

Vignette 1 - Understanding the Results of MCIA

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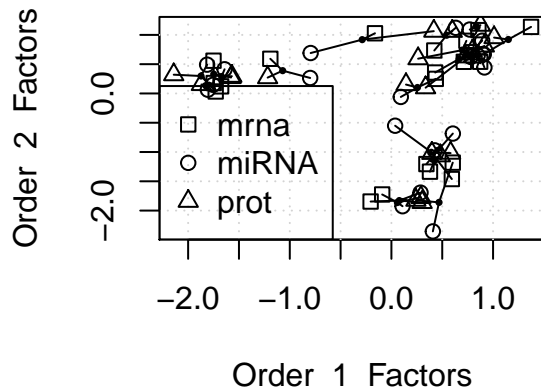
Running MCIA on NCI60 Data and Basic Visualization

NIPALS-MCIA includes a sample multi-omics dataset modified from data collected on the NCI-60 cancer cell lines [CITE: Meng, 2016]. This can be used to illustrate low-dimensional plotting with the global factors.

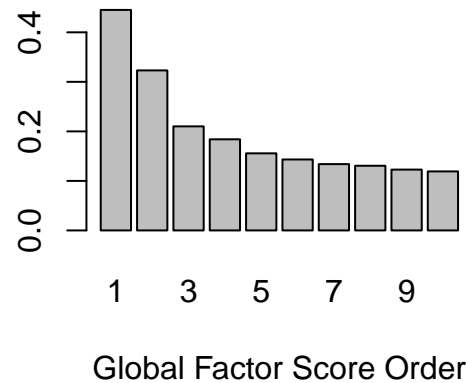
First, we compute the first 10 global factors for the dataset:

```
data(NCI60) # this creates the dataset as `data_blocks`  
mcia_results <- nipals_multiblock(data_blocks, preprocMethod='colprofile', num_PCs = 10, tol=1e-12)
```

Factor Plot



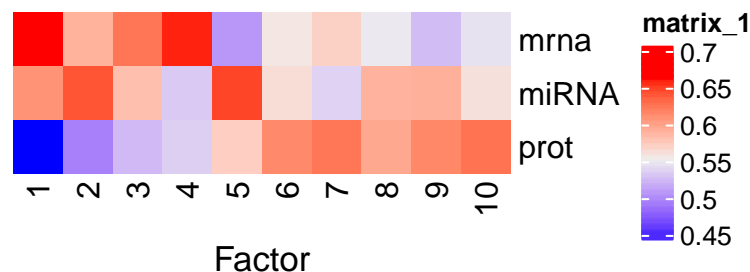
Global Factor Score Eigenvalues



Part 2: Interpreting Global Loadings

Pseudoeigenvalues representing the contribution of each omic to the global factor score

```
## Should make into a function
bs_weights<-as.matrix(data.frame(mcia_results$block_score_weights))
colnames(bs_weights)<-c(1:10)
ComplexHeatmap::Heatmap(bs_weights, cluster_columns=FALSE, cluster_rows=FALSE,
                          column_title_side = "bottom", column_title = "Factor")
```



Scree Plot: Visualizing the top features per factor

```
# make into function
gl<-mcia_results$global_loadings
omic_list<-gsub("^.*_", "", rownames(gl))
factor<-4
gl_f<-data.frame(gl[,factor])
gl_f$omic<-omic_list
colnames(gl_f)<-c("weight", 'omic')
gl_f_ord<-gl_f[order(gl_f$weight, decreasing=TRUE),]

# look at mRNA (need to filter since issue with miRNA omic name)
gl_f_ord_mRNA<-gl_f_ord[gl_f_ord$omic=="mrna",]
omic_name<-sub("_.*", "", rownames(gl_f_ord_mRNA) )
gl_f_ord_mRNA$omic_name<-sub("_.*", "", rownames(gl_f_ord_mRNA) )
ggplot(gl_f_ord_mRNA[0:30,], aes(x=factor(omic_name, level=omic_name), y=weight))+
  geom_point()+
  theme_bw()+
  xlab('Feature')+
  theme(axis.text.x=element_text(angle=45, hjust=1))
```

