NA

1.0e+00

3.1e-03

4.9e-01

1.0e+00

3.1e-02

1.0e+00

4.9e-01

NA

NA

1.0e+00

1.0e+00

9.7e-01

1.0e+00

1.0e+00

1.0e+00

Rivera

NA

NA

4.9e-01

1.0e+00

7.2e-01

1.0e+00

1.0e+00

FVA_BD

NA

Abs - BD vs Li+

NA

NA

NA

NA

NA

1.0e+00

1.2e-09

7.2e-01

FVA_BD_NR

NA

NA

NA

NA

NA

NA

1.0e+00

1.8e-53

NA

NA

NA

NA

NA

NA

NA

1.0e+00

MTA_BD_R

NA

NA

NA

NA

1.0e+00

4.7e-01

1.0e+00

9.7e-01

FVA_BD_R

Lanz

Rivera

FVA_BD

FVA_BD_R

FVA BD NR

MTA_BD

MTA_BD_R

MTA_BD_NR

NA

NA

NA

NA

NA

NA

NA

NA

MTA_BD_NR

Lanz NA NA NA NA NA NA NA NA 1.0e+00 NA NA NA NA NA NA NA Rivera

1.7e-02 1.0e+00 NA NA NA NA NA NA FVA_BD

Norm_T1 - BD vs Li+

1.0e+00

1.0e+00

1.0e+00

1.0e+00

FVA_BD_R

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e+00

2.4e-01

2.7e-01

6.0e-01

Rivera

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e+00

FVA_BD

NA NA NA

NA

1.0e+00

1.0e+00

1.0e+00

FVA_BD_NR

NA

NA

3.1e-42

1.0e+00

FVA_BD_R

FVA BD NR

MTA_BD

MTA_BD_R

MTA_BD_NR

NA

NA

NA

NA

1.0e+00

NA

NA

NA

NA

NA

MTA_BD_NR

Lanz NA NA NA NA NA NA NA NA 1.0e+00 NA NA NA NA NA NA NA Rivera

NA

NA

1.0e+00

1.0e+00

1.0e+00

FVA_BD_NR

NA

NA

NA

1.0e+00

1.0e+00

NA

NA

NA

NA

1.0e+00

MTA_BD_R

NA

NA

NA

NA

NA

MTA_BD_NR

FVA_BD_R

FVA BD NR

MTA_BD

MTA_BD_R

MTA_BD_NR

1.0e+00 1.0e+00 NA NA NA NA NA NA FVA_BD

Norm_T2 - BD vs Li+

NA

1.0e+00

1.0e+00

1.0e+00

6.9e-02

FVA_BD_R

1.0e+00

1.0e+00

2.2e-03

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e+00

1.0e-01

1.0e+00

Rivera

1.1e-54

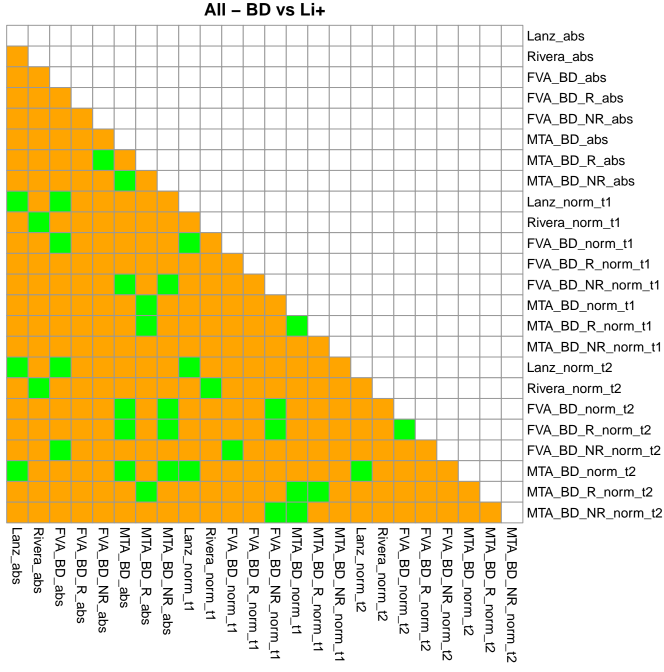
1.0e+00

1.0e+00

1.0e+00

1.0e-01

FVA_BD



Abs 0 0 0 0 0 0 0 0 0

0

0

0

1

1

1

1

Transport, endoplasmic reticular Keratan sulfate synthesis Transport, extracellular ROS detoxification Starch and sucrose metabolism Fructose and mannose metabolism Valine, leucine, and isoleucine metabolism Miscellaneous

Tyrosine metabolism

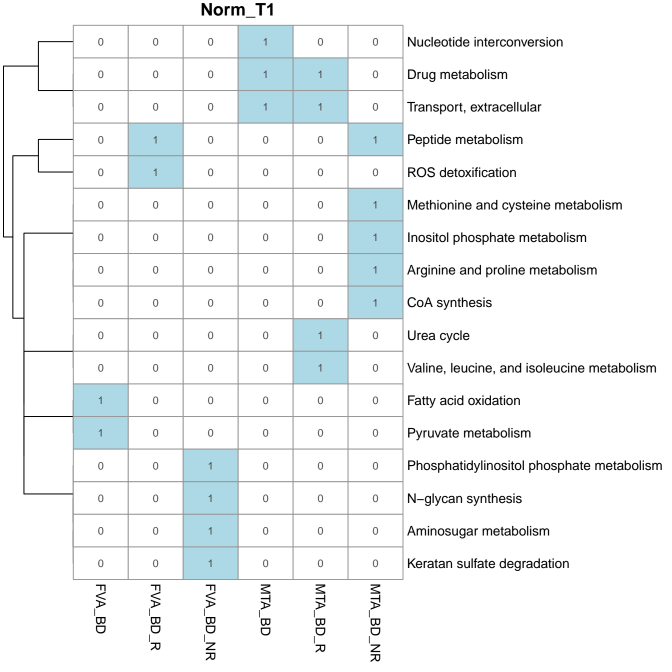
Keratan sulfate degradation

Chondroitin sulfate degradation

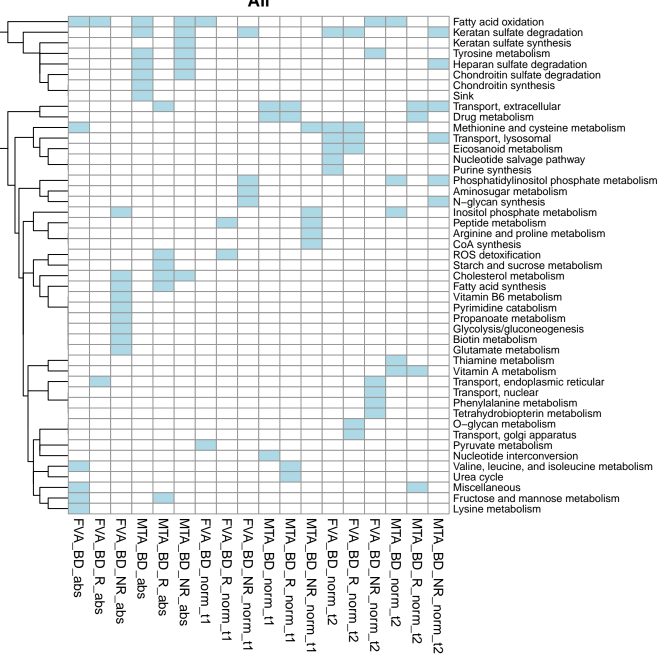
Sink Fatty acid oxidation Lysine metabolism Methionine and cysteine metabolism Cholesterol metabolism Fatty acid synthesis Vitamin B6 metabolism Pyrimidine catabolism Propanoate metabolism Inositol phosphate metabolism Glycolysis/gluconeogenesis

Heparan sulfate degradation Chondroitin synthesis Biotin metabolism Glutamate metabolism

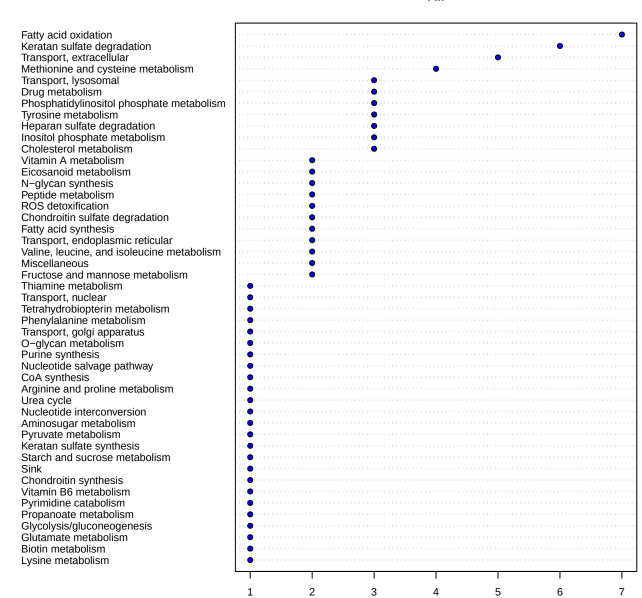
	FVA_BD	FVA_BD_R	FVA_BD_NR	MTA_BD	MTA_BD_R	MTA_BD_NR
	0	0	1	0	0	0
	0	0	1	0	0	0
	0	0	1	0	0	0
	0	0	1	0	0	0
	0	0	1	0	0	0
	0	0	1	0	0	0
4	0	0	1	0	0	0
	0	0	1	0	1	0
	0	0	1	0	1	1
	1	0	0	0	0	0
	1	0	0	0	0	0
	1	0	0	0	0	0
4	1	0	0	0	0	0
	- 1	0	0	0	1	0
	0	0	0	0	1	0
	0	0	0	0	1	0
	0	0	0	0	1	0
	- 0	0	0	0	0	1
	0	1	0	0	0	0
	1	1	0	1	0	1
	0	0	0	1	0	0
	0	0	0	1	0	0
1	0	0	0	1	0	1
					9	



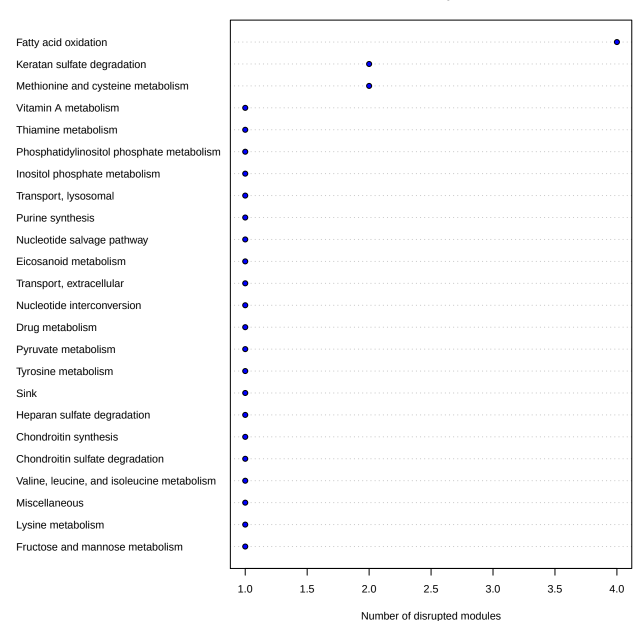


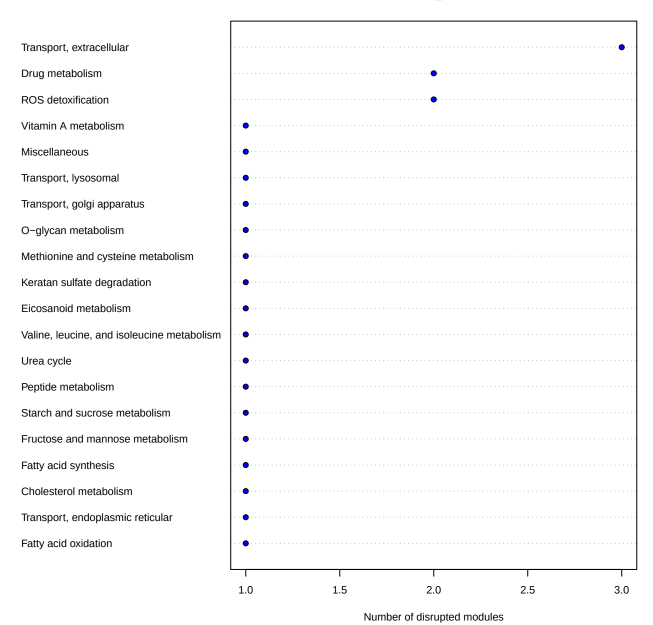


Number of disrupted modules



BD_Lumped





BD_NR

