# Xeva Tutorial

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Load Xeva library and KRAS/P53 PDX data

```
library(Xeva)
data(lpdx)
To see all the model.id
lpdx.mod = modelInfo(lpdx)
head(lpdx.mod$model.id)
## [1] "PHLC1106_P5.501.A1" "PHLC1106_P5.504.A4" "PHLC1106_P5.506.B1"
## [4] "PHLC1106 P5.507.B2" "PHLC1106 P5.508.B3" "PHLC1106 P5.511.C1"
To get the data for one model.id
modId = lpdx.mod$model.id[82]
df = getExperiment(lpdx, model.id = modId)
head(df)
##
              model.id
                                drug.join.name time
                                                        volume width length
## 1 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                      81.20558
                                                               5.18
                                                                        5.82
                                                   Ω
## 2 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                   8
                                                      93.24844
                                                                5.57
                                                                        5.78
## 3 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                      90.13298
                                                                5.16
                                                                        6.51
## 4 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                  19 213.92906
                                                                6.99
                                                                        8.42
## 5 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                  22 252.04349 7.43
                                                                        8.78
## 6 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                  26 375.84838 8.65
                                                                        9.66
##
     dose body.weight
                             date
                                           comment volume.change
     0.0
               19.762 2014-09-25
                                              <NA>
                                                         0.00000
## 1
## 2 0.0
               20.424 2014-10-03
                                                        14.83010
                                     clip removed
## 3 0.0
               21.130 2014-10-10
                                              <NA>
                                                        10.99359
## 4 75.4
               21.103 2014-10-14 Start Treatment
                                                       163.44135
## 5 74.1
               20.761 2014-10-17
                                              <NA>
                                                       210.37708
## 6 72.1
               20.178 2014-10-21
                                              <NA>
                                                       362.83569
##
     average.response
## 1
             0.000000
## 2
             7.415048
## 3
             8.607894
## 4
            47.316257
## 5
            79.928421
## 6
           127.079632
In the data fram df you will see that for first 3 time points dose is 0, which indicate no treatment is given
```

during this time. If you want the data only during the treatment periode specify treatment only = TRUE

```
df = getExperiment(lpdx, modId, treatment.only = TRUE)
head(df)
```

```
##
              model.id
                               drug.join.name time
                                                      volume width length dose
## 4 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                                     8.42 75.4
                                                 19 213.9291
                                                              6.99
## 5 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                              7.43
                                                 22 252.0435
                                                                     8.78 74.1
## 6 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                 26 375.8484 8.65
                                                                     9.66 72.1
```

```
## 7 PHLC191 P5.503.A3 Vinorelbine+ Cisplatin
                                                29 526.0954 9.40 11.45 73.3
## 8 PHLC191_P5.503.A3 Vinorelbine+ Cisplatin
                                                33 683.3432 10.43 12.08 73.3
## 9 PHLC191 P5.503.A3 Vinorelbine+ Cisplatin
                                                36 807.8725 10.97 12.91 75.9
##
     body.weight
                                    comment volume.change average.response
                       date
## 4
          21.103 2014-10-14 Start Treatment
                                                 163.4413
                                                                  47.31626
## 5
                                                                  79.92842
          20.761 2014-10-17
                                       <NA>
                                                 210.3771
## 6
                                                                 127.07963
          20.178 2014-10-21
                                       <NA>
                                                 362.8357
## 7
          20.528 2014-10-24
                                       <NA>
                                                 547.8563
                                                                  187.19059
## 8
          20.534 2014-10-28
                                       <NA>
                                                 741.4979
                                                                  256.47900
## 9
                                       <NA>
          21.257 2014-10-31
                                                 894.8486
                                                                 327.40896
```

Models which are replicates are stored togather in expDesign slot. To get the data for all the replicates pass the 'batch.name' in the getExperiment function.

```
print(batchNames(lpdx))
   [1] "PHLC1106 P5" "PHLC111 P7"
                                                   "PHLC153 P6"
                                                                 "PHLC181_P7"
                                    "PHLC119 P5"
  [6] "PHLC189_P5" "PHLC191_P5"
                                    "PHLC191_P7"
                                                   "PHLC196 P5"
                                                                 "PHLC215 P5"
## [11] "PHLC229_P6"
                      "PHLC235_P4"
                                    "PHLC655_P7"
                                                   "PHLC82_P5"
df = getExperiment(lpdx, batch.name = batchNames(lpdx)[1], treatment.only = TRUE)
head(df)
##
                model.id
                                 drug.join.name time
                                                        volume width length
## 8 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                   39 167.5273
                                                                6.46
                                                                       7.72
## 9 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                                6.48
                                                                       7.91
                                                   40 172.7149
## 10 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                   43 164.4621
                                                                6.38
                                                                       7.77
## 11 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                                6.76
                                                                       7.89
                                                   47 187.4881
## 12 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                   50 264.0641 7.64
                                                                       8.70
  13 PHLC1106_P5.502.A2 Vinorelbine+ Cisplatin
                                                   54 174.6014
                                                                6.66
                                                                       7.57
##
      dose body.weight
                             date
                                           comment volume.change
                22.182 2014-12-08 Start Treatment
## 8 79.2
                                                        126.5905
## 9 78.8
                22.051 2014-12-09
                                                        133.6070
                                              <NA>
## 10 78.6
                21.995 2014-12-12
                                              < NA >
                                                        122.4445
## 11 78.0
                21.827 2014-12-16
                                              <NA>
                                                        153.5886
## 12 80.2
                22.467 2014-12-19
                                              <NA>
                                                        257.1621
                22.185 2014-12-23
                                              <NA>
                                                        136.1586
## 13 79.2
##
      average.response exp.type
              47.16826 treatment
## 8
## 9
              56.77256 treatment
## 10
              63.33976 treatment
## 11
              71.54420 treatment
## 12
              87.01235 treatment
## 13
              90.79283 treatment
```

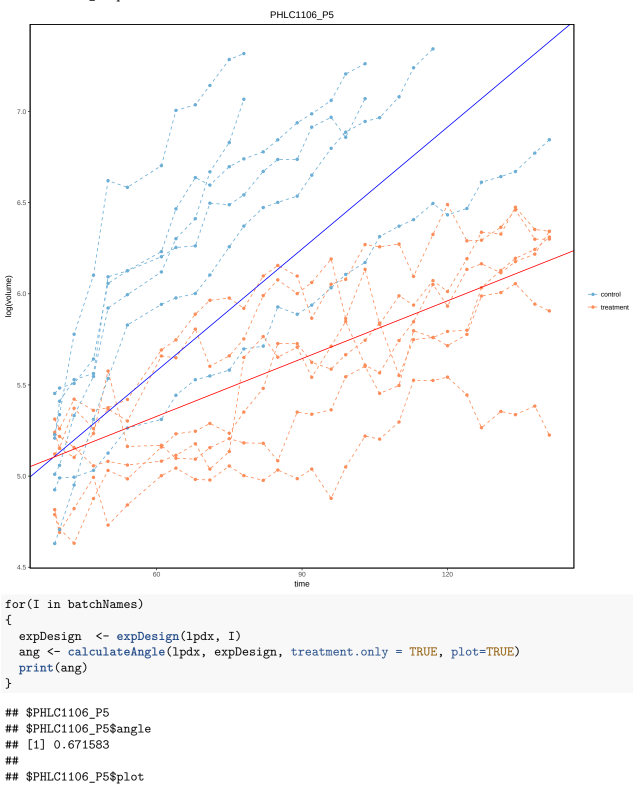
Here the data fram contaions an extra column 'exp.type'. This indicates if this is treatment or control.

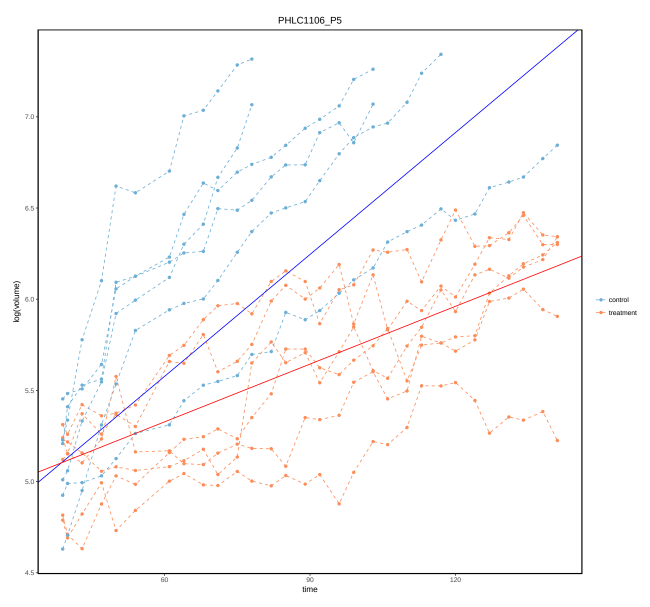
To calculate angle between the treatment and control samples of this batch

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

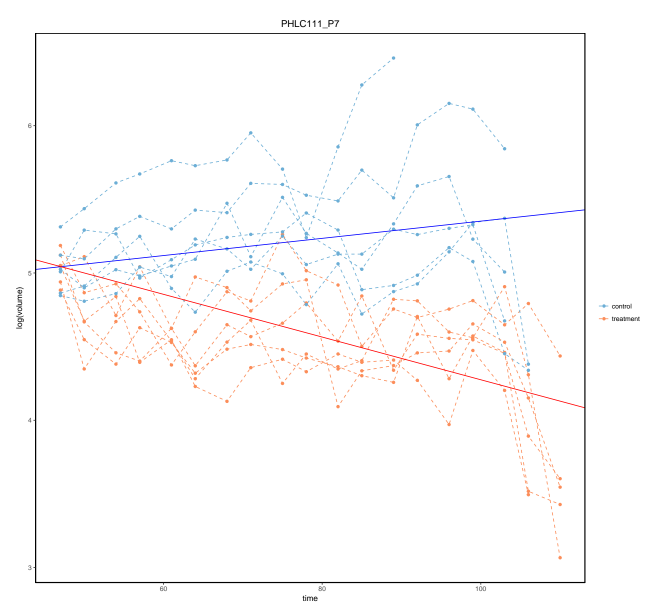
## $PHLC1106_P5
## $PHLC1106_P5$angle
## [1] 0.671583
##</pre>
```

## ## \$PHLC1106\_P5\$plot

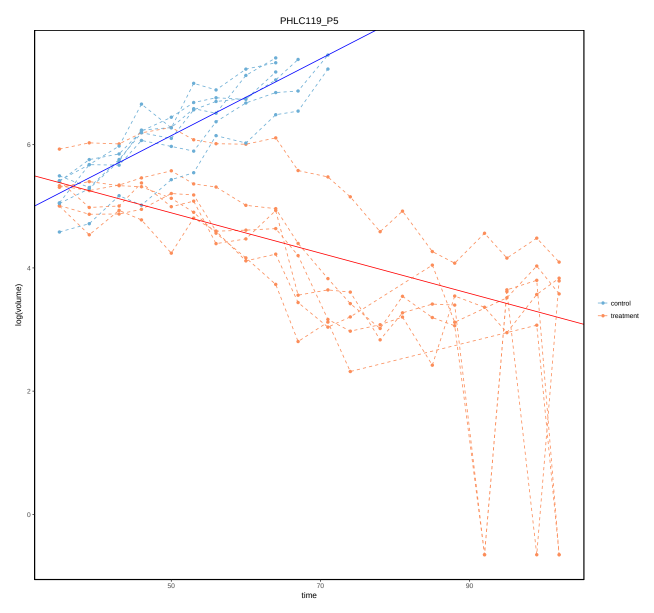




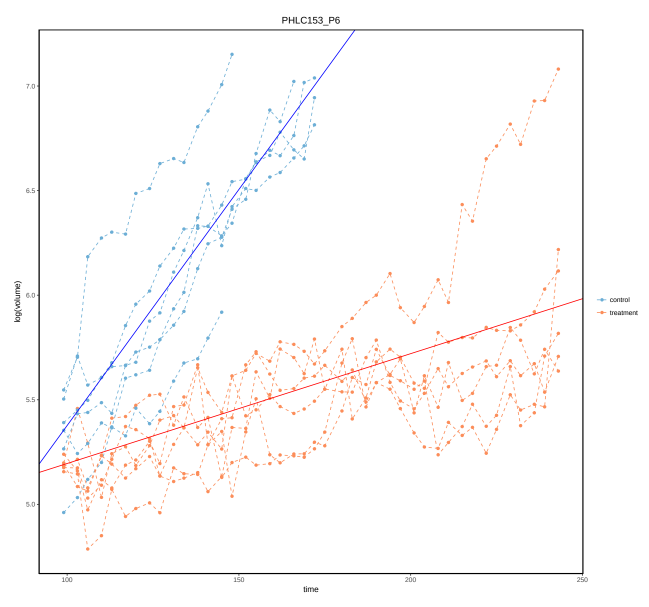
```
##
##
## $PHLC111_P7
## $PHLC111_P7$angle
## [1] 1.165361
##
## $PHLC111_P7$plot
```



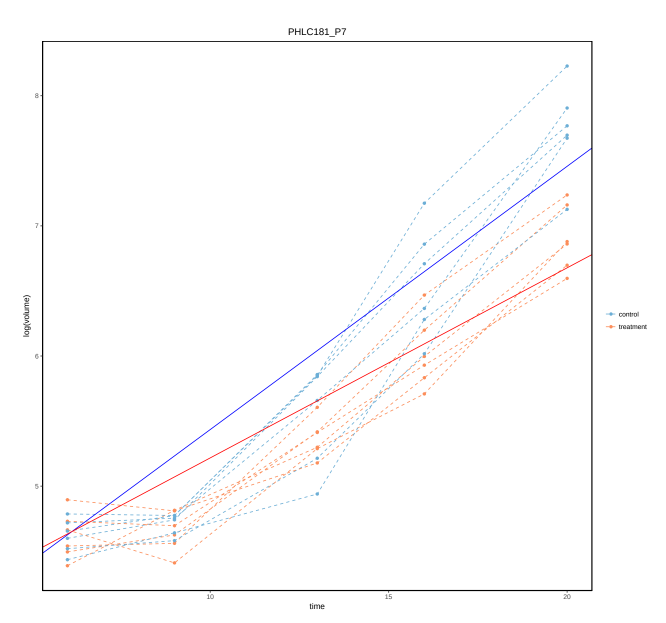
```
## ## ## $PHLC119_P5
## $PHLC119_P5$angle
## [1] 5.437795
## ## $PHLC119_P5$plot
```



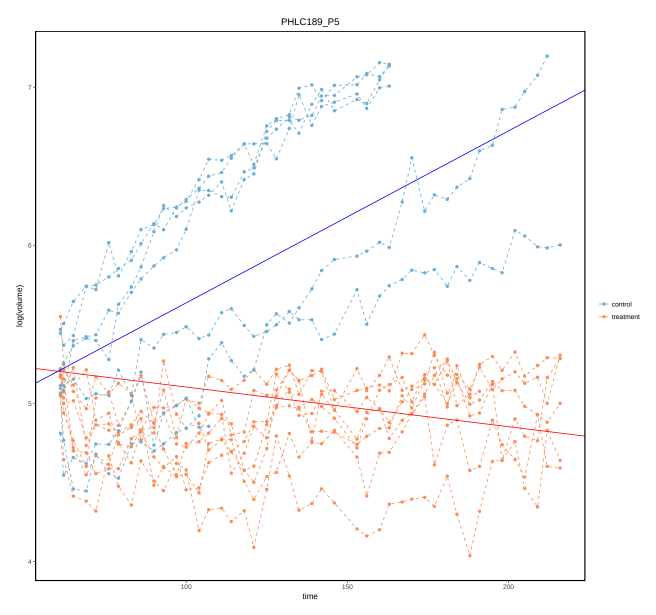
```
##
##
## $PHLC153_P6
## $PHLC153_P6$angle
## [1] 0.9921273
##
## $PHLC153_P6$plot
```



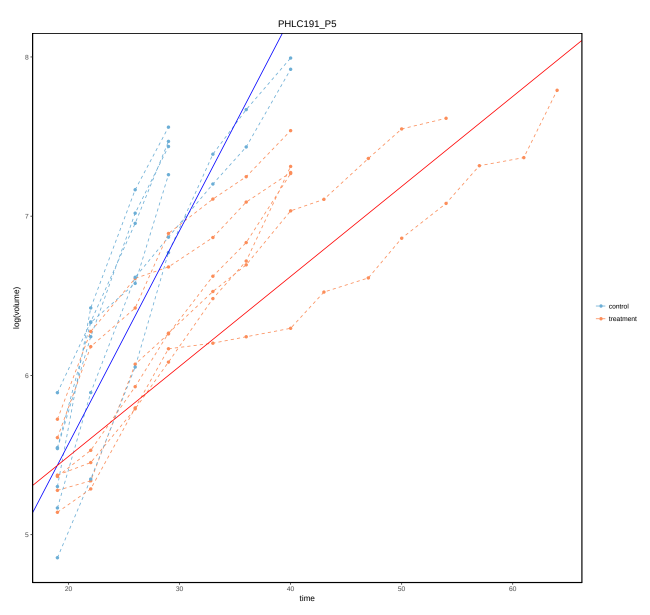
```
##
##
## $PHLC181_P7
## $PHLC181_P7$angle
## [1] 3.125739
##
## $PHLC181_P7$plot
```



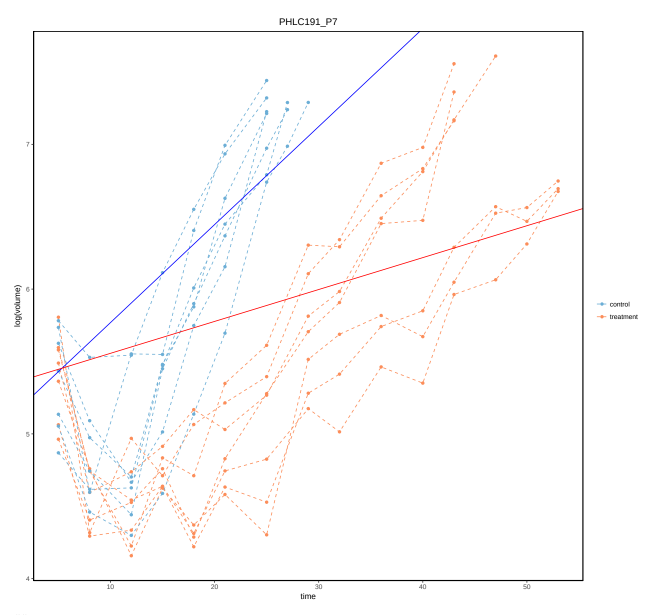
```
##
##
## $PHLC189_P5
## $PHLC189_P5$angle
## [1] 0.7670435
##
## $PHLC189_P5$plot
```



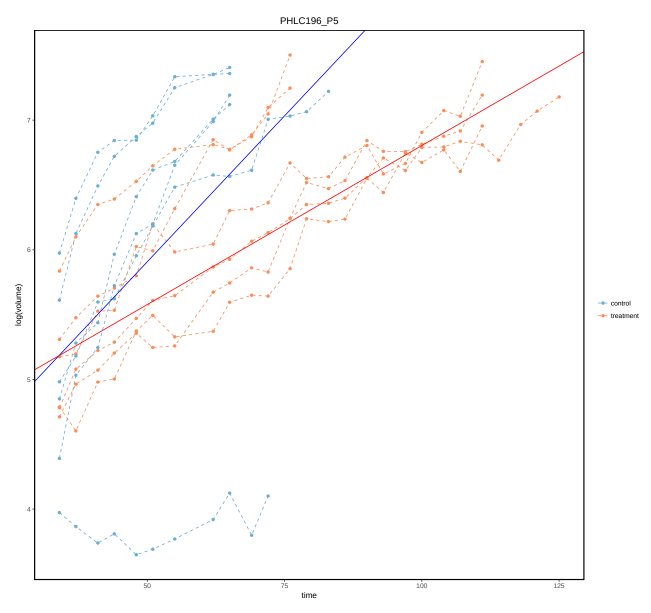
```
##
##
## $PHLC191_P5
## $PHLC191_P5$angle
## [1] 4.391481
##
## $PHLC191_P5$plot
```



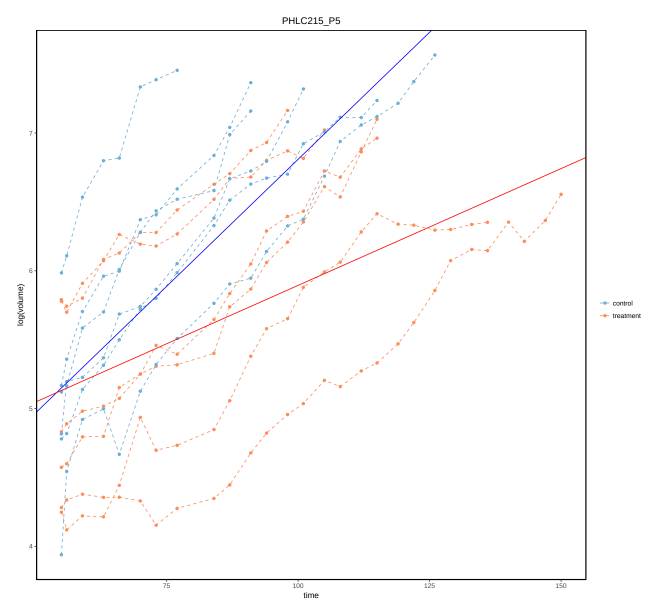
```
##
##
## $PHLC191_P7
## $PHLC191_P7$angle
## [1] 2.610878
##
## $PHLC191_P7$plot
```



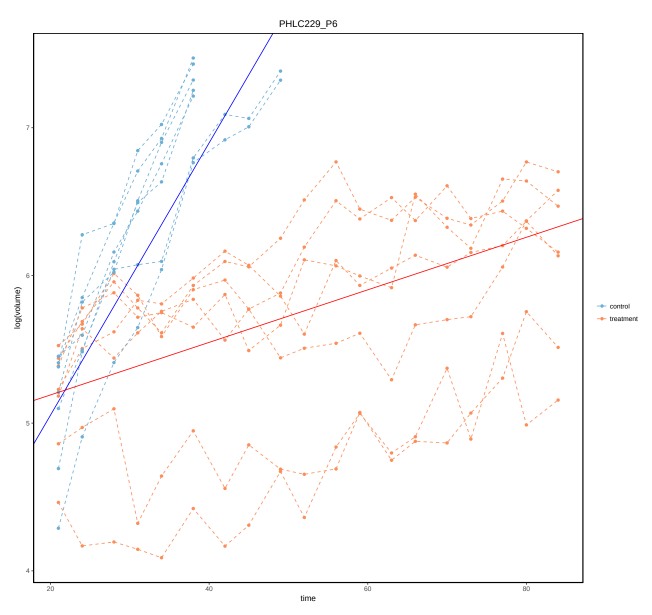
```
##
##
## $PHLC196_P5
## $PHLC196_P5$angle
## [1] 1.172456
##
## $PHLC196_P5$plot
```



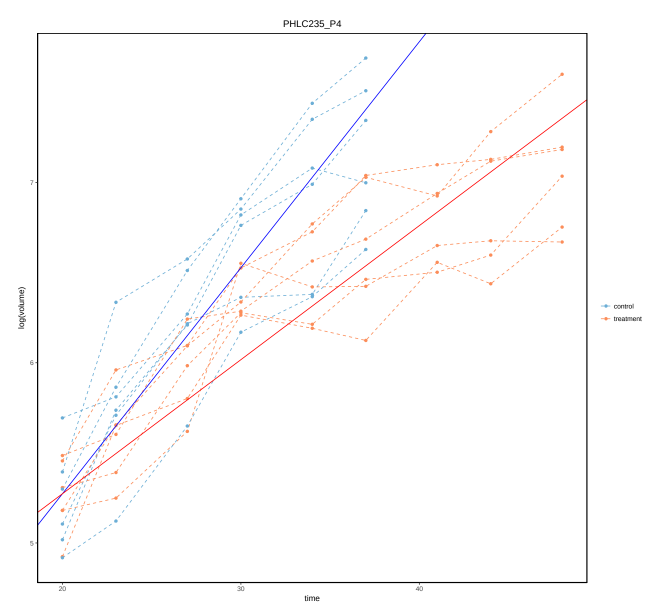
```
##
##
## $PHLC215_P5
## $PHLC215_P5$angle
## [1] 1.140868
##
## $PHLC215_P5$plot
```



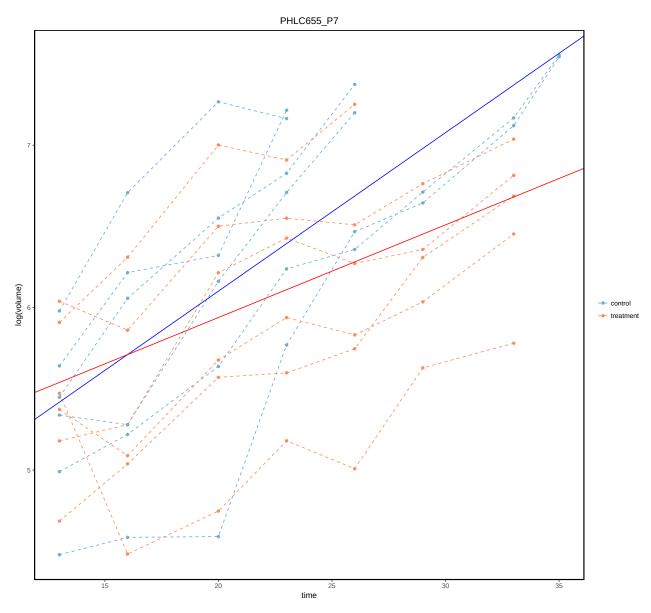
```
## ## $PHLC229_P6 ## $PHLC229_P6$angle ## [1] 4.245984 ## ## $PHLC229_P6$plot
```



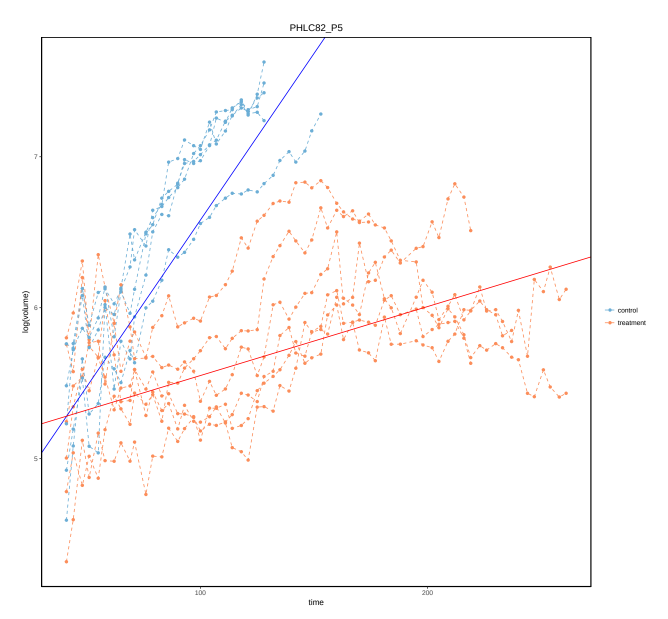
```
##
##
## $PHLC235_P4
## $PHLC235_P4$angle
## [1] 2.89404
##
## $PHLC235_P4$plot
```



```
##
##
## $PHLC655_P7
## $PHLC655_P7$angle
## [1] 2.309245
##
## $PHLC655_P7$plot
```



```
##
##
## $PHLC82_P5
## $PHLC82_P5$angle
## [1] 1.00194
##
##
## $PHLC82_P5$plot
```



Summarize Response of PDXs Get slop of each model and combine summarize all model slop which belongs to same patient by "mean"

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

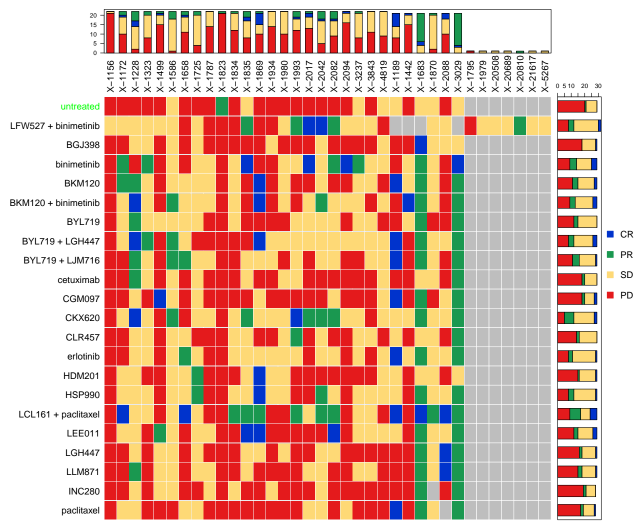
```
lpdx_angle <- summarizeResponse(lpdx, response.measure = "angle")</pre>
```

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

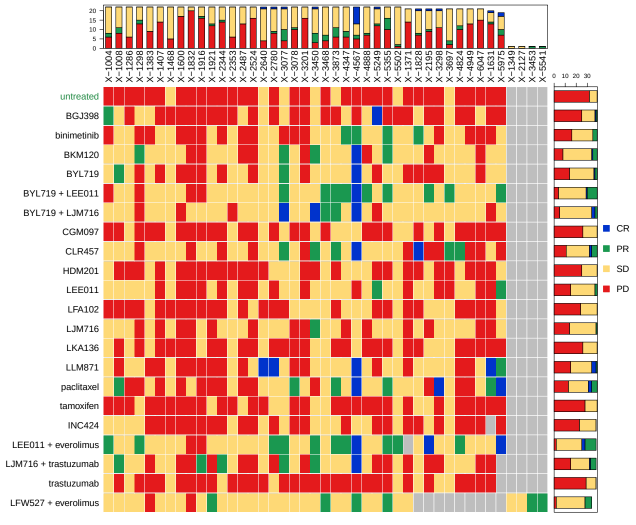
```
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 16116 features, 12 samples
     element names: exprs
## protocolData: none
## phenoData
##
     sampleNames: PHLC1106 PHLC111 ... PHLC82 (12 total)
     varLabels: PHLC.ID X.ID
     varMetadata: labelDescription
##
## featureData
     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)
sn <- Biobase::sampleNames(ldxe_mut)</pre>
smap <- mapModelSlotIds(lpdx, id=sn, id.name = "biobase.id", map.to = "model.id")</pre>
head(smap)
                                            model.id
##
                      biobase.id
## PHLC1106_P5.501.A1 PHLC1106 PHLC1106_P5.501.A1
## PHLC1106_P5.504.A4 PHLC1106 PHLC1106_P5.504.A4
## PHLC1106_P5.506.B1 PHLC1106_P5.506.B1
## PHLC1106_P5.507.B2 PHLC1106 PHLC1106_P5.507.B2
## PHLC1106_P5.508.B3
                        PHLC1106 PHLC1106_P5.508.B3
## PHLC1106_P5.511.C1
                        PHLC1106 PHLC1106_P5.511.C1
What should we do here
df = getExperiment(lpdx, "PHLC119_P5.506.B1")
```

 $\#print(df[df\$time>85\ \&\ df\$time<109,\ c("time", "width", "length", "volume", "comment", "dose")])$ 

#### Create mRECIST plot for PDXE Lung Cancer data



Create mRECIST plot for PDXE Breast Cancer data



Creat mR vs slop bar-plot

### mRECIST vs slope

