

HowTo Use the Bioconductor `edd` package

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1 Introduction

edd is a package that assists with one aspect of exploratory data analysis for microarrays. The basic question addressed in *edd* is the variety of shapes of gene-specific distributions of expression in collections of microarrays. Use of the package is most sensible when there are numerous arrays obtained under the same experimental condition or for a given clinical condition. The key idea is that marginal gene-specific distributions may have a relatively number of different qualitative shapes, some of which may be of considerable substantive interest (e.g., multimodal shapes), and some of which may be of methodologic importance (e.g., when one group of subjects has a skewed distribution for a gene, and another has a symmetric distribution for the same gene, use of a log transform is counterindicated).

In this brief HOWTO, we illustrate directly the use of the *edd* package. We will investigate the diversity of distributions in the two main groups of Golub's leukemia dataset.

2 Important caveat

The `edd` function will transform all gene-specific expression distributions to have common location and scale. This process can make noise have the appearance of signal. Before using `edd`, remove all genes that have small variability. See the next section for an example of this filtering process.

3 Distributional shapes in Golub's data

First we attach the necessary libraries and data frames. `edd` will require the `golubEsets` library.

```
> library(edd)
> library(golubEsets)
> library(xtable)
> data(Golub_Merge)
```

3.1 Filtering out genes with low variation

Next we filter the Golub data to require reasonable dispersion (confine attention to upper half sample defined by size of MAD) and reasonable expression (confine attention to genes with minimum expression level 300).

```
> madvec <- apply(exprs(Golub_Merge), 1, mad)
> minvec <- apply(exprs(Golub_Merge), 1, min)
> keep <- (madvec > median(madvec)) & (minvec > 300)
> gmfilt <- Golub_Merge[keep == TRUE, ]
```

3.2 Forming stratum-specific ExpressionSets

Finally we split the dataset into the ALL and AML samples:

```
> ALL <- gmfilt$ALL.AML == "ALL"
> gall <- gmfilt[, ALL == TRUE]
> gaml <- gmfilt[, ALL == FALSE]
> show(gall)
```

```
ExpressionSet (storageMode: lockedEnvironment)
assayData: 540 features, 47 samples
  element names: exprs
phenoData
  sampleNames: 39, 40, ..., 27 (47 total)
  varLabels and varMetadata description:
```

```

Samples: Sample index
ALL.AML: Factor, indicating ALL or AML
...: ...
Source: Source of sample
(11 total)
featureData
  featureNames: hum_alu_at, AFX-HUMGAPDH/M33197_3_at, ..., X03068_f_at (540 total)
  fvarLabels and fvarMetadata description: none
experimentData: use 'experimentData(object)'
  pubMedIds: 10521349
Annotation: hu6800

```

3.3 Running edd

We will apply edd using an nnet classifier with the default reference catalog. See the edd-Details vignette for information about the reference catalog.

```

> set.seed(12345)
> alldists <- edd(gall, meth = "nnet", size = 10, decay = 0.2)

```

```

# weights: 579
initial value 2078.664028
iter 10 value 1087.941692
iter 20 value 727.152339
iter 30 value 566.315287
iter 40 value 474.576473
iter 50 value 432.376177
iter 60 value 393.533019
iter 70 value 360.720544
iter 80 value 347.231530
iter 90 value 343.157228
iter 100 value 340.265424
final value 340.265424
stopped after 100 iterations

```

```

> amlldists <- edd(gaml, meth = "nnet", size = 10, decay = 0.2)

```

```

# weights: 359
initial value 2194.061594
iter 10 value 1280.052370
iter 20 value 968.639334
iter 30 value 855.773083
iter 40 value 764.281046

```

```

iter 50 value 707.744888
iter 60 value 688.798967
iter 70 value 674.180641
iter 80 value 661.971814
iter 90 value 654.615602
iter 100 value 649.977795
final value 649.977795
stopped after 100 iterations

```

An example of the results is given by the classification calls for the first 5 genes in the filtered ExpressionSet:

```

          hum_alu_at AFX-HUMGAPDH/M33197_3_at AFX-HSAC07/X00351_5_at
".75N(0,1)+.25N(4,1)"          "t(3)"          "t(3)"
AFHX-HSAC07/X00351_3_at      AFX-M27830_M_at
          "t(3)"          "logN(0,1)"

```

We can use edd with other classification methods.

```

> set.seed(123)
> alldistsKNN <- edd(gall, meth = "knn", k = 1, l = 0)
> alldistsTEST <- edd(gall, meth = "test", thresh = 0.3)

```

The agreement between nnet and knn procedures is not exact. See table 1. Choice between these methods and selection of tuning parameters is context-dependent.

```

> cap <- "Comparison of distribution shape classification by nnet (rows) and by knn (
> print(try(xtable(latEDtable(table(alldists, alldistsKNN), reorder = greo),
+   digits = rep(0, length(table(alldists)) + 1), caption = cap,
+   label = "conc1"))))

```

The test procedure is the only one at present that allows an outcome of 'doubt'.

```

> print(table(alldistsTEST))

```

```

alldistsTEST
.25N(0,1)+.75N(4,1) .75N(0,1)+.25N(4,1)          B(2,8)          B(8,2)
                9                91                169                26
                logN(0,1)          N(0,1)          outlier          t(3)
                40                68                4                104
                U(0,1)          X^2(1)
                26                3

```

	Φ	t_3	$LN_{\{0,1\}}$	$\chi^2_{2,1}$	$\beta_{\{8,2\}}$	$U_{\{0,1\}}$	$\beta_{\{2,8\}}$	$\frac{3}{4}\Phi + \frac{1}{4}\Phi_{\{4,1\}}$	$\frac{1}{4}\Phi + \frac{3}{4}\Phi_{\{4,1\}}$
Φ	49	6	0	0	0	0	0	0	0
t_3	25	54	11	0	0	0	0	0	0
$LN_{\{0,1\}}$	0	3	65	12	0	0	0	0	0
$\chi^2_{2,1}$	0	0	1	1	0	0	0	0	0
$\beta_{\{8,2\}}$	1	0	0	0	0	0	0	0	0
$U_{\{0,1\}}$	0	0	0	0	0	0	0	0	0
$\beta_{\{2,8\}}$	24	0	8	0	0	0	0	0	0
$\frac{3}{4}\Phi + \frac{1}{4}\Phi_{\{4,1\}}$	0	0	6	3	0	0	0	0	0
$\frac{1}{4}\Phi + \frac{3}{4}\Phi_{\{4,1\}}$	0	0	0	0	0	0	0	0	0

Table 1: Comparison of distribution shape classification by nnet (rows) and by knn (columns) methods in edd.

3.4 Assessing the results

We can assess the relative frequencies of the different shapes in the ALL samples with a table, see Table 2.

```
> cap <- "Frequencies of distributional shapes in filtered ALL data."
> print(xtable(latEDtable(table(alldists), reorder = greo), digits = rep(0,
+   length(table(alldists)) + 1), caption = cap, label = "marg1"))
```

Φ	t_3	$LN_{\{0,1\}}$	$\chi^2_{2,1}$	$\beta_{\{8,2\}}$	$U_{\{0,1\}}$	$\beta_{\{2,8\}}$	$\frac{3}{4}\Phi + \frac{1}{4}\Phi_{\{4,1\}}$	$\frac{1}{4}\Phi + \frac{3}{4}\Phi_{\{4,1\}}$
69	140	89	2	12	3	167	0	0

Table 2: Frequencies of distributional shapes in filtered ALL data.

We can use barplots also; see Figure 1.

Discordance between distributional shapes in gene expression for the AML and ALL groups can be assessed using the cross-classification, see Table 3.

```
> cap <- "Rows are gene-specific distribution shapes for ALL, columns for AML, and ce
> print(xtable(latEDtable(table(alldists, amldists), reord = greo),
+   cap = cap, label = "disco1"))
```

Let's see what these discordances mean. To begin, let's get some indices for genes with bimodally shaped expression distribution for ALL, but approximately gaussian expression distribution for AML:

```
> print((1:540)[alldists == ".75N(0,1)+.25N(4,1)" & amldists ==
+   "N(0,1)"][1:5])
```

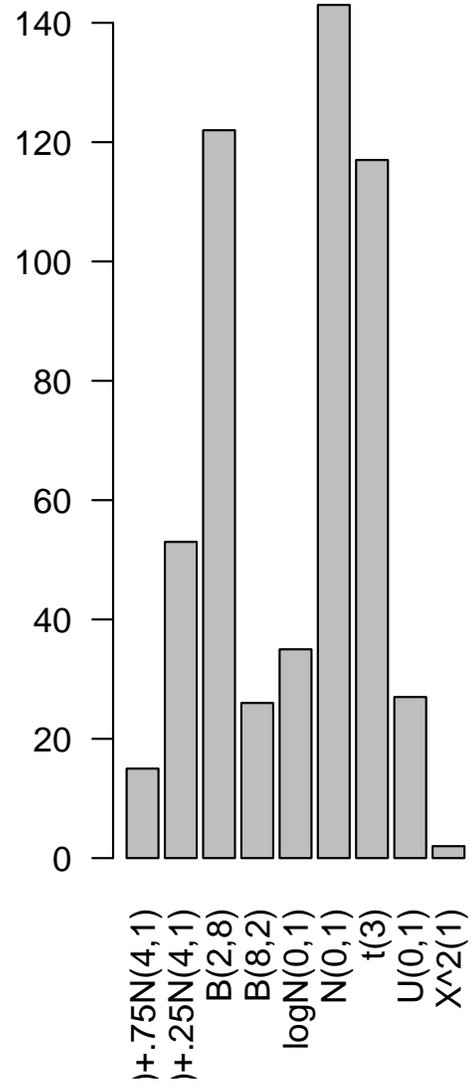
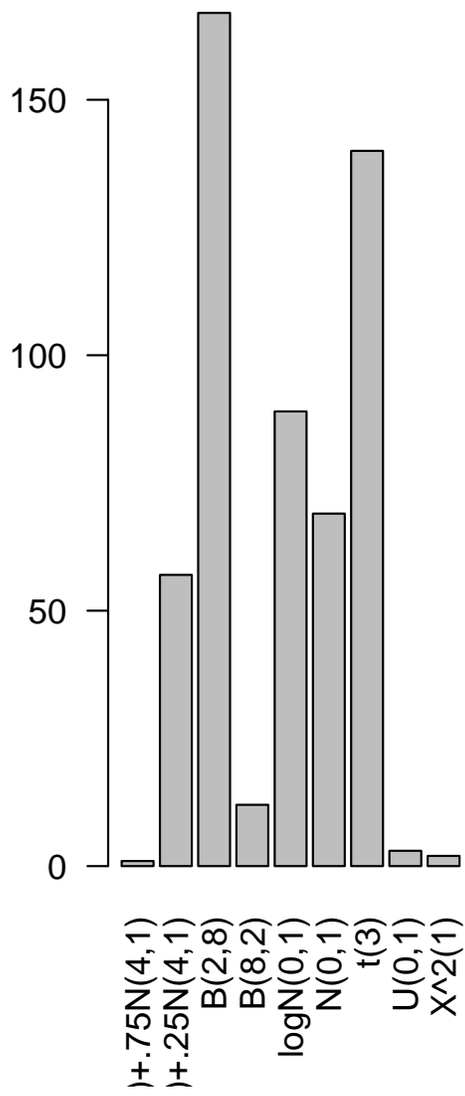


Figure 1: Compositions of distributional shapes within strata.

	Φ	t_3	$LN_{\{0,1\}}$	$\chi^2_{.1}$	$\beta_{\{8,2\}}$
Φ	25	16	2	0	0
t_3	33	42	8	0	0
$LN_{\{0,1\}}$	20	14	11	1	1
$\chi^2_{.1}$	0	1	0	0	0
$\beta_{\{8,2\}}$	2	3	0	0	0
$U_{\{0,1\}}$	0	0	0	0	0
$\beta_{\{2,8\}}$	46	25	10	1	1
$\frac{3}{4}\Phi + \frac{1}{4}\Phi_{\{4,1\}}$	17	15	4	0	0
$\frac{1}{4}\Phi + \frac{3}{4}\Phi_{\{4,1\}}$	0	1	0	0	0

Table 3: Rows are gene-specific distribution shapes for ALL, columns for AML, and cell entries are counts of genes.

[1] 65 78 135 141 191

We consider the gene with probe D87953_at. The top left panel gives the model (solid density trace) and a kernel density estimate applied to the expression levels among ALL patients, and the top right is the corresponding histogram.

While the specific mixture model used as reference is not a perfect fit to the ALL data, the neural net classifier was sensitive to the bimodality. The Gaussian model does not seem particularly appropriate for the AML data, but was the closest match in the reference catalog.

4 Extending the reference catalog

The reference catalog supplied with edd has components

```
> names(eddDistList)
```

```
[1] "N01" "T3" "LN01" "CS1" "B82" "U01" "B28" "MIXN1" "MIXN2"
```

There is nothing sacred about this set. Let's consider its scope (we'll look at 8 of nine reference distributions):

From the example above we see that it might be useful to have a mixture of Gaussians with modes separated by 6SD. To add such a model we construct an instance of the eddDist class:

```
> MIXN3 <- new("eddDist", stub = "mixnorm", parms = c(p1 = 0.75,
+   m1 = 0, s1 = 1, m2 = 6, s2 = 1), median = 0.43, mad = 1.55,
+   tag = ".75N(0,1)+.25N(6,1)", plotlim = c(-3, 11), latexTag = "$\\frac{3}{4}\\Phi + \\frac{1}{4}\\Phi_{\\{4,1\\}}$")
> eddDistList[["MIXN3"]] <- MIXN3
> set.seed(12345)
> alldists2 <- edd(gall, meth = "nnet", size = 10, decay = 0.2)
```

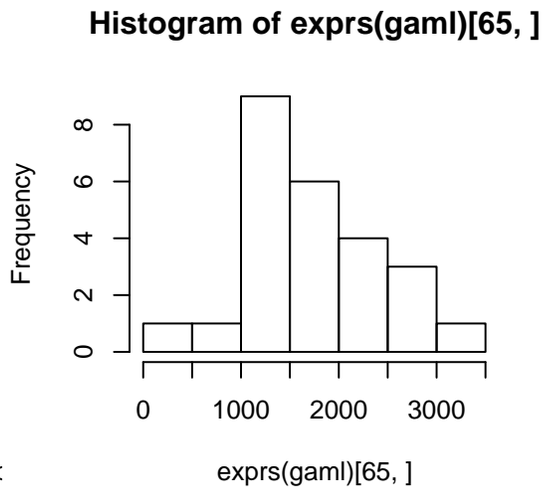
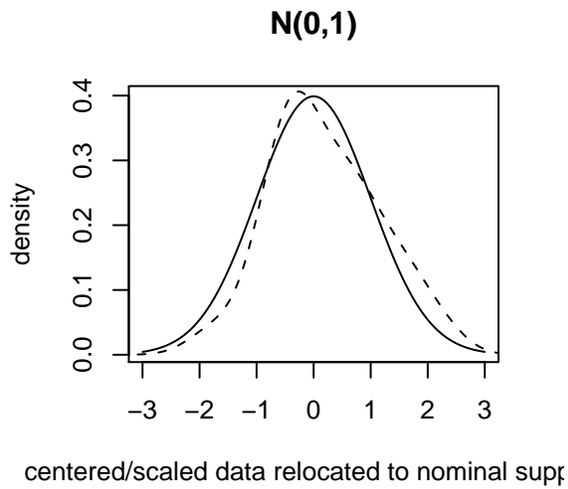
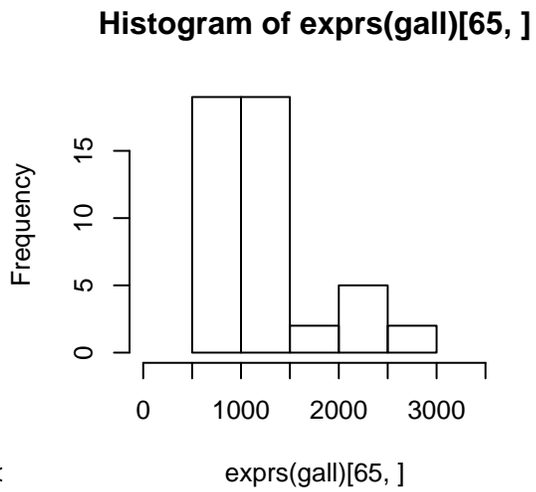
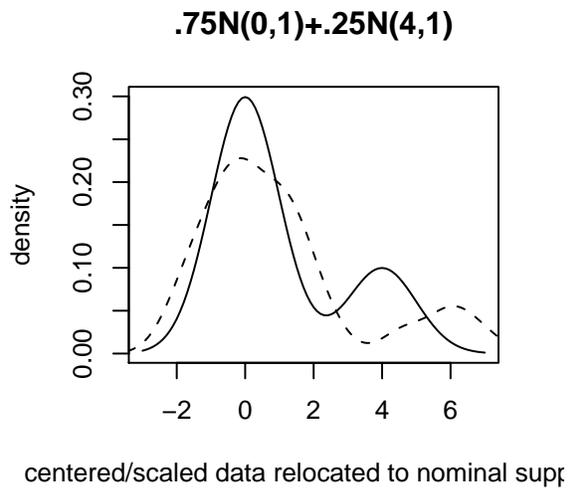


Figure 2: Two models for D87953_at in ALL and AML patients.

```
> par(mfrow = c(4, 2))
> for (i in 1:8) plotED(eddDistList[[i]])
```

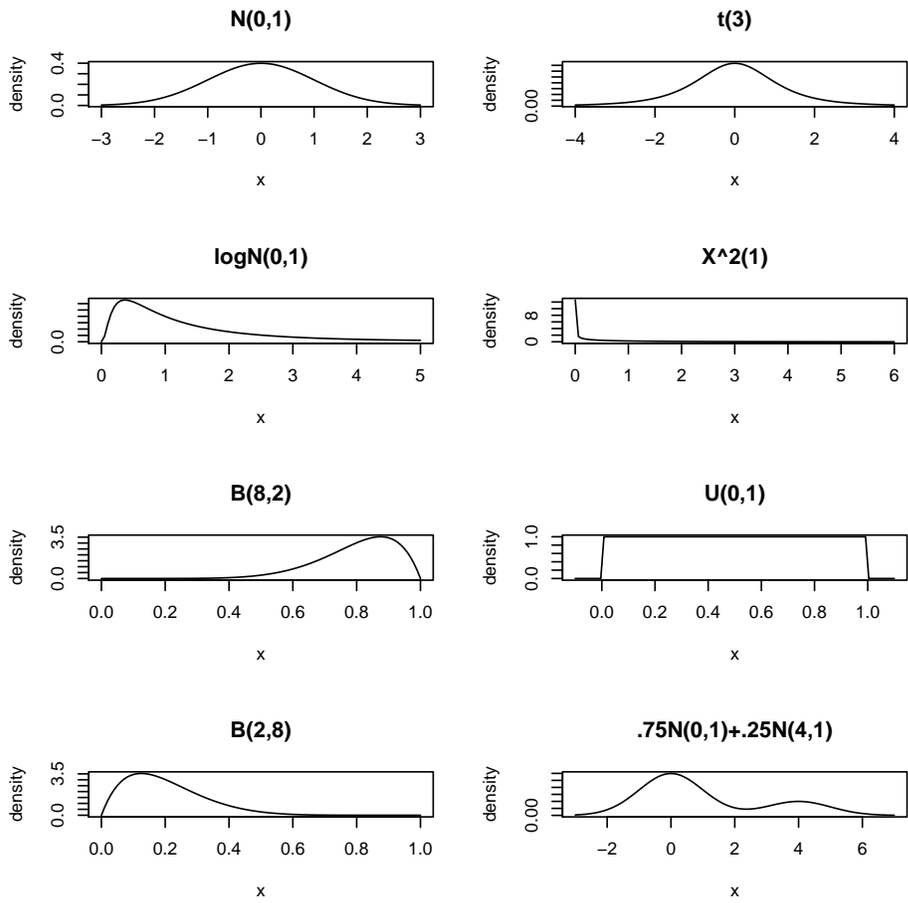


Figure 3: Eight of the reference distributions in the `eddDistList` supplied with `edd`.

```
# weights: 579
initial value 2078.664028
iter 10 value 1087.941692
iter 20 value 727.152339
iter 30 value 566.315287
iter 40 value 474.576473
iter 50 value 432.376177
iter 60 value 393.533019
iter 70 value 360.720544
iter 80 value 347.231530
iter 90 value 343.157228
iter 100 value 340.265424
final value 340.265424
stopped after 100 iterations
```

```
> print(alldists2[65])
```

```
[1] ".75N(0,1)+.25N(4,1)"
```

The symbol MIXN3 used to name the list element is arbitrary, as are the values of the tag and latexTag slots. But the user should choose meaningful values for those items. The new reference distribution is used for classification of probe D87953_at. The two fits for the different mixtures are shown in Figures 4, 5.

```
> plotED(MIXN3, data = exprs(gall)[65, ])
```

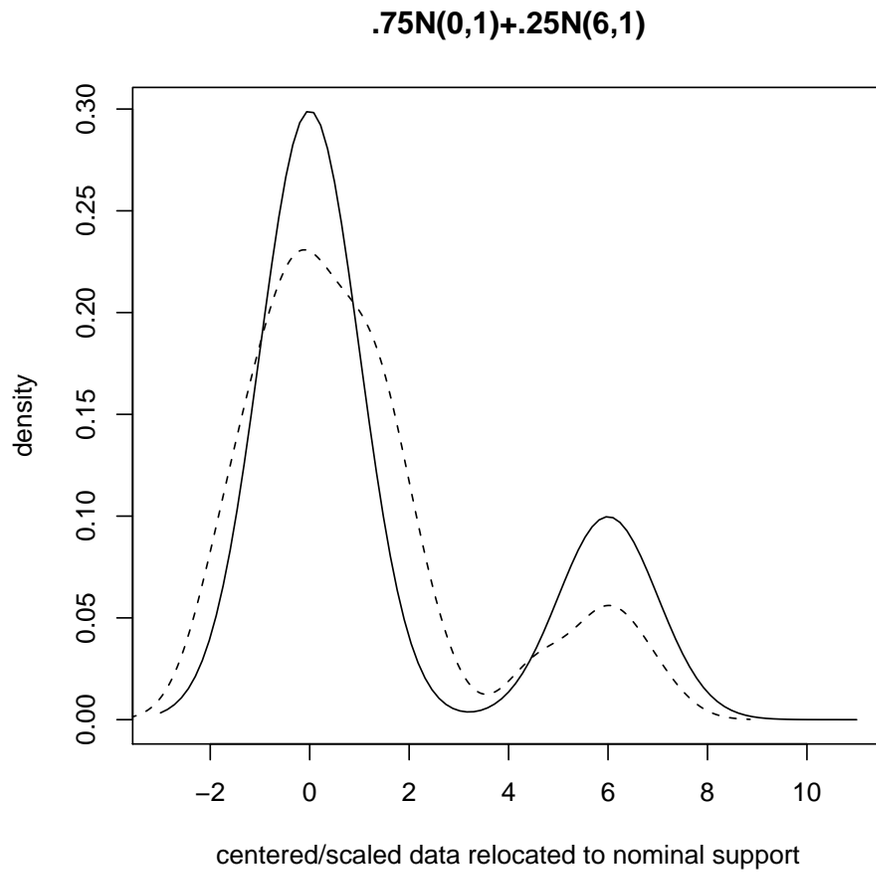


Figure 4: Reference catalog element: mixture with modes separated by 6SD. Superimposed is the kernel smooth of centered/scaled and then translated data for D87953_at.

```
> plotED(MIXN1, data = exprs(gall)[65, ])
```

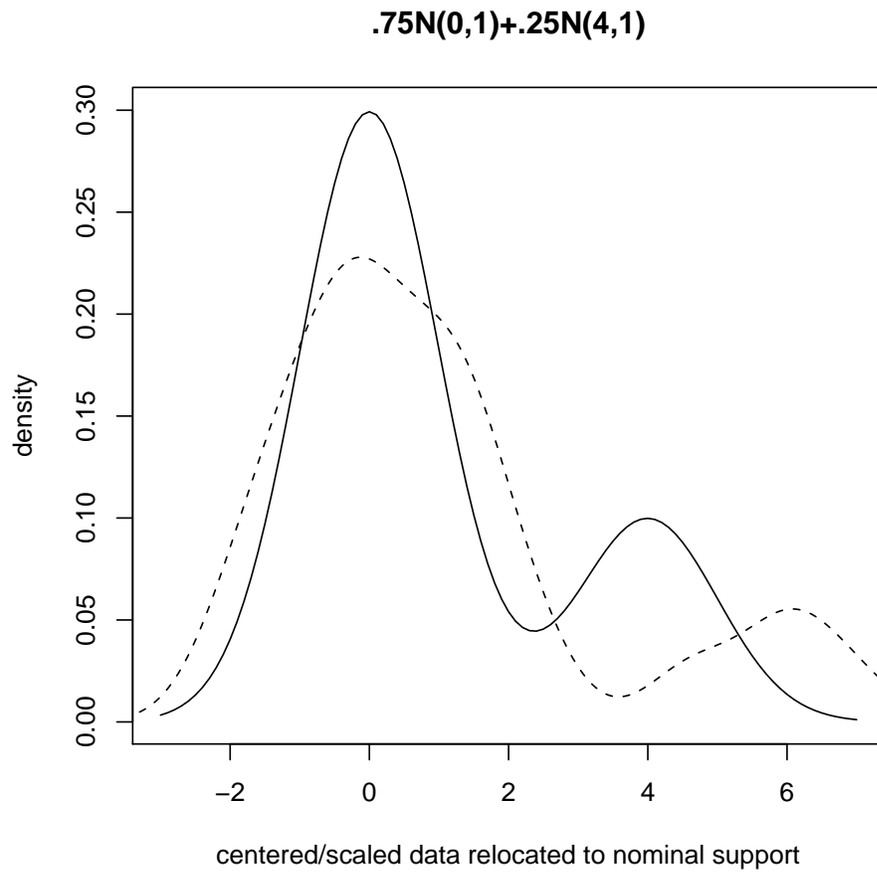


Figure 5: Reference catalog element: mixture with modes separated by 3SD. Superimposed is the kernel smooth of centered/scaled and then translated data for D87953_at.