genefu: a package for breast cancer gene expression analysis

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October 13, 2014

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1 Introduction

The *genefu* package is providing relevant functions for gene expression analysis, especially in breast cancer.

2 Case Study 1: comparing risk prediction models

For computing the risk scores, estimates of the performance of the risk scores, combining the estimates and comparing the estimates we have to load the *genefu* and *survcomp* packages into the workspace. We also load the *xtable* package to display results insight this document.

- > library(genefu)
- > library(xtable)
- > library(rmeta)

The five data sets that we use in the case study are publicly available as experimental data packages on Bioconductor.org. In particular we used:

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- breastCancerMAINZ: http://www.bioconductor.org/packages/release/data/experiment/
 html/breastCancerMAINZ.html
- breastCancerUPP: http://www.bioconductor.org/packages/release/data/experiment/
 html/breastCancerUPP.html
- breastCancerUNT: http://www.bioconductor.org/packages/release/data/experiment/
 html/breastCancerUNT.html
- breastCancerNKI: http://www.bioconductor.org/packages/release/data/experiment/
 html/breastCancerNKI.html
- breastCancerTRANSBIG: http://www.bioconductor.org/packages/release/data/experiment/ html/breastCancerTRANSBIG.html

We don't use the breastCancerVDX experimental package in the case study since it has been used as training data set for GENIUS [?]:

breastCancerVDX: http://www.bioconductor.org/packages/release/data/experiment/ html/breastCancerVDX.html

These experimental data packages can be installed from Bioconductor version 2.8 or higher in R version 2.13.0 or higher. For the experimental data packages the commands for installing the data sets are:

- > source("http://www.bioconductor.org/biocLite.R")
- > biocLite("breastCancerMAINZ")
- > biocLite("breastCancerTRANSBIG")
- > biocLite("breastCancerUPP")
- > biocLite("breastCancerUNT")
- > biocLite("breastCancerNKI")

And for loading the data sets into the current workspace the commands are:

- > library(breastCancerMAINZ)
- > library(breastCancerTRANSBIG)
- > library(breastCancerUPP)
- > library(breastCancerUNT)
- > library(breastCancerNKI)

Table 1: Detailed overview for the data sets used in the case study

Dataset	Patients [#]	ER+[#]	HER2+ [#]	Age [years]	Grade $[1/2/3]$	Platform
MAINZ	200	155	23	25-90	29/136/35	HGU133A
TRANSBIG	198	123	35	24-60	30/83/83	HGU133A
UPP	251	175	46	28-93	67/128/54	HGU133AB
UNT	137	94	21	24-73	32/51/29	HGU133AB
NKI	337	212	53	26-62	79/109/149	Agilent
Overall	1123	759	178	25-73	237/507/350	Affy/Agilent

Table 1 shows an overview of the data sets and the patients. From those 1123 breast cancer patients we selected only the patients that are node negative and didn't receive any treatment (except local radiotherapy), which results in 722 patients.

Since there are duplicated patients in the five data sets, we have to identify the duplicated patients and we subsequently store them in a vector. We then compute the risk scores for NPI, AURKA, GGI and GENIUS with the npi(), sig.score(), ggi() and genius() functions within genefu, respectively.

```
> dn <- c("transbig", "unt", "upp", "mainz", "nki")</pre>
> dn.platform <- c("affy", "affy", "affy", "affy", "agilent")
> res <- ddemo.all <- ddemo.coln <- NULL
> for(i in 1:length(dn)) {
    ## load dataset
    dd <- get(data(list=dn[i]))</pre>
    ddata <- t(exprs(dd))</pre>
    ddemo <- phenoData(dd)@data
    dannot <- featureData(dd)@data
    ddemo.all <- c(ddemo.all, list(ddemo))</pre>
    if(is.null(ddemo.coln)) { ddemo.coln <- colnames(ddemo) } else { ddemo.coln <- interse
    rest <- NULL
    ## NPI
    ss <- ddemo[ , "size"]</pre>
    gg <- ddemo[ , "grade"]</pre>
    nn <- rep(NA, nrow(ddemo))</pre>
    nn[complete.cases(ddemo[ , "node"]) & ddemo[ , "node"] == 0] <- 1</pre>
    nn[complete.cases(ddemo[ , "node"]) & ddemo[ , "node"] == 1] <- 3
    names(ss) <- names(gg) <- names(nn) <- rownames(ddemo)</pre>
    rest <- cbind(rest, "NPI"=npi(size=ss, grade=gg, node=nn, na.rm=TRUE)$score)
    ## AURKA
    ## if affy platform consider the probe published in Desmedt et al., CCR, 2008
```

```
if(dn.platform[i] == "affy") { domap <- FALSE } else { domap <- TRUE }</pre>
   modt <- scmgene.robust$mod$AURKA
   ## if agilent platform consider the probe published in Desmedt et al., CCR, 2008
   if(dn.platform[i] == "agilent") {
      domap <- FALSE
      modt[ , "probe"] <- "NM_003600"
   }
   rest <- cbind(rest, "AURKA"=sig.score(x=modt, data=ddata, annot=dannot, do.mapping=dom
   if(dn.platform[i] == "affy") { domap <- FALSE } else { domap <- TRUE }</pre>
   rest <- cbind(rest, "GGI"=ggi(data=ddata, annot=dannot, do.mapping=domap)$score)
   if(dn.platform[i] == "affy") { domap <- FALSE } else { domap <- TRUE }</pre>
   rest <- cbind(rest, "GENIUS"=genius(data=ddata, annot=dannot, do.mapping=domap)$score)
   res <- rbind(res, rest)</pre>
+ }
> names(ddemo.all) <- dn
```

For further analysis and handling of the data we store all information in one object. We also remove the duplicated patients from the analysis and take only those patients into account, that have complete information for nodal, survival and treatment status.

```
> ddemot <- NULL
> for(i in 1:length(ddemo.all)) {
    ddemot <- rbind(ddemot, ddemo.all[[i]][ , ddemo.coln, drop=FALSE])</pre>
> res[complete.cases(ddemot[ ,"dataset"]) & ddemot[ ,"dataset"] == "VDX", "GENIUS"] <- NA
> ## select only untreated node-negative patients with all risk predictions
> myx <- complete.cases(res, ddemot[ , c("node", "treatment")]) & ddemot[ , "treatment"] =
> res <- res[myx, , drop=FALSE]</pre>
> ddemot <- ddemot[myx, , drop=FALSE]</pre>
   To compare the risk score performances, we compute the concordance index<sup>1</sup>:
```

```
> cc.res <- complete.cases(res)</pre>
> datasetList <- c("MAINZ", "TRANSBIG", "UPP", "UNT", "NKI")</pre>
> riskPList <- c("NPI", "AURKA", "GGI", "GENIUS")</pre>
> setT <- setE <- NULL
> resMatrix <- as.list(NULL)</pre>
> for(i in datasetList){
    dataset.only <- ddemot[,"dataset"] == i</pre>
    patientsAll <- cc.res & dataset.only
```

 $^{^{1}}$ The same analysis could be performed with D index and hazard ratio by using the functions D.index and hazard.ratio from the survcomp respectively

```
## set type of available survival data
   if(i == "UPP") {
     setT <- "t.rfs"
     setE <- "e.rfs"
   } else {
      setT <- "t.dmfs"</pre>
      setE <- "e.dmfs"
   }
+
+
   ## cindex computation
   cindexN <- t(apply(X=t(res[patientsAll,"NPI"]), MARGIN=1, function(x, y, z) {</pre>
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=TRUE
      return(c("cindex"=tt$c.index, "cindex.se"=tt$se, "lower"=tt$lower, "upper"=tt$upper)
      y=ddemot[patientsAll,setT], z=ddemot[patientsAll, setE]))
   cindexA <- t(apply(X=t(res[patientsAll, "AURKA"]), MARGIN=1, function(x, y, z) {</pre>
+
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=TRUE
      return(c("cindex"=tt$c.index, "cindex.se"=tt$se, "lower"=tt$lower, "upper"=tt$upper)
      y=ddemot[patientsAll,setT], z=ddemot[patientsAll, setE]))
   cindexM <- t(apply(X=t(res[patientsAll, "GGI"]), MARGIN=1, function(x, y, z) {</pre>
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=TRUE
      return(c("cindex"=tt$c.index, "cindex.se"=tt$se, "lower"=tt$lower, "upper"=tt$upper)
      y=ddemot[patientsAll, setT], z=ddemot[patientsAll, setE]))
   cindexG <- t(apply(X=t(res[patientsAll, "GENIUS"]), MARGIN=1, function(x, y, z) {</pre>
      tt <- concordance.index(x=x, surv.time=y, surv.event=z, method="noether", na.rm=TRUE
      return(c("cindex"=tt$c.index, "cindex.se"=tt$se, "lower"=tt$lower, "upper"=tt$upper)
      y=ddemot[patientsAll, setT], z=ddemot[patientsAll, setE]))
   resMatrix[["NPI"]] <- rbind(resMatrix[["NPI"]], cindexN)</pre>
   resMatrix[["AURKA"]] <- rbind(resMatrix[["AURKA"]], cindexA)</pre>
   resMatrix[["GGI"]] <- rbind(resMatrix[["GGI"]], cindexM)</pre>
   resMatrix[["GENIUS"]] <- rbind(resMatrix[["GENIUS"]], cindexG)</pre>
+ }
```

Using a random-effects model we combine the dataset-specific performance estimated into overall estimates for each risk prediction model:

```
> for(i in names(resMatrix)){
+ ceData <- combine.est(x=resMatrix[[i]][,"cindex"], x.se=resMatrix[[i]][,"cindex.se"], if
+ cLower <- ceData$estimate + qnorm(0.025, lower.tail=TRUE) * ceData$se
+ cUpper <- ceData$estimate + qnorm(0.025, lower.tail=FALSE) * ceData$se
+ cindex0 <- cbind("cindex"=ceData$estimate, "cindex.se"=ceData$se, "lower"=cLower, "upper cindex.se"=ceData$se, "upper c
```

```
+ resMatrix[[i]] <- rbind(resMatrix[[i]], cindex0)
+ rownames(resMatrix[[i]]) <- c(datasetList, "Overall")
+ }</pre>
```

In order to compare the different risk prediction models we compute one-sided p-values of the meta-estimates:

```
> pv <- sapply(resMatrix, function(x) { return(x["Overall", c("cindex", "cindex.se")]) })
> pv <- apply(pv, 2, function(x) { return(pnorm((x[1] - 0.5) / x[2], lower.tail=x[1] < 0.5
> printPV <- matrix(pv,ncol=4)
> rownames(printPV) <- "P-value"
> colnames(printPV) <- names(pv)

> xtable(printPV, digits=c(0, rep(-1,ncol(printPV))))
```

	NPI	AURKA	GGI	GENIUS
P-value	5.5E-16	1.6E-08	2.2E-15	2.9E-24

The following five figures represent the risk score performances measured by the concordance index for NPI, AURKA, GGI and GENIUS. The last figure represents the overall estimates.

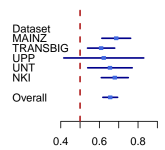
```
> par(mfrow=c(2,2))
> datasetListF <- c("MAINZ", "TRANSBIG", "UPP", "UNT", "NKI", NA, "Overall")
> myspace <- " "
> ## NPI Forestplot
> tt <- rbind(resMatrix[["NPI"]][1:5,],</pre>
             "NA"=NA,
             "Overall"=resMatrix[["NPI"]][6,])
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset", datasetListF))</pre>
> r.mean <- c(NA,tt$cindex)</pre>
> r.lower <- c(NA,tt$lower)
> r.upper <- c(NA,tt$upper)</pre>
> metaplot.surv(mn=r.mean, lower=r.lower, upper=r.upper, labels=labeltext, xlim=c(0.4,0.9)
> #@
> #<<forestplotAURKA,fig=TRUE,echo=FALSE>>=
> ## AURKA Forestplot
> tt <- rbind(resMatrix[["AURKA"]][1:5,],
             "NA"=NA,
             "Overall"=resMatrix[["AURKA"]][6,])
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset", datasetListF))</pre>
> r.mean <- c(NA,tt$cindex)</pre>
```

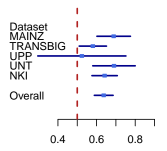
> r.lower <- c(NA,tt\$lower)</pre>

```
> r.upper <- c(NA,tt$upper)</pre>
> metaplot.surv(mn=r.mean, lower=r.lower, upper=r.upper, labels=labeltext, xlim=c(0.4,0.9)
> #@
> #
> #<<forestplotGGI,fig=TRUE,echo=FALSE>>=
> ## GGI Forestplot
> tt <- rbind(resMatrix[["GGI"]][1:5,],
             "NA"=NA,
             "Overall"=resMatrix[["GGI"]][6,])
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset", datasetListF))</pre>
> r.mean <- c(NA,tt$cindex)
> r.lower <- c(NA,tt$lower)
> r.upper <- c(NA,tt$upper)</pre>
> metaplot.surv(mn=r.mean, lower=r.lower, upper=r.upper, labels=labeltext, xlim=c(0.4,0.9)
> #
> #<<forestplotGENIUS,fig=TRUE,echo=FALSE>>=
> ## GENIUS Forestplot
> tt <- rbind(resMatrix[["GENIUS"]][1:5,],</pre>
             "NA"=NA,
             "Overall"=resMatrix[["GENIUS"]][6,])
> tt <- as.data.frame(tt)
> labeltext <- cbind(c("Dataset", datasetListF))</pre>
> r.mean <- c(NA,tt$cindex)
> r.lower <- c(NA,tt$lower)</pre>
> r.upper <- c(NA,tt$upper)</pre>
> metaplot.surv(mn=r.mean, lower=r.lower, upper=r.upper, labels=labeltext, xlim=c(0.4,0.9)
```

NPI Concordance Index

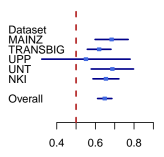
AURKA Concordance Index

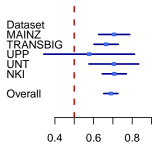




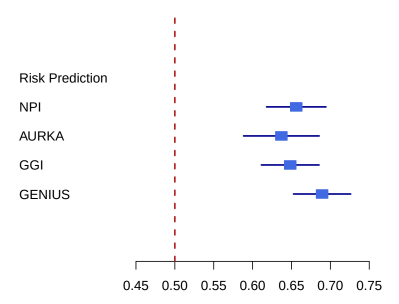
GGI Concordance Index

GENIUS Concordance Index





Overall Concordance Index



In order to assess the difference between the risk scores, we compute the concordance indices with their p-values and compare the estimates with the cindex.comp.meta with a paired student t test.

Table 2 displays the for multiple testing uncorrected p-values for the comparison of the different methods:

> xtable(ccmData, digits=c(0, rep(-1,ncol(ccmData))))

	NPI	AURKA	GGI	GENIUS
NPI	1.0E+00	2.5E-01	3.7E-01	9.1E-01
AURKA	7.5E-01	1.0E+00	7.0E-01	9.8E-01
GGI	6.3E-01	3.0E-01	1.0E + 00	9.7E-01
GENIUS	9.0E-02	2.4E-02	3.1E-02	1.0E+00

We correct the p-value with Holms method:

- > ccmDataPval <- matrix(p.adjust(data.matrix(ccmData), method="holm"),ncol=4,dimnames=list
 Table 3 displays the corrected p-values:
- > xtable(ccmDataPval, digits=c(0, rep(-1,ncol(ccmDataPval))))

	NPI	AURKA	GGI	GENIUS
NPI	1.0E+00	1.0E+00	1.0E+00	1.0E+00
AURKA	1.0E+00	1.0E+00	1.0E+00	1.0E+00
GGI	1.0E + 00	1.0E + 00	1.0E + 00	1.0E+00
GENIUS	1.0E + 00	3.8E-01	4.7E-01	1.0E+00

3 Session Info

- R Under development (unstable) (2014-10-07 r66723), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, 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
- Base packages: base, datasets, grDevices, graphics, grid, methods, parallel, splines, stats, utils
- Other packages: Biobase 2.27.0, BiocGenerics 0.13.0, biomaRt 2.23.0, breastCancerMAINZ 1.3.0, breastCancerNKI 1.3.0, breastCancerTRANSBIG 1.3.0, breastCancerUNT 1.3.0, breastCancerUPP 1.3.0, genefu 1.17.0, mclust 4.4, prodlim 1.4.5, rmeta 2.16, survcomp 1.17.0, survival 2.37-7, xtable 1.7-4
- Loaded via a namespace (and not attached): AnnotationDbi 1.29.0, DBI 0.3.1, GenomeInfoDb 1.3.0, IRanges 2.1.0, KernSmooth 2.23-13, RCurl 1.95-4.3, RSQLite 0.11.4, S4Vectors 0.5.0, SuppDists 1.1-9.1, XML 3.98-1.1, amap 0.8-12, bootstrap 2014.4, lava 1.2.6, stats4 3.2.0, survivalROC 1.0.3, tools 3.2.0