# Fitting a bivariate normal distribution to a 2D scatterplot 

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## 1 Overview

Using FACS (fluorescence-activated cell sorter) one can measure certain properties of each individual cell in a population of cells. Examples for these properties:

- Forward light scatter (FSC): this measures a cell's size
- Sideward light scatter (SSC): this measures a cell's granularity
- Several fluorescence channels (typically 3 to 4 ) that measure the abundance of fluorophores, which may be bound to specific antibodies for surface or intracellular markers, or be encoded by a GFP-tagged transcript.

First, we load example data from a FACS analysis that was performed by Mamatha Sauermann at the German Cancer Research Center in Heidelberg.

```
> library(prada)
> sampdat <- readFCS(system.file("extdata", "fas-Bcl2-plate323-04-04.A01",
+ package = "prada"))
> fdat <- exprs(sampdat)
```

The scatterplot of FSC vs SSC is often used for quality control. It is shown in Fig. 1.

```
> plot(fdat[, "FSC-H"], fdat[, "SSC-H"], pch = 20, col = "#303030",
+ xlab = "FSC", ylab = "SSC", main = "Scatter plot FSC vs SSC")
```

The cell population is often contaminated by cell debris or conjugates. These can be identified by their size: they are either much smaller or much larger than the main population, or they have an unusual degree of granularity. Segmentation is often performed manually by looking at the FSC-SCC scatterplot.

Here we describe an automated algorithm for this task.

## 2 Fitting

The package prada provides the functions fitNorm2 and plotNorm2. We assume that the shape of the main population in the FSC vs SSC plot can be approximated by a normal distribution. The function fitNorm2 fits a bivariate normal distribution into the data (by robust estimation


Figure 1: Scatter plot FACS data: FSC vs SSC.
of its covariance matrix). Contours of equal probability of a bivariate normal are ellipses. We select those cells as being part of the main population that lie within such an ellipse. Its size is controled by the parameter scalefac. The function returns a list.

```
> nfit <- fitNorm2(fdat[, "FSC-H"], fdat[, "SSC-H"], scalefac = 2)
```

We can plot this with the function plotNorm2 (see Fig 2). It shows the ellipse, and the set of discarded points is marked by a red dot. Also the center of the normal distribution is marked by the red cross.

```
> plotNorm2(nfit, selection = TRUE, ellipse = TRUE)
> nfit3 <- fitNorm2(fdat[, "FSC-H"], fdat[, "SSC-H"], scalefac = 3)
> plotNorm2(nfit3, selection = TRUE, ellipse = TRUE)
```

To select the cells from within the ellipse, the list item nfit\$sel is a logical vector with the same length as the number of data points.

```
> cleanfdat <- fdat[nfit$sel, ]
```

Fig. 3 shows again a scatter plot of the two fluorescense channels FL1 and FL4 this time using the 'clean' data set cleanfdat.

```
> par(mfrow = c(1, 2))
> xlim <- range(fdat[, "FL1-H"])
> ylim <- range(fdat[, "FL4-H"])
> plot(fdat[, "FL1-H"], fdat[, "FL4-H"], pch = 20, col = "#303030",
```



Figure 2: Selection of the main population, using two different values of the parameter scalefac.

```
+ xlab = "FL1", ylab = "FL4", main = "all data", xlim = xlim,
+ ylim = ylim)
> plot(cleanfdat[, "FL1-H"], cleanfdat[, "FL4-H"], pch = 20, col = "#303030",
+ xlab = "FL1", ylab = "FL4", main = "clean data only", xlim = xlim,
+ ylim = ylim)
```


## 3 Scatterplots

If you think that scatterplots with thousands of points are hard to read and annoying to view in a PDF viewer, have a look at the function smoothScatter (see Fig. 4):

```
> require(geneplotter)
> smoothScatter(fdat[, c("FSC-H", "SSC-H")], nrpoints = 50)
```



Figure 3: Scatter plots of FL1 vs FL4.


Figure 4: Smooth scatter plots.

