Analysis of Multi-Omics Data with Multiple Co-Inertia Analysis via Nonlinear Iterative Partial Least Squares (NIPALS)

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Quick Start

The package can be installed directly from Github via the devtools package:

```
install.packages("devtools",repos = "http://cran.us.r-project.org")
library(devtools)# required for install_github

install_github("Muunraker/NIPALS-MCIA")
library(nipalsMCIA)
```

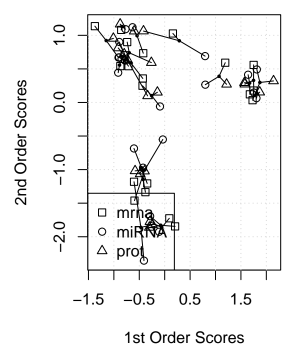
A test dataset of NCI60 cancer cell line data with 3 blocks (mRNA, proteins, and micro-RNA) is included to illustrate how the main MCIA function works with multi-omics data:

```
data(NCI60) # loads data as data_blocks
summary(data_blocks)
```

```
## Length Class Mode
## mrna 12895 data.frame list
## miRNA 537 data.frame list
## prot 7016 data.frame list
```

The main NIPALS function includes a pre-processing option for the data. The results include a plot of the first two global scores as well as an eigenvalue scree plot to aid variable selection.

First and Second Order Scores



Global Score Eigenvalues

