Visualization of data from assays in microtitre plate or slide format

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March 31, 2012

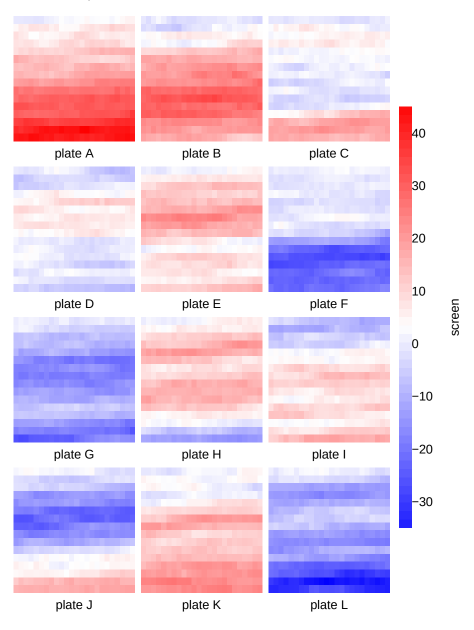
The splots package provides a single function, plotScreen, for visualising data in microtitre plate or slide format. Here we show a short demo of its parameters. First, we generate example data:

- > library("splots")
 > screen = lapply(1:12, function(i) cumsum(rnorm(384)))
- > names(screen) = paste("plate", LETTERS[seq(along=screen)])

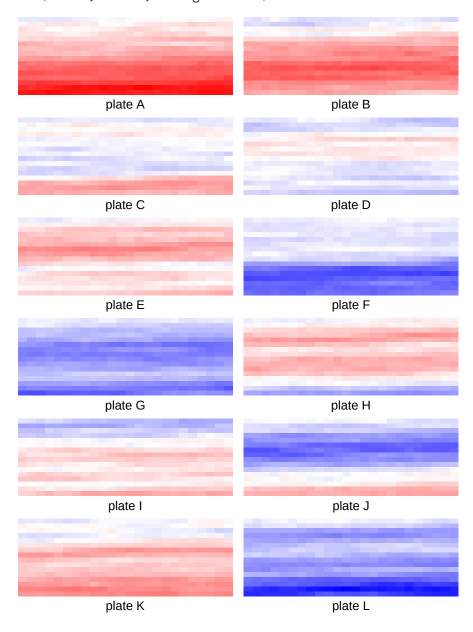
- > toLatex(sessionInfo())
 - R version 2.15.0 RC (2012-03-22 r58802), i386-apple-darwin9.8.0
 - Locale: C
 - Base packages: base, datasets, grDevices, graphics, methods, stats, utils
 - Other packages: splots 1.22.0
 - Loaded via a namespace (and not attached): RColorBrewer 1.0-5, grid 2.15.0, tools 2.15.0

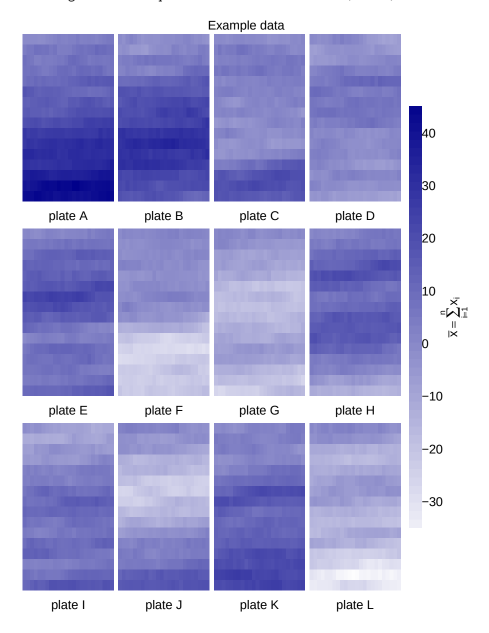
 Table 1: The output of sessionInfo on the build system after running this vignette.

> plotScreen(screen, ncol=3)



> plotScreen(screen, ncol=2, do.legend=FALSE)





- > for(i in seq(along=screen))
 + screen[[i]][sample(384, 5)] = NA
- > plotScreen(screen, ncol=4, do.names=FALSE,
- main="Example data", legend.label="Legend label")

