## Pan Cancer MK2 Survival Analysis

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Updates in Revision 2:

- 1. We updated the Cox PH model to the model that was used for the LUAD dataset.
- 2. We updated the figures for the HR across cancer datasets to also include metrics of the Cox PH models used in each dataset.

The MasterMK2data.csv file contains MK2 transcript levels + clinical data across a variety of cancer datasets. Please see the MK2\_panCA\_annotation pdf for the code that we used to pull the clinical data from the TCGA. The MK2 transcript data was pulled from OncoLnc (manually).

```
library(knitr)
library(formatR)
opts_chunk$set(tidy.opts=list(width.cutoff=50), tidy=TRUE)
```

## Create MK2 categories (low vs. high)

An added issue here is that we need to define quantiles WITHIN tumor groups. So the quantile for each tumor group is different.

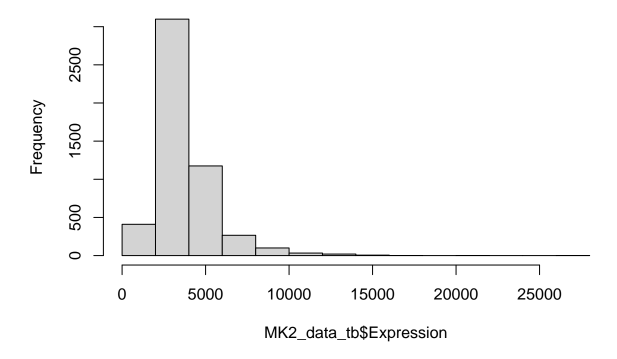
```
MK2 data <- MK2 data[!is.na(MK2 data$Expression), ]</pre>
# create MK2 quantiles
bottom_quant <<- 0.33
top_quant <<- 0.66
BelowBottomQuantile <- function(cancer.type, x) {</pre>
    if (x < quantile(MK2_data[MK2_data$cancer.type.x ==</pre>
        cancer.type, ]$Expression, bottom quant))
        return(1) else return(0)
AboveTopQuantile <- function(cancer.type, x) {
    if (x > quantile(MK2_data[MK2_data$cancer.type.x ==
        cancer.type, ]$Expression, top_quant))
        return(1) else return(0)
MK2_data$MK2_Expression_bottomQ <- mapply(BelowBottomQuantile,
    MK2_data$cancer.type.x, MK2_data$Expression)
MK2_data$MK2_Expression_topQ <- mapply(AboveTopQuantile,
    MK2_data$cancer.type.x, MK2_data$Expression)
MK2_data_tb <- MK2_data
MK2_data_tb$MK2_Expression_topQ <- as.numeric(MK2_data_tb$MK2_Expression_topQ)
MK2_data_tb$MK2_Expression_bottomQ <- as.numeric(MK2_data_tb$MK2_Expression_bottomQ)
```

```
# re-categorize the variable as MK2_tv where 0 =
# lower 10th, 1= top 10th
MK2_data_tb$MK2_tv <- mapply(function(bottomQ, topQ) if (topQ ==
    1) return("High") else return("Low"), MK2_data_tb$MK2_Expression_bottomQ,
    MK2_data_tb$MK2_Expression_topQ)
MK2_data_tb$MK2_tv <- as.factor(MK2_data_tb$MK2_tv)</pre>
MK2_data_tb <- MK2_data_tb[!is.na(MK2_data_tb$time),</pre>
# create a 2 year censor
MK2_data_tb$death_at_2year <- mapply(function(dead,</pre>
    time) if (dead == 1 & time < 366 * 2) return(1) else if (dead ==
    0 & time > 366 * 2) return(0) else if (dead ==
    0 & time < 366 * 2) return(0) else if (dead ==
    1 & time > 366 * 2) return(0), MK2_data_tb$censor,
    as.numeric(MK2_data_tb$time))
MK2_data_tb$death_at_2year <- as.numeric(unlist(as.character(MK2_data_tb$death_at_2year)))
## Warning: NAs introduced by coercion
MK2_data_tb <- MK2_data_tb[MK2_data_tb$death_at_2year ==
    0 | MK2_data_tb$death_at_2year == 1, ]
```

#### Run Cox PH on the entire dataset

So, here we look at the effect of MK2 on the entire dataset, adjusting for cancer type and stage. hist(MK2\_data\_tb\$Expression)

# **Histogram of MK2\_data\_tb\$Expression**



```
MK2_data_tb$Expression_logt <- log(MK2_data_tb$Expression)</pre>
coxph_full <- coxph(Surv(time, death_at_2year) ~ Expression_logt +</pre>
    ajcc_pathologic_stage_combined + age_at_diagnosis +
   gender + smoking + as.factor(cancer.type.x), data = MK2_data_tb)
cox.zph(coxph_full)
##
                                  chisq df
                                                р
## Expression logt
                                  0.453 1
                                              0.50
## ajcc_pathologic_stage_combined 0.557 1
                                             0 46
## age_at_diagnosis
                               17.907 1 2.3e-05
                                 1.237 1
## gender
                                             0.27
## smoking
                                  1.343 1
                                             0.25
## as.factor(cancer.type.x)
                                 46.437 12 5.8e-06
## GLOBAL
                                 67.180 17 6.6e-08
summary(coxph_full)
## Call:
## coxph(formula = Surv(time, death_at_2year) ~ Expression_logt +
      ajcc_pathologic_stage_combined + age_at_diagnosis + gender +
##
      smoking + as.factor(cancer.type.x), data = MK2_data_tb)
##
    n= 5078, number of events= 989
##
##
      (35 observations deleted due to missingness)
##
                                               coef exp(coef) se(coef)
##
## Expression_logt
                                           0.17235 1.18810 0.09891 1.742
## ajcc_pathologic_stage_combinedLate Stage 1.03678
                                                     2.82011 0.07077 14.650
                                           0.02365 1.02393 0.00299 7.910
## age at diagnosis
## gendermale
                                           0.05464
                                                     1.05616 0.06985 0.782
## smoking1
                                           -0.08319 0.92018 0.09720 -0.856
## as.factor(cancer.type.x)BRCA
                                          ## as.factor(cancer.type.x)COAD
                                                     0.47101 0.16045 -4.692
                                          -0.75287
## as.factor(cancer.type.x)ESCA
                                           0.29541
                                                     1.34367 0.18692 1.580
## as.factor(cancer.type.x)KIRC
                                          -0.72921
                                                     0.48229 0.15386 -4.739
## as.factor(cancer.type.x)KIRP
                                          -1.06740 0.34390 0.23383 -4.565
                                                     1.16859 0.15296 1.019
## as.factor(cancer.type.x)LIHC
                                           0.15580
## as.factor(cancer.type.x)LUAD
                                          -0.13029
                                                     0.87784 0.14033 -0.928
## as.factor(cancer.type.x)LUSC
                                          0.12708 1.13551 0.13467 0.944
## as.factor(cancer.type.x)PAAD
                                                     3.09736 0.15347 7.367
                                          1.13055
## as.factor(cancer.type.x)READ
                                          -1.31750
                                                     0.26780 0.29513 -4.464
## as.factor(cancer.type.x)SKCM
                                          -0.64071
                                                     0.52692 0.15691 -4.083
## as.factor(cancer.type.x)STAD
                                           0.20768
                                                     1.23082 0.13384 1.552
##
                                          Pr(>|z|)
## Expression logt
                                            0.0814 .
## ajcc_pathologic_stage_combinedLate Stage < 2e-16 ***
## age_at_diagnosis
                                           2.57e-15 ***
## gendermale
                                            0.4340
## smoking1
                                            0.3921
## as.factor(cancer.type.x)BRCA
                                           < 2e-16 ***
## as.factor(cancer.type.x)COAD
                                          2.70e-06 ***
## as.factor(cancer.type.x)ESCA
                                            0.1140
```

```
## as.factor(cancer.type.x)KIRC
                                            2.14e-06 ***
## as.factor(cancer.type.x)KIRP
                                            5.00e-06 ***
## as.factor(cancer.type.x)LIHC
                                              0.3084
## as.factor(cancer.type.x)LUAD
                                              0.3532
## as.factor(cancer.type.x)LUSC
                                              0.3454
## as.factor(cancer.type.x)PAAD
                                            1.75e-13 ***
## as.factor(cancer.type.x)READ
                                            8.04e-06 ***
                                            4.44e-05 ***
## as.factor(cancer.type.x)SKCM
## as.factor(cancer.type.x)STAD
                                              0.1207
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
                                            exp(coef) exp(-coef) lower .95
                                                                     0.9787
## Expression_logt
                                               1.1881
                                                           0.8417
                                               2.8201
                                                           0.3546
                                                                     2.4549
## ajcc_pathologic_stage_combinedLate Stage
## age_at_diagnosis
                                               1.0239
                                                           0.9766
                                                                     1.0179
## gendermale
                                               1.0562
                                                           0.9468
                                                                     0.9210
## smoking1
                                               0.9202
                                                           1.0867
                                                                     0.7606
                                                           6.1829
                                                                     0.1080
## as.factor(cancer.type.x)BRCA
                                               0.1617
## as.factor(cancer.type.x)COAD
                                               0.4710
                                                           2.1231
                                                                     0.3439
                                               1.3437
## as.factor(cancer.type.x)ESCA
                                                          0.7442
                                                                     0.9315
## as.factor(cancer.type.x)KIRC
                                               0.4823
                                                           2.0734
                                                                     0.3567
## as.factor(cancer.type.x)KIRP
                                                          2.9078
                                                                     0.2175
                                               0.3439
## as.factor(cancer.type.x)LIHC
                                                           0.8557
                                                                     0.8659
                                               1.1686
## as.factor(cancer.type.x)LUAD
                                                                     0.6668
                                               0.8778
                                                          1.1392
## as.factor(cancer.type.x)LUSC
                                               1.1355
                                                          0.8807
                                                                     0.8721
## as.factor(cancer.type.x)PAAD
                                               3.0974
                                                           0.3229
                                                                     2.2928
## as.factor(cancer.type.x)READ
                                               0.2678
                                                           3.7341
                                                                     0.1502
## as.factor(cancer.type.x)SKCM
                                               0.5269
                                                          1.8978
                                                                     0.3874
## as.factor(cancer.type.x)STAD
                                               1.2308
                                                           0.8125
                                                                     0.9468
##
                                            upper .95
## Expression_logt
                                               1.4423
## ajcc_pathologic_stage_combinedLate Stage
                                               3.2397
                                               1.0299
## age_at_diagnosis
## gendermale
                                               1.2111
## smoking1
                                               1.1133
## as.factor(cancer.type.x)BRCA
                                               0.2423
## as.factor(cancer.type.x)COAD
                                               0.6451
## as.factor(cancer.type.x)ESCA
                                               1.9382
## as.factor(cancer.type.x)KIRC
                                               0.6520
## as.factor(cancer.type.x)KIRP
                                               0.5438
## as.factor(cancer.type.x)LIHC
                                               1.5771
## as.factor(cancer.type.x)LUAD
                                               1.1558
## as.factor(cancer.type.x)LUSC
                                               1.4785
## as.factor(cancer.type.x)PAAD
                                               4.1843
## as.factor(cancer.type.x)READ
                                               0.4776
## as.factor(cancer.type.x)SKCM
                                               0.7166
## as.factor(cancer.type.x)STAD
                                               1.6000
## Concordance= 0.741 (se = 0.007)
## Likelihood ratio test= 751.5 on 17 df,
                                             p=<2e-16
## Wald test
                       = 629.1 on 17 df,
                                             p=<2e-16
## Score (logrank) test = 753.2 on 17 df,
                                             p=<2e-16
```

## Collect Cox PH info

Now, we run our pre-specified Cox PH model iteratively, for each dataset, and the functions below return either the model result or model metrics.

```
# Some functions to pull CoxPH models and metrics
# for all cancers.
returnModel <- function(cancer.type) {</pre>
    surv_df <- MK2_data_tb[MK2_data_tb$cancer.type.x ==</pre>
        cancer.type, ]
    MK2_model <- coxph(Surv(time, death_at_2year) ~</pre>
        MK2_Expression_topQ + ajcc_pathologic_stage_combined +
            age_at_diagnosis + gender + smoking, data = surv_df)
    return(MK2_model)
}
returnModel_metrics <- function(cancer.type) {</pre>
    surv_df <- MK2_data_tb[MK2_data_tb$cancer.type.x ==</pre>
        cancer.type, ]
    MK2_model <- coxph(Surv(time, death_at_2year) ~</pre>
        MK2 Expression topQ + ajcc pathologic stage combined +
            age_at_diagnosis + gender + smoking, data = surv_df)
    return(list(anova(MK2_model), cox.zph(MK2_model)))
}
returnModel_MK2cont <- function(cancer.type) {</pre>
    surv_df <- MK2_data_tb[MK2_data_tb$cancer.type.x ==</pre>
        cancer.type, ]
    MK2_model <- coxph(Surv(time, death_at_2year) ~</pre>
        Expression_logt + ajcc_pathologic_stage_combined +
            age_at_diagnosis + gender + smoking, data = surv_df)
    return(MK2_model)
}
returnModel MK2cont metrics <- function(cancer.type) {</pre>
    surv_df <- MK2_data_tb[MK2_data_tb$cancer.type.x ==</pre>
        cancer.type, ]
    MK2_model <- coxph(Surv(time, death_at_2year) ~</pre>
        Expression_logt + ajcc_pathologic_stage_combined +
            age_at_diagnosis + gender + smoking, data = surv_df)
    return(list(anova(MK2_model), cox.zph(MK2_model)))
```

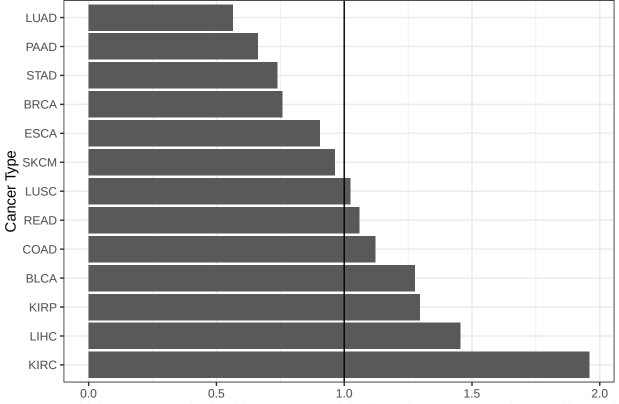
Now that we have some prepared functions, we use them to call the model and retrieve model metrics.

Cox PH Model 1: MK2 as a dichotomous variable + covariates (including stage) Cox PH Model 2: MK2 as a continuous variable + covariates (including stage)

Cancer type is not included because, in essence, we're stratifying by cancer type here.

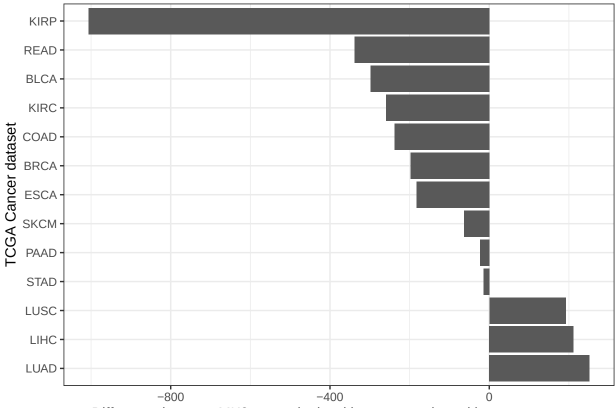
```
# return model fit parameters (cox.zph)
MK2_model_df <- MK2_means_wide</pre>
# model MK2 as a dichotomous var
MK2_model_df$model1.results <- lapply(MK2_model_df$cancer.type.x,
    returnModel)
MK2_model_df$model1.metrics <- lapply(MK2_model_df$cancer.type.x,
    returnModel_metrics)
MK2 model df$cox.zph MK2 <- sapply(MK2 model df$model1.metrics,
    function(x) return(x[[2]][[1]][13]))
MK2_model_df$cox.zph_global <- sapply(MK2_model_df$model1.metrics,
    function(x) return(x[[2]][[1]][18]))
MK2_model_df$wald_pval <- sapply(MK2_model_df$model1.results,</pre>
    function(x) return(broom::glance(x)$p.value.wald))
MK2_model_df$MK2 <- lapply(MK2_model_df$model1.results,</pre>
    function(x) return((x$coefficients[[1]][1])))
MK2_model_df$MK2.lci <- lapply(MK2_model_df$model1.results,
    function(x) return(exp(confint(x)[1])))
MK2_model_df$MK2.uci <- lapply(MK2_model_df$model1.results,
    function(x) return(exp(confint(x)[6])))
MK2_model_df$MK2 <- exp(as.numeric(MK2_model_df$MK2))</pre>
# model MK2 as a continuous var
MK2_model_df$model2.results <- lapply(MK2_model_df$cancer.type.x,
    returnModel_MK2cont)
MK2_model_df$MK2_model2 <- lapply(MK2_model_df$model2.results,
    function(x) return(x$coefficients[[1]][1]))
MK2_model_df$MK2_model2.lci <- lapply(MK2_model_df$model2.results,
    function(x) return(exp(confint(x)[1])))
MK2_model_df$MK2_model2.uci <- lapply(MK2_model_df$model2.results,
    function(x) return(exp(confint(x)[6])))
MK2_model_df$MK2_model2 <- exp(as.numeric(MK2_model_df$MK2_model2))</pre>
MK2_model_df$model2.metrics <- lapply(MK2_model_df$cancer.type.x,</pre>
    returnModel MK2cont metrics)
MK2_model_df$cox.zph_MK2_model2 <- sapply(MK2_model_df$model2.metrics,
    function(x) return(x[[2]][[1]][13]))
MK2_model_df$cox.zph_global_model2 <- sapply(MK2_model_df$model2.metrics,
    function(x) return(x[[2]][[1]][18]))
MK2_model_df$wald_pval_model2 <- sapply(MK2_model_df$model2.results,
    function(x) return(broom::glance(x)$p.value.wald))
```

# Graphical output: graphs of MK2 transcript levels and HR for MK2 across datasets



HR for high MK2 transcript level in Cox PH model adjusted for stage, gender, smoking and

```
ggplot(MK2_model_df, aes(x = reorder(cancer.type.x,
    -diff), y = diff)) + geom_bar(stat = "identity") +
    theme_bw() + coord_flip() + ylab("Difference in mean MK2 transcript level between early and late st
    xlab("TCGA Cancer dataset")
```

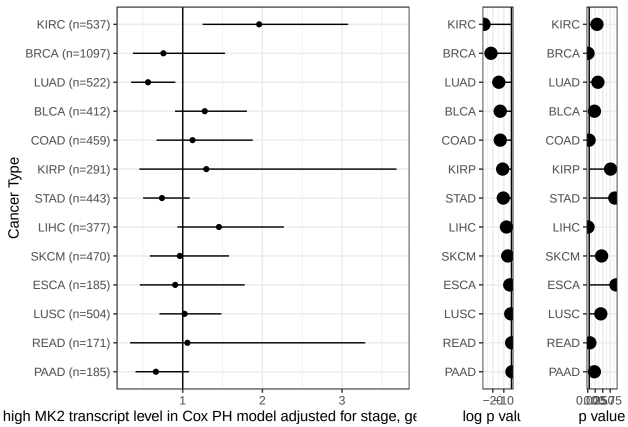


Difference in mean MK2 transcript level between early and late stage cancer

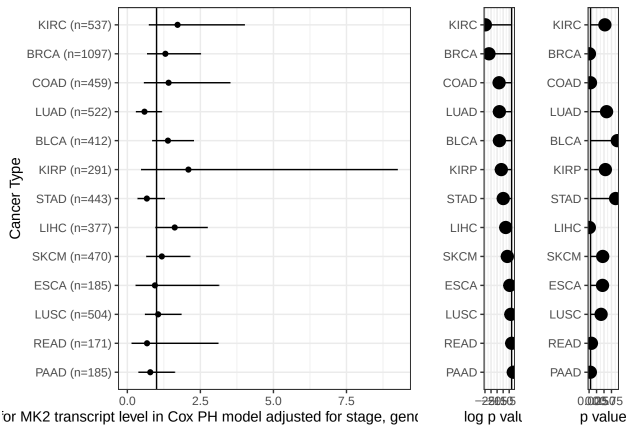
```
MK2_model_df$cancer.type.x.label <- sapply(MK2_model_df$cancer.type.x,
    function(x) return(pts_percatype[pts_percatype$cancer_type ==
        x, ]$num))
MK2_model_df$cancer.type.x.label <- paste0(MK2_model_df$cancer.type.x,
    " (n=", MK2_model_df$cancer.type.x.label, ")")
# Model 1: MK2 as a dichotomous variable
HR_plot <- ggplot(MK2_model_df, aes(x = reorder(cancer.type.x.label,</pre>
    -wald_pval), y = as.numeric(MK2))) + geom_point() +
    geom_errorbar(aes(ymin = as.numeric(MK2.lci), ymax = as.numeric(MK2.uci)),
        width = 0) + geom_hline(yintercept = 1) + theme_bw() +
    ylab("HR for high MK2 transcript level in Cox PH model adjusted for stage, gender, smoking and age"
    xlab("Cancer Type") + coord_flip()
model_stats_df <- MK2_model_df[c("cancer.type.x", "cox.zph_MK2",</pre>
    "wald pval")]
model_stats_df <- arrange(model_stats_df, wald_pval)</pre>
metrics_plot_wald <- ggplot(model_stats_df, aes(x = reorder(cancer.type.x,</pre>
    -wald_pval), y = log(wald_pval))) + geom_point(size = 4) +
    geom_hline(yintercept = log(0.05)) + geom_segment(aes(x = cancer.type.x,
    xend = cancer.type.x, y = log(0.05), yend = log(wald_pval))) +
    theme_bw() + xlab("") + ylab("log p value") + coord_flip()
metrics_plot_PH <- ggplot(model_stats_df, aes(x = reorder(cancer.type.x,</pre>
```

```
-wald_pval), y = cox.zph_MK2)) + geom_point(size = 4) +
geom_hline(yintercept = 0.05) + geom_segment(aes(x = cancer.type.x,
xend = cancer.type.x, y = 0.05, yend = cox.zph_MK2)) +
theme_bw() + xlab("") + ylab("p value") + coord_flip()

gridExtra::grid.arrange(HR_plot, metrics_plot_wald,
metrics_plot_PH, ncol = 3, widths = c(4, 1, 1))
```



theme\_bw() + xlab("") + ylab("log p value") + coord\_flip()



#### R Markdown

This is an R Markdown document. Markdown is a simple formatting syntax for authoring HTML, PDF, and MS Word documents. For more details on using R Markdown see http://rmarkdown.rstudio.com.

When you click the **Knit** button a document will be generated that includes both content as well as the output of any embedded R code chunks within the document. You can embed an R code chunk like this:

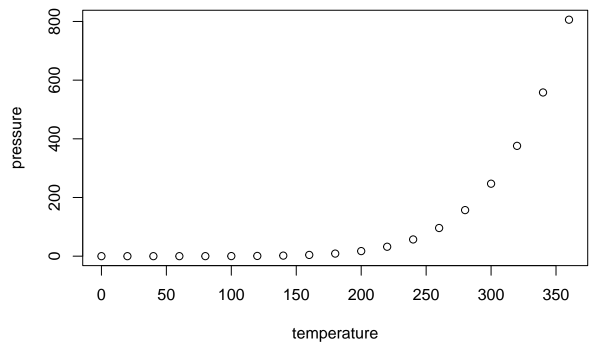
## summary(cars)

```
##
        speed
                         dist
##
           : 4.0
                           : 2.00
    Min.
                   Min.
   1st Qu.:12.0
                   1st Qu.: 26.00
##
   Median:15.0
                   Median : 36.00
##
##
    Mean
           :15.4
                   Mean
                           : 42.98
    3rd Qu.:19.0
                   3rd Qu.: 56.00
```

## Max. :25.0 Max. :120.00

## **Including Plots**

You can also embed plots, for example:



Note that the echo = FALSE parameter was added to the code chunk to prevent printing of the R code that generated the plot.