Supplemental

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Packages:

```
# install the development version of vivid:
#devtools::install_github("AlanInglis/vivid")

# Load relevant packages:
library("vivid") # for visualisations
library("ranger") # to create model
library("ggplot2") # for visualisations
```

Create Data:

Here we use Friedman's Benchmark problem 1^1 to create data from:

```
y = 10\sin(\pi x 1x2) + 20(x3 - 0.5)^2 + 10x4 + 5x5 + \epsilon
```

where, $x_n \sim U(0,1)$ and $\epsilon \sim N(0,1)$ We simulate 10 variables and 1000 observations and fit our model.

```
genFriedman <- function(noFeatures = 10,</pre>
                         noSamples = 100,
                         sigma = 1,
                         bins = NULL,
                         seed = NULL,
                         showTrueY = FALSE) {
  if (!is.null(seed)) {
    set.seed(seed)
  }
  # Set Values
  n <- noSamples # no of rows
  p <- noFeatures # no of variables
  e <- rnorm(n, sd = sigma)
  # Equation:
  # y = 10sin(x1x2) + 20(x3-0.5)^2 + 10x4 + 5x5 +
  xValues <- matrix(runif(n*p, 0, 1), nrow = n)
  colnames(xValues)<- paste0("x",1:p)</pre>
```

¹Friedman, Jerome H. (1991) Multivariate adaptive regression splines. The Annals of Statistics 19 (1), pages 1-67.

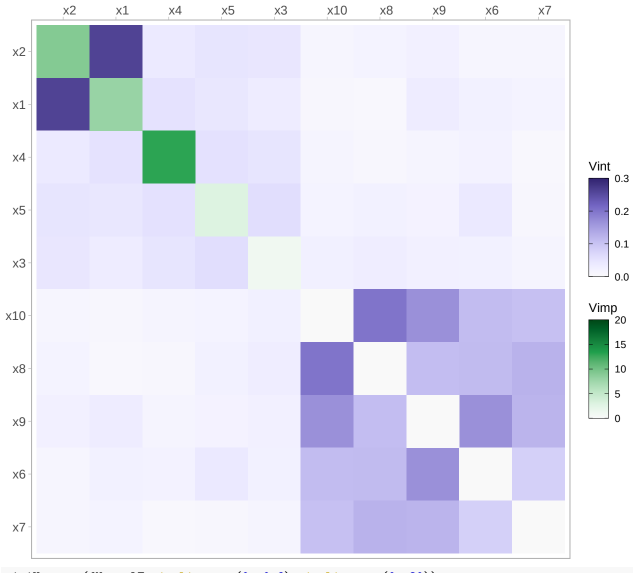
```
yTrue \leftarrow 10 * sin(pi * xValues[, 1] * xValues[, 2]) + 20 * (xValues[, 3] - 0.5)^2 + 10 * xValues[, 4]
  y <- yTrue + e
  if (showTrueY) {
    df <- data.frame(xValues, y, yTrue)</pre>
    df <- data.frame(xValues, y)</pre>
  # Function to bin a numberic vector
  bin <- function(x, bins) {</pre>
    x \leftarrow df$y
    quantiles <- quantile(x, probs = seq(from = 0, to = 1, length = bins + 1))
    bins <- cut(x, breaks = quantiles, label = FALSE, include.lowest = TRUE)</pre>
    as.factor(paste0("class", bins))
  }
  if (!is.null(bins)) {
    bins <- as.integer(bins)</pre>
    if (bins < 2) {</pre>
      stop("bins should be an integer greater than 1.", call. = FALSE)
    df$y <- bin(df$y, bins = bins)</pre>
  }
  df
}
# Data:
fData <- genFriedman(noFeatures = 10, noSamples = 1000, seed = 1701)
# Fit:
set.seed(1701)
fFit <- ranger(y ~ ., data = fData, importance = "permutation")</pre>
```

Create vivid matrix:

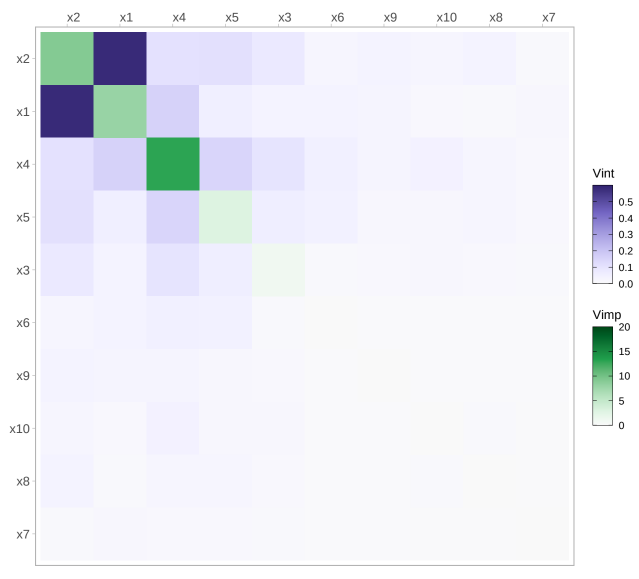
```
# vivi Normalized & Unnormalized:
set.seed(1701)
fNormalT <- vivi(fit = fFit, data = fData, response = "y", normalized = TRUE)
fNormalF <- vivi(fit = fFit, data = fData, response = "y", normalized = FALSE)</pre>
```

Figure 10 (a) & (b)

```
viviHeatmap(fNormalT, impLims = c(0, 20))
```



viviHeatmap(fNormalF, intLims = c(0, 0.6), impLims = c(0, 20))



Now we simulate data with a correlation between x4 and x5 and refit our model:

Create VIVI matrix

```
# vivi Normalized & Unnormalized:
set.seed(1701)
corrNormF <- vivi(fit = corrFit, data = dfCorr, response = "y", normalize = FALSE)</pre>
```

Figure 10 (c)

```
# Heatmap:
viviHeatmap(corrNormF, intLims = c(0, 0.6), impLims = c(0, 20))
```

