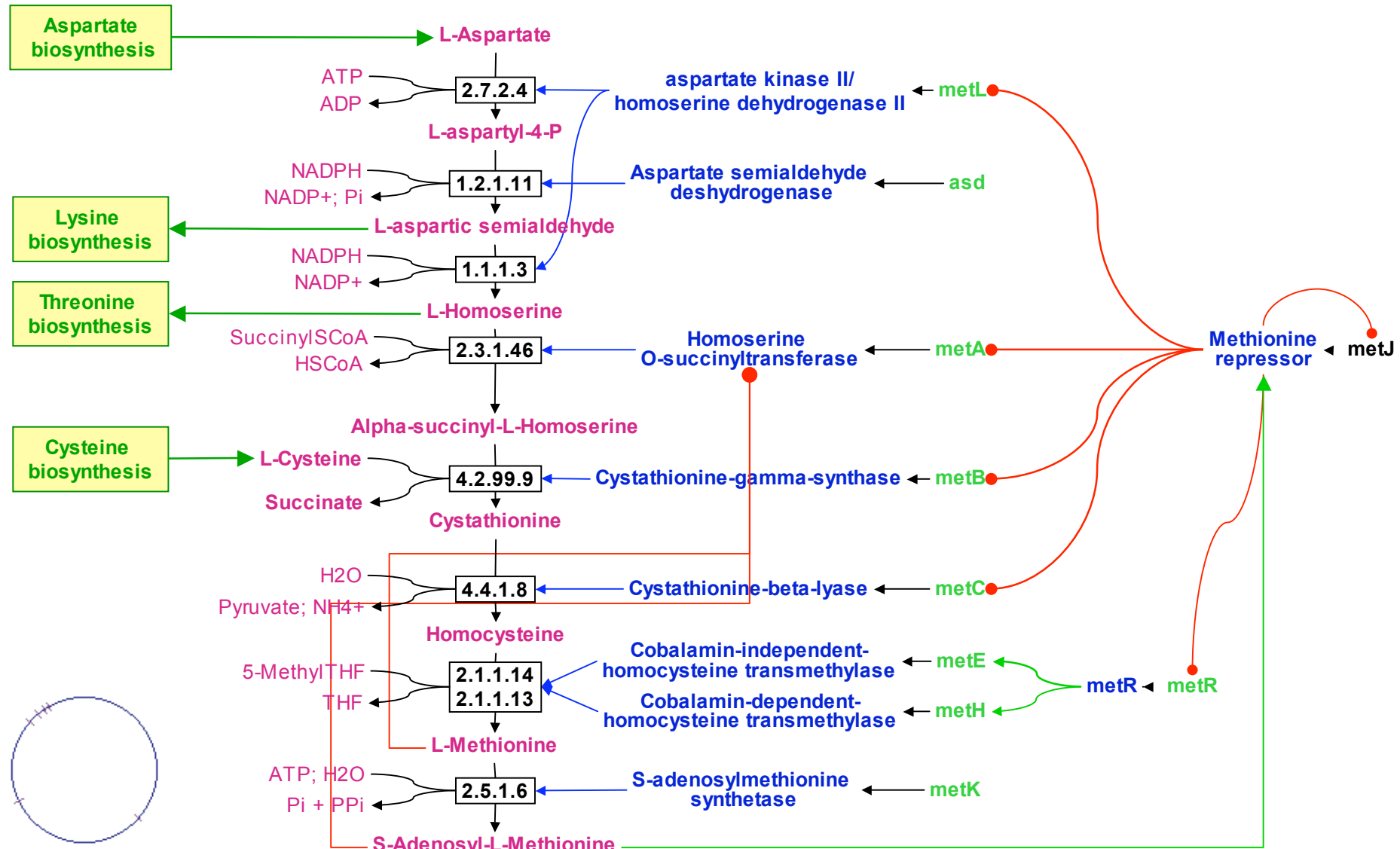


Regulatory Sequence Analysis

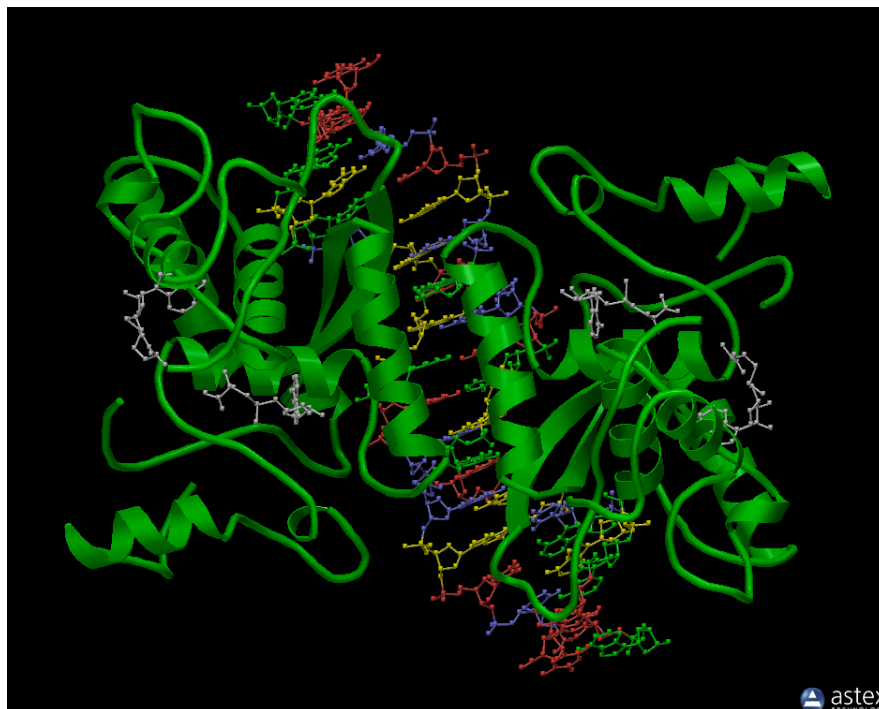
***From phylogenetic footprints
to co-regulation networks***

Methionine Biosynthesis in E.coli

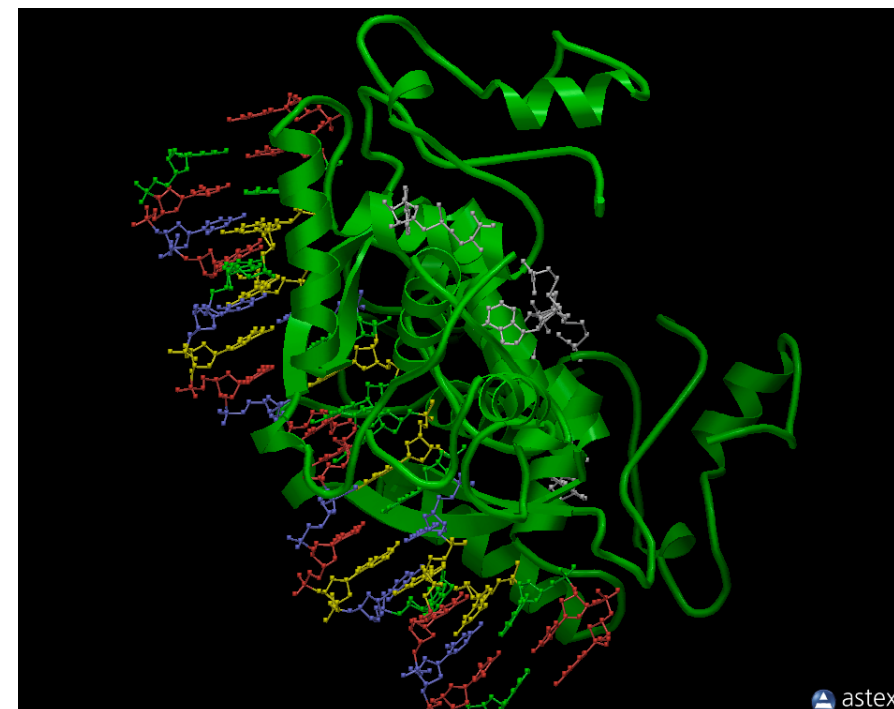


Methionine repressor

- Structure cristalline du répresseur de la méthionine chez *Escherichia coli*.
- En vert: la protéine MetJ forme un dimère qui se lie à l'ADN
- La séquence d'ADN est colorée par nucléotide
- En gris: molécules de méthionine liées au répresseur (elles activent le répresseur)



astex
TECHNOLOGY

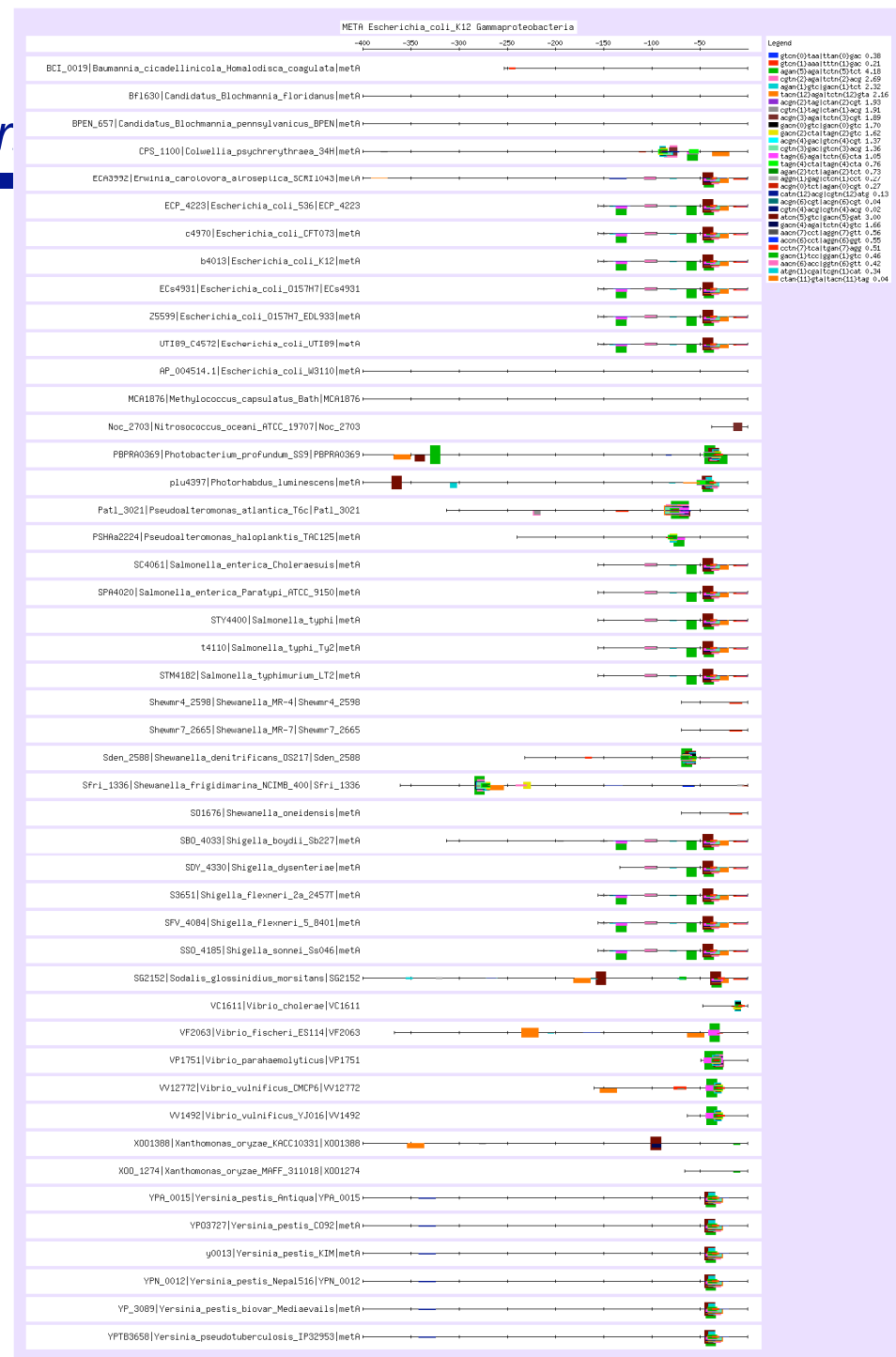


astex
TECHNOLOGY

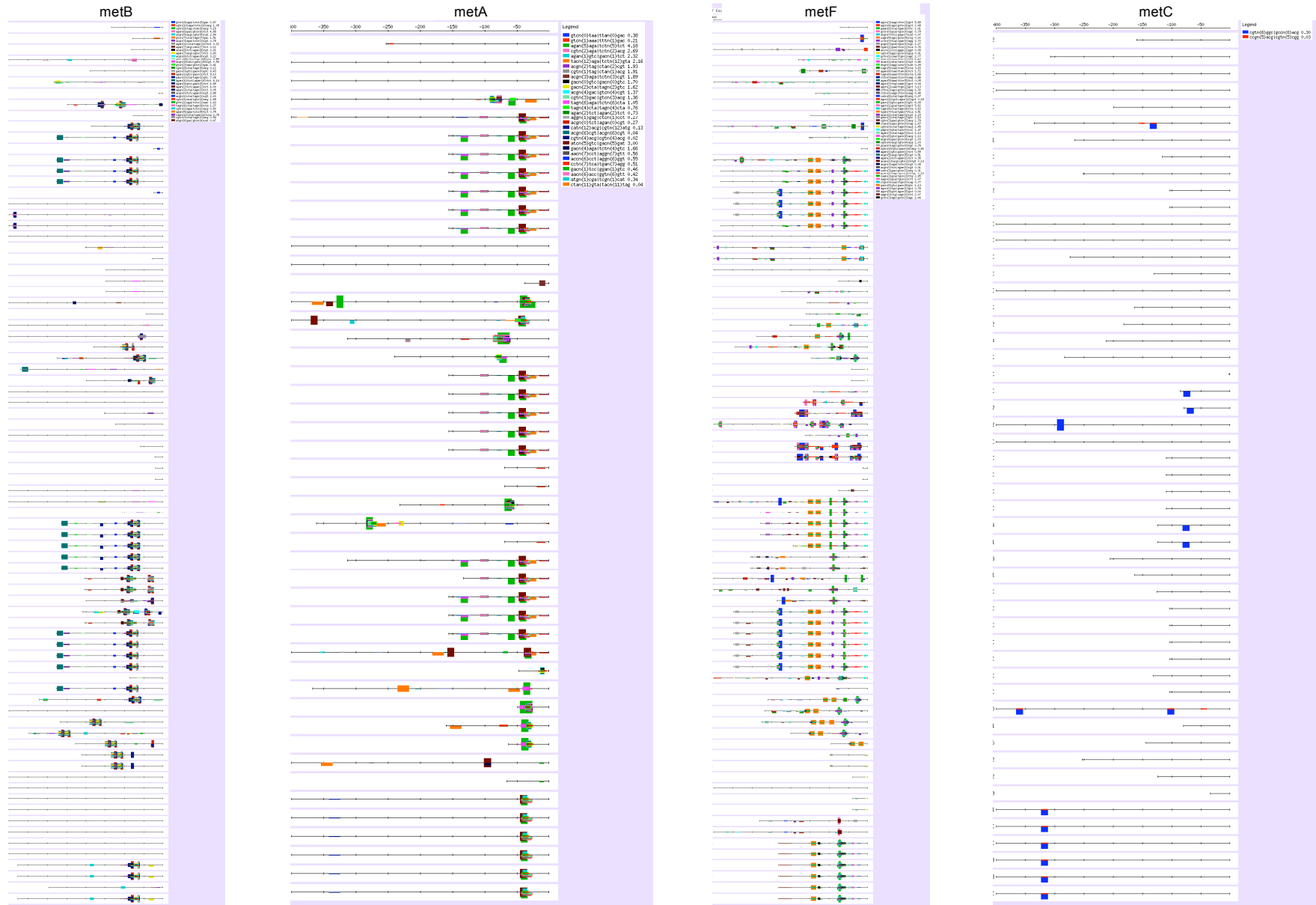
Phylogenetic footprints

MetA orthologs in Gammaproteobacteria

atcnnnnnngtc.....gacnnnnngat	3.00
.tctnnnnntct....agannnnnaga.	4.18
.tctnnnacg.....cgtnnaga.	2.69
.tctnnncgt.....acgnnnaga.	1.89
.tctnnnnngtc.....gacnnnnaga.	1.66
.tctnnnnnncta...	...tagnnnnnaga.	1.05
..ctanncgt.....acgnntag..	1.93
..ctanacg.....cgtn>tag..	1.91
...tagnngtc.....gacnncta...	1.62
...tagnnnnnnaga.	.tctnnnnnncta...	1.05
...tagnnnnncta...	...tagnnnncta...	0.76
....agannnnnaga.	.tctnnnnntct....	4.18
....agangtc.....gacntct....	2.32
....aganntct....agannntct....	0.73
....agacgt.....acgtct....	0.27
....gacnnnnngat	atcnnnnnngtc.....	3.00
....gacntct....agangtc.....	2.32
....gacgtc.....gacgtc.....	1.70
....gacnnnnnaga.	.tctnnnnngtc.....	1.66
....gacnncta...	...tagnngtc.....	1.62
....acgnntag..	..ctanncgt.....	1.93
....acgnnnnaga.	.tctnnncgt.....	1.89
....acgtct....agacgt.....	0.27
....cgtnnaga.	.tctnnnacg.....	2.69
....cgtn>tag...	..ctanacg.....	1.91
atctagacgtctagat	atctagacgtctagat	4.18

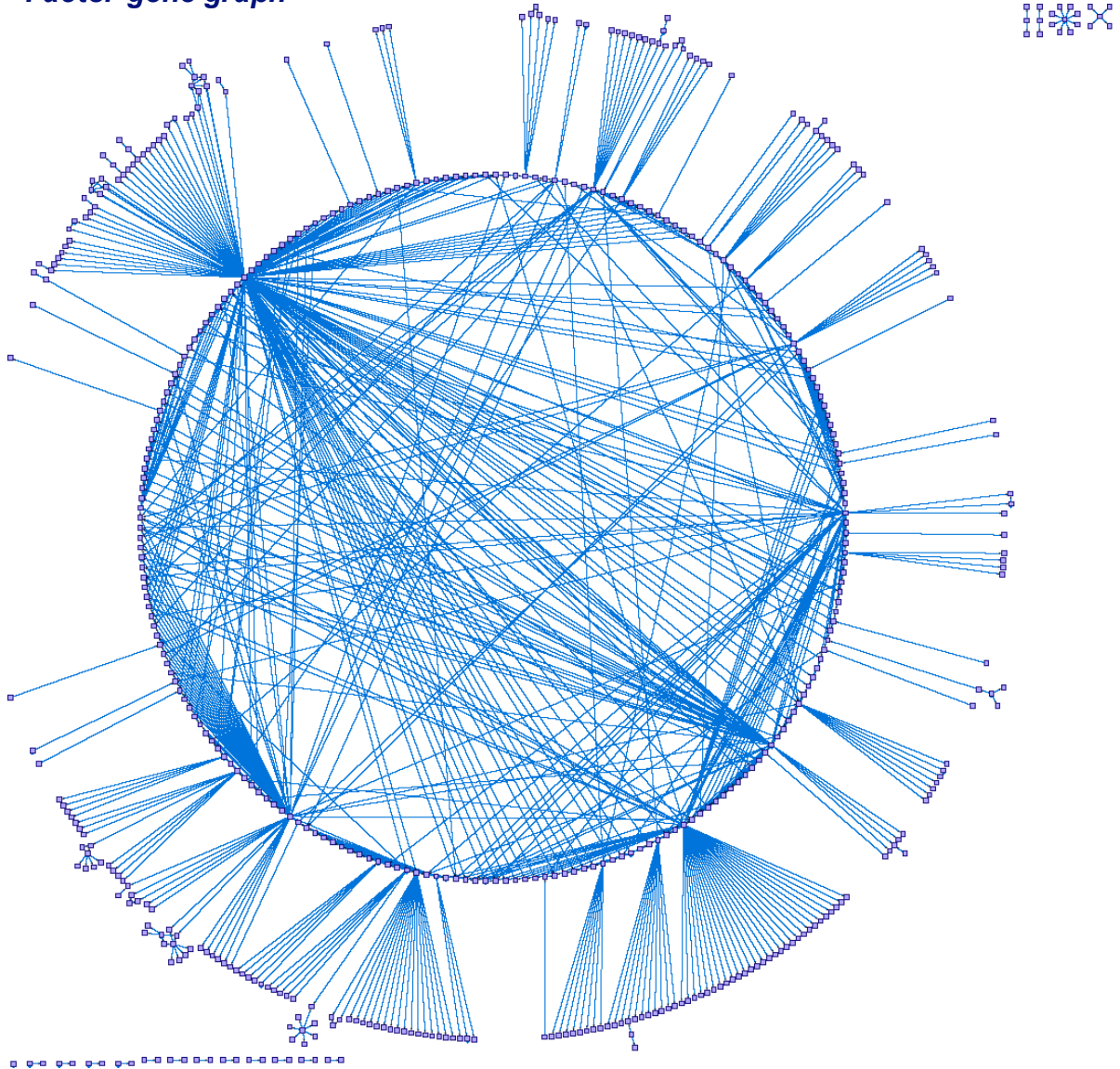


Phylogenetic footprints *MET genes in Gammaproteobacteria*



RegulonDB factor -> gene network

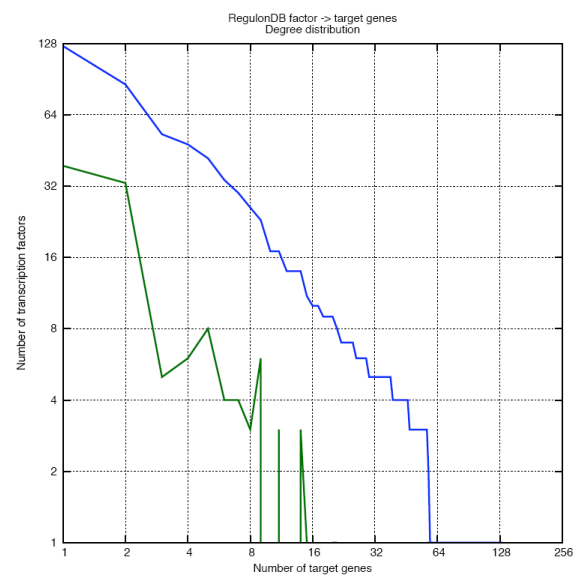
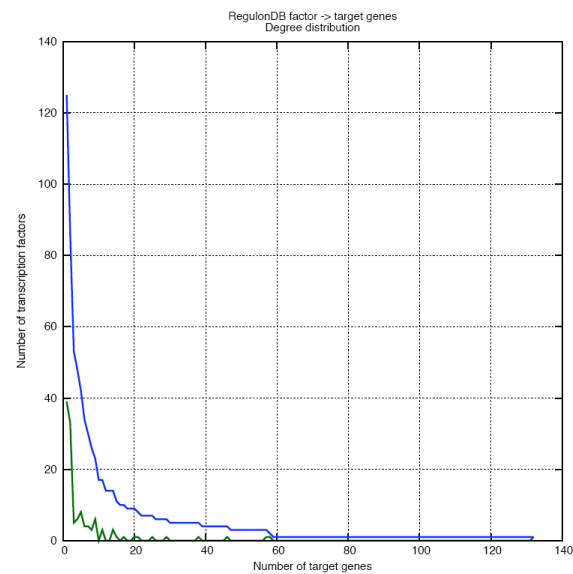
Factor-gene graph



- RegulonDB (Oct. 2005 version)
- The graph represents the relationships between factors and their target genes (factor -> gene graph)
 - 125 transcription factors
 - 467 target genes
 - 847 factor->gene interactions
 - 45 self-regulations
 - Note: CRP alone regulates 132 target genes.

Number of target genes per transcription factor

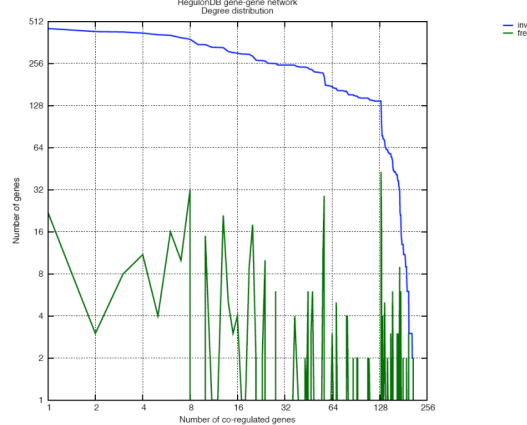
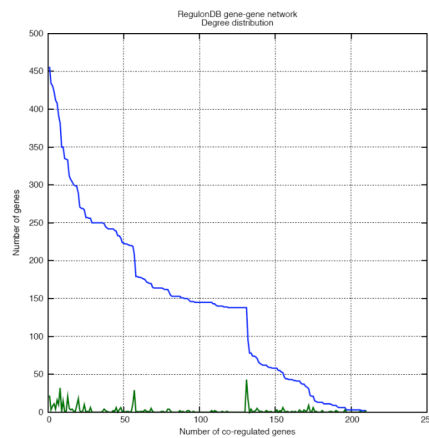
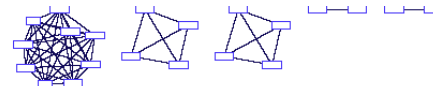
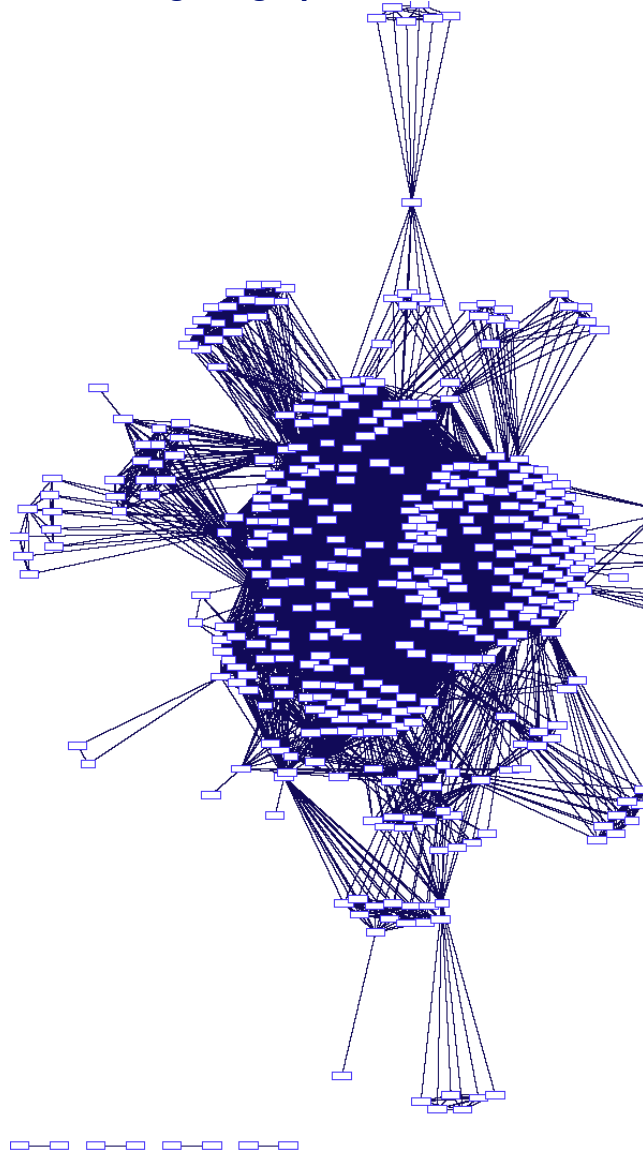
Factor	Targets
CRP	132
FIS	58
IHF	57
FNR	46
ARCA	38
H-NS	29
LRP	25
FUR	21
NARL	20
SOXS	17
GADE	15
PURR	14
NTRC	14
MARA	14
GADX	11
FRUR	11
CPXR	11
PHOB	9
OXYR	9
OMPR	9
LEXA	9
FLHD	9
ARGR	9
TYRR	9
FADR	8
CYTR	8
ROB	7
NAC	7
METJ	7
CYSB	7
NAGC	6
MODE	6
GNTR	6
DNAA	6
TRPR	5
TORR	5



- The number of target genes per factor (out-degree) shows important variations:
 - **Specific factors:** most transcription factors have a very few target genes (typically 1 to 3)
 - **General factors:** a few factors have a very large number of target genes.
- The log-scale representation shows that the distribution of degree more or less follows a power law.

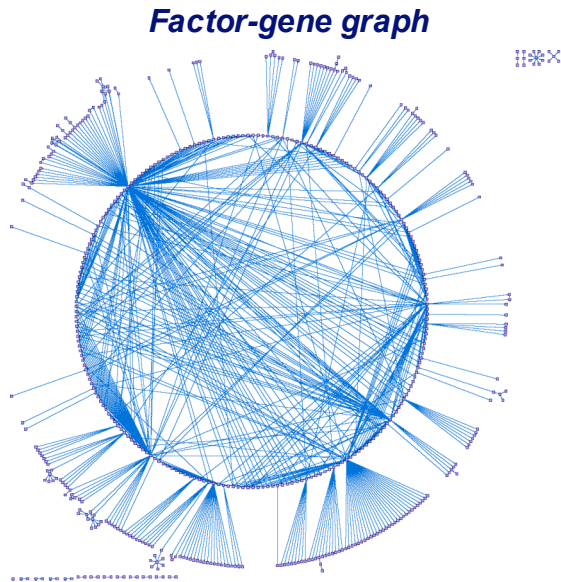
RegulonDB gene <-> gene network

Gene-gene graph

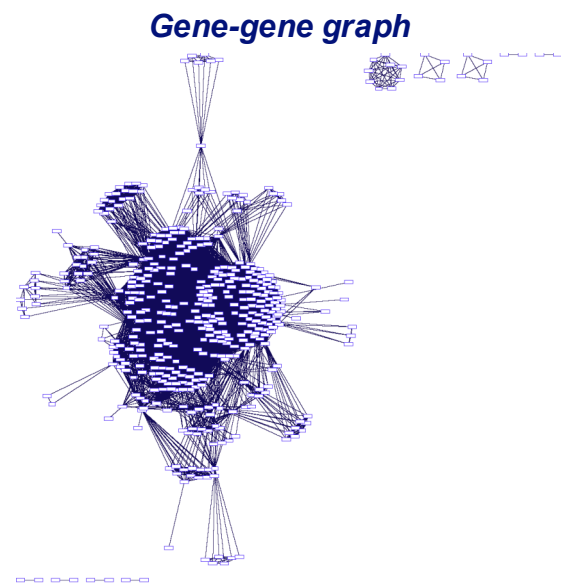


- The factor \rightarrow gene graph can be converted in a gene \leftrightarrow gene graph, with an edge joining each pair of co-regulated genes (bottom).
- The gene-gene co-regulation graph contains
 - 458 nodes (genes).
 - Note: some genes are the only target of a given factor, they are thus not present in this graph.
 - 14,939 arcs (co-regulation between a pair of genes).
- Note
 - $(132 \times 131)/2 = 8,646$ of these arcs correspond to links via CRP.
 - This network **does not** follow a power-law: there are many genes having many neighbours (all those regulated by some general transcription factor)

RegulonDB networks



- RegulonDB (Oct. 2005 version)
- The top figure displays the interaction graph between factors and genes (factor -> gene graph)
 - 125 transcription factors
 - 467 target genes
 - 847 factor->gene interactions
 - 45 self-regulations
 - Note: CRP alone regulates 132 target genes.



- The factor -> gene graph can be converted in a gene <-> gene graph, with an edge joining each pair of co-regulated genes (bottom).
- The gene-gene co-regulation graph contains
 - 458 nodes (genes).
 - Note: some genes are the only target of a given factor, they are thus not present in this graph.
 - 14,939 arcs (co-regulation between a pair of genes).
 - Note : $(132 \times 131)/2 = 8,646$ of these arcs correspond to links via CRP.

Testing the principle with RegulonDB

Gene	Factor	GI
metN	MetJ	1786398
ahpC	MetJ	1786822
metK	MetJ	1789311
metC	MetJ	1789383
metB	MetJ	1790375
metF	MetJ	1790377
metA	MetJ	1790443
glyA	MetR	1788902
metH	MetR	1790450
mhpA	MhpR	1786543
purE	PurR	1786734
pyrD	PurR	1787177
pyrC	PurR	1787301
hflD	PurR	1787377
purR	PurR	1787948
cvpA	PurR	1788652
purC	PurR	1788820
purM	PurR	1788845
guaB	PurR	1788855
glyA	PurR	1788902
glnB	PurR	1788904
gcvT	PurR	1789272
purH	PurR	1790439
purL	PurR	48994899
putA	PutA	1787250
...

- Test
 - Discover motifs in upstream sequences of its orthologous genes of each gene annotated in regulonDB.
 - Link genes with similar discovered motifs.
 - Compare the inferred links with the regulon membership
- Small test case
 - Genes regulated by methionine (MetJ and MetR targets), purine (PurR targets) and proline (PutR, PutA targets)
- Complete test case
 - All the genes annotated as target gene or TF-coding in regulonDB (537 genes)

Dyad-profiles

- We can collect the significant dyads found for each gene, and display this information as “dyad profiles”:
 - Each row represents one gene
 - Each column represents one dyad found significant in at least one gene
 - Dots indicate that a dyad was not significant for a given gene (`occ_sig < 0`)
 - Note that the majority of the cells are empty.

	aaan{0}aat	aagn{0}ggg	aagn{1}gga	aagn{1}tgc	aagn{2}gag	aagn{2}gca	aagn{3}agc	aagn{4}gcc	aagn{6}cca	acgn{0}caa	acgn{0}tct	acgn{1}cta	acgn{1}ttg	acgn{2}aac	acgn{2}tgc	acgn{3}acg	acgn{4}cgt	actn{9}gag	agan{10}tct	0.7	2.7	...	
META
METB	2.6	2.2	...	
METF	0.0	0.5	
METH
METJ	2.7	2.1	1.4	3.5	.	.	.	
METK	5.5	
METN	0.5	
METR	0.1	
PURA	
PURC	.	1.2	0.9	0.9	0.7	0.1	0.6	
PURE	0.1	.	1.5	.	0.1	
PURH	0.2	.	0.9	2.5	
PURL	1.0	
PURM	
PURR	0.2	
PUTA	.	.	0.6	4.2	
PUTP	1.8	.	.	2.1	
...	

Compare-profiles

- The program ***compare-profiles*** was originally designed to compare binary profiles.
- It takes as input a profile file and compares each pair of profiles Q and R.
 - Profile intersection (QR),
 - union (QvR),
 - differences (Q!R, R!Q),
 - common exclusion (!Q!R)
 - Jaccard similarity = intersection/union
 - Significance of the intersection QR
 - hypergeometric significance test on the right tail : P-value + E-value + sig
 - Significance of the common exclusion !Q!R
 - hypergeometric significance test on the left tail : P-value + E-value + sig
 - Mutual information
- This program can be used to detect co-occurrence or mutual exclusion of genes across phylogenetic profiles.
- We can also use it to compare dyad profiles
 - Convert significance scores to a Boolean value

Converting dyad profiles to dyad classes

- Analysis of RegulonDB (537 genes)
 - 422 genes with at least one motif
 - 4944 dyads found significant in at least one gene
 - The profile table contains 422 rows x 4944 columns = 2,086,368 cells.
 - No more than 6,709 of these 2,086,368 cells are non-empty.
- Complete analysis of Escherichia coli (4,200 genes)
 - 2,844 genes with at least one significant motif
 - $2,844 \times 17,153$ dyads = 48,783,132 cells
 - No more than 33,370 of these 48,783,132 cells are non-empty.
- The profile profile matrices are very sparse.
- A (much) cheapest structure to store the same information is a 3 column file
 - dyad
 - gene
 - significance score

dyad	gene	occ_sig
gacn{0}gtc	META	0.16
agan{1}gtc	META	0.65
atcn{5}gtc	META	0.75
agan{5}aga	META	2.74
agan{1}gtc	METB	2.17
agan{10}tct	METB	2.56
gacn{0}gtc	METB	2.96
aagn{2}gca	METF	0.01
caan{3}gca	METF	0.35
gcan{0}agc	METF	0.35
acgn{1}cta	METF	0.52
agcn{2}caa	METF	0.64
gacn{0}gtc	METF	1.64
tagn{5}taa	METF	1.95
agcn{3}aag	METF	2.15
tgan{7}tca	METF	2.83
agcn{1}gca	METF	3.02
tgan{7}tca	METH	0.11
agan{6}gac	METJ	0.15
agan{9}gtc	METJ	0.15
cgtn{0}cta	METJ	0.48
cgtn{2}aga	METJ	1.04
agan{10}tct	METJ	1.35
gacn{2}cta	METJ	1.9
acgn{1}cta	METJ	2.05
acgn{0}tct	METJ	2.73
gacn{0}gtc	METJ	3.1
agan{1}gtc	METJ	3.46
taan{0}aaa	METK	0.1
gacn{0}gtc	METK	0.44
cgtn{2}aga	METK	0.51
...

Compare-classes

- The program **compare-classes** was designed to compare clusters
 - Clusters of co-expressed genes versus GO classes
 - Clusters of co-expressed genes versus annotated regulons
 - Annotated regulons versus annotated regulons (to detect synergy between transcription factors)
 - Chip-on-chip data versus chip-on-chip data
- We will now use compare-classes to compare the sets of dyads between genes.
- For each pair of genes Q and R, compare-classes calculates
 - number of dyads found significant in Q
 - number of dyads found significant in R
 - number of dyads in the intersection (QR)
 - number of dyads in the union (QvR)
 - Jaccard similarity = intersection/union
 - dot product (see next slide)
 - significance test on this intersection (hypergeometric, right tail)
 - P-value, E-value, sig

Gene pairs with all the RegulonDB genes

Top scoring by dot product

Rank	gene R	gene Q	R	Q	QR	QvR	jac_sim	dotprod	P_val	E_val	sig
1	PUTP	PUTA	38	37	25	50	0.5	2542.1	7.10E-49	6.30E-44	43.2
2	RECA	LEXA	57	101	14	144	0.09722	1889.8	3.30E-12	2.90E-07	6.53
3	GLNL	GLNA	51	88	32	107	0.29907	1713.11	7.00E-46	6.20E-41	40.21
4	UDP	MTLA	15	54	11	58	0.18966	1043.88	1.20E-19	1.10E-14	13.98
5	UVRD	LEXA	41	101	11	131	0.08397	831.49	2.80E-10	2.50E-05	4.6
6	TYRP	AROF	25	32	11	46	0.23913	815.729	5.10E-19	4.50E-14	13.34
7	PUTP	GLNA	38	88	3	123	0.02439	628.522	0.02936	2608.27	-3.42
8	MTLA	MLC	54	11	5	60	0.08333	448.632	5.70E-08	0.00503	2.3
9	PSPF	PSPA	64	64	64	64	1	437.011	#####	6.40E-143	142.19
10	PUTA	GLNA	37	88	3	122	0.02459	431.504	0.02739	2432.93	-3.39
11	ILVY	ILVC	93	75	51	117	0.4359	386.541	9.00E-77	8.00E-72	71.1
12	SULA	RECA	41	57	11	87	0.12644	358.099	4.20E-13	3.70E-08	7.43
13	UVRD	RECA	41	57	10	88	0.11364	347.532	1.60E-11	1.40E-06	5.86
14	UVRA	LEXA	12	101	5	108	0.0463	346.642	2.30E-06	0.20236	0.69
15	PUTP	PDHR	38	59	1	96	0.01042	330.398	0.36741	32637.392	-4.51
16	POLA	DNAA	39	32	15	56	0.26786	296.35	6.80E-25	6.10E-20	19.22
17	BIOB	BIOA	31	36	22	45	0.48889	287.556	4.70E-44	4.20E-39	38.38
18	METJ	METB	45	23	19	49	0.38776	286.685	1.70E-37	1.50E-32	31.81
19	MTLA	CYDA	54	19	3	70	0.04286	281.577	0.00105	93.699	-1.97
20	PUTA	PDHR	37	59	1	95	0.01053	269.359	0.35971	31953.518	-4.5
21	NAGE	MTLA	19	54	6	67	0.08955	227.63	3.10E-08	0.00276	2.56
22	UDP	MLC	15	11	5	21	0.2381	200.044	5.60E-11	5.00E-06	5.3
23	MTLA	ADHE	54	8	3	59	0.05085	197.458	6.60E-05	5.896	-0.77
24	MALK	MALE	21	73	19	75	0.25333	197.254	2.70E-34	2.40E-29	28.62
25	NRDD	NRDA	39	63	20	82	0.2439	192.165	2.60E-29	2.30E-24	23.63
26	YFID	MTLA	29	54	5	78	0.0641	183.569	1.30E-05	1.113	-0.05
27	YFID	CYDA	29	19	6	42	0.14286	165.44	6.10E-10	5.40E-05	4.27
28	SULA	LEXA	41	101	4	138	0.02899	164.636	0.00931	826.926	-2.92
29	MTLA	CDD	54	14	4	64	0.0625	162.407	1.20E-05	1.043	-0.02
30	UVRA	RECA	12	57	5	64	0.07812	160.535	1.30E-07	0.01127	1.95
31	SSB	LEXA	22	101	2	121	0.01653	159.836	0.07326	6507.387	-3.81
32	CUER	COPA	43	47	29	61	0.47541	157.892	2.40E-53	2.20E-48	47.67
33	UHPT	MTLA	12	54	3	63	0.04762	157.165	0.00025	22.457	-1.35
34	MTLA	AGAR	54	31	8	77	0.1039	154.984	7.70E-10	6.80E-05	4.17
35	PUTP	GLNL	38	51	2	87	0.02299	150.447	0.05794	5147.255	-3.71
36	TYRR	AROF	29	32	6	55	0.10909	147.935	1.90E-08	0.0017	2.77
37	XSEA	GUAB	29	37	13	53	0.24528	139.955	1.30E-21	1.20E-16	15.92
38	ULAA	MTLA	27	54	3	78	0.03846	132.93	0.00299	265.909	-2.42
39	MTLA	GLPA	54	15	5	64	0.07812	130.283	3.60E-07	0.03161	1.5
40	PSTS	PHOB	28	31	7	52	0.13462	130.01	2.00E-10	1.80E-05	4.75
41	GLNK	GLNA	14	88	6	96	0.0625	125.801	7.20E-08	0.00637	2.2
42	MALP	MALE	19	73	11	81	0.1358	124.547	2.30E-16	2.00E-11	10.69
43	ARGE	ARGC	38	24	14	48	0.29167	122.146	3.10E-25	2.70E-20	19.56
44	PDHR	GLCC	59	13	7	65	0.10769	118.735	3.90E-11	3.40E-06	5.46
45	TYRR	TYRP	29	25	6	48	0.125	114.252	3.90E-09	0.00034	3.47
46	METN	FPR	79	24	13	90	0.14444	113.54	3.40E-18	3.10E-13	12.51
47	MTLA	MALE	54	73	8	119	0.06723	112.418	9.20E-07	0.08208	1.09
48	PDHR	FADB	59	14	5	68	0.07353	110.459	3.80E-07	0.03338	1.48
49	TREB	MTLA	36	54	3	87	0.03448	107.764	0.00682	605.69	-2.78
50	PUTA	GLNL	37	51	2	86	0.02326	103.141	0.05525	4907.985	-3.69

Dot product

$$dp = \sum_{i=1}^p (x_{Ai} \cdot x_{Bi})$$

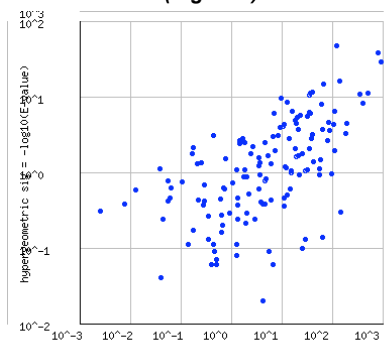
Hypergeometric significance

$$P(X = x) = \frac{C_m^x C_{n-m}^{k-x}}{C_m^k}$$

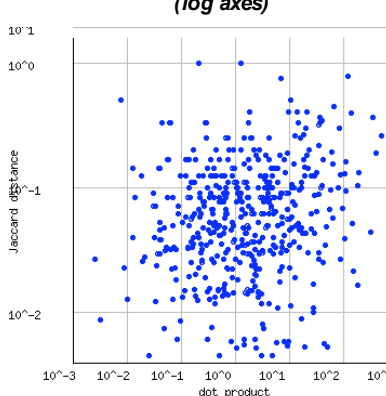
Jaccard similarity

$$S_J = \frac{|Q \cap R|}{|Q \cup R|}$$

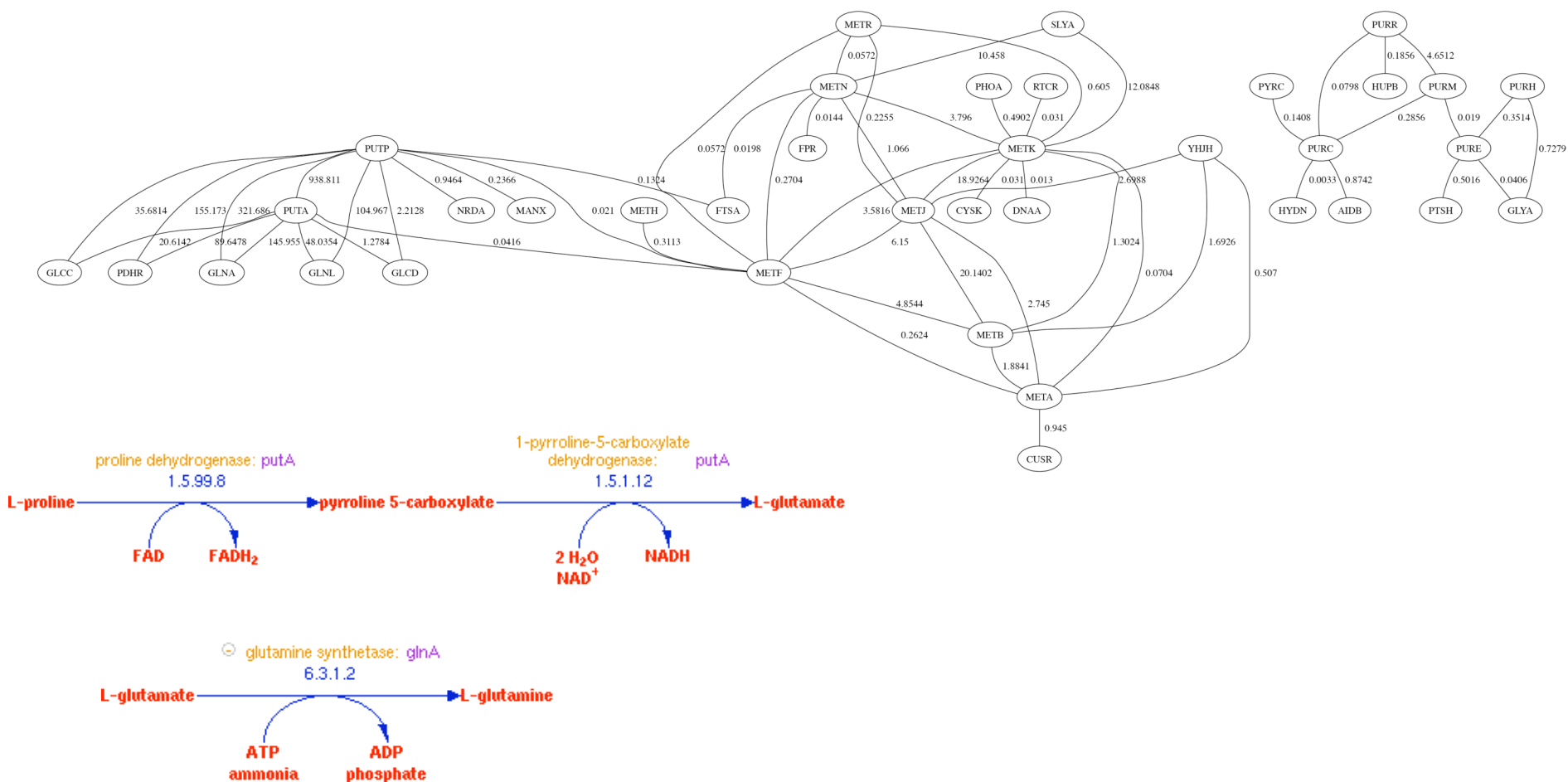
Dot product versus hypergeometric (log axes)



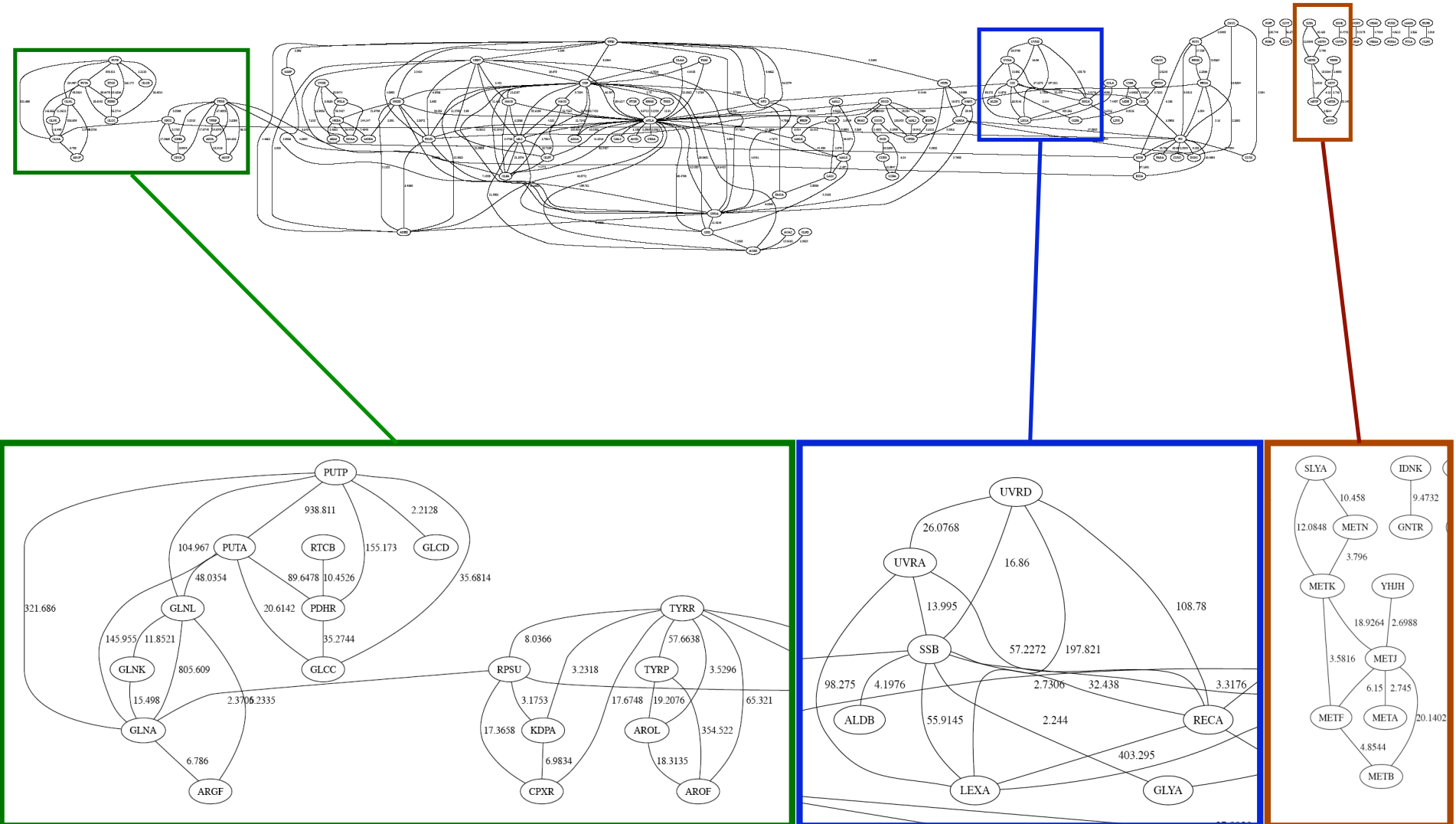
Dot product versus Jaccard (log axes)



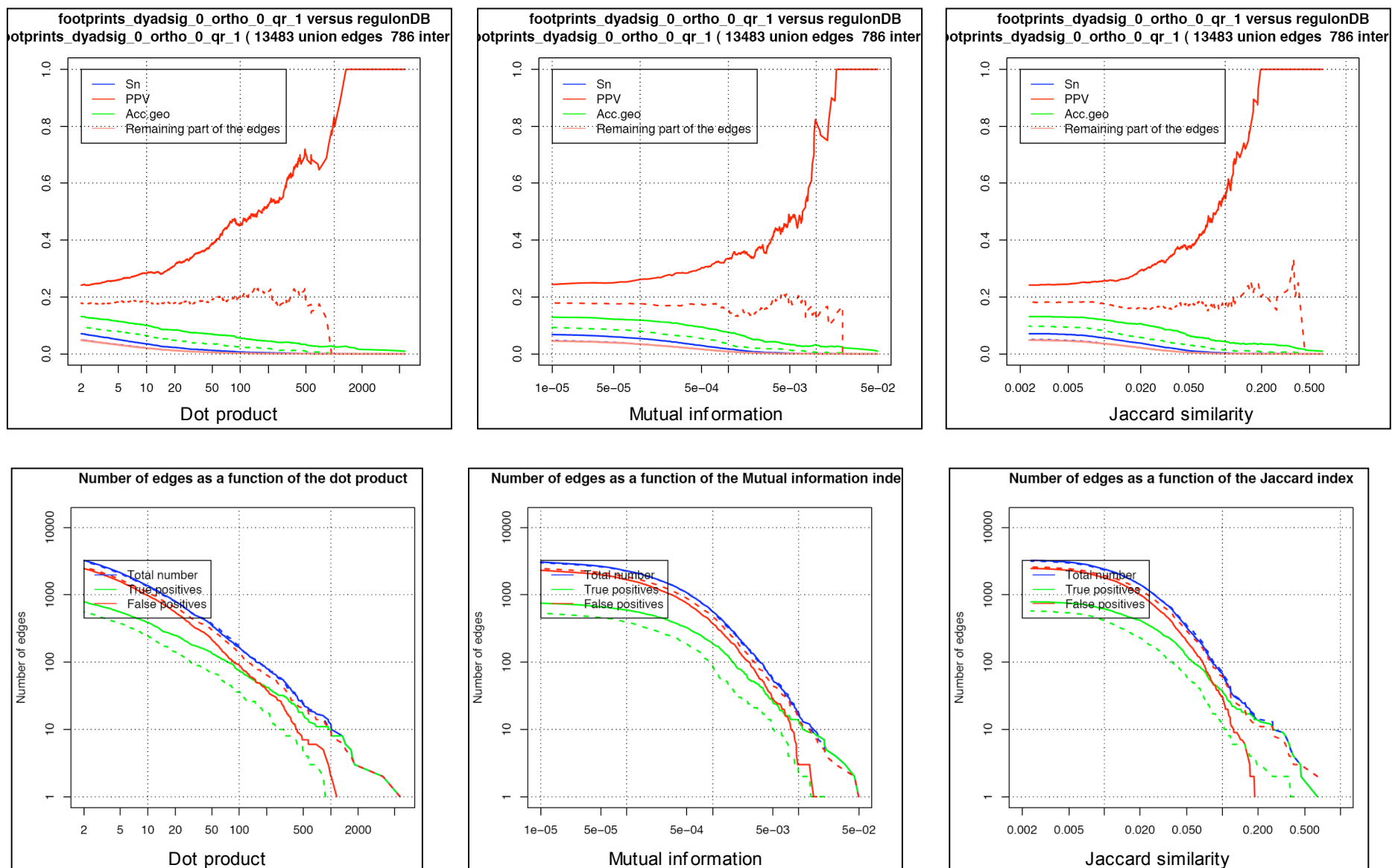
Inferred co-regulation graph (selection)



All the pairs of dyad profiles with $dp \geq 2$



Validation of inferred co-regulation network



Predicted versus annotated gene pairs

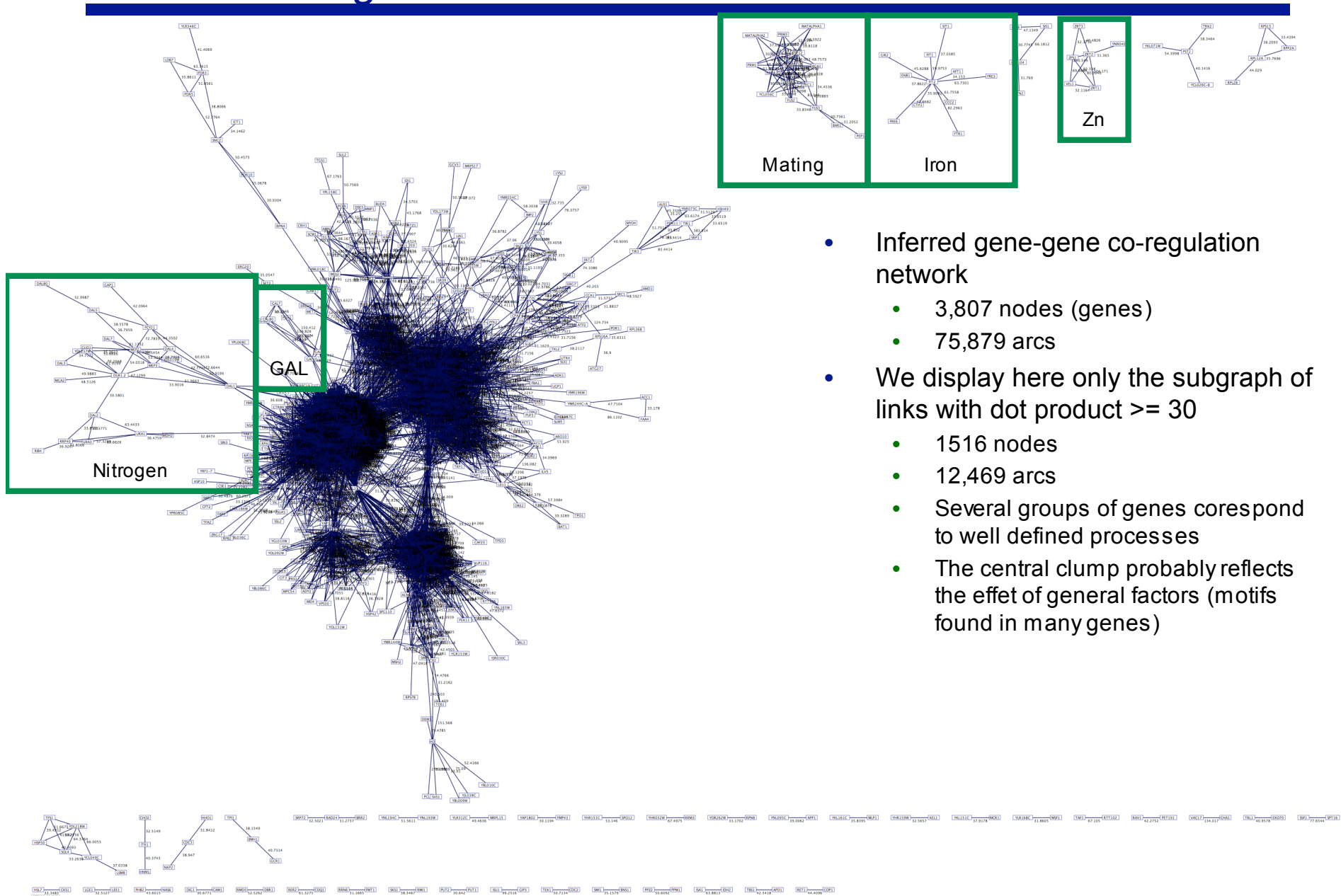
Gene pair	Annot	Score	Sum pred	Sum annot	P.and.A	P.not.A	A.not.P	PPV	Sn
PUTP_PUTA	0	2542.1	1	0	0	1	17076	0.000	0.000
RECA_LEXA	1	1889.8	2	1	1	1	17075	0.500	0.000
GLNL_GLNA	1	1713.1	3	2	2	1	17074	0.667	0.000
UDP_MTLA	1	1043.9	4	3	3	1	17073	0.750	0.000
UVRD_LEXA	1	831.5	5	4	4	1	17072	0.800	0.000
TYRP_AROF	1	815.7	6	5	5	1	17071	0.833	0.000
PUTP_GLNA	1	628.5	7	6	6	1	17070	0.857	0.000
MTLA_MLC	1	448.6	8	7	7	1	17069	0.875	0.000
PSPF_PSPA	0	437.0	9	7	7	2	17069	0.778	0.000
PUTA_GLNA	0	431.5	10	7	7	3	17069	0.700	0.000
ILVY_ILVC	1	386.5	11	8	8	3	17068	0.727	0.001
SULA_RECA	1	358.1	12	9	9	3	17067	0.750	0.001
UVRD_RECA	1	347.5	13	10	10	3	17066	0.769	0.001
UVRA_LEXA	1	346.6	14	11	11	3	17065	0.786	0.001
PUTP_PDHR	1	330.4	15	12	12	3	17064	0.800	0.001
POLA_DNAA	1	296.4	16	13	13	3	17063	0.813	0.001
BIOB_BIOA	1	287.6	17	14	14	3	17062	0.824	0.001
METJ_METB	0	286.7	18	14	14	4	17062	0.778	0.001
MTLA_CYDA	0	281.6	19	14	14	5	17062	0.737	0.001
PUTA_PDHR	0	269.4	20	14	14	6	17062	0.700	0.001
NAGE_MTLA	1	227.7	21	15	15	6	17061	0.714	0.001
UDP_MLC	1	200.0	22	16	16	6	17060	0.727	0.001
MTLA_ADHE	1	197.5	23	17	17	6	17059	0.739	0.001
MALK_MALE	1	197.3	24	18	18	6	17058	0.750	0.001
NRDD_NRDA	0	192.2	25	18	18	7	17058	0.720	0.001
YFID_MTLA	0	183.6	26	18	18	8	17058	0.692	0.001
YFID_CYDA	1	165.4	27	19	19	8	17057	0.704	0.001
SULA_LEXA	1	164.6	28	20	20	8	17056	0.714	0.001
MTLA_CDD	1	162.4	29	21	21	8	17055	0.724	0.001
UVRA_RECA	1	160.5	30	22	22	8	17054	0.733	0.001
SSB_LEXA	1	159.8	31	23	23	8	17053	0.742	0.002
CUER_COPA	0	157.9	32	23	23	9	17053	0.719	0.002
UHPT_MTLA	1	157.2	33	24	24	9	17052	0.727	0.002
MTLA_AGAR	0	155.0	34	24	24	10	17052	0.706	0.002
PUTP_GLNL	0	150.4	35	24	24	11	17052	0.686	0.002
TYRR_AROF	1	147.9	36	25	25	11	17051	0.694	0.002
XSEA_GUAB	1	140.0	37	26	26	11	17050	0.703	0.002
ULAA_MTLA	1	132.9	38	27	27	11	17049	0.711	0.002
MTLA_GLPA	1	130.3	39	28	28	11	17048	0.718	0.002
PSTS_PHOB	1	130.0	40	29	29	11	17047	0.725	0.002
GLNK_GLNA	1	125.8	41	30	30	11	17046	0.732	0.002
MALP_MALE	1	124.5	42	31	31	11	17045	0.738	0.002
ARGE_ARGC	1	122.1	43	32	32	11	17044	0.744	0.002
PDHR_GLCC	0	118.7	44	32	32	12	17044	0.727	0.002
TYRR_TYRP	1	114.3	45	33	33	12	17043	0.733	0.002
METN_FPR	0	113.5	46	33	33	13	17043	0.717	0.002
MTLA_MALE	1	112.4	47	34	34	13	17042	0.723	0.002
PDHR_FADB	0	110.5	48	34	34	14	17042	0.708	0.002
TREB_MTLA	1	107.8	49	35	35	14	17041	0.714	0.002

- Results with 536 genes annotated in RegulonDB
 - Among the 537*536/2=~143,916 gene pairs considered here (between 537 genes annotated in regulonDB), 2,790 show a common motif.
 - Among the 50 top-scoring predicted gene pairs, 35 are annotated as co-regulated in RegulonDB.
 - Among the 15 top-scoring predicted which are non-annotated, at least 8 are actually involved in related pathways
 - Some are already known to be co-regulated, but the annotation is not yet entered in RegulonDB.
 - Some others are very likely to be co-regulated.
- Whole genome analysis (4200 genes)
 - Among the ~16M possible gene pairs 54,099 show a common motif

Analysis of regulatory sequences

***Inferring co-regulation network
for the yeast *Saccharomyces cerevisiae****

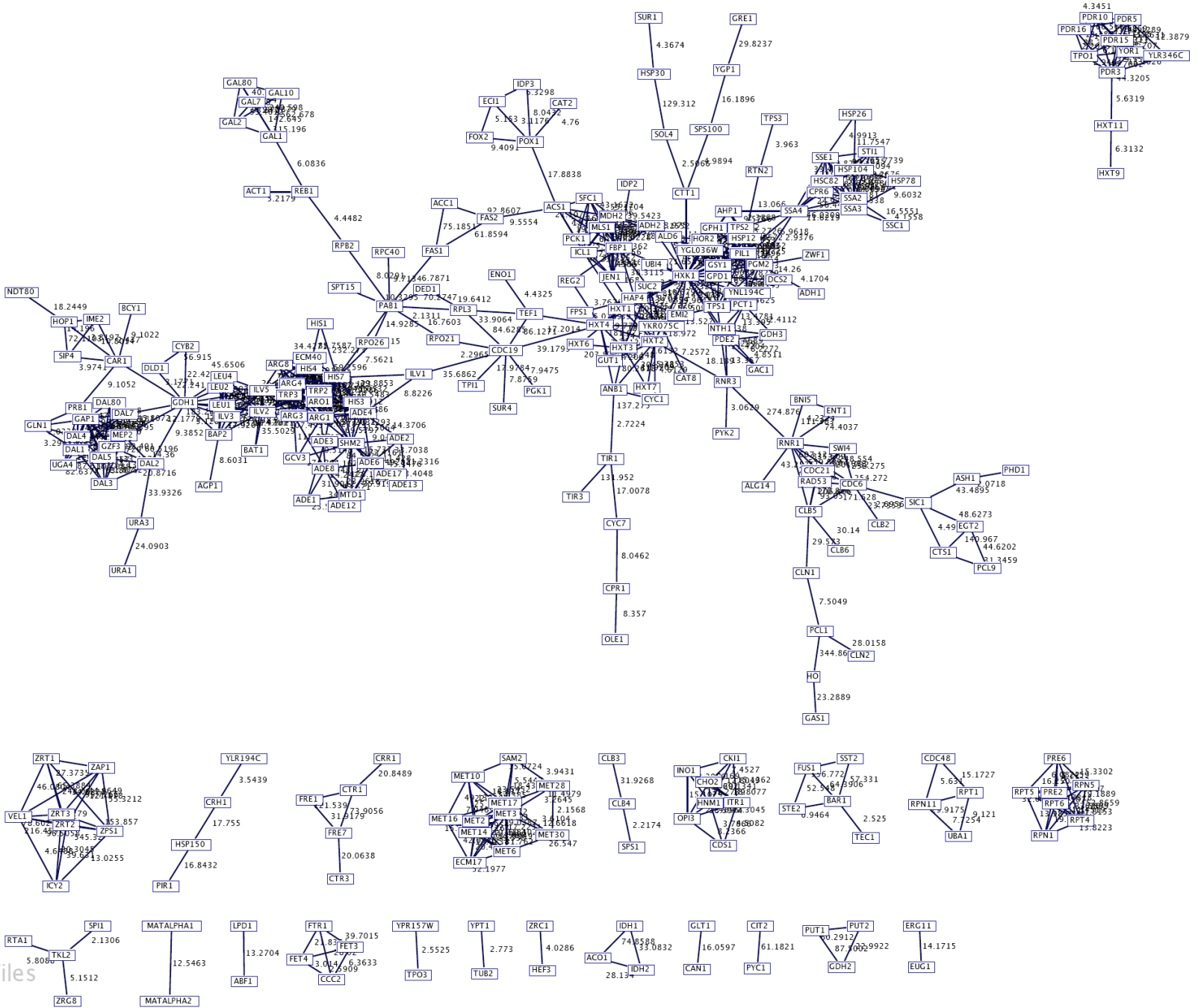
Saccharomyces cerevisiae, *Saccharomycetaceae* Inferred co-regulation network



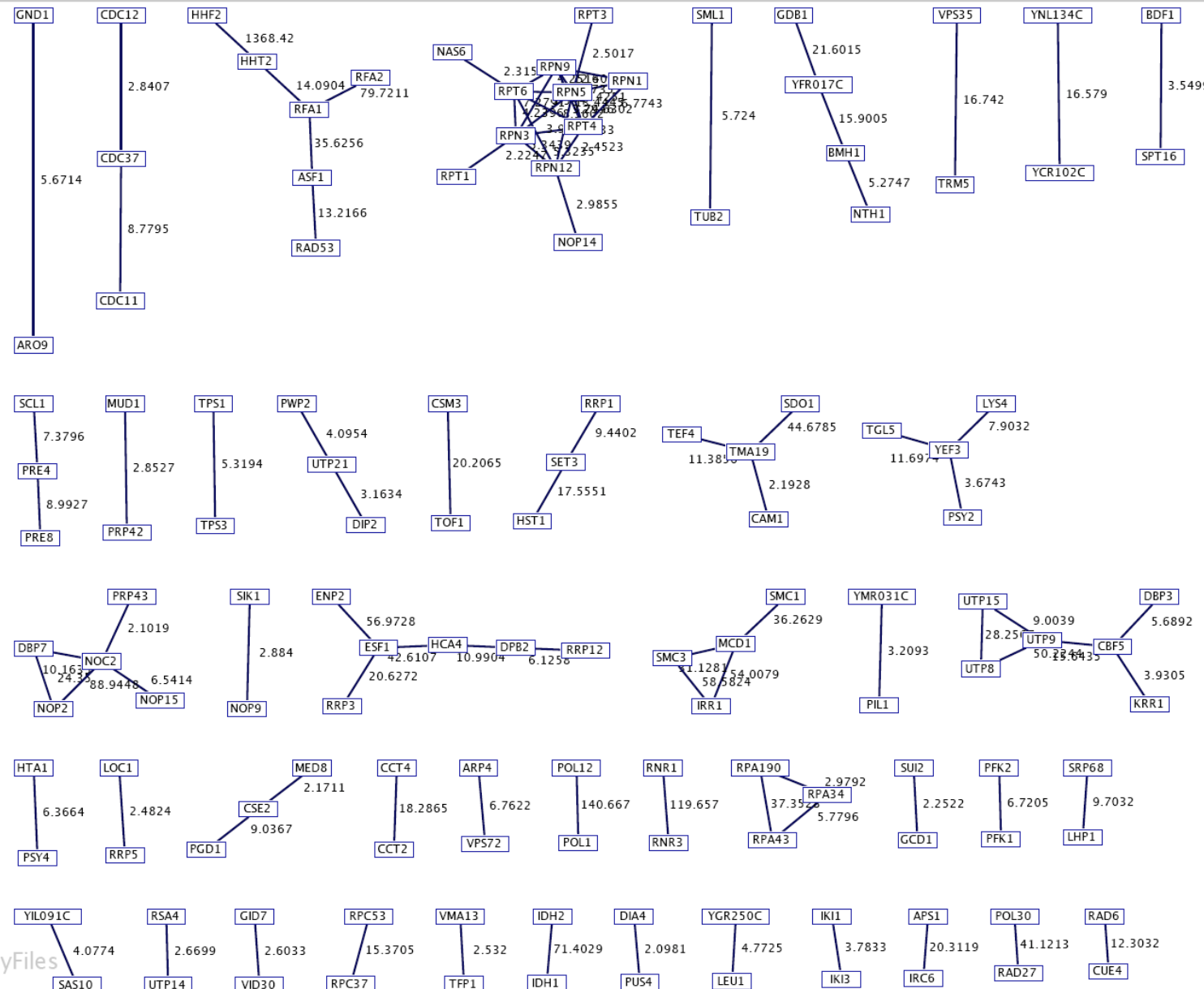
Regulon-wise sensitivity

Regulon	Nodes	Max edges	Predicted edges	Extra edges	Sn	Rand ER Sn	Rand ER extra edges	Rand ND Sn	Rand ND extra edges
MSN4	58	1653	89	1983	0.05	0.01	2128.53	0.01	1606.20
MSN2	56	1540	82	1877	0.05	0.01	2011.60	0.01	1571.57
ZAP1	54	1431	25	785	0.02	0.00	833.20	0.00	1154.13
TEC1	42	861	6	726	0.01	0.00	737.00	0.00	836.43
GCN4	40	780	101	1139	0.13	0.01	1343.60	0.01	1056.67
ABF1	37	666	12	1399	0.02	0.01	1415.60	0.01	976.53
RAP1	32	496	9	1008	0.02	0.01	1032.73	0.00	767.97
YAP1	32	496	8	423	0.02	0.00	438.07	0.00	524.50
GLN3	31	465	69	852	0.15	0.01	983.13	0.01	915.67
MIG1	26	325	37	1377	0.11	0.03	1438.47	0.01	673.77
UME6	26	325	20	519	0.06	0.00	596.40	0.01	703.60
RLM1	25	300	6	339	0.02	0.00	354.20	0.01	725.50
OAF1	24	276	10	218	0.04	0.00	239.53	0.01	637.03
HSF1	21	210	29	443	0.14	0.00	500.00	0.00	492.27
PHO2	21	210	34	506	0.16	0.00	576.13	0.01	629.03
DAL80	19	171	39	428	0.23	0.01	504.13	0.01	675.60
REB1	19	171	8	575	0.05	0.01	602.13	0.01	596.70
INO2	19	171	22	131	0.13	0.00	174.67	0.01	531.30
INO4	19	171	22	131	0.13	0.00	174.67	0.01	531.30
PIP2	19	171	10	152	0.06	0.00	173.80	0.01	458.63
GCR1	18	153	6	808	0.04	0.02	815.80	0.00	415.60
BAS1	17	136	34	425	0.25	0.00	495.73	0.01	559.40
PDR1	16	120	32	367	0.27	0.01	429.60	0.01	457.73
CBF1	16	120	23	432	0.19	0.01	477.67	0.01	499.97
MOT3	15	105	15	975	0.14	0.03	1043.13	0.02	535.20
MIG2	15	105	10	738	0.10	0.02	758.07	0.01	428.67
ROX1	15	105	5	768	0.05	0.01	819.87	0.01	462.67
HAP3	15	105	3	348	0.03	0.00	353.33	0.01	457.60
MCM1	14	91	3	480	0.03	0.01	487.67	0.01	393.80
HAP4	14	91	2	287	0.02	0.00	290.33	0.00	390.67
HAP2	14	91	3	328	0.03	0.00	333.53	0.01	423.57
STE12	13	78	4	131	0.05	0.00	139.93	0.00	282.90
RTG1	12	66	5	422	0.08	0.01	433.80	0.01	425.60
NDT80	11	55	2	610	0.04	0.02	613.53	0.01	352.37
RPN4	11	55	20	270	0.36	0.01	308.93	0.01	316.83
GCR2	11	55	3	367	0.05	0.01	372.07	0.00	281.03
ADR1	11	55	4	143	0.07	0.00	151.87	0.01	351.57
SKO1	11	55	0	89	0.00	0.00	90.00	0.00	143.07
SWI6	10	45	20	890	0.44	0.08	926.07	0.01	352.20
CAT8	10	45	31	448	0.69	0.03	510.73	0.01	353.07
MET4	10	45	27	441	0.60	0.02	496.40	0.00	354.53
HAA1	10	45	2	460	0.04	0.01	463.13	0.01	353.57
DAL81	10	45	11	349	0.24	0.01	371.40	0.01	359.80
PDR3	10	45	16	189	0.36	0.00	220.60	0.01	317.00
GAL4	9	36	10	246	0.28	0.01	265.60	0.01	283.90
GAT1	9	36	7	181	0.19	0.00	194.73	0.01	286.57
UPC2	9	36	3	101	0.08	0.00	150.80	0.01	213.40
RCS1	9	36	6	203	0.17	0.00	214.87	0.01	244.10
MAC1	9	36	5	65	0.14	0.00	75.00	0.01	281.13
HAP5	9	36	0	57	0.00	0.00	57.00	0.00	211.90
THI2	9	36	0	20	0.00	0.00	20.00	0.00	141.10
LEU3	8	28	28	459	1.00	0.03	516.13	0.01	283.53
VID30	8	28	6	406	0.21	0.02	416.87	0.00	286.17
HOG1	8	28	3	296	0.11	0.01	301.40	0.00	211.17
SWI4	8	28	7	333	0.25	0.01	346.47	0.01	281.90
SWI5	8	28	8	312	0.29	0.01	331.67	0.01	286.07
PHO4	8	28	1	148	0.04	0.00	150.73	0.00	175.83
TUP1	7	21	4	732	0.19	0.08	739.73	0.00	211.30
HAC1	7	21	1	188	0.05	0.01	191.73	0.00	174.50
HAP1	7	21	0	237	0.00	0.00	237.80	0.01	248.30
IME1	7	21	5	192	0.24	0.00	237.80	0.00	215.50
RIM101	7	21	0	47	0.00	0.00	46.93	0.01	178.17
MBP1	6	15	10	589	0.67	0.10	609.07	0.00	176.23
CYC8	6	15	7	526	0.47	0.06	539.33	0.01	175.70
SNF2	6	15	4	406	0.27	0.04	413.87	0.01	207.40
TYE7	6	15	1	246	0.07	0.01	247.67	0.01	173.00
SKN7	6	15	0	143	0.00	0.01	142.80	0.00	140.43
DAL82	6	15	7	186	0.47	0.00	199.87	0.00	215.60
RTG3	6	15	3	139	0.20	0.00	146.93	0.02	213.30
GRR1	5	10	8	459	0.80	0.04	474.13	0.01	175.77
RTG1	5	10	8	459	0.80	0.04	474.13	0.01	175.77
RFX1	5	10	1	391	0.10	0.03	394.33	0.00	139.50
SWI1	5	10	4	231	0.40	0.02	239.67	0.01	172.63
MET31	5	10	5	166	0.50	0.01	175.80	0.00	177.63
XBP1	5	10	0	165	0.00	0.01	164.87	0.01	181.80
HMRA1	5	10	0	91	0.00	0.01	90.87	0.00	103.13
SOK2	5	10	0	110	0.00	0.00	112.93	0.03	180.17
NRG1	5	10	1	102	0.10	0.00	105.93	0.02	178.00
RFA2	5	10	2	138	0.20	0.00	142.93	0.00	139.80
ACE2	5	10	5	220	0.50	0.00	233.00	0.02	181.47

Inferred co-regulation versus annotated regulons

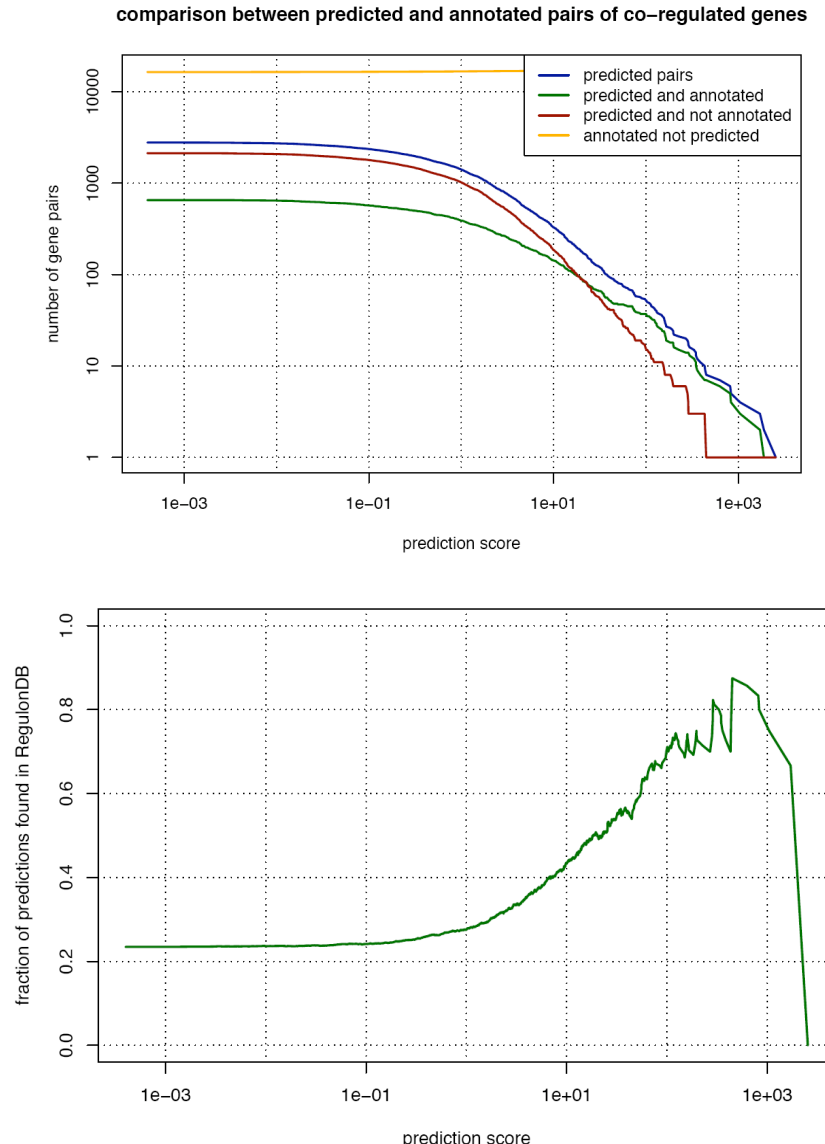


Inferred co-regulation versus protein interactions (Krogan 2006)



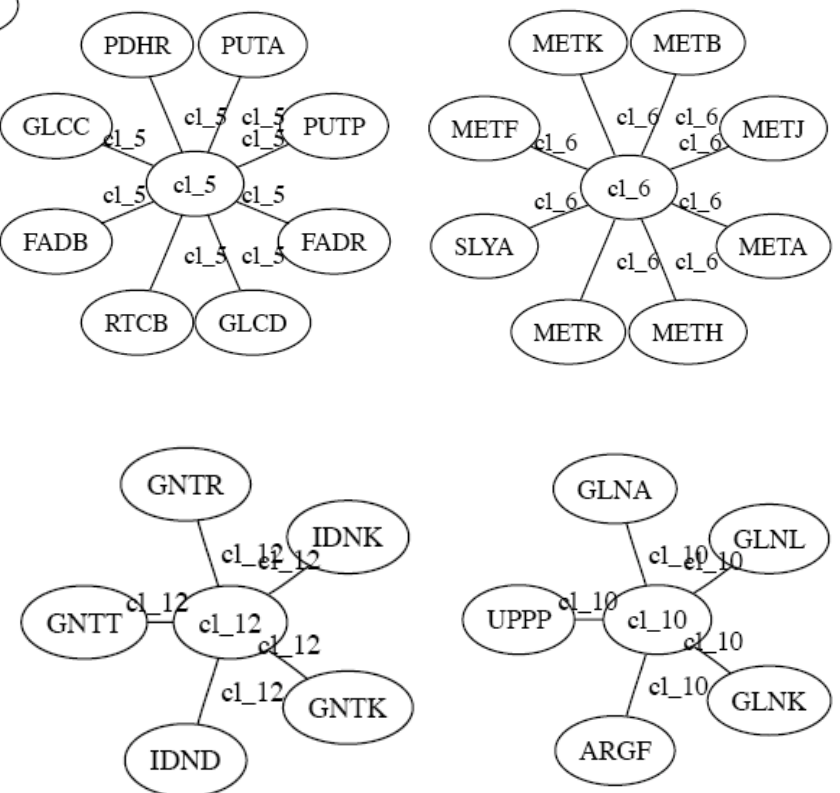
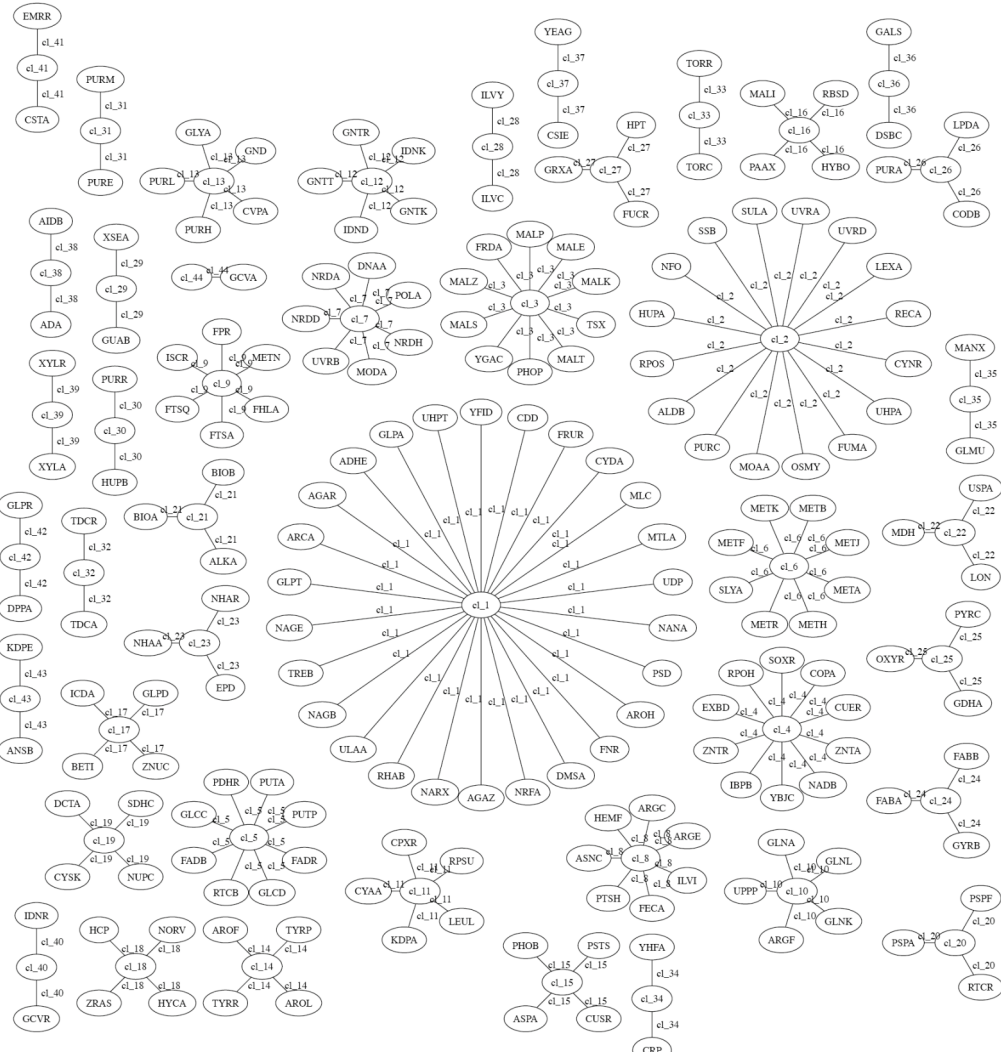
Supplementary material

Predicted versus annotated gene pairs



- We performed a systematic estimation of the fraction of predictions found in RegulonDB, as a function of the prediction score (dot product).
- More than 40% of the predictions with a score $dp \geq 10$ are annotated in RegulonDB.
- Note that predicted pairs which are not found in RegulonDB might nevertheless be correct, it is hard to estimate the rate of "false positive" in such a case.
- The sensitivity can however be estimated, and it is quite low.
 - Among the 14,939 gene-gene co-regulated pairs in RegulonDB, the method detects no more than 652 (4.4%).
 - Note that more than 50% of the annotated gene pairs are linked via CRP.
- To do : analyze the sensitivity per TF, to check whether some regulons are preferentially detected.

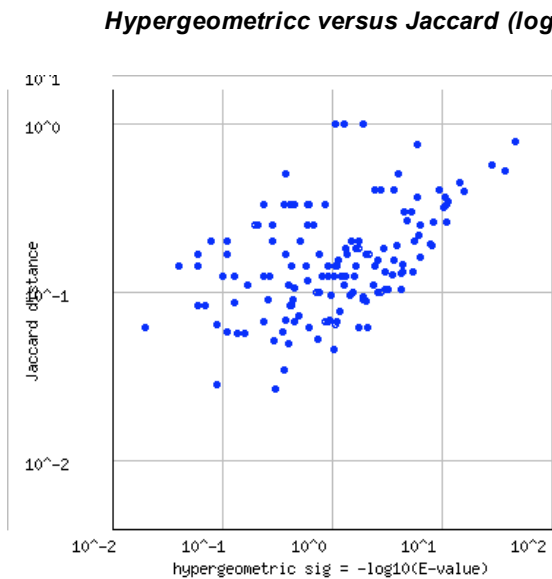
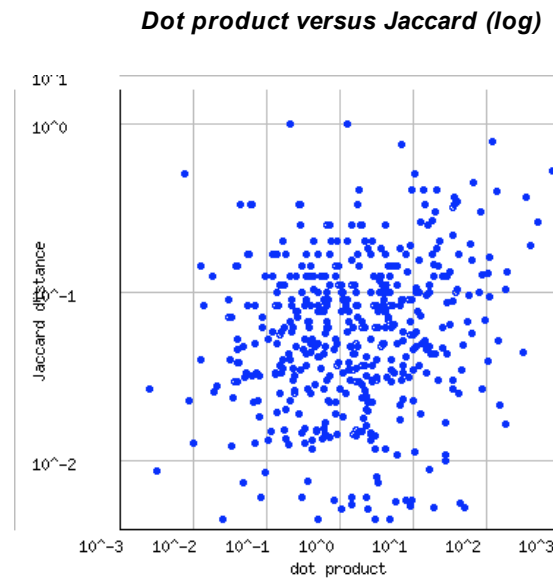
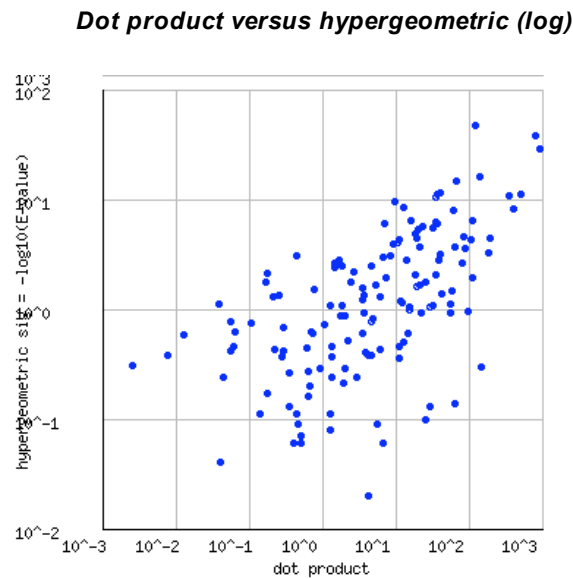
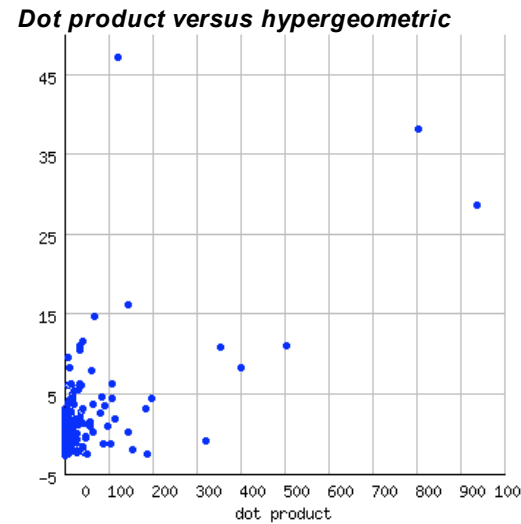
Clustering



Clusters versus regulons

cluster	12	11	10	6	4	2	8	9	7	3	5	1	clust
CRP	22	4	10	1	6					4			47
FNR	4	1	1	7	2		3						18
IHF	5	2	1		1		2	2					13
ARCA	5	1	1	2									9
FIS	4			1			1			1		1	8
LEXA		1							1		6		8
NARL	4				1			3					8
PURR	7												7
FLHD	3				1		1		1				6
LRP	3	1								1			5
MALT			5										5
METJ								5					5
MODE	2			1			2						5
DNAA	1								2			1	4
MLC	2	1			1								4
NARP	3					1							4
TYRR						4							4
CPXR	2								1				3
CYTR	1		1							1			3
FRUR	1			1						1			3
GADX	2	1											3
GLPR	1			2									3
GNTR	3												3
MARA		1		1				1					3
NAGC	1			1						1			3
NTRC	3						3						3
PHOB						3			1				3
ARGP								3			1		2
BIRA							2			1			2
FADR	2												2
FHLA	2												2
GADE	1								1				2
GLCC		2											2
H-NS	1			1									2
IDNR	2												2
ILVY	2												2
MALI	2												2
METR	1							1					2
NAC	2												2

Compare-classes result (selection)



Discovered motifs in promoters of glpD orthologs from Enterobacteriales

Dyad	exp_freq	occ	exp_occ	occ_P	occ_E	sig	rank
atgn{1}tcg cgan{1}cat	0.00069	11	1.02	1.30E-08	2.40E-04	3.62	1
aacn{0}att aatn{0}gtt	0.00131	13	1.95	1.60E-07	3.00E-03	2.52	2
cgan{0}aca tgtn{0}tcg	0.00056	9	0.84	2.90E-07	5.20E-03	2.28	3
agtn{10}tcg cgan{10}act	0.00026	6	0.35	2.20E-06	3.90E-02	1.41	4
acan{13}acg cgtn{13}tgt	0.00044	7	0.6	3.50E-06	6.40E-02	1.19	5
aacn{2}aca tgtn{2}gtt	0.00082	9	1.21	5.60E-06	1.00E-01	0.99	6
aatn{2}tcg cgan{2}att	0.00089	9	1.31	1.00E-05	1.90E-01	0.73	7
atgn{0}ttc gaan{0}cat	0.00094	9	1.4	1.70E-05	3.10E-01	0.52	8
acgn{2}cat atgn{2}cgt	0.00054	7	0.79	2.10E-05	3.80E-01	0.42	9
aacn{3}cat atgn{3}gtt	0.00101	9	1.47	2.60E-05	4.60E-01	0.33	10
aacn{8}aag cttn{8}gtt	0.00080	8	1.11	2.60E-05	4.70E-01	0.33	11
cctn{4}cac gtgn{4}agg	0.00023	5	0.34	2.90E-05	5.30E-01	0.28	12
aacn{0}gaa ttcn{0}gtt	0.00106	9	1.58	4.40E-05	7.90E-01	0.1	13
actn{7}aag cttn{7}agt	0.00027	5	0.37	4.90E-05	8.90E-01	0.05	14

;assembly # 1	seed: atgntcg	13 words
aaangtt....aacnntt	0.82
.aatgtt....aacatt.	3.38
.aatnntcg..	..cgannatt.	2.1
..atgntcg..	..cgancat..	7.55
..atgttc...	...gaacat..	3.29
..atgnnnngt	aacnnncat..	1.63
..atgnncgt.	.acgnncat..	1.46
...tgttcg..	..cgaaca...	4.58
...tgtnnngt	aacnnaca...	2.4
...tgtncgt.	.acgnaca...	1.32
....gttnngt	aacnaac....	1.05
....gttcgt.	.acgaac....	0.47
.....ttcgtt	aacgaa.....	1.53
aaatgttcgtt	aacgaacattt	7.55
;assembly # 2	seed: atgnnnnnnnntcg	19 words
aaangtt.....aacnntt	0.82
.aatgtt.....aacatt.	3.38
.aatnntcg.....cgannatt.	2.1
..atgntcg.....cgancat..	7.55
..atgttc.....gaacat..	3.29
..atgnnnnnnnntcg.	.cgannnnnnncat..	1.82
..atgnnnngt....aacnnncat..	1.63
..atgnncgt....acgnncat..	1.46
..atgnnnnnnnnncga	tcgnnnnnnnnncat..	0.93
...tgttcg.....cgaaca...	4.58
...tgtnnngt....aacnnaca...	2.4
...tgtncgt....acgnaca...	1.32
...tgtnnnnnnntcg.	.cgannnnnnnaca...	0.74
....gttnngt....aacnaac....	1.05
....gttcgt....acgaac....	0.47
....gttnnnnnntcg.	.cgannnnnaac....	0.35
.....ttcgtt....aacgaa.....	1.53
.....tcgnnntcg.	.cgannncga.....	0.6
.....tcgnnnnncga	tcgnnnnncga.....	0.06
aaatgttcgttntcg	tcganaacgaacattt	7.55
;assembly # 3	seed: agtnnnnnnnnnntcg	3 words
atgnnnnnnnnnntcg	cgannnnnnnnnnact	1.52
.gtttnnnnnnnntcg	cgannnnnnnnnnnaac.	0.9
..ttcnnnnnnnntcg	cgannnnnnnnnngaa..	0.78
agttcnnnnnnnntcg	cgannnnnnnnnngaact	1.52
;assembly # 4	seed: acannnnnnnnnnnnnacg	3 words
acannnnnnnnnnnnnacg.	.cgtnnnnnnnnnnnntgt	1.26
acannnnnnnnnnnnncga	tcgtnnnnnnnnnnnntgt	0.48
acannnnnnnnnnnnaac..	..gttnnnnnnnnnntgt	0.01
acannnnnnnnnnnnaacga	tcgttnnnnnnnnnntgt	1.26

Selected inferred clusters

id	name	description [synonyms]
YGL255W	ZRT1	High-affinity zinc transporter of the plasma membrane, responsible for the majority of zinc uptake; transcription is induced under low-zinc conditions by the Zap1p transcription factor [YGL255W;ZRT1;ZRT1;YGL255W;852637;6321182;NP_011259]
YKL175W	ZRT3	Vacuolar membrane zinc transporter, transports zinc from storage in the vacuole to the cytoplasm when needed; transcription is induced under conditions of zinc deficiency [YKL175W;ZRT3;ZRT3;YKL175W;853679;6322673;NP_012746]
YLR130C	ZRT2	Low-affinity zinc transporter of the plasma membrane; transcription is induced under low-zinc conditions by the Zap1p transcription factor [YLR130C;ZRT2;ZRT2;YLR130C;850821;6323159;NP_013231]
YOL154W	ZPS1	Putative GPI-anchored protein; transcription is induced under low-zinc conditions, as mediated by the Zap1p transcription factor, and at alkaline pH [YOL154W;ZPS1;ZPS1;YOL154W;854011;6324419;NP_014488]
YGL258W	VEL1	Protein of unknown function; highly induced in zinc-depleted conditions and has increased expression in NAP1 deletion mutants [YGL258W;VEL1;VEL1;YGL258W;852634;6321179;NP_011256]
YNR040W	YNR040W	Putative protein of unknown function; the authentic, non-tagged protein is detected in highly purified mitochondria in high-throughput studies [YNR040W;YNR040W;855776;6324368;NP_014438]

Phylogenetic footprints MET genes in Gammaproteobacteria

