# An Introduction to the 'PICS' Package, Version 1.0

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### 1 Overview

This vignette provides basic information about the PICS [1] package. PICS stands for "Pathway-guided Identification of Cancer Subtypes". The proposed approach improves identification of molecularly-defined subgroups of cancer patients by utilizing information from pathway databases in the following four aspects.

- (1) Integration of genomic data at the pathway-level improves robustness and stability in identification of cancer subgroups and driver molecular features;
- (2) Summarizing multiple genes and genomic platforms at the pathway-level can potentially improve statistical power to identify important driver pathways because moderate signals in multiple genes can be aggregated;
- (3) In PICS, we consider this "cooperation" or "interaction" between pathways, instead assuming that each pathway operates independently during the cancer progression, which may be unrealistic;
- (4) PICS allows simultaneous inference in multiple biological layers (pathway clusters, pathways, and genes) within a statistically rigorous and unified framework without any additional laborious downstream analysis.

The package can be loaded with the command:

R> library("PICS")

# 2 Input Data

The package requires that the response consist of 4 components: (1) z-scores in the form of a either data frame or matrix; and (2) survival time and censoring indicator in the form of vectors; and (3) pathway information is in the form of a list where gene names are included in pathway lists respectively.

The Cancer Genome Atlas (TCGA) will be used to illustrate the 'PICS' package. The TCGA data was downloaded from the cBio Portal (http://www.cbioportal.org/) using the R package 'cgdsr', and we used z scores for the mRNA expression data.

The 'PICS' package provided an example data 'TCGA'. The 'TCGA' is a list object with four elements, including the 'geneexpr' data frame of z scores for the mRNA expression, the 't' vector of the survival time, the 'd' vector of the survival status indicator, and the 'pathList' list of the pathway information. The 'pathList' has four elements, each of which contains names of genes belonging to each pathway. Z scores for the mRNA expression data of 389 genes are provided for 50 randomly selected high-grade serous ovarian cancer patients, along with survival times and censoring statuses.

This dataset can be loaded as follows:

```
R> data(TCGA)
```

R> TCGA\$geneexpr[1:5,1:5]

```
ACLY AC01 AC02 CS DLAT
1 -2.2410125 -0.48445531 -1.6346455 0.1378804 -3.5310321
2 -2.1301362 0.82116427 -0.9533701 0.6213512 0.6689948
3 -2.9122727 -0.08790649 -1.0975096 -0.2454025 -0.9433900
4 -1.1721514 -0.24249825 -0.7212639 0.1842386 -0.6188785
5 0.5383438 0.98012739 -0.7396043 -0.0699680 1.9573767
```

R> TCGA\$t[1:5]

[1] 43.89 40.97 49.12 2.00 46.59

R> TCGA\$d[1:5]

[1] 1 1 0 1 0

R> TCGA\$pathList[1]

#### \$KEGG\_CITRATE\_CYCLE\_TCA\_CYCLE

[1]	"IDH3B"	"DLST"	"PCK2"	"CS"	"PDHB"	"PCK1"
[7]	"PDHA1"	"L0C642502"	"PDHA2"	"L0C283398"	"FH"	"SDHD"
[13]	"OGDH"	"SDHB"	"IDH3A"	"SDHC"	"IDH2"	"IDH1"
[19]	"ACO1"	"ACLY"	"MDH2"	"DLD"	"MDH1"	"DLAT"
[25]	"OGDHL"	"PC"	"SDHA"	"SUCLG1"	"SUCLA2"	"SUCLG2"
[31]	"IDH3G"	"ACO2"				

# 3 Pre-filtering

To refine the candidate set of genes, we first conduct a supervised pre-filtering by fitting a Cox regression model of each mRNA expression measure on patient survival in the TCGA dataset. Only the gene expressions associated with patient survival at p-values smaller than a pre-specified cut-off are included in the subsequent analysis. By default, p = 0.5 is used as cut-off point.

```
Summary: Pre-filtering results (class: Prefiltered)
-----
Number of genes before prefiltering: 389
Number of genes after prefiltering: 213
```

## 4 Gene Selection

In order to select key genes associated with patient survivals and effectively summarize them by taking into account correlation among them, we fit a sparse partial least squares (SPLS) Cox regression model [3] of patient survivals on gene expression measurements for each pathway.

Using the 'prefilter.results', gene-level analysis result can be generated with 'selectGene' function.

```
R> gene.results <- selectGene(prefilter.results)

R> gene.results

Summary: Gene-level analysis results (class: FitGene)
------

Number of prefiltered genes: 213

Number of selected genes: 132
```

The list of the SPLS regression coefficients of cancer related genes can be generated using the function coef().

R> head(coef(gene.results)[[1]])

```
colnames.xx. spls.mod.betahat
          ACLY
                       0.000000
1
2
             CS
                       0.000000
3
          DLAT
                       0.000000
4
           DLD
                       0.000000
5
          MDH1
                      -0.3560516
6
          PCK1
                      -0.1988449
```

There are two main tuning parameters: 'eta' represents the sparsity tuning parameter and 'K' is the number of hidden (latent) components. Parameters can be chosen by (v-fold) cross-validation. The user can search the range for these parameters and the cross-validation procedure searches within these ranges. Note that 'eta' should have a value between 0 and 1. 'K' is integer valued and can range between 1 and  $min\{p,(v-1)n/v\}$ , where p is the number of genes and n is the sample size. For example, if 10-fold cross-validation is used (default), 'K' should be smaller than  $min\{p,0.9n\}$ . For the TCGA data, we set fold as 5, 'K' as 5, and search for 'eta' between 0.1 and 0.9 with the following command:

## 5 Pathway Selection

Next, in order to identify a parsimonious set of pathways associated with patient survivals, we fit a LASSO-penalized Cox regression [4] on latent components derived from all the pathways. Specifically, a pathway was selected if at least one of its latent components had non-zero LASSO coefficient estimate.

This approach has the following two strengths: First, the latent components generated from the SPLS step preserve pathway structure and also reflect correlation among genes and their association

with survival outcomes. Second, this approach can potentially improve the stability of estimation in the subsequent analysis.

Using the 'gene.results', pathway-level analysis result can be generated with 'selectPath' function.

LASSO regression coefficients of cancer related pathways can be generated using the function coef().

R> head(coef(path.results))

```
rep.pathways..cols. path.beta

KEGG_CITRATE_CYCLE_TCA_CYCLE 0.44388849

KEGG_CITRATE_CYCLE_TCA_CYCLE 0.00000000

KEGG_CITRATE_CYCLE_TCA_CYCLE 0.00000000

KEGG_CITRATE_CYCLE_TCA_CYCLE 0.00000000

KEGG_MAPK_SIGNALING_PATHWAY 0.08378862

KEGG_TGF_BETA_SIGNALING_PATHWAY 0.37094205
```

Hazard ratio plot associated with each latent component in selected pathways can be generated using the function plot() with the argument type="HR".

```
R> plot(path.results, type="HR")
```

Figure 1 shows the hazard ratio (HR) associated with each latent component in the pathways selected by the PICS. Based on the TCGA data, pathways with the largest effect on survival ( $HR \ge 1.15$ ) are KEGG\_CITRATE\_CYCLE\_TCA\_CYCLE and KEGG\_TGF\_BETA\_SIGNALING\_PATHWAY pathways.

## 6 Risk Group Prediction

Risk group predictions can be made using the function predict()

```
R> predicted <- predict(path.results)</pre>
```

The function 'predict' returns the following output: (1) risk.index: number of pathways with elevated activity. (2) riskcat: predicted risk group. (3) cuts: cut off to determine low, intermediate and high risk group.

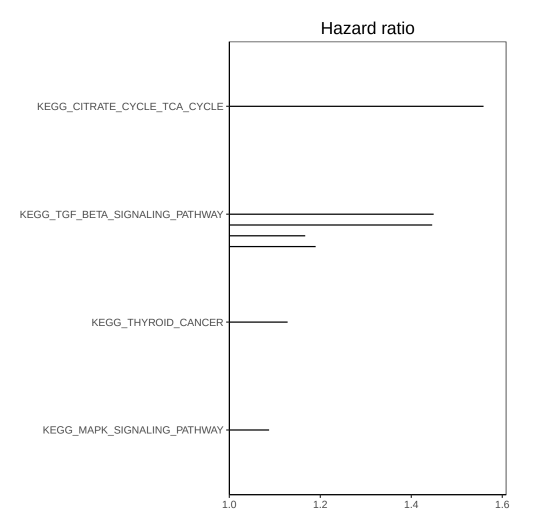


Figure 1: Hazard ratio (HR) associated with each latent component in the pathways selected by the PICS

#### R> predicted[1:3]

#### \$risk.index

[1] 3 4 0 4 2 4 4 0 0 1 4 0 2 3 3 3 0 1 0 2 1 0 1 1 2 2 2 0 0 0 2 0 1 0 0 0 1 0 [39] 4 4 2 4 4 4 4 3 4 3 3

### \$riskcat

- [1] "med" "high" "low" "high" "med" "high" "high" "low" "low" "med"
- [11] "high" "low" "med" "med" "med" "low" "med" "low" "med"
- [21] "med" "low" "med" "med" "med" "med" "low" "low" "low"
- [31] "med" "low" "med" "low" "low" "low" "med" "low" "high" "high"
- [41] "med" "high" "high" "high" "high" "med" "high" "med" "med"

#### \$cuts

[1] 0.00 3.75

## 7 Survival Curve

The predictive performance of PICS method can be presented by Kaplan-Meier curves. Kaplan-Meier curves of predicted patient subgroups based on the PICS approach can be generated with plot() function with argument type="KM".

### R> plot(path.results,type="KM")

Figure 2 shows the Kaplan-Meier curves of predicted patient subgroups based on the PICS approach. The PICS approach successfully distinguish the high, intermediate and low risk group from each other.

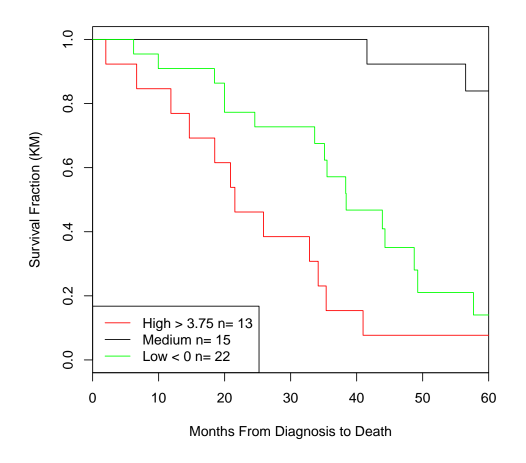


Figure 2: The observed survival curves for patient subgroups identified by the PICS

### 8 Survival ROC

The predictive performance of PICS method can be further evaluated based on area under the time dependent receiver operating curve (ROC). ROC plot can be generated using plot() function with argument type="ROC".

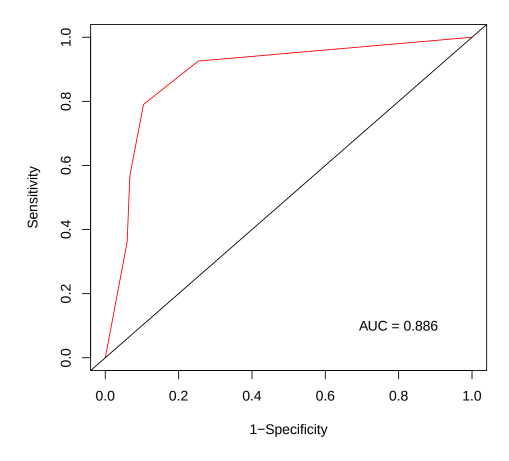


Figure 3: Time dependent receiver operating curve

Figure 3 shows the ROC curves for survival, and for the TCGA data, the area under curve (AUC) associated with the PICS approach was 0.886.

### References

- [1] Wei W., Zequn S., Willian S., Zhenning Y., Andrew L., Gary H., Linda K., Dongjun C. (2017), "PICS: Pathway-guided identification of cancer subtypes". (submitted).
- [2] Cancer Genome Atlas Research Network (2011), "Integrated genomic analyses of ovarian carcinoma". Nature, 474(7353), 609-615.
- [3] Bastien, P., Bertrand, F., Meyer, N., Maumy-Bertrand, M. (2014), "Deviance residuals-based sparse PLS and sparse kernel PLS regression for censored data". Bioinformatics, 31(3), 397-404.
- [4] Tibshirani, R. (1997), "The lasso method for variable selection in the cox model". Statistics in Medicine, 16(4), 385-395.