# Penalty function learning

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## 1 Introduction

In supervised changepoint detection, we are given n labeled data sequences. For each data sequence  $i \in \{1, ..., n\}$  we have a vector of noisy data  $\mathbf{z}_i \in \mathbb{R}^{d_i}$ , a set of labels  $L_i$  which indicates appropriate changepoint positions, and an input/feature  $\mathbf{x}_i \in \mathbb{R}^p$ . Note that the number of data points to segment  $d_i$  can be different for every data sequence i, but the number of features p is constant.

The optimal changepoint model for data sequence i, and a penalty parameter  $\lambda$ , is given by

$$\mathbf{m}_{i}^{\lambda} = \underset{\mathbf{m} \in \mathbb{R}^{d_{i}}}{\min} \, \ell(\mathbf{m}, \mathbf{z}_{i}) + \lambda \mathcal{C}(\mathbf{m}), \tag{1}$$

where

- $\ell$  is a loss function, e.g. the square loss  $\ell(\mathbf{m}, \mathbf{z}_i) = \sum_{j=1}^{d_i} (m_j z_{ij})^2$ ,
- C is a model complexity function, e.g. the number of changes  $C(\mathbf{m}) = \sum_{j=1}^{d_i-1} I(m_j \neq m_{j+1}),$
- $\lambda \geq 0$  is a penalty constant. Larger values penalize more complex models, and result in few changepoints ( $\lambda = \infty$  means no changes). Smaller values penalize less and result in more changepoints ( $\lambda = 0$  means  $d_i 1$  changes).

In this context, we would like to learn a different penalty constant  $\log \lambda_i = f(\mathbf{x}_i)$  for every data sequence, where  $f: \mathbb{R}^p \to \mathbb{R}$  is a function that we will learn by minimizing the number of incorrect labels in the training data:

$$\min_{f} \sum_{i=1}^{n} e[\mathbf{m}_{i}^{\exp f(\mathbf{x}_{i})}, L_{i}]. \tag{2}$$

The function e is the number of labels in  $L_i$  which are incorrectly predicted by the changepoint model  $\mathbf{m}_i^{\exp f(x_i)}$ .

After having learned f on training data, it can be used to predict a changepoint model for a test data sequence  $\mathbf{z} \in \mathbb{R}^d$  which has features  $\mathbf{x} \in \mathbb{R}^p$ . First compute a predicted penalty value  $\lambda = \exp f(\mathbf{x})$ , and then use it to compute a predicted segmentation model  $\mathbf{m}^{\lambda}$ .

## 2 Details

## 2.1 Changepoint model fitting

For each of several labeled segmentation problems (data sequences that are separate but have a similar signal/noise pattern), use your favorite changepoint detection package to compute a sequence of models of increasing complexity (say from 0 to 20 changepoints). In contrast to unsupervised changepoint detection (where we usually compute just one changepoint model per data sequence), it is essential to compute several models in supervised changepoint detection (so that we can learn which models and penalty values result in changepoints with minimum error with repect to the labels). Below we use the Segmentor function to compute a maximum likelihood Gaussian model.

> library(data.table)
> data(neuroblastoma, package="neuroblastoma")
> ids.str <- paste(c(1, 4, 6, 8, 10, 11))
> someProfiles <- function(all.profiles){
+ data.table(all.profiles)[profile.id %in% ids.str, ]</pre>

```
+ }
> profiles <- someProfiles(neuroblastoma$profiles)</pre>
> labels <- someProfiles(neuroblastoma$annotations)</pre>
> problem.list <- split(profiles, profiles[, paste(profile.id, chromosome)])</pre>
> segs.list <- list()
> loss.list <- list()</pre>
> for(problem.i in seq_along(problem.list)){
    problem.name <- names(problem.list)[[problem.i]]</pre>
    pro <- problem.list[[problem.name]]</pre>
    meta <- pro[1, .(profile.id, chromosome)]</pre>
    max.segments <- min(nrow(pro), 10)</pre>
    fit <- Segmentor3IsBack::Segmentor(</pre>
      pro$logratio, model=2, Kmax=max.segments)
    rss.vec <- rep(NA, max.segments)</pre>
    for(n.segments in 1:max.segments){
      end <- fit@breaks[n.segments, 1:n.segments]</pre>
      data.before.change <- end[-n.segments]</pre>
      data.after.change <- data.before.change+1
      pos.before.change <- as.integer(</pre>
      (pro$position[data.before.change]+pro$position[data.after.change])/2)
      start <- c(1, data.after.change)</pre>
      chromStart <- c(pro$position[1], pos.before.change)</pre>
      chromEnd <- c(pos.before.change, max(pro$position))</pre>
      seg.mean.vec <- fit@parameters[n.segments, 1:n.segments]</pre>
      data.mean.vec <- rep(seg.mean.vec, end-start+1)</pre>
      residual.vec <- pro$logratio - data.mean.vec
      rss.vec[n.segments] <- sum(residual.vec * residual.vec)
      segs.list[[paste(problem.name, n.segments)]] <- data.table(</pre>
        meta,
        n.segments,
        start,
        end.
+
        chromStart,
+
        chromEnd,
        mean=seg.mean.vec)
+
    loss.list[[paste(problem.name, n.segments)]] <- data.table(</pre>
      n.segments=1:max.segments,
      loss=rss.vec,
      lik=as.numeric(fit@likelihood))
+ }
> loss <- do.call(rbind, loss.list)</pre>
> segs <- do.call(rbind, segs.list)</pre>
```

Note that since we have saved the segment starts and ends, we can easily derive the changepoint positions of each model:

### > print(segs)

```
profile.id chromosome n.segments start end chromStart chromEnd
                                   1
                                         1 474
  1:
              1
                        1
                                                   809681 249063592
  2:
              1
                         1
                                    2
                                         1 438
                                                   809681 212809180
  3:
                         1
                                    2
                                      439 474 212809180 249063592
              1
  4:
                         1
                                    3
                                         1 437
                                                   809681 212280934
              1
                         1
                                    3
                                       438 460 212280934 234068672
  5:
7871:
                         Y
                                    3
                                         4
                                             4
                                                 16036249 25746303
```

```
7872:
                8
                           Y
                                                  1
                                       4
                                              1
                                                         293556
                                                                  1443054
                           Y
                                              2
7873:
                8
                                       4
                                                  2
                                                        1443054
                                                                  4459374
                8
                           Y
                                       4
                                              3
                                                  3
7874:
                                                        4459374
                                                                 16036249
7875:
                8
                           Υ
                                       4
                                              4
                                                  4
                                                       16036249
                                                                 25746303
             mean
   1:
       0.31345083
   2:
       0.35186515
   3: -0.15392338
      0.35242544
       0.02954558
   5:
7871: -1.51045706
7872: -0.09080294
7873: -0.60384051
7874: -2.25842515
7875: -1.51045706
```

Also note that we have saved the loss and model complexity (n.segments) of each model,

### > print(loss)

3:

|       | <pre>profile.id</pre> | chromosome | n.segments | loss       | lik         |
|-------|-----------------------|------------|------------|------------|-------------|
| 1:    | 1                     | 1          | 1          | 15.9149875 | 364.634368  |
| 2:    | 1                     | 1          | 2          | 7.4048569  | -221.666054 |
| 3:    | 1                     | 1          | 3          | 5.5191996  | -351.577300 |
| 4:    | 1                     | 1          | 4          | 4.3030047  | -435.366330 |
| 5:    | 1                     | 1          | 5          | 4.0235352  | -454.620216 |
|       |                       |            |            |            |             |
| 1430: | 8                     | Х          | 10         | 0.2985018  | -130.032751 |
| 1431: | 8                     | Y          | 1          | 2.7740679  | 5.174541    |
| 1432: | 8                     | Y          | 2          | 0.4113319  | 4.163931    |
| 1433: | 8                     | Y          | 3          | 0.1316038  | 4.044283    |
| 1434: | 8                     | Y          | 4          | 0.0000000  | 3.987992    |

### 2.2 Model selection functions

8 3.557869 -486.7020

We use penaltyLearning::modelSelection to compute the exact path of models that will be selected for every possible non-negative penalty value,

```
> selection <- loss[, {
    penaltyLearning::modelSelection(.SD, "loss", "n.segments")
+ }, by=list(profile.id, chromosome)]
> print(selection[profile.id==1 & chromosome==1])
   profile.id chromosome min.lambda max.lambda min.log.lambda max.log.lambda
                        1 0.00000000 0.07175223
1:
            1
                                                           -Inf
                                                                     -2.6345364
                                                                     -2.0817655
2:
            1
                        1 0.07175223 0.12470984
                                                     -2.6345364
                        1 0.12470984 0.12800334
                                                                     -2.0556989
3:
            1
                                                     -2.0817655
4:
            1
                        1 0.12800334 0.20965938
                                                     -2.0556989
                                                                     -1.5622711
5:
            1
                        1 0.20965938 0.27946950
                                                     -1.5622711
                                                                     -1.2748621
6:
            1
                        1 0.27946950 1.21619490
                                                     -1.2748621
                                                                      0.1957271
7:
            1
                        1 1.21619490 1.88565729
                                                      0.1957271
                                                                      0.6342765
8:
            1
                        1 1.88565729 8.51013055
                                                      0.6342765
                                                                      2.1412573
            1
                        1 8.51013055
                                                      2.1412573
                                                                            Inf
   n.segments
                               lik
                   loss
1:
           10
               3.361407 -500.2371
2:
            9
               3.433159 -495.2938
```

```
4: 6 3.813876 -469.0646

5: 5 4.023535 -454.6202

6: 4 4.303005 -435.3663

7: 3 5.519200 -351.5773

8: 2 7.404857 -221.6661

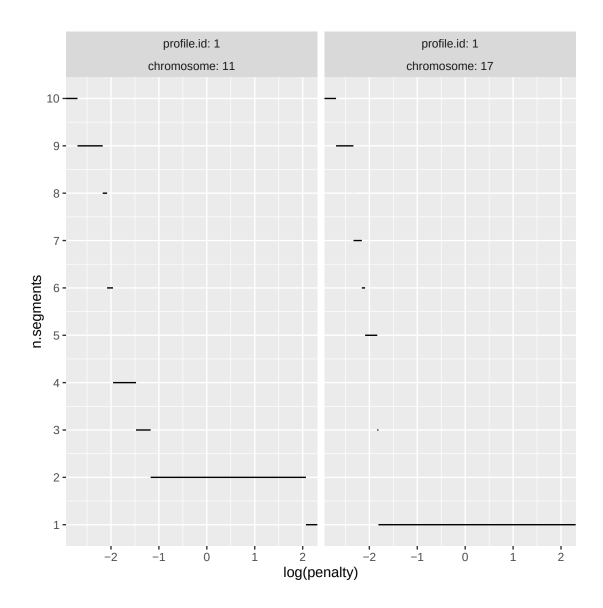
9: 1 15.914987 364.6344
```

Note that the model selection function  $s_i(\lambda)$  is the number of segments that is selected for a given penalty  $\lambda$  and data sequence i,

$$s_i(\lambda) = \underset{s}{\operatorname{arg \, min}} \, \ell(\mathbf{m}, \mathbf{z}_i) + \lambda \mathcal{C}(\mathbf{m}).$$
 (3)

These functions are piecewise constant, and specific to each data sequence:

```
> some.selection <- selection[profile.id==1 & chromosome %in% c(11, 17)]
> some.selection[, list(
   pid.chr=paste(profile.id, chromosome),
   min.log.lambda, max.log.lambda, n.segments, loss
+ )]
   pid.chr min.log.lambda max.log.lambda n.segments
                                                           loss
 1:
       1 11
                      -Inf
                                -2.698480
                                                   10
                                                       1.514946
2:
       1 11
                 -2.698480
                                                    9
                                -2.172301
                                                       1.582253
3:
       1 11
                 -2.172301
                                -2.080750
                                                    8 1.696169
4:
       1 11
                 -2.080750
                                -1.957421
                                                    6 1.945842
                 -1.957421
5:
       1 11
                                -1.477048
                                                    4 2.228286
                                                    3 2.456597
6:
       1 11
                 -1.477048
                                -1.172320
7:
                                 2.067398
                                                    2 2.766245
       1 11
                 -1.172320
       1 11
8:
                  2.067398
                                                    1 10.670475
                                       Inf
       1 17
                                -2.694380
                                                   10 1.398609
9:
                      -Inf
       1 17
                                                    9 1.466194
10:
                 -2.694380
                                -2.332349
                                                    8 1.563261
                                -2.330538
11:
       1 17
                 -2.332349
12:
       1 17
                 -2.330538
                                -2.157175
                                                    7
                                                      1.660504
13:
       1 17
                 -2.157175
                                -2.087849
                                                    6 1.776156
                                                    5 1.900109
14:
       1 17
                 -2.087849
                                -1.834952
       1 17
                 -1.834952
                                -1.810653
                                                    3 2.219352
15:
16:
       1 17
                 -1.810653
                                       Inf
                                                    1 2.546446
> library(ggplot2)
> gg.selection <- ggplot()+
    theme_grey()+
    facet_grid(. ~ profile.id + chromosome, labeller=label_both)+
   geom_segment(aes(
      min.log.lambda, n.segments,
      xend=max.log.lambda, yend=n.segments),
      data=some.selection)+
    scale_y_continuous(breaks=1:max(some.selection$n.segments))+
    xlab("log(penalty)")
> print(gg.selection)
```



## 2.3 Label error

To compute the label error function  $E_i(\lambda)$  (number of incorrect labels for data sequence i as a function of penalty  $\lambda$ ), we need to first get the positions of predicted changepoints:

- > changes <- segs[1 < start, ]</pre>
- > changes[, list(profile.id, chromosome, n.segments, changepoint=chromStart)]

|       | profile.id | chromosome | n.segments | changepoint |
|-------|------------|------------|------------|-------------|
| 1:    | 1          | 1          | 2          | 212809180   |
| 2:    | 1          | 1          | 3          | 212280934   |
| 3:    | 1          | 1          | 3          | 234068672   |
| 4:    | 1          | 1          | 4          | 40348010    |
| 5:    | 1          | 1          | 4          | 212280934   |
|       |            |            |            |             |
| 6437: | 8          | Y          | 3          | 4459374     |
| 6438: | 8          | Y          | 3          | 16036249    |
| 6439: | 8          | Y          | 4          | 1443054     |
| 6440: | 8          | Y          | 4          | 4459374     |
| 6441: | 8          | Y          | 4          | 16036249    |

Then, we use penaltyLearning::labelError to compute the number of incorrect labels for each model, for each labeled data sequence. The named arguments specify how the three input data tables are used:

change.var is the column name of changes that will be used as the changepoint position.

label.vars are the column names of the start and end of the labeled regions.

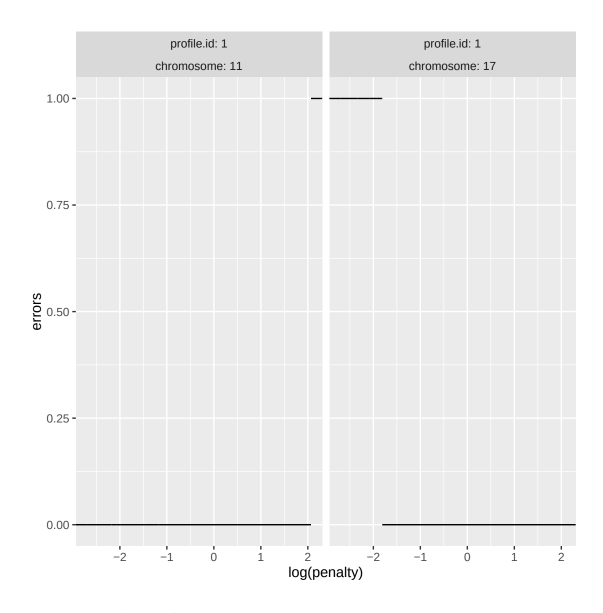
problem.vars are the column names (common to all data tables) that are used to identify independent data sequences.

```
> errors <- penaltyLearning::labelError(
+ selection, labels, changes,
+ change.var="chromStart",
+ label.vars=c("min", "max"),
+ problem.vars=c("profile.id", "chromosome"))</pre>
```

The labelError function returns a list of two data tables: label.errors has one row for each label and each model, and model.errors has one row for each data sequence and each model. In this case they are the same size because the neuroblastoma data set has only one label per data sequence. The model.errors  $E_i(\lambda)$  is a piecewise constant function for every data sequence i (the number of incorrect labels as a function of penalty  $\lambda$ ):

$$E_i(\lambda) = e[\mathbf{m}_i^{\lambda}, L_i]. \tag{4}$$

```
> some.errors <- errors$model.errors[some.selection, on=list(
+ profile.id, chromosome, n.segments)]
> gg.err <- ggplot()+
+ theme_grey()+
+ facet_grid(. ~ profile.id + chromosome, labeller=label_both)+
+ geom_segment(aes(
+ min.log.lambda, errors,
+ xend=max.log.lambda, yend=errors),
+ data=some.errors)+
+ xlab("log(penalty)")
> print(gg.err)
```



## 2.4 Target intervals

Now, let's perform a computational cross-validation experiment to train and evaluate a learned penalty function. We use labels in chromosome 11 as a test set, and use all other labels as a train set.

```
> all.errors <- data.table(errors$model.errors)
> all.errors[, set := ifelse(chromosome=="11", "test", "train")]
```

|      | <pre>profile.id</pre> | ${\tt chromosome}$ | ${\tt min.lambda}$ | $\verb max.lambda $ | ${\tt min.log.lambda}$  | <pre>max.log.lambda</pre> |
|------|-----------------------|--------------------|--------------------|---------------------|-------------------------|---------------------------|
| 1:   | 1                     | 1                  | 0.00000000         | 0.07175223          | -Inf                    | -2.6345364                |
| 2:   | 1                     | 1                  | 0.07175223         | 0.12470984          | -2.6345364              | -2.0817655                |
| 3:   | 1                     | 1                  | 0.12470984         | 0.12800334          | -2.0817655              | -2.0556989                |
| 4:   | 1                     | 1                  | 0.12800334         | 0.20965938          | -2.0556989              | -1.5622711                |
| 5:   | 1                     | 1                  | 0.20965938         | 0.27946950          | -1.5622711              | -1.2748621                |
|      |                       |                    |                    |                     |                         |                           |
| 252: | 11                    | 17                 | 0.34732889         | 0.46128451          | -1.0574831              | -0.7737403                |
| 253: | 11                    | 17                 | 0.46128451         | 0.61549695          | -0.7737403              | -0.4853253                |
| 254: | 11                    | 17                 | 0.61549695         | 0.72402388          | -0.4853253              | -0.3229309                |
| 255: | 11                    | 17                 | 0.72402388         | 2.82055129          | -0.3229309              | 1.0369324                 |
| 256: | 11                    | 17                 | 2.82055129         | Inf                 | 1.0369324               | Inf                       |
|      | ${\tt n.segments}$    | loss               | lik p              | possible.fp         | ${\tt fp\ possible.fn}$ | fn labels                 |

```
1:
              10
                  3.361407 -500.23714
                                                    1
                                                       1
                                                                     0
                                                                        0
                                                                                1
  2:
                  3.433159 -495.29382
                                                    1
                                                                     0
                                                                         0
                                                                                1
               9
                                                       1
  3:
               8
                  3.557869 -486.70201
                                                    1
                                                       1
                                                                     0
                                                                         0
                                                                                1
  4:
                  3.813876 -469.06458
                                                    1
                                                       1
                                                                     0
                                                                        0
                                                                                1
               6
  5:
                  4.023535 -454.62022
                                                    1
                                                        1
                                                                     0
                                                                         0
                                                                                1
252:
                   5.021862
                              -63.02933
                                                    1
                                                        1
                                                                     0
                                                                         0
                                                                                1
                                                                     0
                                                                         0
253:
                  5.483146
                              -56.10827
                                                    1
                                                        1
                                                                                1
               6
254:
                  6.098643
                              -46.87343
                                                    1
                                                       1
                                                                     0
                                                                         0
                                                                                1
                              -14.28391
                                                                     0
                                                                        0
255:
               2
                  8.270715
                                                    1
                                                       1
                                                                                1
256:
               1 11.091266
                               28.03533
                                                    1
                                                       0
                                                                         0
                                                                                1
     errors
               set
  1:
           1 train
  2:
           1 train
  3:
           1 train
  4:
           1 train
  5:
           1 train
252:
           1 train
253:
           1 train
254:
           1 train
255:
           1 train
256:
           0 train
```

To train a penalty learning model, we compute target intervals of log(penalty) values that achieve the minimal number of incorrect labels (for each problem independently). This is the "output" in the machine learning problem, which is a (n x 2) matrix that we compute using the penaltyLearning::targetIntervals function.

```
> target.dt <- penaltyLearning::targetIntervals(
    all.errors[set=="train", ],
    c("profile.id", "chromosome"))
> target.mat <- target.dt[, cbind(min.log.lambda, max.log.lambda)]</pre>
> rownames(target.mat) <- target.dt[, paste(profile.id, chromosome)]</pre>
> print(head(target.mat))
     min.log.lambda max.log.lambda
1 1
          0.1957271
1 2
         -1.1087554
                                 Inf
1 3
         -1.0006564
                                 Inf
1 4
         -0.9951232
                                 Inf
1 17
         -1.8106532
                                 Inf
4 1
                -Inf
                           3.265185
```

Note that the first column is the lower limit (possibly -Inf) of acceptable log(penalty) values, and the second column is the upper limit (possibly Inf).

### 2.5 Feature matrix

Then we compute a feature matrix (problems x features), which is the "input" in the machine learning problem. Here we compute a simple matrix with just 2 columns/features (log.var =  $\log \sigma_i$  is a variance estimate, and  $\log data = \log d_i$  is  $\log data = \log data$  points to segment).

```
> feature.dt <- profiles[, list(
+ log.data=log(.N),
+ log.var=log(median(abs(diff(logratio))))
+ ), by=list(profile.id, chromosome)]
> all.feature.mat <- feature.dt[, cbind(log.data, log.var)]
> rownames(all.feature.mat) <- feature.dt[, paste(profile.id, chromosome)]</pre>
```

You can also use penaltyLearning::featureMatrix for computing a larger feature matrix in real data sets where you aren't sure what features are relevant.

## 2.6 Learn a penalty function

Then we use penalty-Learning::IntervalRegressionUnregularized to learn a penalty function for this small data set (see Hocking et al ICML'13 for details). Because we only computed two features  $\log d_i$ ,  $\log \sigma_i$ , the function that we learn here is  $f(x_i) = \beta + w_1 \log d_i + w_2 \log \sigma_i$  (we learn the intercept/bias  $\beta$  and feature weights  $w_1, w_2$ ). Remember that the prediction function models  $\log(\text{penalty})$  values  $f(x_i) = \log \lambda_i$ , so the penalty function we learn here is  $\lambda_i = e^{\beta} d_i^{w_1} \sigma_i^{w_2}$ .

For data sets with more labels and features, penaltyLearning:;IntervalRegressionCV would be preferable, since it also performs variable selection using L1-regularization.

### 2.7 Prediction

Then we use the model to predict log(penalty) values for each segmentation problem (even those in the test set).

> feature.dt[, pred.log.lambda := predict(fit, all.feature.mat)]

|      | <pre>profile.id</pre> | ${\tt chromosome}$ | log.data | log.var   | <pre>pred.log.lambda</pre> |
|------|-----------------------|--------------------|----------|-----------|----------------------------|
| 1:   | 8                     | 1                  | 6.013715 | -2.877904 | 0.41081127                 |
| 2:   | 8                     | 2                  | 5.375278 | -2.928180 | -0.10122396                |
| 3:   | 8                     | 3                  | 5.081404 | -2.917816 | -0.27701599                |
| 4:   | 8                     | 4                  | 4.882802 | -3.045320 | -0.63627765                |
| 5:   | 8                     | 5                  | 5.164786 | -2.833040 | -0.07032902                |
|      |                       |                    |          |           |                            |
| 140: | 10                    | 20                 | 4.828314 | -2.882404 | -0.38106243                |
| 141: | 10                    | 21                 | 4.110874 | -2.239610 | 0.29365581                 |
| 142: | 10                    | 22                 | 4.543295 | -2.847312 | -0.50679475                |
| 143: | 10                    | X                  | 5.231109 | -2.847312 | -0.05198766                |
| 144: | 10                    | Y                  | 3.295837 | -2.900422 | -1.42660240                |

Note that this is the predict method for the IntervalRegression class.

#### 2.8 Evaluation

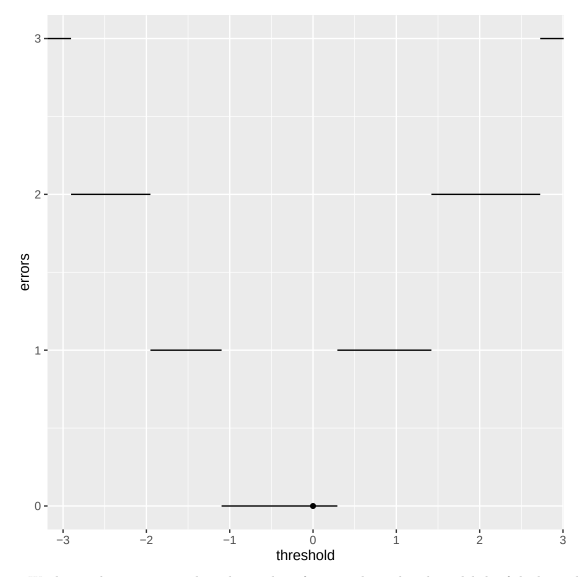
We can use penaltyLearning::ROChange to compute test ROC curves, and Area Under the Curve (AUC), which in this case is 1, indicating a function which can perfectly discriminate between positive and negative labels in the test set.

```
> test.pred <- feature.dt[chromosome=="11",]</pre>
> roc <- penaltyLearning::ROChange(</pre>
    all.errors, test.pred, c("profile.id", "chromosome"))
> pred.thresh <- roc$thresholds[threshold=="predicted"]</pre>
> gg.roc <- ggplot()+</pre>
    geom_path(aes(
      FPR, TPR),
      data=roc$roc)+
    geom_point(aes(
      FPR, TPR),
      data=pred.thresh)
> print(gg.roc)
> print(pred.thresh)
   threshold labels possible.fp possible.fn min.thresh max.thresh fp fn errors
                                             3 -1.097374 0.2923082 0 0
1: predicted
   FPR tp TPR error.percent
     0 3
            1
   1.00
   0.75 -
A 0.50 -
   0.25 -
   0.00 -
                                                               0.75
                            0.25
                                             0.50
                                                                                1.00
          0.00
```

Using the ROC output you can also easily plot the number of incorrect labels as a function of threshold  $\tau \in \mathbb{R}$  added to the predicted log(penalty) function  $f(x_i) + \tau$ ,

**FPR** 

```
> gg.thresh <- ggplot()+
+ geom_segment(aes(
+ min.thresh, errors,
+ xend=max.thresh, yend=errors),
+ data=roc$roc)+
+ geom_point(aes(
+ 0, errors),
+ data=pred.thresh)+
+ xlab("threshold")
> print(gg.thresh)
```



We draw a dot at  $\tau = 0$  to show the number of incorrectly predicted test labels of the learned prediction function (0 in this case).

## 2.9 Visualizing predictions

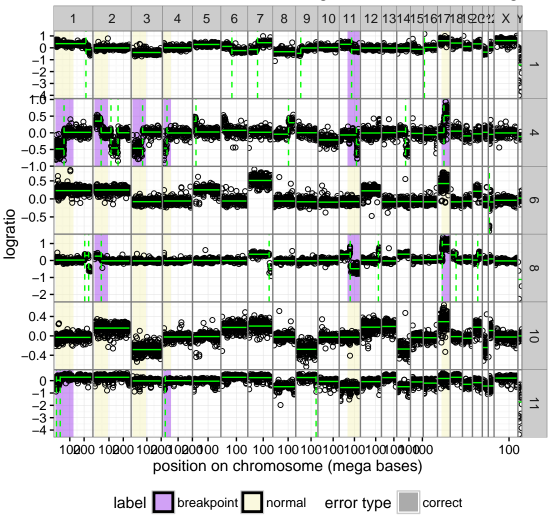
Finally we use a non-equi join of feature.dt (which contains the predicted penalty values) with selection (which contains the number of segments for each penalty value). Then we plot the predicted models along with the data and labels.

```
> pred.models <- feature.dt[selection, nomatch=OL, on=list(
+ profile.id, chromosome,</pre>
```

<sup>+</sup> pred.log.lambda < max.log.lambda,

```
pred.log.lambda > min.log.lambda)]
> pred.segs <- segs[pred.models, on=list(profile.id, chromosome, n.segments)]</pre>
> pred.changes <- pred.segs[1 < start, ]</pre>
> pred.labels <- errors$label.errors[pred.models, nomatch=OL, on=list(
    profile.id, chromosome, n.segments)]
> breakpoint.colors <- c(</pre>
    "breakpoint"="#a445ee",
    "normal"="#f6f4bf")
> viz.learned <- ggplot()+
    ggtitle("data + labels + learned model segment means and changes")+
    theme_bw()+
    theme(
      legend.position="bottom",
      legend.box="horizontal",
      panel.margin=grid::unit(0, "lines"))+
    facet_grid(profile.id ~ chromosome, scales="free", space="free_x")+
    penaltyLearning::geom_tallrect(aes(
      xmin=min/1e6, xmax=max/1e6, fill=annotation, linetype=status),
      data=pred.labels)+
    scale_linetype_manual("error type",
                          values=c(correct=0,
+
                                    "false negative"=3,
+
                                    "false positive"=1))+
    scale_fill_manual("label", values=breakpoint.colors)+
    geom_point(aes(position/1e6, logratio),
               data=profiles,
               shape=1)+
    scale_x_continuous(
      "position on chromosome (mega bases)",
      breaks=c(100, 200))+
    geom_segment(aes(chromStart/1e6, mean, xend=chromEnd/1e6, yend=mean),
+
                 data=pred.segs,
                 color="green")+
    geom_vline(aes(xintercept=chromStart/1e6),
               data=pred.changes,
               color="green",
               linetype="dashed")
> print(viz.learned)
```





The plot above can be interpreted in the following manner

- black dots are the noisy data points in which we have attempted to detect changepoints.
- each panel is a separate data sequence (a separate multiple changepoint detection problem).
- horizontal solid green lines are the learned segment means.
- vertical dashed green lines are the predicted changepoint positions.
- colored rectangles are the labels. yellow normal labels should have no changepoints, and purple breakpoint labels should have at least one changepoint.

It is clear that in these data, all of the predicted models are consistent with the labels. You can also see that the model makes reasonable predictions for the unlabeled chromosomes.

### 2.10 Comparison with BIC

The Bayesian Information Criterion (BIC) corresponds to using the penalty function  $f(x_i) = \log \log d_i$  which is unsupervised (nothing to learn). Below we compute the ROC curve for the same test set (chromosome 11) as we used for the learned model.

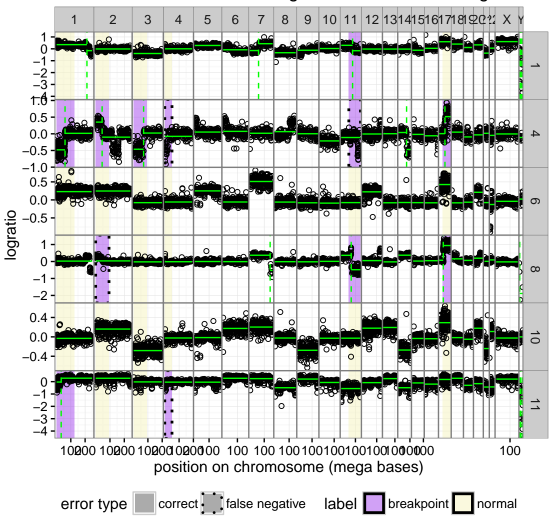
```
> bic.dt <- profiles[, list(</pre>
```

<sup>+</sup> pred.log.lambda=log(log(.N))

It is clear that the BIC suffers from a false negative in chromosome 11 (a purple breakpoint region with no predicted changepoint). We visualize the model along with the data and labels in the plot below,

```
> bic.models <- bic.dt[selection, nomatch=OL, on=list(</pre>
    profile.id, chromosome,
   pred.log.lambda < max.log.lambda,</pre>
   pred.log.lambda > min.log.lambda)]
> bic.segs <- segs[bic.models, on=list(profile.id, chromosome, n.segments)]</pre>
> bic.changes <- bic.segs[1 < start, ]</pre>
> bic.labels <- errors$label.errors[bic.models, nomatch=0L, on=list(</pre>
    profile.id, chromosome, n.segments)]
> viz.bic <- ggplot()+</pre>
    ggtitle("data + labels + BIC model segment means and changes")+
    theme_bw()+
    theme(
      legend.position="bottom",
      legend.box="horizontal",
      panel.margin=grid::unit(0, "lines"))+
    facet_grid(profile.id ~ chromosome, scales="free", space="free_x")+
    penaltyLearning::geom_tallrect(aes(
      xmin=min/1e6, xmax=max/1e6, fill=annotation, linetype=status),
      data=bic.labels)+
+
    scale_linetype_manual("error type",
                           values=c(correct=0,
                                     "false negative"=3,
                                     "false positive"=1))+
+
    scale_fill_manual("label", values=breakpoint.colors)+
+
+
    geom_point(aes(position/1e6, logratio),
               data=profiles,
+
               shape=1)+
    scale_x_continuous(
+
      "position on chromosome (mega bases)",
+
      breaks=c(100, 200))+
    geom_segment(aes(chromStart/1e6, mean, xend=chromEnd/1e6, yend=mean),
                 data=bic.segs,
                  color="green")+
    geom_vline(aes(xintercept=chromStart/1e6),
               data=bic.changes,
+
               color="green",
               linetype="dashed")
> print(viz.bic)
```

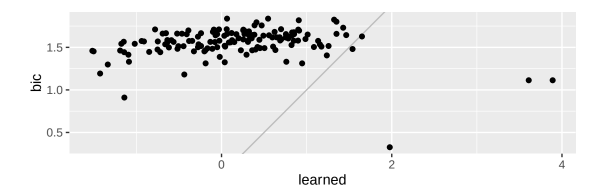




It is clear from the plot above that there are many false negatives – purple breakpoint labels in which there should be at least one change, but the BIC model predicts none.

Finally we can make the scatterplot that compare the predicted penalty values of the learned model and the BIC:

```
> scatter.dt <- data.table(
+ bic=bic.dt$pred.log.lambda,
+ learned=feature.dt$pred.log.lambda)
> gg.scatter <- ggplot()+
+ geom_abline(
+ slope=1, intercept=0, color="grey")+
+ geom_point(aes(
+ learned, bic),
+ data=scatter.dt)+
+ coord_equal()
> print(gg.scatter)
```



It is clear from the scatterplot that the learned model always predicts larger log(penalty) values than the BIC.

# 3 Conclusion

We have shown an application of the penaltyLearning package to the neuroblastoma data set. We showed how to learn a penalty function based on set of data sequences which have labels that indicate presence or absence of changes in specific regions. We showed that the learned penalty function is more accurate than an unsupervised penalty function (BIC).