# Gaussian Graphical Models in Metabolomics

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Graphical models in medicine

Data

Introduction to network analysis in R

Gaussian Graphical Models (GGM) in R

# Graphical models in medicine

## NETWORK MEDICINE

- **Fundamental principle**: disease module hypothesis that disease variants are connected.
- Evidence in literature: 10-fold increase in products of genes associated with a disorder when compared to expectation under random chance.
- References: Su and Clish, Metabolomics and Network Medicine, 2017; Goh, K. I., Cusick, M. E. et. al., The human disease network, 2007.

#### METABOLITES AS NETWORKS

Metabolites are naturally represented as networks:

- Nodes: represent individual metabolites.
- Edges (undirected): denote pairwise metabolite relationships.

## EXAMPLE NETWORK

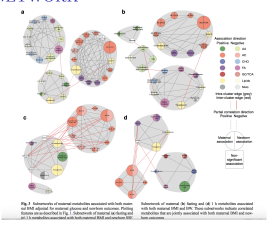


Figure 1: Maternal BMI and newborn SSF associated metabolite networks from Sandler, V., Reisetter, A. C. et. al., Diabetologia, 2017.

#### CORRELATION NETWORKS

- Correlation networks are established methods for constructing metabolite networks.
- Edges in correlation networks depict pairwise correlations between metabolite pairs.
- Networks are often created by thresholding on a correlation cut-off.
- Recent example from literature: A network analysis of biomarkers for Type 2 Diabetes in the Nurses Health Study.

<sup>&</sup>lt;sup>1</sup>Huang, T., Glass, K. et al., Diabetes, 2018.

# CORRELATION NETWORKS

- Drawback: Correlations between metabolite pairs can be driven by direct and indirect relationships.
- Drivers of high correlation include shared or common enzymatic activities. <sup>2</sup>.
- Large number of non-zero pairwise correlations are usually observed.
- Absence of an edge results from satisfying a strong criterion of marginal independence between metabolite pairs.

<sup>&</sup>lt;sup>2</sup>Su and Clish, Metabolomics and Network Medicine, 2017

<sup>&</sup>lt;sup>3</sup>Strimmer, K., Notes on Gaussian Graphical Models.

# Gaussian graphical models (GGM)

- Model: Metabolites are multivariate Gaussian with mean  $\mu$  and covariance matrix  $\Sigma$ .
- The precision (concentration) matrix  $\Omega = \Sigma^{-1}$ .
- If  $\Omega_{jk} = 0$ , then the *i*th metabolite is independent of the *j*th metabolite, given all other variables.

## GGM ESTIMATION

- Meinshausen and Buhlmann (2006): estimates  $\Omega_{jk} = 0$  by fitting a lasso to each metabolite, using all others as predictors.
- $\hat{\Omega}_{jk} \neq 0$ : if the estimated coefficients of metabolite i on j AND vice-versa are non-zero.
- Friedman et al. (2007): Glasso and variants for exact maximization of the penalized log-likelihood.

### MODEL SELECTION

- Gaussian graphical model estimation involves a process to estimate the **optimal regularization parameter**  $(\lambda)$ .
- Large values of  $\lambda$  correspond to increasing sparsity of the resulting graph.

- Stability approach for regularization selection (StARS): uses a subsampling approach to estimate the optimal  $\lambda$ .
- Rotation information criterion (RIC): uses a permutation approach to estimate  $\lambda$ .

## CORRELATION NETWORK VERSUS GGM

 Correlation network: An edge between metabolite pairs can result from both direct AND indirect relationships.

 GGM: An edge exists ONLY if the metabolite pair is dependent after accounting for all other indirect relationships.

# Data

- Hyperglycemia and Adverse Pregnancy Outcome (HAPO) Study conducted during 2000 - 2006 at 15 international field centers.
- Blood samples were obtained during a 75-g oral glucose tolerance test (OGTT) between 24 and 32 weeks gestation.
- Metabolites were measured in maternal fasting and 1-h serum samples from 400 mothers in each ancestry group (Afro-Caribbean, Mexican American, Northern European, Thai).
- Mothers were sampled to span the range of maternal glucose and BMI.

#### Data Format:

- Column 1: ID
- Column 2: Ancestry Group
- Column 3: Fasting glucose
- Columns 4-54: 51 metabolites

### Loading data ..

#PC users

```
#setwd("/cloud/project/Metabolomics R Ladies Strasbourg 2020/")
#mac users
setwd("/cloud/project/Metabolomics R Ladies Strasbourg 2020")
mydat <- read.csv(file = "Data/hapo_metabolomics_2020.csv")</pre>
print(mydat[1:3,1:10])
        id anc_gp fpg mt1_1 mt1_2 mt1_3 mt1_4
##
                                                               mt1 5
## 1 hm0001
             ag3 75.6 218.2223 76.99525 19.06366 14.23091 86.75162
## 2 hm0002 ag3 84.6 292.6314 136.41320 43.14854 17.77549 120.17344
## 3 hm0003
              ag4 79.2 361.1135 79.98370 22.15848 13.05497 74.75441
##
       mt1 7
## 1 64.00578
## 2 91.30156
## 3 83.67878
```

Three groups of metabolites:

- Prefix mt1: Amino Acids (AA)
- Prefix mt2: Acyl carnitines (AC)
- Prefix mt3: Other

## Let's take a look at the numbers by **ancestry group**:

```
ag <- mydat[,2]
table(ag)</pre>
```

```
## ag1 ag2 ag3 ag4
## 400 400 400 400
```

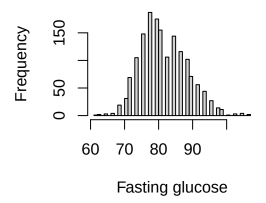
Let's take a look at the distribution of **fasting glucose**:

```
fg <- mydat[,3]
summary(fg)</pre>
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 61.20 77.40 81.00 81.63 86.40 106.20
```

Let's take a look at the distribution of **fasting glucose**:

# Histogram of fg



Introduction to network analysis in R

# **PRELIMINARIES**

• **igraph R** package: provides a way of representing graphs and various tools for working with graphs.

### **Preliminaries**

- Let's work with a small (p=6) set of metabolites sampled from the HAPO dataset.
- As an example, we start with a simple correlation network of 6 metabolites

```
mx <- mydat[,-c(1:3)]
mx.1 <- mx[ag == "ag1", c(1,2,16,17,34,35)]
cor.1 <- round(cor(mx.1, use="pairwise.complete.obs"), digits=2)

### Create an adjacency matrix using a threshold of 0.1
adj.1 <- matrix(0, nrow(cor.1), nrow(cor.1))
adj.1[abs(cor.1) > 0.1] <- 1
colnames(adj.1) <- rownames(adj.1) <- colnames(cor.1)</pre>
```

### Defining network objects in R

Let *p* denote the number of metabolites in our network.

 Adjacency matrix: p × p matrix, where i, j element is 1 if there is an edge between metabolite i and metabolite j, and 0 otherwise.

```
### Adjacency matrix
print(adj.1)
```

#### IGRAPH R PACKAGE

#### We can convert an adjacency matrix to an igraph object.

```
library(igraph)
igraph.obj <- graph.adjacency(adj.1,mode="undirected",weighted=NULL,diag=FALSE)

## Extracting nodes and edges from igraph object
V(igraph.obj)

## + 6/6 vertices, named, from b6868a8:
## [1] mt1_1 mt1_2 mt2_1 mt2_2 mt3_1 mt3_2
E(igraph.obj)

## + 5/5 edges from b6868a8 (vertex names):
## [1] mt1_1--mt1_2 mt1_1--mt2_1 mt1_1--mt2_2 mt1_2--mt3_2 mt2_1--mt2_2</pre>
```

# VISUALIZING OUR NETWORK

Let's assign metabolite class to each of our nodes and an associated color.

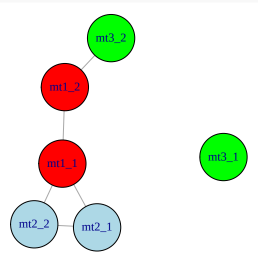
```
### Assigning attributes to the list of nodes

V(igraph.obj)$MxClass <- c(rep("AA",2), rep("AC", 2), rep("Oth",2))
V(igraph.obj)$color <- c(rep("red", 2), rep("light blue",2), rep("green",2))
V(igraph.obj)$size <- 50
V(igraph.obj)$label.cex <- 0.75</pre>
```

# VISUALIZING OUR NETWORK

#### Visualize the network..

### Visualizing network
plot.igraph(igraph.obj,vertex.label=colnames(adj.1),layout=layout.fruchterman.reingold)



## CHANGING NODE ATTRIBUTES

Let's change node size in proportion to significance of association with fasting glucose..

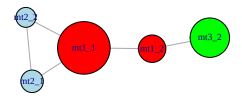
```
### Changing the node size to match the level
### of signficance with outcome (fasting glucose)
myfun <- function(metabolite, outcome){</pre>
    mymod <- lm(outcome ~ metabolite)</pre>
    minuslogp <- -log(summary(mymod)$coef[2,4])
    return(minuslogp)
fg1 \leftarrow fg[ag == "ag1"]
vals <- apply(mx.1, 2, myfun, fg1)</pre>
### scaling the node size
### changing the font fize
### of the vertex label
V(igraph.obj)$size <- vals*3+20
V(igraph.obj) $label.cex <- 0.6
```

# VISUALIZING OUR NETWORK

## Visualize the network after changing node attributes..

### Visualizing network
plot.igraph(igraph.obj,vertex.label=colnames(adj.1),layout=layout.fruchterman.reingold)



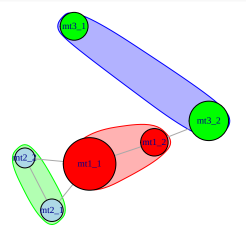


#### GROUPING NODES

We can also visually depict metabolite classes (Amino acids, Acyl carnitines, Other) in our network ..

```
### Visualizing network with node groups
mylist <- list(c("mt1_1","mt1_2"), c("mt2_1","mt2_2"), c("mt3_1","mt3_2"))</pre>
```

# GROUPING NODES



### NETWORKS IN R.

There are a myriad of options available for visualizing networks. For more, see help associated with plot.igraph() in the igraph package.

```
### Other layouts (Kamada-Kawai)
### For other options -- Check ?plot.igraph

1 <- layout_with_kk(igraph.obj)
plot.igraph(igraph.obj,vertex.label=colnames(adj.1),layout=1, mark.groups=mylist)</pre>
```

# Gaussian Graphical Models (GGM) in R

## GGM IN R.

We illustrate estimation of the Gaussian graphical model using the R package huge.

#### To keep in mind:

- Missing values of metabolite levels need to be imputed prior to invoking the functions in huge.
- Each metabolite should be standardized to render them of unit variance.

### **PRELIMINARIES**

We prepare metabolite data in ancestry group ag1 for graphical model estimation.

```
### Prepping data for GGM
### Impute missing values
### Standardize
standardizeMetabolite = function(x)
  x[x == Inf] \leftarrow NA
  x[is.na(x)] \leftarrow min(x, na.rm=T)/2
  return((x-mean(x, na.rm=T))/sd(x, na.rm=T))
mx.1 <- mx[ag == "ag1",]
mx1.s <- apply(mx.1, 2, standardizeMetabolite)</pre>
summary(apply(mx1.s,2,sd))
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 1 1 1 1 1 1
```

## GGM ESTIMATION

#### The key functions involved are:

- huge: estimates GGM over a range of penalty parameters (can be left unspecified).
- huge.select: implements regularization parameter selection.
   Reference: T. Zhao and H. Liu (2012). The huge Package for High-dimensional Undirected Graph Estimation in R. Journal of Machine Learning Research.

## GGM ESTIMATION

#### Regularization parameter selection options include:

- StARS: tends to overselects edges.
- RIC: more computationally efficient, tends to underselect edges.
- Reference: T. Zhao and H. Liu (2012). The huge Package for High-dimensional Undirected Graph Estimation in R. Journal of Machine Learning Research.

#### GGM ESTIMATION

Let's estimate the GGM network for our data...

```
library(huge)
### creates the GGM model object
mbModel <- huge(mx1.s, method="mb")</pre>
## Conducting Meinshausen & Buhlmann graph estimation (mb)....done
### Optimal parameter selection using ric
mbOptRIC = huge.select(mbModel, criterion="ric")
## Conducting rotation information criterion (ric) selection....done
## Computing the optimal graph....done
### extract the graph corresponding to optimal param
mbOptRICGraph = mbOptRIC$refit
```

## GGM

#### Visualize our estimated GGM ..

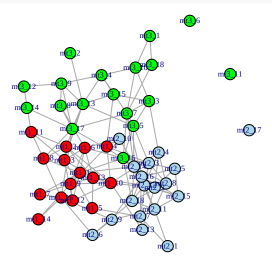
#### Let's estimate the GGM network for our data..

```
myg <- graph_from_adjacency_matrix(mbOptRICGraph, mode="undirected")
### Assigning attributes to the list of nodes

V(myg)$MxClass <- c(rep("AA",15), rep("AC", 18), rep("Oth",18))
V(myg)$color <- c(rep("red", 15), rep("light blue",18), rep("green",18))
V(myg)$size <- 10
V(myg)$label.cex <- 0.5</pre>
```

# GGM

### Visualizing network
plot.igraph(myg,vertex.label=colnames(mx.1),layout=layout.fruchterman.reingold)



#### OTHER OPTIONS

- Method: can be changed to glasso; huge(.., method="glasso").
- Selecting λ: in huge.select(.., criterion="stars").
- Relaxing Gaussian assumption: using nonparanormal (npn) transformation; huge.npn() will return a transformed data matrix.

NEXT ..

## Telling stories with GGMs

- Detecting communities within networks
- Differential networks
- Case studies

#### REFERENCES

- Su, J. and Clish, C. (2018). Metabolomics and Network Medicine, Network Medicine: Complex Systems in Human Disease and Therapeutics, Harvard University Press.
- Go, KI, Cusick, ME, Valle, D, Childs B, Vidal M, Barabási AL (2007). The human disease network, PNAS, 104(21):8685-90.
- Sandler, V., Reisetter, A. C., Bain, J.R., ..., Scholtens, D.M., Lowe, W.L.Jr (2018) Associations of maternal BMI and insulin resistance with the maternal metabolome and newborn outcomes, Diabetologia, 60(3):518-530.
- Meinshausen, N. and Buhlmann, P. (2006). High-dimensional graphs and variable selection with the Lasso, Annals of Statistics, Vol. 34, No. 3, 1436-1462.
- Friedman, J., Hastie, T. and Tibshirani, R. (2008). Sparse inverse covariance estimation with the graphical lasso, Biostatistics, 9(3):432-441.
- Roeder, K., Lafferty, J., Wasserman, L., Zhao, T., Liu, H. (2012) The huge package for high-dimensional undirected graph estimation in R. Journal of Machine Learning Research, (13):1059–1062.

# Gaussian Graphical Models in Metabolomics -Part 2

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- 1) Subnetworks associated with phenotype
- 2) Differential network analysis

#### BEYOND SIMPLE NETWORKS

- Graphical lasso identifies conditional dependence between pairs of metabolites and applies a node-and-edge graph representation of these dependencies
- While estimating conditional dependencies among metabolite pairs is interesting, for most investigations, these dependencies are not of primary interest.
- More complex questions:
  - Which subnetworks are associated with a phenotype?
  - Do networks vary across groups?

1) Subnetworks associated with phenotype

#### Subnetworks associated with phenotype

- Prior to network analyses, investigators often perform per-metabolite association analyses with a phenotype of interest
- How can per-metabolite and network analyses be linked?
- Some existing approaches:
  - Dittrich et al. (2008) Bioinformatics. Identifying functional modules in protein–protein interaction networks: an integrated exact approach.
  - Ben-Hamo et al. (2014) Bioinformatics. PhenoNet: identification of key networks associated with disease phenotype.
  - Soul et al. (2015) Scientific Reports. PhenomeExpress: A refined network analysis of expression datasets by inclusion of known disease phenotypes.

## Subnetworks associated with phenotype

- A simple approach using graphical lasso
  - Identify a set of metabolites,  $\mathcal{M}_p$ , associated with phenotype
  - Identify additional metabolites,  $\mathcal{M}_c$ , with Pearson correlation exceeding some threshold (say 0.25) with at least one member of  $\mathcal{M}_p$
  - Run graphical lasso on  $\mathcal{M}_p \cup \mathcal{M}_c$

In case you'd like to start a new R session, let's reload the libraries and set the working directory.

```
#PC users
#setwd("/cloud/project/Desktop/Metabolomics R Ladies Strasbourg 2020/")
#mac users
setwd("/cloud/project/Metabolomics R Ladies Strasbourg 2020/")
library(igraph)
library(ggplot2)
library(iDINGO)
library(huge)
```

Now read in the data and review some simple descriptors.

```
mydat <- read.csv("Data/hapo_metabolomics_2020.csv")</pre>
rownames(mydat) <- mydat$id</pre>
dim(mydat)
## [1] 1600
             54
head(colnames(mydat))
## [1] "id"
                "anc_gp" "fpg"
                                   "mt1 1" "mt1 2" "mt1 3"
table(mydat$anc_gp)
##
## ag1 ag2 ag3 ag4
## 400 400 400 400
```

Perform simple ancestry-group specific mean imputation of missing metabolite values.

```
hapo_ag <- split(mydat,f=mydat$anc_gp)
length(hapo_ag)
## [1] 4
sapply(hapo_ag,FUN=dim)
##
       ag1 ag2 ag3 ag4
## [1,] 400 400 400 400
## [2,] 54 54 54 54
hapo_ag_m_i <- lapply(hapo_ag,
        FUN=function(x) apply(x[,grep("mt",colnames(x),value=TRUE)],
        MARGIN=2.
        FUN=function(y) ifelse(is.na(y),mean(y,na.rm=TRUE),y)))
```

Check to make sure imputation worked as planned.

```
hapo_m_i <- do.call("rbind",hapo_ag_m_i)</pre>
hapo_i <- data.frame(mydat[rownames(hapo_m_i),c("id", "anc_gp", "fpg")],
                     hapo_m_i)
tapply(mydat[, "mt3_4"], INDEX=mydat$anc_gp, FUN=mean, na.rm=TRUE)
##
        ag1
              ag2
                          ag3
                                   ag4
## 18.11342 22.06506 20.54547 19.95429
tapply(mydat[,"mt3_12"],INDEX=mydat$anc_gp,FUN=mean,na.rm=TRUE)
##
        ag1
             ag2
                          ag3
                                   ag4
## 26.41744 29.66998 29.01828 26.97278
```

Check to make sure imputation worked as planned.

```
mydat[c(1,2,3,6),c("anc_gp","mt3_4","mt3_12")]
##
         anc_gp mt3_4 mt3_12
## hm0001
            ag3 20.50824 29.37834
## hm0002 ag3
                     NA 29.51101
## hm0003 ag4 19.89055 27.85653
## hm0006
           ag4 20.04486
hapo_i[rownames(mydat)[c(1,2,3,6)],c("anc_gp","mt3_4","mt3_12")]
         anc_gp mt3_4 mt3_12
##
         ag3 20.50824 29.37834
## hm0001
         ag3 20.54547 29.51101
## hm0002
           ag4 19.89055 27.85653
## hm0003
## hm0006
            ag4 20.04486 26.97278
```

## ANCESTRY-SPECIFIC NETWORKS ASSOC WITH FPG

Find subset of metabolites within each ancestry associated with fpg.

```
myfun <- function(metabolite,outcome){</pre>
    mymod <- lm(outcome~metabolite)</pre>
    minuslogp <- -log(summary(mymod)$coef[2,4])
    return(minuslogp)
hapo_i_ag <- split(hapo_i,f=hapo_i$anc_gp)
m_fpg_p_ag <- lapply(hapo_i_ag,</pre>
             FUN=function(x){
                 x_m <- x[,grep("mt",colnames(x))]</pre>
                 ans <- apply(x_m,MARGIN=2,FUN=myfun,outcome=x$fpg)
                 return(ans)
                 7)
```

## Find subset of metabolites within each ancestry associated with fpg.

```
sig_m_ag <- lapply(m_fpg_p_ag,
       FUN=function(x) names(x[which(x>-log(.05))]))
sig m ag
## $ag1
## [1] "mt1 1" "mt1 2" "mt1 3" "mt1 5" "mt1 11" "mt1 12" "mt2 3" "mt2 8"
## [9] "mt2 11" "mt3 1" "mt3 2" "mt3 3" "mt3 4" "mt3 5" "mt3 10" "mt3 15"
##
## $ag2
## [1] "mt1 1" "mt1 2" "mt1 3" "mt1 5" "mt1 11" "mt1 12" "mt2 10" "mt3 4"
## [9] "mt3_6" "mt3_9" "mt3_13" "mt3_16"
##
## $ag3
## [1] "mt1 1" "mt1 2" "mt1 3" "mt1 5" "mt1 8" "mt1 11" "mt1 12" "mt1 15"
## [9] "mt2_4" "mt2_8" "mt2_13" "mt2_14" "mt3_1" "mt3_6" "mt3_10" "mt3_13"
##
## $ag4
## [1] "mt1 1" "mt1 5" "mt1 12" "mt1 15" "mt2 2" "mt2 8" "mt2 14" "mt3 5"
## [9] "mt3_12"
```

Find other metabolites correlated with significant metabolites.

```
m_cor_ag <- lapply(hapo_ag_m_i,FUN=cor,use="pairwise.complete.obs")</pre>
sig_cor_ag <- vector("list",length=4)</pre>
names(sig_cor_ag) <- names(sig_m_ag)</pre>
for (i in 1:4){
    sig_m_cor_pairs <- m_cor_ag[[i]][sig_m_ag[[i]],]</pre>
    sig_m_cor <- names(which(colSums(abs(sig_m_cor_pairs)>=.25)>0))
    sig_m_cor_vals <- hapo_ag_m_i[[i]][,sig_m_cor]</pre>
    sig_m_cor_vals_s <- apply(sig_m_cor_vals,MARGIN=2,FUN=scale)</pre>
    sig_cor_ag[[i]] <- sig_m_cor_vals_s</pre>
sapply(sig_cor_ag,FUN=dim)
##
        ag1 ag2 ag3 ag4
## [1,] 400 400 400 400
## [2,] 42 40 44 31
```

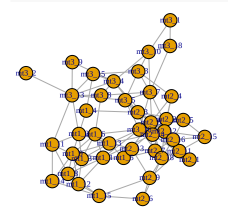
Now apply graphical lasso for these subsets of metabolites.

```
mbModel_ag <- lapply(sig_cor_ag,FUN=huge,method="mb")</pre>
## Conducting Meinshausen & Buhlmann graph estimation (mb)....done
mb_opt_ag <- lapply(mbModel_ag,FUN=huge.select,criterion="ric")</pre>
## Conducting rotation information criterion (ric) selection....done
## Computing the optimal graph....done
## Conducting rotation information criterion (ric) selection....done
## Computing the optimal graph....done
## Conducting rotation information criterion (ric) selection....done
## Computing the optimal graph....done
## Conducting rotation information criterion (ric) selection....done
## Computing the optimal graph....done
```

## Generate the igraph objects.

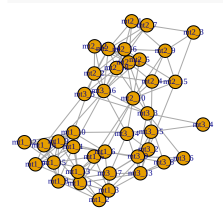
Now plot the graphs - Ancestry group 1 (layout may vary)

```
plot(ggm_ag_g[["ag1"]],vertex.label=V(ggm_ag_g[["ag1"]])$label,
    vertex.label.cex=.5)
```



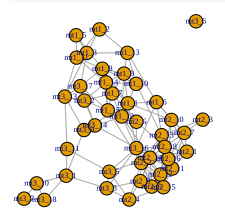
#### Ancestry group 2

```
plot(ggm_ag_g[["ag2"]],vertex.label=V(ggm_ag_g[["ag2"]])$label,
    vertex.label.cex=.5)
```



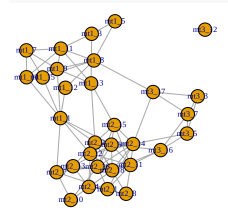
#### Ancestry group 3 - note the singleton node

```
plot(ggm_ag_g[["ag3"]],vertex.label=V(ggm_ag_g[["ag3"]])$label,
    vertex.label.cex=.5)
```

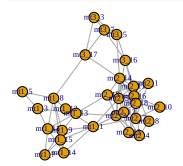


#### Ancestry group 4 - note the singleton node

```
plot(ggm_ag_g[["ag4"]],vertex.label=V(ggm_ag_g[["ag4"]])$label,
    vertex.label.cex=.5)
```



#### Drop the singletons.



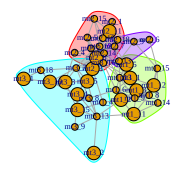
- Visual inspection and biological interpretation of these networks is challenging
- Pick out pairwise relationships? Then what?
- Community detection helps tell a story
- igraph package
  - cluster\_spinglass (Newman and Girvan, 2004)
  - cluster\_fast\_greedy
  - cluster\_label\_prop
  - cluster\_walktrap
  - etc.

Spinglass clustering on all four graphs

ggm\_ag\_g\_spg <- lapply(ggm\_ag\_g,FUN=cluster\_spinglass)</pre>

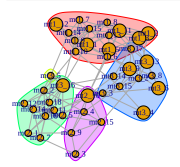
#### Spinglass clustering - ancestry group 1

```
plot(ggm_ag_g[["ag1"]],
    vertex.label=V(ggm_ag_g[["ag1"]])$label,
    vertex.label.cex=.5,
    mark.groups=ggm_ag_g_spg[["ag1"]],
    vertex.size=ifelse(V(ggm_ag_g[["ag1"]])$label %in%
        sig_m_ag[["ag1"]],20,10))
```



#### Spinglass clustering - ancestry group 2

```
plot(ggm_ag_g[["ag2"]],
    vertex.label=V(ggm_ag_g[["ag2"]])$label,
    vertex.label.cex=.5,
    mark.groups=ggm_ag_g_spg[["ag2"]],
    vertex.size=ifelse(V(ggm_ag_g[["ag2"]])$label %in%
        sig_m_ag[["ag2"]],20,10))
```



## EXAMPLE FROM HAPO METABOLOMICS

- Investigation of associations between maternal metabolites at 28 weeks gestation with newborn phenotypes at birth
- Examined associations within and across four ancestry groups Afro-Caribbean, European, Mexican-American, Thai
- Used a similar approach to that described here
- For graphical lasso, used residuals from a linear model for each metabolite with predictors for covariates of interest
- Kadakia et al. (2019) Diabetologia Maternal metabolites during pregnancy are associated with newborn outcomes and hyperinsulimaemia across ancestries.

## Example from HAPO Metabolomics

Maternal fasting metabolites associated with newborn sum of skinfolds under 2 covariate adjustment models

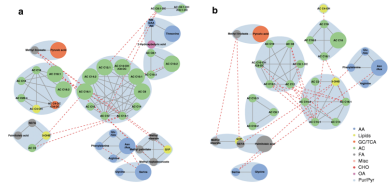


Figure 1: Kadakia et al. (2019)

# 2) Differential network analysis

## DIFFERENTIAL NETWORK ANALYSIS

- Visual inspection suggests there are differences in the ancestry-specific networks we just generated
- But are the differences 'statistically significant'?
- One approach to differential network analysis:
  - iDINGO R package
  - Ha et al. Bioinformatics (2015) DINGO: differential network analysis in genomics.
  - Class et al. Bioinformatics (2018) iDINGO integrative differential network analysis in genomics with Shiny application.

- DINGO estimates a 'global' component of the network,  $\mathcal{G}$ , that represents edges that are common across groups
- DINGO also estimates 'local' group-specific components,  $\mathcal{L}(x)$ , that represent unique relationships in each group depending on the value of a categorical variable x.
- For two groups, group-specific edges are identified using a Differential Score:

$$\delta_{ab}^{(12)} = \frac{\hat{\phi}_{ab}^{(1)} - \hat{\phi}_{ab}^{(2)}}{s_{ab}^{B}}$$

where  $\hat{\phi}_{ab}^{(1)}$  and  $\hat{\phi}_{ab}^{(2)}$  are Fisher's Z transformation of the estimates of group-specific partial correlations between metabolites a and b in groups 1 and 2, and  $s_{ab}^B$  is the bootstrap estimate of the standard error.

```
Let's work with the first two ancestry groups.

hapo_2ag <- subset(hapo_i,anc_gp %in% c("ag1","ag2"))
hapo_2ag <- droplevels(hapo_2ag)
hapo_2ag_mt <- hapo_2ag[,grep("mt",colnames(hapo_2ag),value=TRUE)]
dim(hapo_2ag)

## [1] 800 54
dim(hapo_2ag_mt)

## [1] 800 51
```

The commented code below would perform the DINGO algorithm. The bootstrapping takes a long time. So we will just load an R object of the results that should be in your working directory.

```
#hapo_2ag_dn <- dingo(hapo_2ag_mt,x=hapo_2ag$anc_gp,B=50)
load("Data/hapo_2ag_dn_B50.rda")</pre>
```

Let's look at the various components of the output.

## [1] 1275

```
names(hapo_2ag_dn)
    [1] "genepair" "levels.x" "R1"
##
                                              "R2"
                                                          "boot.diff"
##
    [6] "diff.score" "p.val"
                                 "rho"
                                              "P"
                                                          "0"
## [11] "Psi" "step.times"
head(hapo_2ag_dn$genepair)
## gene1 gene2
## 1 mt1_1 mt1_2
## 2 mt1_1 mt1_3
## 3 mt1_2 mt1_3
## 4 mt1_1 mt1_4
## 5 mt1_2 mt1_4
## 6 mt1_3 mt1_4
dim(hapo_2ag_dn$genepair)
```

```
More components of the output.
hapo_2ag_dn$levels.x
## [1] ag1 ag2
## Levels: ag1 ag2
length(hapo_2ag_dn$R1)
## [1] 1275
length(hapo_2ag_dn$R2)
## [1] 1275
dim(hapo_2ag_dn$boot.diff)
## [1] 1275
              50
```

```
More components of the output.

length(hapo_2ag_dn$diff.score)

## [1] 1275

length(hapo_2ag_dn$p.val)

## [1] 1275
```

#### Create a data frame of some of the output

# Create a data frame of some of the output.

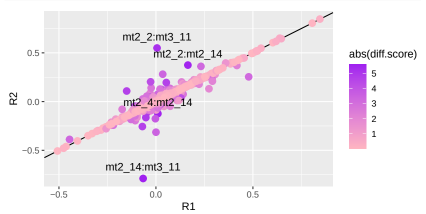
```
head(hapo_2ag_dn_df)
```

```
## 1 mt1_1 mt1_2 mt1_1:mt1_2 0.07809638 0.07832990 -0.016040253 0.9986491 ## 2 mt1_1 mt1_3 mt1_2:mt1_3 0.01951538 0.02186509 -0.135858912 0.8148208 ## 3 mt1_2 mt1_3 mt1_2:mt1_3 0.40212136 0.40158482 0.047007443 0.9039632 ## 4 mt1_1 mt1_4 mt1_1:mt1_4 -0.25814119 -0.25921713 0.092097027 0.8351067 ## 5 mt1_2 mt1_4 mt1_2:mt1_4 0.29185984 0.29178087 0.005269656 0.9683637 ## 6 mt1 3 mt1 4 mt1_3:mt1 4 -0.24110807 -0.24006858 -0.076546532 0.9051876
```

### Identify extremely different scores with diff.score>5 or <-5.

```
hapo_2ag_dn_df$high_ds <- ifelse(abs(hapo_2ag_dn_df$diff.score)>5,
                                as.character(hapo_2ag_dn_df$genepair),"")
hapo 2ag dn df[which(!hapo 2ag dn df$high ds=="").]
##
       gene1 gene2
                         genepair
                                           R1
                                                      R2 diff.score p.val
## 395 mt2_2 mt2_14 mt2_2:mt2_14 0.164750400 0.3744255 -5.521262
## 397 mt2_4 mt2_14 mt2_4:mt2_14 0.010156435 -0.1238160
                                                         5.522768
## 920 mt2_2 mt3_11 mt2_2:mt3_11 0.005775354 0.5498016 -5.201985
## 932 mt2_14 mt3_11 mt2_14:mt3_11 -0.065983987 -0.7890289 5.598957
                                                                        0
##
            high_ds
## 395 mt2 2:mt2 14
## 397 mt2 4:mt2 14
## 920 mt2_2:mt3_11
## 932 mt2 14:mt3 11
```

Compare R1 and R2, colored by diff.score.

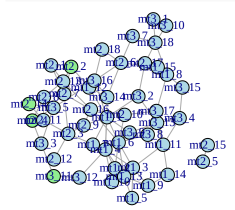


Plot of diff.score by p.val, colored by diff.score.

```
ggplot(hapo_2ag_dn_df,aes(x=diff.score,y=p.val)) +
         geom_point(aes(color=abs(diff.score)),size=3) +
         scale_color_gradient(low="lightpink",high="purple")
  1.00 -
                                                             abs(diff.score)
  0.75 -
                                                                 5
o.50 -
                                                                 3
                                                                 2
  0.25 -
  0.00 -
                            diff.score
```

Explore the global component of the dingo graph.

Explore the global component of the dingo graph.



Explore the local components of the dingo graphs.

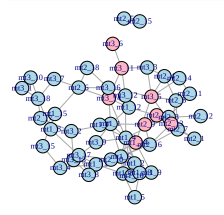
```
hapo_2ag_dn_df$local_ag1 <- ifelse(
          (abs(hapo_2ag_dn_df$R1)>dingo_rho_thresh) &
                    (abs(hapo_2ag_dn_df$R2)<dingo_rho_thresh) &
                    (hapo_2ag_dn_df$p.val<.05),1,0)
hapo_2ag_dn_df$local_ag2 <- ifelse(
          (abs(hapo_2ag_dn_df$R2)>dingo_rho_thresh) &
                    (abs(hapo_2ag_dn_df$R1)<dingo_rho_thresh) &
                    (hapo 2ag dn df$p.val<.05),1,0)
table(hapo_2ag_dn_df$local_ag1,hapo_2ag_dn_df$local_ag2)
##
##
##
    0 1259 11
##
```

Explore the local components of the dingo graphs.

### Explore the local components of the dingo graphs.

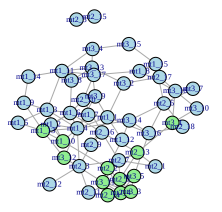
Local component for ancestry group 1

plot(local\_g\_ag1,vertex.label.cex=.5)



Local component for ancestry group 2

plot(local\_g\_ag2,vertex.label.cex=.5)



# SUMMARY

- Networks are very helpful for 'story telling' in metabolomics (and other omics) settings
- Graphical lasso and related methods focus on conditional dependence
- Gives some assurance that edges aren't simply an artifact of sharing common correlations between a pair of nodes with a third node
- Focusing on subnetworks related to phenotype can place per-metabolite associations into context
- Differential network analyses based on graphical models can point to meaningful differences between groups
- Graphics take a while...be patient and use Google!

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