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# Exploring the *Mycoplasma capricolum* genome: a minimal cell reveals its physiology

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#### Summary

We report on the analysis of 214kb of the parasitic eubacterium Mycoplasma capricolum sequenced by genomic walking techniques. The 287 putative proteins detected to date represent about half of the estimated total number of 500 predicted for this organism. A large fraction of these (75%) can be assigned a likely function as a result of similarity searches. Several important features of the functional organization of this small genome are already apparent. Among these are (i) the expected relatively large number of enzymes involved in metabolic transport and activation, for efficient use of host cell nutrients; (ii) the presence of anabolic enzymes; (iii) the unexpected diversity of enzymes involved in DNA replication and repair; and (iv) a sizeable number of orthologues (82 so far) in Escherichia coli. This survey is beginning to provide a detailed view of how M. capricolum manages to maintain essential cellular processes with a genome much smaller than that of its bacterial relatives.

#### Introduction

Mycoplasmas are a diverse group of Gram-positive bacteria with a characteristically low GC content (Herrmann, 1992) that lack a cell wall. Their genomes vary between about 600 kb and 1800 kb (Wenzel *et al.*, 1992) and are the smallest genomes of living cells (Razin, 1992). As

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deduced from sequence comparison of 16S rRNA, Bacillus subtilis is a well-characterized close relative (Woese, 1987; Weisburg et al., 1989) but has a genome size of about 4.2 Mb, about four or five times larger than that of the mycoplasmas. The main driving force for the condensation of its genome appears to be the parasitic life of mycoplasmas; the use of metabolic products of the host appears to allow a drastic reduction in the number or components of biosynthetic pathways. Although considerable amounts of biochemical data provide an insight into this highly specialized metabolism (for a comprehensive review see Maniloff et al., 1992), direct analysis of the genome is a powerful approach to complement existing knowledge. Large-scale genomic sequencing of Mycoplasma species has already successfully begun for Mycoplasma genitalium (Peterson et al., 1993; Nowak, 1995), Mycoplasma pneumoniae (R. Herrmann, personal communication) and Mycoplasma capricolum (Gillevet, 1993). Here we report the initial analysis of 214 kb from M. capricolum. Although this is a preliminary data set consisting of multiple short contigs, the analysis has already yielded many insights into the physiology of this small parasitic bacterium.

#### **Results and Discussion**

Data and data quality

One-pass fluorescent sequencing was used to produce 1505 random clones as starting points for genomic walking techniques (Gillevet, 1993). At the current stage, more than 870 477 bases have been sequenced from M. capricolum. (California kid) strain ATCC 27343 and finally assembled into 372 non-overlapping contigs covering 214528 bp. The length of the contigs varies from 63 to 2049 bases with an average length of 658. The data consist of 13091 bases (6%) from one-pass fluorescent sequencing that cover 1198 of the 1505 random clones and 201 437 bases (94%) produced by genomic walking. Of the latter, 45 632 bases (23%) are single-stranded regions at the end of the growing contig, 155 805 bases (77%) have multiple coverage on at least one strand while 112621 bases (56%) are covered on both strands. There is a total of 968 ambiguous calls (nucleotide assignments) in the data set (~0.5%).

We have compared 8868 bases of our data with  $\it M.$  capricolum sequences stored in public databases and

note less than 0.7% difference between the two data sets (these include ambiguous calls, insertions, deletions and mismatches). It should be noted that the strand coverage of this latter data set is essentially identical to that of the entire data set. Many of the discrepancies noted between the published data and our data set are in regions where we have multiple coverage on both strands and, therefore, they may be attributed to strain differences. As a result, the true error rate of the walking data may be much lower than 0.7%. Furthermore, we have identified only 97 frameshifts and aberrant stop codons in the 103 000 bases contained in the open reading frames (ORFs) that we have analysed, which indicates that the error rate of this type is probably less than 10<sup>-3</sup> (details of the sequencing methodology will be published elsewhere; P. M. Gillevet et al., in preparation).

#### Identification of proteins by similarity searches

Similarity searches are insensitive to error rates at least up to 1% (States and Botstein, 1992) and can even be used for detection of sequencing errors (Posfai and Roberts, 1992). Therefore, a preliminary data set consisting of multiple short contigs with appreciable sequencing errors can still yield many insights into the functionality of putative proteins, as has the one-pass fluorescent data published for M. genitalium (Peterson et al., 1993). After correction of frameshifts (see the Experimental procedures) and masking compositionally biased regions (low-entropy regions, Claverie and States, 1993), relatively stringent cut-offs were applied in similarity searches (BLAST Altschul et al., 1990):  $p \leqslant 10^{-9}$ ; FASTA (Pearson and Lipman, 1988): opt score ≥ 150) in order to identify matches with proteins that are clearly related to the putative gene products of M. capricolum. We have identified 176 distinct regions within the 372 contigs that fulfilled these criteria; their translated primary structures have obvious homologues. Among the 176 regions that match database proteins, several pairs were identified that correspond to non-overlapping segments of a single database protein. Although it is somewhat difficult to judge whether multiple hits (e.g. in the N-terminal and C-terminal portions of a database protein) characterize only a single gene product or different mycoplasma proteins of a multigene family, 22 of these pairs were considered to be from the same protein and were merged and considered in the statistics as a single entry (resulting in 154 unique protein hits, Table 1). In addition to this first fully automatic procedure, various sequence analysis tools (Bork et al., 1992; Koonin et al., 1994) were used to verify more subtle relationships, which revealed another 61 regions of similarity. The 215 distinct proteins thus identified (Tables 1 and 2) represent the minimum number of proteins encoded by the ORFs in the 372 contigs. Not a single gene was found that overlaps

Table 1. Statistics and calculation of gene density in M. capricolum.

Contigs	Characteristics	Size (kb)
202	Contain at least one protein and maybe RNA genes	
+43	Contain DNA repeats	
+10	Contain at least one RNA gene	
+117	Have no special characteristics — no sequence similarity	
=372	Total number of contigs	214
Proteins 154 +61 =215 +72 =287	Have easily identifiable homologues Have distant homologues Total number of identified proteins Other, non-overlapping ORFS >100 aa Estimated number of ORFs	≈103 +≈34 ≈137
Capacity 64%	Coding density (137 kb/214 kb)	
Function 75%	Function prediction rate (215/287)	

significantly with, or is contained in, another gene. There are 72 ORFs longer than 300 bases that do not match any other sequence in the databases and do not overlap with known genes. Most of these might code for additional proteins (Table 1) with as yet unidentified function.

At the DNA level, numerous matches were found (Table 2) with tRNA, rRNA and, surprisingly, snRNA-like sequences (Ushida and Muto, 1993). Also, the six conceptual translations of 43 contigs do not match any protein in the database but do match another M. capricolum contig (Table 2). Most of these cases appear to be DNA repeats, a striking feature of mycoplasma genomes. A relatively high number of repetitive non-coding regions has also been reported for M. pneumoniae (Wenzel and Herrmann, 1988; Ruland et al., 1990) and M. genitalium (Peterson et al., 1993). In spite of the large number of contigs characterized by sequence similarity with DNA repeats, RNA genes or proteins, 117 contigs still remain dissimilar to any sequence in public databases and, therefore, have no special characteristics, i.e. they do not have any ORFs longer than 300 bp or other obvious structural features. These contigs comprise  $\sim$  77 kb of sequence data.

### Coding density

The regions with similarity to database proteins or fragments cover 103 kb, which is a lower limit for estimating the coding density. For a more realistic estimate, the additional 72 ORFs (larger than 100 amino acids) with no database similarity were taken into account (an additional 34 kb). Some of these putative ORFs, though, may not code for proteins. This possible overestimate is compensated by the fact that shorter ORFs may have escaped identification in the remaining 77 kb with no obvious features. Therefore, the coding density can be roughly

estimated as (103kb coding+34kb putative coding)/ 214 kb, which is about 64% (Table 1). Of course, longer contigs are desirable for such an estimate, but the calculation is supported by the low coding density of M. genitalium, estimated at 55% (based on a much smaller data set; Peterson et al., 1993). It appears that the protein coding density of mycoplasmas is much lower than the 82% of Escherichia coli (Daniels et al., 1992; Blattner et al., 1993) and, surprisingly, even lower than the approximately 72% coding density in yeast (Fickett and Guigo, 1993; Dujon et al., 1994). Therefore, reduction in genome size does not necessarily lead to an increase in coding density. Instead, M. capricolum has apparently retained or accumulated non-coding DNA, the function of which remains to be elucidated.

#### Genome size and its coverage

Almost 30% of published rare restriction sites are present in the random sample of 214 kb (Table 3). This would seem to support earlier estimates of the genome size based on two-dimensional denaturating gradient gel electrophoresis, i.e. 724 kb (Poddor and Maniloff, 1989). Calculations based on pulsed-field gel electrophoresis (PFGE) (Whitely et al., 1991) predict a genome size of 1070 kb, a possible overestimate caused by the aberrant movement of ATrich restriction fragments in pulsed-field gels (Poddor and Maniloff, 1989; Maniloff, 1989). This explanation has been rebutted, however (Robertson et al., 1990; Neimark and Lange, 1990), and a final conclusion can only be drawn when the entire genome has been sequenced.

If the estimate of 500 proteins in M. capricolum (Muto et al., 1987) is correct (which would correlate with the smaller genome size estimate), then the approximately 287 proteins identified in this work represent about half of the total number of proteins in this organism. Yet, this fraction (287 proteins in 214kb) cannot be extrapolated to the whole genome, as most ORFs analysed here contain only partial sequences, i.e. future contigs will often contain sequences of genes already identified.

#### Prediction of function by similarity

The 215 proteins identified as a result of similarity searches reflect a high success rate in predicting function by sequence analysis in randomly sequenced genomes. Considering the additional 72 putative protein-coding ORFs larger than 100 amino acids, the percentage of proteins with homologues in public databases (215/287) is about 75% of those sequenced (Table 1). This level is much higher than in other large-scale sequencing projects, where the percentage varies between 33% (Wilson et al., 1994) and 61% (Koonin et al., 1994), with an average of about 50% throughout all phyla (Blattner et al., 1993; Adams et al., 1993; Honoré et al., 1993; Glaser et al., 1993; Bork et al., 1994). This high percentage is probably the result of the reduction of a parasitic genome to the genes coding for essential enzymes, which are usually well characterized and therefore over-represented in current databases, as well as the extensive characterization of other prokaryotes such as E. coli.

Finding homologous proteins in databases does not necessarily imply a precise functional prediction (Bork et al., 1994). Yet, for most of the proteins shown in Table 2, the best matching protein or protein family permits transfer of at least some functional information. This, in turn, enables us to draw conclusions about metabolic pathways and to obtain some insight into how M. capricolum manages to maintain cellular processes with such a small genome.

#### From ORFs to physiology

As long as the complete genome sequence is unavailable. only the presence of a certain protein but not its absence is conclusive. The enzymes already identified are useful markers of biochemical pathways of M. capricolum and closely related species.

Intermediary metabolism. As predicted by biochemical methods, glycolysis, as one of the major catabolic pathways, is certainly present in M. capricolum. Partial ORFs similar to nearly all glycolytic enzymes were found as well as components of the pyruvate dehydrogenase complex (Table 2). However, only one protein, transketolase, of the pentose phosphate cycle was identified. Together with the fact that no enzymes of the pentose phosphate cycle have been found yet in M. genitalium (Peterson et al., 1993), the intactness of this important pathway remains to be verified. The absence so far of enzymes of the citric acid cycle tentatively confirms that mycoplasmas take other routes for energy storage. A major enzyme, ornithine carbamoyltransferase, of the arginine hydrolase pathway was identified and may be utilized for ATP production, although other roles of carbamoylphosphates are possible (Pollak, 1992). Energy metabolism is often associated with the production of reduced NADH (e.g. in glycolysis). NADH might be further processed by flavoproteins and an NADH oxidase (Pollak, 1992). This is an alternative route to the respiratory chain involving cytochromes transferring released protons to oxygen (Pollak, 1992). Indeed, we found at least one protein similar to NADH oxidase (encoded by an ORF on the contig with EMBL accession number Z33133, Table 2). This alternative pathway might be coupled to ATP synthesis. As in M. genitalium (Peterson et al., 1993), several components of a membrane-associated ATP synthase were also identified.

 Table 2. Sequence similarities of M. capricolum contigs with database entries.

Metabolite trans	Bases	Best hit	P-value	Frame	Function
	port and activa	tion			
Z33018	1201	LKTB_ACTAC	$1.5 \times 10^{-22}$	+2	ABC transporter
Z33019_2	1444	PT2S_STRMU	$6.1 \times 10^{-11}$	+1	Phosphotransferase Ell (sucrose)
Z33047	855	PT2X_ECOLI	$1.7 \times 10^{-9}$	+1*	Phosphotransferase Ell (sucrose) Phosphotransferase Ell (maltose)
Z33074_1	1623	PT2G_ECOLI	$9.3 \times 10^{-31}$	+3	Phosphotransferase Ell (glucose)
Z33097	808	ATMB_SALTY	$9.6 \times 10^{-30}$	+1*	Mostransporting ATPose District
Z33098_2	848	GYLA_STRCO	$4.1 \times 10^{-20}$	+3	Mg-transporting ATPase, P-type
Z33100	803	PT2G_BACSU	$1.9 \times 10^{-15}$	+3	Glycerol facilitator protein
Z33102	1006	ATMB_SALTY	$6.9 \times 10^{-33}$	-1*	Phosphotransferase Ell (glucose)
233105	1174	ARAG_ECOLI	$3.3 \times 10^{-22}$	-3*	Mg-transporting ATPase
33112	271	PT2G_BACSU	$6.5 \times 10^{-7}$	-3	Arabinose-transport protein
33141	284	PT2G_BACSU	$3.0 \times 10^{-10}$	-3 +1	Phosphotransferase Ell (glucose)
33144	620	GLNQ_BACST	$2.0 \times 10^{-19}$		Phosphotransferase Ell (glucose)
33157	495	HMT1_SCHPO	$7.9 \times 10^{-27}$	+3	ABC transporter (glutamine)
33174	203	P29_MYCHR	$1.4 \times 10^{-13}$	-1*	ABC transporter (MDR subfamily)
33178	579	MDR_PLAFF		+1*	ABC transporter
33179	586		$4.8 \times 10^{-48}$	-3	ABC transporter (MDR subfamily)
33187	625	BRAF_PSEAE	$1.8 \times 10^{-9}$	<b>–</b> 1	ABC transporter (amino acids)
33210		PT3M_ENTFA	$1.7 \times 10^{-17}$	+3*	Phosphotransferase EIII (mannitol)
33222	408	PT2N_ECOLI	$1.1 \times 10^{-6}$	<b>-2</b>	Phosphotransferase EII (NAG)
	589	AMIE_STRPN	$6.1 \times 10^{-8}$	-3	ABC transporter (oligopeptide)
33224	353	PT2G_BACSU	$7.4 \times 10^{-4}$	+2	Similar to phosphotransferase EII
33225	188	GLPF_BACSU	$7.0 \times 10^{-2}$	+3	Glycerol facilitator protein
33229	422	OPPD_BACSU	$1.1 \times 10^{-21}$	-3	ABC transporter (oligopeptide)
33249	265	CYSA_SYNP7	$2.1 \times 10^{-7}$	+2	ABC transporter (sulphate)
33251	406	ATC2_YEAST	$2.1 \times 10^{-15}$	+3	Ca <sup>2+</sup> -transporting ATPase
33253	208	PT2M_ECOLI	$4.9 \times 10^{-7}$	-1	Phosphotranferase EII (mannitol)
33266	580	ATC1_YEAST	$1.2 \times 10^{-39}$	<b>-1*</b>	Ca <sup>2+</sup> -transporting ATPases
33293	242	PT2G_BACSU	$9.3 \times 10^{-13}$	+2	Phosphotransferase Ell (glucose)
33307	183	AROP_ECOLI	$1.6 \times 10^{-4}$	-3	Aromatia transport metalia (giucose)
33314	283	GLPF_BACSU	$2.3 \times 10^{-19}$	-3 + <b>1</b> *	Aromatic transport protein (permease)
33339	539	CYSA_SYNP7	$1.4 \times 10^{-23}$	-2	Glycerol facilitator protein
33340	353	PT2S_VIBAL	$5.2 \times 10^{-12}$		ABC transporter (sulphate)
33348_2	1708	YHBI_ECOLI	$1.7 \times 10^{-16}$	-1	Phosphotransferase EII (sucrose)
33348_3	1708	PT2F_ECOLI		+2	Similar to phosphotransferase EII
33353	829	PT2G_BACSU	$1.6 \times 10^{-10}$	+2	Phosphotransferase Ell (fructose)
33361	1191		$2.5 \times 10^{-34}$	+3*	Phosphotransferase EII (glucose)
33363	463	YBBA_ECOLI PT3M_ECOLI	$7.9 \times 10^{-23}$	-2	ABC transporter (MDR subfamily)
		F TOIM_ECOLI	$3.5 \times 10^{-6}$	+2	Similar to phosphotransferase EIII
mino acid meta	<u>bolism</u>				
33027	1144	AQPETBDCP_3	$9.5 \times 10^{-5}$	+1	Similar to carino protects DE
33032_1	1250	DAPE_ECOLI	1.2 × 10 <sup>-9</sup>	2*	Similar to serine protease RE
33281	179	METX_ECOLI	$2.0 \times 10^{-25}$	-2	SDAP desuccinylase
33292	295	LON_ECOLI	$1.5 \times 10^{-32}$	-2 -1	S-adenosyl-Met synthetase 2
33299	451	METK_HUMAN	$3.9 \times 10^{-27}$		ATP-binding endopeptidase La
33311	157	OTCC_NEICI	$3.9 \times 10^{-15}$	-3	S-adenosyl-Met synthetase (alpha)
33349	1106		3.0 × 10 10	+1	Ornithine carbamoyltransferase
33357		OTC_ASPNG	$3.2 \times 10^{-38}$	+3	Ornithine carbamoyltransferase
30007	922	LON_ECOLI	$3.6 \times 10^{-74}$	+3	ATP-binding endopeptidase LA
ucleotide metal	oolism				
33022	1759	PYRG_ECOLI	$3.1 \times 10^{-123}$	+2*	CTP synthase
33033	566	YC42_CAEEL	$6.6 \times 10^{-9}$	-3	
33044	999	C1TM_YEAST	$7.8 \times 10^{-40}$	-3 +2*	Similar to dCTP deaminases
33066	900	DEOK_HUMAN	$3.8 \times 10^{-4}$		C1-THF synthase
33079	971	KTHY_HUMAN	5.2 × 10 <sup>-5</sup>	+1*	Pyrimidine kinase
33170	533	RNC_ECOLI		+3	Pyrimidine kinase
33218	124	KPRS_ECOLI	$1.5 \times 10^{-16}$	+3	Ribonuclease III
33263	127		$2.7 \times 10^{-6}$	+1	Phosphoribosyl-PP-kinase
33279	297	UPP_ECOLI	$2.3 \times 10^{-9}$	+2	Uracil-phosphoribosyltransferase
		KITH_AMEPV	$2.1 \times 10^{-6}$	+3*	Thymidine kinase
	363	PURA_HUMAN	$3.8 \times 10^{-33}$	-3	Adenylosuccinate synthetase
33336					Adenylate kinase
33336 ad_Mycca					
33336 ad_Mycca pid metabolism					
33336 ad_Mycca pid metabolism 33013_1	1904	PGSA_ECOLI	$6.2 \times 10^{-10}$	+3	Phosphatidylalycerol P cynthese
33336 ad_Mycca pid metabolism 33013_1 33059_2		PGSA_ECOLI TPES_PSEPU		+3 -2	Phosphatidylglycerol-P synthase
33336 ad_Mycca ipid metabolism 33013_1 33059_2 33081	1904		$1.2 \times 10^{-4}$	-2	Similar to lipase/esterase family
	1904 1484	TPES_PSEPU			Phosphatidylglycerol-P synthase Similar to lipase/esterase family Similar to GIpE and PspE proteins Lipoamide acyltransferase (E2)

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Table 2. Continued

Contig	Bases	Best hit	P-value	Frame	Function
Carbohydrate m	etabolism/ener	gy storage			
Z33030_1	1748	OUTB_BACSU	$9.8 \times 10^{-18}$	-1	ATP pyrophosphatase <sup>a</sup>
Z33051	710	ATPB_PECFR	$2.5 \times 10^{-44}$	+1*	ATP synthase (beta)
Z33053_1	627	THIO_BACSU	$1.6 \times 10^{-19}$	+3	Thioredoxin
Z33055	1061	NAGB_ECOLI	$3.0 \times 10^{-30}$	-2	Glucosamine-6-isomerase
Z33072_1	2057	K1PF_ECOLI	$6.0 \times 10^{-11}$	-2	1-Phosphofructokinase
Z33074_2	1623	YICI_ECOLI	$8.0 \times 10^{-14}$	+1	Similar to isomaltase-sucrases
Z33075	225	ALF2_RHOSH	$6.4 \times 10^{-6}$	<b>-1*</b>	Fructose-bisphosphate aldolase
Z33087_2	859 710	PMGY_MAIZE	$3.8 \times 10^{-17}$	+3	Phosphoglyceromutase
Z33089 Z33096	710	NADO_THEBR	$1.3 \times 10^{-4}$	<b>-1</b>	Similar to NADH oxidase
Z33098_1	875 848	DHAC_RAT	$4.7 \times 10^{-26}$	+3	Aldehyde dehydrogenase
Z33104	1036	GLPK_ECOLI	$6.8 \times 10^{-20}$	+1*	Glycerol kinase
Z33109	1300	O16G_BACCE TSR_BACSU	$3.0 \times 10^{-79}$ $1.5 \times 10^{-24}$	+3*	Cytoplasmic oligo-1,6-glucosidase
Z33110	545	ODPB_BACSU	$2.0 \times 10^{-57}$	-1 -2	RNA synthesis protein/fructose-Bi-P-aldolase
Z33126	534	MGLC_ECOLI	$4.3 \times 10^{-7}$	+3 +2	Pyruvate DH E1 (beta)
Z33133	153	S26965P	8.8 × 10 <sup>-13</sup>	+2 -1	Membrane forming protein
Z33226	486	YLP2_PSEPU	$1.1 \times 10^{-17}$	-3	Similar to Streptococcus faecalis NADH oxidase Similar to dihydrolipoamide DH
Z33228	372	YINL_LISMO	$1.7 \times 10^{-13}$	+2*	Similar to diriydrolipoamide DH Similar to 7-alpha-hydroxysteroid DH
Z33230	121	TKT_RHOSH	$1.2 \times 10^{-11}$	+1	Transketolase
Z33232	267	ENO_ARATH	$5.4 \times 10^{-22}$	-2*	Enolase
Z33234	178	YIEK ECOLI	$2.8 \times 10^{-7}$	-3	Similar to glucosamine isomerase
Z33235_1	681	ATPD_SYNY3	$1.1 \times 10^{-6}$	+1*	ATP synthase (delta)
Z33235_2	681	ATPA_BACME	$2.0 \times 10^{-22}$	+1	ATP synthase (alpha)
Z33250	427	PT1_STACA	$4.8 \times 10^{-21}$	-1*	PEP P-transferase
Z33255	187	ENO_ECOLI	$1.4 \times 10^{-20}$	+1	Enolase
Z33265	152	PT1_STACA	$1.9 \times 10^{-5}$	+3	PEP P-transferase
Z33277	420	XYLK_STAXY	$2.6 \times 10^{-4}$	-1	Sugar kinase (hexokinase family)
Z33286	510	IPYR_THEP3	$1.9 \times 10^{-28}$	+2	Inorganic pyrophosphatase
Z33288	291	PGKY_WHEAT	$1.2 \times 10^{-35}$	+1	Cytoplasmic phosphoglycerate kinase
Z33313	166	ODP2_BACSU	$1.9 \times 10^{-15}$	-2	Dihydrolipoamide acetyltransferase (E2)
Z33330	319	ATPA_ECOLI	$2.4 \times 10^{-12}$	3	ATP synthase (alpha)
Z33331	176	K6PF_BACST	$1.6 \times 10^{-8}$	+3	6-Phosphofructokinase
Z33348_1	1708	K1PF_RHOCA	$1.5 \times 10^{-9}$	+3	1-Phosphofructokinase
Z33368	2167	MGLA_ECOLI	$6.4 \times 10^{-15}$	<b>– 1</b>	Galactoside-binding protein
Z33370	1329	ATPB_THEP3	$9.6 \times 10^{-143}$	<b>-1*</b>	ATP synthase (beta)
Other metabolic	enzymes				
Z33006	897	YIDA_ECOLI	$1.3 \times 10^{-16}$	+3	Hydrolase (HAD family <sup>b</sup> )
Z33015	1156	BAIC_EUBSP	$3.8 \times 10^{-14}$	+2	Similar to trimethylamine DH
Z33083	567	AMID_RHOER	$4.6 \times 10^{-9}$	-3	Amidase
Z33087_1	859	YIDA_ECOLI	$5.2 \times 10^{-10}$	+1	hydrolase (HAD family <sup>2</sup> )
Z33177	214	UGPQ_ECOLI	$7.8 \times 10^{-3}$	-2	Similar to phosphodiesterase
Z33272	283	GSHR_HUMAN	$2.4 \times 10^{-4}$	+2	FAD/NAD-binding reductase
Z33273	257	PCR_AVESA	$8.4 \times 10^{-3}$	<b>–1</b>	NAD-binding oxidoreductase
Z33290	236	YHDG_ECOLI	$1.2 \times 10^{-21}$	-3	Similar to DH subfamily (BAIC_EUBSP)
Z33346	778	FOLC_ECOLI	$8.2 \times 10^{-5}$	<b>-</b> 1*	Similar to folylpoly-Glu synthetase
Z33362_2	1259	YHDG_ECOLI	$5.5 \times 10^{-7}$	+2	Similar to DH subfamily (BAIC_EUBSP)
DNA replication	, repair and rec	ombination			
Z33035	2049	GYRA_BACSU	$1.7 \times 10^{-54}$	-2*	DNA gyrase (alpha)
Z33054	580	A30868P	$2.9 \times 10^{-9}$	+2	Transposase
Z33057	752	DNAA_MYCCA	$1.6 \times 10^{-119}$	+3	DnaA protein
Z33071	1102	GYRA_STAAU	$1.8 \times 10^{-13}$	-3	DNA gyrase (alpha)
Z33091	594	DP3A_ECOLI	$3.6 \times 10^{-30}$	+1	DNA polymerase III (alpha)
Z33108	752	GYRB_BACSU	$2.3 \times 10^{-35}$	<b>2*</b>	DNA gyrase (beta)
Z33173	307	TOP1_SYNP7	$1.5 \times 10^{-2}$	1*	Topoisomerase I
Z33193	234	DPO1_STRPN	$1.0 \times 10^{-12}$	1	DNA polymerase I
Z33201	495	DP3A_BACSU	$5.1 \times 10^{-5}$	<u> </u>	DNA polymerase III (alpha)
Z33239	367	RNH2_ECOLI	$7.9 \times 10^{-14}$	-3	Ribonuclease HII
Z33252	286	DP3A_BACSU	$1.0 \times 10^{-8}$	-1	DNA polymerase III (alpha)
Z33256	54	GYRB_ECOLI	$1.9 \times 10^{-1}$	+3	DNA gyrase (beta)
Z33302	129	DNLJ_ECOLI	$3.5 \times 10^{-8}$	-3	DNA ligase
Z33305	417	GYRA_BACSU	$1.2 \times 10^{-39}$	-1	DNA gyrase (alpha)
Z33335	385	DP3A_BACSU	$2.5 \times 10^{-30}$	+3*	DNA polymerase III (alpha)
Dp3b_Mycca Z33060_1	1308	ADMI VEACT	7040=3	,	DNA polymerase III (beta)
200000_I	1300	APN1_YEAST	$7.9 \times 10^{-3}$	-1	Similar to AP-endonucleases

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Table 2. Continued

Contig	Bases	Best hit	P-value	Frame	Function
Z33120	621	UVRA_MICLU	9.1 × 10 <sup>-65</sup>	+1	Excinuclease ABC (A)
Z33128	483	RECM_BACSU	$1.3 \times 10^{-11}$	+1	DNA repair protein RecM
Z33191	510	UVRC_BACSU	$9.9 \times 10^{-7}$	+2	Excinuclease ABC (C)
233233	585	MTS1_STRSA	1.9 × 10 <sup>-9</sup>	-1*	
33254	340	APN1_YEAST	$2.4 \times 10^{-4}$	+1	Modification methylase
233278	208	S18707P	$6.2 \times 10^{-5}$		Similar to AP-endonucleases
233332	263	JQ0894P		-1	UvrD protein (also in M. genitalium)
233345			$1.4 \times 10^{-13}$	+2*	P115 protein, RecF/RecN family
	1065	UVRB_ECOLI	$3.9 \times 10^{-127}$	+3	Excinuclease ABC (B)
Z33355	1401	YAT3_RHORU	$1.1 \times 10^{-28}$	+2	RecF/RecN family
33373	299	UVRA_MICLU	$6.5 \times 10^{-31}$	+3	Excinuclease ABC (A)
Cell division					
233086	245	FTSH_ECOLI	$3.1 \times 10^{-24}$	+2	Cell division protein FTSH
33209	186	FTSH_ECOLI	$1.0 \times 10^{-5}$	+2	Cell division protein FTSH
33322	149	FTSZ_BACSU	$6.5 \times 10^{-18}$	-2	Cell division protein FTSZ
ranscription fac	ctors				
33019_1	1444	LACR_STAAU	$2.7 \times 10^{-24}$	+2	Sugar repressor (HTH family)
33041	1410	RPOB PSEPU	$1.2 \times 10^{-20}$	+2*	
33050_2	1490	RPOA_BACSU			DNA-directed RNA polymerase (beta)
33052			$7.7 \times 10^{-57}$	-1	DNA-directed RNA Polymerase (alpha)
	880	PILB_NEIGO	$1.3 \times 10^{-43}$	+3	Transcription repressor
33063	448	PHNF_ECOLI	$7.4 \times 10^{-2}$	-2	HTH motif (HTH family)
33072_2	2057	LACR_LACLA	$2.0 \times 10^{-15}$	-2	lac repressor
33085	1500	RPOB_ECOLI	$3.3 \times 10^{-81}$	3*	DNA-directed RNA polymerase (beta)
33123	325	HMGT_ONCMY	$9.6 \times 10^{-4}$	+3	Similar to HMG proteins
33203	290	GREA_ECOLI	$3.6 \times 10^{-13}$	+1*	Transprintion standard for
33275	209	XYLR_BACSU	$1.6 \times 10^{-1}$		Transcription elongation factor
33297	160	_		+2	Similar to xylose repressor (HTH family)
33338		RPOC_MYCLE	$4.1 \times 10^{-19}$	<b>-1</b>	DNA-directed RNA polymerase (beta)
	278	RPOD_NOSCO	$8.8 \times 10^{-9}$	-3	DNA-directed RNA polymerase (delta)
	ein biosynthesi	s and ribosomal proteins	3		
33016_1	1476	EFTS_ECOLI	$8.5 \times 10^{-27}$	-3	Elongation factor EF-TS
33034_2	840	IF1_BACSU	$5.0 \times 10^{-35}$	-2	Initiation factor IF-1
33068	852	EFG_SPIPL	$1.7 \times 10^{-95}$	+1*	
33076_2	1299	SPOU_ECOLI	$9.0 \times 10^{-10}$	-2	Elongation factor G
33271	377	FMT_ECOLI	1.1 × 10 <sup>-13</sup>		rRNA methylase
		I WII_ECOLI		+2*	Met-tRNA formyltransferase
33384		DE1 FOOL			
	487	RF1_ECOLI	$8.8 \times 10^{-18}$	<b>-2*</b>	Peptide chain release factor 1
33284 33303	487 131	EFTU_ECOLI	$1.4 \times 10^{-12}$	+3*	Peptide chain release factor 1 Elongation factor TU
33303 33352_1	487 131 1629				Elongation factor TU
33303 33352_1	487 131	EFTU_ECOLI	$1.4 \times 10^{-12}$	+3* +3	Elongation factor TU Initiation factor IF-3
33303 33352_1 33371	487 131 1629	EFTU_ECOLI IF3_BACST IF2_BACST	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$	+3* +3 -2*	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2
33303 33352_1 33371 33008	487 131 1629 1151 878	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI	$   \begin{array}{r}     1.4 \times 10^{-12} \\     3.7 \times 10^{-32} \\     2.2 \times 10^{-87} \\     4.3 \times 10^{-32}   \end{array} $	+3* +3 -2* +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase
33303 33352_1 33371 33008 33021	487 131 1629 1151 878 865	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$	+3* +3 2* +3 +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase
33303 33352_1 33371 33008 33021 33026	487 131 1629 1151 878 865 678	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$	+3* +3 -2* +3 +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3	487 131 1629 1151 878 865 678 840	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$	+3* +3 -2* +3 +3 +1	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase
33303 33352_1 33371 33008 33021 33026 33034_3 33042	487 131 1629 1151 878 865 678 840 1582	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$	+3* +3 -2* +3 +3 +1 -3 -3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33034_3 33042 33048	487 131 1629 1151 878 865 678 840 1582 1474	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$	+3* +3 -2* +3 +3 +1	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33042 33048 33053_2	487 131 1629 1151 878 865 678 840 1582 1474 627	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$	+3* +3 -2* +3 +3 +1 -3 -3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33042 33048 33053_2 33056	487 131 1629 1151 878 865 678 840 1582 1474 627 1031	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33042 33048 33053_2 33056	487 131 1629 1151 878 865 678 840 1582 1474 627	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3 -2	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI SYM_ECOLI SYM_BACSU	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$	+3* +3 -2* +3 +1 -3 -3 +1 -3 -2 -3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Val-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYT1_ECOLI SYT1_BACSU SYT1_BACSU SYV_BACST SYI_METTH	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$	+3* +3 -2* +3 +1 -3 -3 +1 -3 -2 -3 +1*	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127 33137_1	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYM1_ECOLI SYM1_ECOLI SYM1_ECOLI SYM1_ECOLI SYM1_ECOLI SYM1_BACSU SYV_BACST SYI_METTH SYH_STREQ	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$ $3.6 \times 10^{-14}$	+3* +3 -2* +3 +1 -3 -3 +1 -3 -2 -3 +1* +1	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase His-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127 33137_1 33137_2	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYT1_BACSU SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-14}$ $1.8 \times 10^{-13}$	+3* +3 -2* +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase His-tRNA synthetase Asp-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127 33137_1 33137_1 33137_2 33153	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYK1_ECOLI SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$ $3.6 \times 10^{-14}$ $1.8 \times 10^{-13}$ $3.8 \times 10^{-53}$	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3 +1*	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase His-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127 33137_1 33137_1 33137_2 33153 33154	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735 63	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYK1_ECOLI SYK1_ECOLI SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU SYE_BACST	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$ $3.6 \times 10^{-14}$ $1.8 \times 10^{-13}$ $3.8 \times 10^{-53}$ $4.7 \times 10^{-1}$	+3* +3 -2* +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Val-tRNA synthetase His-tRNA synthetase His-tRNA synthetase Asp-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33127 33137_1 33137_1 33137_2 33153 33154 33244	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735 63 306	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYK1_ECOLI SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$ $3.6 \times 10^{-14}$ $1.8 \times 10^{-13}$ $3.8 \times 10^{-53}$ $4.7 \times 10^{-1}$ $5.6 \times 10^{-11}$	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3 +1*	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase His-tRNA synthetase His-tRNA synthetase Phe-tRNA synthetase Phe-tRNA synthetase Phe-tRNA synthetase Phe-tRNA synthetase Glu-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33157 33137_1 33137_1 33137_2 33153 33154	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735 63	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYK1_ECOLI SYK1_ECOLI SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU SYE_BACST	$1.4 \times 10^{-12}$ $3.7 \times 10^{-32}$ $2.2 \times 10^{-87}$ $4.3 \times 10^{-32}$ $7.1 \times 10^{-14}$ $2.5 \times 10^{-57}$ $5.8 \times 10^{-43}$ $3.7 \times 10^{-31}$ $7.4 \times 10^{-63}$ $2.8 \times 10^{-17}$ $7.3 \times 10^{-41}$ $4.3 \times 10^{-38}$ $1.6 \times 10^{-10}$ $3.6 \times 10^{-14}$ $1.8 \times 10^{-13}$ $3.8 \times 10^{-53}$ $4.7 \times 10^{-1}$ $5.6 \times 10^{-11}$	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3 +1* +1	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase Ile-tRNA synthetase His-tRNA synthetase His-tRNA synthetase Asp-tRNA synthetase Phe-tRNA synthetase Calu-tRNA synthetase
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33303 33352_1 33371 33008 33021 33026 33034_3 33042 33048 33053_2 33056 33122 33157 33137_1 33137_1 33153 33154 33244 33261 33276	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735 63 306 192 401	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYD_ECOLI SYT1_BACSU SYV_BACST SYI_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU SYE_BACST SYL_ECOLI SYE_BACST SYL_ECOLI	1.4 × 10 <sup>-12</sup> 3.7 × 10 <sup>-32</sup> 2.2 × 10 <sup>-87</sup> 4.3 × 10 <sup>-32</sup> 7.1 × 10 <sup>-14</sup> 2.5 × 10 <sup>-57</sup> 5.8 × 10 <sup>-43</sup> 3.7 × 10 <sup>-63</sup> 2.8 × 10 <sup>-17</sup> 7.3 × 10 <sup>-41</sup> 4.3 × 10 <sup>-38</sup> 1.6 × 10 <sup>-10</sup> 3.6 × 10 <sup>-14</sup> 1.8 × 10 <sup>-13</sup> 3.8 × 10 <sup>-53</sup> 4.7 × 10 <sup>-1</sup> 5.6 × 10 <sup>-1</sup> 5.6 × 10 <sup>-11</sup> 7.2 × 10 <sup>-10</sup> 3.5 × 10 <sup>-28</sup>	+3* +3 -2* +3 +3 +1 -3 -3 +1 -3 -2 -3 +1* +1 +3 -3* +3	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase His-tRNA synthetase His-tRNA synthetase Asp-tRNA synthetase Glu-tRNA synthetase Clu-tRNA synthetase Clu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase
33303 33352_1 33371 33008 33021 33026 33034_3 33042 33053_2 33056 33122 33157 33137_1 33137_1 33137_2 33153 33154 33244 33261 33276 33289	487 131 1629 1151 878 865 678 840 1582 1474 627 1031 790 298 700 700 735 63 306 192 401 622	EFTU_ECOLI IF3_BACST IF2_BACST SYL_ECOLI SYL_ECOLI SYN_ECOLI AMPM_BACSU SYA_ECOLI SYD_ECOLI SYT1_BACSU SYL_BACST SYL_METTH SYH_STREQ SYD_ECOLI SYFA_BACSU SYE_BACST SYL_ECOLI SYE_BACST SYL_ECOLI SYE_BACST SYL_ECOLI SYE_BACST SYL_ECOLI	1.4 × 10 <sup>-12</sup> 3.7 × 10 <sup>-32</sup> 2.2 × 10 <sup>-87</sup> 4.3 × 10 <sup>-32</sup> 7.1 × 10 <sup>-14</sup> 2.5 × 10 <sup>-57</sup> 5.8 × 10 <sup>-43</sup> 3.7 × 10 <sup>-31</sup> 7.4 × 10 <sup>-63</sup> 2.8 × 10 <sup>-17</sup> 7.3 × 10 <sup>-41</sup> 4.3 × 10 <sup>-38</sup> 1.6 × 10 <sup>-10</sup> 3.6 × 10 <sup>-14</sup> 1.8 × 10 <sup>-13</sup> 3.8 × 10 <sup>-53</sup> 4.7 × 10 <sup>-1</sup> 5.6 × 10 <sup>-11</sup> 7.2 × 10 <sup>-10</sup> 3.5 × 10 <sup>-28</sup> 8.5 × 10 <sup>-33</sup>	+3* +3 -2* +3 +1 -3 -1 -3 -1 -3 +1* +1 +3 -3* +3 +3*	Elongation factor TU Initiation factor IF-3 Initiation factor IF-2 Leu-tRNA synthetase Leu-tRNA synthetase Asn-tRNA synthetase M-aminopeptidase Ala-tRNA synthetase Asp-tRNA synthetase Lys-tRNA synthetase Thr-tRNA synthetase Thr-tRNA synthetase Val-tRNA synthetase His-tRNA synthetase His-tRNA synthetase Asp-tRNA synthetase Glu-tRNA synthetase Clu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Glu-tRNA synthetase Ile-tRNA synthetase
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Table 2. Continued

Contig	Bases	Best hit	P-value	Frame	Function
Z33050_3	1490	RS11_BACST	$1.7 \times 10^{-7}$	-1	30S ribosomal protein
Z33076_1	1299	RL33_BACST	$1.3 \times 10^{-9}$	-2	50S ribosomal protein
Z33236	396	RS18_BACST	$3.3 \times 10^{-23}$	-2*	30S ribosomal protein
Z33246	180	RL10_STRAT	$1.4 \times 10^{-2}$	+3	Similar to 50S ribosomal protein
Z33291	324	RL6_MYCCA	$5.8 \times 10^{-55}$	-2*	50S ribosomal protein
Z33312	244	RS11_BACSU	$1.2 \times 10^{-27}$	<b>~1*</b>	30S ribosomal protein
Z33327_1	422	RL23_MYCCA	$8.6 \times 10^{-32}$	<b>~2*</b>	50S ribosomal protein
Z33327_2	422	RL4_MYCCA	$2.3 \times 10^{-38}$	-3	50S ribosomal protein
Z33333	273	RL34_PSEAE	$5.2 \times 10^{-15}$	-3	59S ribosomal protein
Z33343_1	339	RS9_BACST	$7.1 \times 10^{-25}$	-3	30S ribosomal protein
Z33343_2	339	RL13_HAESO	$7.2 \times 10^{-4}$	<b>-1</b>	50S ribosomal protein
Z33352_2	1629	RL35_BACST	$1.2 \times 10^{-7}$	+1*	50S ribosomal protein
Z33352_3	1629	RL20_BACST	$5.1 \times 10^{-7}$	+3	50S ribosomal protein
Z33366_1	1299	RL11_THEMA	$2.3 \times 10^{-42}$	+2	50S ribosomal protein
Z33366_2	1299	RL1_BACST	$2.6 \times 10^{-95}$	+1	50S ribosomal protein
RI2_Mycca					50S ribosomal protein
Rl3_Mycca					50S ribosomal protein
RI5_Mycca					50S ribosomal protein
Rs8_Mycca					30S ribosomal protein
RI15_Mycca					50S ribosomal protein
RI24_Mycca					50S ribosomal protein
Rs10_Mycca					30S ribosomal protein
Rs17_Mycca					30S ribosomal protein
Rs19_Mycca					30S ribosomal protein
Protein folding					<b>F. 1.1.1</b>
<del></del>			100		
Z33106	753	DNAK_BACME	$5.6 \times 10^{-102}$	+1	DnaK protein
Z33267	364	YGRP_BACSU	$2.7 \times 10^{-11}$	-1	CLPA/CLPB family
Z33269	228	CLPX_ECOLI	$1.1 \times 10^{-3}$	+3	CLPA/CLPB family
Z33364_1	754	DNAJ_BACSU	$4.2 \times 10^{-19}$	-1	DnaJ protein
Z33364_2	754	DNAK_BORBU	$2.9 \times 10^{-20}$	-3*	DnaK protein
Z33365	569	TIG_ECOLI	$4.9 \times 10^{-19}$	+1	Trigger factor
Z33376	488	CLPB_ECOLI	$4.8 \times 10^{-50}$	-3	CLPA/CLPB family
Internal transport	and transloca	ation			
Z33092	549	SECA_BACSU	$2.6 \times 10^{-59}$	+3	Preprotein translocase SECA subunit
Z33142	595	PSTC_ECOLI	$2.7 \times 10^{-11}$	+1	Phosphate transport protein
Z33270	198	SECA_BACSU	$7.8 \times 10^{-7}$	-2	Preprotein translocase SECA subunit
Z33315	221	AMIC_STRPN	$1.2 \times 10^{-4}$	- <b>2</b>	Oligopeptide transporter
Z33328	396	SECY_MYCCA	$1.1 \times 10^{-72}$	-1	Preprotein translocase SECY subunit
Z33334	254	PSTA_ECOLI	$6.8 \times 10^{-9}$	+3	Phosphate transporter PstA
Z33358	350	SRP5_MYCMY	$8.3 \times 10^{-50}$	3*	Signal recognition particle
Z33374_1	2095	AMIE_STRPN	$2.8 \times 10^{-96}$	-2*	Oligopeptide transporter
Z33374_2	2095	AMID_STRPN	$5.5 \times 10^{-30}$	-1*	Oligopeptide transporter
Z33375	241	AMIF_STRPN	$1.1 \times 10^{-36}$	+1	Oligopeptide transporter
Unclassified					
	070	S14881P	$6.5 \times 10^{-16}$	+2	
Z33012 1			0.5 X IU	+4	
Z33012_1 Z33012_2	872 872				Hypothetical yeast protein 1
Z33012_2	872	S14882P	$3.2 \times 10^{-8}$	+3	Hypothetical yeast protein 2
Z33012_2 Z33013_2	872 1904	S14882P GIDB_BACSU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$	+3 +1	Hypothetical yeast protein 2 Methyltransferases
Z33012_2 Z33013_2 Z33013_3	872 1904 1904	S14882P GIDB_BACSU OBG_BACSU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$	+3 +1 +1	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family
Z33012_2 Z33013_2 Z33013_3 Z33023	872 1904 1904 808	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$	+3 +1 +1 2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024	872 1904 1904 808 788	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$	+3 +1 +1 -2 -2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2	872 1904 1904 808 788 1748	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$	+3 +1 +1 -2 -2 -3	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33032_2	872 1904 1904 808 788 1748	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$	+3 +1 +1 -2 -2 -3 -1	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33032_2 Z33040	872 1904 1904 808 788 1748 1250	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$	+3 +1 +1 -2 -2 -3 -1 +1	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33030_2 Z33030_2 Z33040 Z33045_2	872 1904 1904 808 788 1748 1250 1019 640	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$	+3 +1 +1 -2 -2 -3 -1 +1 -3#	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33030_2 Z33032_2 Z33045_2 Z33045_2 Z33058	872 1904 1904 808 788 1748 1250 1019 640 719	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33032_2 Z33032_2 Z33045_2 Z33045_2 Z33058 Z33060_2	872 1904 1904 808 788 1748 1250 1019 640 719 1308	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33030_2 Z33040 Z33045_2 Z33058 Z33060_2 Z33103	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33032_2 Z33032_2 Z33040 Z33045_2 Z33058 Z33060_2 Z33103 Z33138	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR Small protein B
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33032_2 Z33032_2 Z33040 Z33045_2 Z33045_2 Z33060_2 Z33103 Z33138 Z33140	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815 583	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI YJIF_ECOLI	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$ $3.6 \times 10^{-6}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1 -2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR
Z33012_2 Z33013_2 Z33023 Z33024 Z33024 Z33030_2 Z33030_2 Z33040 Z33045_2 Z33058 Z33060_2 Z33103 Z33138 Z33140 Z33167	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815 583 935	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI YJIF_ECOLI OBG_BACSU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$ $3.6 \times 10^{-6}$ $5.7 \times 10^{-12}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1 -2 -2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR Small protein B
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33030_2 Z33040 Z33045_2 Z33058 Z33060_2 Z33103 Z33138 Z33140 Z33167 Z33192	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815 583 935 443	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI YJIF_ECOLI OBG_BACSU YP15_STAAU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$ $3.6 \times 10^{-6}$ $5.7 \times 10^{-12}$ $4.5 \times 10^{-3}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1 -2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR Small protein B Secreted protein downstream SMP
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2 Z33032_2 Z33045_2 Z33045_2 Z33058 Z33060_2 Z33103 Z33138 Z33140 Z33140 Z33167 Z33192 Z33195	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815 583 935 443 617	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI YJIF_ECOLI OBG_BACSU YP15_STAAU LEPA_ECOLI	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$ $3.6 \times 10^{-6}$ $5.7 \times 10^{-12}$ $4.5 \times 10^{-3}$ $9.7 \times 10^{-21}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1 -2 -2	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR Small protein B Secreted protein downstream SMP Similar to GTP-binding GTP1/OBG family Unknown function (see YDEU_BACSU) GTP-binding proteins
Z33012_2 Z33013_2 Z33013_3 Z33023 Z33024 Z33030_2	872 1904 1904 808 788 1748 1250 1019 640 719 1308 1306 815 583 935 443	S14882P GIDB_BACSU OBG_BACSU HP9ORFS_7 BSGENR_1GP OBG_BACSU HIS4_METVA YMG1_MYCGE A44803P Y311_BACSU YAAC_PSEFL A43577P SMPB_ECOLI YJIF_ECOLI OBG_BACSU YP15_STAAU	$3.2 \times 10^{-8}$ $1.0 \times 10^{-32}$ $1.8 \times 10^{-12}$ $3.6 \times 10^{-2}$ $4.5 \times 10^{-11}$ $2.4 \times 10^{-3}$ $2.2 \times 10^{-1}$ $6.3 \times 10^{-8}$ $6.9 \times 10^{-20}$ $8.8 \times 10^{-17}$ $1.2 \times 10^{-5}$ $4.5 \times 10^{-12}$ $2.1 \times 10^{-9}$ $3.6 \times 10^{-6}$ $5.7 \times 10^{-12}$ $4.5 \times 10^{-3}$	+3 +1 +1 -2 -2 -3 -1 +1 -3# +2 -3 -1 +1 -2 -1* +3	Hypothetical yeast protein 2 Methyltransferases Similar to GTP-binding GTP1/OBG family Hypothetical yeast protein 7 Unknown function Similar to GTP-binding GTP1/OBG family Similar to isomerases/dehydrogenases Unknown function Human pG1 protein GTP binding CD48/PAS1/SEC18 family Similar protein X in RPST-ILES region Regulatory Clostridium protein PfoR Small protein B Secreted protein downstream SMP Similar to GTP-binding GTP1/OBG family Unknown function (see YDEU_BACSU)

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Table 2. Continued

Contig	Bases	Best hit	P-value	Frame	Function
Z33342	344	P37_MYCHR	3.5×10 <sup>-3</sup>	-3	Surface protein (transport-associated)
Z33354	1538	A43863P	3.3×10 <sup>-25</sup>	-1	Similar to a haemolysin

#### RNAs

Z33007 (tRNA), Z33009 (tRNA), Z33010\_1 (tRNA), Z33025 (5S rRNA), Z33045\_1 (16S rRNA), Z33060\_3 (tRNA), Z33070 (snRNA-like), Z33094 (16S/23S rRNA), Z33172 (23S rRNA), Z33190 (snRNA-like), Z33223 (rRNA), Z33231 (23S rRNA)

#### Repeats

Z33005, Z33010\_2, Z33014, Z33036, Z33037, Z33038, Z33043, Z33046, Z33059\_1, Z33065, Z33078, Z33082, Z33101, Z33116, Z33117, Z33119, Z33124, Z33130, Z33131, Z33152, Z33164, Z33175, Z33185, Z33197, Z33198, Z33200, Z33204, Z33206, Z33211, Z33212, Z33217, Z33227, Z33228, Z33248, Z33248, Z33257, Z33258, Z33248, Z33257, Z33258, Z33259, Z33359, Z33360

Contig: EMBL accession number; an underscore followed by a number denotes a particular segment of a contig that has several distinct proteins or other features. The contigs are sorted according to functional categories.

Bases: length of contig in number of base pairs.

Best hit: SWISSPROT codes or PIR accession numbers for the database protein with the best sequence similarity.

P-value: significance of the similarity from the BLASTP program. All similarities with p-values above  $10^{-9}$  were verified by other techniques such as multiple alignments.

Frame: position of first codon and orientation of ORF relative to the beginning of the contig DNA sequence.

Similar to: denotes a statistically significant homology without precise assignment of function.

Repeat: used for all contigs with no homologues in protein databases that resemble at least one other *M. capricolum* contig or match known DNA repeats in nucleic acid sequence databases. *M. capricolum* proteins already in sequence databases have the SWISSPROT identifiers (Bairoch and Boeckmann, 1993) in column Contig. The longest continuous DNA segment known so far includes the origin of replication (Miyata *et al.*, 1993); a. This ATP pyrophosphatase domain is usually associated with other domains within one peptide chain (e.g. in GMP synthetase). Therefore, it might be only a subunit of a larger complex (data not shown).

b. HAD is a superfamily of enzymes that contains functionally diverse hydrolases.

#Possibly erroneous stop codons.

Anabolism. While the presence of catabolic enzymes is expected, the detection of putative anabolic enzymes is somewhat surprising for such a highly specialized parasite. Indeed, only a few were found. (i) One of them appears to be involved in amino acid biosynthesis: succinyl-diaminopimilate desuccinylase catalyses a step in the synthesis of diaminopimeline and lysine. (ii) Another anabolic enzyme is a phosphatidyl transferase involved in phospholipid synthesis. (iii) A third one is tetrahydrofolate (THF) synthase. THF and its derivatives are essential for numerous C1 unit (carbon) transfers that occur in a living

**Table 3.** Identification of rare restriction sites in the 214 kb of *M. capricolum*.

Enzyme	Recognition site	Total <sup>a</sup>	Observed
Fspl	TGCGCA	5	2
<i>Bg</i> /∣	GCCNNNNGCC	6	Õ
Apal	GGGCCC	2	1
<i>Bss</i> HII	GCGCGC	1	ò
Sal I	GTCGAC	2	2
Smal	cccggg	2	1
Xhol	CTCGAG	2	1
Total		20	7 (35%)

a. The restriction sites mapped onto the *M. capricolum* genome were taken from Whitely *et al.* (1991) and Miyata *et al.* (1991). *Bam*HI was omitted as reciprocal 2D-PFGE was used to map the sites relative to the *BgI*I site (see Whitely *et al.*, 1991 and refs therein). Extrapolating from the number of observed cutting sites (even by including *Bam*HI for which 1 out of 9 sites were found), a genome size of about 780 kb would result

cell. The last steps in THF biosynthesis involve the reduction of folate, a vitamin for many organisms including the majority of mycoplasmas (Finch and Mitchell, 1992). All three anabolic enzymes act at the level of metabolic intermediates, so the complete pathways may not be necessarily present.

DNA repair. Several DNA repair mechanisms exist, and the lack of at least some of them in *M. capricolum* has been claimed (for review see Labarere, 1992). We have found excinucleases and homologues of at least two proteins, RecM and RecN, that are thought to be involved in SOS DNA repair mechanisms (Alonso *et al.*, 1990; van Hoy and Hoch, 1990). RecN of *B. subtilis* and *E. coli* and their *M. capricolum* homologue show high similarity to a yeast chromosome segregation protein (Fig. 1). Indeed, internal coiled coil domains (data not shown) suggest a mechanochemical role of this family. Interestingly, the entire N-terminal ATP-binding domain of the family is similar to that of RecF, yet another *E. coli* protein involved in DNA repair (Gorbalenya and Koonin, 1990).

DNA replication. Although it was thought that Mycoplasma species have only one DNA polymerase (Razin, 1992; Labarere, 1992), ORFs with similarity to E. coli DNA polymerases I and III reveal the presence of at least two distinct enzymes (Table 2). Therefore, replication in M. capricolum may resemble the replication mechanisms of other eubacteria more closely than previously thought.

<sup>\*</sup>Putative frameshifts.

```
family:
                                                                                                                                                                                                                  h hhhG G GKoohh h
 Recf_Ecoli
                                                                       1 \  \, \text{SLTRLLIRD} \\ \textbf{FRNI}. \\ \textbf{ETADLALS}. \\ \textbf{PGFNFLV} \\ \textbf{GANGS} \\ \textbf{GK} \\ \textbf{TSVLE} \\ \textbf{A} \\ \textbf{IYTLGHGRAFRSLQIGRVIRHEQEAFVLHG} \\ \textbf{GANGS} \\ \textbf{GK} \\ \textbf{TSVLE} \\ \textbf{A} \\ \textbf{IYTLGHGRAFRSLQIGRVIRHEQEAFVLHG} \\ \textbf{GANGS} \\ \textbf{GK} \\ \textbf{GNGS} \\ \textbf{GK} \\ \textbf{GNGS} \\ 
Recf_Promi
                                                                        2 ILSRLLIRHFRNI.EQADLPLA.DGFNFLVGPNGSGKTSILEAIYTLGHGRAFRSAQANRVIQHDENAFILHG
Recf_Actpl
Recf_Bacsu
                                                                       2 PLSRLIINNFRNL.QSLDLELS.PNFNFIVGHNGSGKTSLLEAIFYLGHGRSFKSHISNRIIHYQAEDFVLHA
                                                                        2 YIQNLELTSYRNY.DHAELQFE.NKVNVIIGENAQGKTNLMEAIYVLSMAKSHRTSNDKELIRWDKDYAKIEG
Recn_Ecoli
                                                                       1 MLAQLTISNFAIV.RELEIDFH.SGMTVITGETGAGKSIAIDALGLCLGGRAE.....ADMVRTG
                                                                       1 MLAELSIKNFAII.EELTVSFE.RGLTVLTGETGAGKSIIIDAISLLVGGRGS......SEFVRYG
Recn Bacsu
 P115/Mychy
                                                                        {\tt 3~KLIKIEIEGFKSFADPISINFD.GSVVGIV\textbf{GPNGSGKSNIND}\textbf{A}IRWVLGEQSAKQLRGLNM.....DDVIFA\textbf{G}}
Z33332
                                                                                                      IRASGFKXFADLTVMDFN.YDMTGVVGPNGSGKSNITDAIRWTLGXQSTKTLRGSKM.....ADIVXSG
Smc1 Yeast
                                                                        3 RLVGLELSNFKSYRGVTKVGFGESNFTSIIGPNGSGKSNMMDAISFVLGVRS.NHLRSNIL....KDLIYRG
Nam1/Yeast
                                                                        ? YIKKVILRNFMCH.EHFELEG.SRLNFIVGNNGSGKSAILTAITIGLGAKASETNRGSSL.....KDLIREG
consensus:
                                                                                       h h h tF h t h htht th hhGttGtGKo hhtAh hh tttt
 В
 family:
Recn_Ecoli
                                                             394 IDVKFDEHHLGADG.ADRIEFRVTTNPGQPMQPI..SKVASGGELSRIALAIQVITARKMETPALIFDEVDVG
Recn_Bacsu
                                                             410 PLVNGQPVQLTEQG.IDLVKFLISTNTGEPLKSL..AKVASGGELSRVMLAIKSIFSSOODVTSIIFDEVDTG
P115/Mychy
                                                             839 KMFGGKAEIHFTDKNDILNSGVEISAQPPGKTIKNLRLFSGGEKAIIAISLLFAILKARPIPLCILDEVEAA
Z33355
                                                                                \texttt{KMFG} \textbf{\textit{G}} \textbf{\textit{G}}} \textbf{\textit{G}} \textbf{\textit
 Yat3_Rhoru
                                                                         ? KLFGGGRAHLTLIESDDPLEAGLEIMASPPGKRLQSLGLLSGGEQALTATALLFAVFLTNPAPICVLDEVDAP
{\tt Smc1\_Yeast~1090~Velaggnasltiededepfnagikyhatpplk} {\tt RfkDmeylsggektvarlallfainsyopspffvldevdaa}
consensus:
                                                                                                                               hh t t ht hth ttP Kth hthhSGGE hhAh h
 family:
                                                                                                                                                                                                             hhh H
Recn_Ecoli
                                                                                  ISGPTAAVVGKLLRQLGE..STQVMCVTHLPOVAGCGHOHYFVSKETDGAMTETHMOSLNKKAR
                                                                                                                                                                                                                                                                                                                                                                                                                                         P05824
 Recn_Bacsu
                                                                                  {\tt VSGRVAQAIAEKIHKVSI..GS{\color{red}Q}{\color{blue}VLCIT}{\color{blue}H}{\color{blue}LPQVAAMA}{\color{blue}D}{\color{blue}THLYIAKELKDGRTTTRVKPLSKQEK}
                                                                                                                                                                                                                                                                                                                                                                                                                                         P17894
 P115/Mychy
                                                                                  \verb|LDESNVIRYVEFLKLLKE..NTQFLIITHRSGTMSRVDQLLGVTMQKRG.VTSIFSVELSKAKE|
                                                                                                                                                                                                                                                                                                                                                                                                                                         M34956
 Z33355
                                                                                  LDPVNVERFARYVRHFSD..NTQFIIVTHREGTMTQCDSLFGVTMQTKG.ITKIINVKLVEAKN
                                                                                                                                                                                                                                                                                                                                                                                                                                         Z33355
 Yat3_Rhoru
                                                                                  LDDANVDRFCAMLRHLTDTTGTRFLVVTHHRMTMARMDRLFGVTMAERG. VSSLVSVDLCOAED
                                                                                                                                                                                                                                                                                                                                                                                                                                         P15016
 Smc1_Yeast
                                                                                  LDITNVQRIAAYIRRHRN.PDLQFIVISLKNTMFEKSDALVGVYRQQQENSSKIITLDLSNYAE
                                                                                                                                                                                                                                                                                                                                                                                                                                         P32908
 consensus:
                                                                                                          h hh hht t
                                                                                                                                                                                     t OhhhhoH t hh
                                                                                                                                                                                                                                                                               D hhhh t tt othh tL t tt
```

Fig. 1. Multiple alignment of two ORFs in contigs Z33332 and Z33355, with RecN and related proteins. Based on the strong similarity to the P115 proteins of other mycoplasmas, Z33332 and Z33355 might be fragments of the same protein. The role of P115 remains, however, to be elucidated, although the similarity to different repair proteins (RecN, RecF) is striking. SMC1 is involved in chromosome segregation. As in SMC1, RecN and Z33355, the N- and C-terminus is separated by coiled coil regions (cc) of different length, as predicted using the algorithm of Lupas et al. (1991). All the proteins shown in the alignments are members of a vast group of functionally diverse ATP-binding proteins (Gorbalenya and Koonin, 1990).

A. Alignment of the ORF in contig Z33332 with the N-terminal (ATP-binding) domain of the closest relatives.

B. Alignment of Z33355 with the closest relatives. The first line (family) indicates the consensus of the superfamily whereas the bottom line (consensus) summarizes conserved features of the alignment in the RecN subfamily: capitals, amino acids conserved in at least all but one sequence; h, hydrophobic position; o, S or T, negatively charged. An X within the alignment denotes an unidentified residue. The first column shows the SWISS-PROT codes (Bairoch and Boeckmann, 1993) of the respective proteins if available (underscore in protein name); the second row indicated the position of the aligned sequences in the respective proteins.

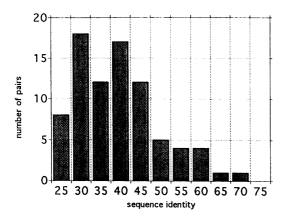
Metabolism of a parasite. The analysis further shows the presence of proteins involved in protein biosynthesis. translation and transcription regulation, protein folding and intracellular protein transport (Table 2). As expected for a parasite, a relatively large number of proteins is involved in the transport of intermediate metabolites and energy sources through the membrane, and their modification (first category, Table 2). As an even more surprising fact for a 'perfect' parasite, several carbohydrate- and proteindegrading enzymes have apparently been retained by M. capricolum to produce usable intermediates in metabolism (Table 2).

Quantitative sequence analysis: comparison with M. genitalium and E. coli

Quantitative sequence analysis can also reveal how closely related two organisms are, how protein function is conserved and which proteins appear to be most constrained in evolution. The high percentage of M. capricolum proteins with homologues of known function in other organisms supports the notion that mainly essential and conserved genes have been retained.

A set of contigs from M. genitalium that cover about 101 kb (Peterson et al., 1993) provide a good basis for species comparisons. Although M. capricolum and M. genitalium both belong to the Mollicutes, mycoplasmas are a polyphyletic group, separated by other species including those with intact cell walls (Weisburg et al., 1989). Indeed, the two data sets are complementary to each other with surprisingly little overlap: comparing our data with the contigs from M. genitalium, only 25 protein matches were observed, their sequence identity ranging between 40 and 80%. Because of the few and very short matches, however, quantification and generalization are difficult. Although it is still possible that all M. genitalium proteins are present in M. capricolum, the considerable variability among mycoplasmas is striking.

Numerous homologues were detected in E. coli. In order to quantify these similarities, all matches with E. coli



**Fig. 2.** Histogram of sequence identity of 82 *M. capricolum* proteins with their putative orthologues in *E. coli.* All hits above 50% amino acid identity are shown in Table 4.

proteins were studied in a first attempt to compare considerable fractions of two prokaryotic genomes. As many as 107 out of the 215 *M. capricolum* proteins functionally identified in this study have a homologue in *E. coli*. As it can only be estimated whether these are orthologues (the equivalent gene in another species) or paralogues (implying descent after gene duplication; Fitch, 1970), we have selected 95 *E. coli* sequences that can be readily aligned to their mycoplasma homologues above a certain similarity threshold (see the *Experimental procedures*). Lower scoring proteins, such as ABC transporters and members of the helix-turn-helix DNA repressor family, were excluded from the set of 95 homologous pairs. The similarity between the remaining 82 putative orthologues

from E. coli and M. capricolum ranges from 26% to 72% amino acid identity, with an average of 41% (Fig. 2). Most of the 'genetic' proteins (e.g. in translation, transcription, replication, repair and cell division) and those involved in protein biosynthesis are the most similar, the highest being the cell division protein FtsH with 72.3% amino acid identity over 83 residues (Table 4). Although metabolic enzymes in general have considerably lower similarity levels, enolase also appears highly conserved (Table 4). This conclusion is tentative, as the two short enolase fragments sequenced so far (ORFs in contigs Z33232 and Z33255) may be parts of the most conserved regions. A longer strongly conserved match (59% identity over 95 residues) was found for protease LA (Table 4). Although it is known that protease LA plays an important role in controlled degradation of short-lived or abnormal proteins (Suzuki et al., 1994), the precise reason for the constraints at the sequence level remains to be discovered.

Quantitative sequence analysis may also reveal unusual modes of protein evolution, i.e. contradictions between the generally accepted phylogenetic position of an organism and the evolutionary tree of a particular protein family. One of those cases is a putative *M. capricolum* prolyltRNA synthetase with high similarity to mammalian multifunctional aminoacyl-tRNA synthetases but only modest similarity to *E. coli* and other prolyl-tRNA synthetases (Fig. 3). One explanation for this anomaly would be that prokaryotic orthologues have not been sequenced yet. However, more than 60% of the *E. coli* genome is already stored in public databases and several other prokaryotic genome projects have also produced large amounts of

**Table 4.** Proteins or protein fragments most conserved between *E. coli* and *M. capricolum* (pairwise amino acid identities above 50%).

Contig	E. coli	DIST	OPT	%IDE	LEN	Description
Z33086 Z33255 Z33376 Z33013 Z33068 Z33345 Z33034 Z33034 Z33292 Z33370 Z33195 Z33330 Z33011	ftsh_ecoli eno_ecoli clpb_ecoli ychf_ecoli efg_ecoli uvrb_ecoli if1_ecoli lon_ecoli atpb_ecoli atpa_ecoli rl22_ecoli	47.5 36.8 38.4 36.9 36.7 35.2 32.5 34.1 32.3 29.8 24.1 27.0	284 175 400 228 608 980 240 263 917 197 161 195	72.3 65.6 63.2 63.0 61.5 60.0 59.4 58.9 57.1 56.3 52.4 51.8	83 61 125 73 200 305 69 95 345 71 63 85	Cell division protein Enolase clpb (stress protection) OBG-related protein Elongation factor EF-G Excinuclease subunit B Initiation factor IF-1 Cytoplasmic protease LA ATP synthase beta chain GTP-binding protein LepA ATP synthase alpha chain 50S ribosomal protein L22
Z33026 Z33011 Z33120	syn_ecoli rl16_ecoli uvra_ecoli	26.9 25.9 25.7	459 367 510	51.7 50.7 50.5	172 134 200	Asn-tRNA synthetase 50S ribosomal protein L16 Excinuclease subunit A

Contig: EMBL/GenBank accession numbers.

E. coli: SWISSPROT codes for E. coli proteins.

DIST: distance (in percentage points) to the length-dependent threshold of structural homology (25% for length 80 or more residues, higher for shorter alignments (Sander and Schneider, 1991).

OPT: FASTA 'optimized' scores (Pearson and Lipman, 1988).

%IDE: percentage of identical residues.

LEN: extent of the pairwise alignment number of amino acid residues.

support:	.* :
Syep_Human	YHDISGCYILRPWAYAIWEAIKDFFDAEIKKLGVENCYFPMFVSQSALEKEKTHVADFAPEVAWVTRSGKTELAEPIAIRPTSETVMYPAYAKWVQSHRDLPIKLNQWCN
-1-F	TERT WHAT THE ATROPP DAE TRACEVENCIPPMF VSQSALERERTHVADFAPEVAWVTRSGKTELAEPIAIRPTSETVMY PAYAKWVOSHRDLPIKINOWCN
DJ CPDI OMC	TID VOGCTION QUARTET INFORESTITATION DAESTITATION DE LA TRANSPORTATION D
Z33341	LWSVKGTMIFRPYGYRIWELIQKYLDEEFKKVNVDNVYFPLLIPESLFNKEKDHIDGFSPEIATVTRVGQKQLEENLFIRPTSEVLMMDYFSNEINSYRDLPLIYNQWCN
	ZHOYNOJIHI KI IGIKIWEDIQKI EDEBIKKVINVDINVIPPLLI PESLINKEKDHI DGFSPEIATVTRVGOKOLEENLI IRPTSEVI MMDYFSNEINSYRDLPI. IVNOMON
Syp_Ecoli	KLA.SGLYTWLPTGVRVLKKVENIVREEMNNAGAIEVSMPVVQPADLWQ.ESGRWEQYGPELLRFVDRGERPFVLGPTHEEVITDLIRNELSSYKQLPLNFYQIQT
Syt1_Bacsu	WYC OCI DIEN DYCAMIDDULEN CONTROL OF ADDRESS
Syci_Dacsu	KVG.QGLPLWLPKGATIRRVIERYIVDKEISLGYEHVYTPVLGSKELYE.TSG#WDHYQEGMFPPMEMDNETLVLRPMNCPHHMMIYKQDIHSYRELPIRIAELGT
consensus:	
	Gillian in I I am E aga ay Pah ha E Ha h PEha G haharpt hah ha syr Lph h oh

Fig. 3. Alignment of tRNA synthetases similar to an ORF in contig Z33341. The ORF in contig Z33341 is closer to the animal proteins than to other prokaryotic synthetases, which suggests the possibility that horizontal transfer has been involved in the acquisition of the gene for prolyl-tRNA synthetase. The first line shows positions that support grouping with the animal (+) or with the prokaryotic (-) sequences; characteristic insertions are denoted by an exclamation mark (!).

sequence data (for review, see Bork et al., 1994). Therefore, the chance of finding relatively well-characterized proteins which have not yet been described in other prokaryotes is relatively low. A tempting hypothesis for this surprising similarity would be the acquisition of the animal gene by horizontal gene transfer (the lateral transfer of genetic material across species). Although the acquisition of animal genes by bacteria is extremely rare (Heinemann, 1991; Doolittle and Bork, 1993), the transfer of genetic material between bacteria (e.g. via plasmids) appears to be a frequent event and it is known that parasites are able to acquire genes from their hosts (Heinemann, 1991). Interestingly, a similar phenomenon has been independently proposed for the other (N-terminal) part of the multifunctional aminoacyl-tRNA synthetase. This part of the protein codes for a Glu-tRNA synthetase which is more similar to Gln-tRNA than Glu-tRNA synthetase in E. coli (Lamour et al., 1994). It has been suggested that GIn-tRNA synthetase in various eubacteria has evolved from eukaryotic Glu-tRNA synthetase after having been acquired by horizontal gene transfer (Lamour et al., 1994). Other candidates for a horizontal transmission of genetic material in M. capricolum are U6 snRNA-like sequences (Ushida and Muto, 1993), which are involved in pre-mRNA splicing in eukaryotes, while no such phenomena have been reported for eubacteria.

#### Conclusion

The detection of at least 215 proteins in 214 kb DNA shows the power of the current sequencing and analysis approaches. The function of many proteins can be predicted by similarity, and biochemical pathways can be reconstructed. Quantitative sequence analysis also indicates an unorthodox similarity between some eukaryotic proteins and M. capricolum proteins. As mycoplasmas have been shown to have faster mutation rates than many other prokaryotes (Woese et al., 1984), the conservation profile of numerous proteins provides a wealth of data about functional and structural constraints at the protein sequence level, with proteins acting on DNA or involved in protein biosynthesis being the most constrained.

Comparison of complete genomes of prokaryotes such

as E. coli, as well as eukaryotes such as Saccharomyces cerevisiae and Caenorhabditis elegans, will soon enable us to follow in detail the evolution of genomes on the basis of sequence similarity. The study presented here is a first attempt to include M. capricolum in such systematic comparisons.

#### **Experimental procedures**

#### Sequencing of the contias

The sequences were obtained by the genomic walking technique (Gillevet, 1990), i.e. the direct sequencing of the organism by probing Southern blots of sequencing gels (Ohara et al., 1989). These blots contain the DNA of the entire organism that has been cut with restriction enzymes, treated with chemical sequencing reactions, run out on a sequencing gel and transferred to a charged nylon membrane. When such a membrane is probed with a labelled oligonucleotide, the resulting autoradiograph displays sequencing patterns in those lanes in which the oligonucleotide has hybridized near a restriction cut. About 1000 random clones were sequenced using fluorescent technology to supply starting points for the walking process (Gillevet, 1993). We also picked starting points from 64 segments taken from the EMBL nucleotide sequence database. The resulting 372 contigs have been deposited in the EMBL database (accession numbers Z33005 to Z33376).

## First-round sequence analysis

The computational and database analysis comprises the following steps.

- 1. The assembled consensus sequence from the contigs was subjected to BLASTX searches (Gish and States, 1993) using the mycoplasma genetic code in which UGA codes for tryptophan. BLASTX performs a rapid search of all six conceptual translations of the contig consensus against protein sequence databases.
- 2. In order to check for frameshifts and artificial stop codons within the contigs, BLASTX output was also automatically parsed (program FRAMESHIFT by G. Casari, unpublished) and critical regions extracted.
- 3. All possible ORFs longer than 10 amino acids were translated. The requirement for recording ORFs within the contigs was the presence of start and stop codons; in terminal fragments, only start (C-terminus) or stop codons (Nterminus) were required.
- 4. With the resulting 1845 putative ORFs, BLAST similarity

- searches (Altschul et al., 1990) against sequence databases were carried out.
- For ORFs longer than 30 amino acids without clear similarity, other search methods (Bork et al., 1992; Koonin et al., 1994) were applied and possible 'twilight zone' hits studied in detail.
- Finally, DNA sequence databases were screened for nonprotein coding elements such as RNA genes and internal repeats.
- 7. All results were merged (Table 2) and stored in a relational database (Scharf *et al.*, 1994) for further evaluation.

The last database search was done in January 1994. We realize that the increasing number of characterized proteins from other organisms is a valuable source of information (Bork *et al.*, 1994) and the searches will be repeated in updated databases with a more complete *M. capricolum* data set that we are currently obtaining.

#### Quantitative comparison

For verifying the significance of subtle similarities or detecting the possibility of horizontal gene transfers, multiple alignments and phylogenetic trees (as implemented in CLUSTALW, a new version of CLUSTALV; Higgins *et al.*, 1992) were used.

The comparison with *E. coli* and *M. genitalium* proteins involved the following steps: (i) extracting all *E. coli* and *M. genitalium* proteins from sequence databases; (ii)• allagainst-all comparison with the mycoplasma contigs using TFASTA (Pearson and Lipman, 1988); (iii) extraction of putative orthologues; and (iv) ranking the hits according to several scoring schemes such as FASTA 'opt' scores (Pearson and Lipman, 1988), amino acid identity and distance to a length-dependent threshold for structural similarity (Sander and Schneider, 1991).

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