# R implementation

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2 Loaded functions:

<sup>1</sup> Project started Dec 10 2017

```
#source("/media/Data/Dropbox/humanR/01funcs.R")
rm(list=ls())
#setwd("/media/Data/Dropbox/humanR/PD/")
#setwd("~/Dropbox/humanR/PD/")
###load("PD.Rdata", .GlobalEnv)
#lsos(pat="")
```

Load packages.

#### 4 1 Data structure

- Data is from patients with Lymphoma tumors, either undergone or not a Rituximab CHOP treatment.
- 6 Some patients show relapse after treatment. Tumors migrate though nodal (lymphnodes) or extranodal
- 7 tissues. Tumors involve two different subtypes of cells of origin, ABC or GCB. The first aim is to find
- 8 correlation genes that respond differently to treatment, nodal transmission, and cell subtypes.

```
metadata <- read.table("data/phenodata", sep = "\t", header = T)</pre>
```

```
head (metadata)
  SAMPLE_ID PATIENT_ID Timepoint OTHER_ID res_id INCLUDE_MATCHING
1 CNR1001T1 CNR1001 T1 01-18186 YES
2 CNR1002T1 CNR1002 T1 01-26575
3 CNR1002T2 CNR1002 T2 01-26575
4 CNR1003T1 CNR1003 T1 02-10117
5 CNR1006T1 CNR1006 T1 DLC_0304 03-11110
                                    01-26575
6 CNR1007T1 CNR1007 T1 DLC_0193 03-26640
  INCLUDED_SUBMISSION_TCAG GROUP SITE Normalization Score
                    YES CNS_RELAPSE_RCHOP SO 37 789
                     YES CNS_RELAPSE_RCHOP GA
2
                                                      60 3548
3
                    YES CNS_RELAPSE_RCHOP CNS
                                                      62 3941
                                                      79
4
                     YES CNS_RELAPSE_RCHOP SO
                                                          -355
                    YES CNS_RELAPSE_RCHOP LN
YES CNS_RELAPSE_RCHOP SO
5
                                                     843 -245
                                                    143 3469
  ABClikelihood Prediction BCL2_BA BCL6_BA MYC_BA DH COMMENT CODE_OS
1 0 GCB 0 0 1 0 1
                   ABC
                            0
2
           1
                                    0
                                          0 0
                                                            1
                            0
3
           1
                   ABC
                                   0
                                          0 0
                                                           1
           0
                   GCB
                            1
                                   1
                                          1 1
           0 GCB 1 0 0 0
1 ABC 0 0 0 0
  CODE_DSS CODE_PFS CODE_TTP CODE_CNS Overall.survival..y.
  1 1 1 0.87
               1
                       1
2

    1
    1
    1

    1
    1
    1

    1
    1
    1

    1
    1
    1

    1
    1
    1

    1
    1
    1

        1
                                1
                                                2.98
                               1
3
                                                2.98
                                1
5
                                1
                       1
                               1
Disease.specific.survival..y. Progression.free.survival..y.
                      0.87
1
2
                        2.98
                                                   0.38
3
                        2.98
                                                   0.38
                        0.60
                                                   0.31
                        0.42
                        4.64
  Time.to.progression..y. Time.to.CNS.relapse..y. SEX AGE STAGE
                 0.52
                                       0.52 F 82 4B
1
                                        0.38 F 77
2
                  0.38
                                        0.38 F 77
3
                  0.38
                                                      4A
                                              F 54
4
                  0.31
                                        0.31
5
                  0.13
                                        0.15
                                               M 59
                                              M 62
                   0.54
                                        0.45
                                                      1AE
  STAGEGRP E4SITE PS LDH LDHNORML LDHRATIO MASS IPI IPI_GROUP
  ADV BoSo 0 997 415 2.40 14 4 3
1
                                       1 -1
     ADV GaKi 1 -1
                          210
                                 -1.00
2.
                         210 -1.00
                                        1 -1
3
     ADV GaKi 1 -1
                         210
                                 4.73 11 4
     ADV SoOvUt 4 993
     ADV Gi 2 861 540
LIM BoSo 1 424 210
5
                                 1.59 5 2
                                 2.02
                                        7 3
 CNS.RiskScore CNS.RiskGrp Rehyb
           4 3 NO
2
                      -1 YES
           -1
3
           -1
                      -1 YES
                      3
4
            4
                           NO
5
            2
                       2
                           NO
```

In the first steps of the analysis, the samples will be classified (supervised) into the following categories.

```
metadata <- read.table("data/phenodata", sep = "\t", header = T) %>%
```

```
dplyr::select(SAMPLE_ID, Timepoint, GROUP, SITE, Score, Prediction, ABClikelihood) %>%
    filter(Timepoint != "T2") %>%
    mutate(Groups = case_when(GROUP %in% c("CNS_RELAPSE_RCHOP",
                                           "CNS_RELAPSE_CHOPOREQUIVALENT",
                                           "CNS_DIAGNOSIS") ~ "CNS",
                              GROUP %in% c("TESTICULAR_NO_CNS_RELAPSE", "NO_RELAPSE") ~ "NOREL",
                              GROUP == "SYTEMIC_RELAPSE_NO_CNS" ~ "SYST",
                              TRUE ~ "CTRL")) %>%
   mutate(ABClassify = case_when(ABClikelihood >= .9 ~ "ABC",
                                ABClikelihood <= .1 ~ "GCB",
                                TRUE ~ "U")) %>%
    mutate(ABCScore = case_when(Score > 2412 ~ "ABC",
                              Score <= 1900 ~ "GCB",
                               Score == NA ~ "NA",
                              TRUE ~ "U")) %>%
   mutate(Nodes = case_when(SITE == "LN" ~ "LN",
                            SITE == "TO" ~ "LN",
                            SITE == "SP" ~ "LN",
                            TRUE ~ "EN")) %>%
    mutate(Lymphnodes = case_when(Nodes == "LN" ~ 1, TRUE ~ 0))
# make sure all samples preserve their ID
metadata$Groups <- as.factor(metadata$Groups)</pre>
metadata$ABClassify <- as.factor(metadata$ABClassify)</pre>
metadata$ABCScore <- as.factor(metadata$ABCScore)</pre>
metadata$Nodes <- as.factor(metadata$Nodes)</pre>
metadata$Lymphnodes <- as.factor(metadata$Lymphnodes)</pre>
summary (metadata)
                                                  GROUP
    SAMPLE_ID Timepoint
 CNR1001T1: 1 T1:236 NO_RELAPSE
                                                   :96
 CNR1002T1: 1 T2: 0 SYTEMIC_RELAPSE_NO_CNS
                   CNS_RELAPSE_RCHOP :39
TESTICULAR_NO_CNS_RELAPSE :12
CNS_DIAGNOSIS :11
 CNR1003T1: 1
 CNR1006T1: 1
 CNR1007T1: 1
 CNR1008T1: 1
                        CNS_RELAPSE_CHOPOREQUIVALENT: 8
 (Other) :230
                         (Other)
     SITE Score Prediction ABClikelihood Groups
    :127 Min. :-881 ABC : 92 Min. :0.00 CNS : 58
      : 20 1st Qu.: 676 GCB :103 1st Qu.:0.00 CTRL : 6
       : 18 Median :2106 U : 39 Median :0.02 NOREL:108
             Mean :1820 NA's: 2 Mean :0.47
       : 16
                                                     SYST : 64
 GI : 11
SP : 7
              3rd Qu.:2941
                                       3rd Qu.:1.00
             Max. :4323
                                       Max. :1.00
 (Other): 37 NA's :2
                                       NA's :4
 ABClassify ABCScore Nodes Lymphnodes
 ABC:103 ABC: 92 EN: 86 0: 86
 GCB:117 GCB:103 LN:150 1:150
 U: 16 U: 41
```

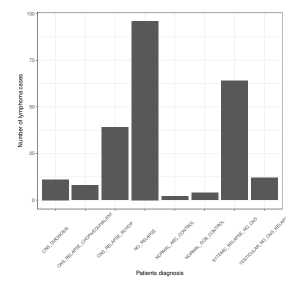
Difference in cases being indexed based on their *cell-of-origin* association subtypes using either of the following features: prediction, ABClassify, ABCScore.

```
metadata %>%
```

```
select (Prediction, ABClassify, ABCScore) %>%
summary

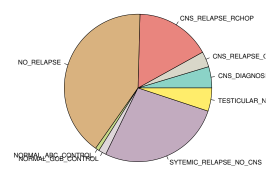
Prediction ABClassify ABCScore
ABC: 92 ABC:103 ABC: 92
GCB:103 GCB:117 GCB:103
U : 39 U : 16 U : 41
NA's: 2
```

Distribution of samples with different treatments.

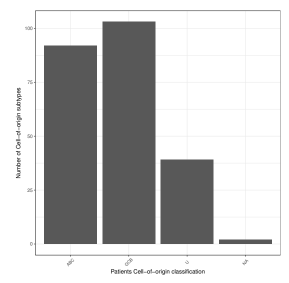


Or as a pie chart.

```
palette.pies <- brewer.pal(12, name = "Set3")
palette.pies.adj <- colorRampPalette(palette.pies)(length(unique(metadata$GROUP)))
pie(table(metadata$GROUP), col=palette.pies.adj)</pre>
```



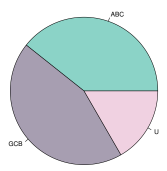
Distribution of samples with different cells of origin subtypes.



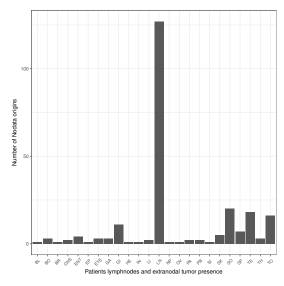
8 Or as pie chart.

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```
palette.pies <- brewer.pal(12, name = "Set3")
palette.pies.adj <- colorRampPalette(palette.pies)(length(unique(metadata$Prediction)))
pie(table(metadata$Prediction), col=palette.pies.adj)</pre>
```



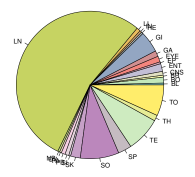
Distribution of samples with different lymphnodes and extranodal cancer metastasis.



Or as a pie chart.

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```
palette.pies <- brewer.pal(12, name = "Set3")
palette.pies.adj <- colorRampPalette(palette.pies) (length(unique(metadata$SITE)))
pie(table(metadata$SITE), col=palette.pies.adj)</pre>
```



# 2 Differential expression

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Genes have been fitted in a model that is based on an Empirical Bayes approach. Ranking of the genes determine if they are statistically significant. Bonferroni correction is used to control the false discovery rate (FDR). Moderated t-statistics, FDR, and fold change (log2) are implemented to reduce selection of false positives.

- adjpval is the adjusted P-value to control the FDR using Bonferroni correction. Genes selected here based on their adjpval are also greater than or equal to the bstat threshold.
- avgex is the average expression the ordinary arithmetic average of the log2-expression values for the probe, across all arrays. Genes selected here based on their avgex are also greater than or equal to the bstat threshold.
- bstat is the moderated t-statistics using an Empirical Bayes approach generating B-statistics scores.

```
expression <- read.table("data/summary.full.90800.txt", sep = "\t", header = T) %>%
   select(Design, Model, Bthreshold, adjPval, Category, Parameter, Transcripts) %>%
   filter(Category == "total")
summary (expression)
                                                  Model
                  Design
CNSvsNOREL_ABC
                 : 54 systemicRelapse
                                                 : 54
CNSvsNOREL_GCB
                    : 54 systemicRelapseCOOclasses :162
                    : 54 systemicRelapseCOOprediction:162
CNSvsSYST_ABC
               : 54 systemicRelapseCOOscores :162
CNSvsSYST_GCB
diffCNSvsNOREL_ABCvsGCB: 54 systemicRelapseNodes
diffCNSvsSYST ABCvsGCB : 54
(Other) :378
Bthreshold adjPval Category
                                         Parameter
Min. :-2.00 Min. :0.049 down : 0 adjpval:234
1st Qu.:-1.00 1st Qu.:0.049 total:702 avgex :234
             Median :0.049
                                       bval :234
Median: 0.25
                             up : 0
Mean : 0.00
              Mean :0.049
3rd Ou.: 1.00
              3rd Ou.:0.049
Max. : 1.50
              Max. :0.049
 Transcripts
Min. : 0
           2
1st Qu.:
Median: 46
Mean : 580
3rd Qu.: 463
Max. :10578
```

Number of transcripts when comparing B-statistics scores, which represent confidence in selecting each significantly expressed gene.

```
aggregate ( Transcripts ~ Bthreshold, data=expression, FUN=range)
 Bthreshold Transcripts.1 Transcripts.2
       -2.0
1
2
       -1.0
                      0
                                6448
                                 3618
3
       0.0
                      0
4
        0.5
                       0
                                 2688
5
                       0
        1.0
                                 1976
6
        1.5
                       0
                                 1429
```

Number of transcripts when samples are classed into groups, which are based on clinical data (e.g., cell-of-origin, CNS relapse, and nodal/extranodal tumor transmission).

```
aggregate( Transcripts ~ Model, data=expression, FUN=range)
                       Model Transcripts.1 Transcripts.2
             systemicRelapse 0
2
  systemicRelapseCOOclasses
                                       0
                                                 10578
3 systemicRelapseCOOprediction
                                       0
                                                 10578
                                       0
4
    systemicRelapseCOOscores
                                                 10578
5
    systemicRelapseNodes
                                        0
                                                  6609
```

Number of transcripts found when comparing different sample cases indexed based on their clinical data.

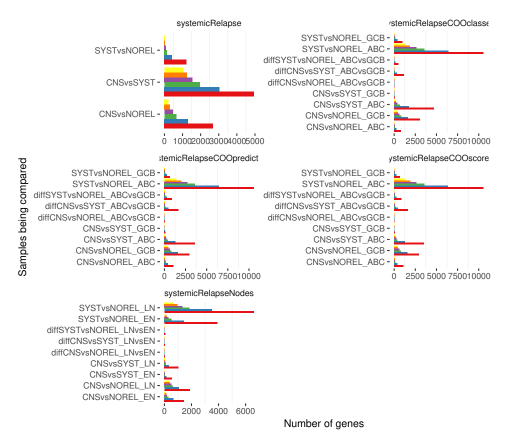
| aggregate ( Transcripts ~ De:      | sign, data=express | ion, FUN=range) |  |
|------------------------------------|--------------------|-----------------|--|
| Design Transcripts.1 Transcripts.2 |                    |                 |  |
| 1 CNSvsNOREL                       | 116                | 2678            |  |
| 2 CNSvsNOREL_ABC                   | 2                  | 1082            |  |
| 3 CNSvsNOREL_EN                    | 51                 | 1442            |  |
| 4 CNSvsNOREL_GCB                   | 130                | 3019            |  |
| 5 CNSvsNOREL_LN                    | 125                | 1873            |  |
| 6 CNSvsSYST                        | 441                | 4938            |  |
| 7 CNSvsSYST_ABC                    | 2                  | 4691            |  |
| 8 CNSvsSYST_EN                     | 3                  | 547             |  |
| 9 CNSvsSYST_GCB                    | 0                  | 98              |  |
| 10 CNSvsSYST_LN                    | 0                  | 1014            |  |
| 11 diffCNSvsNOREL_ABCvsGCB         | 0                  | 58              |  |
| 12 diffCNSvsNOREL_LNvsEN           | 0                  | 37              |  |
| 13 diffCNSvsSYST_ABCvsGCB          | 1                  | 1640            |  |
| 14 diffCNSvsSYST_LNvsEN            | 0                  | 23              |  |
| 15 diffSYSTvsNOREL_ABCvsGCB        | 0                  | 868             |  |
| 16 diffSYSTvsNOREL_LNvsEN          | 0                  | 85              |  |
| 17 SYSTVSNOREL                     | 0                  | 1214            |  |
| 18 SYSTVSNOREL_ABC                 |                    | 10578           |  |
| 19 SYSTVSNOREL_EN                  | 35                 | 3907            |  |
| 20 SYSTvsNOREL_GCB                 |                    | 994             |  |
| 21 SYSTVSNOREL_LN                  | 295                | 6609            |  |

Number of genes that respond to treatment, cell subtypes, and nodal transmission.

```
expression %>%
```

```
ggplot (aes (
    x = Design,
    y = Transcripts,
    fill = factor(Bthreshold))) +
theme_bw() +
geom bar(stat = "identity",
        position = "dodge") +
coord_flip() +
facet_wrap( ~ Model,
         ncol = 2,
          scales = "free") +
scale_fill_brewer(type = "qual", palette = 6) +
labs(x = "Samples being compared",
     y = "Number of genes") +
theme(legend.position = "top",
      strip.background = element_rect(linetype = "blank",
                                       fill = "white"),
      panel.border = element_rect(linetype = "blank",
                                   fill = NA),
      panel.grid.major = element_line(linetype = "blank"))
```





#### 3 Networks

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The number of clusters and modules per networks are assigned by designing first a similarity matrix between differentially expressed gene for any two conditions (eg., relapse vs no relapse patient cases). An adjacency matrix is then constructed by weighting the previously inferred measures. The data is transformed to increase the correlation coefficient therefore improving detection of strong correlated patterns. (Example of the strength of data transformation and correlation, visit the following online page).

- MaxEdgesPerGene, maximum number of correlations per genes
- · NbNodes, number of genes found for each edge connection bracket

- **Normalization**, method that focuses on creating complete clusters. We tested methods ranging from Complete clustering, Average, and Ward. Each method is detailed here. Only Complete clustering was retained. All other methods overfitted the data.
- 10verfitting is a source of bias.
- Correlation, finding ranges from linear to non-linear trends. We tested Pearson and Spearman correlation.

†Effect of correlation methods is seen on module content

- **Standardization**, data transformation method. We tested transformation by Hellinger, Standardize, Range, and Logarithmic scaling. Each method is detailed here.
- MaxGenePerModule, how many genes assigned by cluster (module)
- SimilaritySize, number of initial differentially expressed genes
- EdgeThreshold, parameter to limit the weight of the edges
- CorrelationPower, power transformation of the data

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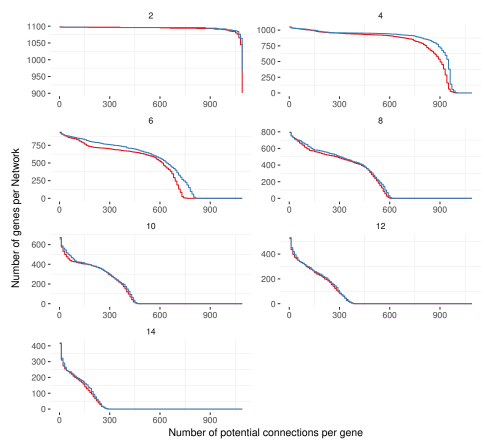
60

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```
ns <- read.table("./data/networks.summary.104795.txt", header = T)</pre>
summary(ns)
MaxEdgesPerGene NbNodes
                              Normalization Correlation
Min. : 1 Min. : 0
                               complete:4620 spearman:4620
1st Ou.: 271
               1st Qu.:
                          0
              Median: 244
Median : 546
Mean : 546 Mean : 406
3rd Qu.: 821 3rd Qu.: 862
Max. :1091 Max. :1098
  Standardization MaxGenesPerModule SimilaritySize EdgeThreshold
hellinger :2310 Min. :26 Min. :1099 Min. :0.5
standardize:2310 1st Qu.:36
Median:55
                                   1st Qu.:1099 1st Qu.:0.5
                                   Median:1099 Median:0.5
                   Mean :57 Mean :1099 Mean :0.5
3rd Qu::79 3rd Qu::1099 3rd Qu::0.5
Max. :91 Max. :1099 Max. :0.5
CorrelationPower
Min. : 2
1st Qu.: 4
Median: 8
Mean : 8
3rd Ou.:12
```

Difference between methods used for network inference. Are we able to generate convergence of the output of all iterations across all methods?

```
ns %>%
    ggplot (aes (
       x = MaxEdgesPerGene,
       y = NbNodes,
       fill = Standardization)) +
    theme bw() +
    geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
              scales = "free") +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
    theme(legend.position = "top",
         strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



Showing the number of modules per network and the number of genes per module. Each module contains differing number of nodes based on their correlation strength. Each cluster contains at least one module. Each network contains at least one cluster. One module can be assigned to nodes that belong to more than one cluster. The Lowess curves show if the trend in the data is linear or not. The wave around Lowess curves represents the level of confidence of the data points (the narrower the interval the better, less variability = more accuracy).

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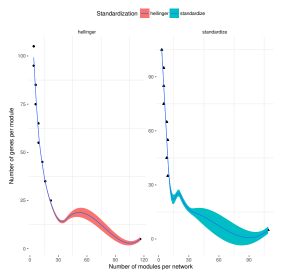
68

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¹Points=iterations. With less iterations comes high variability of the curve

```
read.table("./data/modules.summary.104795.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



# 3.1 Network analysis for Spearman-related correlations

Thresholds based on the Empirical Bayes approach to rank genes and determine if a gene is significantly expressed. Limma implementation.

- Average Expression: 5
- Adjusted P-value: equal or less than 0.045
- · Log Fold Change: 1
- B-statisitcs: 1.5

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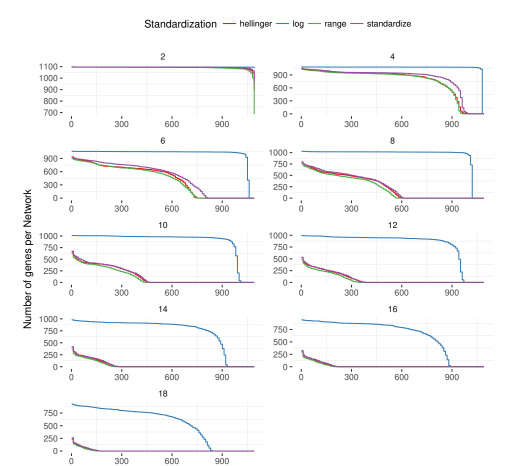
78

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#### 3.1.1 Nodal versus extra-nodal lymphoma

Genetic networks from differentially expressed genes selected by comparing sample cases with nodal and extranodal lymphoma.

```
read.table("./data/networks.summary.104859.txt", header = TRUE) %>%
    ggplot (aes (
       x = MaxEdgesPerGene,
       y = NbNodes,
        fill = Standardization)) +
    theme_bw() +
    geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```

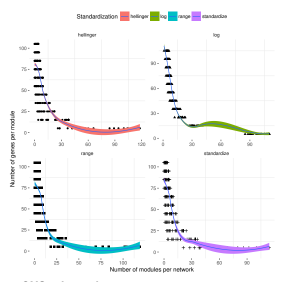


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```
read.table("./data/modules.summary.104859.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

Number of potential connections per gene



## 3.1.2 Relapsed versus no CNS relapsed cases

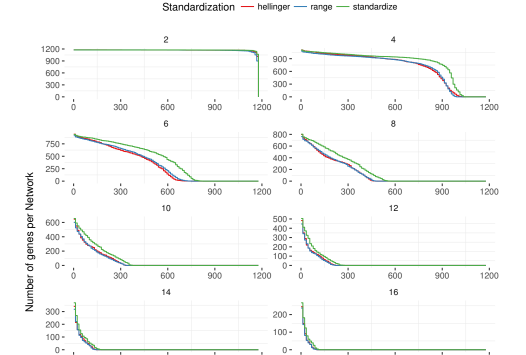
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Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

```
read.table("./data/networks.summary.114018.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```



1200

Ö

300

600

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1200

50 - 0 - 0 - 300 600 900 1200

Number of potential connections per gene

300

600

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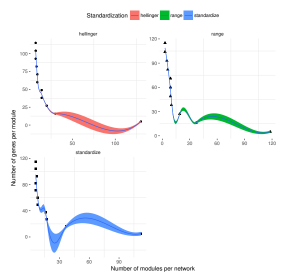
0

200 **-**150 **-**100 **-**

89 Showing the number of modules per network and the number of genes per module.

900

```
read.table("./data/modules.summary.114018.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



## 3.1.3 Lymphoma cases classified by Cell-of-origin subtypes

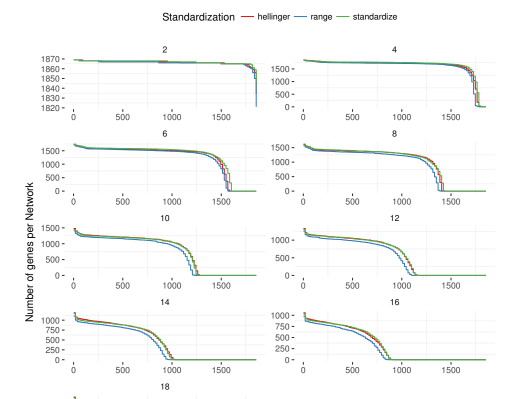
90

91

92

Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

```
read.table("./data/networks.summary.114017.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```



1500

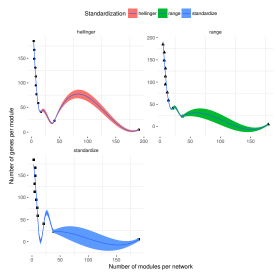
1000

750 - 500 - 250 - 0 - 0

500

```
read.table("./data/modules.summary.114017.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

Number of potential connections per gene



## 3.2 Network analysis for Pearson-related correlations

Thresholds based on the Empirical Bayes approach to rank genes and determine if a gene is significantly expressed. Limma implementation.

"With pearson, we can only raise the data to power 10. All are discarded after 10.

• Average Expression: 5

• Adjusted P-value: equal or less than 0.045

Log Fold Change: 1

• B-statisitcs: 1.5

96

97

98

99

100

101

102

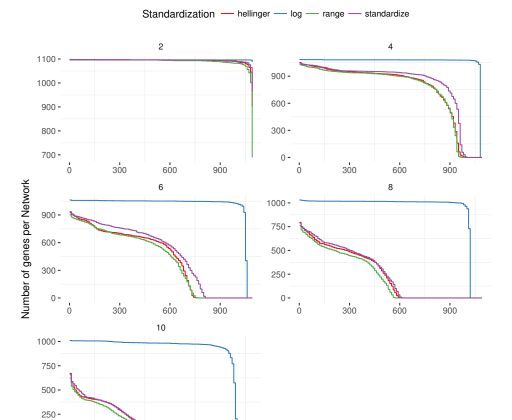
103

104

## 3.2.1 Nodal versus extra-nodal lymphoma

Genetic networks from differentially expressed genes selected by comparing sample cases with nodal and extranodal lymphoma.

```
read.table("./data/networks.summary.104862.txt", header = TRUE) %>%
    ggplot (aes (
        x = MaxEdgesPerGene,
        y = NbNodes,
        fill = Standardization)) +
    theme_bw() +
    geom step(aes(color = Standardization),
              stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



600

900

Number of potential connections per gene

0 -

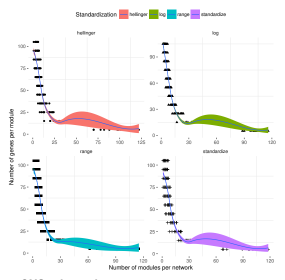
107

108

300

```
read.table("./data/modules.summary.104862.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

nSince Lowess ranks by confidence, Log transformation seems the best, ie, low variability. For this, Log is removed from further tests.



## 3.2.2 Relapsed versus no CNS relapsed cases

109

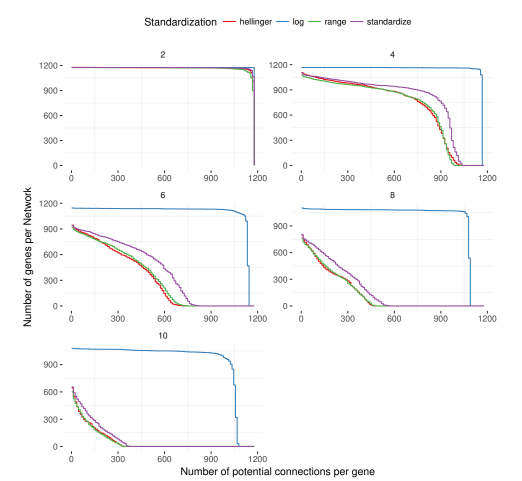
110

111

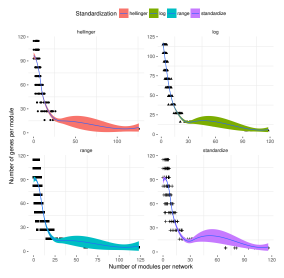
112

Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

```
read.table("./data/networks.summary.104863.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```



```
read.table("./data/modules.summary.104863.txt", header = TRUE) %>%
   ggplot (aes (
       x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
   theme_bw() +
   geom_point(aes(shape = Standardization)) +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of modules per network",
         y = "Number of genes per module") +
   facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
   geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



# 3.2.3 Lymphoma cases classified by Cell-of-origin subtypes

115

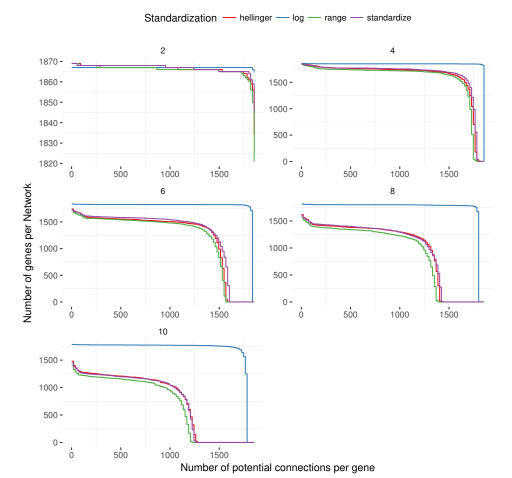
116

117

118

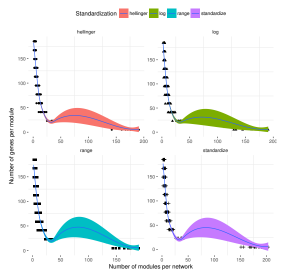
Genetic networks from differentially expressed genes selected by comparing sample cases with cell of origin classification based on ABC or GCB subtypes.

```
read.table("./data/networks.summary.104864.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```



119

```
read.table("./data/modules.summary.104864.txt", header = TRUE) %>%
   ggplot (aes (
       x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
   theme_bw() +
   geom_point(aes(shape = Standardization)) +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of modules per network",
        y = "Number of genes per module") +
   facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
   geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



# 3.3 Network analysis for Spearman-related correlations

Thresholds based on the Empirical Bayes approach to rank genes and determine if a gene is significantly expressed. Limma implementation.

†Same analysis with more stringent parameters

- Average Expression: 10
- · Adjusted P-value: equal or less than 0.030
- · Log Fold Change: 1
- B-statisitcs: 2

121

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129

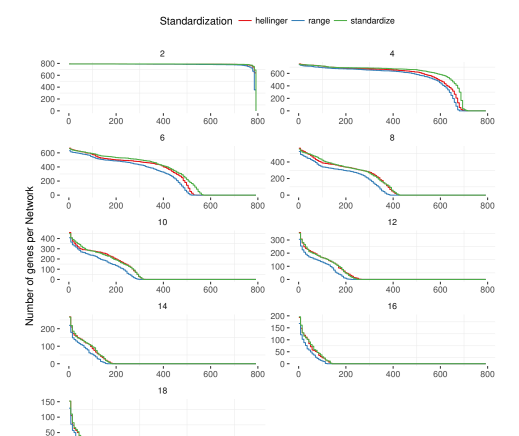
130

131

#### 3.3.1 Nodal versus extra-nodal lymphoma

Genetic networks from differentially expressed genes selected by comparing sample cases with nodal and extranodal lymphoma.

```
read.table("./data/networks.summary.119759.txt", header = TRUE) %>%
    ggplot (aes (
       x = MaxEdgesPerGene,
        y = NbNodes,
        fill = Standardization)) +
    theme_bw() +
    geom_step(aes(color = Standardization),
              stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



600

400

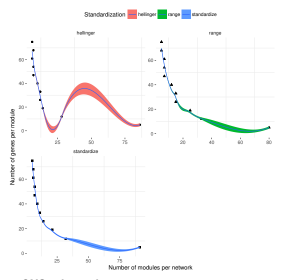
0 -0

132

200

```
read.table("./data/modules.summary.119759.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

800 Number of potential connections per gene



## 3.3.2 Relapsed versus no CNS relapsed cases

134

135

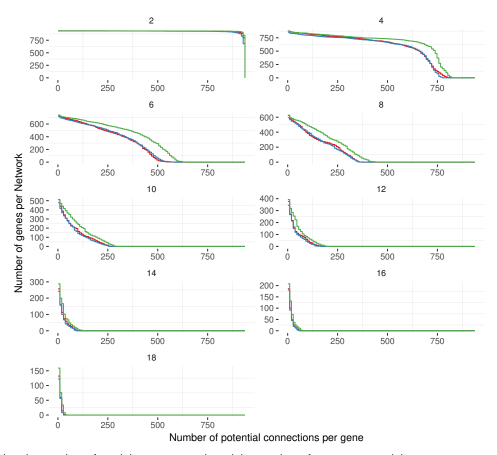
136

137

Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

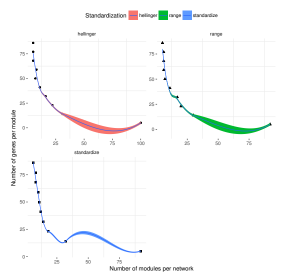
```
read.table("./data/networks.summary.119760.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
   facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```





138

```
read.table("./data/modules.summary.119760.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



## 3.3.3 Lymphoma cases classified by Cell-of-origin subtypes

140

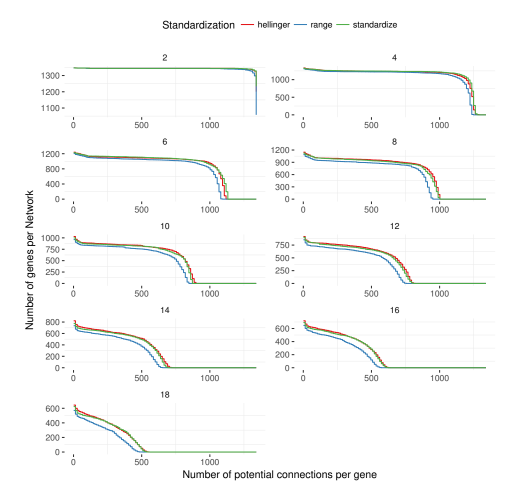
141

142

143

Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

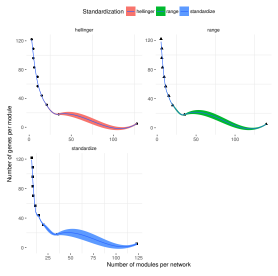
```
read.table("./data/networks.summary.119758.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



Showing the number of modules per network and the number of genes per module.

144

```
read.table("./data/modules.summary.119758.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



## 3.4 Network analysis for Pearson-related correlations

Thresholds based on the Empirical Bayes approach to rank genes and determine if a gene is significantly expressed. Limma implementation.

†Same analysis with more stringent parameters

• Average Expression: 10

Adjusted P-value: equal or less than 0.030

· Log Fold Change: 1

• B-statisitcs: 2

146

147

148

150

151

152

153

154

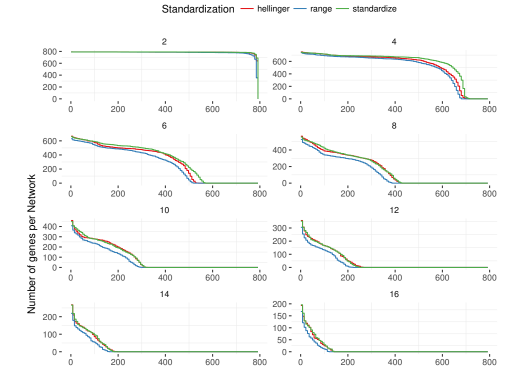
155

156

#### 3.4.1 Nodal versus extra-nodal lymphoma

Genetic networks from differentially expressed genes selected by comparing sample cases with nodal and extranodal lymphoma.

```
read.table("./data/networks.summary.119755.txt", header = TRUE) %>%
    ggplot (aes (
       x = MaxEdgesPerGene,
        y = NbNodes,
        fill = Standardization)) +
    theme_bw() +
    geom_step(aes(color = Standardization),
              stat = "identity") +
    facet_wrap( ~ CorrelationPower,
               ncol = 2,
               scales = "free") +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



157 Showing the number of modules per network and the number of genes per module. 158

600

18

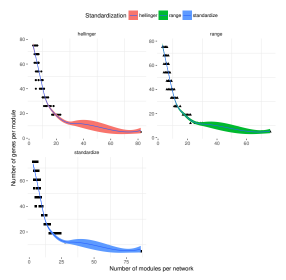
400

200

150 **-**100 -50 -0 -0

```
read.table("./data/modules.summary.119755.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

800 Number of potential connections per gene



## 3.4.2 Relapsed versus no CNS relapsed cases

159

160

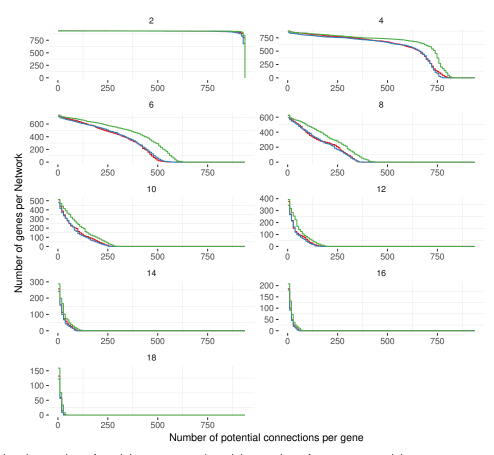
161

162

Genetic networks from differentially expressed genes selected by comparing sample cases with systemic or no CNS relapse lymphoma.

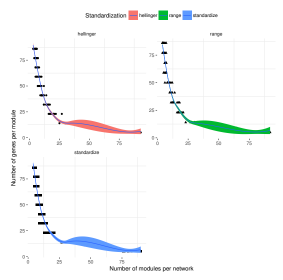
```
read.table("./data/networks.summary.119754.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
   facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
         panel.grid.major = element_line(linetype = "blank"))
```





163

```
read.table("./data/modules.summary.119754.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```



## 3.4.3 Lymphoma cases classified by Cell-of-origin subtypes

165

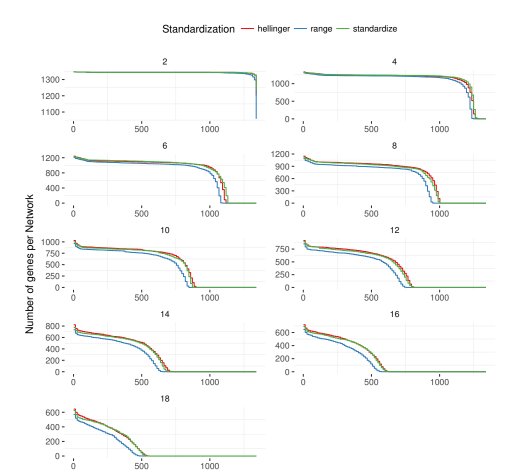
166

167

168

Genetic networks from differentially expressed genes selected by comparing sample cases with cell of origin classification based on ABC or GCB subtypes.

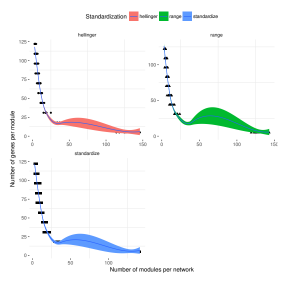
```
read.table("./data/networks.summary.119757.txt", header = TRUE) %>%
   ggplot (aes (
       x = MaxEdgesPerGene
       y = NbNodes,
       fill = Standardization)) +
   theme_bw() +
   geom_step(aes(color = Standardization),
             stat = "identity") +
    facet_wrap( ~ CorrelationPower,
              ncol = 2,
               scales = "free") +
   scale_color_brewer(type = "qual", palette = 6) +
   labs(x = "Number of potential connections per gene",
        y = "Number of genes per Network") +
   theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                          fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                      fill = NA),
          panel.grid.major = element_line(linetype = "blank"))
```



169

```
read.table("./data/modules.summary.119757.txt", header = TRUE) %>%
    ggplot (aes (
        x = NbModules,
        y = MaxGenesPerModule,
        fill = Standardization)) +
    theme_bw() +
    geom_point(aes(shape = Standardization)) +
    scale_color_brewer(type = "qual", palette = 6) +
    labs(x = "Number of modules per network",
         y = "Number of genes per module") +
    facet_wrap( ~ Standardization,
               ncol = 2,
               scales = "free") +
    theme(legend.position = "top",
          strip.background = element_rect(linetype = "blank",
                                           fill = "white"),
          panel.border = element_rect(linetype = "blank",
                                       fill = NA),
          panel.grid.major = element_line(linetype = "blank")) +
    geom_smooth(method = 'loess', size = .5, level = 0.5, alpha=1)
```

Number of potential connections per gene



## 4 System Information

171

The version number of R and packages loaded for generating the vignette were:

```
###save(list=ls(pattern=".*|.*"),file="PD.Rdata")
sessionInfo()
R version 3.4.4 (2018-03-15)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: elementary OS 0.4.1 Loki
Matrix products: default
BLAS: /usr/lib/libblas/libblas.so.3.6.0
LAPACK: /usr/lib/lapack/liblapack.so.3.6.0
locale:
[1] LC CTYPE=en US.UTF-8
                              LC NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
                              LC_COLLATE=en_US.UTF-8
 [5] LC_MONETARY=en_US.UTF-8
                              LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                               LC_NAME=C
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[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] stats
          graphics grDevices utils
                                          datasets methods
[7] base
other attached packages:
                        DescTools_0.99.23
[1] bindrcpp_0.2
                                          igraph_1.1.2
 [4] tidyr_0.7.2
                       dplyr_0.7.4
                                           ggplot2_2.2.1
 [7] latticeExtra_0.6-28 RColorBrewer_1.1-2 lattice_0.20-35
[10] leaps_3.0
                        gdata_2.18.0
                                           knitr_1.17
loaded via a namespace (and not attached):
 [1] Rcpp_0.12.13 pillar_1.1.0 compiler_3.4.4
 [4] plyr_1.8.4
                                     bindr_0.1
                    highr_0.6
                    digest_0.6.12 boot_1.3-20
 [7] tools_3.4.4
[10] evaluate_0.10.1 tibble_1.4.2
                                     manipulate_1.0.1
                  pkgconfig_2.0.1 rlang_0.1.2
[13] gtable_0.2.0
                   expm_0.999-2
                                   mvtnorm_1.0-6
[16] Matrix_1.2-11
[19] stringr_1.2.0 gtools_3.5.0
                                    grid_3.4.4
[22] glue_1.2.0
                   R6_2.2.2
                                    foreign_0.8-69
                   magrittr_1.5 scales_0.5.0
[25] purrr_0.2.4
                   assertthat_0.2.0 colorspace_1.3-2
[28] MASS_7.3-47
[31] labeling_0.3
                   stringi_1.1.5 lazyeval_0.2.1
[34] munsell_0.4.3
```