male (n = 3 at each timepoint for each condition) and
270 female (n = 3 at each timepoint for each condition) mice under standard lighting (normal circadian, n =
271 72) and chronic jetlag (n = 72) conditions. Chronic jetlag is known to affect the phase of gene expression
272 but not the relative amplitude (rAMP) or the correlation of the core clock genes in the pancreas.²³
273 Correspondingly, we found that the core clock relationships remained intact on the correlation matrix
274 when compared to the baseline clock correlation matrix for both normal and chronic jetlag conditions ($p <$
275 0.001, z-statistic = 54.96 vs. $p < 0.001$, z-statistic = 54.08) (**Supplemental Figure 1A**).^{31,32} Further, the
276 nCV of 11 clock genes remained unchanged ($p = 0.76$) between the normal circadian (mean nCV (\pm
277 standard error) = 1.78 (\pm 0.27)) and chronic jetlag (mean nCV = 1.9 (\pm 0.28)) conditions (**Supplemental**
278 **Figure 1B**) – consistent with the murine pancreatic clock being strongly rhythmic and intact, matching
279 precisely what we had found in our prior longitudinal analysis.²³

280 CYCLOPS was then used to reorder the mouse pancreas samples by their predicted phase, and cluster
281 reordering of samples by CYCLOPS was analyzed for appropriate phase progression compared to the
282 known zeitgeber time (ZT) of sample collection. Notably, CYCLOPS accurately (phase appropriately)
283 reordered both the normal ($p < 0.01$; $\text{Met}^{\text{smooth}} = 0.90$) and the chronic jetlag conditions ($p < 0.01$;
284 $\text{Met}^{\text{smooth}} = 0.97$), validating the pipeline for use in the pancreas (**Supplemental Figure 2B-C**).
285 Collectively, these results confirmed the ability of nCV, correlation matrix, and CYCLOPS to determine
286 the robustness of the circadian clock in the pancreas.

287 Considerable data support the tumor suppressor role of the circadian clock and the assertion that circadian
288 disruption is present in human cancer.⁴⁷ In human PDAC, the supposition is that the clock may be altered,
289 but supporting data is limited to the relative expression of single genes and is consequently
290 inconclusive.^{18,21,48} We therefore applied our pipeline to determine clock health in PDAC using the TCGA
291 and CPTAC-3 pancreatic datasets.^{22,26-30} After processing and batch correction of the 50 available normal
292 and 318 available PDAC samples (**Supplemental Figure 3**), we examined the correlation matrix between
293 the core clock genes (CCGs) and found that there was a weaker relationship amongst the CCGs in PDAC
294 compared to normal (z-statistic PDAC = 13.77 vs. z-statistic normal = 22.42) (**Figure 1A**). Concordantly,
295 we found that there was a clear decrease ($p = 0.04$) in the nCV between PDAC (mean nCV = 0.69 (\pm
296 0.08)) and normal (mean nCV = 1.03 (\pm 0.13)) (**Figure 1B**), indicating a weaker clock in PDAC
297 compared to normal. We then applied our pancreatic CYCLOPS analysis (**Supplemental Figure 4**)
298 which significantly reordered the normal samples ($p < 0.01$; Met^{smooth} = 0.99), indicating an intact clock.
299 This was further reinforced when evaluating the predicted phase of CCGs and clock-controlled genes
300 using CYCLOPS. We found that the phase sequence of normal pancreatic samples was similar to the
301 phase sequence of mouse pancreatic samples (**Figure 2A-B**), with concordant phase in 12 out of 17 genes
302 and ROR-phased genes largely peaking before E-box phase genes. Based on the CYCLOPS ordering, we
303 analyzed the proportion of genes that were rhythmically expressed in normal pancreas. We found that
304 4,034/18,196 (22.17%) of normal pancreatic genes were rhythmic, (**Supplemental Data Files 2**). We
305 found that several key CCGs and clock-controlled genes were rhythmically expressed in normal samples
306 on cosinor analysis, including *CLOCK*, *PER1*, *PER3*, *NR1D2*, *RORC*, *NFIL3*, *TEF*, *NR1D1*, *BHLHE40*,
307 and *NPAS2* (**Figure 3A and Supplemental Table 1**). The CCGs *BMAL1* ($p = 0.07$, rAMP = 0.8) and
308 *CRY1* ($p = 0.08$, rAMP = 0.48) were nearly rhythmic in the normal samples based on the pre-defined
309 cutoff. Finally, phase set enrichment analysis (PSEA) was used to determine phase-dependent gene
310 enrichment. The top 15 significant pathways were selected for analysis (**Figure 3C**). Consistent with our
311 prior murine pancreas data, we found that normal pancreas was associated with time-dependent metabolic

312 gene pathway enrichment.^{23,49} In aggregate, this data confirms the health of the circadian clock in normal
313 pancreatic tissue, which was expected but not previously shown.

314 Despite the validation steps and success in ordering normal pancreas, CYCLOPS could not reorder PDAC
315 samples ($p = 0.43$; $\text{Met}^{\text{smooth}} = 0.99$), revealing a less robust (or disrupted) clock in PDAC compared to
316 normal. This inability to reorder strongly supports the attenuated circadian clock in PDAC identified with
317 nCV and clock correlation analysis. We selected the best reordering available based on the eigengenes
318 (**Supplemental Figure 4**) to enable visual representation of the PDAC samples compared to normal
319 pancreas samples. This also enabled us to estimate as best as possible the rAMP of the PDAC samples
320 (**Supplemental Data Files 3**), to coincide with the visual representation. Consistent with the nCV
321 analysis, we found that rAMP decreased substantially in every core clock gene from normal to tumor
322 (**Supplemental Table 1**), as evidenced by *PER1* and *PER3* expression (**Figure 3A**). Collectively, the
323 weaker clock correlation, markedly reduced nCV, and CYCLOPS analysis of human population-level
324 data convincingly show for the first time that the circadian clock is significantly disrupted in PDAC.

325 *Creation of a Circadian Dysfunction Pancreatic Cancer Model*

326 After demonstrating that the clock in human PDAC was dysfunctional, we sought to understand the
327 potential clinical implications of clock disruption. Identifying an aberrant clock in other cancers has
328 consistently demonstrated accelerated cancer progression and worse prognosis, while simultaneously
329 revealing novel therapeutic targets.¹⁵ Moreover, there appears to be a putative correlation between
330 suppressed *BMAL1* expression and poor prognosis in patients with PDAC.^{17,18} Therefore, we
331 hypothesized that disruption of the circadian clock in PDAC would cause accelerated cancer progression.
332 To test this hypothesis, we used a syngeneic *Kras*- and *Trp53*-mutant pancreas cancer cell line (KPC) and
333 employed CRISPR/Cas9 technology to functionally knock out *Bmal1* (KPC-BKO) so that we could
334 examine the effects of clock dysfunction *in vitro* and *in vivo*.³⁷ The reason for mutating *Bmal1* in the
335 PDAC cells was several-fold: i) KPC cells demonstrated an intact clock (see below) necessitating core
336 clock gene modulation, ii) *BMAL1* is a central transcriptional regulator of the circadian clock machinery

337 and suppressed gene expression has been found in several human cancers, correlating with worse
338 prognosis,^{13,15,17,18,20,21,50} iii) *BMAL1* expression is decreased in tumor compared to normal tissue in human
339 PDAC,¹⁸ iv) we similarly found a substantially dampened nCV (rAMP) of *BMAL1* expression in PDAC
340 compared to normal tissue (*PER1*, *PER3*, *NR1D1*, and *BMAL1* displayed the greatest decrease in nCV),
341 and v) segregating patients with PDAC into high and low *BMAL1* expression appears to be prognostic for
342 survival outcomes.^{17,18,21}

343 Consequently, guide RNAs were designed to target exon 8 of the *Bmal1* gene, which resulted in the
344 insertion of adenine on one allele and a 2 base pair deletion on the other between nucleotide 81,074 and
345 81,075 at amino acid 73 (GRCm38) (**Figure 4A**). The result of both mutations was a frameshift just
346 upstream of the basic helix loop helix (bHLH) domain known to assist the PAS A domain in
347 heterodimerization with its binding partner CLOCK (**Figure 4B**).^{51,52} Confirmatory western blotting
348 revealed the presence of protein in wild-type (WT) cells and an absence of protein in *Bmal1* knockout
349 (BKO) KPC cells (**Figure 4C**). To evaluate clock functionality, mRNA was isolated at 4-hour intervals (n
350 = 3 per condition) for 24 hours after cell synchronization, and we performed RT-qPCR for *Per1*, a core
351 downstream repressor of the positive arm of the molecular clock and a gene that demonstrated
352 rhythmicity in our human normal CYCLOPS analysis (**Figure 4D**). Rhythmicity was assessed with the
353 meta2d function of the Metacycle package.³⁴ KPC (WT) cells exhibited an intact and robust circadian
354 clock with a $q = 0.0004$ and rAMP = 0.11 for *Per1*. To further evaluate the functionality of the clock in
355 KPC cells, luciferase activity driven by the *Per2* promoter was measured. Consistent with our evaluation
356 of mRNA expression, luciferase activity in two separate clones was highly rhythmic (Clone 1: $q = 5.51E-$
357 5, rAMP = 0.44; Clone 2: $q = 1.95E-7$, rAMP = 0.39) (**Supplemental Figure 5**). Conversely, KPC-BKO
358 cells exhibited no detectable clock function with a $q = 0.81$ and rAMP = 0.003 for *Per1* expression
359 (**Figure 4D**). Consistent with the known cross-talk between the circadian clock and cell cycle, KPC-BKO
360 cells demonstrated alterations in the cell cycle compared to KPC cells (**Figure 4E**).²⁰ BKO cells were
361 found to have a higher mean (\pm standard error) percentage of cells in S phase (48.66 (\pm 0.13) vs. 36.96 (\pm

362 2.04); $p = 0.029$) and a lower percentage in G2 (18.03 (± 0.27) vs. 27.49 (± 0.89); $p = 0.0053$). There
363 were no differences between KPC and KPC-BKO cells in G1 (35.56 (± 1.19) vs. 33.31 (± 0.17); $p = 0.2$).
364 Taken together, these data demonstrate the successful creation of a novel PDAC cell line with an
365 abolished clock (*Bmal1* gene mutation), which can be used to elucidate the repercussions of clock
366 disruption in PDAC.

367 *Clock Dysfunction Accelerates Pancreatic Cancer Growth*

368 After establishing the model, we sought to understand how a dysfunctional clock impacted cell growth.
369 KPC and KPC-BKO cells were grown in culture and injected into the right flanks of syngeneic C57BL/6J
370 mice. Tumor growth was measured every 3-4 days for a total of 28 days to understand differences in
371 primary tumor growth. We found that BKO caused an accelerated growth pattern, resulting in higher
372 mean (\pm standard error) weight tumors at the study conclusion compared to KPC-derived tumors (438.02
373 ± 48.84 mg vs. 280.11 ± 42.73 mg; $p = 0.02$) (**Figure 5A-B**). These findings were evaluated and
374 independently confirmed in a second identically created *Bmal1* mutant (functional knock-out) clone
375 (**Supplemental Figure 6**). Perhaps more relevant to human PDAC – given the aggressiveness of this
376 cancer – we assessed the effect of clock disruption on survival (i.e. metastatic spread) by implanting KPC
377 and KPC-BKO heterotopic tumors and observing the mice until moribund status or lethality (**Figure 5C**).
378 Notably, the terminal endpoint occurred from metastatic spread to the diaphragm, lungs, or abdominal
379 cavity, or uncommonly from locally advanced tumors. On Kaplan Meier log-rank analysis, mice
380 harboring tumors derived from KPC-BKO cells experienced worse survival than mice bearing KPC-
381 derived tumors (median survival: 52 versus 75 days, $p = 0.04$). Thus, clock dysfunction resulted in
382 accelerated tumor growth and earlier onset of tumor metastases (i.e., PDAC progression) *in vivo*, leading
383 to worse overall survival.

384 *Loss of Bmal1 Promotes the Enrichment of Cell Growth Pathways*

385 To determine the possible etiologies of clock disruption causing accelerated PDAC progression, we
386 compared the transcriptomic profiles of KPC and KPC-BKO cells. The principal component analysis
387 revealed clear transcriptional profile differences between each condition (**Figure 6A**). Differential gene
388 expression was quantified with edgeR and of the 15,110 genes, 5,235 (34.65%) were upregulated and
389 5,113 (33.84%) were downregulated in KPC-BKO compared to KPC cells (**Figure 6B, Supplemental**
390 **Data File 4**).⁴³ When we examined the CCGs known to control the circadian cycle, we found that 12/15
391 examined clock genes were differentially expressed, including *Per1*, *Per2*, *Per3*, *Cry1*, *Cry2*, *Nr1d1*,
392 *Nr1d2*, *Blhhe40*, *Blhhe41*, *Npas2*, *Arntl2*, and *Dbp* (All q < 0.05).⁵¹ We then performed a KEGG pathway
393 analysis to examine for enrichment of pathways as a result of *Bmal1* knock-out in the PDAC cells
394 (**Figure 5C**). We found that there was an enrichment of pathways important for cellular adhesion, such as
395 ECM-Receptor Interaction, Cell Adhesion, and Focal Adhesion, as well as several cellular growth
396 pathways including PI3K-AKT Signaling Pathway, MAPK Signaling Pathway, and Rap1 Signaling
397 Pathway. Collectively, these data indicate that clock disruption in the KPC cells results in significant core
398 clock gene changes and a transcriptional shift that alters key cancer promotion-related pathways such as
399 cellular attachment and proliferation.

400 *Cell Survival is Increased with the Loss of Bmal1 through Alterations in Multiple Cell Death Pathways*

401 A hallmark of cancer progression is the inhibition of apoptosis.⁵³ This phenotype is readily apparent in
402 response to chemotherapy. In particular, the mechanism of action of gemcitabine (inhibition of DNA
403 synthesis) and paclitaxel (microtubule stabilization) – backbones in PDAC therapy – is to ultimately
404 induce apoptosis.^{54,55} To understand the potential implications of clock disruption for PDAC patients
405 undergoing treatment, we subjected KPC and KPC-BKO cells to 72 hours of chemotherapeutic treatment
406 with either gemcitabine or paclitaxel.⁵⁶ We found that the activity of cell death pathways through
407 apoptosis (measured by Caspase 3/7 activity) was blunted in response to both chemotherapeutic agents as
408 a result of clock dysfunction (**Figure 7A**). We also assessed cytotoxic cell death, as measured by Dead-
409 Cell Protease activity, and found that clock disruption promoted resistance to cytotoxic cell death induced

410 by gemcitabine and paclitaxel (**Figure 7B**). Prior work has implicated alteration of *Trp53* signaling to
411 impact cell survival by suppressed apoptosis,²⁰ but KPC cells (both WT and BKO) are *Trp53* mutant
412 (*Trp53*^{R172H}) indicating alternative mechanisms of heightened resistance in the KPC-BKO cells.
413 Gemcitabine resistance in PDAC often occurs due to the downregulation of the channel protein hENT1
414 (*SLC29A1* gene), or deoxycytidine kinase (*DCK* gene) which activates gemcitabine once inside the cell.⁵⁷
415 However, these were only marginally downregulated (1.05-fold, $q = 0.04$ and 1.08-fold, $q = 0.02$) due to
416 *Bmal1* knock-out, and thus unlikely to contribute to the differences seen with clock disruption.
417 Furthermore, these were non-rhythmic when examining our human samples, suggesting a lack of
418 circadian control. Meanwhile, paclitaxel drug resistance is thought to occur mostly through upregulation
419 of drug efflux transporter proteins (P-glycoprotein also known as multidrug-resistance associated protein),
420 but these ATP-binding cassette transporter proteins (*Abcb1a* and *Abcb1b* genes) were instead
421 significantly *downregulated* (8.2-fold, $q < 0.0001$ and 2.1-fold, $q < 0.0001$) in KPC-BKO vs KPC cells.⁵⁵
422 Concordantly, the *ABCB1* gene was rhythmic with elevated expression in the human normal samples ($p =$
423 0.015, rAMP = 0.95, mean expression = 73) compared to the human PDAC samples (mean expression =
424 23). Thus, the commonly described resistance mechanisms for gemcitabine and paclitaxel were not
425 identified, underscoring the complexity of clock disruption induced chemotherapeutic resistance in
426 cancer. Regardless, these data demonstrate that clock dysfunction promotes broad resistance in PDAC
427 including multiple cell death pathways in the setting of two different PDAC backbone agents.

428 **Discussion:**

429 Our work herein used matched normal and tumor samples from TCGA and CPTAC-3 to provide the first
430 substantive evidence that the circadian clock is disrupted in PDAC while the adjacent normal pancreatic
431 clock is intact. To evaluate clock functionality in the pancreas, we assessed the clock with three
432 complementary analyses not previously performed in pancreatic tissues: nCV, clock correlation, and
433 CYCLOPS. The nCV was pioneered and validated in several tissues by Wu and colleagues.²⁵ It is a
434 measurement that is directly related to the relative amplitude (rAMP) of the oscillation, and assesses

435 overall and individual clock gene robustness.²⁵ For example, Wu and colleagues demonstrated that the
436 nCV of core clock genes (CCGs) was consistently and significantly diminished in clock-disrupted tissues
437 versus wild-type tissues (e.g. *Bmal1* knockout adipocytes vs wild-type adipocytes) and human datasets of
438 tumor compared to matched (adjacent) normal tissue, where the timing of sample collection was
439 unspecified.²⁵ It is therefore extraordinarily beneficial for understanding the rAMP of the circadian clock
440 genes – a key measure of clock health – in population-level data where sample time acquisition is
441 unknown. In conjunction with nCV, Shilts *et al.* developed the method of clock correlation to determine
442 the progression of the clock in time-indeterminate datasets.²² By capitalizing on the concept of co-
443 expression of CCGs (intrinsic to the transcriptional translational feedback loop), Shilts and colleagues
444 were able to computationally discern clock progression in transcriptomic data from numerous human
445 datasets. Importantly, this technique is not dependent on full coverage of the period by samples, providing
446 a beneficial approach for our normal data set which included only 50 samples and did not have phase
447 representation across the 24-hour period. Furthermore, a direct comparison between heatmaps can be
448 made (i.e. murine vs human or tumor vs normal) because each heatmap has the same corresponding color
449 to rho correlation value.²² Thus, when combining nCV and clock correlation analysis in unordered human
450 samples, the health of the circadian clock in population-level data can be ascertained even when working
451 with low sample numbers (at least 30).

452 Wu and colleagues applied nCV analysis to several paired tumor-normal datasets,²⁵ while Shilts *et al.*
453 examined clock correlation in normal tissues and paired tumor-normal datasets.²² However, we are the
454 first group to analyze the pancreas, possibly because our recent publication was the first to characterize
455 the diurnal expression of mouse pancreatic genes over 48 hours.²³ Thus, an appropriate mouse reference
456 group had not been published for comparison. In the study by Shilts *et al.*, the human liver clock
457 correlation heatmap (a similarly metabolic organ) demonstrated a weaker clock correlation versus the
458 mouse reference, which they attributed to increased noise in the liver dataset.²² Therefore, in our human
459 data, we expected to identify an apparent ‘diminished health’ of the clock (lower nCV and weaker clock

460 correlation) in the human pancreatic tissue versus mouse samples due to significant variation in the
461 normal samples. This variability is contributed by human factors, such as type and timing of diet,^{58,59}
462 underlying genetic differences, and alterations in the ‘normal pancreas’ that surrounds the tumor
463 (pancreatic atrophy, fibrosis, inflammation, etc.).⁶⁰ These can modulate the relative amplitude of
464 oscillation or CCG correlation which contributes to significant noise in the data, as compared to the
465 genetically identical, age-matched, and environmentally matched mouse samples.^{22,28} Regardless of these
466 limitations, the nCV and clock correlation revealed two key components of a healthy clock in human
467 pancreas, which was an important component of the pipeline.

468 CYCLOPS is a machine learning method developed by Anafi *et al.*,²⁸ with subsequent elegant studies by
469 Wu *et al.*,^{26,27} to demonstrate how clock gene relationships can be used to infer and reorder samples by
470 their predicted phase of expression to understand circadian biology in population-level data. The authors
471 who developed CYCLOPS recommend roughly 250 samples for a complete phase distribution,²⁸ which is
472 dependent on differences in time of surgery (presume specimens acquired between 6 AM and 6 PM) as
473 well as differences in genetics and environmental factors (e.g. differences in sleep-wake cycle, or shift
474 worker status).⁶¹ CYCLOPS has been applied to the lung, liver, brain, hepatocellular carcinoma, and skin,
475 but not the pancreas.^{27,28} This is possibly due to known challenges associated with CYCLOPS, which
476 includes optimizing the seed gene list for appropriate ordering.²⁸ We used the modified CYCLOPS
477 approach by Wu *et al.* and were able to leverage our murine pancreatic longitudinal expression data to
478 generate the seed gene list for use with human data, which increases the signal-to-noise ratio to optimize
479 ordering capability.^{23,27,28} Thus, despite the expected variability in human normal pancreas samples,
480 CYCLOPS significantly ordered the samples across the period, with several clock genes, including
481 *CLOCK*, *PER1*, *PER3*, *NR1D2*, *RORC*, *NFIL3*, *TEF*, *NR1D1*, *BHLHE40*, and *NPAS2* demonstrating
482 rhythmic expression. While *BMAL1* and *CRY1* did not pass our predetermined cutoff for rhythmicity in
483 the normal samples, they demonstrated a robust rAMP with near-significant p values likely reflecting
484 limitations with our sample number (i.e., distribution of samples across the period) than true lack of

485 rhythmicity in the normal samples. As evidenced by the apparent gap in phase in the normal data, there
486 was not uniform distribution of normal samples across the 24-hour period, which signifies a limitation of
487 our analysis. However, despite these limitations, CYCLOPS statistically ordered the human normal data,
488 and the phase-set enrichment analysis and predicted phase of expression (of most CCGs) aligned well
489 with our prior mouse pancreas transcriptomic data (reference dataset). When combined with nCV and
490 clock correlation, the data clearly demonstrate an intact clock in the human normal pancreas samples,
491 which was an essential premise for evaluating the clock in human PDAC.

492 We then proceeded to use nCV, clock correlation, and CYCLOPS to provide compelling evidence of
493 circadian clock disruption in PDAC. While the concept of circadian disruption in PDAC has been posited
494 by others, past studies examining global clock function in human PDAC have been limited by detecting
495 binary differences in overall RNA and protein expression between normal pancreas and PDAC.^{17,18,20,21}
496 Relles *et al.* found that several CCGs demonstrated decreased expression in PDAC compared to benign
497 tumors or normal pancreas,¹⁸ while Li *et al.* found that low *Bmal1* expression (compared to ‘higher
498 expression’) was associated with worse disease-free and overall survival in patients with PDAC.¹⁷ Similar
499 studies have been repeated with concordant findings.^{20,21} However, binary comparisons of expression are
500 unlikely to capture the complexities of clock health such as rhythmicity, phase changes, changes in
501 rAMP, or changes in clock progression. With the 318 available PDAC samples, we showed that there was
502 markedly diminished nCV and a much weaker correlation among clock genes in PDAC compared to
503 normal pancreas, depicting a loss of clock health in the cancer tissue. Further, although the inability to
504 reorder PDAC samples may be a limitation of CYCLOPS, it more likely indicates clock dysfunction.⁶² In
505 the PDAC samples, there was a sufficient sample number (n = 318) for phase distribution, but the rAMP
506 of nearly all CCGs was markedly decreased (as visualized through best reordering), with many
507 manifesting an arrhythmic pattern. Collectively, our approach to human data (nCV, clock correlation, and
508 CYCLOPS) provided convincing evidence of clock disruption in PDAC.

509 The main limitation of our current strategy is the inability to discern why there was a loss of circadian
510 signatures in human PDAC population data. These observations could be the consequence of i)
511 dampening of the clock to a significant extent in every patient (i.e., global phenomena), ii) differences in
512 the extent to which the clock is disrupted (i.e., substantial clock dysfunction in a certain subset of
513 patients), or iii) biological heterogeneity in the PDAC samples surpassing circadian variability. The last
514 consideration seems unlikely given the marked decrease in rAMP (nCV) and clock correlation showing
515 significantly diminished clock health in PDAC. Furthermore, as seen with the CYCLOPS ordered *PER1*
516 expression data over the 24-hr period (Figure 3A), enhanced variability did not lead to the apparent
517 arrhythmicity in PDAC. Unfortunately, we were unable to discriminate between the first two possibilities.
518 The literature in cancer would suggest that cohorts of patients exhibit a differential extent of clock
519 disruption in their tumors (i.e. some clock intact vs some markedly disrupted),^{15,17,18,50} which would result
520 in an overall loss of circadian signatures seen in the nCV, clock correlation, and CYCLOPS analysis. Our
521 assessment of clock function in the KPC cells would also support this hypothesis, given that the clock
522 was intact in the *Kras*- and *Trp53*-mutant pancreas cancer cells and necessitated *Bmal1* mutagenesis to
523 generate clock disruption. Further, we assessed the cycling of the *Per1* gene and *Per2*-luciferase in the
524 KPC cells and identified a robust rhythm; by comparison, the human PDAC samples demonstrated
525 arrhythmic *PER1* expression. Therefore, we generated the novel murine KPC-BKO cell line as a basis to
526 evaluate clock disruption in PDAC, and intend for this cell line to be a useful approach for investigators
527 evaluating the clock in PDAC.

528 Using our KPC and KPC-BKO cells, we found accelerated tumor growth in our syngeneic *in vivo* model
529 with *Bmal1* functional knock-out. Our preference was to use an immune-competent model given the
530 known impact of the circadian clock on the immune system (another strength of our model).^{63,64}
531 Interestingly, our primary tumor growth results were similar to results by Jiang and colleagues, who used
532 implanted human PDAC cells (BxPC-3 and PANC-1) into immunocompromised mice after shRNA
533 knockdown of *Bmal1*.^{20,21} However, patients with PDAC ultimately succumb to distant metastatic spread,

534 rather than local tumor growth, and so examining the contribution of clock disruption to overall survival
535 was of greater importance.⁶⁵ We found that *Bmal1* mutation promoted earlier metastatic spread of tumors
536 resulting in earlier lethality, which has not been shown before. Concordant with the aggressive tumor
537 phenotype, *Bmal1* mutation also caused resistance to chemotherapy. While resistance to gemcitabine has
538 been shown,²¹ we found chemoresistance to two different backbone anti-cancer agents (gemcitabine and
539 paclitaxel), including suppressed apoptosis and cytotoxicity, indicating a more broad resistance to
540 standard PDAC chemotherapy. Although suppressed *Bmal1* expression in PDAC has been suggested to
541 modulate *Trp53* to promote a tumor suppressor effect, this was unlikely the case in our study considering
542 KPC cells are a *Trp53*-mutant cell line.²⁰ Other work indicates the transcription factor YY1 is
543 significantly overexpressed in PDAC and ultimately causes BMAL1 suppression with consequent PDAC
544 progression and resistance to gemcitabine (unclear mechanism of resistance).²¹ Yet, when we examined
545 our human PDAC and human normal samples, we found that *YY1* expression was equivalent between the
546 groups (mean expression 101.61 versus 102.87). Turning to other cancers where BMAL1 is suppressed,
547 *MYC* amplification in neuroblastoma alters *BMAL1* mRNA expression through induction of *NR1D1*.^{50,66}
548 Notably, this is associated with poor prognosis and is *BMAL1* dependent since ectopic expression of
549 *BMAL1* inhibits tumor growth. This mechanism is unlikely to be occurring in PDAC since there is no
550 upregulation of *MYC*, *MYC-N*, or *NR1D1* gene expression in human PDAC compared to normal samples.
551 While common resistance mechanisms of gemcitabine and paclitaxel were not reflected in the human
552 PDAC versus human normal data or the KPC versus KPC-BKO cells (e.g. channel proteins), RNA
553 sequencing identified several enriched pathways integral to cancer progression and chemoresistance in the
554 *Bmal1* functional knock-out cells, such as the PI3K-AKT pathway.^{67,68} The PI3K-AKT pathway is
555 inextricably linked to cancer cell proliferation and resistance to apoptosis, indicating a plausible
556 mechanism for inhibition of programmed cell death to multiple agents seen in our study.^{69,70} Further,
557 resistance to paclitaxel is associated with the activation of the PI3K-AKT pathway,⁷¹ and similar
558 correlations have been identified with gemcitabine resistance.^{72,73} However, the mechanism of
559 chemoresistance is quite complex and the etiology for suppressed apoptosis and cytotoxicity due to clock

560 disruption remains unclear. In future work, we can employ a series of clock-manipulated PDAC cell lines
561 to better understand the contributions of clock disruption to chemoresistance and PDAC progression.

562 In conclusion, we used a comprehensive approach (nCV, clock correlation, and CYCLOPS) to evaluate
563 the health of the circadian clock in human normal pancreas and demonstrated clock disruption in human
564 PDAC. Additionally, we developed novel cell lines to evaluate the repercussions of clock disruption in
565 PDAC and identified factors associated with poor prognosis (i.e., earlier metastases, resistance to
566 chemotherapy, and enrichment of cancer-related pathways). Future work will expand on this foundation
567 to disentangle the clock-dependent effects of PDAC and consequently focus therapeutic efforts.

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754 **Figures:**

755 **Figure 1:** *The human pancreatic clock robustness is diminished after malignant transformation.* **A.** Clock
756 correlation matrices were created for normal (left; n = 50) and pancreatic ductal adenocarcinoma (right; n
757 = 318) samples. **B.** The normalized coefficient of variation (nCV) was calculated to determine the
758 robustness of the pancreatic clock for the normal (circles) and PDAC (triangles) samples. Each color
759 indicates a different gene evaluated.

760 **Figure 2:** *Clock phase predictions demonstrate conserved clock relationships:* **A.** Graphical
761 representation and **B.** table of the CCG and clock-controlled gene phase prediction from the cosinor
762 analysis of CYCLOPS reordered human pancreas samples (n = 50) and Metacycle predicted phases of
763 mouse pancreas samples (n = 72), relative to *BMAL1* (human) and *Bmal1* (mouse) phase.

764 **Figure 3:** *The human pancreatic cancer tumor clock is dysfunctional relative to the normal pancreas.* **A.**
765 CYCLOPS was used to reorder samples from normal (n = 50) and pancreatic ductal adenocarcinoma (n =
766 318) TCGA and CPTAC-3 samples. Plots from several core clock genes reordered by CYCLOPS in
767 normal tissue as compared to best reordering in PDAC (**A**) are shown – ordered from 0 to 2π . Shading
768 around the blue regression line indicates the 95% confidence interval. **B.** Graphical representation of
769 Phase Set Enrichment Analysis (PSEA) of normal pancreatic samples, ordered by phase of expression.

770 **Figure 4:** *Creation of a mouse clock dysfunction cell line model.* **A.** The core clock gene *Bmal1* was
771 mutated (functional knock out) in KPC murine pancreatic cells with CRISPR-Cas9 genome editing
772 technology (BKO) with the frameshift site indicated by the blue line. **B.** The frameshift mutation was
773 induced upstream of the basic helix loop helix (bHLH) domain (red star). The resulting protein lacked
774 critical downstream elements for functionality, including the PAS-A, PAS-B, and PAC domains. **C.**

775 Western blot demonstrating the loss of BMAL1 protein across 24 hours for wild-type (WT) and BKO
776 cells (M = marker lane). **D.** The mean (\pm standard error) *Per1* mRNA expression of synchronized KPC (n
777 = 3 at each time point; black dashed) and KPC-BKO (n = 3 at each time point; grey) cells from ZT0-
778 ZT24. **E.** Graphs comparing the percent of cells in G1, S, and G2 in WT (n = 3; white) and BKO (n = 3;
779 grey) KPC cells.

780 **Figure 5:** *Mutation of Bmal1 promotes pancreatic cancer progression.* **A.** Comparison of mouse tumor
781 weight at 28 days for WT (n = 20; white) and BKO (n = 20) KPC tumors. **B.** WT (n = 20; black) and BKO
782 (n = 20) mean (\pm standard error) tumor size (in mm³) over the 28-day growth period. **C.** Kaplan Meier
783 survival curve for mice implanted with WT (n = 7; black) and BKO (n = 7; grey) KPC tumors. The dotted
784 lines indicate the median survival. [ns = not significant, * = p < 0.05, ** = p < 0.01]

785 **Figure 6:** *Functional knockout of Bmal1 promotes widespread transcriptomic alterations and activation*
786 *of multiple oncogenic pathways.* **A.** Principal Component Analysis (PCA) demonstrating overall
787 differences in expression between WT(green) and BKO (purple) samples (n = 6 each) **B.** Volcano plot
788 showing genes that were significantly upregulated (red) and downregulated (blue) on differential gene
789 expression (DGE) analysis (q < 0.05). All significant core clock genes are highlighted **C.** KEGG analysis
790 was then performed, and the top 9 pathways ordered by significance are shown. For each pathway, the
791 size of each dot corresponds to the number of genes involved in each pathway.

792 **Figure 7:** *Clock dysfunction diminishes the chemotherapeutic cell death response.* KPC wildtype (WT)
793 and KPC-BKO cells were treated with increasing doses of gemcitabine and paclitaxel. Fold change
794 differences (\pm standard error) in Caspase 3/7 activity in response to either **A.** gemcitabine or **B.** paclitaxel.
795 Fold change differences (\pm standard error) in dead-cell protease activity in response to **C.** gemcitabine or
796 **D.** paclitaxel. [ns = not significant, * = p < 0.05, ** = p < 0.01, *** = p < 0.001]

797 **Supplementary Files:**

798 **Supplementary Data Files:**

799 **Supplemental Data File 1:** CYCLOPS seed gene list.

800 **Supplemental Data File 2:** Significant genes on cosinor analysis for normal CYCLOPS reordered
801 normal pancreas

802 **Supplemental Data File 3:** Significant genes on cosinor analysis for normal CYCLOPS reordered PDAC

803 **Supplemental Data File 4:** KPC wildtype versus BKO edgeR differential gene expression

804 **Supplementary Figures:**

805 **Supplemental Figure 1:** *The wildtype mouse pancreatic clock is intact and robust.* **A.** Clock correlation
806 matrices were created for normal (left, n = 72) and pancreatic ductal adenocarcinoma (right, n = 72)
807 samples. **B.** The normalized coefficient of variation (nCV) was calculated to determine the robustness of
808 the pancreatic clock for the normal circadian (circles) and chronic jetlag (triangles) pancreas samples.
809 Each color indicates a different gene evaluated.

810 **Supplemental Figure 2:** *CYCLOPS accurately reorders wildtype mouse pancreas samples.* Eigengenes
811 identified by Oscope for use in CYCLOPS in **A.** normal circadian (upper) and chronic jetlag (lower)
812 samples. Shading around the blue regression line indicates the 95% confidence interval for each plot. **B.**
813 The normal circadian (left) and chronic jetlag (right) clusters ordered by CYCLOPS demonstrate accurate
814 reordering for both conditions **C.** Normal circadian (left) and chronic jetlag (right) genes found to be
815 significantly rhythmic on both CYCLOPS reordered cosinor analysis and rhythmicity testing based on the
816 known sample collection time with the Metacycle meta3d function are ordered by their predicted phase of
817 expression. Clock genes are shown in orange.

818 **Supplemental Figure 3:** *Normal pancreatic and pancreatic ductal adenocarcinoma (PDAC) samples
819 were processed and filtered.* TCGA and CPTAC-3 samples were processed and batched corrected. An
820 MDS plot is shown demonstrating differences between the matched normal (green; n = 50) and PDAC
821 (purple; n = 318) samples.

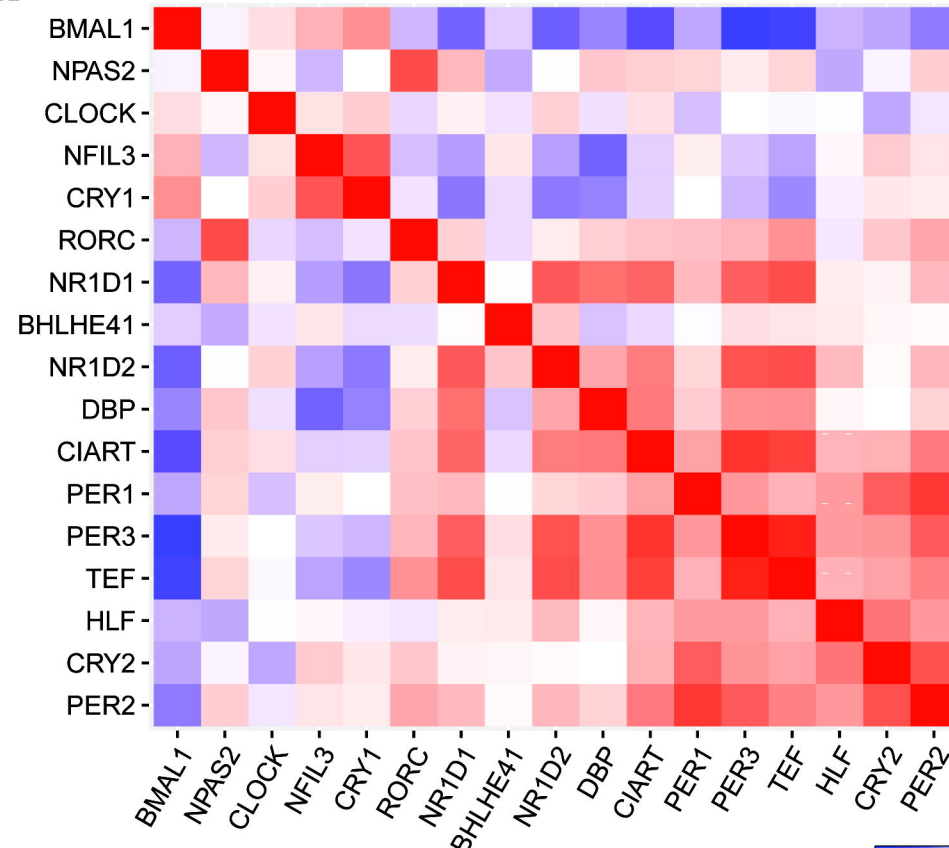
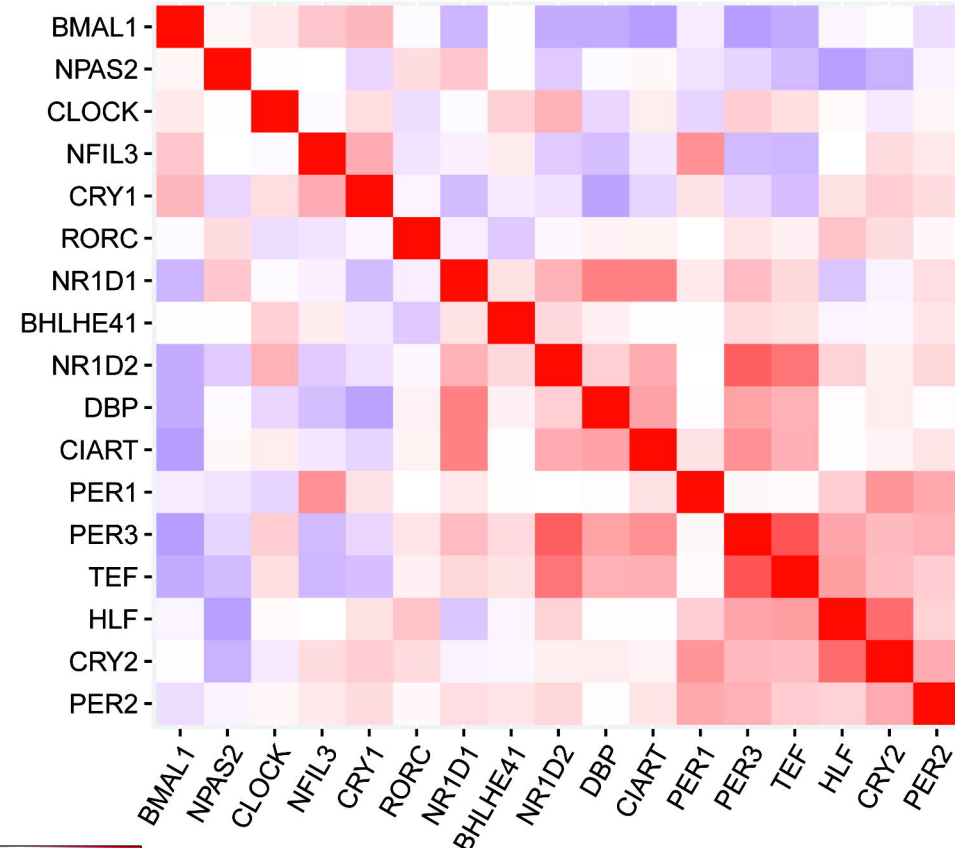
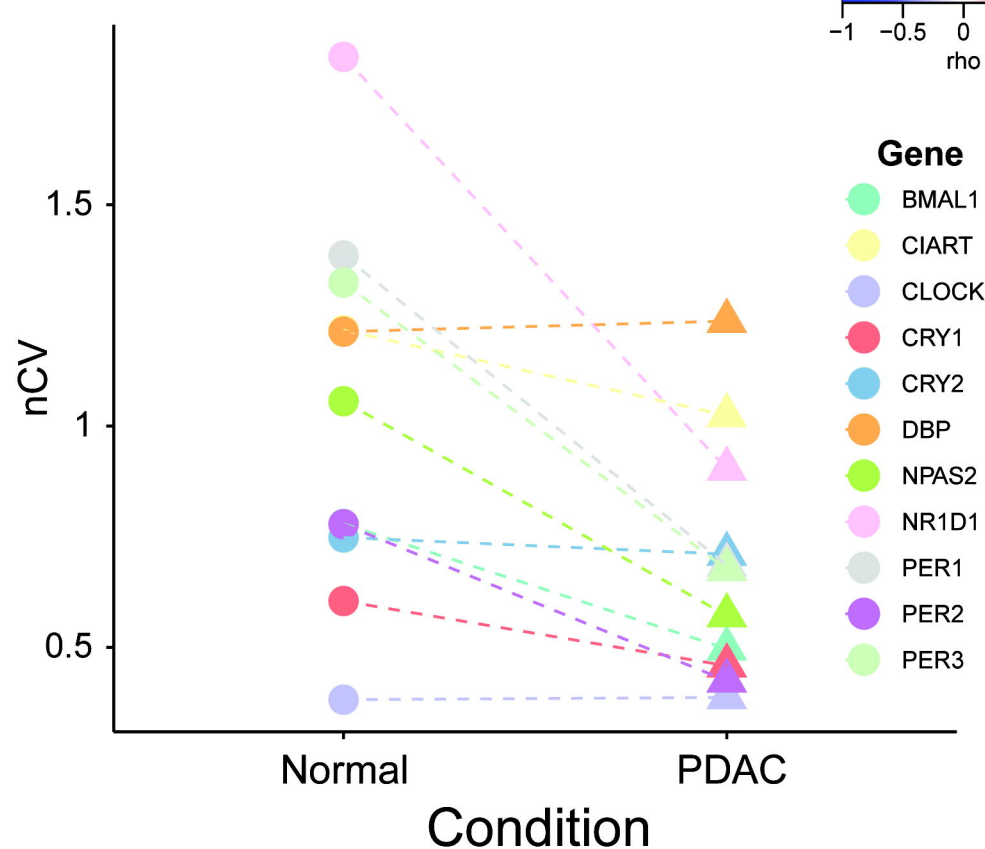
822 **Supplemental Figure 4:** Eigengenes for the human normal pancreas and pancreatic ductal
823 adenocarcinoma (PDAC) samples. Eigengenes were selected by Oscope for normal (upper) and PDAC
824 (lower) samples for use in CYCLOPS. Shading around the blue regression line indicates the 95%
825 confidence interval for each plot.

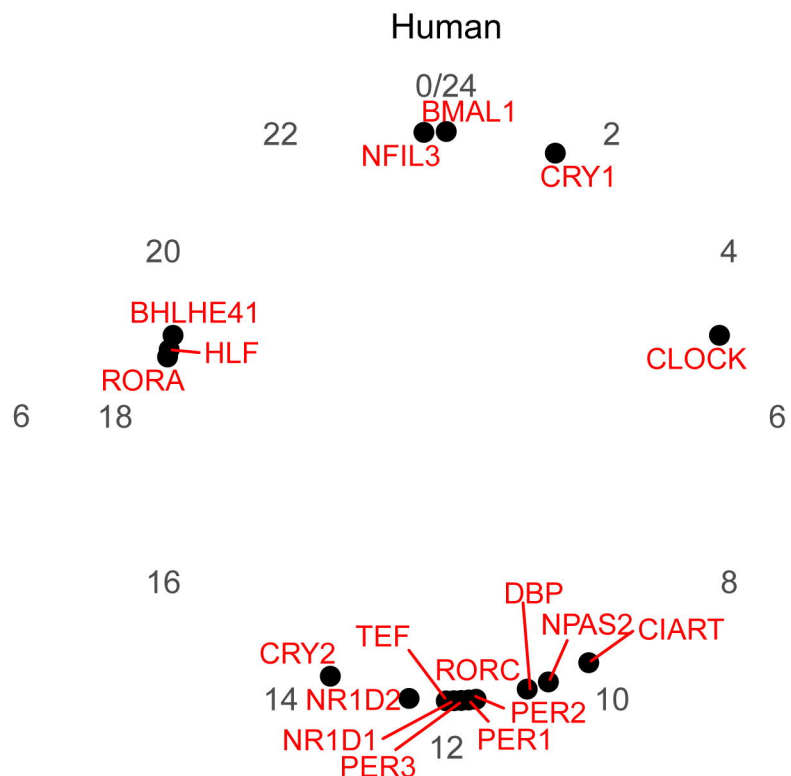
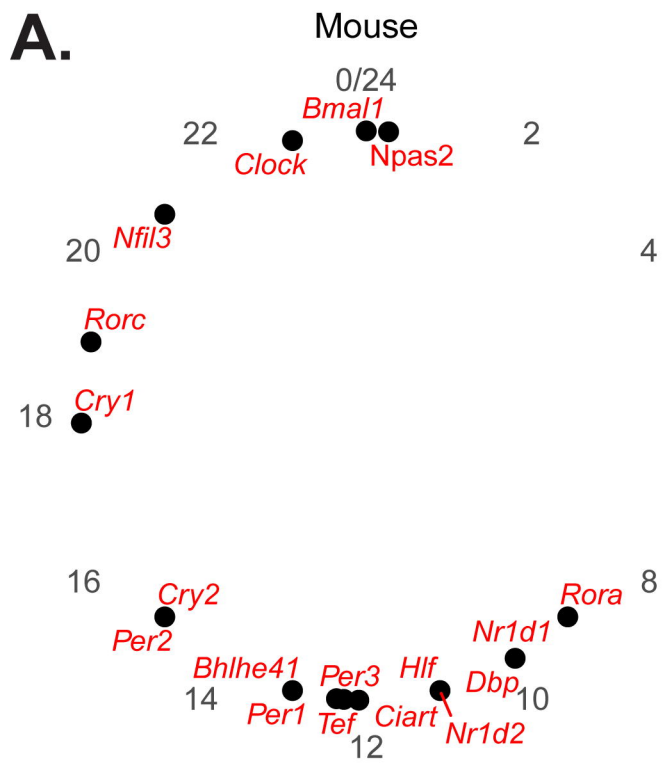
826 **Supplemental Figure 5:** KPC cells expressing dLuciferase driven by the Per2 promoter exhibit circadian
827 activity. Two independent clones were examined, and luciferase activity was measured at 4-hour
828 intervals. Rhythmicity was calculated with Metacycle. Clone 1 (left) was found to be rhythmic with a $q =$
829 5.51E-5 and rAMP = 0.44. Clone 2 (right) was found to be rhythmic with a $q = 1.95\text{E-}7$, rAMP = 0.39.

830 **Supplemental Figure 6:** Cross-validation of mutant *Bmal1* (functional knockout) growth in a second
831 independent clone. A second independent *Bmal1* knockout (BKO) clone with an identical mutation to the
832 first was validated with **A.** western blot analysis. We then heterotopically implanted wildtype (WT) and
833 BKO cells into C57Bl/6J mice and followed **B.** tumor weight and **C.** cell growth. BKO_2 (dark grey; n =
834 20) had a similar mean (\pm standard error) tumor weight to BKO_1 (light grey; n = 20) (426.09 (\pm 40.07)
835 mg vs 438.02 (\pm 48.84) mg; p = 0.85), and both BKO_2 (426.09 (\pm 40.07) mg vs 280.11 (\pm 42.73) mg; p
836 = 0.017) and BKO_1 (438.02 (\pm 48.84) mg vs 280.11 (\pm 42.73) mg; p = 0.02) were larger than WT
837 (white; n = 20). BKO_1 and BKO_2 had similar significantly faster growth trajectories. [ns = not
838 significant, * = p < 0.05, ** = p < 0.01, *** = p < 0.001]

839 **Supplementary Tables**

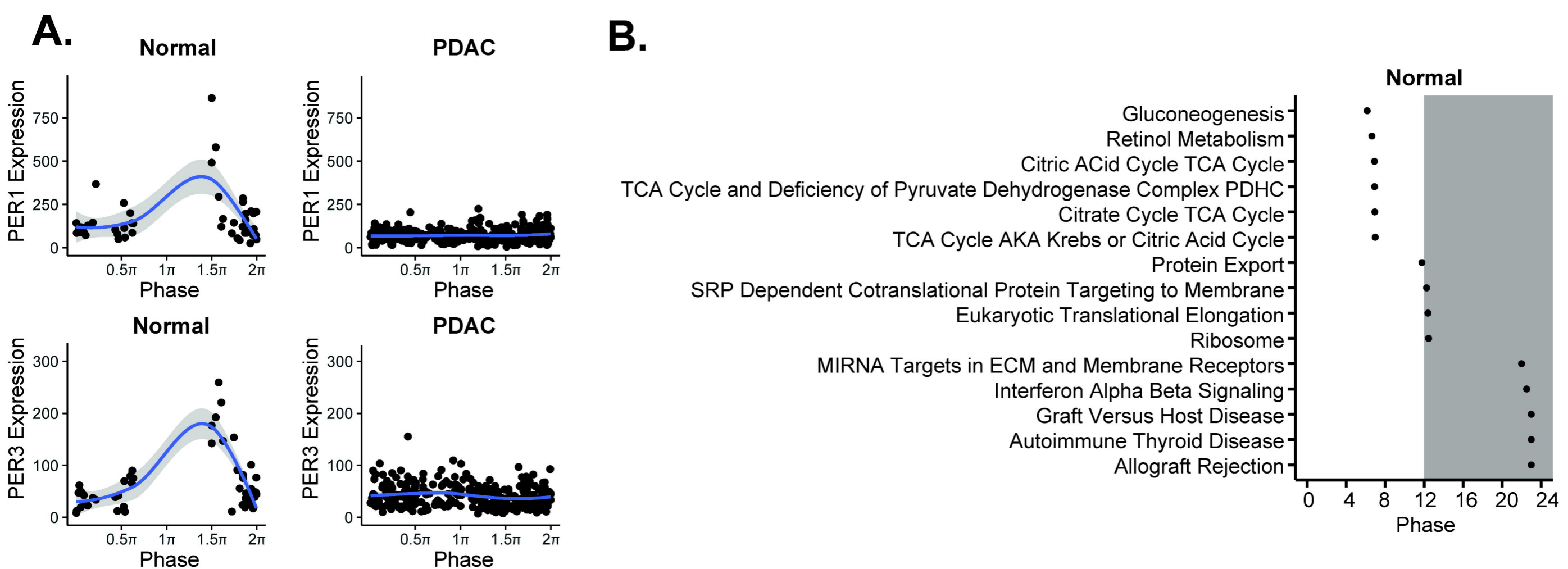
840 **Supplemental Table 1:** Cosinor rhythmicity analysis of CYCLOPS ordered normal pancreas and
841 PDAC

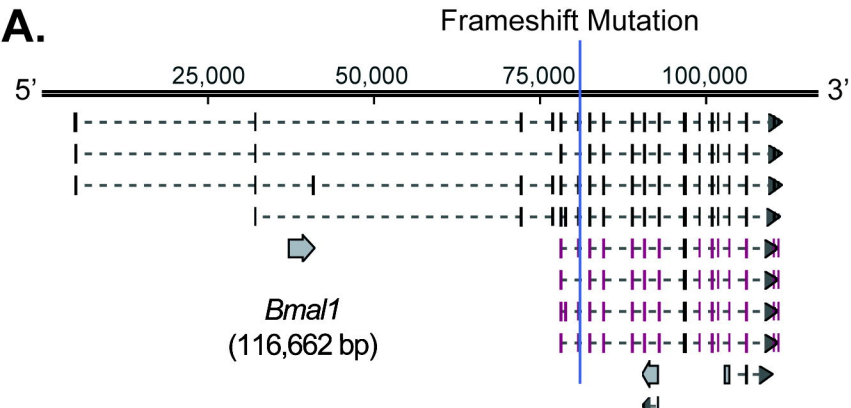
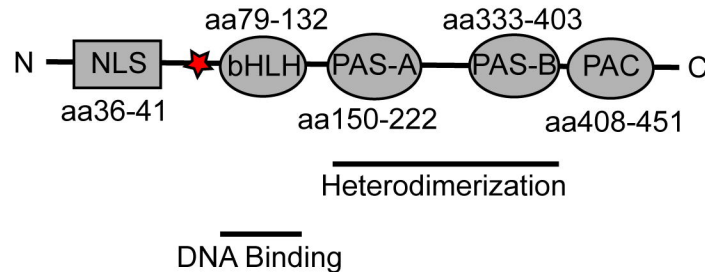
A.**Normal****PDAC****B.**

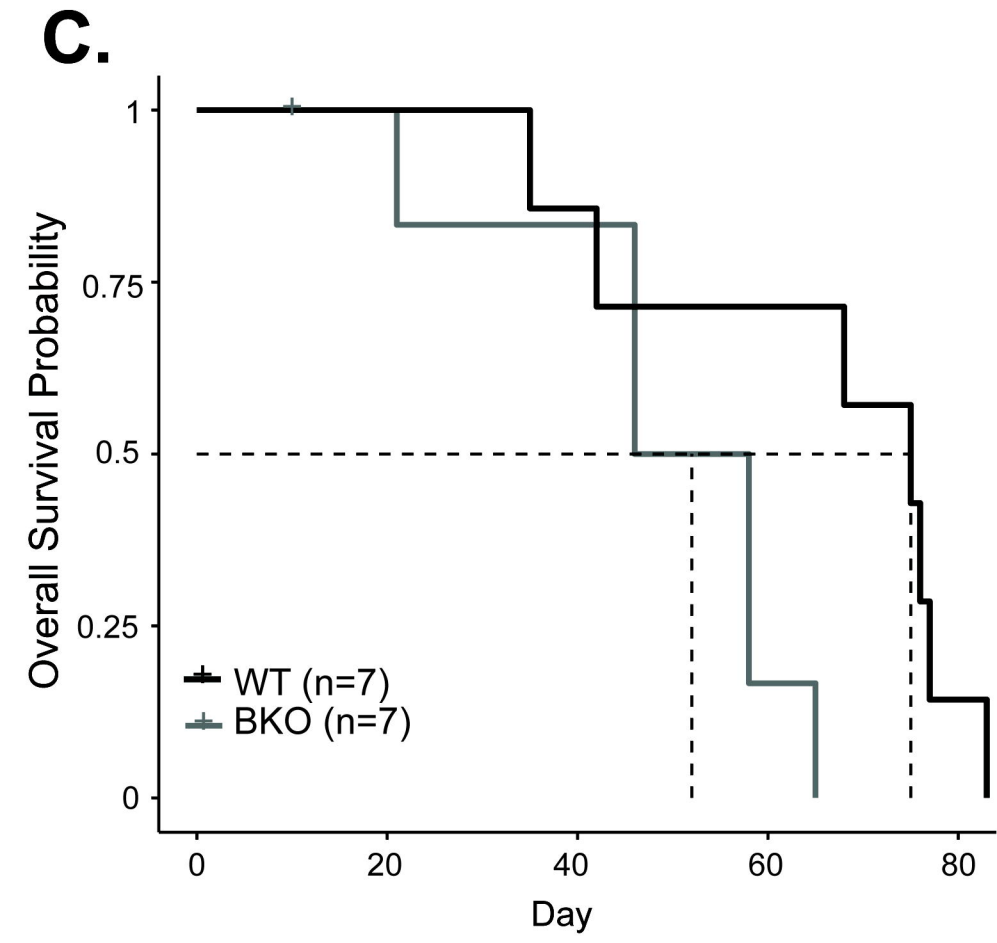
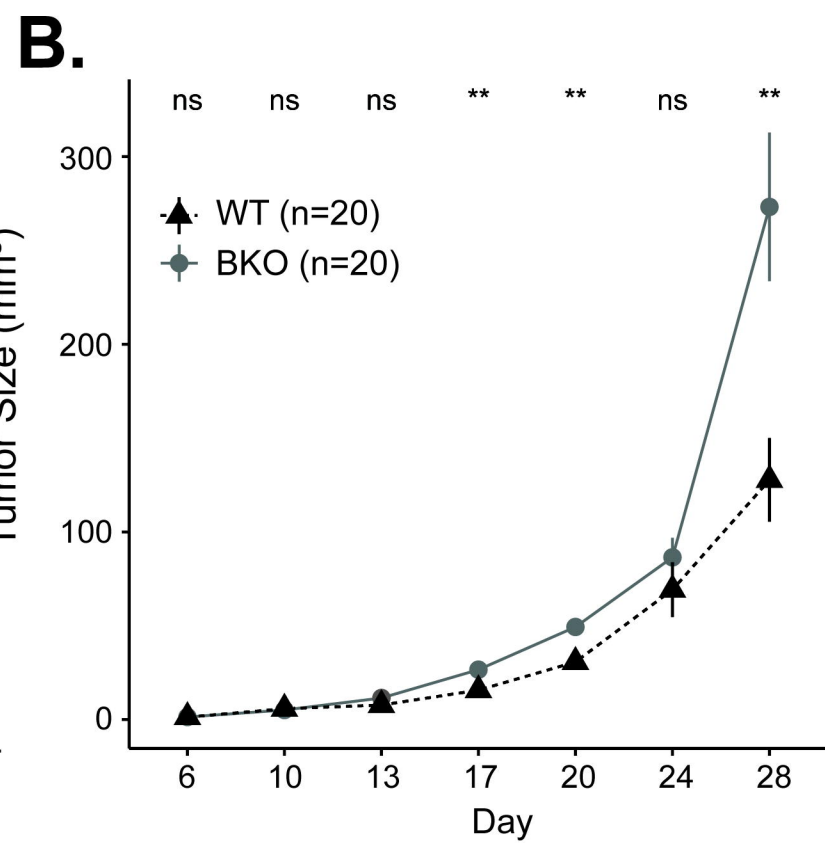
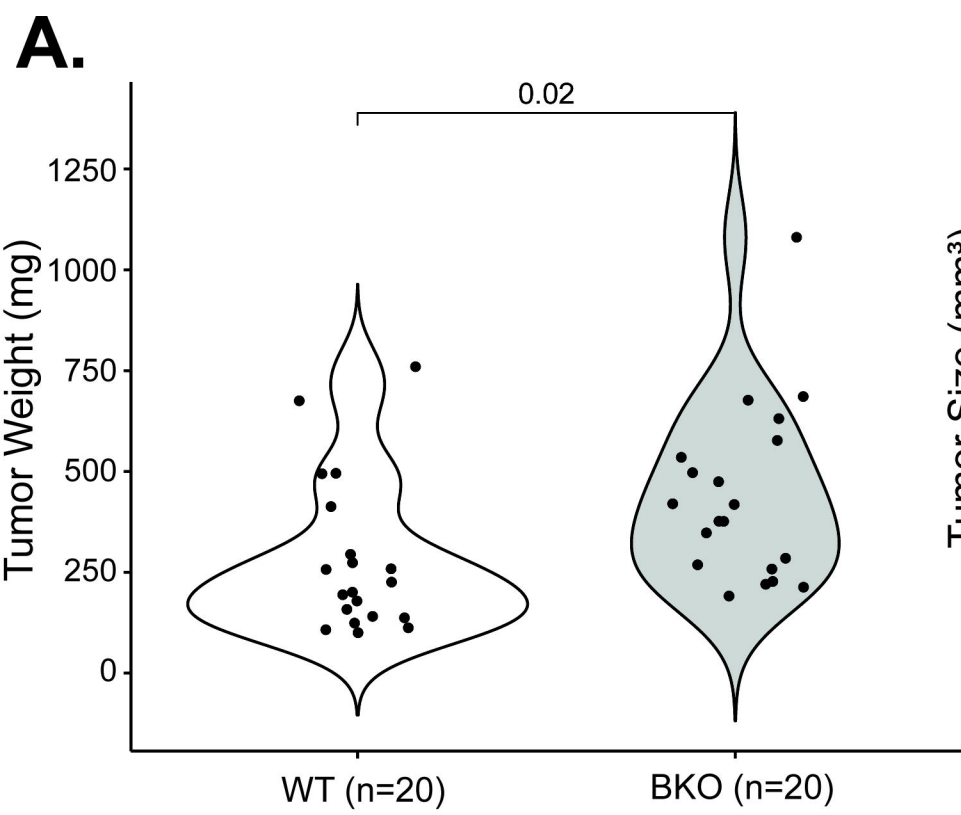


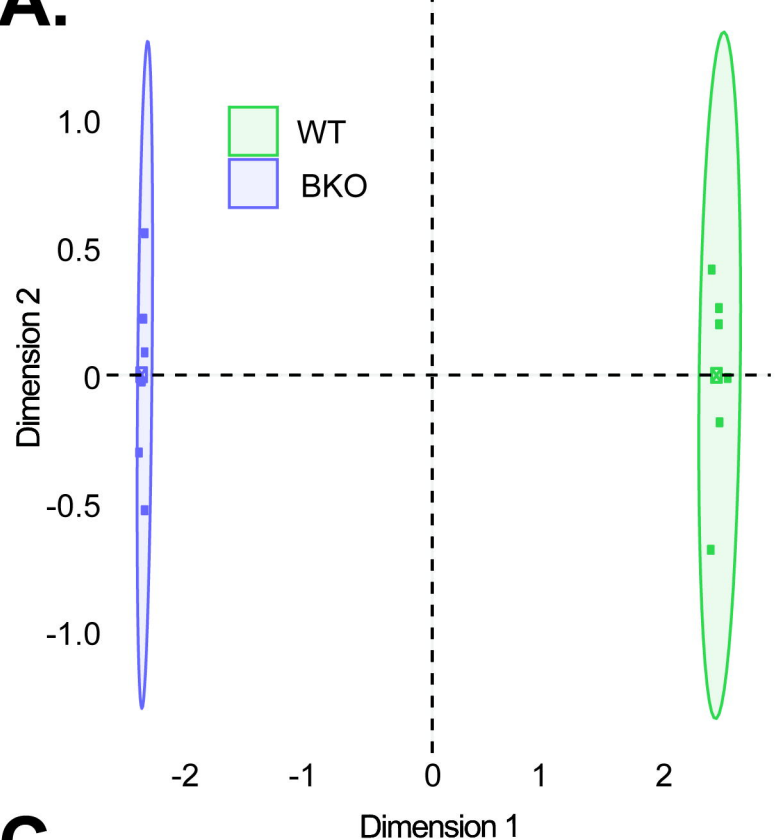
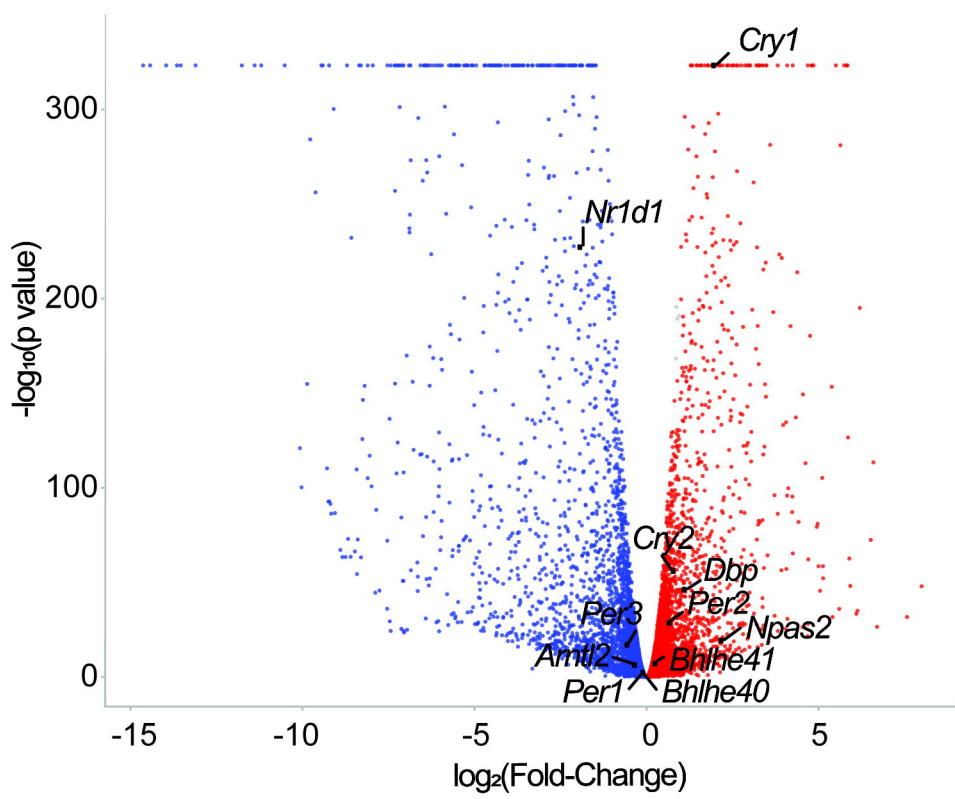
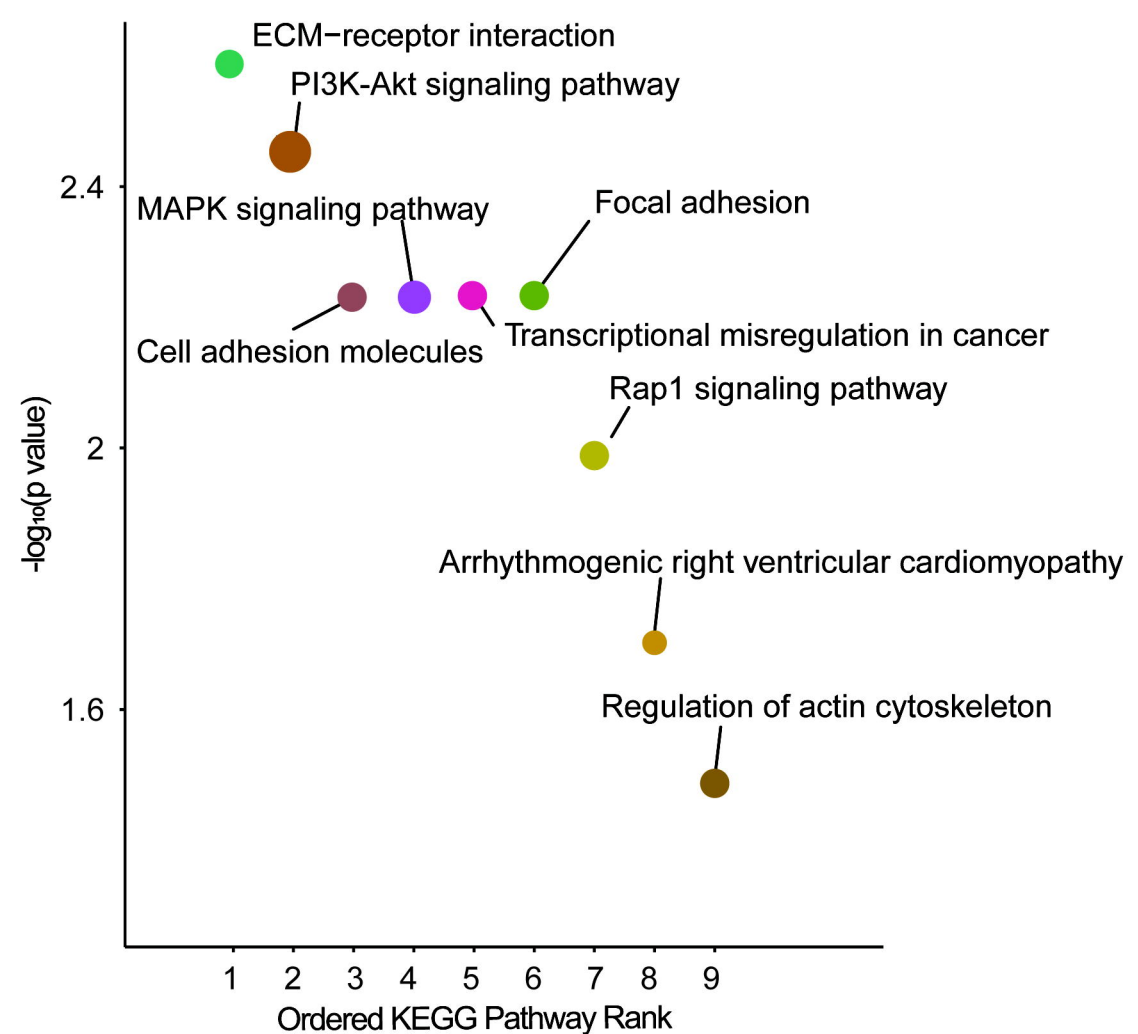
B.

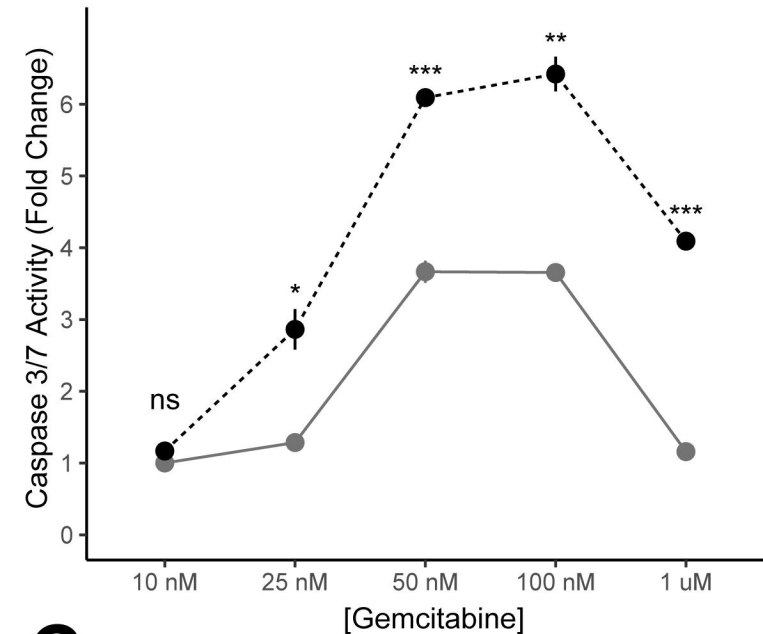
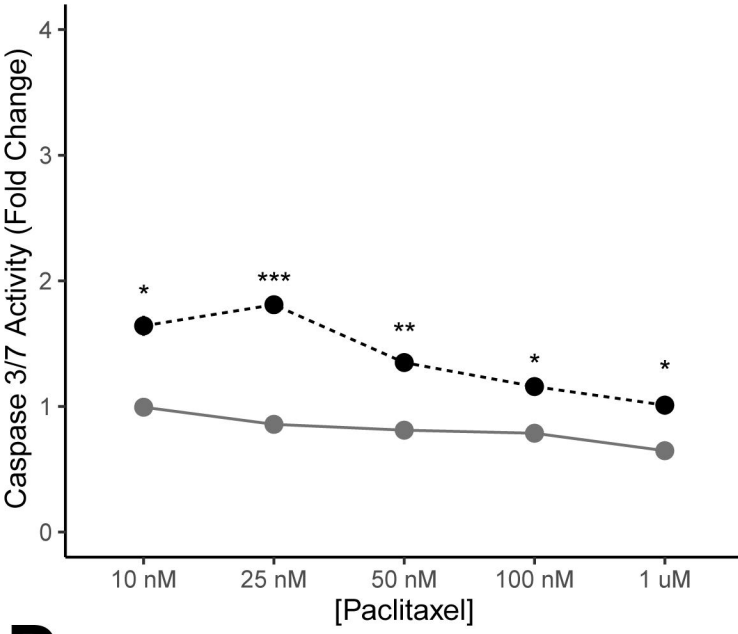
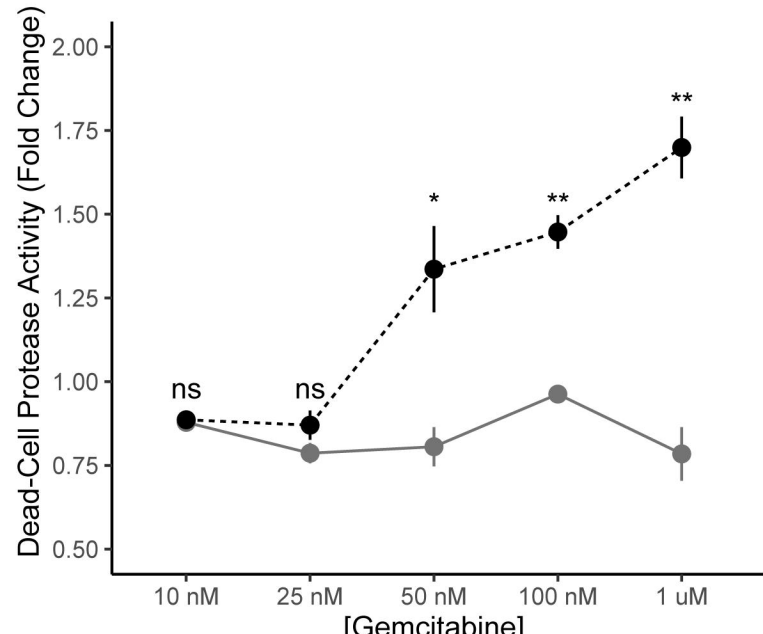
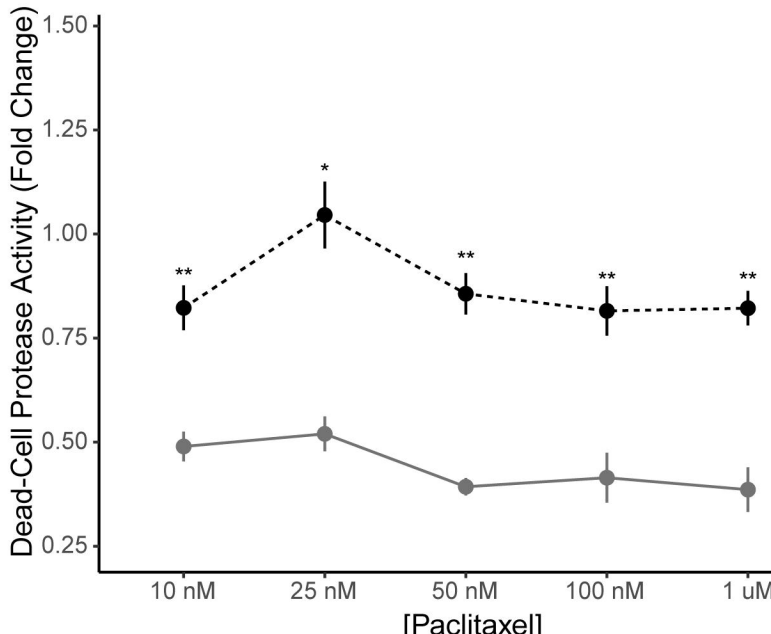
Mouse Gene	Phase	Human Gene	Phase
Bmal1	0	BMAL1	0
Clock	23	CLOCK	4.9
Cry1	17.9	CRY1	1.5
Cry2	15	CRY2	13.6
Per1	13	PER1	11.7
Per2	15	PER2	11.6
Per3	12.4	PER3	11.8
Nr1d1	9	NR1D1	11.9
Nr1d2	11	NR1D2	12.5
Rora	9	RORA	18.8
Rorc	19	RORC	11.7
Npas2	0.3	NPAS2	10.6
Nfil3	21	NFIL3	23.7
Bhlhe41	13	BHLHE41	19.1
Dbp	9.9	DBP	10.9
Ciart	12.1	CIART	10
Tef	12.3	TEF	12
Hlf	11	HLF	18.9



A.**B.**



A.**B.****C.**

A.**B.****C.****D.**

WT (n=20)
BKO (n=20)