

**Table 3.** (R/G)<sub>median</sub> of 86 genes over 11 experiments

UNIQID	Name	a=glnLup(Cy5) vs glnLup nac(Cy3) [NH3], 2nd PCR =>Nac control	b=glnLup(Cy5) vs glnLup nac(Cy3) [Gln], 2nd PCR =>Nac control	c=glnLup(Cy5) vs glnG(Cy3) [NH3], 1st PCR =>NtrC&Nac control	d=glnLup(Cy5) vs glnG(Cy3) [NH3], 2nd PCR =>NtrC&Nac control	e=glnLup(Cy5) vs glnG(Cy3) [Gln], 1st PCR =>NtrC&Nac control	f=glnLup(Cy5) vs glnG(Cy3) [Gln], 2nd PCR =>NtrC&Nac control	g=glnLup nac(Cy5) vs glnG(Cy3) [NH3], 1st PCR =>NtrC control	h=glnLup nac(Cy5) vs glnG(Cy3) [Gln], 2nd PCR =>NtrC control	i=Gln(Cy5) vs NH3(Cy3) [glnG], 1st PCR =>other control	j=NH3(Cy5) vs Gln(Cy3) [glc], 1st PCR	k=NH3(Cy5) vs Gln(Cy3) [gly], 1st PCR
b0336	b0336 codB cytosine permease/transport	1.8	6.3	3.1	2.1	11.9	6	0.9	1.1	0.3	0.2	0.1
b0337	b0337 codA cytosine deaminase	1.8	3.4	2.8	2.4	7.3	3.7	0.7	1	0.4	0.3	0.1
b0450	b0450 glnK nitrogen regulatory protein P-II 2	0.9	0.7	10.3								
b0451	b0451 amtB probable ammonium transporter	0.9	1	6.2	0.9	7.9	0.9	1.3	1	1.2	0.1	0.3
b0652	b0652 gltL ATP-binding protein of glutamate/aspartate transport system	1.1	0.8	1.5	1.2	1.7	1.3	1.5	1.4	1.1	0.8	0.9
b0653	b0653 gltK glutamate/aspartate transport system permease	1	0.9	1.6	1.1	2	1.3	1.1	1.6	0.8	1	1.1
b0654	b0654 gltJ glutamate/aspartate transport system permease	1.1	1.1	1.7	1.1	2.2		1.4	1.2	0.6	0.8	0.7
b0655	b0655 ybeJ putative periplasmic binding transport protein	1.3	1.4	3.4	0.8	6.5	1.1	1.6	1	0.9	0.8	0.7
b0809	b0809 glnQ ATP-binding component of glutamine high-affinity transport system	1.1	0.4	2.7	1.9	3.6	2	1.3	4.9	0.8	0.5	0.5
b0810	b0810 glnP glutamine high-affinity transport system; membrane component	1.2	0.6	3.5	2.1	6	2.9	2.3	6.4	1	0.4	0.5
b0811	b0811 glnH periplasmic glutamine-binding protein; permease	1	0.5	4.5	5.8	9.4	4.7	5.9	11.3	1.1	0.3	0.4
b0854	b0854 potF periplasmic putrescine-binding protein; permease protein	1.4	0.9	4.8	3.4	6.3	2.6	2.6	3.7	1.3	0.3	0.3
b0855	b0855 potG ATP-binding component of putrescine transport system	1.3	0.8	1.9	2.1	4	1.6	1.2	2.3	1	0.4	0.4
b0856	b0856 potH putrescine transport protein; permease	1.2	0.9	2.5	1.5	2.6	1.2	1.2	1.9	0.9	0.5	0.5
b0857	b0857 potI putrescine transport protein; permease	1.1	0.9	1.2	1.1	1.7	0.9	0.6	1.3	0.9	0.7	0.7
b0929	b0929 ompF outer membrane protein 1a (la;b;F)	2.6	8.8	4.1	4.9	24.9	28.8	2.1	2.8	0.2	0.5	0.6
b1006	b1006 ycdG putative transport protein	1.1	0.6	5.1	2.6	6.5	2.4	1.5	5.9	1	0.2	0.2
b1007	b1007 orf, hypothetical protein	1	0.6	1.8	2	6.6	3.1	2.3	6.4	1	0.3	0.2
b1008	b1008 putative enzyme	1	0.7	16	2.4	16.9	3	2.4	6.1	1.5	0.3	0.1
b1009	b1009 putative acetyltransferase	1	0.7	7.1	4.4	17.1	3.5	3.3	9.3	1	0.3	0.1
b1010	b1010 orf, hypothetical protein	1	0.6	4	1.9	2.8	2.6	2.4	4.3	0.8	0.6	0.3
b1011	b1011 putative synthetase	1.2	0.6	25	5.5	32.9	3.4	2.7	8.4	1	0.2	0.1
b1012	b1012 orf, hypothetical protein	1.2	0.7	26.3	10.6	51.8	6.7	4.3	15.8	1.1	0.1	0.1
b1034	b1034 ycdX orf, hypothetical protein	1	1.1	3.6	1.1	4.9	1.1	0.9	1.1	1.1	0.7	0.6
b1035	b1035 ycdY putative oxidoreductase component	0.9	0.9	0.7	1.1	0.9	1	1.4	1.1	0.9	1.3	1.5
b1217	b1217 chaB cation transport regulator	1.1	0.9	1	1.2	1.1	0.8	1	0.9	0.9	1	1
b1218	b1218 chaC cation transport regulator	1.2	0.7	3.8	1.7	5.3	1.5	1.1	2.5	1.4	0.2	0.4
b1243	b1243 oppA oligopeptide transport; periplasmic binding protein	1.4	1.7	2	2.8	5.8	3.7	1.4	2.7	0.3	2.4	0.9
b1244	b1244 oppB oligopeptide transport permease protein	1.3	1.6	1.4	1.6	3.3	2.2	0.9	1.4	0.3	1.2	0.9
b1245	b1245 oppC homolog of <i>Salmonella</i> oligopeptide transport permease protein	1.2	1.4	1.3	1.2	3		0.9	1.4	0.4	1.1	1
b1246	b1246 oppD homolog of <i>Salmonella</i> ATP-binding protein of oligopeptide ABC transport system	1.3	1.6	1.6	1.3	3.9	1.7	1.3	1.2	0.4	0.8	0.7
b1296	b1296 ycjY putative amino acid/amine transport protein	1.2	1.1	3	1.2	2.7	0.9	0.8	1.1	1.8	0.5	0.4
b1384	b1384 feaR regulatory protein for 2-phenylethylamine catabolism	1.3	1.3	2.7	1.9	3.6	1.1	0.8	0.9	0.7	0.3	0.9
b1440	b1440 putative transport protein	2.5	6.1	3.8	3	7.4	4.8	0.7	0.9	1	0.4	0.3
b1441	b1441 putative ATP-binding component of a transport system	1.8	2.3	2.1	1.8	3.3	2.3	2.2	1	1.3	0.6	0.5
b1442	b1442 putative transport system permease protein	1.3	2.1	2.6	1.4	3.9	1.8	0.8	1	1.1	0.6	0.6
b1443	b1443 putative transport system permease protein	1.9	2	1.6	1.8	2.6	1.8	0.9	1	0.9	0.7	0.5
b1444	b1444 putative aldehyde dehydrogenase	1.9	2.2	3.8	4.3	5.3	2.6	1.2	1.2	1.2	0.3	0.3
b1483	b1483 putative ATP-binding component of a transport system	0.9	0.9	5.3	2	4.5	1.7	1.9	2.5	1.3	0.6	0.3
b1484	b1484 putative ATP-binding component of a transport system	1.1	0.9	6.9	2.4	6.7	2.9	1.8	3.6	0.9	0.5	0.3
b1485	b1485 putative transport protein	1.1	0.9	4.9	2.1	7.2	2.1	1.5	2.9	1	0.5	0.5
b1486	b1486 putative transport system permease protein	1.1	0.9	1	1.9	1.2	2.2	2.5	3.2	0.8	1.1	0.8
b1487	b1487 putative hemin-binding lipoprotein	1	0.6	50.4	18.3	65.3	7.2	4.9	15.2	0.9	0	0.1
b1488	b1488 orf, hypothetical protein	0.8	0.8	9.7	4.7	14.6	3.1	4.1	4.9	0.8	0.1	0.1
b1744	b1744 ydiS orf, hypothetical protein	0.9	0.8	6.9	1.6	6.3	1.3	1.4	1.9	1.2	0.3	0.3
b1745	b1745 orf, hypothetical protein	0.9	0.7	4	1.8	5.3	1.7	1.5	2.6	1.3	0.4	0.4
b1746	b1746 putative aldehyde dehydrogenase	0.9	0.7	11.5	2	7.3	1.6	1.4	2.4	1.4	0.3	0.2
b1747	b1747 orf, hypothetical protein	0.8	0.8	2.3	2.9	8.2	1.8	1.8	3.6	0.8	0.3	0.2

UNIQID	Name		a=glnLup (Cy5)	b=glnLup (Cy5)	c=glnLup (Cy5)	d=glnLup (Cy5)	e=glnLup (Cy5)	f=glnLup (Cy5)	g=glnLup nac(Cy5) vs glnG(Cy3) [NH3]	h=glnLup nac(Cy5) vs glnG(Cy3) [Gln]	i=Gln(Cy5) vs NH3(Cy3)	j=NH3(Cy5) vs Gln(Cy3) [glc]	k=NH3(Cy5) vs Gln(Cy3) [gly]		
b1748	b1748	cstC	acetylornithine delta-aminotransferase	0.8	0.6	7.7	3.6	10.9	3.4	2.2	7.9	0.7	0.2	0.2	
b1783	b1783	yeaG	orf, hypothetical protein	1.4	0.5	4.5	4.4	2.9	1.1	1.7	2.4	3	0.6	0.3	
b1784	b1784	yeaH	orf, hypothetical protein	1.2	0.6	4.1	3.1	2.5	0.9	1.2	1.7	2.4	0.4	0.3	
b1932	b1932	yedL	orf, hypothetical protein	3	3.6	5.2	7.5	4.6	3.3	1.2	0.8	0.8	0.1	0.2	
b1987	b1987	cbl	transcriptional regulator cys regulon; accessory regulatory circuit affecting cysM	3.1	2.3	29.6	9.9	19.3	2.4	0.7	1	0.7	0.1	0.1	
b1988	b1988	nac	nitrogen assimilation control protein	15.5	25	48.2	25.4	56.2	22.7	0.7	0.8	1	0	0	
b2306	b2306	hisP	ATP-binding component of histidine transport	1	0.6	3.2	1.8	8.4	2	1.7	4.3	0.7	0.7	0.5	
b2307	b2307	hisM	histidine transport, membrane protein M	1	0.9	1.8	1.6	5.3	1.8	1.5	2.6	0.6	0.5	0.5	
b2308	b2308	hisQ	histidine transport system permease protein	0.9	0.8	2.1	2.2	3.1	1.4	1.3	1.9	0.6	0.6	0.4	
b2309	b2309	hisJ	histidine-binding periplasmic protein of high-affinity histidine transport system	0.8	0.6	4.1	2.5	7.4	5.2	2.7	6.7	0.5	0.4	0.4	
b2310	b2310	argT	lysine-, arginine-, ornithine-binding periplasmic protein	1.1	0.6	10.5	5.5	18.1	12	6.9	26.3	0.5	0.2	0.3	
b2393	b2393	nupC	permease of transport system for 3 nucleosides	2	2.9	3.9	3.5	5.3	2.6	0.8	1.2	0.6	0.8	0.3	
b2661	b2661	gabD	succinate-semialdehyde dehydrogenase, NADP-dependent activity	1.7	1.5	2.6	2	1.9	1.4	1	0.8	1.4	0.7	0.8	
b2662	b2662	gabT	4-aminobutyrate aminotransferase activity	1.3	1.2	4.3	1.5	2.6	1.1	0.9	0.8	1.4	0.3	0.5	
b2663	b2663	gabP	transport permease protein of gamma-aminobutyrate	2	2	5.2	2.7	2.9	1.4	0.8	0.8	1.1	0.4	0.5	
b2664	b2664	ygaE	putative transcriptional regulator	1.2	1	2.2	1.6	1.2	0.8	1	0.8	1.7	0.4	0.6	
b2875	b2875	b2875	putative synthases	1.1	0.5	4.8	1.7	7.4	1.8	1.2	4.9	1.1	0.4	0.3	
b2876	b2876	orf	hypothetical protein	1	0.9	3.2		3.1	1.2	1	2	0.5	0.6	0.8	
b2882	b2882	ygfO	putative transport protein	1	0.9	0.9	1	0.7	0.9	0.9	1	1.4	0.8	1.2	
b2883	b2883	ygfP	orf, hypothetical protein	1.2	1.3	5.1	2.3	4.2	1.4	1.3	1.1	1.2	0.4	0.3	
b2884	b2884	ygfQ	orf, hypothetical protein	1.2	1.2	1.4	1	1.3	1	0.9	1	1	0.8	1.1	
b2885	b2885	ygfR	putative oxidoreductase	1	1.1	1.7	1	1.2	1	1	1	1.1	1	0.7	0.8
b3073	b3073	ygjG	probable ornithine aminotransferase	1.4	0.8	4.8	4.9	3.6	1.8	1.6	1.7	2.2	0.2	0.2	
b3268	b3268	yhdW	putative periplasmic binding transport protein	1.1	0.8	1	9.2	1.4	4.4	2.9	6.7	1	1.5	1	
b3269	b3269	yhdX	putative transport system permease protein	1.1	1	3	1.5	1.7	1.1	0.8	1.3	1.1	0.6	0.7	
b3270	b3270	yhdY	putative transport system permease protein	1.1	0.8	1	1.1	1.2	0.9	0.9	1.2	1.7	0.8	0.8	
b3271	b3271	yhdZ	putative ATP-binding component of a transport system	0.9	0.9	4.4	1.7	4.2	1.5	1.8	1.7	1.3	0.4	0.5	
b3512	b3512	yhiE	orf, hypothetical protein	3.1		3.1	3.2	1		0.7	0.4	8.6	0.1	0	
b3540	b3540	dppF	putative ATP-binding component of dipeptide transport system	1.8	2	3.3	2.4	5	2.8	1.1	1.4	0.5	0.6	0.3	
b3541	b3541	dppD	putative ATP-binding component of dipeptide transport system	1.5	2.2	3.1	1.8	3.9	2.5	0.8	1.3	0.4	0.7	0.5	
b3542	b3542	dppC	dipeptide transport system permease protein 2	1.6	1.9	2.7	2.1	2.6	2.3	1	1.2	0.5	0.7	0.5	
b3543	b3543	dppB	dipeptide transport system permease protein 1	1.4	2.6	3.5	1.9	4.4	2.4	0.6	1.2	0.3	0.8	0.7	
b3544	b3544	dppA	dipeptide transport protein	3.3	5.9	5.5	8.4	10.4	9.6	1.7	2.5	0.5	1.3	0.5	
b3868	b3868	glnG	response regulator for gln (sensor glnL) (nitrogen regulator I, NR1)	0.9	0.7	2.5	1.7	1.2	1.3	1.1	1.7	1.7	0.5	0.3	
b3869	b3869	glnL	histidine protein kinase sensor for GlnG regulator (nitrogen regulator II, NR2)	0.9	0.9	3.4	2.6	1.4	1.6	2.7	2.1	2	0.5	0.2	
b3870	b3870	glnA	glutamine synthetase	1.2	1.1	11.6	9.3	24.4	13.1	5.2	16.7	0.5	0.5	0.2	
b4207	b4207	fklB	FKBP-type 22KD peptidyl-prolyl cis-trans isomerase (rotamase)	1.3	1.7	1.6	2.1	3.5	1.8	0.9	1.2	0.7	0.6	0.5	
b4208	b4208	cycA	transport of D-alanine, D-serine, and glycine	2.5	3.1	5.2	2.8	7.9	3	1	1	0.8	0.2	0.3	

Operons were selected by the following: For those genes that met the criteria for Table 2, selected were those genes that were >2.5-fold higher in *gln* (Up) compared to *gln* when both were grown on glutamine. See *Materials and Methods* for details.