## Supervised Multiblock analyses

#### Florence Mehl

### March 11, 2024

- · Nutrimouse dataset
- · Discriminant analysis of genotypes
  - Question 1: based on lipids and genes data, can we discriminate wt vs ppar samples?
  - · Question 2: Choose optimal number of latent variables?
  - · Question 3: Is the model statistically significant?
  - Question 4: what is the variance explained for each block by each latent variable and globally?
  - Question 5: observe the samples distributions in the space of the latent variables.
  - Question 6: which genes and lipids are discriminant for genotype?
- Consensus OPLS Discriminant analysis of genotypes
  - Question 1: based on lipids and genes data, can we discriminate wt vs ppar samples?
  - · Question 2: Is the model statistically significant?
  - Question 3: What is the contribution of each data block?
  - Question 5: Show the loadings of variables in the space of the predictive and orthogonal latent variables?
  - Question 6: Show the importance of variables in the model?

### Nutrimouse dataset

The data sets come from a nutrigenomic study in the mouse (Martin et al., 2007) in which the effects of five regimens with contrasted fatty acid compositions on liver lipids and hepatic gene expression in mice were considered.

Two sets of variables were acquired on forty mice: - genes: expressions of 120 genes measured in liver cells, selected (among about 30,000) as potentially relevant in the context of the nutrition study. These expressions come from a nylon macroarray with radioactive labelling - lipids: concentrations (in percentages) of 21 hepatic fatty acids measured by gas chromatography

Biological units (mice) were cross-classified according to two factors experimental design (4 replicates): - genotype: 2-levels factor, wild-type (WT) and PPARalpha -/- (PPAR) - diet: 5-levels factor. Oils used for experimental diets preparation were corn and colza oils (50/50) for a reference diet (REF), hydrogenated coconut oil for a saturated fatty acid diet (COC), sunflower oil for an Omega6 fatty acid-rich diet (SUN), linseed oil for an Omega3-rich diet (LIN) and corn/colza/enriched fish oils for the FISH diet (43/43/14)

```
data("nutrimouse")
genes <- nutrimouse$gene
lipids <- nutrimouse$lipid
metadata <- data.frame(genotype = nutrimouse$genotype, diet = nutrimouse$diet)
metadata$sample_name <- paste0(rownames(metadata), "_", metadata$genotype, "_", metadata$diet)
rownames(genes) <- metadata$sample_name
rownames(lipids) <- metadata$sample_name</pre>
```

## Discriminant analysis of genotypes

# Question 1: based on lipids and genes data, can we discriminate wt vs ppar samples?

Run block.plsda analysis with block.plsda() from mixomics package

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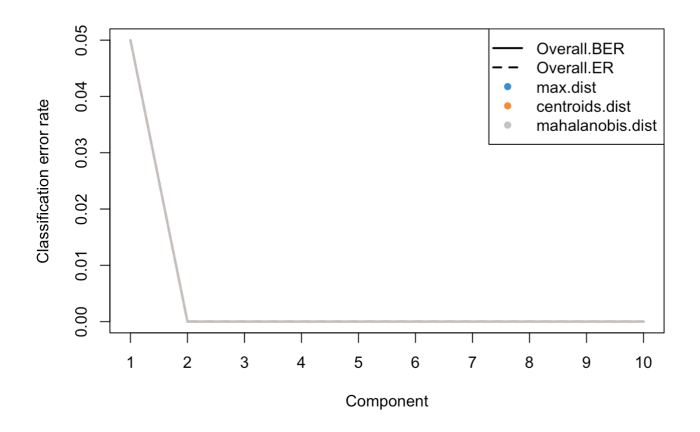
```
# prepare data
blockPLS_data <- list(genes=genes, lipids=lipids)
genotype <- as.factor(metadata$genotype)

# run analysis
blockPLS_res <- block.plsda(X = blockPLS_data, Y = genotype, design = "full", all.outputs
= T, ncomp = 10)</pre>
```

## Question 2: Choose optimal number of latent variables?

Run perf() plot the results with plot() Run the analysis with optimal number of latent variables

```
blockPLS_perf <- perf(blockPLS_res, validation = 'Mfold', folds = 7, nrepeat = 1, auc = TR
UE, cpus=2, progressBar = FALSE)
plot(blockPLS_perf)</pre>
```



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```
blockPLS_res <- block.plsda(X = blockPLS_data, Y = genotype, design = "full", all.outputs
= T, ncomp = 2)</pre>
```

## Question 3: Is the model statistically significant?

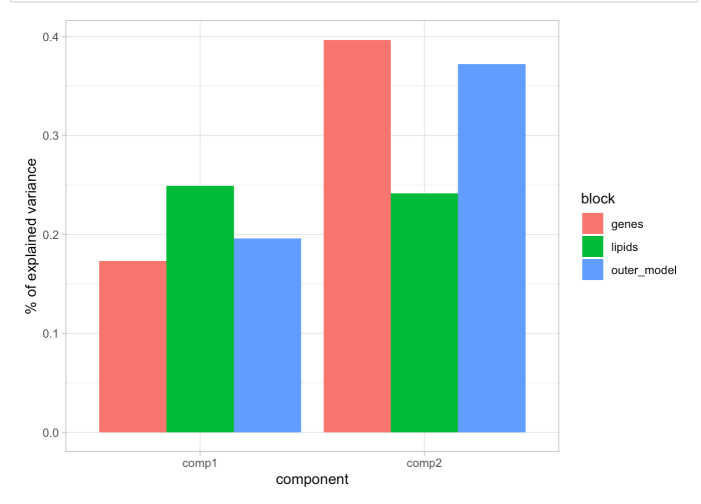
Run a permutation test with DIABLO.test() from RVAideMemoire package

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```
blockPLS_permtest <- DIABLO.test(blockPLS_res, progress = FALSE)</pre>
```

# Question 4: what is the variance explained for each block by each latent variable and globally?

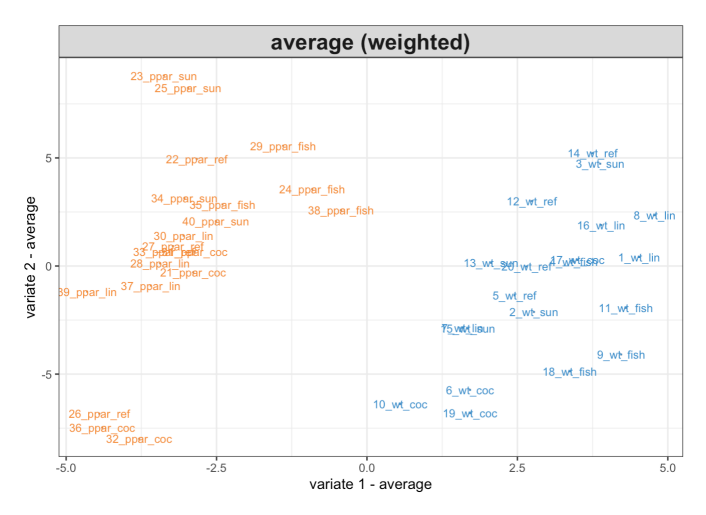
- for each block: AVE\_X
- global: AVE[["AVE\_outer"]]



# Question 5: observe the samples distributions in the space of the latent variables.

plot scores with plotIndiv()

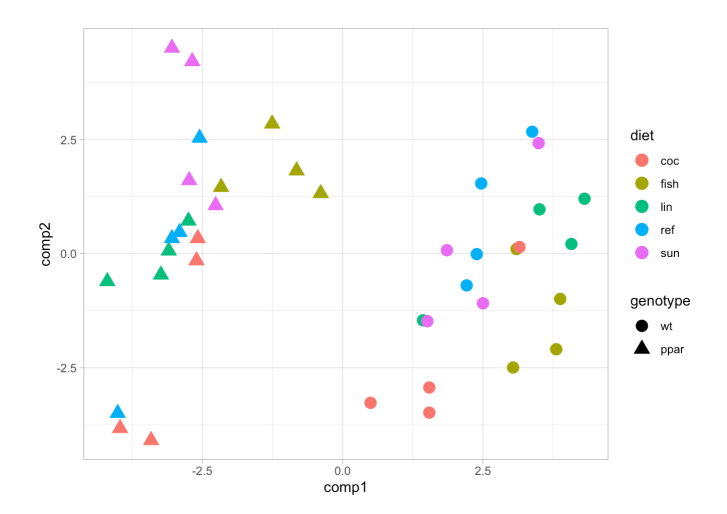
```
plotIndiv(blockPLS_res, block = "weighted.average")
```



```
blockPLS_variates.weighted <- blockPLS_res$variates[c("genes", "lipids")]
for(omic in c("genes", "lipids")){
    for(comp in c("comp1", "comp2")){
        blockPLS_variates.weighted[[omic]][,comp] <- blockPLS_variates.weighted[[omic]][,com
p] * blockPLS_res$weights[omic, comp]
    }
}
blockPLS_scores.weighted <- abind(blockPLS_variates.weighted[c("genes", "lipids")], along
    = 3)
blockPLS_scores.weighted <- apply(blockPLS_scores.weighted, c(1,2), mean)

blockPLS_scores.weighted <- data.frame(metadata, blockPLS_scores.weighted)

ggplot(blockPLS_scores.weighted, aes(x=comp1, y=comp2, col=diet, shape = genotype)) +
    geom_point(size=4) +
    theme_light()</pre>
```

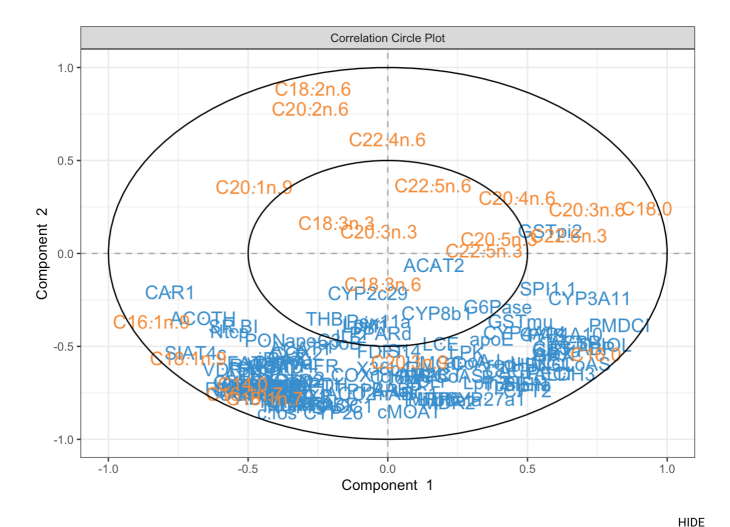


# Question 6: which genes and lipids are discriminant for genotype?

• plot loadings with plotVar()

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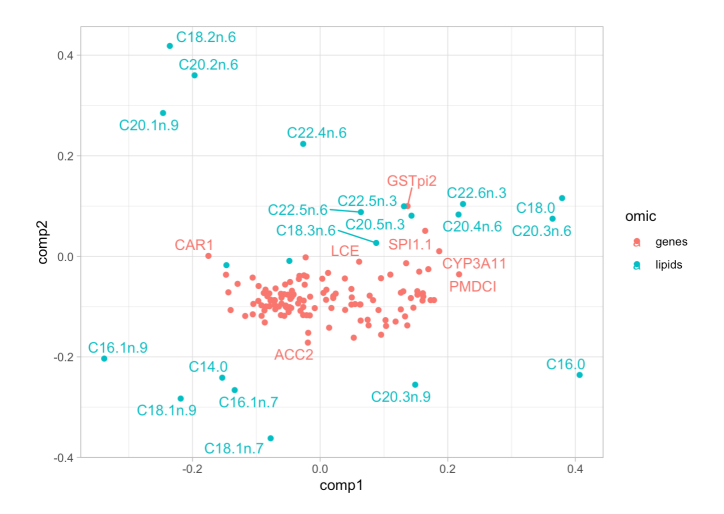
plotVar(blockPLS\_res)



```
# ou

blockPLS_loadings_genes <- blockPLS_res$loadings$genes
blockPLS_loadings_lipids <- blockPLS_res$loadings$lipids
blockPLS_loadings <- rbind.data.frame(blockPLS_loadings_genes, blockPLS_loadings_lipids)
blockPLS_loadings$omic <- c(rep("genes", dim(genes)[[2]]), rep("lipids", dim(lipids)
[[2]]))
blockPLS_loadings$variable <- rownames(blockPLS_loadings)

ggplot(blockPLS_loadings, aes(x=comp1, y=comp2, col=omic, label=variable)) +
    geom_point() +
    geom_text_repel() +
    theme_light()</pre>
```



# Consensus OPLS Discriminant analysis of genotypes

# Question 1: based on lipids and genes data, can we discriminate wt vs ppar samples?

Run ConsensusOPLS-DA analysis with ConsensusOPLS() from ConsensusOPLS package

```
COPLS_data <- list(genes=as.matrix(genes), lipids=as.matrix(lipids))
COPLS_data <- lapply(COPLS_data, scale)
dummy_genotype <- as.matrix(data.frame(wt = ifelse(genotype == "wt", 1, 0),ppar = ifelse(genotype == "ppar", 1, 0)))

COPLS_res <- ConsensusOPLS(
    data = COPLS_data,
    Y = dummy_genotype,
    maxPcomp = 1,
    maxOcomp = 1,
    nfold = 40,
    cvType = "nfold",
    nperm = 100,
    modelType = "da",
    mc.cores = 1,
    verbose = FALSE
)</pre>
```

## Question 2: Is the model statistically significant?

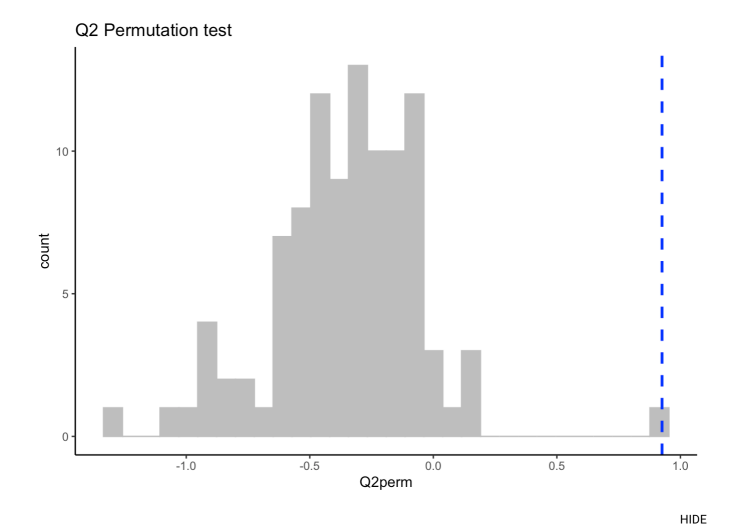
The results of permutations can be found in COPLS\_res

 $perm Stats.\ The results for the optimal model can be found in COPLS_resoptimal model CV and COPLS_resoptimal model CV cv$ 

- · plot Q2 permutations
- plot DQ2 permutations
- · plot R2Y permutations

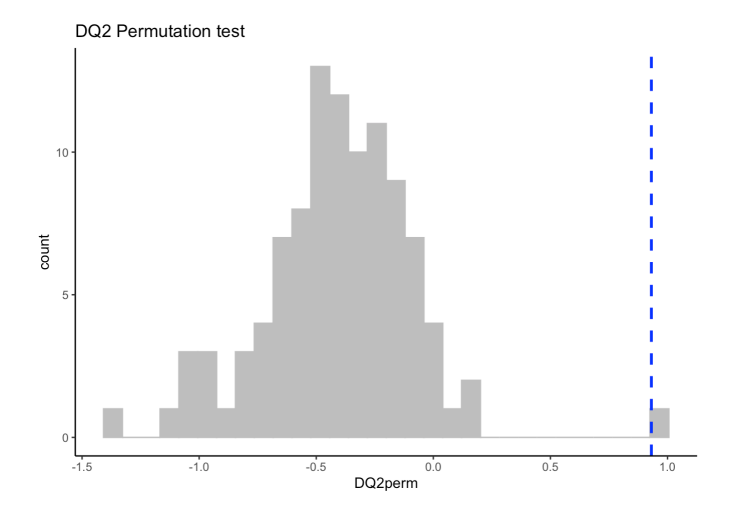
```
Q2perm <- data.frame(Q2perm = COPLS_res$permStats$Q2Yhat)

ggplot(data = Q2perm, aes(x = Q2perm)) +
  geom_histogram(color="grey", fill="grey") +
  geom_vline(aes(xintercept=COPLS_res$optimal$modelCV$cv$Q2Yhat[2]),color="blue", linetype
="dashed", size=1) +
  theme_classic() +
  ggtitle("Q2 Permutation test")</pre>
```



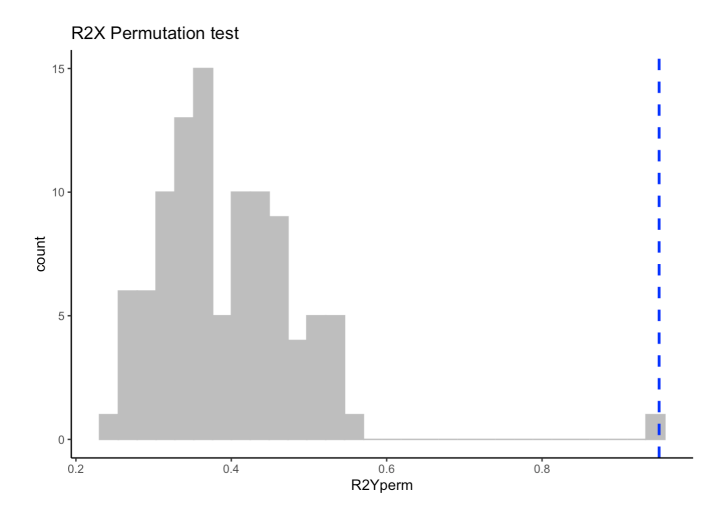
```
DQ2perm <- data.frame(DQ2perm = COPLS_res$permStats$DQ2Yhat)

ggplot(data = DQ2perm, aes(x = DQ2perm)) +
    geom_histogram(color="grey", fill="grey") +
    geom_vline(aes(xintercept=COPLS_res$optimal$modelCV$cv$DQ2Yhat[2]),color="blue", linetype="dashed", size=1) +
    theme_classic() +
    ggtitle("DQ2 Permutation test")</pre>
```



```
R2Yperm <- data.frame(R2Yperm = COPLS_res$permStats$R2Yhat)

ggplot(data = R2Yperm, aes(x = R2Yperm)) +
    geom_histogram(color="grey", fill="grey") +
    geom_vline(aes(xintercept=COPLS_res$optimal$modelCV$Model$R2Yhat[2]),color="blue", linet
ype="dashed", size=1) +
    theme_classic() +
    ggtitle("R2X Permutation test")</pre>
```



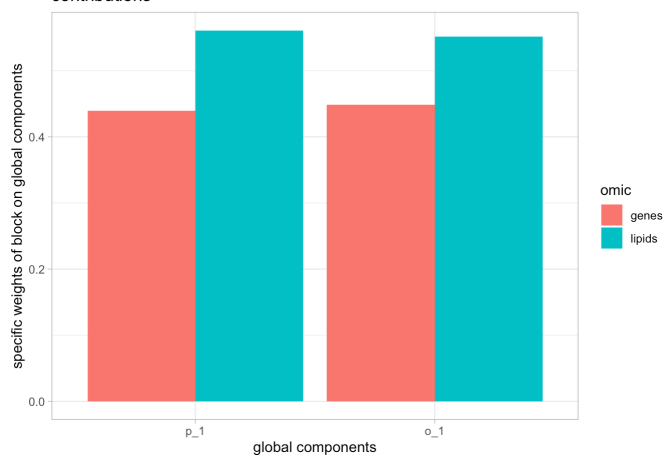
## Question 3: What is the contribution of each data block?

• plot blockContribution of the optimal model

```
contributions <- COPLS_res$optimal$modelCV$Model$blockContribution
contributions <- melt(contributions)
colnames(contributions) <- c("dataset", "Dim", "value")

ggplot(contributions, aes(x=Dim, y=value, fill=dataset)) +
    geom_bar(stat = "identity", position=position_dodge()) +
    theme_light() +
    labs(x = "global components", y = "specific weights of block on global components", fill
= "omic",
    title = "contributions")</pre>
```

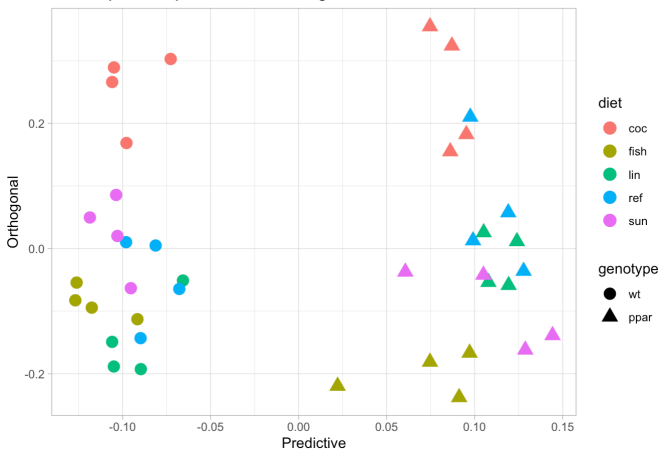
#### contributions



## Question 4: Show the disctribution of samples in the space of the predictive and orthogonal latent variables?

· plot scores of the optimal model

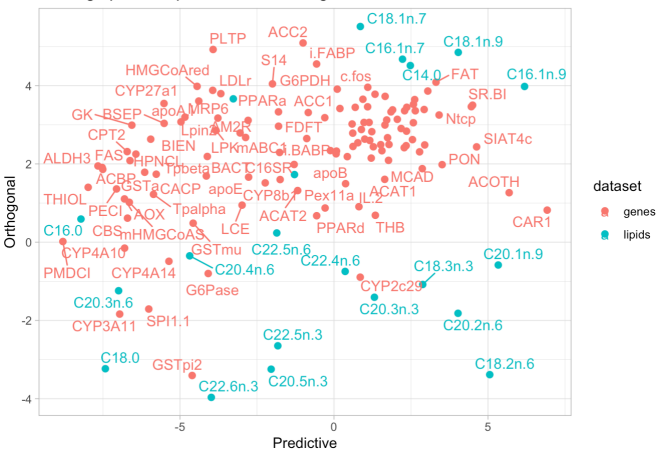
### scores plots on predictive vs orthogonal latent variables



# Question 5: Show the loadings of variables in the space of the predictive and orthogonal latent variables?

· plot loadings of the optimal model

### loadings plots on predictive vs orthogonal latent variables



## Question 6: Show the importance of variables in the model?

• plot loadings and VIP of the optimal model

### loadings plots on predictive vs orthogonal latent variables

