survcomp: a package for performance assessment and comparison for survival analysis

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4 Functions within *SurvComp*

1 Introduction

The *SurvComp* package is providing functions to assess and to statistically compare the performance of risk prediction (survival) models. It includes (i) implementation of state-of-the-art statistics developed to measure the performance of risk prediction models and (ii) to combine these statistics estimated from multiple datasets using a meta-analytical framework, functions (iii) to visualize those measurements in a clear and compact way, and (iv) to statistically compare the performance of competitive models.

1.1 Installation

SurvComp requires that survival, ipred, prodlim, survivalROC, SuppDists, bootstrap and R (>= 2.3.0) are installed. These should be installed automatically when you install SurvComp. To install SurvComp:

```
> if (!requireNamespace("BiocManager", quietly=TRUE))
+    install.packages("BiocManager")
> BiocManager::install("survcomp")

Load the SurvComp, into your current workspace:
> library(survcomp)
```

1.2 Further help

To view the *SurvComp* description and a summary of all the functions within *Surv-Comp*, type the following:

```
> library(help=survcomp)
```

1.3 Citing

We are delighted if you use this package. Please do email us if you find a bug or have a suggestion. We would be very grateful if you could cite:

B. Haibe-Kains, C. Desmedt, C. Sotiriou and G. Bontempi (2008) A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all? *Bioinformatics* **24(19)**:2200-2208.

2 A use case: from expression data to survival analysis

We will very briefly demonstrate some of the functions in *SurvComp*. We use the breastCancerData datafile for demonstration purposes, it includes subsets of the datasets *breastCancerMAINZ*, *breastCancerTRANSBIG*, *breastCancerUPP*, *breastCancerUNT*, *breastCancerVDX* and *breastCancerNKI*, available as experimental datapackages on Bioconductor. The six datasets in breastCancerData contain the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as uPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008 [1]. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

2.1 Overview

To use the ExpressionSet object we have to load the *Biobase* package. We also make use of the package *xtable* in order to visualize some of the results as tables in this Vignette.

```
> library(Biobase)
> library(xtable)
> library(rmeta)
> library(xtable)
```

Loading the breastCancerData object will results in 6 new objects. If you execute ls() you will see mainz7g,transbig7g, upp7g, unt7g, vdx7g and nki7g. More details about these datasets are available in the breastCancerData manpage (?breastCancerData).

```
> data(breastCancerData)
> mainz7g

ExpressionSet (storageMode: lockedEnvironment)
assayData: 7 features, 200 samples
   element names: exprs
protocolData: none
phenoData
   sampleNames: MAINZ_BC6001 MAINZ_BC6002 ... MAINZ_BC6232 (200 total)
   varLabels: samplename dataset ... e.os (21 total)
   varMetadata: labelDescription
featureData
   featureNames: 205225_at 216836_s_at ... 202763_at (7 total)
   fvarLabels: probe Gene.title ... GO.Component.1 (22 total)
```

-	Gene Symbol	Gene ID	Probes Agilent	Probes Affy
1	esr1	2099	NM_000125	205225_at
2	erbb2	2064	NM_004448	216836_s_at
3	aurka	6790	NM_003600	208079_s_at
4	plau	5328	NM_002658	211668_s_at
5	vegfa	7422	NM_003376	211527_x_at
6	stat1	6772	NM_007315	209969_s_at
7	casp3	836	NM_004346	202763_at

Table 1: Overview of the annotation of the seven genes.

```
fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
  pubMedIds: 18593943
Annotation: hgu133a
```

Before we can start the analysis, we have to define the annotation for the mentioned seven genes, the datasets we use and a few help-variables. We define the gene symbol list (gsList), the entrez-gene ID list (gidList), the probe names for the Agilent microarray (probesNKI), the probe names for the Affymetrix microarray (probesAffy), a list containing the dataset names (datasetList), spaces for displaying the text in the forestplot at the right place (myspace and mybigspace) and to for setting the censored time to 10 years. We converted the gene symbols for each gene to lowercase for better separation from the datasets. Table 1 gives an overview of the gene annotation.

```
> gsList <- tolower(fData(mainz7g)[,"Gene.symbol"])
> gidList <- fData(mainz7g)[,"Gene.ID"]
> probesNKI <- as.character(fData(nki7g)[,"probe"])
> probesAffy <- fData(mainz7g)[,"probe"]
> datasetList <- c("MAINZ","TRANSBIG","UPP","UNT","VDX","NKI","","Overall")
> myspace <- " "
> mybigspace <- " "
> tc <- 10 * 365</pre>
```

2.2 Computing concordance index, D index and hazard ratio

To compute the concordance index [2, 3] for each gene in each dataset, we have to call the <code>concordance.index()</code> function for each dataset. See '?concordance.index' for details. The following command shows the computation of the concordance index for each gene in the <code>mainz7g</code> dataset.

	Min	Max
MAINZ	4.05	14.57
TRANSBIG	4.87	15.18
UPP	4.13	11.22
UNT	-5.04	3.77
VDX	2.77	15.61
NKI	-1.62	0.93

Table 2: Overview of the gene expression ranges in the six datasets.

To compute the D index [4] for each gene in each dataset, we have to call the D.index() function. See '?D.index' for details. The following command shows the computation of the D index for each gene in the mainz7g dataset.

```
> dindexall.mainz.small <- t(apply(X=exprs(mainz7g), MARGIN=1, function(x, y, z) {
+ tt <- D.index(x=x, surv.time=y, surv.event=z, na.rm=TRUE);
+ return(c("dindex"=tt$d.index, "dindex.se"=tt$se, "lower"=tt$lower, "upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=tt$upper"=
```

To compute the hazard ratio [5] for each gene in each dataset, we have to call the hazard.ratio()) function. See ?hazard.ratio for details. Before we compute the hazard ratio, we have to rescale the gene expression data for each dataset to a comparable scale, since the Affymetrix and Agilent microarrays have a different range of their gene expression, which would affect the hazard ratio computation. Table 2 gives an overview of the gene expression ranges in the six datasets that are included in breastCancerData.

Therefore we use the following function to rescale the gene expression values to lie approximately in [-1,1], robust to extreme values (possibly outliers).

```
> rescale <- function(x, na.rm=FALSE, q=0.05) {
+    ma <- quantile(x, probs=1-(q/2), na.rm=na.rm)
+    mi <- quantile(x, probs=q/2, na.rm=na.rm)
+    x <- (x - mi) / (ma - mi)
+    return((x - 0.5) * 2)
+ }</pre>
```

The following command shows the rescaling and the computation of the hazard ratio for each gene in the mainz7g dataset.

```
> hratio.mainz.small <- t(apply(X=rescale(exprs(mainz7g) , q=0.05, na.rm=TRUE), MARGIN=
+ tt <- hazard.ratio(x=x, surv.time=y, surv.event=z, na.rm=TRUE);</pre>
```

```
+ return(c("hratio"=tt$hazard.ratio, "hratio.se"=tt$se, "lower"=tt$lower, "upper"=t
+ y=pData(mainz7g)[ ,"t.dmfs"], z=pData(mainz7g)[ ,"e.dmfs"]))
```

To get an overall estimate over all datasets for the concordance index from each gene, we iterate over all the concordance indices of all datasets and combine them with the combine.est() function [6] and recalculate the lower- and upper border accordingly. We do that for the D indices and the hazard ratios in the same way.

2.3 Combining estimations across datasets

```
> tt <- as.data.frame(NULL)</pre>
> for(i in 1:7){
    tt <- rbind(
      tt, combine.est(
      x=cbind( cindexall.mainz.small[i,"cindex"],
          cindexall.transbig.small[i, "cindex"],
+
          cindexall.upp.small[i, "cindex"],
          cindexall.unt.small[i, "cindex"],
          cindexall.vdx.small[i, "cindex"],
          cindexall.nki.small[i, "cindex"]),
+
      x.se=cbind (cindexall.mainz.small[i, "cindex.se"],
          cindexall.transbig.small[i, "cindex.se"],
          cindexall.upp.small[i, "cindex.se"],
          cindexall.unt.small[i, "cindex.se"],
          cindexall.vdx.small[i, "cindex.se"],
          cindexall.nki.small[i, "cindex.se"]), na.rm=TRUE)
      )
+ }
> tt$lower <- tt$estimate + qnorm(0.025, lower.tail=TRUE) * tt$se
> tt$upper <- tt$estimate + qnorm(0.025, lower.tail=FALSE) * tt$se
> rownames(tt) <- gsList
> colnames(tt) <- c("cindex", "cindex.se", "lower", "upper")</pre>
> ccindex <- tt
```

The combined concordance indices for the six datasets are shown in table 3.

The combined log2 D indices for the six datasets are shown in table 4.

The combined log2 hazard ratios for the six datasets are shown in table 5.

2.4 The forestplot.surv

To display the combined concordance indices of each genes over all datasets, we use the forestplot.surv() function [7]. The resulting forestplot for all concordance indices is:

	cindex	cindex.se	lower	upper
esr1	0.46	0.02	0.43	0.49
erbb2	0.50	0.02	0.47	0.53
aurka	0.64	0.01	0.62	0.67
plau	0.52	0.01	0.49	0.55
vegfa	0.56	0.01	0.53	0.59
stat1	0.53	0.01	0.51	0.56
casp3	0.52	0.01	0.50	0.55

Table 3: Combined concordance indices of each gene for the six datasets.

dindex	dindex.se	lower	upper
-0.17	-3.64	-0.45	0.07
0.09	-3.62	-0.14	0.29
0.96	-3.66	0.84	1.07
0.24	-3.63	0.03	0.42
0.45	-3.65	0.28	0.61
0.19	-3.69	-0.01	0.37
0.19	-3.66	-0.02	0.37
	-0.17 0.09 0.96 0.24 0.45 0.19	-0.17 -3.64 0.09 -3.62 0.96 -3.66 0.24 -3.63 0.45 -3.65 0.19 -3.69	-0.17 -3.64 -0.45 0.09 -3.62 -0.14 0.96 -3.66 0.84 0.24 -3.63 0.03 0.45 -3.65 0.28 0.19 -3.69 -0.01

Table 4: Combined log2 D indices of each gene for the six datasets.

	hratio	hratio.se	lower	upper
esr1	-0.27	-3.63	-0.57	-0.01
erbb2	0.38	-3.11	0.10	0.61
aurka	2.07	-2.59	1.95	2.18
plau	0.84	-2.34	0.50	1.13
vegfa	0.93	-2.79	0.69	1.13
stat1	0.48	-2.68	0.12	0.76
casp3	3.24	-1.23	3.11	3.36

Table 5: Combined log2 hazard ratios of each gene for the six datasets.

```
> labeltext <- cbind(c("Gene Symbol",gsList),c(rep(myspace,8)))
```

> bs <- rep(0.5, nrow(labeltext))</pre>

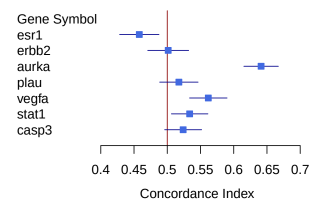
> r.mean <- c(NA,ccindex\$cindex)</pre>

> r.lower <- c(NA,ccindex\$lower)</pre>

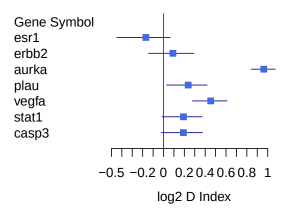
> r.upper <- c(NA,ccindex\$upper)</pre>

> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero= + align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(0.4,0.7,0.05), xlab

+ col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, c

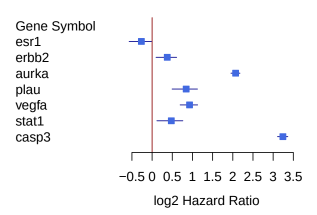


The resulting forestplot for all D indices is:



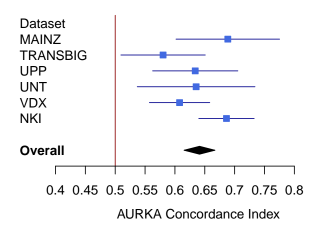
The resulting forestplot for all hazard ratios is:

```
> labeltext <- cbind(c("Gene Symbol",gsList),c(rep(mybigspace,8)))
> bs <- rep(0.5, nrow(labeltext))
> r.mean <- c(NA,log2(chratio$hratio))
> r.lower <- c(NA,log2(chratio$lower))
> r.upper <- c(NA,log2(chratio$upper))
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero= align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(-0.5,3.5,0.5), xlatecol=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, col=meta.colors(box="royalblue", line="darkblue", zero="darkblue", zero="darkblue")
```

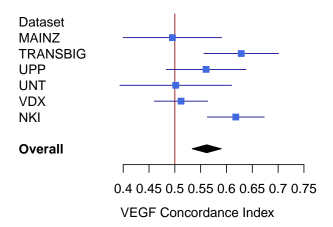


Taking a more specific look, e.g. at the genes AURKA and VEGF, we create the forestplot the same way as before, showing the concordance indices for both genes in each dataset and the combined estimation over all datasets.

```
> r.upper <- c(NA,tt$upper)
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=
+ align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(0.4,0.8,0.05), xla
+ col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, colors("l")
```

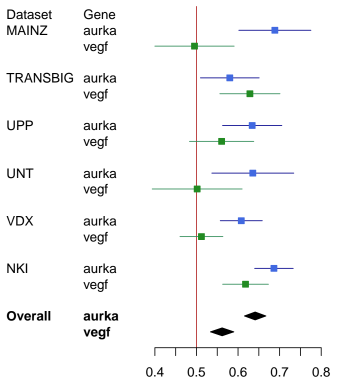


```
> r.mean <- c(NA,tt$cindex)
> r.lower <- c(NA,tt$lower)
> r.upper <- c(NA,tt$upper)
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=
+ align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(0.4,0.75,0.05), x2
+ col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, col=meta.colors(box="royalblue", line="darkblue", zero="darkred")
```



More advanced displaying of the genes AURKA and VEGF in a single forestplot with different colors and labels is possible:

```
NA,
+
              cindexall.unt.small[3,],
              cindexall.unt.small[5,],
+
              NA,
              cindexall.vdx.small[3,],
              cindexall.vdx.small[5,],
              NA,
              cindexall.nki.small[3,],
              cindexall.nki.small[5,],
              NA,
              as.numeric(ccindex[3,]),
              as.numeric(ccindex[5,]))
> rownames(tt) <- c("MAINZa", "MAINZv", "a", "TRANSBIGa", "TRANSBIGv", "b", "UPPa", "UP
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset", "MAINZ", NA, NA, "TRANSBIG", NA, NA, "UPP", NA, NA, '
                      c("Gene", rep(c("aurka", "vegf", NA), length(datasetList)-2), c("aur
> bs <- rep(0.5, nrow(labeltext))</pre>
> r.mean <- c(NA,tt$cindex)</pre>
> r.lower <- c(NA,tt$lower)</pre>
> r.upper <- c(NA,tt$upper)</pre>
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=
      align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(0.4,0.8,0.05), xlaign=c("l")
      col=meta.colors(line=c(rep(c(NA, "darkblue", "seagreen"),7)), zero="firebrick", k
      clip=c(0.3,1), is.summary=(c(rep(FALSE,19), TRUE, TRUE)))
```

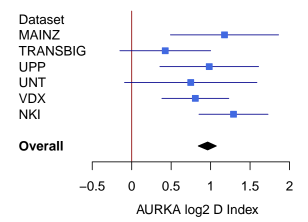


AURKA and VEGF Concordance Index

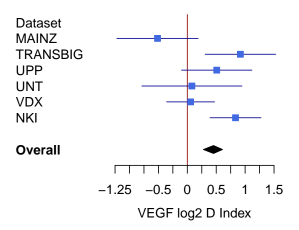
We display the D indices for both genes in each dataset and the combined estimation over all datasets in the same way.

```
> tt <- rbind(dindexall.mainz.small[3,],</pre>
               dindexall.transbig.small[3,],
+
               dindexall.upp.small[3,],
               dindexall.unt.small[3,],
               dindexall.vdx.small[3,],
               dindexall.nki.small[3,],
               NA,
               as.numeric(cdindex[3,]))
> rownames(tt) <- datasetList
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset",datasetList),c(rep(mybigspace,length(datasetList)+1)))
> bs <- rep(0.5, nrow(labeltext))</pre>
> r.mean <- c(NA,log2(tt$dindex))</pre>
> r.lower <- c(NA,log2(tt$lower))</pre>
> r.upper <- c(NA,log2(tt$upper))</pre>
```

```
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=
+ align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(-0.5,2,0.5), xlab=
+ col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, col=meta.colors(box="royalblue", line="darkblue", zero="darkblue")
```

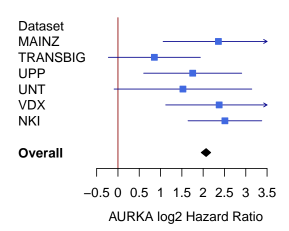


```
> r.lower <- c(NA,log2(tt$lower))
> r.upper <- c(NA,log2(tt$upper))
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=
+ align=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(-1.25,1.5,0.25), x2
+ col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, colemeta.colors(box="royalblue")
```



And at last the hazard ratio for the gene AURKA in each dataset and the combined estimation over all datasets.

```
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset",datasetList),c(rep(myspace,length(datasetList)+1)))
> bs <- rep(0.5, nrow(labeltext))
> r.mean <- c(NA,log2(tt$hratio))
> r.lower <- c(NA,log2(tt$lower))
> r.upper <- c(NA,log2(tt$upper))
> forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=talign=c("l"), graphwidth=grid::unit(2, "inches"), x.ticks=seq(-0.5,3.5,0.5), xlabeltext=col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, col=meta.colors(box="royalblue", line="darkblue", zero="darkblue")
```



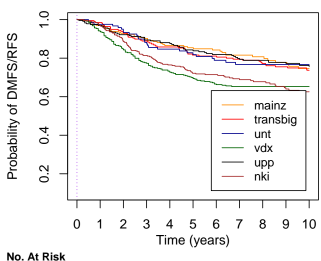
The following small loop shows an easy way for creating several forestplots showing the concordance indices for a single gene for all datasets and the combined estimation over all datasets. The same can be done for the D indices and hazard ratios. Since it is not yet possible to combine several forestplots in one figure (e.g. with par(mfrow=c(2,2))), we don't display the results of the following loop.

```
> for(i in 1:length(gsList)) {
+ tt <- rbind(cindexall.mainz.small[i,],</pre>
```

```
cindexall.transbig.small[i,],
+
                cindexall.upp.small[i,],
                cindexall.unt.small[i,],
+
                cindexall.vdx.small[i,],
                cindexall.nki.small[i,],
                NA,
                as.numeric(ccindex[i,]))
+
    rownames(tt) <- datasetList
+
    tt <- as.data.frame(tt)</pre>
    labeltext <- cbind(c("Dataset",datasetList), c(rep(myspace,length(datasetList)+1)))
    bs <- rep(0.5, nrow(labeltext))</pre>
+
    r.mean <- c(NA,tt$cindex)</pre>
    r.lower <- c(NA,tt$lower)
+
    r.upper <- c(NA,tt$upper)
+
    x.ticks.lower \leftarrow (floor((min(r.mean, na.rm=TRUE) - 0.1) * 10)/10)
+
    x.ticks.upper <- (floor((max(r.mean, na.rm=TRUE) + 0.2) * 10)/10)
+
    forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zer
+
        align=c("1"), graphwidth= grid::unit(2, "inches"), x.ticks=seq(x.ticks.lower,x.
        col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs,
+ }
```

2.5 Kaplan Meier survival curves

To display a Kaplan Meier curve [8] for all datasets you can use:

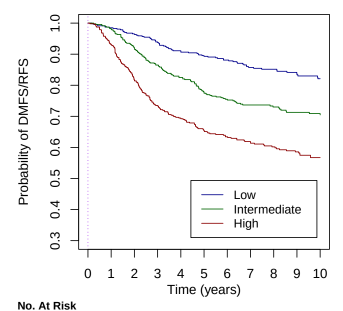


main 200 191 179 171 159 153 130 107 86 76 60 transbig 98 195 185 175 166 154 151 146 136 132 123 un 125 119 112 101 93 86 76 68 61 48 38 vd 844 324 292 267 252 231 207 184 136 89 53 up 234 223 208 195 188 177 167 159 152 146 134 nk 319 305 278 249 232 213 179 148 118 94 77

If you want do display the survival curve for a single gene using the data of all six datasets, we have to concatenate the survival and expression data of all datasets (see surv.time.all, surv.event.all and aurka.exprs below). After that we split the patients in each dataset into three parts acording to their gene expression. We use the function quantile() for that. In the end we have three groups, representing the low gene expression group (lowest 33% of the gene expression), intermediate gene expression group (gene expression between 33% and 66%) and high gene expression group (over 66%).

```
> aurkaGs <- "AURKA"
> aurkaGid <- 6790
> aurkaPaf <- "208079_s_at"
> aurkaPagi <- "NM_003600"
> surv.time.all <- c(pData(mainz7g)[ ,"t.dmfs"], pData(transbig7g)[ ,"t.dmfs"], pData(v)
> surv.event.all <- c(pData(mainz7g)[ ,"e.dmfs"], pData(transbig7g)[ ,"e.dmfs"], pData(v)
> aurka.exprs <- c(exprs(mainz7g)[aurkaPaf,], exprs(transbig7g)[aurkaPaf,], exprs(unt7g)
> aurka.exprs.length <- c(length(exprs(mainz7g)[aurkaPaf,]), length(exprs(transbig7g))</pre>
```

```
> mygroup <- NULL
> for(i in aurka.exprs.length){
    qq <- aurka.exprs[(pos+1):(pos+i)]
    myq <- quantile(qq, probs=c(0.33, 0.66), na.rm=TRUE)</pre>
+
    qq[aurka.exprs[(pos+1):(pos+i)] < myq[1]] <- 1
    qq[aurka.exprs[(pos+1):(pos+i)] >= myq[1] & aurka.exprs[(pos+1):(pos+i)] < myq[2]]
    qq[aurka.exprs[(pos+1):(pos+i)] > myq[2]] <- 3
    qq <- factor(x=qq, levels=1:3)</pre>
    mygroup <- c(mygroup,qq)</pre>
+
   pos <- pos + i
> surv.data <- censor.time(surv.time=surv.time.all / 365, surv.event=surv.event.all, ta
> dd <- data.frame("time"=surv.data[[1]], "event"=surv.data[[2]], "gg"=mygroup)
> gg <- factor(c(rep("mainz", nrow(pData(mainz7g))), rep("transbig", nrow(pData(transbig))))
> km.coxph.plot(formula.s=formula(Surv(time, event) ~ gg), data.s=dd, sub.s="all", x.la
```



Low 467 459 446 431 411 394 351 314 268 220 178 Intermediate 468 455 424 391 367 333 297 263 222 193 166 High 478 437 378 330 307 282 257 230 194 167 137

2.6 Meta analysis of estimation values

The *SurvComp* package integrates functions for meta-analysis of risk-prediction models, e.g. for the concordance index or the D index. The following example shows the cindex.comp.meta() function [9]. Table 6 shows the p-values representing the difference between the cindices of two genes using the cindices of all six datasets. For example, the cindex of the gene AURKA is with a p-value of 0.00001 significantly different from the cindex of the gene VEGF using the six datasets.

```
> cindexMetaMainz <- t(apply(X=exprs(mainz7g), MARGIN=1, function(x, y, z) {
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
      return(tt); }, y=pData(mainz7g)[ ,"t.dmfs"], z=pData(mainz7g)[ ,"e.dmfs"]))
> cindexMetaTransbig <- t(apply(X=exprs(transbig7g), MARGIN=1, function(x, y, z) {
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
      return(tt); }, y=pData(transbig7g)[ ,"t.dmfs"], z=pData(transbig7g)[ ,"e.dmfs"]);
> cindexMetaUpp <- t(apply(X=exprs(upp7g), MARGIN=1, function(x, y, z)) 
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
      return(tt); }, y=pData(upp7g)[ ,"t.rfs"], z=pData(upp7g)[ ,"e.rfs"]))
> cindexMetaUnt <- t(apply(X=exprs(unt7g), MARGIN=1, function(x, y, z)) {
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
      return(tt); }, y=pData(unt7g)[ ,"t.dmfs"], z=pData(unt7g)[ ,"e.dmfs"]))
> cindexMetaVdx <- t(apply(X=exprs(vdx7g), MARGIN=1, function(x, y, z)) 
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
      return(tt); }, y=pData(vdx7g)[ ,"t.dmfs"], z=pData(vdx7g)[ ,"e.dmfs"]))
> ccNki <- complete.cases(exprs(nki7g)[1,], exprs(nki7g)[2,], exprs(nki7g)[3,], exprs(1
> cindexMetaNki <- t(apply(X=exprs(nki7g)[,ccNki], MARGIN=1, function(x, y, z) {
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=1
     return(tt); }, y=pData(nki7g)[ccNki ,"t.dmfs"], z=pData(nki7g)[ccNki ,"e.dmfs"]);
> ccmData <- tt <- rr <- NULL
> for(i in 1:7){
   tt <- NULL
   listOne <- list("mainz" = cindexMetaMainz[[i]],</pre>
                     "transbig" = cindexMetaTransbig[[i]],
+
                     "upp" = cindexMetaUpp[[i]],
                     "unt" = cindexMetaUnt[[i]],
                     "vdx" = cindexMetaVdx[[i]],
                     "nki" = cindexMetaNki[[i]])
+
+
   for(j in 1:7){
      listTwo <- list("mainz" = cindexMetaMainz[[j]],
                      "transbig" = cindexMetaTransbig[[j]],
+
+
                      "upp" = cindexMetaUpp[[j]],
                      "unt" = cindexMetaUnt[[i]],
```

	esr1	erbb2	aurka	plau	vegfa	stat1	casp3
esr1	1.00000	0.96698	1.00000	0.99489	1.00000	0.99967	0.99860
erbb2	0.03302	1.00000	1.00000	0.79944	0.99855	0.94848	0.87718
aurka	0.00000	0.00000	1.00000	0.00000	0.00001	0.00000	0.00000
plau	0.00511	0.20056	1.00000	1.00000	0.98988	0.80287	0.62407
vegfa	0.00000	0.00145	0.99999	0.01012	1.00000	0.07944	0.02743
stat1	0.00033	0.05152	1.00000	0.19713	0.92056	1.00000	0.28393
casp3	0.00140	0.12282	1.00000	0.37593	0.97257	0.71607	1.00000

Table 6: cindex.comp.meta() results showing the significance of the difference between concordance indices.

3 Session Info

- R version 4.3.1 (2023-06-16), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_GB, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Time zone: America/New_York
- TZcode source: system (glibc)
- Running under: Ubuntu 22.04.3 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
- LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: Biobase 2.62.0, BiocGenerics 0.48.0, prodlim 2023.08.28, rmeta 3.0, survcomp 1.52.0, survival 3.5-7, xtable 1.8-4
- Loaded via a namespace (and not attached): KernSmooth 2.23-22, Matrix 1.6-1.1, Rcpp 1.0.11, SuppDists 1.1-9.7, bootstrap 2019.6, codetools 0.2-19, compiler 4.3.1, data.table 1.14.8, digest 0.6.33, future 1.33.0, future.apply 1.11.0, globals 0.16.2, grid 4.3.1, lattice 0.22-5, lava 1.7.2.1, listenv 0.9.0, parallel 4.3.1, parallelly 1.36.0, splines 4.3.1, survivalROC 1.0.3.1, tools 4.3.1

4 Functions within SurvComp

For references to the following functions, please see [2]-[21].

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FUNCTION DESCRIPTION

D.index Function to compute the D index

breastCancerDataSample data containing six datasets for gene expression, annotations

and clinical data

censor.time Function to artificially censor survival data cindex.comp Function to compare two concordance indices cindex.comp.metaFunction to compare two concordance indices

combine.est Function to combine estimates combine.test Function to combine probabilities

concordance.indexFunction to compute the cindex for survival or binary class prediction

cvpl Function to compute the CVPL dindex.comp Function to compare two D indices dindex.comp.metaFunction to compare two D indices

fisherz Function to compute Fisher z transformation

forestplot.surv Forest plots enables to display performance estimates of survival mod-

els

getsurv2 Function to retrieve the survival probabilities at a specific point in time

hazard.ratio Function to estimate the hazard ratio through Cox regression

hr.comp Function to statistically compare two hazard ratios hr.comp.meta Function to compare two concordance indices

hr.comp2 Function to statistically compare two hazard ratios (alternative inter-

face)

iauc.comp Function to compare two IAUCs through time-dependent ROC curves

ibsc.comp Function to compare two IBSCs

km.coxph.plot Function to plot several Kaplan-Meier survival curves

logpl Function to compute the log partial likelihood of a Cox model

metaplot.surv Meta plots enables to display performance estimates of survival mod-

els

no.at.risk Function to compute the number of individuals at risk

sbrier.score2probaFunction to compute the BSCs from a risk score, for all the times of

event occurrence

score2proba Function to compute the survival probabilities from a risk score survcomp
Performance Assessment and Comparison for Survival Analysis

package

td.sens.spec Function to compute sensitivity and specificity for a binary classifica-

tion of survival data

tdrocc Function to compute time-dependent ROC curves test.hetero.est Function to test the heterogeneity of set of probabilities test.hetero.test Function to test the heterogeneity of set of probabilities