

TWAS Analysis of Diversity Outbred Strain RNAseq Data

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Contents

1 Purpose	1
2 Experimental Details	1
3 Raw Data	2
4 Analysis	2
4.1 Cholesterol Levels	2
5 TWAS with Cholesterol Levels for NCD	4
6 TWAS with Cholesterol Levels for HFD	7
6.1 TWAS Modification by Sex	10
7 Comparason with mouse GWAS	10
8 Integrated TWAS Analysis	11
9 Pathway Analyses for HFD	26
10 Specific Associations	37
10.1 Bile Acid Metabolism	39
10.2 SCD1	41
10.3 Serpina3k	43
10.4 Ugt1a5	45
10.5 Lasp1	47
11 Session Information	49

1 Purpose

2 Experimental Details

The RNA expression data was downloaded from GSE72759 as a matrix file, and compared with genotypes from the Svenson-183 dataset.

3 Raw Data

```
library(readr) #loads the readr package
expression.filename <- "GSE72759_D0192_RNAseq_UpperQuartileNormalized_n21454genes_forGEOSubmission.txt"
expression.data <- read_tsv(expression.filename, show_col_types = F) %>%
  dplyr::rename(ENSEMBL.ID=1)

genotype.filename <- 'Svenson-183_Svenson_D0-MegaMUGA-calls.csv'
genotype.data <- read_csv(genotype.filename,
                          col_types = cols(
                            .default = col_character(),
                            chr = col_factor(levels=NULL),
                            pos = col_double()
                          ))

phenotype.filename <- 'Svenson_HFD_D0_phenotype_V12.csv'
phenotype.data <- read_csv(phenotype.filename)

phenotype.data[phenotype.data=='-999999'] <- NA

mean.expression <-
  expression.data %>%
  dplyr::select(contains(phenotype.data$mouse.id))%>%
  rowMeans()

expression.data <-
  expression.data %>%
  filter(mean.expression>10)
```

4 Analysis

Only evaluated gene expression for genes with >10 TPM

4.1 Cholesterol Levels

Regressed cholesterol levels by diet and sex

```
summary.data <-
  phenotype.data %>%
  group_by(sex,diet) %>%
  summarize_at(.vars=vars(chol2), .funs=list(mean=mean,
                                             se=se,
                                             sd=sd,
                                             n=length))

kable(summary.data,caption="Summary statistics for cholesterol levels at 8 weeks")
```

Table 1: Summary statistics for cholesterol levels at 8 weeks

sex	diet	mean	se	sd	n
F	chow	NA	1.48	NA	225
F	hf	NA	2.40	NA	198
M	chow	NA	1.57	NA	224
M	hf	NA	2.33	NA	193

```
library(broom)
lm(chol2~sex*diet, data=phenotype.data) %>%
  tidy %>%
  kable(caption="Global interactions between sex and diet")
```

Table 2: Global interactions between sex and diet

term	estimate	std.error	statistic	p.value
(Intercept)	78.67	1.88	41.887	0.000
sexM	17.80	2.68	6.648	0.000
diethf	34.64	2.75	12.615	0.000
sexM:diethf	-1.86	3.94	-0.474	0.636

```
chol.lm <- lm(chol2~sex+diet, data=phenotype.data)
chol.lm.ncd <- lm(chol2~sex, data=filter(phenotype.data, diet=='chow'))

cholesterol.data.ncd <-
  phenotype.data %>%
  filter(!is.na(chol2)) %>%
  filter(diet=='chow') %>%
  mutate(adj.chol.ncd=residuals(chol.lm.ncd)+coefficients(chol.lm.ncd)['(Intercept)'])

summary.data <-
  phenotype.data %>%
  group_by(sex,diet) %>%
  summarize_at(.vars=vars(chol2), .funs=list(mean=mean,
                                             se=se,
                                             sd=sd,
                                             n=length))

kable(summary.data,caption="Summary statistics for cholesterol levels at 8 weeks")
```

Table 3: Summary statistics for cholesterol levels at 8 weeks

sex	diet	mean	se	sd	n
F	chow	NA	1.48	NA	225
F	hf	NA	2.40	NA	198
M	chow	NA	1.57	NA	224
M	hf	NA	2.33	NA	193

```
library(broom)
lm(chol2~sex*diet, data=phenotype.data) %>%
```

```
tidy %>%
kable(caption="Global interactions between sex and diet")
```

Table 4: Global interactions between sex and diet

term	estimate	std.error	statistic	p.value
(Intercept)	78.67	1.88	41.887	0.000
sexM	17.80	2.68	6.648	0.000
diethf	34.64	2.75	12.615	0.000
sexM:diethf	-1.86	3.94	-0.474	0.636

```
chol.lm <- lm(chol2~sex+diet, data=phenotype.data)
chol.lm.hf <- lm(chol2~sex, data=filter(phenotype.data, diet=='hf'))

cholesterol.data.hfd <-
  phenotype.data %>%
    filter(!is.na(chol2)) %>%
    filter(diet=='hf') %>%
    mutate(adj.chol.hf=residuals(chol.lm.hf)+coefficients(chol.lm.hf)['(Intercept)'])
```

5 TWAS with Cholesterol Levels for NCD

```
library("org.Mm.eg.db")

library(purrr)
possible.lm <- possibly(.f = lm, otherwise=NULL) # to catch errors when we only have one sex and a cont

twas.data.ncd <-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.ncd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.ncd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.ncd, by='mouse.id') %>%
  dplyr::select(chol2, sex, mouse.id, ENSEMBL.ID, expression) %>%
  group_by(ENSEMBL.ID) %>%
  group_modify(~ broom::tidy(possible.lm(expression ~ chol2+sex, data = .x))) %>%
  filter(term=='chol2') %>%
  arrange(p.value) %>%
  mutate(p.adj=p.adjust(p.value, method="BH"))

twas.data.ncd.all <-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.ncd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.ncd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.ncd, by='mouse.id') %>%
  dplyr::select(chol2, sex, mouse.id, ENSEMBL.ID, expression) %>%
  group_by(ENSEMBL.ID) %>%
```

```

group_modify(~ broom::tidy(possible.lm(expression ~ chol2+sex, data = .x))) %>%
filter(term %in% c('chol2', '(Intercept)')) %>%
dplyr::select(ENSEMBL.ID,term,estimate,std.error) %>%
pivot_wider(id_cols=ENSEMBL.ID,
             names_from = 'term',
             values_from = c(estimate,std.error)) %>%
mutate(estimate.rel = estimate_chol2/`estimate_(Intercept)`,
       std.error.rel = std.error_chol2/`estimate_(Intercept)`)

twas.data.ncd.r2 <-
expression.data %>%
dplyr::select(ENSEMBL.ID,one_of(cholesterol.data.ncd$mouse.id))%>%
pivot_longer(cols=one_of(cholesterol.data.ncd$mouse.id),
             names_to='mouse.id',
             values_to='expression') %>%
full_join(cholesterol.data.ncd,by='mouse.id') %>%
group_by(ENSEMBL.ID) %>%
group_modify(~ broom::glance(possible.lm(expression ~ chol2+sex, data = .x))) %>%
arrange(p.value)%>%
mutate(p.adj=p.adjust(p.value,method="BH"))

twas.data.ncd.int <-
expression.data %>%
dplyr::select(ENSEMBL.ID,one_of(cholesterol.data.ncd$mouse.id))%>%
pivot_longer(cols=one_of(cholesterol.data.ncd$mouse.id),
             names_to='mouse.id',
             values_to='expression') %>%
full_join(cholesterol.data.ncd,by='mouse.id') %>%
group_by(ENSEMBL.ID) %>%
group_modify(~ broom::tidy(possible.lm(expression ~ chol2*sex, data = .x))) %>%
filter(term=='chol2:sexM') %>%
arrange(p.value) %>%
mutate(p.adj=p.adjust(p.value,method="BH"))

twas.data.ncd$symbol <- mapIds(org.Mm.eg.db,
                             keys=twas.data.ncd$ENSEMBL.ID,
                             column="SYMBOL",
                             keytype="ENSEMBL",
                             multiVals="first")

twas.data.ncd.int$symbol <- mapIds(org.Mm.eg.db,
                                  keys=twas.data.ncd.int$ENSEMBL.ID,
                                  column="SYMBOL",
                                  keytype="ENSEMBL",
                                  multiVals="first")

twas.data.ncd.all$symbol <- mapIds(org.Mm.eg.db,
                                  keys=twas.data.ncd.all$ENSEMBL.ID,
                                  column="SYMBOL",
                                  keytype="ENSEMBL",
                                  multiVals="first")

```

```
twas.data.ncd %>%
  head(10) %>%
  kable(caption="Top 10 liver TWAS associations with cholesterol levels")
```

Table 5: Top 10 liver TWAS associations with cholesterol levels

ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG00000089943	chol2	0.279	0.067	4.18	0.000	0.000	Ugt1a5
ENSMUSG00000006567	chol2	0.044	0.013	3.52	0.001	0.001	Atp7b
ENSMUSG00000028412	chol2	0.078	0.023	3.33	0.001	0.001	Slc44a1
ENSMUSG00000024981	chol2	0.326	0.100	3.27	0.002	0.002	Acs15
ENSMUSG00000025809	chol2	-0.054	0.016	-3.26	0.002	0.002	Itgb1
ENSMUSG00000083863	chol2	1.847	0.572	3.23	0.002	0.002	NA
ENSMUSG00000026238	chol2	-0.078	0.024	-3.20	0.002	0.002	Ptma
ENSMUSG00000049940	chol2	0.048	0.016	3.09	0.003	0.003	Pgrmc2
ENSMUSG00000024066	chol2	0.105	0.034	3.08	0.003	0.003	Xdh
ENSMUSG00000090555	chol2	-1.371	0.446	-3.08	0.003	0.003	Gm8893

```
twas.data.ncd.int %>%
  head(10) %>%
  kable(caption="Top 10 liver TWAS associations with cholesterol levels that are modified by sex")
```

Table 6: Top 10 liver TWAS associations with cholesterol levels that are modified by sex

ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG00000071644	chol2:sexM	-0.146	0.029	-5.06	0	0	Eef1g
ENSMUSG00000024120	chol2:sexM	-0.179	0.040	-4.43	0	0	Lrp1rc
ENSMUSG00000021595	chol2:sexM	-0.112	0.026	-4.29	0	0	Nsun2
ENSMUSG00000026895	chol2:sexM	-0.053	0.012	-4.29	0	0	Ndufa8
ENSMUSG00000010608	chol2:sexM	-0.069	0.017	-4.15	0	0	Rbm25
ENSMUSG00000069874	chol2:sexM	0.221	0.054	4.12	0	0	Irgm2
ENSMUSG00000034422	chol2:sexM	0.128	0.031	4.08	0	0	Parp14
ENSMUSG00000031877	chol2:sexM	-0.081	0.021	-3.85	0	0	Ces2g
ENSMUSG00000050043	chol2:sexM	-0.072	0.019	-3.80	0	0	Tmx2
ENSMUSG00000021610	chol2:sexM	-0.082	0.022	-3.79	0	0	Clptm11

```
write_csv(twas.data.ncd,
  file="NCD TWAS Results.csv")
write_csv(twas.data.ncd.int,
  file="NCD TWAS Interaction Results.csv")

twas.data.ncd.combined <-
  left_join(twas.data.ncd,
    filter(twas.data.ncd.int, p.value<0.05), #only append interaction values when significant
    by=c('ENSEMBL.ID','symbol'),
    suffix = c('_main','_int'))

write_csv(twas.data.ncd.combined,
  file="NCD TWAS Results - Combined.csv")
```

6 TWAS with Cholesterol Levels for HFD

```
twas.data.hf <-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.hfd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.hfd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.hfd, by='mouse.id') %>%
  group_by(ENSEMBL.ID) %>%
  group_modify(~ broom::tidy(possible.lm(expression ~ chol2+sex, data = .x))) %>%
  filter(term=='chol2') %>%
  arrange(p.value) %>%
  mutate(p.adj=p.adjust(p.value, method="BH"))

twas.data.hf.all<-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.hfd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.hfd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.hfd, by='mouse.id') %>%
  group_by(ENSEMBL.ID) %>%
  group_modify(~ broom::tidy(possible.lm(expression ~ chol2+sex, data = .x))) %>%
  filter(term %in% c('chol2', '(Intercept)')) %>%
  dplyr::select(ENSEMBL.ID, term, estimate, std.error) %>%
  pivot_wider(id_cols=ENSEMBL.ID,
              names_from = 'term',
              values_from = c(estimate, std.error)) %>%
  mutate(estimate.rel = estimate_chol2/`estimate_(Intercept)`*100,
         std.error.rel = std.error_chol2/`estimate_(Intercept)`*100)

twas.data.hf.r2 <-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.hfd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.hfd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.hfd, by='mouse.id') %>%
  group_by(ENSEMBL.ID) %>%
  group_modify(~ broom::glance(possible.lm(expression ~ chol2+sex, data = .x))) %>%
  arrange(p.value) %>%
  mutate(p.adj=p.adjust(p.value, method="BH"))

twas.data.int.hf <-
  expression.data %>%
  dplyr::select(ENSEMBL.ID, one_of(cholesterol.data.hfd$mouse.id)) %>%
  pivot_longer(cols=one_of(cholesterol.data.hfd$mouse.id),
               names_to='mouse.id',
               values_to='expression') %>%
  full_join(cholesterol.data.hfd, by='mouse.id') %>%
  group_by(ENSEMBL.ID) %>%
```

```

group_modify(~ broom::tidy(possible.lm(expression ~ chol2*sex, data = .x))) %>%
filter(term=='chol2:sexM') %>%
arrange(p.value) %>%
mutate(p.adj=p.adjust(p.value,method="BH"))

twas.data.hf$symbol <- mapIds(org.Mm.eg.db,
                             keys=twas.data.hf$ENSEMBL.ID,
                             column="SYMBOL",
                             keytype="ENSEMBL",
                             multiVals="first")

twas.data.int.hf$symbol <- mapIds(org.Mm.eg.db,
                                  keys=twas.data.int.hf$ENSEMBL.ID,
                                  column="SYMBOL",
                                  keytype="ENSEMBL",
                                  multiVals="first")

twas.data.hf.all$symbol <- mapIds(org.Mm.eg.db,
                                  keys=twas.data.hf.all$ENSEMBL.ID,
                                  column="SYMBOL",
                                  keytype="ENSEMBL",
                                  multiVals="first")

twas.data.hf %>%
  head(10) %>%
  kable(caption="Top 10 liver TWAS associations with cholesterol levels for HFD")

```

Table 7: Top 10 liver TWAS associations with cholesterol levels for HFD

ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG00000038366	chol2	-0.044	0.014	-3.21	0.002	0.002	Lasp1
ENSMUSG000000054469	chol2	0.018	0.006	3.19	0.002	0.002	Lclat1
ENSMUSG000000037266	chol2	0.070	0.023	3.10	0.003	0.003	Rsrp1
ENSMUSG000000030662	chol2	0.014	0.004	3.07	0.003	0.003	Ipo5
ENSMUSG000000025003	chol2	0.309	0.103	3.01	0.003	0.003	Cyp2c39
ENSMUSG000000000631	chol2	0.036	0.012	2.94	0.004	0.004	Myo18a
ENSMUSG000000022615	chol2	-0.058	0.020	-2.92	0.004	0.004	Tymp
ENSMUSG000000031762	chol2	0.165	0.057	2.90	0.005	0.005	Mt2
ENSMUSG000000024140	chol2	0.063	0.022	2.88	0.005	0.005	Epas1
ENSMUSG000000079012	chol2	0.235	0.082	2.86	0.005	0.005	Serpina3m

```

twas.data.int.hf %>%
  head(10) %>%
  kable(caption="Top 10 liver TWAS associations with cholesterol levels that are modified by sex for HFD")

```

Table 8: Top 10 liver TWAS associations with cholesterol levels that are modified by sex for HFD

ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG000000087579	chol2:sexM	-0.141	0.038	-3.69	0.000	0.000	Hectd2os

ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG00000060407	chol2:sexM	-0.763	0.222	-3.44	0.001	0.001	Cyp2a12
ENSMUSG00000018102	chol2:sexM	-0.084	0.024	-3.44	0.001	0.001	H2bc4
ENSMUSG00000022010	chol2:sexM	-0.304	0.091	-3.33	0.001	0.001	Tsc22d1
ENSMUSG00000006529	chol2:sexM	-0.670	0.211	-3.18	0.002	0.002	Itih1
ENSMUSG00000026405	chol2:sexM	-0.450	0.149	-3.03	0.003	0.003	C4bp
ENSMUSG00000037780	chol2:sexM	-0.119	0.039	-3.03	0.003	0.003	Mbl1
ENSMUSG00000035540	chol2:sexM	-2.960	0.983	-3.01	0.003	0.003	Gc
ENSMUSG00000038521	chol2:sexM	-0.326	0.112	-2.91	0.005	0.005	C1s1
ENSMUSG00000030695	chol2:sexM	0.034	0.012	2.79	0.007	0.007	Aldoa

```

write_csv(twas.data.hf,
          file="HFD TWAS Results.csv")
write_csv(twas.data.int.hf,
          file="HFD TWAS Interaction Results.csv")

twas.data.combined.hf <-
  left_join(twas.data.hf,
            filter(twas.data.int.hf, p.value<0.05), #only append interaction values when significant
            by=c('ENSEMBL.ID','symbol'),
            suffix = c('_main','_int'))

write_csv(twas.data.combined.hf,
          file="HFD TWAS Results - Combined.csv")

sig.twas.data.hf <-
  twas.data.hf %>%
  filter(p.adj<0.05)

sig.twas.data.ncd <-
  twas.data.ncd %>%
  filter(p.adj<0.05)

sig.twas.data.ncd$symbol %in% sig.twas.data.hf$symbol %>% table

## .
## FALSE TRUE
## 106 7

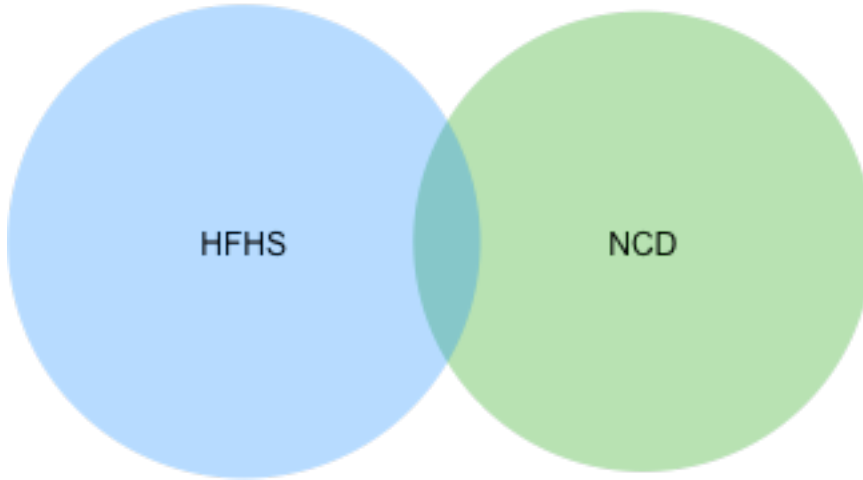
sig.twas.data.ncd$genes %in% sig.twas.data.hf$genes %>% table

## < table of extent 0 >

library(venneuler)
vd.all <- venneuler(c("NCD"=dim(sig.twas.data.ncd)[1],
                      "HFHS"=dim(sig.twas.data.hf)[1],
                      "NCD&HFHS"=intersect(sig.twas.data.hf$symbol,sig.twas.data.ncd$symbol) %>% length),
                    main="Transcripts Associating with Cholesterol")

```

Transcripts Associating with Cholesterol



Our analysis identified **113** nominally significant associations between expression of genes and adjusted cholesterol levels on a normal chow diet. Among those, 68 were positively correlated and 68 were negatively correlated with cholesterol levels.

6.1 TWAS Modification by Sex

By modeling the interactions between sex and expression on cholesterol levels, we identified **526** genes where the cholesterol/expression relationship was modified by sex in a nominally significant manner. This included 14 genes where the relationship was stronger in males, and 512 where it was stronger in females.

7 Comparason with mouse GWAS

```
chr11.genes <- c('Znrf3', 'Xbp1', 'Ccdc117', 'Ankrd36', 'Mrps24', 'Urgcp', 'Dbnl', 'Pgsm2', 'Ael', 'Polm', 'Ael')
genes.of.interest <- c('Cyp7a1', 'Fasn', 'Ldlr', 'Hmgcr')
twas.data.ncd %>%
  filter(symbol %in% c(chr11.genes, genes.of.interest)) %>%
  kable(caption="Genes in the chromosome 11 interval with liver expression")
```

Table 9: Genes in the chromosome 11 interval with liver expression

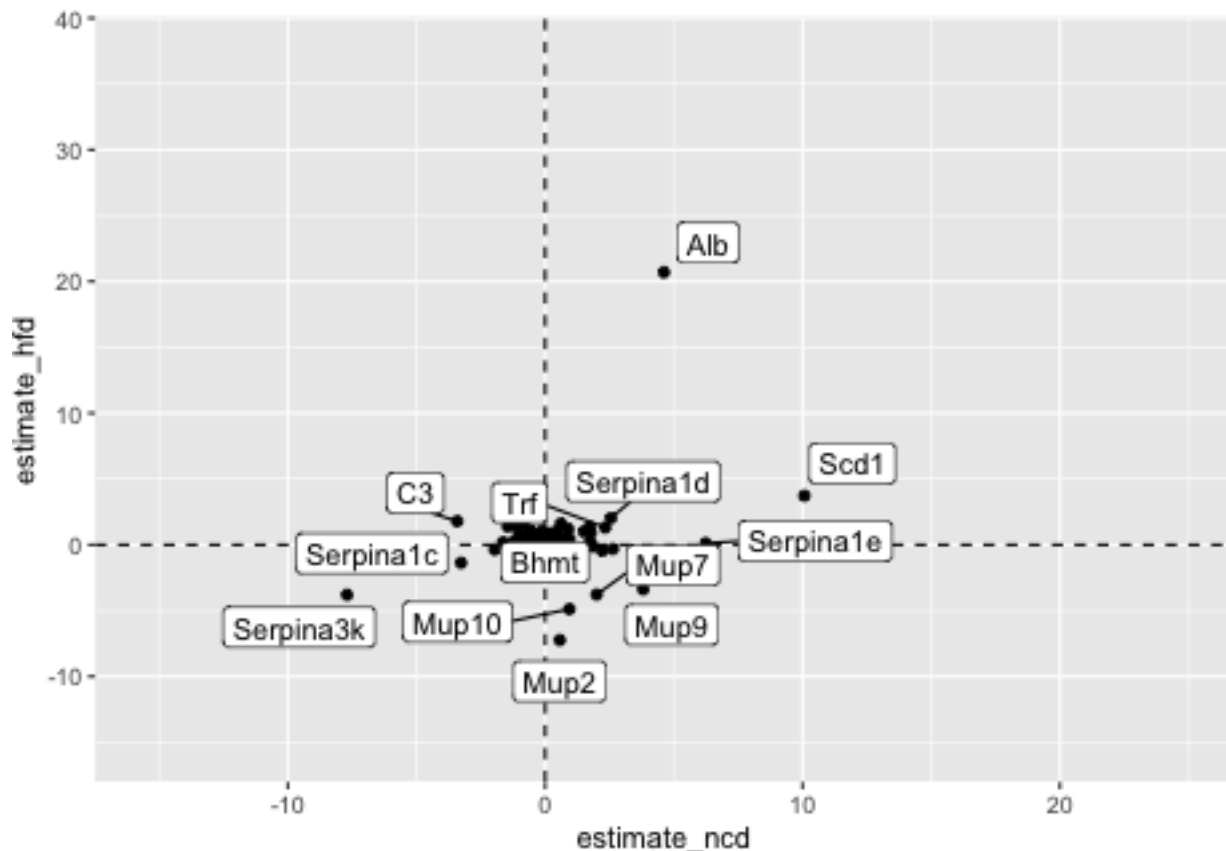
ENSEMBL.ID	term	estimate	std.error	statistic	p.value	p.adj	symbol
ENSMUSG00000002741	chol2	-0.014	0.008	-1.699	0.093	0.093	Ykt6
ENSMUSG00000004394	chol2	-0.008	0.006	-1.381	0.170	0.170	Tmed4
ENSMUSG000000041798	chol2	0.055	0.053	1.027	0.307	0.307	Gck
ENSMUSG000000020484	chol2	-0.064	0.079	-0.812	0.419	0.419	Xbp1
ENSMUSG000000021670	chol2	0.050	0.091	0.556	0.580	0.580	Hmgcr
ENSMUSG000000028240	chol2	-0.049	0.138	-0.355	0.724	0.724	Cyp7a1
ENSMUSG000000032193	chol2	-0.004	0.043	-0.090	0.928	0.928	Ldlr
ENSMUSG000000025153	chol2	0.050	0.636	0.078	0.938	0.938	Fasn

8 Integrated TWAS Analysis

```
combined.twas.data <-
  full_join(twas.data.ncd, twas.data.hf, by=c('ENSEMBL.ID','term','symbol'), suffix=c('_ncd','_hfd'))

combined.twas.data.all <-
  full_join(twas.data.ncd.all, twas.data.hf.all, by=c('ENSEMBL.ID','symbol'), suffix=c('_ncd','_hfd'))

library(ggplot2)
library(ggrepel)
ggplot(combined.twas.data,
  aes(y=estimate_hfd, x=estimate_ncd,
      xmin=estimate_ncd-std.error_ncd, xmax=estimate_ncd+std.error_ncd,
      ymin=estimate_hfd-std.error_hfd, ymax=estimate_hfd+std.error_hfd )) +
  geom_point() +
  #geom_errorbar() +
  #geom_errorbarh() +
  geom_hline(yintercept = 0, lty=2) +
  geom_vline(xintercept = 0, lty=2) +
  geom_label_repel(aes(label=symbol),
    data = subset(combined.twas.data, abs(estimate_hfd) > 2|abs(estimate_ncd) > 2))
```

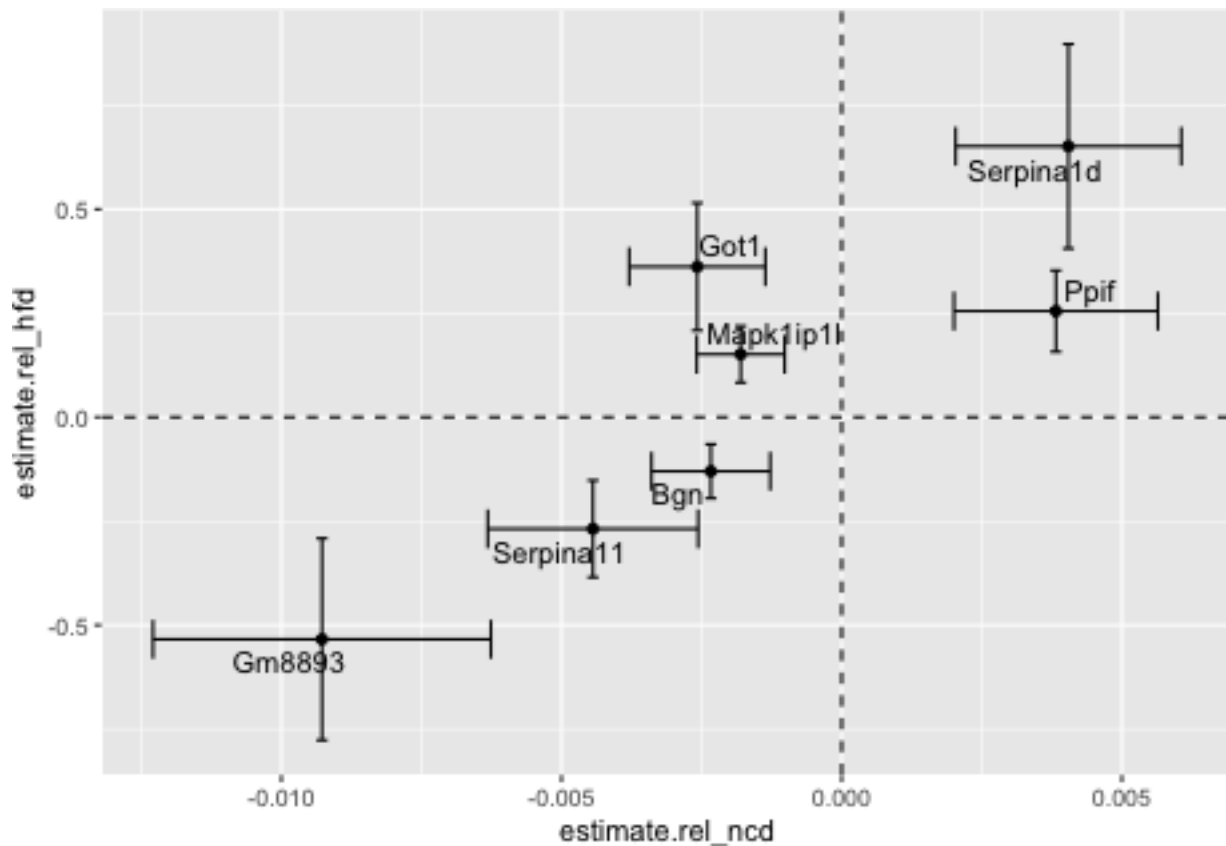


```
sig.genes <- filter(combined.twas.data, p.adj_ncd<0.05&p.adj_hfd<0.05) %>% pull(symbol)
ggplot(combined.twas.data.all %>% filter(symbol %in% sig.genes),
  aes(y=estimate.rel_hfd, x=estimate.rel_ncd,
      xmin=estimate.rel_ncd-std.error.rel_ncd,
```

```

    xmax=estimate.rel_ncd+std.error.rel_ncd,
    ymin=estimate.rel_hfd-std.error.rel_hfd,
    ymax=estimate.rel_hfd+std.error.rel_hfd )) +
geom_point() +
geom_errorbar() +
geom_errorbarh() +
geom_hline(yintercept = 0, lty=2) +
geom_vline(xintercept = 0, lty=2) +
geom_text_repel(aes(label=symbol))

```

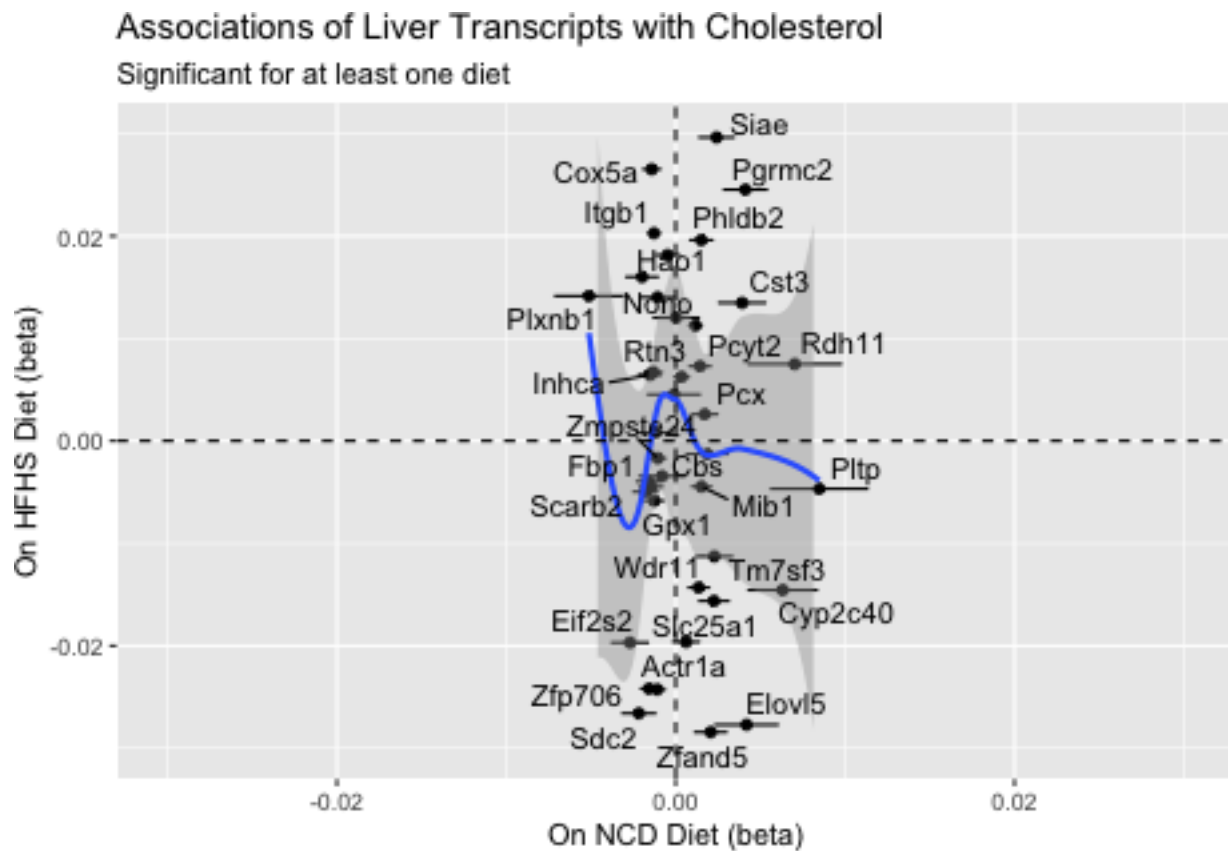


```

sig.genes <- filter(combined.twas.data, p.adj_ncd<0.05|p.adj_hfd<0.05) %>% pull(symbol)
ggplot(combined.twas.data.all %>% filter(symbol %in% sig.genes),
  aes(y=estimate.rel_hfd, x=estimate.rel_ncd,
    xmin=estimate.rel_ncd-std.error.rel_ncd,
    xmax=estimate.rel_ncd+std.error.rel_ncd,
    ymin=estimate.rel_hfd-std.error.rel_hfd,
    ymax=estimate.rel_hfd+std.error.rel_hfd )) +
geom_point() +
geom_errorbar() +
geom_errorbarh() +
geom_smooth() +
geom_hline(yintercept = 0, lty=2) +
geom_vline(xintercept = 0, lty=2) +
xlim(-0.03,0.03) +
ylim(-0.03,0.03) +
geom_text_repel(aes(label=symbol)) +
labs(y="On HFHS Diet (beta)",

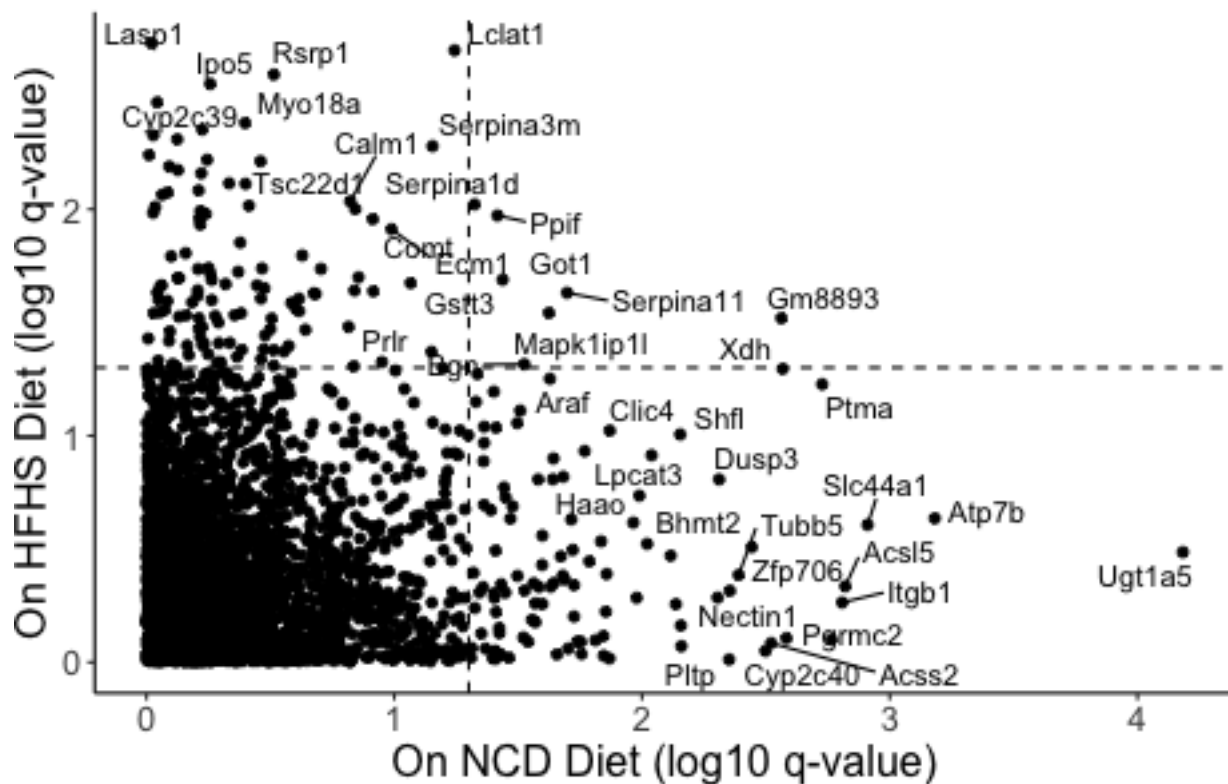
```

```
x="On NCD Diet (beta)",
title="Associations of Liver Transcripts with Cholesterol",
subtitle="Significant for at least one diet")
```



```
cutoff <- -log10(0.05)
ggplot(combined.twas.data,
  aes(y=-log10(p.adj_hfd), x=-log10(p.adj_ncd) )) +
  geom_point() +
  #geom_errorbar() +
  #geom_errorbarh() +
  geom_hline(yintercept = -log10(0.05), lty=2) +
  geom_vline(xintercept = -log10(0.05), lty=2) +
  geom_text_repel(aes(label=symbol),
    data = subset(combined.twas.data, -log10(p.adj_hfd) > cutoff|-log10(p.adj_ncd) > cutoff),
    labs(y="On HFHS Diet (log10 q-value)",
      x="On NCD Diet (log10 q-value)",
      title="Associations of Liver Transcripts with Cholesterol") +
  theme_classic() +
  theme(text=element_text(size=16))
```

Associations of Liver Transcripts with Cholesterol



Comparason with human GWAS

Downloaded human cholesterol associated alleles from <https://t2d.hugeamp.org/phenotype.html?phenotype=CHOL>

```
“{ chol-human-gwas} gwas.filename <- ‘cholesterol-associations.csv’ gwas.data <- read_csv(gwas.filename)
%>% mutate(Symbol=gsub(‘{2}$’, ‘’, nearest)) %>% mutate(Symbol=gsub(“^{0,2}”, “”, Symbol)) #re-
moves first and last characters
```

```
library(biomaRt) human = useMart(“ensembl”, dataset = “hsapiens_gene_ensembl”) mouse = use-
Mart(“ensembl”, dataset = “mmusculus_gene_ensembl”)
```

```
mapping.data <- getLDS(attributes = c(“hgnc_symbol”), filters = “hgnc_symbol”, values =
gwas.data$Symbol, mart = human, attributesL = c(“mgi_symbol”), martL = mouse, uniqueRows=T)
```

```
gwas.data <- full_join(gwas.data, mapping.data, by=c(‘Symbol’=‘HGNC.symbol’)) %>% dplyr::filter(!(is.na(MGI.symbol)))
```

#are gwas alleles enriched in correlation analyses

```
sig.twas.data <- filter(twas.data, p.value<0.05) #sig.twas.datasymbolMGI.symbol %>% table
```

```
combined.twas.data %>% filter(symbol %in% gwas.data$MGI.symbol) %>% arrange(p.value) %>% head
%>% kable(caption=“Most significant TWAS association hits that are also nearby GWAS hits”)
```

```
#checked if a TWAS was a GWAS hit twas.data.matched <- combined.twas.data %>% mu-
tate(hGWAS.match=symbol %in% gwas.data$MGI.symbol)
```

```
with(twas.data.matched, table(hGWAS.match,p.value<0.05)) %>% fisher.test() %>% tidy %>%
kable(caption=‘Fisher test for enrichment of GWAS hits in liver TWAS genes’)
```

```
glm(hGWAS.match~p.value, data=twas.data.matched, family=‘binomial’) %>% tidy %>% kable(caption=“Logistic
regression of TWAS values against likilihood of a GWAS hit.”)
```

```

# Pathway Analyses for NCD

```r
twas.list <- twas.data.ncd %>% arrange(-estimate) %>% pull(estimate)
names(twas.list) <- twas.data.ncd %>% arrange(-estimate) %>% pull(symbol)
twas.list <- sort(twas.list, decreasing = TRUE)
#twas.list <- twas.list[!(is.na(names(twas.list)))]

library(clusterProfiler)
go.twas.bp <- gseGO(geneList=twas.list,
 ont="BP",
 keyType='SYMBOL',
 OrgDb=org.Mm.eg.db,
 pvalueCutoff=0.25,
 verbose=T,
 by='fgsea',
 eps=1E-25)

#enrichement
twas.data.ncd %>%
 filter(p.value<0.05) %>%
 pull(symbol) ->
 twas.sig

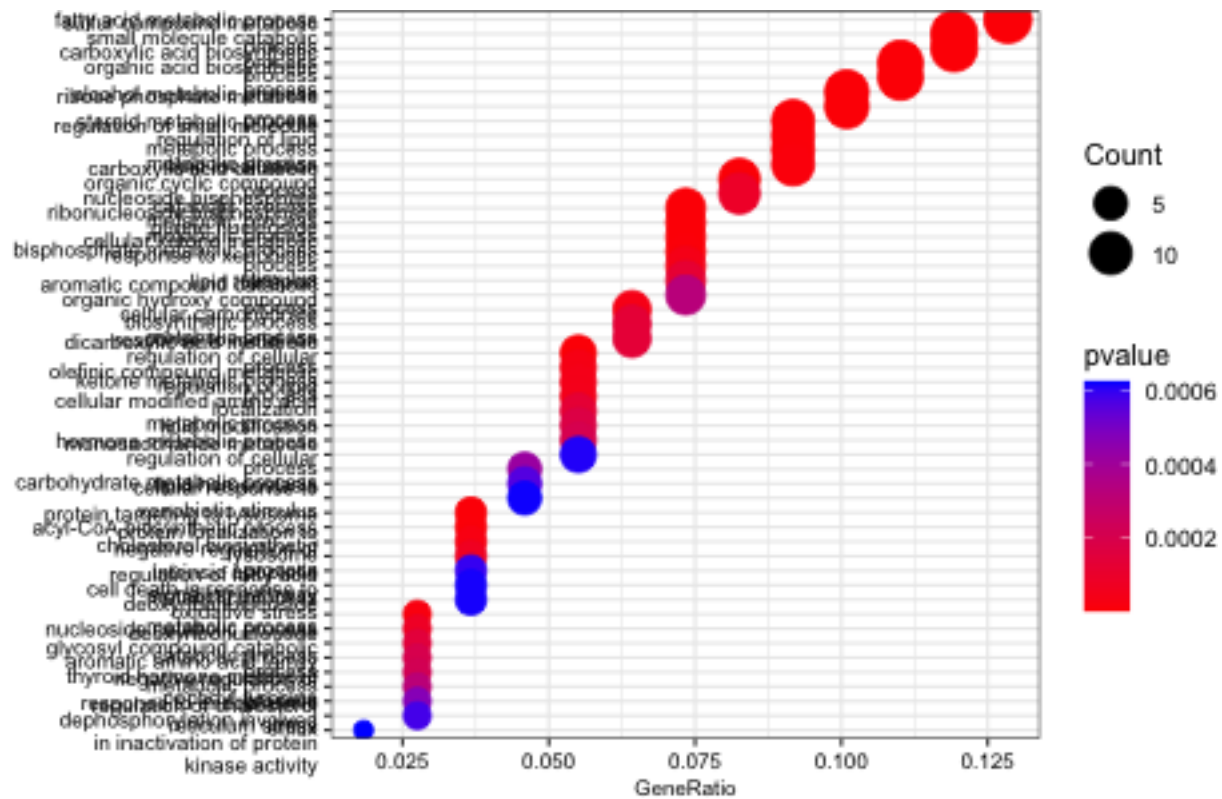
go.twas.bp.enrich <- enrichGO(gene=twas.sig,
 ont="BP",
 keyType='SYMBOL',
 OrgDb=org.Mm.eg.db,
 pvalueCutoff=0.05)

go.twas.bp.enrich.simpl <- simplify(go.twas.bp.enrich,
 cutoff=0.7,
 by="p.adjust",
 select_fun=min)

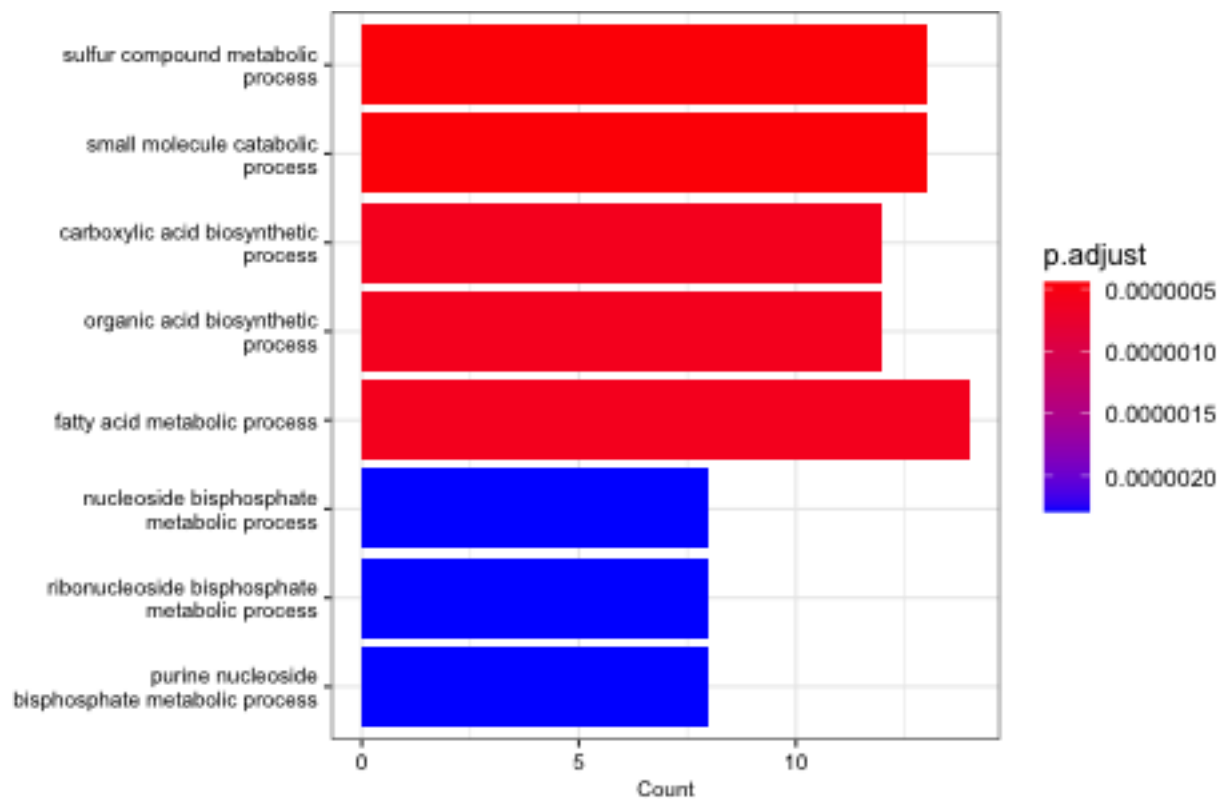
library(enrichplot)

dotplot(go.twas.bp.enrich.simpl,
 showCategory=50,
 color='pvalue',
 font.size=8)

```



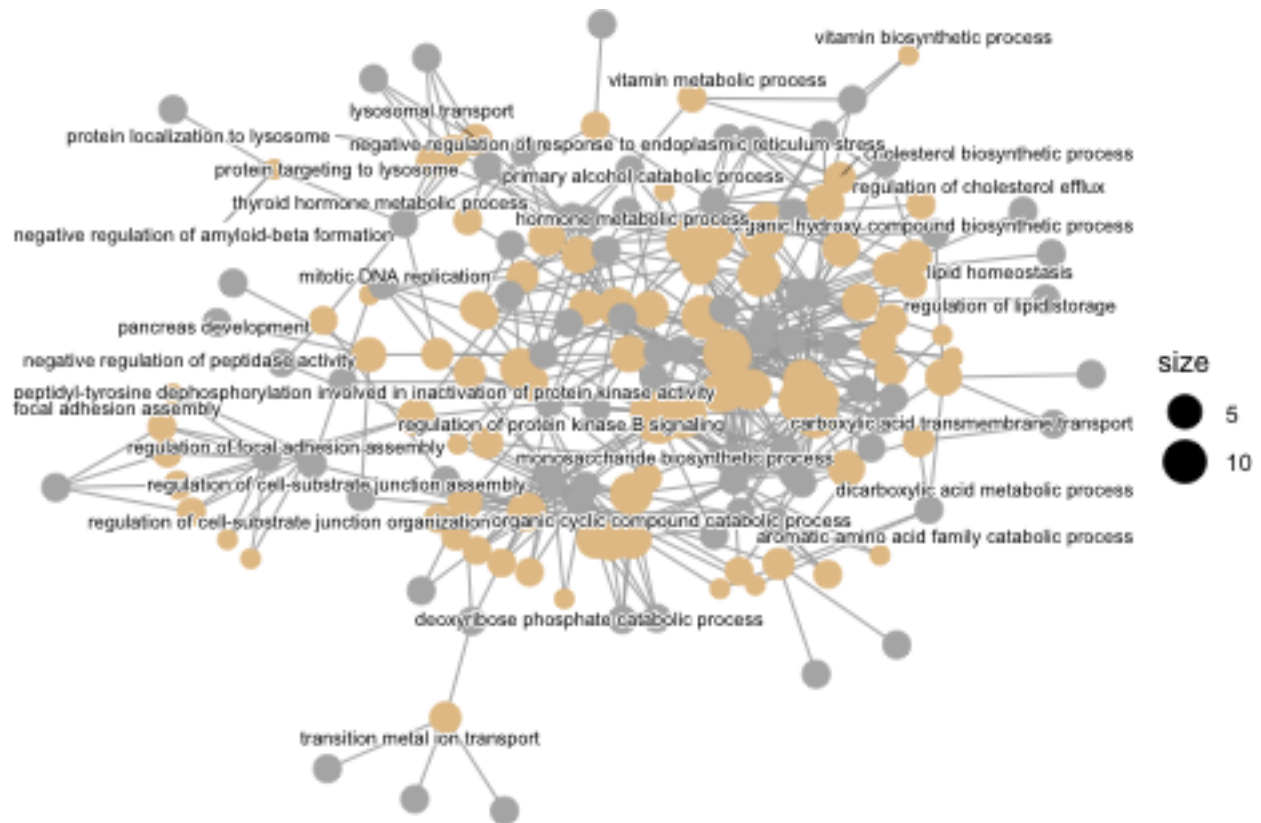
```
barplot(go.twas.bp.enrich.simpl,font.size=8)
```



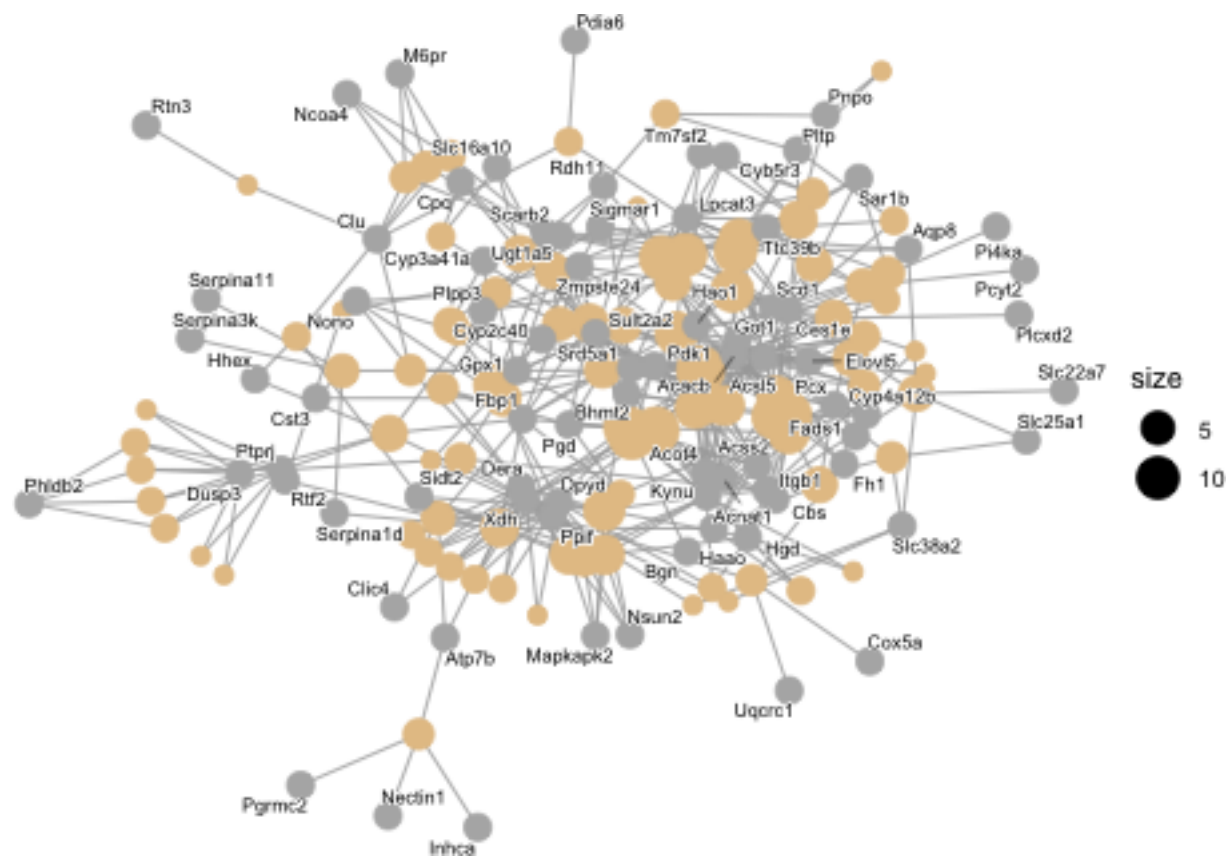
```
cnetplot(go.twas.bp.enrich.simpl,
```



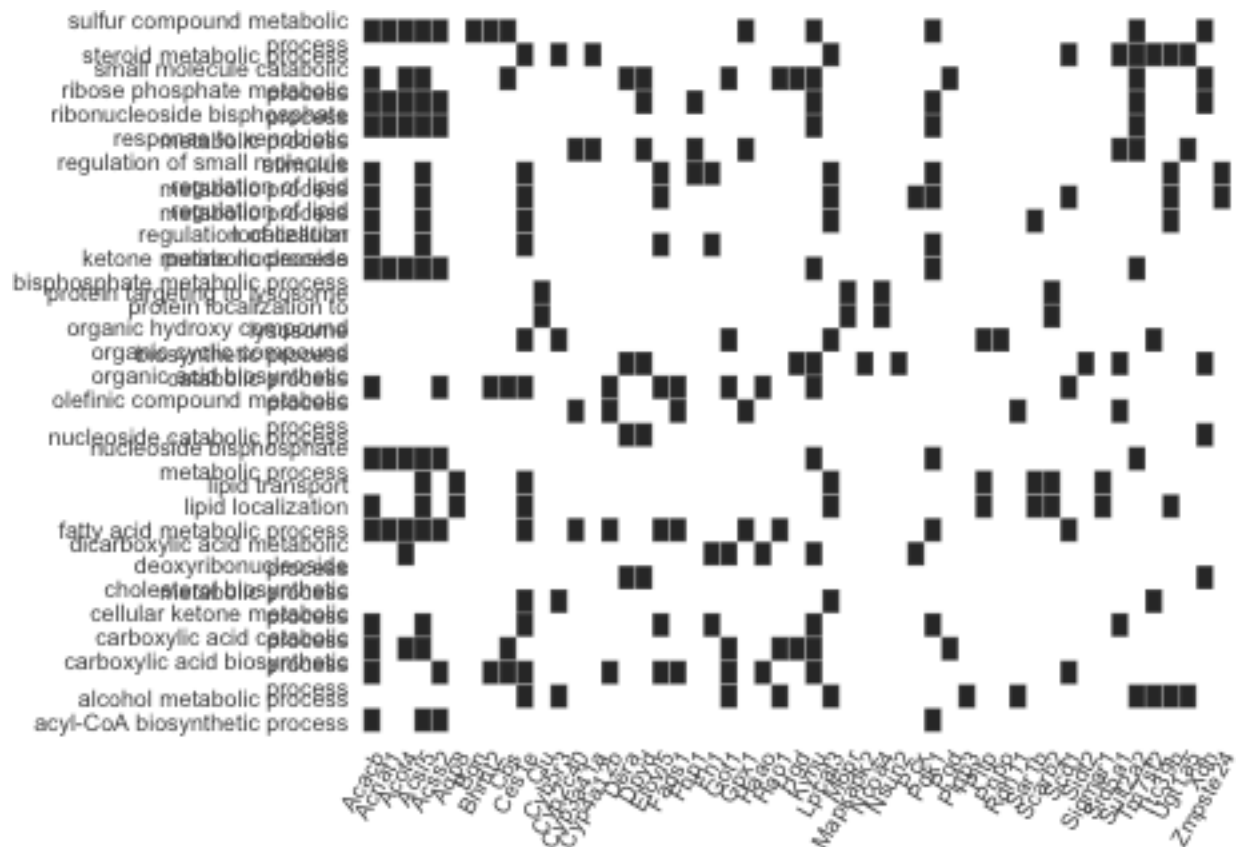
```
showCategory=100,
node_label="gene",
cex_label_gene=0.5)
```



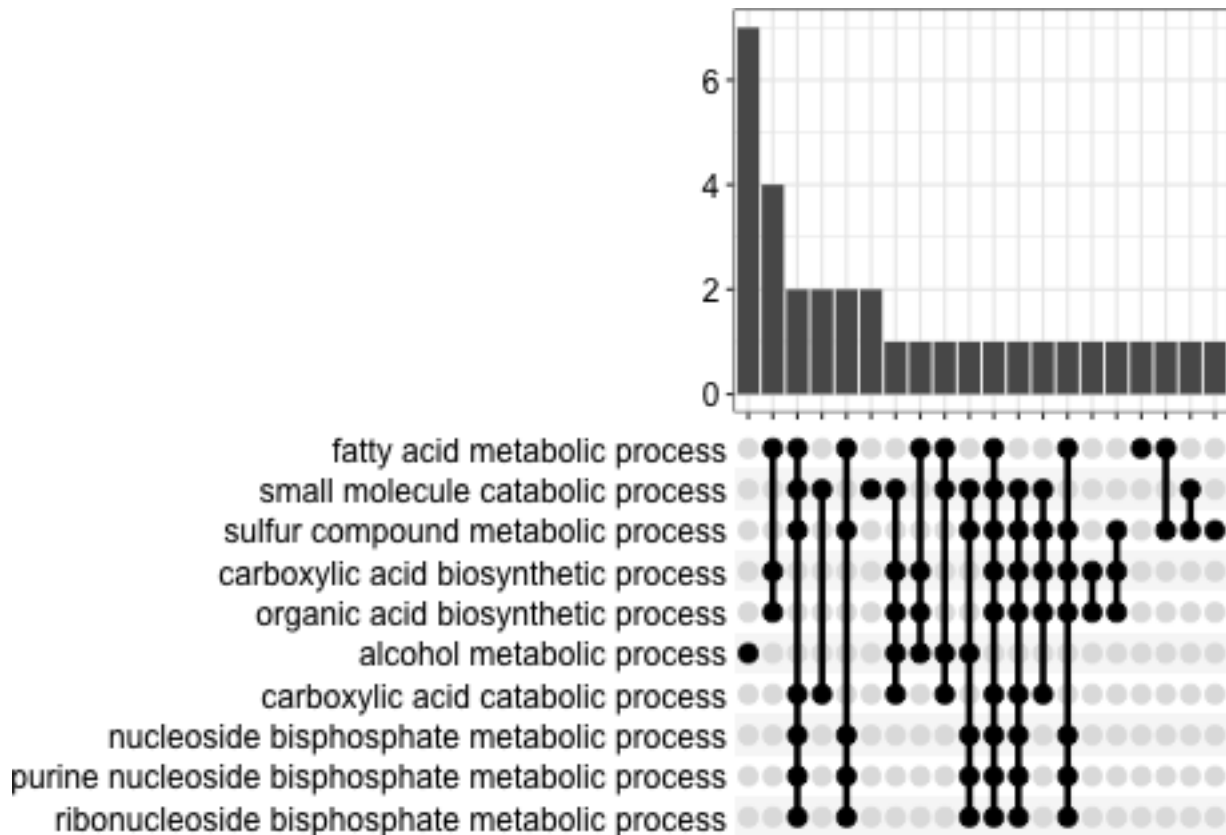
```
cnetplot(go.twas.bp.enrich.simpl,
showCategory=100,
node_label="category",
cex_label_category=0.5)
```



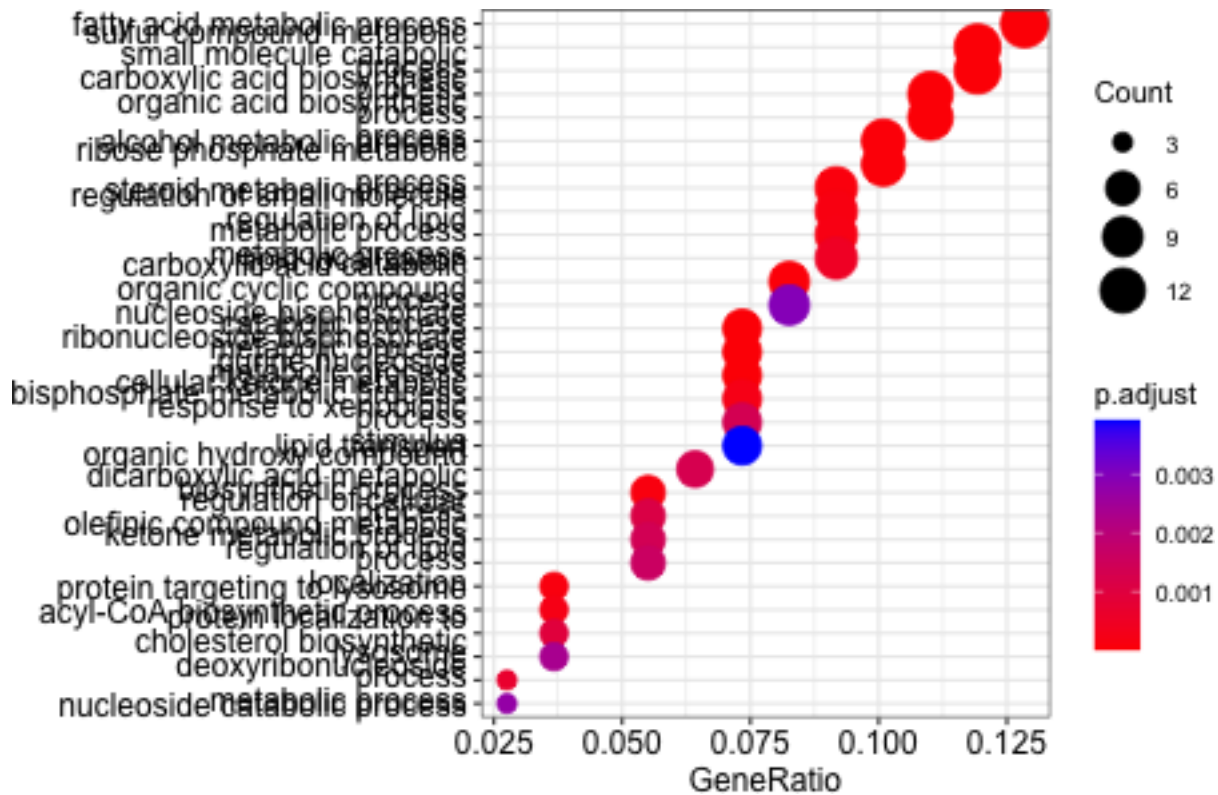
```
heatplot(go.twas.bp.enrich.simpl)
```



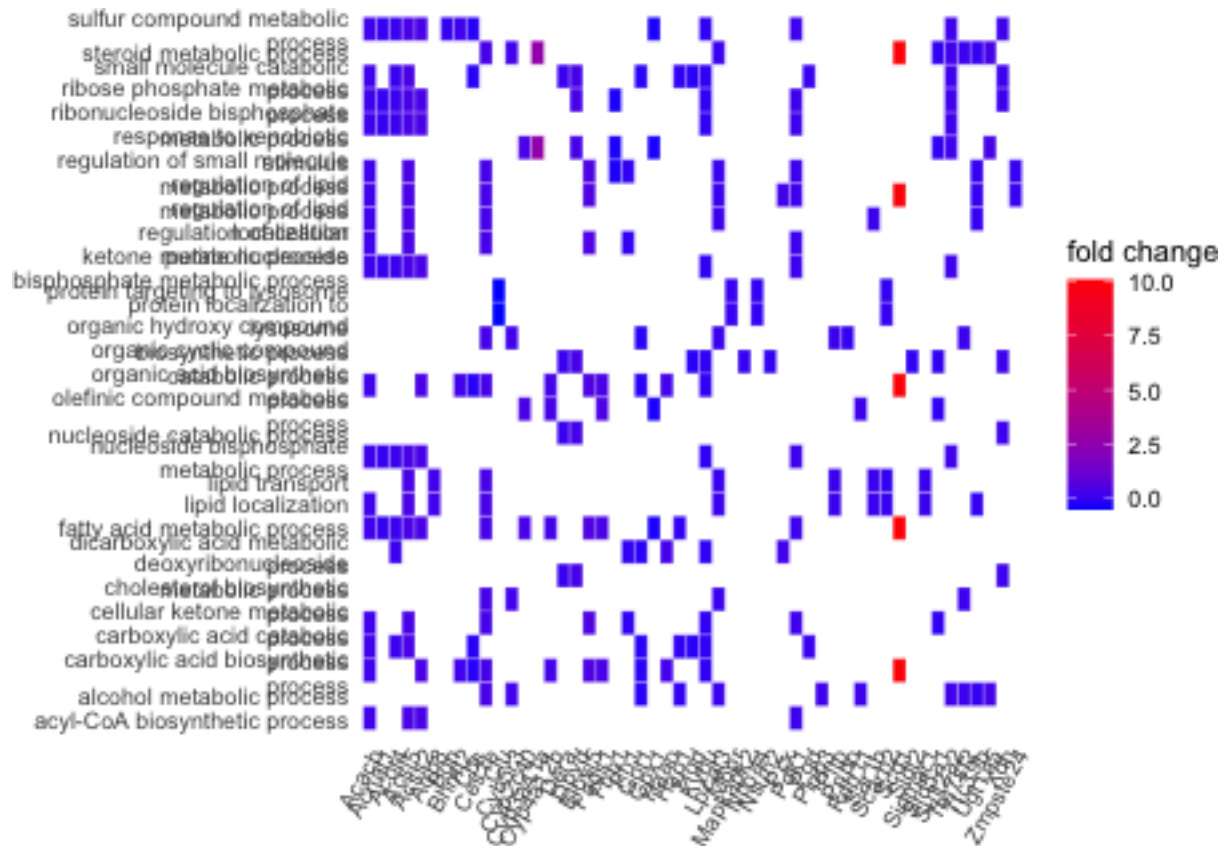
```
library(ggplot2)
upsetplot(go.twas.bp.enrich.simpl)
```



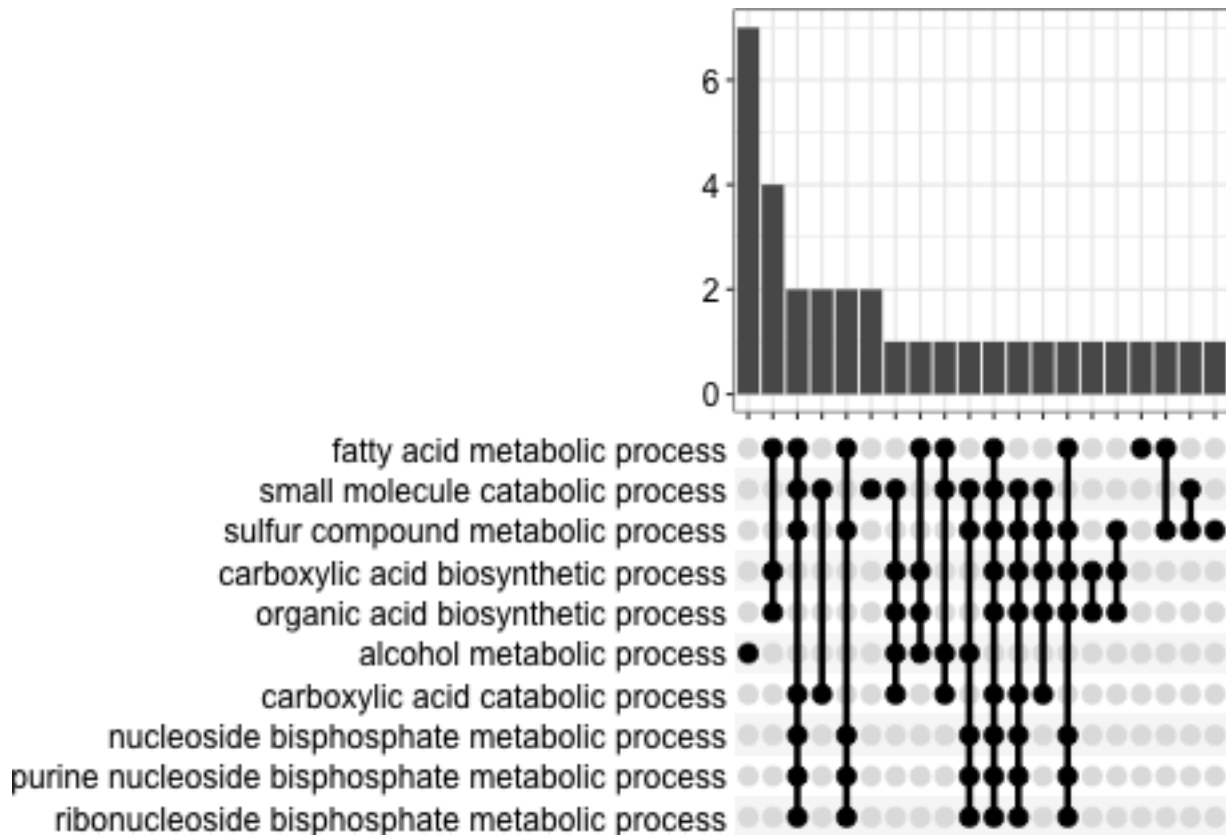
```
dotplot(go.twas.bp.enrich.simpl, showCategory=30)
```



```
heatplot(go.twas.bp.enrich.simpl, foldChange=twas.list)
```



```
upsetplot(go.twas.bp.enrich.simpl)
```



```
as.data.frame(go.twas.bp) %>% select("Description","setSize","enrichmentScore","NES","pvalue","p.adjust")
kable(caption="NCD GO BP Analysis of TWAS Associations from GSEA")
```

Table 10: NCD GO BP Analysis of TWAS Associations from GSEA

Description	setSize	enrichmentScore	NES	pvalue	p.adjust
-------------	---------	-----------------	-----	--------	----------

```
as.data.frame(go.twas.bp) %>%
 write_csv(file="NCD TWAS GO-BP GSEA.csv")

as.data.frame(go.twas.bp.enrich.simpl) %>% select("Description","GeneRatio","Count","pvalue","p.adjust")
kable(caption="NCD GO BP Analysis of TWAS Associations from Enrichment")
```

Table 11: NCD GO BP Analysis of TWAS Associations from Enrichment

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0006790	sulfur compound metabolic process	13/109	13	0.000	0.000
GO: 0044282	small molecule catabolic process	13/109	13	0.000	0.000
GO: 0046394	carboxylic acid biosynthetic process	12/109	12	0.000	0.000
GO: 0016053	organic acid biosynthetic process	12/109	12	0.000	0.000

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0006631	fatty acid metabolic process	14/109	14	0.000	0.000
GO: 0033865	nucleoside bisphosphate metabolic process	8/109	8	0.000	0.000
GO: 0033875	ribonucleoside bisphosphate metabolic process	8/109	8	0.000	0.000
GO: 0034032	purine nucleoside bisphosphate metabolic process	8/109	8	0.000	0.000
GO: 0006066	alcohol metabolic process	11/109	11	0.000	0.000
GO: 0046395	carboxylic acid catabolic process	9/109	9	0.000	0.000
GO: 0019693	ribose phosphate metabolic process	11/109	11	0.000	0.000
GO: 0008202	steroid metabolic process	10/109	10	0.000	0.000
GO: 0043648	dicarboxylic acid metabolic process	6/109	6	0.000	0.000
GO: 0006622	protein targeting to lysosome	4/109	4	0.000	0.000
GO: 0062012	regulation of small molecule metabolic process	10/109	10	0.000	0.000
GO: 0019216	regulation of lipid metabolic process	10/109	10	0.000	0.000
GO: 0071616	acyl-CoA biosynthetic process	4/109	4	0.000	0.000
GO: 0042180	cellular ketone metabolic process	8/109	8	0.000	0.000
GO: 0010876	lipid localization	10/109	10	0.000	0.000
GO: 0009120	deoxyribonucleoside metabolic process	3/109	3	0.000	0.001
GO: 0061462	protein localization to lysosome	4/109	4	0.000	0.001
GO: 0010565	regulation of cellular ketone metabolic process	6/109	6	0.000	0.001
GO: 1901617	organic hydroxy compound biosynthetic process	7/109	7	0.000	0.001
GO: 0009410	response to xenobiotic stimulus	8/109	8	0.000	0.001
GO: 0120254	olefinic compound metabolic process	6/109	6	0.000	0.001
GO: 1905952	regulation of lipid localization	6/109	6	0.000	0.002
GO: 0006695	cholesterol biosynthetic process	4/109	4	0.000	0.002
GO: 0009164	nucleoside catabolic process	3/109	3	0.000	0.003
GO: 1901361	organic cyclic compound catabolic process	9/109	9	0.000	0.003
GO: 0006869	lipid transport	8/109	8	0.000	0.004

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0006575	cellular modified amino acid metabolic process	6/109	6	0.000	0.004
GO: 0044262	cellular carbohydrate metabolic process	7/109	7	0.000	0.004
GO: 0010038	response to metal ion	7/109	7	0.000	0.004
GO: 0009264	deoxyribonucleotide catabolic process	3/109	3	0.000	0.004
GO: 0030258	lipid modification	6/109	6	0.000	0.005
GO: 1901658	glycosyl compound catabolic process	3/109	3	0.000	0.005
GO: 0042445	hormone metabolic process	6/109	6	0.000	0.005
GO: 0009072	aromatic amino acid family metabolic process	3/109	3	0.000	0.006
GO: 0042403	thyroid hormone metabolic process	3/109	3	0.000	0.008
GO: 0019439	aromatic compound catabolic process	8/109	8	0.000	0.008
GO: 0010675	regulation of cellular carbohydrate metabolic process	5/109	5	0.000	0.010
GO: 1903573	negative regulation of response to endoplasmic reticulum stress	3/109	3	0.000	0.011
GO: 0055088	lipid homeostasis	5/109	5	0.001	0.013
GO: 0010874	regulation of cholesterol efflux	3/109	3	0.001	0.013
GO: 2001243	negative regulation of intrinsic apoptotic signaling pathway	4/109	4	0.001	0.013
GO: 0005996	monosaccharide metabolic process	6/109	6	0.001	0.013
GO: 0019217	regulation of fatty acid metabolic process	4/109	4	0.001	0.013
GO: 0036473	cell death in response to oxidative stress	4/109	4	0.001	0.013
GO: 0071466	cellular response to xenobiotic stimulus	5/109	5	0.001	0.013
GO: 1990264	peptidyl-tyrosine dephosphorylation involved in inactivation of protein kinase activity	2/109	2	0.001	0.013
GO: 0008631	intrinsic apoptotic signaling pathway in response to oxidative stress	3/109	3	0.001	0.014
GO: 0000041	transition metal ion transport	4/109	4	0.001	0.016
GO: 0034310	primary alcohol catabolic process	2/109	2	0.001	0.017
GO: 0007041	lysosomal transport	4/109	4	0.001	0.019
GO: 0010883	regulation of lipid storage	3/109	3	0.001	0.020
GO: 0044270	cellular nitrogen compound catabolic process	7/109	7	0.001	0.021



	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0046700	heterocycle catabolic process	7/109	7	0.001	0.021
GO: 1902969	mitotic DNA replication	2/109	2	0.001	0.021
GO: 0016042	lipid catabolic process	6/109	6	0.002	0.025
GO: 0008654	phospholipid biosynthetic process	5/109	5	0.002	0.025
GO: 0009074	aromatic amino acid family catabolic process	2/109	2	0.002	0.025
GO: 0051893	regulation of focal adhesion assembly	3/109	3	0.002	0.027
GO: 0090109	regulation of cell-substrate junction assembly	3/109	3	0.002	0.027
GO: 0071243	cellular response to arsenic-containing substance	2/109	2	0.002	0.027
GO: 1905039	carboxylic acid transmembrane transport	4/109	4	0.002	0.028
GO: 0016052	carbohydrate catabolic process	4/109	4	0.002	0.028
GO: 0071241	cellular response to inorganic substance	5/109	5	0.002	0.030
GO: 0007584	response to nutrient	3/109	3	0.002	0.030
GO: 0015711	organic anion transport	6/109	6	0.002	0.031
GO: 0150116	regulation of cell-substrate junction organization	3/109	3	0.002	0.031
GO: 0009110	vitamin biosynthetic process	2/109	2	0.003	0.033
GO: 1902430	negative regulation of amyloid-beta formation	2/109	2	0.003	0.033
GO: 0071404	cellular response to low-density lipoprotein particle stimulus	2/109	2	0.003	0.038
GO: 1901657	glycosyl compound metabolic process	3/109	3	0.003	0.038
GO: 0010466	negative regulation of peptidase activity	5/109	5	0.003	0.039
GO: 0031998	regulation of fatty acid beta-oxidation	2/109	2	0.003	0.040
GO: 0090201	negative regulation of release of cytochrome c from mitochondria	2/109	2	0.003	0.040
GO: 0009060	aerobic respiration	4/109	4	0.004	0.041
GO: 0006766	vitamin metabolic process	3/109	3	0.004	0.044
GO: 0050860	negative regulation of T cell receptor signaling pathway	2/109	2	0.004	0.044
GO: 0042326	negative regulation of phosphorylation	6/109	6	0.004	0.045
GO: 0048041	focal adhesion assembly	3/109	3	0.004	0.045

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0046386	deoxyribose phosphate catabolic process	2/109	2	0.004	0.045
GO: 0046685	response to arsenic-containing substance	2/109	2	0.004	0.045
GO: 0051896	regulation of protein kinase B signaling	4/109	4	0.005	0.047
GO: 0006979	response to oxidative stress	6/109	6	0.005	0.047
GO: 0031016	pancreas development	3/109	3	0.005	0.049
GO: 0046364	monosaccharide biosynthetic process	3/109	3	0.005	0.049
GO: 0042059	negative regulation of epidermal growth factor receptor signaling pathway	2/109	2	0.005	0.049

```
as.data.frame(go.twas.bp.enrich.simpl) %>%
 write_csv(file="NCD TWAS GO-BP Enrichment.csv")

steroid.enrichment <- as.data.frame(go.twas.bp) %>% filter(ID=='GO:0016126')
```

## 9 Pathway Analyses for HFD

```
twas.list <- twas.data.hf %>% arrange(-estimate) %>% pull(estimate)
names(twas.list) <- twas.data.hf %>% arrange(-estimate) %>% pull(symbol)
twas.list <- sort(twas.list, decreasing = TRUE)
#twas.list <- twas.list[!(is.na(names(twas.list)))]

library(clusterProfiler)
go.twas.bp <- gseGO(geneList=twas.list,
 ont="BP",
 keyType='SYMBOL',
 OrgDb=org.Mm.eg.db,
 pvalueCutoff=0.25,
 verbose=T,
 by='fgsea',
 eps=1E-25)

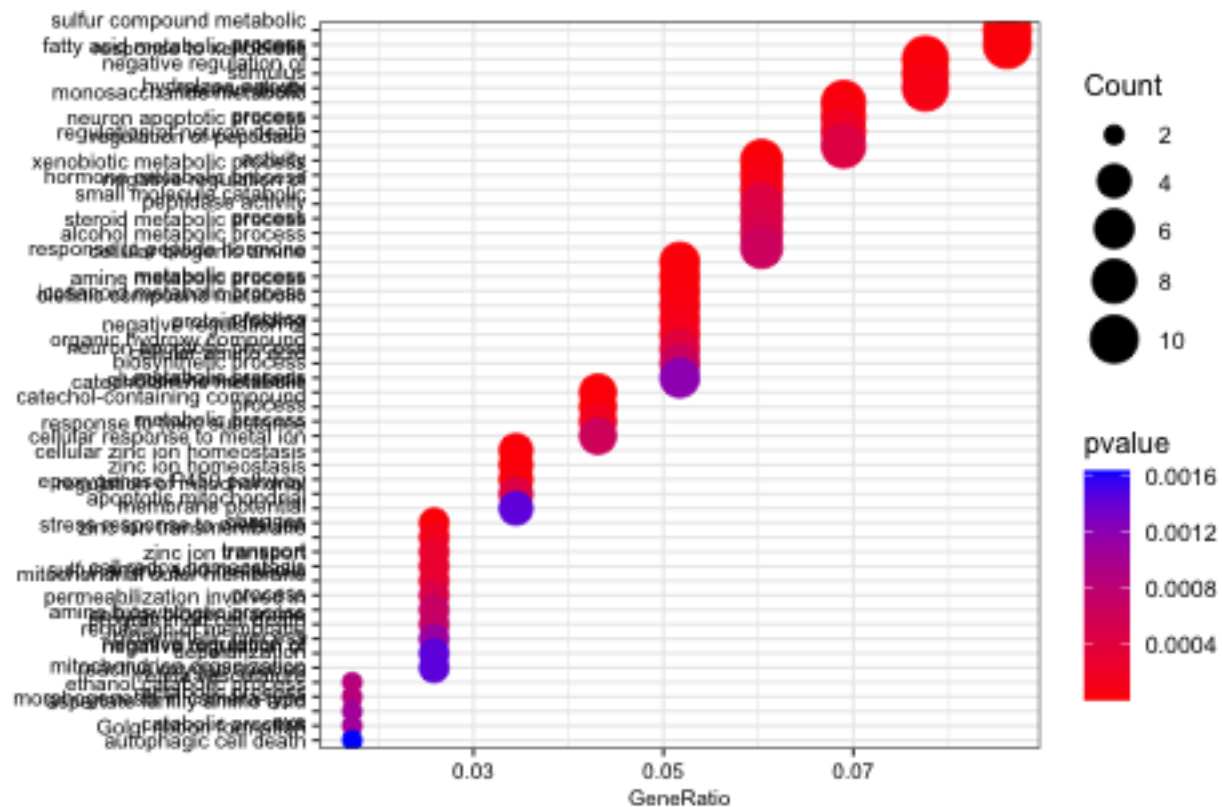
#enrichment
twas.data.hf %>%
 filter(p.value<0.05) %>%
 pull(symbol) ->
 twas.sig

go.twas.bp.enrich <- enrichGO(gene=twas.sig,
 ont="BP",
 keyType='SYMBOL',
 OrgDb=org.Mm.eg.db,
 pvalueCutoff=0.05)
```

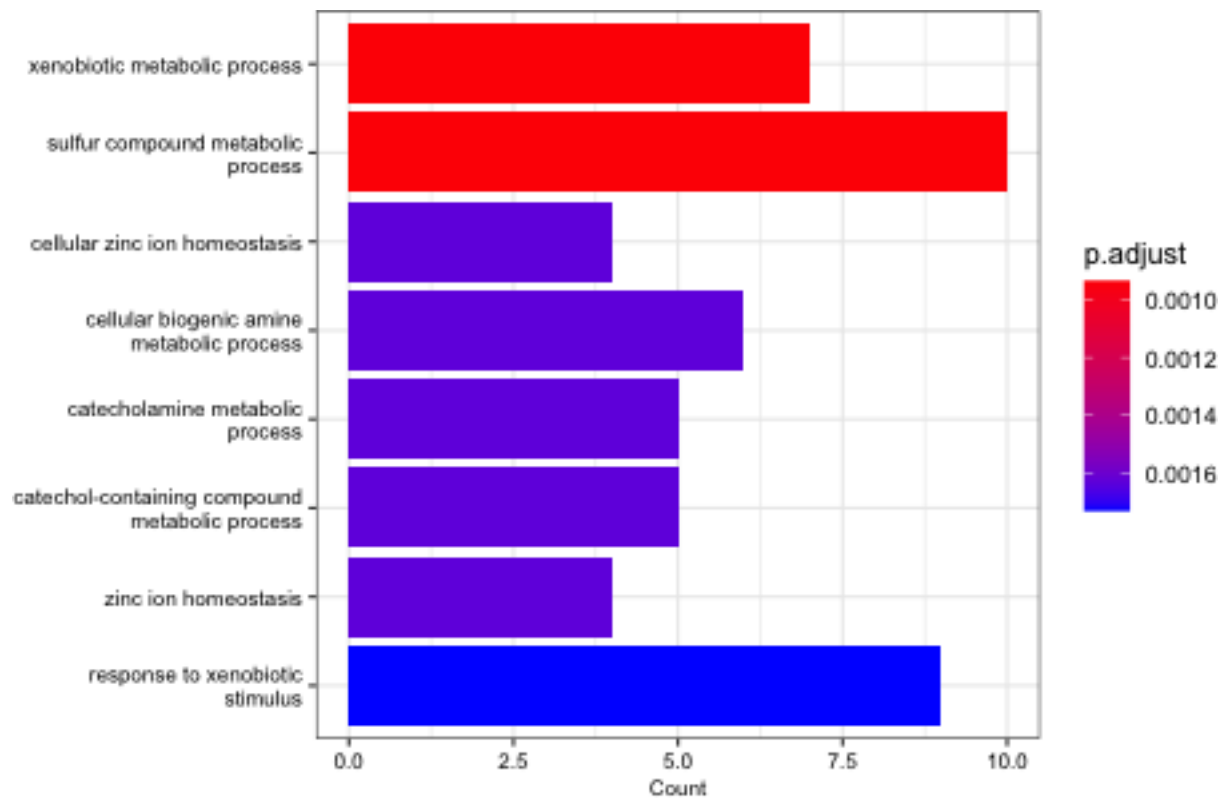
```
go.twas.bp.enrich.simpl <- simplify(go.twas.bp.enrich,
 cutoff=0.7,
 by="p.adjust",
 select_fun=min)
```

```
library(enrichplot)
```

```
dotplot(go.twas.bp.enrich.simpl,
 showCategory=50,
 color='pvalue',
 font.size=8)
```



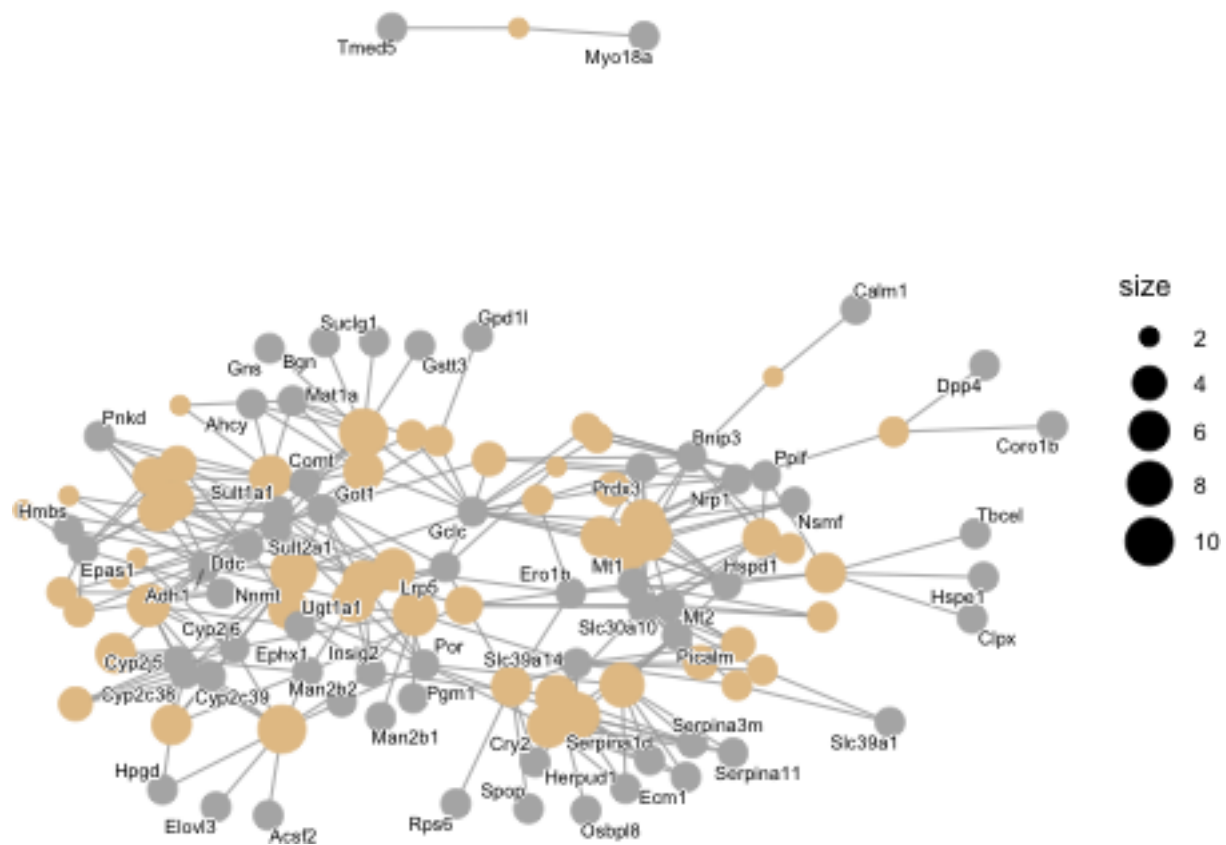
```
barplot(go.twas.bp.enrich.simpl, font.size=8)
```



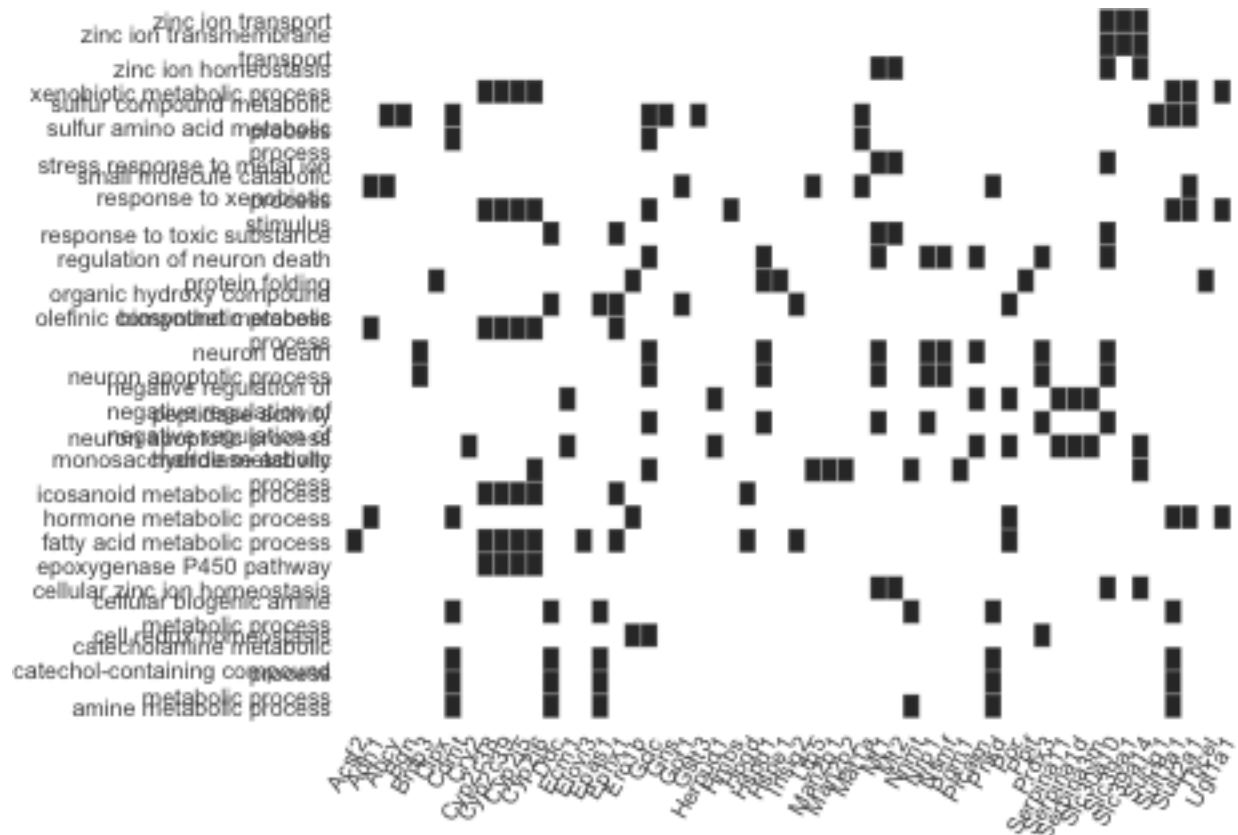
```
cnetplot(go.twas.bp.enrich.simpl,
 showCategory=100,
 node_label="gene",
 cex_label_gene=0.5)
```



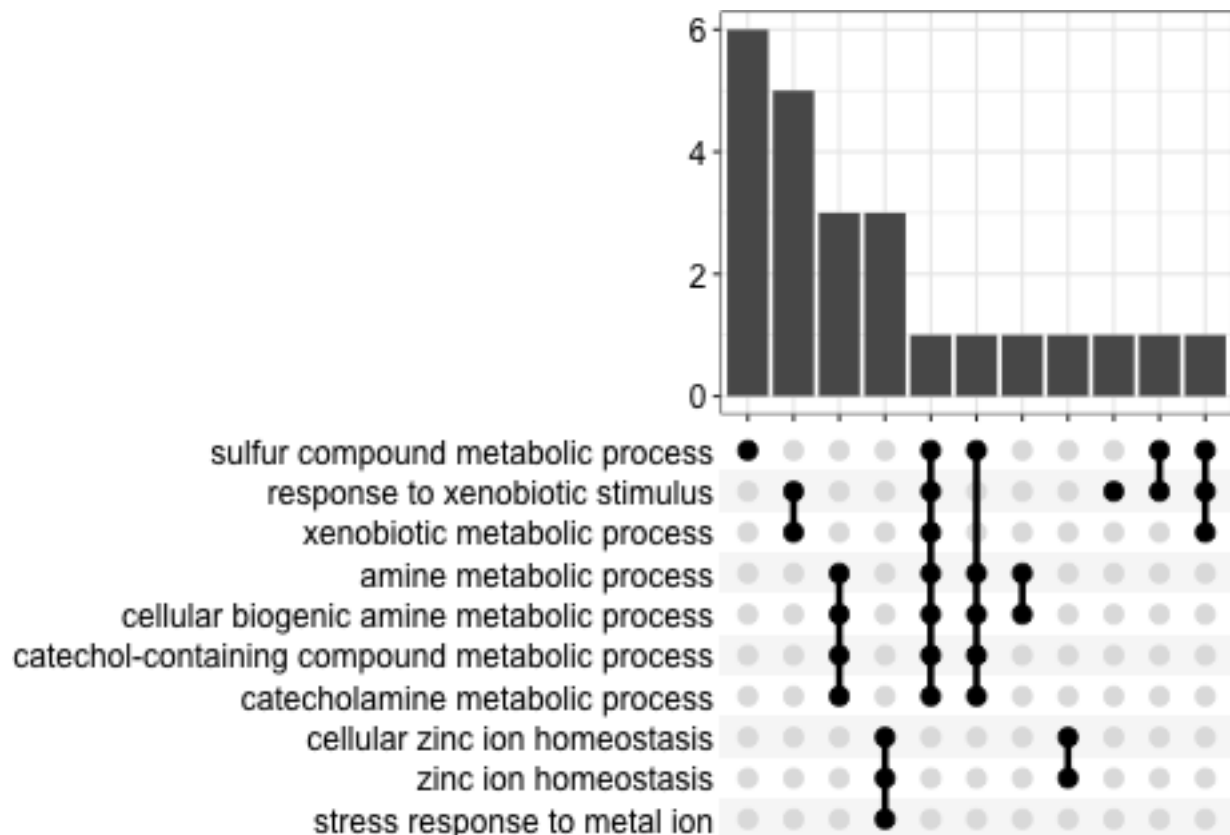
```
cnetplot(go.twas.bp.enrich.simpl,
 showCategory=100,
 node_label="category",
 cex_label_category=0.5)
```



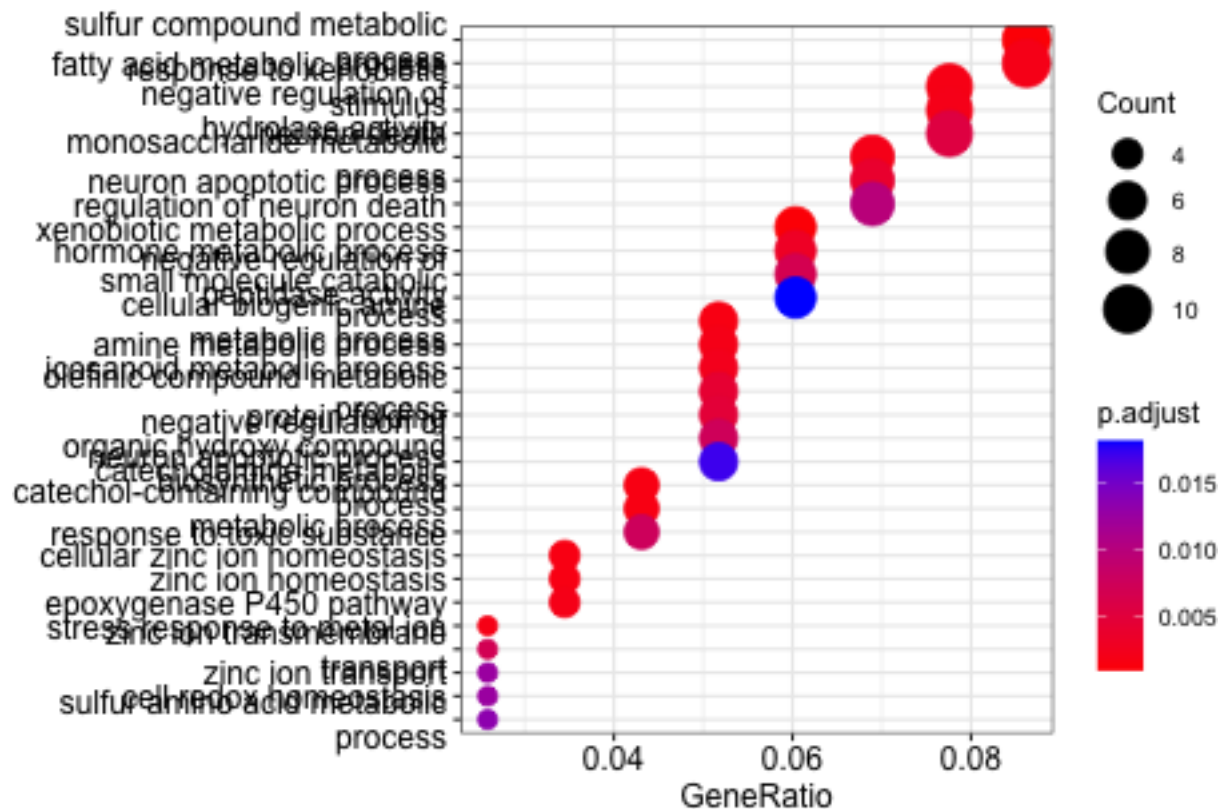
```
heatplot(go.twas.bp.enrich.simpl)
```



```
library(ggupset)
upsetplot(go.twas.bp.enrich.simpl)
```

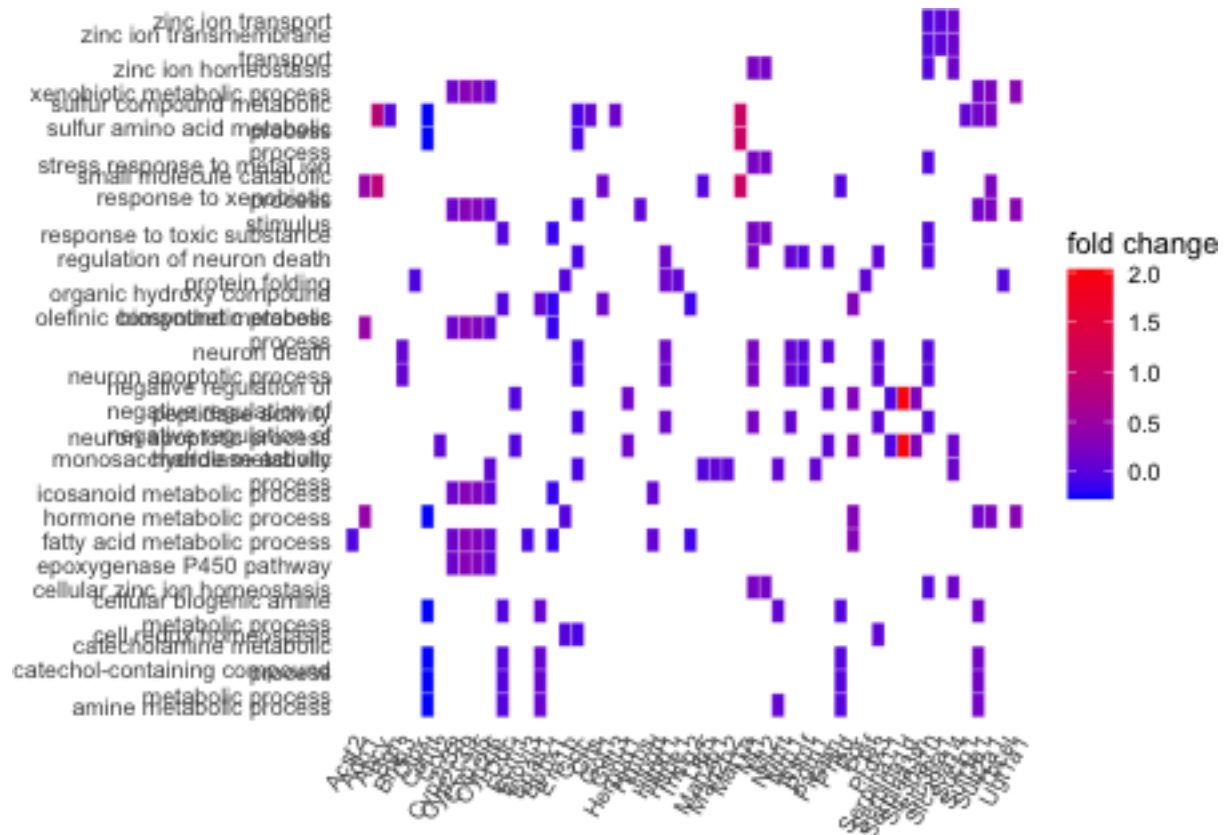


```
dotplot(go.twas.bp.enrich.simpl, showCategory=30)
```

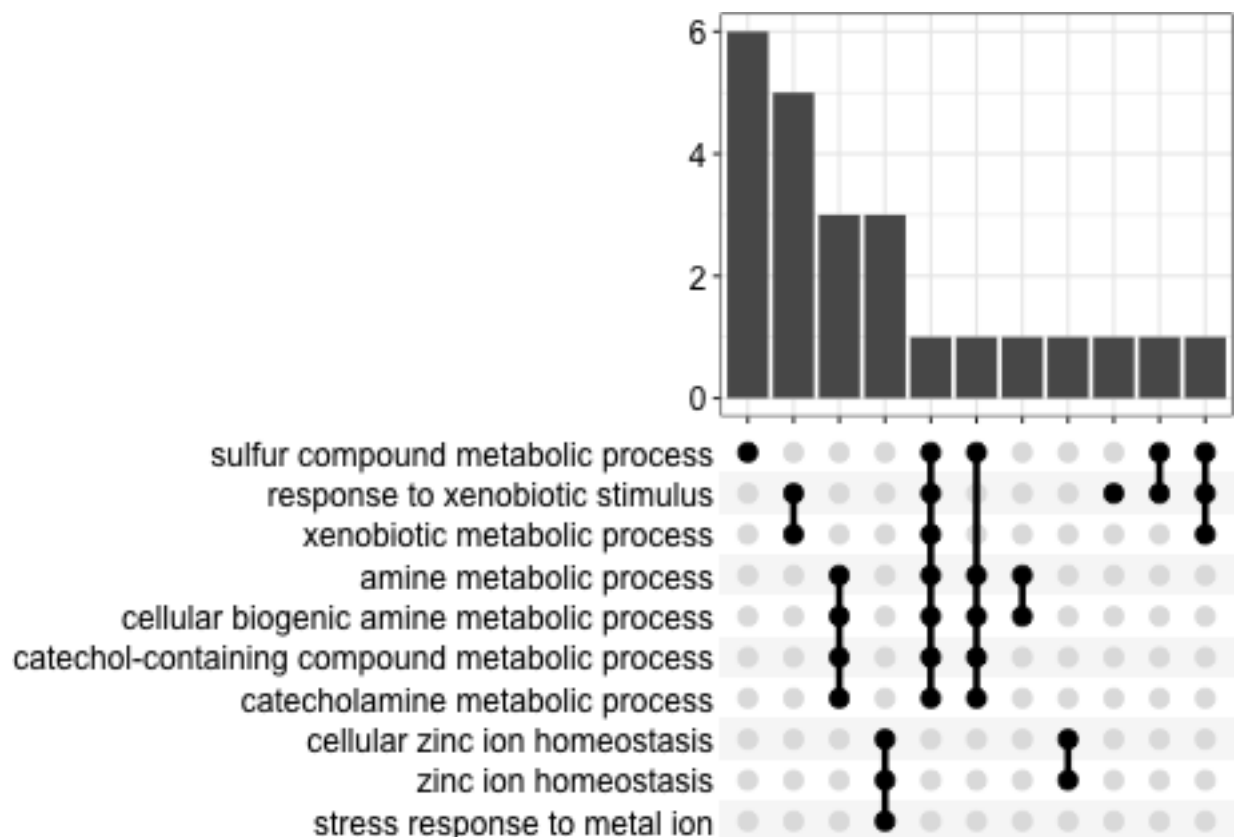




```
heatplot(go.twas.bp.enrich.simpl, foldChange=twas.list)
```



```
upsetplot(go.twas.bp.enrich.simpl)
```



```
as.data.frame(go.twas.bp) %>% dplyr::select("Description","setSize","enrichmentScore","NES","pvalue","p.adjust")
kable(caption="HFD GO BP Analysis of TWAS Associations from GSEA")
```

Table 12: HFD GO BP Analysis of TWAS Associations from GSEA

	Description	setSize	enrichmentScore	NES	pvalue	p.adjust
GO:0051640	organelle localization	113	0.959	1.82	0	0

```
as.data.frame(go.twas.bp) %>%
 write_csv(file="HFD TWAS GO-BP GSEA.csv")

as.data.frame(go.twas.bp.enrich.simpl) %>% dplyr::select("Description","GeneRatio","Count","pvalue","p.adjust")
kable(caption="HFD GO BP Analysis of TWAS Associations from Enrichment")
```

Table 13: HFD GO BP Analysis of TWAS Associations from Enrichment

	Description	GeneRatioCount	pvalue	p.adjust	
GO:0006805	xenobiotic metabolic process	7/116	7	0.000	0.001
GO:0006790	sulfur compound metabolic process	10/116	10	0.000	0.001
GO:0006882	cellular zinc ion homeostasis	4/116	4	0.000	0.002
GO:0006576	cellular biogenic amine metabolic process	6/116	6	0.000	0.002

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0006584	catecholamine metabolic process	5/116	5	0.000	0.002
GO: 0009712	catechol-containing compound metabolic process	5/116	5	0.000	0.002
GO: 0055069	zinc ion homeostasis	4/116	4	0.000	0.002
GO: 0009410	response to xenobiotic stimulus	9/116	9	0.000	0.002
GO: 0097501	stress response to metal ion	3/116	3	0.000	0.002
GO: 0009308	amine metabolic process	6/116	6	0.000	0.002
GO: 0019373	epoxygenase P450 pathway	4/116	4	0.000	0.002
GO: 0006631	fatty acid metabolic process	10/116	10	0.000	0.002
GO: 0005996	monosaccharide metabolic process	8/116	8	0.000	0.002
GO: 0051346	negative regulation of hydrolase activity	9/116	9	0.000	0.002
GO: 0006690	icosanoid metabolic process	6/116	6	0.000	0.002
GO: 0042445	hormone metabolic process	7/116	7	0.000	0.003
GO: 0051402	neuron apoptotic process	8/116	8	0.000	0.004
GO: 0120254	olefinic compound metabolic process	6/116	6	0.000	0.004
GO: 0006457	protein folding	6/116	6	0.000	0.005
GO: 0070997	neuron death	9/116	9	0.000	0.005
GO: 0010466	negative regulation of peptidase activity	7/116	7	0.000	0.007
GO: 0071577	zinc ion transmembrane transport	3/116	3	0.000	0.007
GO: 0043524	negative regulation of neuron apoptotic process	6/116	6	0.000	0.007
GO: 0009636	response to toxic substance	5/116	5	0.000	0.008
GO: 1901214	regulation of neuron death	8/116	8	0.000	0.010
GO: 0006829	zinc ion transport	3/116	3	0.000	0.012
GO: 0045454	cell redox homeostasis	3/116	3	0.000	0.012
GO: 0000096	sulfur amino acid metabolic process	3/116	3	0.000	0.013
GO: 1901617	organic hydroxy compound biosynthetic process	6/116	6	0.000	0.017
GO: 0044282	small molecule catabolic process	7/116	7	0.000	0.018

	Description	GeneRatio	Count	pvalue	p.adjust
GO: 0051881	regulation of mitochondrial membrane potential	4/116	4	0.000	0.018
GO: 0052547	regulation of peptidase activity	8/116	8	0.000	0.018
GO: 0008202	steroid metabolic process	7/116	7	0.000	0.018
GO: 1902686	mitochondrial outer membrane permeabilization involved in programmed cell death	3/116	3	0.000	0.019
GO: 0006066	alcohol metabolic process	7/116	7	0.001	0.020
GO: 0071248	cellular response to metal ion	5/116	5	0.001	0.023
GO: 0043434	response to peptide hormone	7/116	7	0.001	0.023
GO: 0006520	cellular amino acid metabolic process	6/116	6	0.001	0.023
GO: 0009309	amine biosynthetic process	3/116	3	0.001	0.023
GO: 0042401	cellular biogenic amine biosynthetic process	3/116	3	0.001	0.023
GO: 0006068	ethanol catabolic process	2/116	2	0.001	0.027
GO: 0061299	retina vasculature morphogenesis in camera-type eye	2/116	2	0.001	0.027
GO: 0009068	aspartate family amino acid catabolic process	2/116	2	0.001	0.031
GO: 0090161	Golgi ribbon formation	2/116	2	0.001	0.031
GO: 0003254	regulation of membrane depolarization	3/116	3	0.001	0.031
GO: 0042593	glucose homeostasis	6/116	6	0.001	0.034
GO: 0008637	apoptotic mitochondrial changes	4/116	4	0.001	0.037
GO: 0010823	negative regulation of mitochondrion organization	3/116	3	0.001	0.037
GO: 2000378	negative regulation of reactive oxygen species metabolic process	3/116	3	0.001	0.037
GO: 0048102	autophagic cell death	2/116	2	0.002	0.041
GO: 0050922	negative regulation of chemotaxis	3/116	3	0.002	0.044
GO: 0042416	dopamine biosynthetic process	2/116	2	0.002	0.044
GO: 0010713	negative regulation of collagen metabolic process	2/116	2	0.002	0.048
GO: 0032966	negative regulation of collagen biosynthetic process	2/116	2	0.002	0.048

```
as.data.frame(go.twas.bp.enrich.simpl) %>%
 write_csv(file="HFD TWAS GO-BP Enrichment.csv")
```

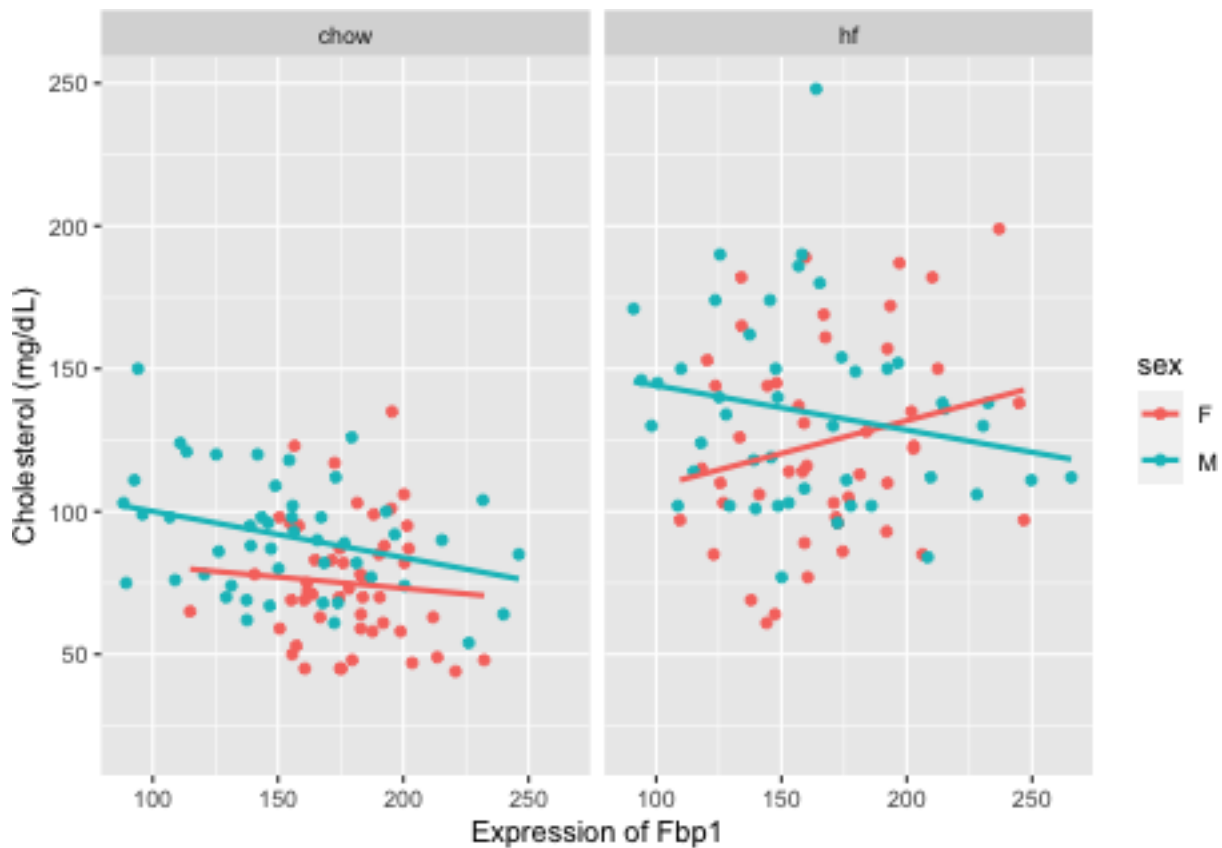
```
steroid.enrichment <- as.data.frame(go.twas.bp) %>% filter(ID=='GO:0016126')
```

## 10 Specific Associations

Fbp1 was one of the most highly associated results on NCD.

```
gene <- 'Fbp1'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)
library(ggplot2)
```

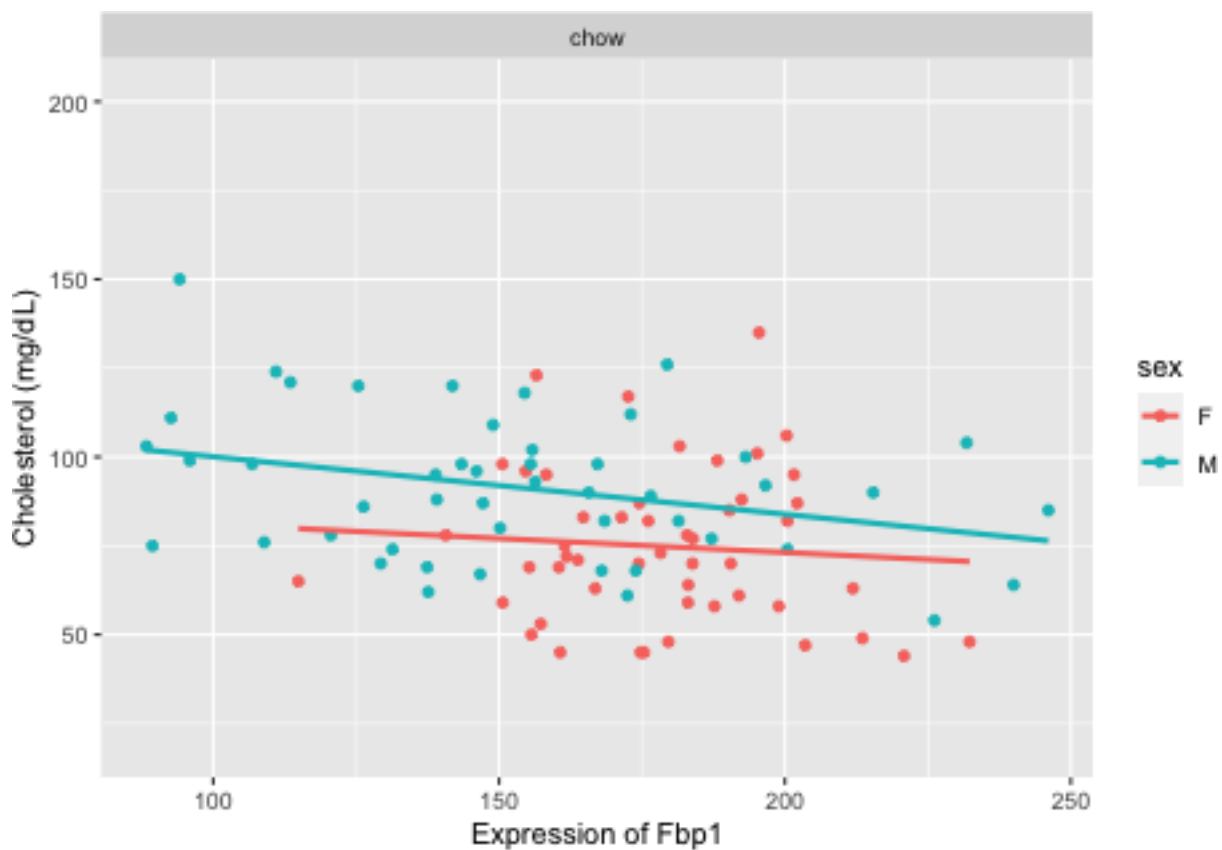
```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))
```



```

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))

```



```

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> fbp1.chow.data

lm(data=fbp1.chow.data, chol2 ~ expression + sex) %>%

```

```
glance
```

```
A tibble: 1 x 12
r.squ~1 adj.r~2 sigma stati~3 p.value df logLik AIC BIC devia~4 df.re~5
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <int>
1 0.176 0.159 20.6 10.2 9.96e-5 2 -434. 876. 886. 40231. 95
... with 1 more variable: nobs <int>, and abbreviated variable names
1: r.squared, 2: adj.r.squared, 3: statistic, 4: deviance, 5: df.residual
```

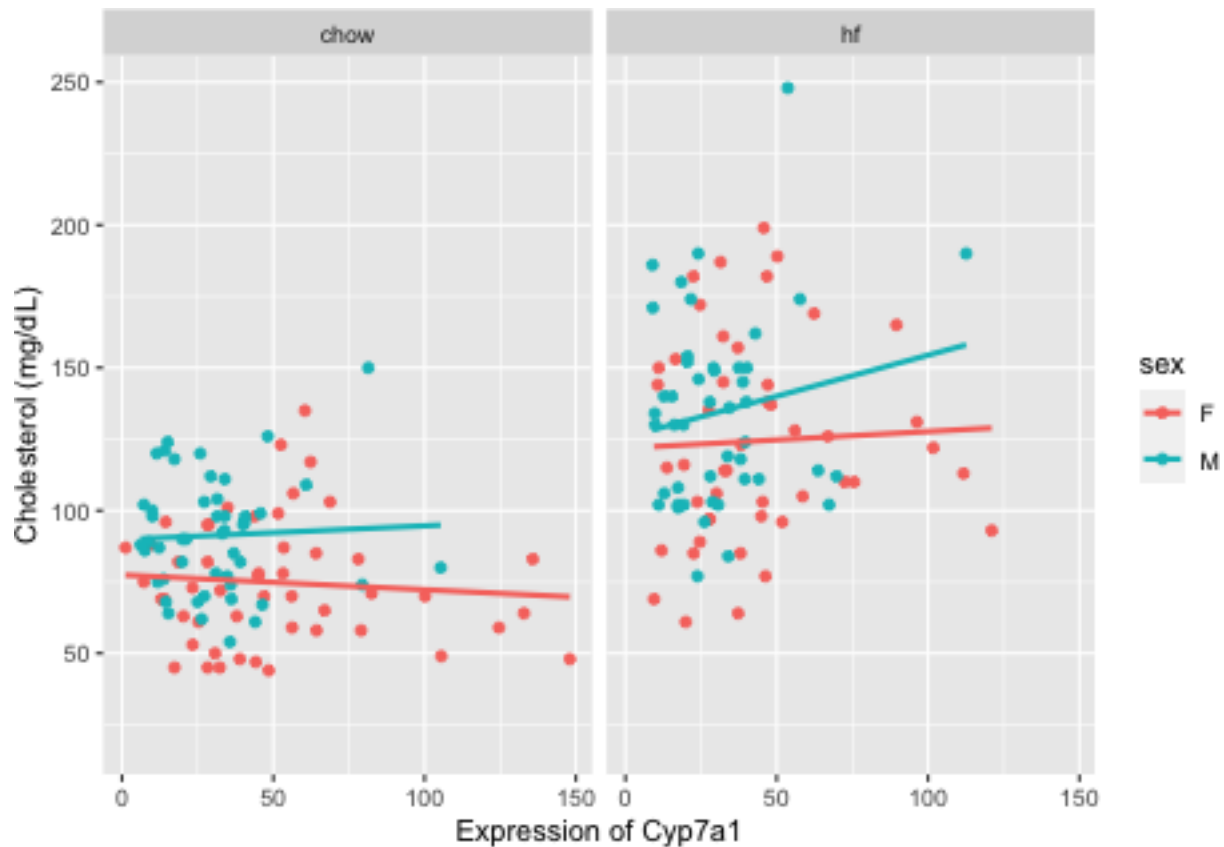
```
expression.data %>%
 filter(ENSEMBL.ID == gene) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) -> fbp.hf.data
```

## 10.1 Bile Acid Metabolism

CYP7A1 is the rate limiting step for de novo *bile acid* biosynthesis

```
gene <- 'Cyp7a1'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))
```



```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> gene.chow.data

lm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Cyp7a1 and cholesterol on chow")
```

Table 14: Summary associations of Cyp7a1 and cholesterol on chow

term	estimate	std.error	statistic	p.value
(Intercept)	76.191	4.881	15.611	0.000
expression	-0.027	0.076	-0.355	0.724
sexM	15.954	4.518	3.531	0.001

```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
```



```

 names_to='mouse.id',
 values_to='expression') %>%
full_join(phenotype.data,by='mouse.id') %>%
filter(!is.na(diet)) -> gene.hf.data

lm(data=gene.hf.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Cyp7a1 and cholesterol on HFD")

```

Table 15: Summary associations of Cyp7a1 and cholesterol on HFD

term	estimate	std.error	statistic	p.value
(Intercept)	100.060	6.108	16.381	0.000
expression	-0.026	0.103	-0.255	0.799
sexM	13.262	5.565	2.383	0.018

## 10.2 SCD1

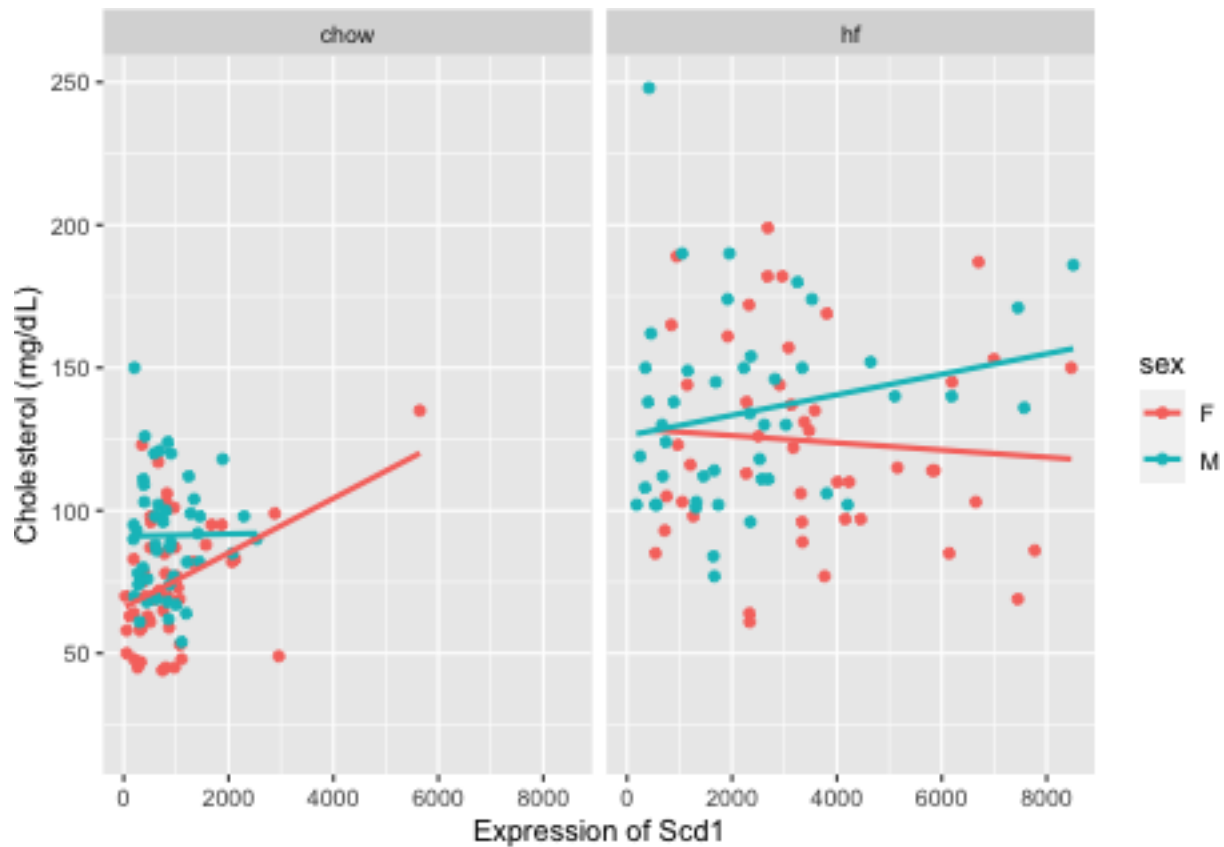
SCD1 is the gene with the biggest effect size on NCD

```

gene <- 'Scd1'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
full_join(phenotype.data,by='mouse.id') %>%
filter(!is.na(diet)) %>%
ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))

```



```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> gene.chow.data

lm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Scd1 and cholesterol on chow")
```

Table 16: Summary associations of Scd1 and cholesterol on chow

term	estimate	std.error	statistic	p.value
(Intercept)	68.093	3.775	18.04	0.000
expression	0.007	0.003	2.74	0.007
sexM	17.118	4.105	4.17	0.000

```
library(MASS)
rlm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Scd1 and cholesterol on chow, using robust linear models")
```

Table 17: Summary associations of Scd1 and cholesterol on chow, using robust linear models

term	estimate	std.error	statistic
(Intercept)	65.735	3.908	16.82
expression	0.009	0.003	3.30
sexM	16.623	4.250	3.91

```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) -> gene.hf.data

lm(data=gene.hf.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Scd1 and cholesterol on HFD")
```

Table 18: Summary associations of Scd1 and cholesterol on HFD

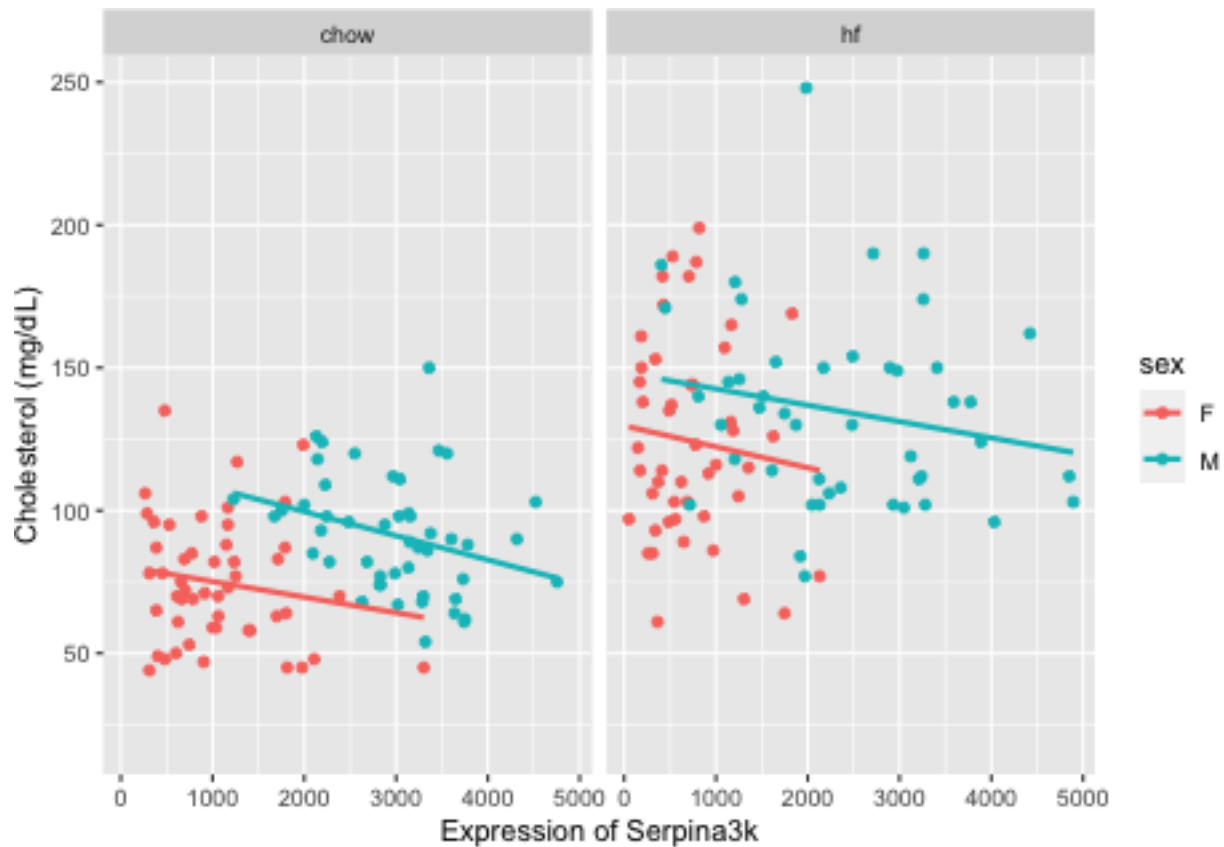
term	estimate	std.error	statistic	p.value
(Intercept)	80.778	4.398	18.37	0
expression	0.008	0.001	6.40	0
sexM	18.589	4.874	3.81	0

### 10.3 Serpina3k

Serpina3k is the gene with the biggest inverse effect size on NCD

```
gene <- 'Serpina3k'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))
```



```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> gene.chow.data

lm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Serpina3k and cholesterol on chow")
```

Table 19: Summary associations of Serpina3k and cholesterol on chow

term	estimate	std.error	statistic	p.value
(Intercept)	82.390	4.331	19.02	0.000
expression	-0.007	0.003	-2.35	0.021
sexM	30.226	7.160	4.22	0.000

```
library(MASS)
rlm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Serpina3k and cholesterol on chow, using robust linear models")
```

Table 20: Summary associations of Serpina3k and cholesterol on chow, using robust linear models

term	estimate	std.error	statistic
(Intercept)	81.991	4.226	19.40
expression	-0.008	0.003	-2.74
sexM	32.690	6.985	4.68

```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) -> gene.hf.data

lm(data=gene.hf.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Serpina3k and cholesterol on HFD")
```

Table 21: Summary associations of Serpina3k and cholesterol on HFD

term	estimate	std.error	statistic	p.value
(Intercept)	111.314	4.516	24.65	0
expression	-0.014	0.003	-4.46	0
sexM	38.748	7.554	5.13	0

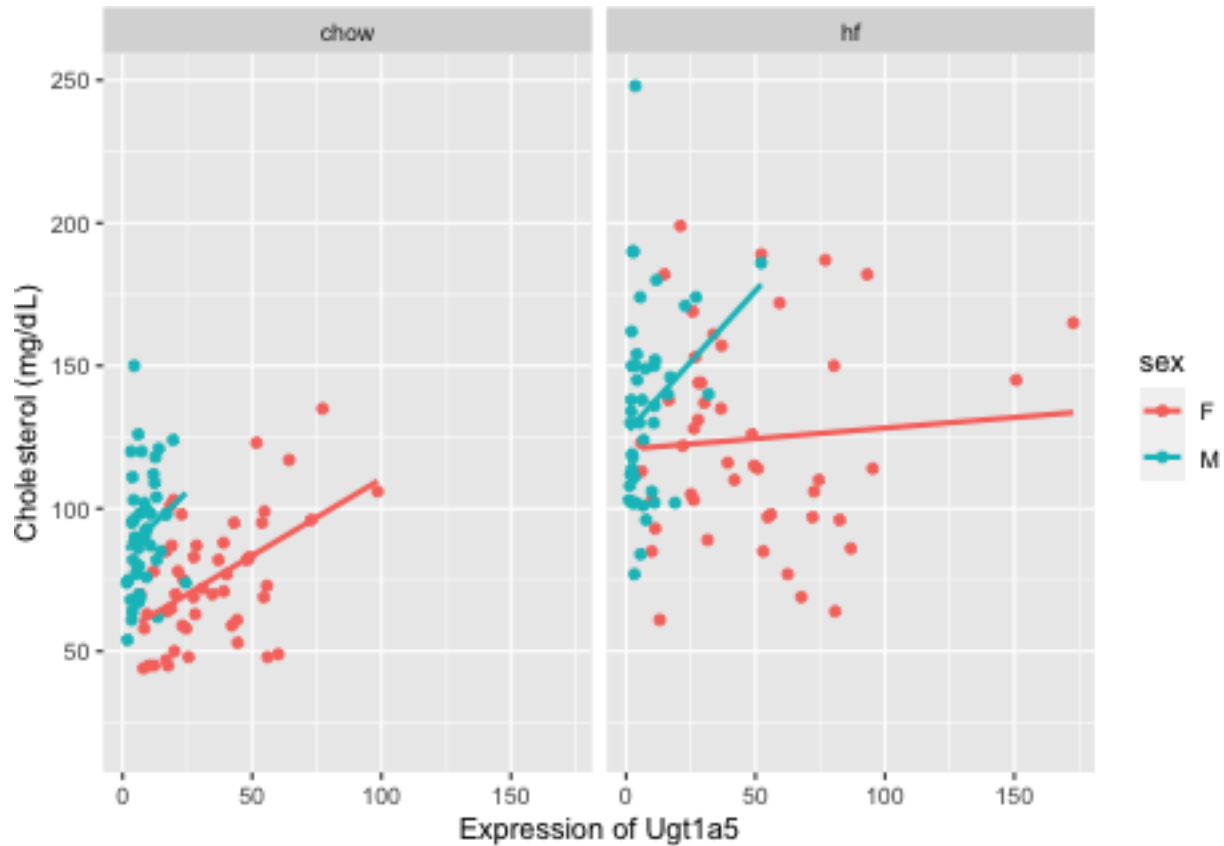
## 10.4 Ugt1a5

Ugt1a5 is the gene with the most significant effect on ncd

```
gene <- 'Ugt1a5'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
 labs(y="Cholesterol (mg/dL)",
```

```
x=paste('Expression of ', gene, sep="")
```



```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> gene.chow.data

lm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Ugt1a5 and cholesterol on chow")
```

Table 22: Summary associations of Ugt1a5 and cholesterol on chow

term	estimate	std.error	statistic	p.value
(Intercept)	55.979	5.277	10.61	0
expression	0.556	0.133	4.18	0
sexM	30.772	5.197	5.92	0

```
library(MASS)
rlm(data=gene.chow.data, chol2 ~ expression + sex) %>%
```

```
tidy %>%
kable(caption="Summary associations of Ugt1a5 and cholesterol on chow, using robust linear models")
```

Table 23: Summary associations of Ugt1a5 and cholesterol on chow, using robust linear models

term	estimate	std.error	statistic
(Intercept)	54.556	5.512	9.90
expression	0.586	0.139	4.22
sexM	30.864	5.428	5.69

```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) -> gene.hf.data

lm(data=gene.hf.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Ugt1a5 and cholesterol on HFD")
```

Table 24: Summary associations of Ugt1a5 and cholesterol on HFD

term	estimate	std.error	statistic	p.value
(Intercept)	80.915	6.12	13.22	0
expression	0.436	0.12	3.62	0
sexM	28.021	6.49	4.32	0

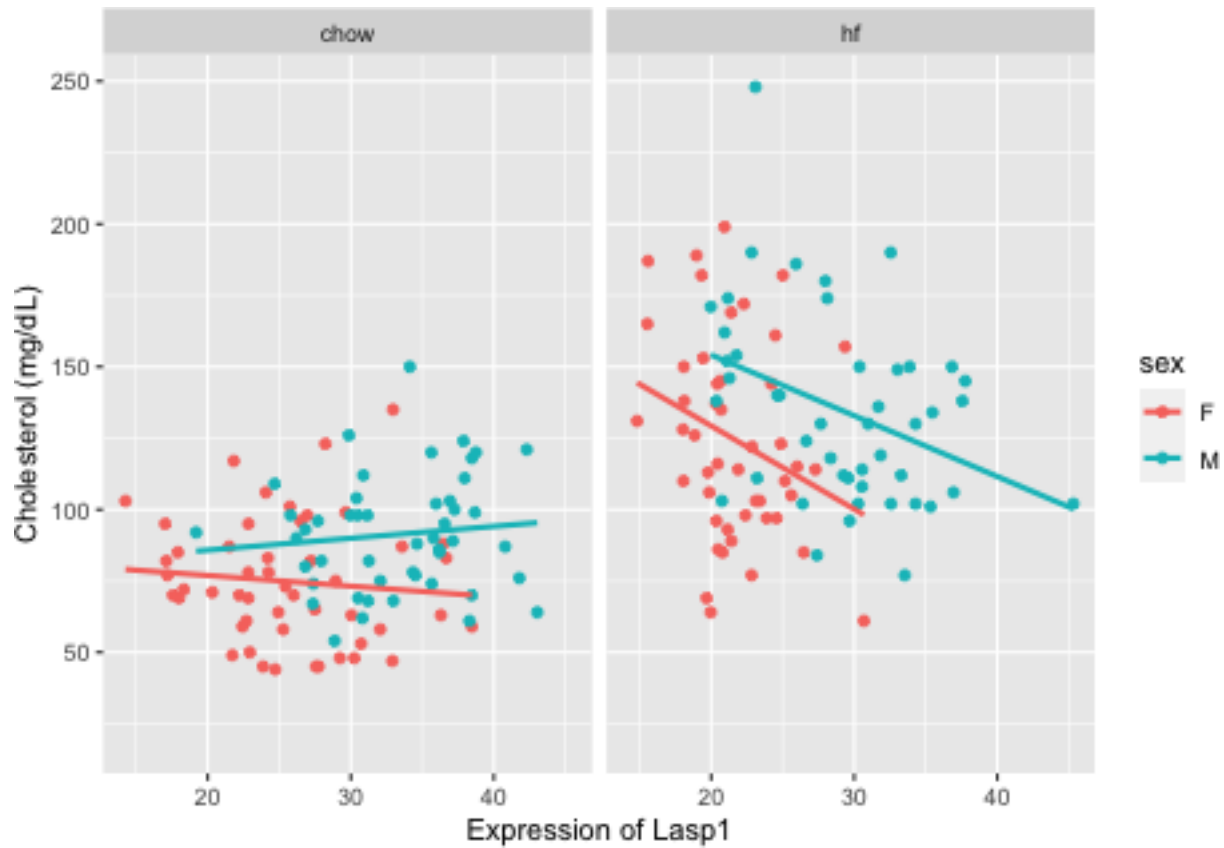
## 10.5 Laspl

Laspl is the gene with the most significant effect on HFD

```
gene <- 'Laspl'
gene.ens <- filter(twas.data.ncd, symbol==gene) %>% pull(ENSEMBL.ID)

expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 ggplot(aes(y=chol2,expression,col=sex)) +
 geom_point() +
 geom_smooth(method='lm',se=F) +
 facet_grid(~diet) +
```

```
labs(y="Cholesterol (mg/dL)",
 x=paste('Expression of ', gene, sep=""))
```



```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) %>%
 filter(diet=='chow') -> gene.chow.data

lm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Lasp1 and cholesterol on chow")
```

Table 25: Summary associations of Lasp1 and cholesterol on chow

term	estimate	std.error	statistic	p.value
(Intercept)	75.387	10.515	7.170	0.000
expression	-0.022	0.394	-0.056	0.955
sexM	16.663	5.225	3.189	0.002



```
library(MASS)
rlm(data=gene.chow.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Lasp1 and cholesterol on chow, using robust linear models")
```

Table 26: Summary associations of Lasp1 and cholesterol on chow, using robust linear models

term	estimate	std.error	statistic
(Intercept)	77.547	11.157	6.951
expression	-0.167	0.418	-0.399
sexM	18.260	5.544	3.294

```
expression.data %>%
 filter(ENSEMBL.ID == gene.ens) %>%
 pivot_longer(cols=c(starts_with('F'),
 starts_with('M')),
 names_to='mouse.id',
 values_to='expression') %>%
 full_join(phenotype.data,by='mouse.id') %>%
 filter(!is.na(diet)) -> gene.hf.data

lm(data=gene.hf.data, chol2 ~ expression + sex) %>%
 tidy %>%
 kable(caption="Summary associations of Lasp1 and cholesterol on HFD")
```

Table 27: Summary associations of Lasp1 and cholesterol on HFD

term	estimate	std.error	statistic	p.value
(Intercept)	156.11	11.333	13.78	0
expression	-2.42	0.455	-5.31	0
sexM	31.96	6.032	5.30	0

## 11 Session Information

```
sessionInfo()

R version 4.2.2 (2022-10-31)
Platform: x86_64-apple-darwin17.0 (64-bit)
Running under: macOS Big Sur ... 10.16
##
Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/4.2/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/4.2/Resources/lib/libRlapack.dylib
##
locale:
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
attached base packages:
```

```

[1] stats4 stats graphics grDevices utils datasets methods
[8] base
##
other attached packages:
[1] MASS_7.3-58.1 ggupset_0.3.0 enrichplot_1.16.2
[4] clusterProfiler_4.4.4 ggrepel_0.9.2 ggplot2_3.4.0
[7] venneuler_1.1-3 rJava_1.0-6 purrr_1.0.1
[10] org.Mm.eg.db_3.15.0 AnnotationDbi_1.58.0 IRanges_2.30.1
[13] S4Vectors_0.34.0 Biobase_2.56.0 BiocGenerics_0.42.0
[16] broom_1.0.2 readr_2.1.3 dplyr_1.0.10
[19] tidyr_1.2.1 knitr_1.41
##
loaded via a namespace (and not attached):
[1] fgsea_1.22.0 colorspace_2.0-3 ggtree_3.4.4
[4] ellipsis_0.3.2 qvalue_2.28.0 XVector_0.36.0
[7] aplot_0.1.9 rstudioapi_0.14 farver_2.1.1
[10] graphlayouts_0.8.4 bit64_4.0.5 scatterpie_0.1.8
[13] fansi_1.0.3 codetools_0.2-18 splines_4.2.2
[16] cachem_1.0.6 GOSemSim_2.22.0 polyclip_1.10-4
[19] jsonlite_1.8.4 GO.db_3.15.0 png_0.1-8
[22] ggforce_0.4.1 compiler_4.2.2 httr_1.4.4
[25] backports_1.4.1 lazyeval_0.2.2 assertthat_0.2.1
[28] Matrix_1.5-3 fastmap_1.1.0 cli_3.6.0
[31] tweenr_2.0.2 htmltools_0.5.4 tools_4.2.2
[34] igraph_1.3.5 gtable_0.3.1 glue_1.6.2
[37] GenomeInfoDbData_1.2.8 reshape2_1.4.4 DO.db_2.9
[40] fastmatch_1.1-3 Rcpp_1.0.9 vctrs_0.5.1
[43] Biostrings_2.64.1 ape_5.6-2 nlme_3.1-161
[46] ggraph_2.1.0 xfun_0.36 stringr_1.5.0
[49] lifecycle_1.0.3 DOSE_3.22.1 zlibbioc_1.42.0
[52] scales_1.2.1 tidygraph_1.2.2 vroom_1.6.0
[55] hms_1.1.2 parallel_4.2.2 RColorBrewer_1.1-3
[58] yaml_2.3.6 memoise_2.0.1 gridExtra_2.3
[61] downloader_0.4 ggfun_0.0.9 yulab.utils_0.0.6
[64] stringi_1.7.12 RSQlite_2.2.20 highr_0.10
[67] tidytree_0.4.2 BiocParallel_1.30.4 GenomeInfoDb_1.32.4
[70] rlang_1.0.6 pkgconfig_2.0.3 bitops_1.0-7
[73] evaluate_0.19 lattice_0.20-45 treeio_1.20.2
[76] patchwork_1.1.2 labeling_0.4.2 shadowtext_0.1.2
[79] bit_4.0.5 tidyselect_1.2.0 plyr_1.8.8
[82] magrittr_2.0.3 R6_2.5.1 magick_2.7.3
[85] generics_0.1.3 DBI_1.1.3 pillar_1.8.1
[88] withr_2.5.0 mgcv_1.8-41 KEGGREST_1.36.3
[91] RCurl_1.98-1.9 tibble_3.1.8 crayon_1.5.2
[94] utf8_1.2.2 tzdb_0.3.0 rmarkdown_2.19
[97] viridis_0.6.2 grid_4.2.2 data.table_1.14.6
[100] blob_1.2.3 digest_0.6.31 gridGraphics_0.5-1
[103] munsell_0.5.0 viridisLite_0.4.1 ggplotify_0.1.0

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