Xeva Tutorial

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Load Xeva

Load Xeva library and data.

```
library(Xeva)
data(lpdx)
head(modelInfo(lpdx))
```

```
##
                                   model.id
                                                                  donor
## PHLC1106 P5.501.A1.1 PHLC1106 P5.501.A1.1 11101S-213RC-312S(F)-412S-
## PHLC1106_P5.504.A4.1 PHLC1106_P5.504.A4.1 11101S-213RC-312S(F)-412S-
## PHLC1106_P5.506.B1.1 PHLC1106_P5.506.B1.1 11101S-213RC-312S(F)-412S-
## PHLC1106_P5.507.B2.1 PHLC1106_P5.507.B2.1 11101S-213RC-312S(F)-412S-
## PHLC1106_P5.508.B3.1 PHLC1106_P5.508.B3.1 11101S-213RC-312S(F)-412S-
## PHLC1106_P5.511.C1.1 PHLC1106_P5.511.C1.1 11101S-213RC-312S(F)-412S-
                            dob sex
                                        PHLC biobase.id patient.id
## PHLC1106_P5.501.A1.1 Aug31.14 F PHLC1106
                                               PHLC1106
                                                          PHLC1106
## PHLC1106_P5.504.A4.1 Aug31.14 F PHLC1106
                                               PHLC1106
                                                          PHLC1106
## PHLC1106_P5.506.B1.1 Aug31.14 F PHLC1106
                                               PHLC1106
                                                          PHLC1106
## PHLC1106_P5.507.B2.1 Aug31.14 F PHLC1106
                                               PHLC1106
                                                          PHLC1106
## PHLC1106_P5.508.B3.1 Aug31.14 F PHLC1106
                                                          PHLC1106
                                               PHLC1106
## PHLC1106 P5.511.C1.1 Sep14.14 F PHLC1106
                                               PHLC1106
                                                          PHLC1106
```

Models which belongs to same batch are in one list which is stored in expDesign slot. For example

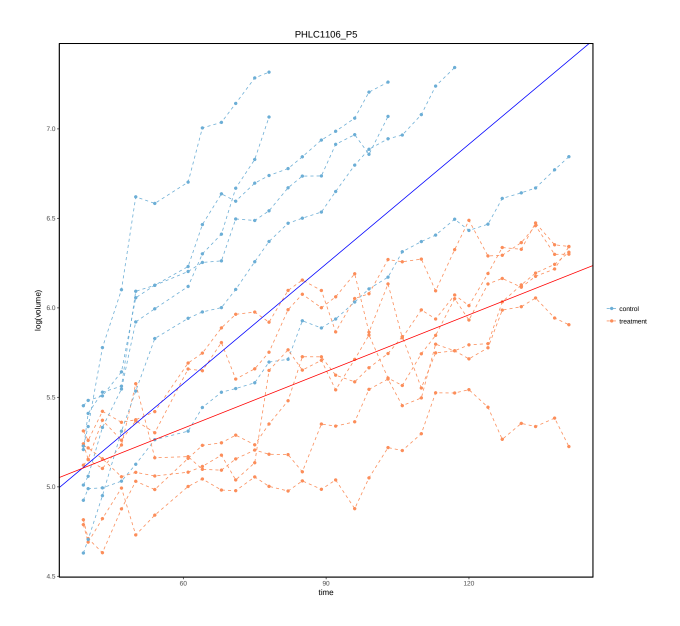
```
print(batchNames(lpdx))
```

```
## [1] "PHLC1106_P5" "PHLC111_P7" "PHLC119_P5" "PHLC153_P6" "PHLC181_P7" 
## [6] "PHLC189_P5" "PHLC191_P5" "PHLC191_P7" "PHLC196_P5" "PHLC215_P5" 
## [11] "PHLC229_P6" "PHLC235_P4" "PHLC655_P7" "PHLC82_P5"
```

To calculate angle between the treatment and control samples of this batch

```
batchNames <- batchNames(lpdx)
expDesign <- expDesign(lpdx, batchNames[1])
ang <- calculateAngle(lpdx, expDesign, treatment.only = TRUE, plot=TRUE)
print(ang)</pre>
```

```
## $PHLC1106_P5
## $PHLC1106_P5$angle
## [1] 0.671583
##
## $PHLC1106_P5$plot
```



```
#par(mfrow=c(5,3))
for(I in batchNames)
{
   expDesign <- expDesign(lpdx, I)
   ang <- calculateAngle(lpdx, expDesign, treatment.only = TRUE, plot=TRUE)
# print(ang)
}</pre>
```

Summarize Response of PDXs Get slop of each model and combine summarize all model slop which belongs to same patient by "mean" $^{\prime\prime}$

Get angle between treatment and control model ids. For each batch it will give one angle value

```
lpdx_angle <- summarizeResponse(lpdx, response.measure = "angle")</pre>
```

Get mutation expression profile

```
ldxe_mut <- getMolecularProfiles(lpdx, data.type="mutation")
print(ldxe_mut)</pre>
```

```
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 16116 features, 12 samples
    element names: exprs
## protocolData: none
## phenoData
     sampleNames: PHLC1106 PHLC111 ... PHLC82 (12 total)
##
     varLabels: PHLC.ID X.ID
##
     varMetadata: labelDescription
## featureData
     featureNames: NOC2L ISG15 ... RNF128 (16116 total)
##
    fvarLabels: probe.Id
##
     fvarMetadata: labelDescription
## experimentData: use 'experimentData(object)'
## Annotation: MUT
```

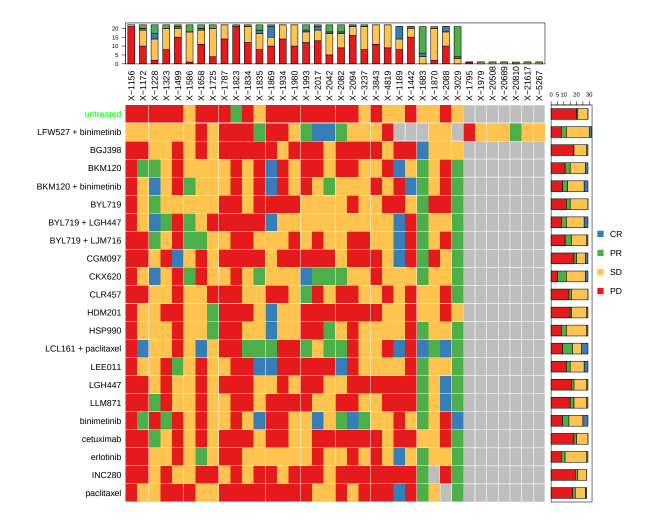
The sample names in expression set are called biobase.id in model slot. Sample names from the expression set canb be be mapped to individual PDX model.ids as

```
# get sample names
library(Biobase)
```

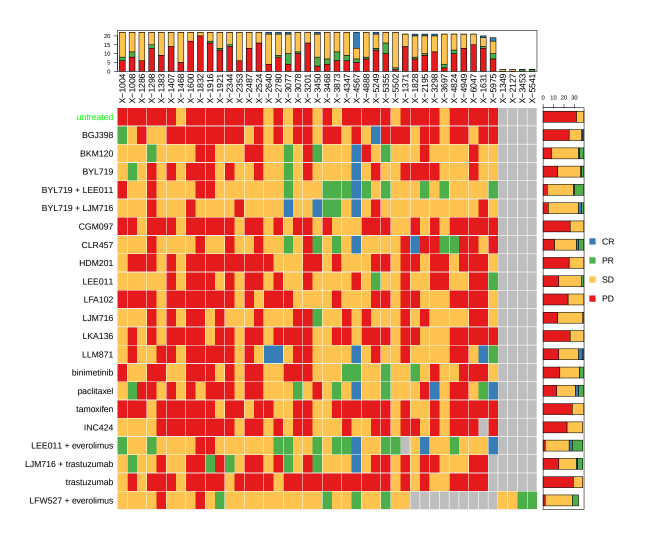
```
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
##
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
##
       clusterExport, clusterMap, parApply, parCapply, parLapply,
##
       parLapplyLB, parRapply, parSapply, parSapplyLB
## The following objects are masked from 'package:stats':
##
       IQR, mad, xtabs
##
## The following objects are masked from 'package:base':
##
##
       Filter, Find, Map, Position, Reduce, anyDuplicated, append,
       as.data.frame, cbind, colnames, do.call, duplicated, eval,
##
```

```
##
       evalq, get, grep, grepl, intersect, is.unsorted, lapply,
##
       lengths, mapply, match, mget, order, paste, pmax, pmax.int,
       pmin, pmin.int, rank, rbind, rownames, sapply, setdiff, sort,
##
##
       table, tapply, union, unique, unsplit, which, which.max,
##
       which.min
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
##
       'browseVignettes()'. To cite Bioconductor, see
##
       'citation("Biobase")', and for packages 'citation("pkgname")'.
sn <- Biobase::sampleNames(ldxe_mut)</pre>
smap <- mapModelSlotIds(lpdx, id=sn, id.name = "biobase.id", map.to = "model.id")</pre>
head(smap)
                        biobase.id
                                               model.id
## PHLC1106_P5.501.A1.1 PHLC1106 PHLC1106_P5.501.A1.1
## PHLC1106_P5.504.A4.1 PHLC1106 PHLC1106_P5.504.A4.1
## PHLC1106_P5.506.B1.1 PHLC1106 PHLC1106_P5.506.B1.1
## PHLC1106_P5.507.B2.1 PHLC1106 PHLC1106_P5.507.B2.1
## PHLC1106_P5.508.B3.1 PHLC1106 PHLC1106_P5.508.B3.1
## PHLC1106_P5.511.C1.1 PHLC1106 PHLC1106_P5.511.C1.1
What should we do here
df = getExperiment(lpdx, "PHLC119 P5.506.B1.3")
print(df[df$time>85 & df$time<109, c("time", "width", "length", "volume", "comment", "dose")])</pre>
     time width length
##
                        volume
## 22
       88 3.20
                 4.01 21.35245
                 1.00 0.52000
## 23
       92 1.00
       95 3.71
                 5.19 37.14655
## 24
## 25
       99 1.00
                 1.00 0.52000
## 26 102 4.13
                 4.98 44.17055
## 27 106 3.32 4.22 24.18755
##
                                            comment
                                                        dose
## 22
                                               <NA> 78.51786
## 23 small bud - put in arbritary "1" measurements 82.25357
                                               <NA> 79.66071
## 25 small bud - put in arbritary "1" measurements 83.27500
## 26
                                               <NA> 81.25357
## 27
                                     Stop treatment 0.00000
```

Create mRECIST plot for PDXE Lung Cancer data



```
#pdf(file="DATA-raw/mRECIST_plot_BRCA.pdf", width=12, height=10)
brDF = df[df$tumor.type=="BRCA", ]
plotmRECIST(brDF, groupBy = "biobase.id", control.name = "untreated")
```



#dev.off()

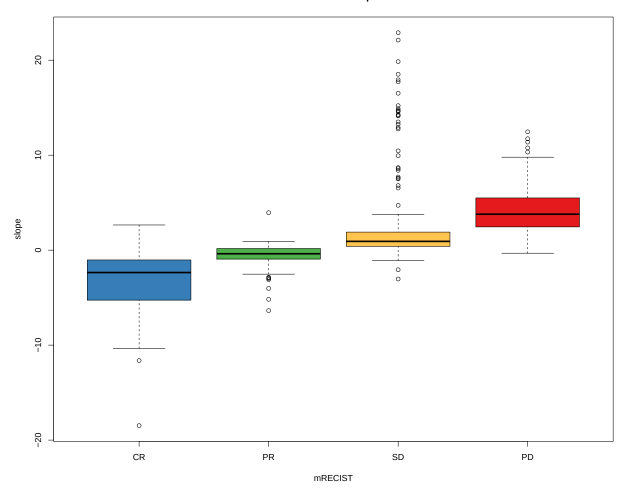
Creat mR vs slop bar-plot

```
slope=c(); mR=c()
for(dn in rownames(lung_pdxe_slope))
{
   for(pi in colnames(lung_pdxe_slope))
   {
      v = c(lung_pdxe_slope[dn,pi], lung_pdxe_mR[dn,pi])
      if(!is.na(v[1]) & !is.na(v[1]))
      { slope = c(slope,v[1]); mR=c(mR,v[2]) }
}

df = data.frame(mR= mR, slope= as.numeric(slope), stringsAsFactors = FALSE)
df$mR= factor(df$mR, c("CR", "PR", "SD", "PD"))

colPalette = c("#377eb8", "#4daf4a", "#fec44f", "#e41a1c")
#pdf(file="DATA-raw/boxplot_lungCancer.pdf", width=12, height=10)
boxplot(slope~mR, data=df, col=colPalette,
   main="mRECIST vs slope", xlab="mRECIST", ylab="slope")
```

mRECIST vs slope



#dev.off()