

Details of strains used to compare RBI algorithm efficiency against others

Strain	The number of				Reference
	Metabolites	Reactions	Genes	TFs	
E. coli core	72	95	137	18	Orth et al. (2010)
iAF1260	1668	2382	1261	49	Feist et al. (2007)
iJO1366	1805	2583	1367	49	Orth et al. (2011)
iMM904	1226	1577	905	99	Mo et al. (2009)
iTO977	1218	1562	961	98	Osterlund et al. (2013)
Yeast7	2220	3498	910	99	Aung et al. (2013)

References

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