The genome sequence of *Rickettsia* prowazekii and the origin of mitochondria

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We describe here the complete genome sequence (1,111,523 base pairs) of the obligate intracellular parasite *Rickettsia prowazekii*, the causative agent of epidemic typhus. This genome contains 834 protein-coding genes. The functional profiles of these genes show similarities to those of mitochondrial genes: no genes required for anaerobic glycolysis are found in either *R. prowazekii* or mitochondrial genomes, but a complete set of genes encoding components of the tricarboxylic acid cycle and the respiratory-chain complex is found in *R. prowazekii*. In effect, ATP production in *Rickettsia* is the same as that in mitochondria. Many genes involved in the biosynthesis and regulation of biosynthesis of amino acids and nucleosides in free-living bacteria are absent from *R. prowazekii* and mitochondria. Such genes seem to have been replaced by homologues in the nuclear (host) genome. The *R. prowazekii* genome contains the highest proportion of non-coding DNA (24%) detected so far in a microbial genome. Such non-coding sequences may be degraded remnants of 'neutralized' genes that await elimination from the genome. Phylogenetic analyses indicate that *R. prowazekii* is more closely related to mitochondria than is any other microbe studied so far.

The *Rickettsia* are α -proteobacteria that multiply in eukaryotic cells only. R. prowazekii is the agent of epidemic, louse-borne typhus in humans. Three features of this endocellular parasite deserve our attention. First, R. prowazekii is estimated to have infected 20-30 million humans in the wake of the First World War and killed another few million following the Second World War (ref. 1). Because it is the descendent of free-living organisms²⁻⁴, its genome provides insight into adaptations to the obligate intracellular lifestyle, with probable practical value. Second, phylogenetic analyses based on sequences of ribosomal RNA and heat-shock proteins indicate that mitochondria may be derived from the αproteobacteria^{5,6}. Indeed, the closest extant relatives of the ancestor to mitochondria seem to be the Rickettsia⁷⁻¹⁰. That modern Rickettsia favour an intracellular lifestyle identifies these bacteria as the sort of organism that might have initiated the endosymbiotic scenario leading to modern mitochondria¹¹. Finally, the genome of R. prowazekii is a small one, containing only 1,111,523 base pairs (bp). Its phylogenetic placement and many other characteristics identify it as a descendant of bacteria with substantially larger genomes^{2–4}. Thus *Rickettsia*, like mitochondria, are good examples of highly derived genomes, the products of several types of reductive

The genome sequence of *R. prowazekii* indicates that these three features may be related. For example, prokaryotic genomes evolving within a cell dominated by a much larger, eukaryote genome and constrained by bottle-necked population dynamics will tend to lose genetic information 12,13 . Predictable sets of expendable genes will tend to disappear from the prokaryotic genome when they are made redundant by the activities of nuclear genes. Likewise, non-essential sequences and otherwise highly conserved gene clusters may be obliterated by deleterious mutations that are fixed in clonal parasite or organelle populations because they cannot be eliminated by selection. This process is ongoing in the *Rickettsia* genomes, as shown by the identification of sequences that have recently become pseudogenes. Also, a large fraction (\sim 25%) of non-coding sequences in this genome may be gene remnants that have been

degraded by mutation and have not yet been removed from the genome. Finally, transfer of genes from a mitochondrial ancestor to the nucleus of the host would both reduce the mitochondrial genome size and stabilize the symbiotic relationship. Phylogenetic reconstructions that identify genes in the *Rickettsia* genome as sister clades to eukaryotic homologues found in the nucleus or the organelle support this interpretation. *Rickettsia* and mitochondria probably share an α -proteobacterial ancestor and a similar evolutionary history.

General features of the genome

The circular chromosome of R. prowazekii strain Madrid E has 1,111,523 bp and an average G+C content of 29.1% (Figs 1, 2). The genome contains 834 complete open reading frames with an average length of 1,005 bp. Protein-coding genes represent 75.4% of the genome and 0.6% of the genome encodes stable RNA. We have assigned biological roles to 62.7% of the identified genes and pseudogenes; 12.5% of the identified genes match hypothetical coding sequences of unknown function and the remaining 24.8% represent unusual genes with no similarities to genes in other organisms (Table 1). Multivariate statistical analysis has shown that there is no major variation in codon-usage patterns among genes that are expressed in different amounts, indicating that codon-usage patterns in R. prowazekii may be dominated mainly by mutational forces¹⁴. G+C-content values at the three codon positions average 40.4, 31.2 and 18.6%, and these values are similar at different positions in the genome. We classified the open reading frames with significant sequence-similarity scores to gene sequences in the public databases into functional categories (Table 1) that allow comparisons with the metabolic profiles of other bacterial genomes¹⁵⁻²³.

Non-coding DNA. The coding content of previously sequenced bacterial genomes is, on average, 91%, ranging from 87% in *Haemophilus influenzae* to 94% in *Aquifex aeolicum*. In comparison, a large fraction of the *R. prowazekii* genome, 24%, represents non-coding DNA (Fig. 3). A small fraction of this corresponds to

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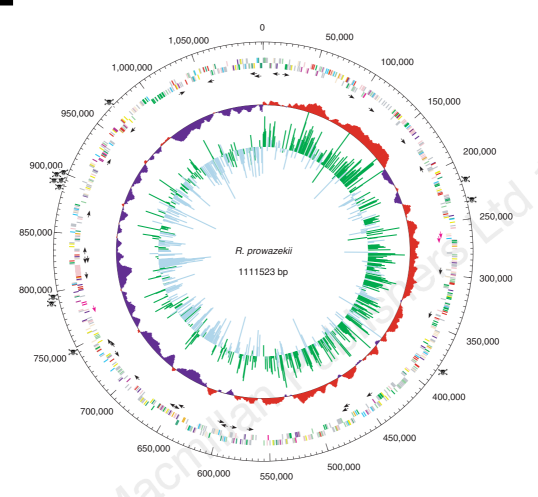


Figure 1 Overall structure of the *R. prowazekii* genome. The putative origin of replication is at 0 kb. The outer scale indicates the coordinates (in base pairs). The positions of pseudogenes are highlighted with death's heads. The distribution of genes is shown on the first two rings within the scale. The location and direction of transcription of rRNA are shown by pink arrows and of tRNA genes by black arrows. The next circle in shows GC-skew values measured over all bases in the genome. Red and purple colours denote positive and negative signs, respec-

tively. The window size was 10,000 nucleotides and the step size was 1,000 nucleotides. The central circles shows GC-skew values calculated for third positions in the codon only. GC-skew values were calculated separately for genes located on the outer strand (green) and on the inner strand (blue). To allow easier visual inspection, the signs of the values calculated for genes located on the inner strand have been reversed.

pseudogenes (0.9% of the genome) and less than 0.2% of the genome is accounted for by non-coding repeats. The remaining 22.9% contains no open reading frames of significant length and it has the low G+C content (mean 23.7%) that is characteristic of spacer sequences in the *R. prowazekii* genome¹⁴. A region of 30 kilobases (kb) located at position 886–916 kb contains as much as 41.6% non-coding DNA and 11.5% pseudogenes. The non-coding DNA in this region has a small, but significantly higher, G+C content (mean 27.3%) than non-coding DNA in other areas of the genome (mean 23.7%) (P < 0.001), indicating that it may correspond to inactivated genes that are being degraded by mutation (Fig. 3).

Origin of replication. The origin of replication has not been experimentally identified in the *R. prowazekii* genome, but we identified *dnaA* at ∼750 kb. However, the genes flanking the *dnaA* gene differ from the conserved motifs found in *Escherichia coli* and *Bacillus subtilis* (*rnpA−rpmH−dnaA−dnaN−recF−gyrB*). In *R. prowazekii*, the genes *rnpA* and *rpmH* are located in the vicinity of *dnaA*, but in the reverse orientation compared to the consensus motif, and *dnaN*, *recF* and *gyrB* are located elsewhere.

The origin and end replication in microbial genomes are often associated with transitions in GC skew (G - C/G + C) values²⁴. In R. prowazekii we observe transitions in the GC skew values at

around 0 and 500–600 kb (Fig. 1). There is a weak asymmetry in the distribution of genes in the two strands, such that the first half of the genome has a 1.6-fold higher gene density on one strand and the second half of the genome has a 1.6-fold higher gene density on the other strand. The shift in coding-strand bias correlates with the shift in GC-skew values. As most genes are transcribed in the direction of replication in microbial genomes, the origin of replication may correspond to the shift in GC-skew values at the position that we have chosen as the start point for numbering. Indeed, several short sequence stretches that are characteristic of *dnaA*-binding motifs are found in the intergenic region of genes *RP001* and *RP885* at 0 kb, supporting this interpretation.

Stable RNA sequences and repeat elements. We identified 33 genes encoding transfer RNA, corresponding to 32 different isoacceptortRNA species. There is a single copy of each of the rRNA genes, with *rrs* located more than 500 kb away from the *rrl-rrf* gene cluster

Figure 2 Linear map of the *R. prowazekii* chromosome. The position and **J** orientation of known genes are indicated by arrows. Coding regions are colour-coded according to their functional roles. The positions of tRNA genes are indicated (inverted triangle on stalk). For additional information, see http://evolution.bmc.uu.se/~siv/gnomics/Rickettsia.html.

(Fig. 1). Comparison of the sequences from ten different *Rickettsia* species indicates that the disruption of the rRNA gene operon preceded the divergence of the typhus group and spotted fever group *Rickettisia* (S.G.E.A. *et al.*, unpublished observations). In addition, the genome contains a short sequence with similarity to a 213-nucleotide RNA molecule in *Bradyrhizobium japonicum* that may regulate transcription²⁵.

There are unusually few repeat sequences in this genome. We identified four different types of repeat sequence: all of these are located in intergenic regions. There is a sequence of 80 bp that is repeated seven times downstream of *rpmH* and *rnpA* in the *dnaA* region. A repetitive sequence of 325 bp is found at two intergenic regions that are more than 80 kb apart, downstream of the genes *ksgA* and *rnh*, respectively. A 440-bp-long repetitive sequence has been identified at two intergenic sites, 140 kb apart; one of these sites is downstream of *rrf* and the others downstream of *pdhA* and

Table 1 Asterisks indicate putative pseudogenes. Abbreviations of species names are: Bacteria: Acinetobacter calcoaceticus (B-Aca), Actonobacillus actinomycetemcomitans (B-Aac), Acyrthosiphon condii (B-Aco), Agrobacterium tumefaciens (B-Atu), Alcaligenes eutrophus (B-Aeu), Anabena sp. PCC7120 (B-Asp), Anabena variabilis (B-Ava), Anacystis nidulans (B-Ani), Azorhizobium caulinodans (B-Aca), Azosprillium brasiliense (B-Abr), Azotobacter vinelandii (B-Avi), Bacillus caldotenax (B-Bca), Bacillus stereothermophilus (B-Bst), Bacillus subtilis (B-Bsu), Bartonella bacilliformis (B-Bba), Bartonella henselae (B-Bhe), Bordetella pertussis (B-Bpe), Borrelia burgdorferi (B-Bbu), Bradyrhizobium japonicum (B-Bja), Brucella abortus (B-Bab), Brucella ovis (B-Bov), Caulobacter crescentus (B-Ccr), Chlamydia trachomatis (B-Ctr), Chloroflexus aurantiacus (B-Cau), Chromatum viosum (B-Cvi), citrus-greening-disease-associated bacterium (B-Cgr), Clostridium acetobutylicum (B-Cac), Clostridium pasteurianum (B-Cpa), Clostridium thermosaccharolyticum (B-Cts), Coxiella burnetii (B-Cbu), Erwinia chrysanthemi (B-Ech), Escherichia coli (B-Eco), Haemophilus influenzae (B-Hin), Helicobacter pylori (B-Hpy), Klebsiella pneumoniae (B-Kpn), Legionella pneumophila (B-Lpn), Leucothrix mucor (B-Lmu), Liberobacter africanum (B-Laf), Methylobacterium extorquens (B-Mex), Micrococcus luteus (Mlu), Moraxella catarrhalis (Mca), Mycobacterium leprae (Mle), Mycobacterium smegmatis (B-Msm), Mycobacterium tuberculosis (B-Mtu), Mycoplasma capricolum (B-Mca), Mycoplasma genitalium (B-Mge), Mycoplasma pneumoniae (B-Mpn), Paracoccus denitrificans (B-Pde), Pasteurella haemolytica (B-Pha), Plectonema boryanum (B-Pbo), Proteus mirabilis (B-Pmi), Proteus vulgaris (BPvu), Pseudomonas aeruginosa (B-Pae), Pseudomonas fluorescens (B-Pfl), Pseudomonas putida (B-Ppu), Pseudomonas syringae (B-Psy), Rhizobium meliloti (B-Rme), Rhizobium sp. NGR234 (B-Rsp), Rhodobacter capsulatus (B-Rca), Rhodobacter sphaeroides (B-Rsp), Rhodobacter sulfidophilus (B-Rsu), Rhodopseudomonas blastica (B-Rbl), Rhodospirillum rubrum (B-Rru), Rickettsia japonicum (B-Rja), Rickettsia rickettsii (B-Rri), Rickettsia typhi (B-Rty), Salmonella typhi (B-Sti), Salmonella typhimurium (B-Sty), Shigella flexneri (B-Sfl), Sprioplasma citri (B-Sci), Staphylococcus aureus (B-Sau), Staphyloccus carnosus (B-Sca), Streptococcus pneumoniae (B-Spn), Streptomyces clavuligerus (B-ScI), Streptomyces coelicor (B-Sco), Synechocystis PCC 6803 (B-Syn), Thermus aquaticus (B-Taq), Thermus thermophilus (B-Tth), Thiobacillus cuprinus (B-Tcu), Treponema hyodysenteriae (B-Thy), Vibrio alginolytics (B-Val), Vibrio cholera (B-Vch), Vibrio parahaemolyticus (B-Vpa), Vibrio proteolyticus (B-Vpr), Wolbachia sp. (B-Wsp), Yersinia entercolitica (B-Yen), Zooglea ramigera (B-Zra), Zymomonas mobilis (B-Zmo). Archaea: Methanococcus jannaschii (A-Mja), Sulfulobus acidocaldarius (A-Sac). Eukaryotes: Apis mellifera (E-Ame), Arabidopsis thaliana (E-Ath), Atratylodes japonica (E-Aja), Bos taurus (E-Bta), Candida albicans (E-Cal), Caenorhabidits elegans (E-Cel), Dictylostelium discoideum (E-Ddi), Flaveria trinervia (E-Ftr), Giardia theta (E-Gth), Glycine max (E-Gma), Haematobia irritans (E-Hir), Homo sapiens (E-Hsa), Marchantia polymorpha (E-Mpa), Mus musculum (E-Mmu), Prototheca wickerhamii (E-Pwi), Petunia hybrida (E-Phy), Pisum sativum (E-Psa), Porphyra purpurea (E-Ppu), Odontella sinensis (E-Osi), Reclinomonas americana (E-Ram), Rattus novergicus (E-Rno), Rhizopus oryzae (E-Ror), Saccharomyces cerevisiae Schizosaccharomyces pombe (E-Spo), Solanum tuberosum (E-Stu), Spinacia oleracea (E-Sol).

pdhB. Finally, two similar sequences of 730 bp are located immediately next to each other at 850 kb.

Paralogous families. We have identified 54 paralogous gene families comprising 147 gene products. Of these, 125 have an assigned function. Most paralogues encode proteins with transport functions, such as the ABC transporters, the proline/betaine transporters and the ATP/ADP transporters. Five paralogous genes located next to each other at 115 kb encode putative integral membrane proteins with unknown functions.

Biosynthetic pathways

A striking feature of the *R. prowazekii* genome is the small proportion of biosynthetic genes compared with free-living proteobacterial relatives (such as *Haemophilus influenzae*, *Helicobacter pylori* and *E. coli*) 15,19,20 . This scarcity of biosynthetic functions is also seen in diverse endocellular and epicellular parasites $^{16-18,23}$. This scarcity of biosynthetic functions is also seen in diverse endocellular and epicellular parasites $^{16-18,23}$.

Amino-acid metabolism. As many as 43 and 69 genes required for amino-acid biosynthesis are found in *Helicobacter pylori* and *Haemophilus influenzae*, respectively. In contrast, *Mycoplasma genitalium* and *Borrelia burgdorferi* contain only *glyA*, which encodes serine hydroxymethyltransferase. This gene is also found in *R. prowazekii* (Table 1). Serine hydroxymethyltransferase catalyses the conversion of serine and tetrahydrofolate into glycine and methylenetetrahydrofolate, respectively. A role in tetrahydrofolate metabolism may account for the ubiquity of *glyA* in bacteria.

Seven genes normally associated with lysine biosynthesis (*lysC*, *asd*, *dapA*, *dapB*, *dapD*, *dapE* and *dapF*) are also present in *R. prowazekii*. The biosynthetic pathways leading to lysine, methionine and threonine share the first two of these (*lysC* and *asd*). However, none of the downstream genes for threonine biosynthesis are found in *R. prowazekii*. Likewise, the lysine pathway is incomplete, and *lysA*, which encodes the enzyme that converts meso-diaminopimelate to lysine, is missing. The likely role of the upstream genes of this pathway in *R. prowazekii* is the biosynthesis of diaminopimelate, an essential envelope component. We have therefore classified these genes as 'cell-envelope' genes (Table 1).

We have identified other genes that are superficially involved in the metabolism of amino acids, but which apparently function in deamination pathways that divert amino acids into the tricarboxylic acid (TCA) cycle. For example, there is *aatA*, encoding aspartate aminotransferase, which catalyses the degradation of aspartate to oxaloacetate and glutamate. *tdcB* encodes threonine deaminase, which converts threonine into α-ketobutyrate. Another gene (*ilvE*) encodes branched-chain-amino-acid aminotransferase, which converts leucine, isoleucine or valine into glutamate. *pccA* and *pccB* encode propionyl-CoA carboxylase, which converts propionyl-CoA, an intermediate in the breakdown of methionine, valine and isoleucine, into succinyl-CoA. The *pccA* and *pccB* gene products show greatest similarity to the eukaryotic proteins that are located in the mitochondrial matrix.

Nucleotide biosynthesis. No genes required for the *de novo* syntheses of nucleosides have been found in the *R. prowazekii* genome. However, four genes required for the conversion of nucleoside monophosphates into nucleoside diphosphates (*adk*, *gmk*, *cmk* and *pyrH*) are present. There are also two genes encoding ribonucleotide reductase, which converts ribonucleoside diphosphates into deoxyribonucleoside diphosphates. Nucleoside diphosphate kinase (encoded by *ndk*), which converts NDPs and dNDPs to NTPs and dNTPs, is also present in *R. prowazekii*. Finally, there is a complete set of genes for the conversion of dCTP and dUTP into TTP, including *thyA*, which codes for thymidylate synthase. Thus, the *R. prowazekii* genome encodes all of the enzymes required for the interconversion of nucleoside monophosphates into all of the other required nucleotides. The nucleoside monophosphates are probably imported from the eukaryotic host.



Energy metabolism

Early in its infectious cycle, *R. prowazekii* uses the ATP of the host with the help of membrane-bound ATP/ADP translocases. However, *R. prowazekii* is also capable of generating ATP, which may compensate for the gradual depletion of cytosolic ATP later in the infection. *R. prowazekii*'s repertoire of genes involved in ATP production and transport include determinants for the TCA cycle, the respiratory-chain complexes, the ATP-synthase complexes and the ATP/ADP translocases (Table 1). Genes to support anaerobic glycolysis are absent.

Pyruvate dehydrogenase. Pyruvate is imported into mitochondria directly from the cytoplasm and converted into acetyl-CoA by pyruvate dehydrogenase. The genes encoding three components (E1–E3) of the pyruvate dehydrogenase complex are found in *R. prowazekii*, indicating that it too uses cytosolic pyruvate. Pyruvate dehydrogenase (E1) consists of two subunits (α and β) in *R. prowazekii*, mitochondria and Gram-positive bacteria; the corresponding genes are clustered in the genome. In contrast, proteobacteria such as *E. coli*, *Haemophilus influenzae* and *Helicobacter pylori* have a single subunit for the E1 component and these have little similarity to the α and β subunits of the E1 component in *R. prowazekii* and mitochondria (data not shown).

Two paralagous genes code for the dihydrolipoamide dehydrogenase (E3) in R. prowazekii. One of these most resembles mitochondrial homologues, whereas the other is most similar to bacterial homologues (data not shown). The presence of several paralogous gene families for pyruvate dehydrogenases complicates attempts to reconstruct a genome phylogeny based on these genes. **ATP production.** Genes encoding all enzymes in the TCA cycle are found in R. prowazekii. Proton translocation is mediated by NADH dehydrogenase (complex I), cytochrome reductase (complex III) and cytochrome oxidase (complex IV). Several clusters of genes code for components of the NADH dehydrogenase complex. Seven of these genes (nuoJKLM and nuoGHO) are located near to each other, but the order of genes is inverted relative to the order of this cluster in E. coli. An additional set of five genes is grouped in the order nuoABCDE, but the single genes nuoF and nuoN are distant from both of these clusters. Several proteins in the cytochrome bc_1 reductase complex, such as ubiquinol-cytochrome c reductase

(encoded by petA), cytochrome b (encoded by cytb) and cytochrome c_1 (encoded by fbhC), are present, as are several subunits of the cytochrome oxidase complex.

The ATP-synthesizing complex is composed of the ATP synthase F_1 component (comprising five polypeptides, α , β , γ , ϵ and δ) and the F_0 component, a hydrophobic segment that spans the inner mitochondrial membrane. The genes encoding these components are normally clustered in one of the most highly conserved operon structures in microbial genomes. In *R. prowazekii*, however, the ATP-synthase genes encoding the α , β , γ , δ and ϵ subunits of the F_1 complex (atpH, atpA, atpG, atpD and atpC) are clustered in the common order, but atpB, atpE and atpF, encoding the A, B and C chains of the F_0 complex, are split from this cluster.

Replication, repair and recombination

R. prowazekii has a smaller set of genes involved in DNA replication than do free-living bacteria such as *E. coli*, *Haemophilus influenzae* and *Heliocobacter pyrlori*. Four genes have been identified that code for the core structure of DNA polymerase III, which includes the α (dnaE), ϵ (dnaQ), β (dnaN), γ and θ (dnaX) subunits. Extra subunits present in the *E. coli* DNA polymerase III are missing from *R. prowazekii*, as well as from *M. genitalium* and *B. burgdorferi*.

Genes encoding DNA-repair mechanisms are similar in the small genomes of the parasites *R. prowazekii*, *M. genitalium* and *B. burgdorferi*. Thus, genes involved in the repair of ultravioletinduced DNA damage (*uvrABCD*) have been identified in all three genomes. In *R. prowazekii*, DNA-excision repair probably occurs by a pathway involving endonuclease III, polI and DNA ligase, as in *B. burgdorferi*.

The *R. prowazekii* genome has a limited capacity for mismatch repair. The DNA-mismatch-repair enzymes encoded by *mutL* and *mutS* are present, but *mutH* and *mutY* are not. There is a complete lack of *mut* genes in *M. genitalium*, but *mutL* and *mutHLY* have been identified in *B. burgdorferi* and *Chlamydia trachomatis*. The transcription-repair coupling factor (encoded by *mfD*) is found in *R. prowazekii*, *B. burgdorferi* and *C. trachomatis* but not in *M. genitalium*.

The *R. prowazekii* genome contains several genes involved in homologous recombination, such as *recA*, *recF*, *recJ*, *recN* and *recR*. A similar set of genes has been found in *A. aeolicus*²¹. The *rec* genes

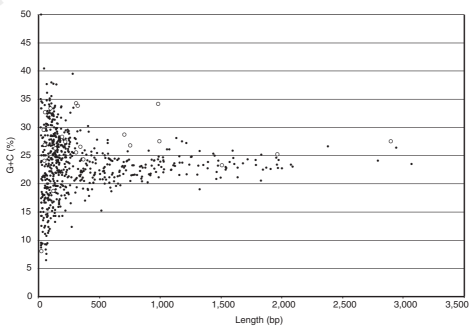


Figure 3 G+C content in intergenic regions longer than 20 bp in the *R. prowazekii* genome. The empty circles correspond to spacer sequences located at 886 to

916kb, a region with an unusually large fraction of non-coding DNA and pseudogenes.

136

are scattered in the other small genomes of parasites. The *RecBCD* complex is missing in *R. prowazekii*, *M. genitalium* and *Helicobacter pylori* but it has been identified in *B. burgdorferi*.

Transcription and translation

R. prowazekii has three subunits (α , β and β') of the core RNA polymerase, as well as σ^{70} and one alternative σ factor, σ^{32} , which controls transcription of the genes encoding heat-shock proteins in *E. coli*. Genes involved in transcription elongation and termination, *nusA*, *nusB*, *nusG*, *greA* and *rho*, are also present. The gene encoding σ^{32} is absent in most other small genomes, such as those of *B. burgdorferi*, *Helicobacter pylori*, *M. genitalium* and *C. trachomatis*, although genes for heat-shock proteins are present.

An unusually large number of genes involved in RNA degradation are found in *R. prowazekii*. Of these, only four appear to be common to the bacterial genomes analysed so far (those encoding polyribonucleotide nucleotidyltransferase and ribonucleases HII, III and P). Four more ribonucleases (D, E, HI and PH) are present in *R. prowazekii*, but in none of the other small parasites.

Of the 33 identified tRNA genes, which code for 32 different tRNA isoacceptor species, two code for tRNA Phe. There are two tRNA species for most of the amino acids that are encoded by four-codon boxes; the exceptions are the four-codon boxes for proline and valine, for which we have identified only one isoacceptor-tRNA species, with U in the first anticodon position. selC, which codes for tRNA Sec, and selABD are missing. R. prowazekii has a set of genes coding for tRNA modifications (tgt, queA, trmD, truA, truB and miaA) which resembles that of Helicobacter pylori, C. trachomatis and B. burgdorferi; M. genitalium has only trmD and truA.

In R. prowazekii, 21 genes encode 18 of the 20 aminoacyl-tRNA synthetases normally required for protein synthesis. There are two genes (gltX) encoding glutamyl-tRNA synthetase. As seen in several bacterial genomes²⁵, the gene coding for glutaminyl-tRNA synthetase, glnS, is missing. Three genes encoding subunits of the glutamyl-tRNA amidotransferase are present, indicating that a glutamyl-tRNA charged with glutamic acid may be transamidated to generate Gln-tRNA. The gene coding for asparaginyl-tRNA synthetase, asnS, is also missing from the R. prowazekii genome as well as from Helicobacter pylori, C. trachomatis and A. aeolicus²⁶. A transamidation process to form Asn-tRNA^{Asn} from Asp-tRNA^{Asn} has been proposed for the archaeon Haloferax volcanii²⁷ and this reaction may also occur in R. prowazekii. The valyl-tRNA synthetase is 38.3% identical to its homologue in Methanococcus jannaschii, but only 27.6% identical to its most similar homologue in bacteria, which is found in Bacillus stearothermophilus, possibly indicating a horizontal transfer event. The lysyl-tRNA synthetase (encoded by lysS) in R. prowazekii is a class I enzyme with no resemblance to the conventional class II lysyl-tRNA synthetases. Class I type of lysyltRNA synthetases have been observed previously in only B. burgdorferi, Pyrococcus woesii, Methanococcus jannaschii and a few other methanogens²⁶.

Regulatory systems

As in other genomes of small parasites, *R. prowazekii* has a reduced set of regulatory genes. There are a few members of two-component regulatory systems, such as the proteins encoded by *barA*, *envZ*, *ntrY*, *ntrX*, *ompR* and *phoR*. *spoT*, which is involved in the stringent response, has been identified in *B. burgdorferi*, *Helicobacter pylori* and *M. genitalium*. Only remnants of genes coding for aminoterminal fragments of proteins similar to those encoded by *spoT* and *relA* are identifiable in *R. prowazekii*. No fragments of *spoT* encoding the carboxy-terminal segments of the protein have been identified in the genome.

Cell division and protein secretion

Proteins involved in detoxification, such as superoxide dismutase, and those involved in thiophen and furan oxidation are present in *R*.

prowazekii. Two genes encoding haemolysins have also been identified, and an *R. typhi* homologue of *tlyC* exhibits haemolytic activities when expressed in *E. coli* (S. Radulovic, J. M. Troyer, B. Noden, S.G.E.A. and A. Azad, unpublished observations).

The data indicate that the basic mechanisms of cell division and secretion in *R. prowazekii* are similar to those in free-living proteobacteria. There is a common set of bacterial chaperones (encoded by *dnaK*, *dnaJ*, *hslU*, *hslV*, *groEL*, *groES* and *htpG*) and genes involved in the *secA*-dependent secretory system (*secABDEFGY*, *ffH* and *ftsY*). *R. prowazekii* has a significantly larger set of genes involved in peptide secretion than does *M. genitalium*.

Membrane-protein analysis

Many studies of *R. prowazekii* have focused on outer-surface membrane proteins because of their potential importance in bacterial detection and vaccination. The superficial lipopolysaccharide (LPS) molecule is important in the pathogenesis of *R. prowazekii*. LPS consists of a polysaccharide that is covalently linked to lipid A, the biosynthesis of which is catalysed by products of *lpxABCD*, all of which are present in the *R. prowazekii* genome. These genes are clustered in *E. coli*, but *lpxA* and *lpxD* are separate from *lpxB* and *lpxC* in *R. prowazekii*. Three genes involved in the biosynthesis of the 3-deoxy-D-manno-octulosonic acid (KDO) residues reside in the *R. prowazekii* genome (*kdsA*, *kdsB* and *kdtA*). Only one gene (*rfaJ*) with a putative function in outer-core biosynthesis has been identified.

We have identified a set of genes involved in the biosynthesis of murein and diaminopimelate and a set involved in the biosynthesis of fatty acids. These includes: fabD, which is involved in the last step of the initiation phase of fatty-acid biosynthesis; four genes involved in the elongation cycle of fatty-acid biosynthesis (fabFGHI); and three genes involved in the first three steps of the synthesis of polar head groups (cdsA, pssA and pgsA). Finally, post-translational processing and addition of lipids to an N-terminal cysteine require the gene products prolipoprotein diacylglycerol transferase (lgt), prolipoprotein signal peptidase (lspA) and apolipoprotein:phosholipid N-acyl transferase (lnt). These are found in the genome with several genes involved in the degradation of fatty acids, such as fadA which encodes the 3-ketoacyl-CoA thiolase.

Virulence

The R. prowazekii genome contains several homologues of the VirB gene operon found in Agrobacterium tumefaciens. This gene family encodes proteins that direct the export of the T-DNA-protein complex across the bacterial envelope to the plant nuclei²⁸. R. prowazekii has two homologues of VirB4 and one homologue each of VirB8, VirB9, VirB10, VirB11 and VirD4. The latter five genes are clustered with the gene trbG, which is involved in conjugation in Agrobactrium tumefaciens. Homologues of the single-stranded DNA-binding proteins VirD2 and VirE2 are missing. In Agrobacterium tumefaciens, these proteins are bound to the transferred T-DNA, indicating different functions for the homologues of the VirB genes in R. prowazekii. Indeed, VirB proteins are homologous to components of the E. coli transport system for plasmids, as well as to components of the Pt1 transport machinery in Bordetella pertussis, which exports pertussis toxin²⁸. A set of genes coding for VirB4 and several other VirB proteins has been identified in the cag pathogenicity island of Helicobacter pylori. In this species, the VirB proteins facilitate export of a factor that induces interleukin-8 secretion in gastric epithelial cells²⁸. Thus, R. prowazekii may encode components of a transport system for both conjugal DNA transfer and protein export.

The virulence of *Staphylococcus aureus* has been correlated with the production of capsular polysaccharides in phagocytic assays and mouse lethality assays^{29,30}. A cluster of ten capsule genes (capA-M) is involved in capsule biosynthesis in *S. aureus* strain M^{31} . We have identified three *R. prowazekii* genes with sequence similarities to *S. aureus cap* genes. Two of these (capD and capM) are separated by ten



genes, most of which are unknown genes or genes involved in the biosynthesis of LPS or techoic acid. Thus, *R. prowazekii* may produce components of a microcapsular layer that is involved in virulence.

Reductive evolution

Genome sequences of organisms enjoying an endosymbiotic lifestyle are at risk. The activities of homologous nuclear genes may render genes of the endosymbiont expendable and as a consequence they become vulnerable to obliteration by mutation. Good candidates for such purged genes in *Rickettsia* and mitochondria are genes required for amino-acid biosynthesis, nucleoside biosynthesis and anaerobic glycolyis. These and other genes would have been deleted when an ancestral genome first lived in a nucleated cell. Once genes essential to a free-living mode are lost, the endosymbiont becomes an obligate resident of its host.

Likewise, small, bottle-necked populations of bacteria infecting a eukaryotic cell will tend to accumulate deleterious mutations because selection cannot remove them from such clonal populations¹³. The accumulation of such harmful but non-lethal mutations is referred to as 'Muller's ratchet'³² or 'near-neutral evolution'^{33,34}. The consequence of accumulation of these mutations will be the inactivation and eventual deletion of non-essential genes.

The first mutation that inactivates an expendable gene is likely to initiate a sequence of events in which subsequent mutations freely transform it, by degrees, from a pseudogene, to unrecognizable sequence, to small fragments, to extinction. In this sequence, mutations are released from amino-acid-coding constraints. Thus nucleotide substitutions will reflect the mutation bias of the genome. This bias can be estimated roughly by frequencies of third-position bases in the codons. For R. prowazekii, the bias of the third-position bases is 18% G+C rather than the 29% G+C average for the genome. So, as sequences age in R. prowazekii, their composition should gradually approach the low G+C content of third codon positions. Nearly one-quarter of the R. prowazekii genome is composed of non-coding sequences, with a G+C content lower than that of coding sequences (25% G+C compared to 30%; P < 0.001). Thus, much of the non-coding sequence may be remnants of coding sequences that are in the process of being eliminated from the genome.

The gene encoding S-adenosylmethionine synthetase (metK), which catalyses the biosynthesis of S-adenosylmethionine (SAM), illustrates the initiation of this process. The metK sequence in the strain of R. prowazekii studied here has a termination codon within a region of the gene that is otherwise highly conserved among

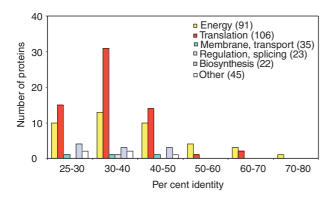


Figure 4 Histogram representation of the similarity of predicted *R. prowazekii* proteins to yeast proteins targeted to the mitochondria. Only protein pairs with per cent identity values greater than 25% are shown. Numbers in parentheses represent the total number of yeast mitochondrial proteins within each category. The yeast mitochondrial protein sequences have been taken from http://www.proteome.com.

bacterial species³⁵. However, a closely related strain does not have the termination codon. Many other defects, such as termination codons, insertions, and a preponderance of small deletions, have also been observed in the *metK* genes in several members of the spotted fever group *Rickettsia* (J.O.A. and S.G.E.A., unpublished observations). This random distribution of lethal mutations among some *metK* alleles from different *Rickettsia* species indicates that the gene may have just entered the extinction process. This distribution, and the identification of 11 more pseudogenes for carboxypeptidase (*ypwA*), penicillin-binding protein (*pbpC*), succinyl CoA-transferase (*scoB*), transposase (*tra3*), resolvase (*pin*), conjugative transfer protein (*taxB*), a hypothetical protein (*yfc*1) and four different fragmented open reading frames for (p)ppGpp 3'-pyrophosphohydrolase, indicates that the *R. prowazekii* genome continues to eliminate genes.

Genome sequences can be purged by a more abrupt mechanism. This consists of intrachromasomal recombination at duplicated sequences, which can result in the deletion of intervening sequences, the loss of a sequence duplication and the rearrangement of flanking

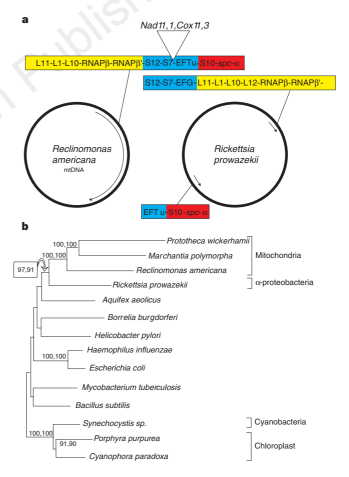


Figure 5 The organization and phylogenetic relationships of gene encoding ribosomal protein from *R. prowazekii* and the mitochondrial genome of *Reclinomonas americana*. **a**, The organization of ribosomal-protein genes. The S10, spc and α-operons are organized similarly in these two genomes, except that several ribosomal-protein genes³⁸ have been deleted from the mitochondrial genome of *Reclinomonas americana*. **b**, The phylogenetic relationships of mitochondria and bacteria were derived from the combined amino-acid sequences of ribosomal proteins S2, S3, S7, S10, S11, S12, S13, S14, S19, L5, L6 and L16. Neighbour-joining and maximum-parsimony methods gave identical topologies. Branch lengths are proportional to those reconstructed by using the neighbour-joining method. Values at nodes are bootstrap values indicating the degree of support for individual clusters under each method (neighbour-joining, maximum parsimony). Only bootstrap values >90% are shown.

sequences. Such a mechanism will account for the presence in *R. prowazekii* of one, unlinked copy of *rrs* and *rrl*, both of which are surrounded by new flanking sequences³⁶. Likewise, *R. prowazekii* has one *tuf* gene and one *fus* gene in atypical clusters that seem to have been created by intrachromosomal recombination between the two *tuf* genes that are normally found in Gram-negative bacteria³⁷. Indeed, rearranged gene operon structures encoding ribosomal proteins are characteristic of all members of the genus *Rickettsia* (H. Amiri, C.A. and S.G.E.A., unpublished observations).

Conserved operons that are found in free-living bacteria are often dispersed throughout the *Rickettsia* genome (see above). The *R. prowazekii* genome contains an unusually small fraction of repeat sequences (<10% of that observed in free-living bacteria). We suggest that the repeat sequences found in the ancestor to the *Rickettsia* have been 'consumed' by the intrachromosomal-recombination mechanism that generated some of the deletions and rearrangements seen in *R. prowazekii*. Such intrachromosomal recombinants arise at a substantial rate in bacteria growing in culture, but here they are eliminated from the populations by selection. That such remnants of intrachromosomal recombination are retained in *R. prowazekii* indicates that purifying selection has been attenuated in this organism.

Mitochondrial affinities

The reduction in genome size in mitochondria and *Rickettsia* is likely to have occurred independently in the two lineages. Most of

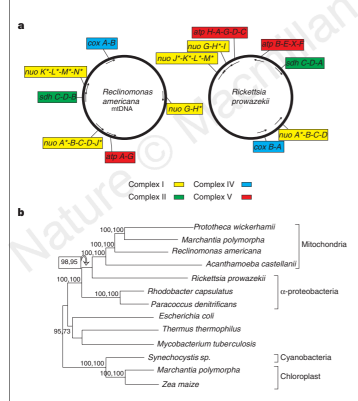


Figure 6 The organization and phylogenetic relationships of genes involved in ATP synthesis from *R. prowazekii* and the mitochondrial genome of *Reclinomonas americana*. **a**, The organization of bioenergetic genes. **b**, The phylogenetic relationships of mitochondria and bacteria were derived from the combined amino-acid sequences of NADH dehydrogenase I chains A, J, K, L, M and N, which are encoded by the genes *nuoA*, *J*, *K*, *L*, *M*, *N*. These genes are highlighted by asterisks in **a**. Neighbour-joining and maximum-parsimony methods gave identical topologies. Branch lengths are proportional to those reconstructed using the neighbour-joining method. Values at nodes are bootstrap values indicating the degree of support for individual clusters under each method (neighbour-joining, maximum parsimony). Only bootstrap values >90% are shown.

the genes supporting mitochondrial activities are nuclear. Many of the 300 proteins encoded in the nucleus of the yeast *Saccharomyces cerevisiae* but destined for service within the mitochondrion are close homologues of their counterparts in *R. prowazekii*. Nearly one-quarter of these proteins are required for bioenergetic processes and another one-third of them are required for the expression of the genes encoded in the mitochondrial genome. In total, more than 150 nucleus-encoded mitochondrial proteins share significant sequence homology with *R. prowazekii* proteins (Fig. 4).

Another group of 58 nucleus-encoded mitochondrial proteins represents components of the mitochondrial transport machinery and regulatory system (Fig. 4). These include proteins found in the mitochondrial outer membrane and others involved in splicing reactions. Such proteins have probably been secondarily recruited to mitochondria from genomes not necessarily related to that of the α -proteobacterial ancestor.

The mitochondrial genome of the early diverging, freshwater protozoan Reclinomonas americana is more like that of a bacterium than any other mitochondrial genome sequenced so far³⁸. This genome contains 67 protein-coding genes, most of which provide components of genetic processes and the bioenergetic system³⁸. Several gene clusters in this mitochondrial genome are reminiscent of those in bacteria (Figs 5a, 6a). Most similarities represent retained, ancestral traits present in the common ancestor of bacteria and mitochondria. For example, the genes rplKAJL and rpoBC are identically organized in R. prowazekii and the mitochondrial genome of Reclinomonas americana. Likewise, the genes encoding the S10, spc and the α -ribosomal protein operons are organized similarly in the two genomes. The immediate proximity of these two clusters in the Reclinomonas americana mitochondrial DNA is reminiscent of the arrangement in free-living bacteria, whereas the physical separation of the two clusters in the R. prowazekii genome is atypical. A further rearrangement event is indicated by the fact that the rpsLrpsGfus cluster is located upstream of the rplKAJLrpoBC cluster in R. prowazekii, rather than downstream as it is in the Reclinomonas americana mtDNA. Phylogenetic reconstructions based on ribosomal proteins within each of these two clusters indicate that there is a close evolutionary relationship between R. prowazekii and mitochondria (Fig. 5b).

Mitochondria and *R. prowazekii* have a similar repertoire of proteins involved in ATP production and transport, including genes encoding components of the TCA cycle, the respiratory-chain complexes, the ATP-synthase complexes and the ATP/ADP translocases. There are some similarities in the gene orders of some functional clusters (Fig. 6a). There are also some rearrangements of clusters that are specific to *Rickettsia*. One example is the inversion of segments corresponding to *nuoJKLM* and *nuoGHI*. Another is the scattered displacement of genes involved in the biogenesis of cytochrome *c*. Nevertheless, phylogenetic reconstructions based on components of the NADH dehydrogenase complexes indicate that there is a close evolutionary relationship between *R. prowazekii* and mitochondria (Fig. 6b).

We have identified as many as five genes coding for ATP/ADP transporters, all of which are expressed (R.M.P. *et al.*, unpublished observations). The *Rickettsia* ATP/ADP translocases are monomers with 12 transmembrane regions each, whereas the mitochondrial translocates are dimers with six transmembrane regions per dimer. We found no relationship between the primary structures of the mitochondrial and *Rickettsia* ATP/ADP translocases, indicating that these transport systems may have originated independently.

The study of the *R. prowazekii* genome sequence supports the idea that aerobic respiration in eukaryotes originated from an ancestor of the *Rickettsia*, as indicated previously by phylogenetic reconstructions based on the rRNA gene sequences^{7,9}. Phylogenetic analyses of the *petB* and *coxA* genes indicate that the respiration systems of *Rickettsia* and mitochondria diverged \sim 1,500–2,000 million years ago¹⁰, shortly after the amount of oxygen in the atmosphere began



to increase. The finding that the ATP/ADP translocases in R. prowazekii and mitochondria are of different evolutionary origin is problematic (R.M.P. et al., unpublished observations). Free-living bacteria do not seem to have homologues of ATP/ADP translocases, which are found only in organelles and in two obligate intracellular parasites, Rickettsia and Chlamydia. Thus it is not known whether the original endosymbiont was capable of efficient exchange of adenosine nucleotides with its host cell. More detailed comparative analysis of the genomes of α -proteobacteria may refine our understanding of the origins of mitochondria.

Methods

Genome sequencing. We prepared genomic DNA from the Madrid E strain of R. prowazekii, which was originally isolated in Madrid from a patient who died in 1941 with epidemic typhus. We propagated R. prowazekii in the yolk sac of embryonated hen eggs and purified DNA according to standard procedures³⁹. We sequenced the R. prowazekii genome by a whole-genome shotgun approach in combination with shotgun sequencing of a selected set of clones from a cosmid library (A.Z. et al., unpublished observations). Genomic and cosmid DNA was sheared by nebulization to an average size of \sim 2 kb. The random fragments were cloned into a modified M13 vector using the double adaptor method 40 . We collected 19,078 sequence reads during the random sequencing phase using Applied Biosystems 377 DNA sequencers (Perkin-Elmer).

The sequences were assembled and the consensus sequence was edited using the STADEN program $^{\rm 41}.$ We verified the structure of the assembled sequence by end-sequencing of 3-kb-insert λ Zap II clones³⁶, 10-kb λ clones and 30-kb cosmid clones. More than 97% of the genome was covered by clones from the three different libraries (A.Z. et al., unpublished observations). Gaps between contigs were closed by direct sequencing of clones from the three libraries or of polymerase chain reaction (PCR) products. The final four gaps were closed by direct sequencing of PCR products generated with the Long Range PCR system (Gene Amp). Regions of ambiguity were identified by visual inspection of the assembly and resequenced. The final assembly contains \sim 20,000 sequences. The genome sequence has eightfold coverage on average and no single region has less than twofold coverage. We estimate the overall error frequency to be $< 1 \times 10^{-5}$.

Informatics. Sequence analysis and annotation was managed by CapDB (T.S.-P. et al., unpublished observations). We identified open reading frames of more than 50 codons as genes on the basis of their characteristic patterns in nucleotide-frequency statistics¹⁴ using BioWish⁴². The identified genes were analysed using the program BLASTX⁴³ to search for sequence similarities in EMBL, TREMBL, SwissProt and in-house databases. We identified tRNA genes with the program tRNA scan-SE⁴⁴. Remaining frameshifts were considered to be authentic and annotated as pseudogenes. Families of paralogues were constructed using BLAST to search for sequence similarities within the R. prowazekii genome. Multiple alignments and phylogenetic trees for genes with significant sequence similarities to genes in the public databases were constructed automatically using CLUSTAL-W⁴⁵, Phylo_win⁴⁶ and GRS⁴⁷. The final annotation was based on manual inspection of the phylogenetic placement of R. prowazekii in the resulting gene trees.

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Table 1. Functional classification of *Rickettsia prowazekii* protein-coding genes. Gene numbers correspond to those in Fig. 2. Percentages represent per cent identitites.

AMINO ACID METABOLISM	Rri, 90.4%) RP321 <i>ipxB</i> lipid A disaccharide synthetase (B- <i>Hin</i> , 27.3%)	RP115 nuoF NADH dehydrogenase I chain F (B- <i>Pde</i> , 69.1%)
RP743 glyA serine hydroxymethyl transferase (B-Mex, 60.9%)	RP254 IpxC UDP-3-O-acyl N-acetylglucosamine deacetylase	RP797 nuoG NADH dehydrogenase I chain G (E- <i>Bta</i> mt, 49.3%) RP796 nuoH NADH dehydrogenase I chain H (E- <i>Ram</i> mt, 63.5%)
RP428 IVE branched-chain amino acid aminotransferase (B- Eco, 36.8%)	(B-Eco, 44.4%) RP009 lpxD UDP-3-O-(R-3-hydroxymyristoyl)-glucosamine N-	RP795 nuol NADH dehydrogenase I chain I (E-Ram mt, 71.4%) RP790 nuol NADH dehydrogenase I chain J (E-Ram mt 42.3%)
Amino acid degradation RP091 aatA aspartate aminotransferase (B-Rme, 55.6%)	acetyltransferase (B-Rri, 92.4%) RP062 kdsA 3-deoxy-d-manno-octulosonic acid 8-phosphate	RP791 nuoK NADH dehydrogenase I chain K (B-Pde, 61.4%) RP792 nuoL1 NADH dehydrogenase I chain L (E-Ram mt, 45.5%)
RP618 pccA propionyl-CoA carboxylase α chain (E-Rno,	synthetase (B- <i>Pha</i> , 45.1%) RP379 kdsB CTP:CMP-3-deoxy-manno-octulosonate-cytidylyl	RP283 nuoL2 NADH dehýdroğenase I chain L (E-Aja cp. 27.0%) RP282 nuoL3 NADH dehydrogenase I chain L (E-Ame mt, 17.0%)
45.0%) RP619 <i>pccB</i> propionyl-CoA carboxylase β chain (E- <i>Hsa</i> ,	transferase (B-Hin, 34.8%)	RP793 nuoM NADH dehydrogenase I chain M (E-Ram mt, 46.6%) RP537 nuoN1 NADH dehydrogenase I chain N (E-Ram mt, 34.6%)
63.3%) RP449 tdcB threonine dehydratase (E-Yeast, 35.3%)	(B- <i>Eco</i> , 28.9%)	RP284 nuoN2 NADH dehydrogenase I chain N (E-Ram mt, 23.2%)
BIOSYNTHESIS OF COFACTORS25	RP505 kpsF polysiliac acid capsule expression protein (B- Eco, 36.9%)	RP270 petA Rieske-I iron sulphur protein (B-Bja, 58.3%) RP271 petB cytochrome b (B-Rru, 65.4%)
Folic acid	RP833 <i>omp</i> cell surface antigen, 17 kD (B- <i>Rty</i> , 46.9%) RP160 <i>omp</i> 1 OMP1 precursor (B- <i>Bab</i> , 29.5%)	RP863 pntAA NAD(P) transhydrogenase α subunit (B-Eco, 37.7%) RP862 pntAB NAD(P) transhydrogenase α subunit (B-Hin, 44.7%)
RP536 folC folylpolyglutamate synthetase (B-Bsu, 34.5%)	RP771 pal peptidoglycan-associated lipoprotein (B-Eco, 37.9%)	RP862 pntAB NAD(P) transhydrogenase α subunit (B-Hin, 44.7%) RP074 pntB NAD(P) transhydrogenase β subunit (B-Hin, 51.5%)
46.3%)	RP476 rfaJ lipopolysaccharide 1,2-glucosyltransferase (B- Sty, 26.1%)	Fermentation RP110 ackA acetate kinase (B-Cts, 38.4%)
RP383 folE GTP cyclohydrolase I (B-Syn, 48.1%)	RP004 rfbA O-antigen export system permease (B-Kpn, 22.3%)	
Haem and porphyrins RP841 hemA delta-aminolevulinate synthase (B-Bja, 49.1%)	RP003 rfbE O-antigen ABC export system, ATP-binding protein (B-Yen, 34.0%)	Glycolysis RP492 ppdK pyruvate, orthophosphate dikinase (E-Ftr, 48.8%)
RP539 hemB delta-aminolevulinate dehydratase (B-Bja, 53.3%) RP466 hemC prophobilinogen deaminase (B-Pmi, 39.1%)	RP334 rffE UDP-N-acetylglucosamine 2-epimerase (B-Bsu, 26.8%)	Phosphate
RP885 hemE uroporhyrinogen decarboxylase (B-Rca, 42.6%) RP882 hemF coproporhyrinogen III oxidase (E-Gma, 43.0%)	RP018 sca1 cell surface antigen (B-Rri, 24.9%) RP081 sca2 cell surface antigen (B-Rri, 27.4%)	RP589 ppa inorganic pyrophosphatase (B-Eco, 59.3%)
RP884 hemH ferrochelatase (B-Syn, 40.4%)	RP451 <i>sca</i> 3 cell surface antigen (B- <i>Rri</i> , 27.6%) RP498 <i>sca</i> 4 cell surface antigen (B- <i>Rja</i> , 67.4%)	Pyruvate dehydrogenase RP261 pdhA pyruvate dehydrogenase E1 component, α sub-
RP847 hemK protoporphyrinogen oxidase (B-Eco, 44.3%) RP175 hemN oxygen-independent coproporphyrinogen II (B-	RP704 sca5 cell surface antigen (B-Rri, 72.5%)	unit (E-Ath, 44.0%)
Bsu, 34.4%) Lipoate		RP262 pdhB pyruvate dehydrogenase E1 component, β sub- unit (E-Sce, 59.7%)
RP742 <i>lipA</i> lipoic acid synthetase (B- <i>Hin</i> , 50.5%) RP876 <i>lipB</i> lipoic acid ligase (B- <i>Mtu</i> , 35.6%)	CELLULAR PROCESSES	RP530 pdhC dihydrolipoamide acetyltransferase E2 component (E-Rno, 45.1%)
Menaquinone and ubiquinonens	Cell division RP251 ftsA cell division protein FtsA (B-Hin, 29.5%)	RP460 pdhD dihydrolipoamide dehydrogenase E3 component (E-Psa, 54.7%)
RP190 coq7 ubiquinone biosynthesis prt Coq7 (E-Rno, 36.9%)	RP043 ftsH cell division protein FtsH (B-Eco, 54.0%) RP163 ftsJ cell division protein FtsJ (B-Eco, 44.4%)	RP805 pdhD dihydrolipoamide dehydrogenase E3 component (Zym, 51.1%)
RP479 ispB octaprenyl-diphosphate synthase (B-Eco,	RP823 ftsK cell division protein FtsK (B-Cbu, 41.5%)	TCA cycle RP799 acnA aconitate hydratase (B-Lpn, 59.1%)
36.5%) RP686 ubiA 4-hydroxybenzoate octaprenyltransferase (B-	RP250 ftsQ cell division protein FtsQ (B-Hin, 17.9%) RP411 ftsW cell division protein FtsW (B-Eco, 33.2%)	RP665 fumC fumarate hydratase (B-Ror, 63.5%)
Eco, 36.1%) RP541 ubiX 3-octaprenyl-4-hydroxybenzoate carboxylase (B-	RP775 ftsY cell division protein FtsY (B-Hin, 43.2%) RP666 ftsZ cell division protein FtsZ (B-Wsp, 65.3%)	RP844 gltA citrate synthase (B-Rty, 97.8%) RP265 icd isocitrate dehydrogenase (B-Tth, 38.6%)
Eco, 53.2%) RP680 ubiE ubiquinone biosynthesis methyltransferase (B-	RP056 gidA glucose inhibited division protein A (B-Eco, 48.8%) RP057 gidB glucose inhibited division protein (B-Ppu, 26.8%)	RP376 mdh malate dehydrogenase (B-Can, 51.5%) RP128 sdhA succinate dehydrogenase, flavoprotein subunit
Eco, 44.0%) RP622 ubiG 3-demethylubiquinone methyltransferase (B-Eco,	RP815 maf MAF protein (B-Bsu, 38.1%) RP042 mesJ cell cycle protein MesJ (B-Eco, 22.1%)	(B- <i>Bja</i> , 70.0%) RP044 <i>sdhB</i> succinate dehydrogenase, iron-sulphur protein (E-
39.1%)	RP768 mreB rod shape-determining protein (B-Eco, 60.5%)	Ram mt, 69.0%) RP126 sdhC succinate dehydrogenase, cytochrome b ₅₅₆
Thio- and glutaredoxin	RP767 mreC rod shape-determining protein (B-Eco, 23.1%) RP280 rodA rod shape-determining protein (B-Eco, 38.1%)	subunit (E- <i>Ram</i> mt, 39.5%) RP127 sdhD succinate dehydrogenase, subunit IV (E- <i>Ram</i>
RP204 <i>grxC</i> 1 glutaredoxin 3 (B- <i>Eco</i> , 50.0%) RP745 <i>grxC</i> 2 glutaredoxin 3 (B- <i>Syn</i> , 50.0%)	Cell killing	mt, 25.6%)
RP327 tdpX1 thioredoxin-peroxidase (B-Hpy, 54.0%) RP002 trxA thioredoxin (B-Ani, 52.8%)	RP555 <i>tlyA</i> hemolysin (B- <i>Thy</i> , 34.3%) RP740 <i>tlyC</i> hemolysin (B- <i>Thy</i> , 28.8%)	RP180 sucA 2-oxoglutarate dehydrogenase (B-Hin, 44.3%) RP179 sucB dihydrolipoamide succinyltransferase (B-Eco, 48.7%)
RP445 trxB1 thioredoxin reductase (B-Hin, 52.0%) RP514 trxB2 thioredoxin reductase (B-Cpa, 28.4%)	Chaperones and stress-induced proteins	RP433 sucC succinyl-CoA synthetase, β subunit (B-Eco, 52.1%) RP432 sucD succinyl-CoA synthetase, α subunit (E-Ddi, 70.7%)
CELL ENVELOPE59	RP670 cspA cold shock protein (B-Scl, 57.6%) RP816 dksA DnaK suppressor protein (B-Hin, 38.8%)	Sugars
	RP184 dnaJ heat shock protein (B-Bov, 49.7%)	RP509 exoC phosphomannomutase (B-Abr, 42.7%) RP299 lacA galactosidase acetyltransferase (B-Mpn, 44.4%)
Diaminopimelate RP316 asd aspartate-semialdehyde dehydrogenase (B-Vch,	RP185 dnaK heat shock protein 70 (B-Rme, 72.7%) RP626 groEL heat shock protein GroEL (B-Rme, 69.4%)	
43.3%) RP429 dapA dihydrodipicolinate synthase (A-Mja, 39.6%)	RP627 groES heat shock protein GroES (B-Rty, 98.9%) RP629 grpE heat shock protein GrpE (B-Ccr, 39.5%)	FATTY ACID AND PHOSPHOLIPID METABOLISM25
RP148 dapB dihydrodipicolinate reductase (B-Hpy, 37.7%) RP194 dapD tetrahydrodipicolinate N-succinyltransferase (B-	RP200 hscA heat shock protein A (B-Hin, 39.6%) RP320 hslU heat shock protein HslU (B-Bsu, 54.8%)	RP620 aas 2-acyl-glycerol-phosphate-ethanolamine (B-Eco, 39.9%)
Eco, 57.9%) RP874 dapE succinyl-diaminopimelate desuccinylase (B-Hin,	RP319 hs/V heat shock prt HsIV (B-Hin, 54.1%) RP273 hsp22 heat shock protein (E-Phy cp, 29.0%)	RP038 aco1 acyl-CoA desaturase (E-Yeast, 27.6%) RP763 acpP acyl carrier protein (B-Lmu, 52.6%)
37.5%)	RP840 htpG heat shock protein C62.5 (B-Eco, 43.1%)	RP577 acpS holo-[acyl carrier protein] synthase (B-Eco, 38.5%)
PD415 danE diaminonimolato onimoraco (PUin 35.0%)	THE TO THE	
RP415 dapF diaminopimelate epimerase (B-Hin, 35.0%) RP753 lysC aspartokinase (B-Bst, 37.3%)	Detoxification	RP533 birA biotin Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cdsA phosphatidate cytidylyltransferase (B-Eco, 31.5%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins		RP533 birA biotin Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cdsA phosphatidate cytidylyltransferase (B-Eco, 31.5%) malonyl-CoA:Acyl carrier protein transacylase (B-Bsu, 40.3%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%)	Detoxification RP555 sodB superoxide dismutase (B-Lpn, 53.4%) RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion	RP533 bi/A biotin Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cdsA phosphalidate cylidylytransferase (B-Eco, 31.5%) RP765 fabb malonyl-CoA:Acyl carrier protein transacylase (B-Bu, 40.3%) RP764 fabF 3-xxxxxy-lacyl-carrier-protein] synthase II(B-
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco,	Detoxification RP535 sodB superoxide dismutase (B-Lpn, 53.4%) RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%)	RP533 birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóAs phosphatidate oylidylthransferase (B-Eo., 31.5%) RP735 fabD malonyl-CoA:Aoyl carrier protein transacylase (B-Bsu, 40.3%) RP64 fabF 3-oxoacyl-(acyl-carrier-protein) synthase II(B-Eo., 53.5%) RP762 fabG 3-oxoacyl-(acyl-carrier-protein) reductase (B-
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 [gtD] prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int apolipoprotein N-acyltransferase (B-Hin, 29.1%)	Detoxification RP535 sod8 RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE	RP533 bir/a biotin Ac-CoA carboxylase synthase (B-P0e, 33.6%) RP424 cdsA phosphatidate cylidylytransferase (B-Eco, 31.5%) RP735 fabb malonyl-CoA:Aoyl carrier protein transacylase (B-But, 40.3%) RP764 fabF 3-oxoacyl-(acyl-carrier-protein) synthase II(B-Eco, 53.5%) RP762 fabb 3-oxoacyl-(acyl-carrier-protein) reductase (B-Fme, 54.8%) RP772 fabb 3-oxoacyl-(acyl-carrier-protein) synthase (B-Foxoacyl-(acyl-carrier-protein)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 [gtD] RP366 Int RP300 nlpD RP390 nlpD RP390 rear lipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 23.9%)	Detoxification RP535 sod8 RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 fth signal recognition particle receptor protein (B-Eco,	RP533 bir/s biotin Ac-CoA carboxylase synthase (B-P0e, 33.6%) RP424 cdsA phosphatidate cylidylytransferase (B-Pc0, 31.5%) RP735 fabb RP735 fabb malonyl-CoA:Aoyl carrier protein transacylase (B-Bcu, 40.3%) RP764 fabF 3-oxoacyl-[acyl-carrier-protein] synthase II(B-Ecc, 53.5%) RP762 fabG 3-oxoacyl-[acyl-carrier-protein] reductase (B-Rep, 54.8%) RP772 fabH 3-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.3%) RP365 fabI enoyl-[acyl-carrier-protein] reductase (B-Asp, 49.0%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP46 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int apolipoprotein (B-Hin, 22.4%)	Detoxification RP535 sod8 RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%)	RP533 birá blotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóAs phosphatidate cylidylytransferase (B-Púe, 33.6%) RP735 fabD malonyl-CoA:Aoyl carrier protein transacylase (B-Bú, 40.3%) RP764 fabF 3-oxoacyl-[acyl-carrier-protein] synthase II(B-Euc, 53.5%) RP762 fabG 3-oxoacyl-[acyl-carrier-protein] reductase (B-Rim, 54.8%) RP772 fabH Reserved (Fabre (B-Riss)) RP365 fabI Reserved (Fabre (B-Riss)) RP008 fabZ 3-Rhydroxymyristoyl acyl carrier protein dehy dratase (B-Ris, 19.7%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP466 [gtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD apolipoprotein (B-Hin, 22.4%) RP390 rlpA rare lipoprotein (B-Hin, 22.4%) RP242 tolC outer membrane protein (B-Eco, 22.9%) RP048 yidC inner membrane protein, 60 kDa (B-Hin, 30.4%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Lin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 seeA preprotein translocase SeeA subunit (B-Rca, 51.8%)	RP533 bira
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP46 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP300 nlpD lipoprotein (B-Hin, 22.4%) RP300 nlpA rare lipoprotein A (B-Hin, 29.9%) RP294 lctC outer membrane protein (B-Eco, 22.9%) RP048 yldC inner membrane protein (B-Eco, 22.9%) Murein sacculus RP055 alr RP389 dacF penicillin binding protein precursor (B-Bsu, 19.5%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Lin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP070 secB preprotein translocase SecB subunit (B-Hin, 30.7%) RP586 secD protein-export membrane protein (B-Eco, 40.4%)	RP533 birá biotín Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cd24 phosphatidate oytidyltransferase (B-Pco, 31.5%) RP735 tabb malonyl-CoA:Aoyl carrier protein transacylase (B-Bst, 40.3%) RP762 tabb a-coacyl-tacyl-carrier-protein] synthase II(B-Eco, 53.5%) RP762 tabb a-coacyl-tacyl-carrier-protein] reductase (B-Rme, 54.8%) RP772 tabb a-coacyl-tacyl-carrier-protein] synthase (B-Ra, 47.3%) enoyl-tacyl-carrier-protein] synthase (B-Ra, 47.3%) enoyl-tacyl-carrier-protein] reductase (B-Asp, 49.0%) RP737 tabb a-coacyl-tacyl-carrier-protein] reductase (B-Asp, 49.0%) renoyl-tacyl-carrier-protein] reductase (B-Asp, 49.0%) acyl-carrier-protein] reductase (B-Asp, 49.0%) acyl-carrier-protein] reductase (B-Asp, 49.0%) acyl-carrier-protein] reductase (B-Asp, 49.0%) acyl-carrier-protein] reductase (B-Asp, 49.0%) reductase (B-Ri, 91.7%) acyl-carrier-protein] reductase (B-Asp, 49.0%) acyl-carrier-protein] reductase (B-
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD lipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 nlpD rafel in training tra	Detoxification RP535 sod8 RP759 thdF thiophene and furan oxidizer (B-Lpn, 53.4%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA RP116 lepB RP575 secA preprotein translocase SecB subunit (B-Rac, 51.8%) RP5070 secB preprotein translocase SecB subunit (B-Rac, 51.8%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP134 secE preprotein translocase SecB subunit (B-Fac, 30.7%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP134 secE preprotein translocase SecB subunit (B-Fac, 37.3%)	RP533 birá biotin Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cd2A phosphatidate cylidylytransferase (B-Eco, 31.5%) RP765 fabb RP764 fabF RP762 fabG RP772 fabH RP762 fabG RP772 fabH RP763 fabB RP773 fabH RP764 fabF RP773 fabH RP765 fabB RP773 fabH RP765 fabB RP773 fabB RP774 fabH RP767 fabB RP775 fabB RP775 fabB RP775 fabB RP775 fabB RP775 fabB RP777 fabB RP777 fabB RP777 fabB RP777 fabB RP777 fabB RP777 fabB RP778 fabB RP778 fabB RP778 fabB RP779 fabB
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD lipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 nlpD rare lipoprotein N-acyltransferase (B-Hin, 29.1%) RP224 tolic RP048 yidc outer membrane protein (B-Eco, 29.9%) RP499 glacF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 3.8%) RP454 glmU UDP-N-acetylglucoseamine pyrophosphorylase	Detoxification RP535 sod8 RP759 thdF thiophene and furan oxidizer (B-Lpn, 53.4%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) signal recognition particle receptor protein (B-Eco, 49.6%) RP375 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP134 secF preprotein translocase SecE subunit (B-Hin, 37.7%) RP114 secF protein-export membrane protein (B-Bsu, 37.3%) RP119 secG protein-export membrane protein (B-Hin, 37.7%) RP079 secG protein-export membrane protein (B-Hin, 37.7%)	RP533 birá biotín Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cd2s phosphatidate cylidyltransferase (B-Eco, 31.5%) RP735 fabb malonyl-CoA:Aoyl carrier protein transacylase (B-Bst, 40.3%) 3-oxoacyl-[acyl-carrier-protein] synthase II (B-Eco, 53.5%) RP762 fabb 3-oxoacyl-[acyl-carrier-protein] reductase (B-me, 54.8%) RP772 fabr RP008 fabb 3-oxoacyl-[acyl-carrier-protein] synthase (B-Re, 47.3%) enoyl-[acyl-carrier-protein] reductase (B-Asp, 49.0%) RP008 fabb RP009 facall staty oxidation complex a subunit (E-Cel, 30.6%) RP389 fb06 facl Br009 fabr RP009 fabb RP009 fabr RP009 fabb RP009 fabr RP009 fab
RP753 IysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 IgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD prolipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 nlpD rlpA rate lipoprotein (B-Hin, 22.4%) RP244 tolC RP048 yidC inner membrane protein (B-Hin, 30.4%) Murein asaculus Allarine racemace (B-Hin, 29.5%) RP399 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddiB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP454 gfmU UDPN-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mraY1 phospho-N-acetylmuramoyl-pentapeptide-trans-	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD rotease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE RP315 the Secretion ATP-binding protein (B-Pae, 40.0%) RP315 the Secretion ATP-binding protein (B-Pae, 40.0%) RP316 the Secretion ATP-binding protein (B-Eco, 49.6%) RP317 the Secretion ATP-binding protein (B-Eco, 49.6%) RP318 the Secretion ATP-binding protein (B-Eco, 40.6%) RP318 the Secretion ATP-binding protein (B-Eco, 40.6%) RP318 the Secretion ATP-binding protein (B-Eco, 40.4%) RP318 the Secretion ATP-binding protein (B-Ec	RP533 birá biotín Ac-CoA carboxylase synthase (B-Pde, 33.6%) RP424 cd24 phosphatidate cylidylytransferase (B-Pco, 31.5%) RP735 fabb malonyl-CoA:Aoyl carrier protein transacylase (B-But, 40.3%) 3-oxoacyl-[acyl-carrier-protein] synthase II (B-Eco, 53.5%) 3-oxoacyl-[acyl-carrier-protein] reductase (B-RP72 fabb 3-oxoacyl-[acyl-carrier-protein] reductase (B-RP72 fabb 3-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.3%) 4-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.3%) 8-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.3%) 8-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.5%) 8-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.5%) 8-oxoacyl-[acyl-carrier-protein] synthase (B-Rea, 47.5%) 8-oxoacyl-(acyl-carrier-protein] synthase (B-Rea, 49.0%) 8-oxoacyl-(acyl-carrier-protein] 8-oxoacyl-(acyl-carrier-protein) 8-oxoacyl-(acyl-carrier-prot
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP300 nlpD lipoprotein (B-Hin, 22.4%) RP300 nlpD lipoprotein (B-Hin, 22.4%) RP300 nlpD lipoprotein (B-Hin, 23.0%) RP300 rlpD lipoprotein (B-Hin, 23.0%) RP300 rlpD lipoprotein (B-Hin, 23.0%) RP244 lglC outer membrane protein (B-Eco, 22.9%) RP245 lglC penicillin binding protein precursor (B-Bsu, 33.8%) RP309 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP454 glmU UDPN-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mra/Y phosphor-N-acetylmuramoyl-pentapeptide-transferase (B-Hin, 49.9%) RP825 mra/Y2 phosphor-Nacetylmuramoyl-pentapeptide-transferase (B-Hin, 49.9%) RP825 mra/Y2 phosphor-Nacetylmuramoyl-pentapeptide-transferase (B-Hin, 49.9%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Irin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE rotease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTT-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secD protein-export membrane protein (B-Eco, 40.4%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP114 secE protein translocase SecB subunit (B-Bsu, 37.3%) RP079 secG protein-export membrane protein (B-Hin, 37.7%) RP079 secG protein-export membrane protein (B-Hin, 32.7%) RP079 secG protein-export membrane protein (B-Hin, 32.7%) RP079 secG protein-export membrane protein (B-Hin, 32.7%) RP079 secG protein-export membrane protein (B-Fup, 32.0%) RP639 secG protein-export membrane protein (B-Fup, 32.0%) RP639 secG protein-export membrane protein (B-Fup, 32.0%) RP639 secG protein-export membrane protein (B-Eco, 60.0%)	RP533 birá RP424 cdz Aphosphatidate cylidylytrans/erase (B-Eco, 31.5%) RP735 fabb RP764 fabF RP762 fabG RP763 fabG RP763 fabG RP763 fabG RP764 fabF RP764 fabF RP765 fabG RP766 fabG RP766 fabG RP766 fabG RP766 fabG RP767
RP753 IysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int RP300 nlpD lipoprotein (B-Hin, 22.4%) RP390 nlpD lipoprotein (B-Hin, 22.4%) RP390 rlpD lipoprotein (B-Hin, 23.9%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP048 yidC outer membrane protein (B-Eco, 22.9%) RP049 yidC alaniner racemace (B-Hin, 29.5%) RP095 air alanine racemace (B-Hin, 29.5%) RP249 ddilb — balanine ligase (B-Hin, 32.8%) RP454 gimU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mra11 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Fin, 34.9%) RP825 mra12 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Sac, 22.0%) RP807 mrcA pencililib indicing protein 1A (B-Eco, 35.6%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP586 secA preprotein translocase SecB subunit (B-Bsu, 37.3%) RP586 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP134 secE protein-export membrane protein (B-Hin, 37.7%) RP144 secF protein-export membrane protein (B-Hin, 37.7%) RP079 secY preprotein translocase SecE subunit (B-Bsu, 37.3%) RP079 secY preprotein translocase SecP subunit (B-Eco, 50.0%) trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion	RP533 birá RP424 cdz Aphosphatidate cylidylytrans/erase (B-Eco, 31.5%) RP735 fabb RP764 fabF RP762 fabG RP763 fabG RP763 fabG RP764 fabF RP764 fabF RP765 fabG RP766 fabG RP767 fabG RP766 fabG RP766 fabG RP767
RP753 IysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP300 nlpD lipoprotein (B-Hin, 22.4%) RP300 nlpD lipoprotein (B-Hin, 22.4%) RP300 nlpD lipoprotein (B-Hin, 23.6%) RP300 nlpD lipoprotein (B-Hin, 24.6%) RP40 ydd lipoprotein (B-Hin, 29.5%) RP404 ddlb D-alanine:D-alanine ligase (B-Hin, 32.8%) RP454 gimU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP559 mraY1 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bac, 22.0%) RP6579 murA UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Aca, 51.6%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) Signal recognition particle receptor protein (B-Eco, 49.6%) RP173 ffh signal recognition particle receptor protein (B-Eco, 49.6%) RP176 lepB Signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP586 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP586 secA preprotein translocase SecA subunit (B-Bau, 37.3%) RP586 secA preprotein translocase SecA subunit (B-Bau, 37.3%) RP588 secA protein-export membrane protein (B-Hin, 37.7%) RP589 secY protein-export membrane protein (B-Hin, 37.7%) RP639 secY protein-export membrane protein (B-Hip, 32.7%) RP639 secY protein-export membrane protein (B-Hip, 32.7%) RP639 secY protein-translocase SecA subunit (B-Eco, 50.0%) RP630 atpa ATP synthase F olubinit a (B-Rru, 65.2%) RP630 atpa ATP synthase F olubinit a (B-Rru, 51.5%)	RP533 birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidythransferase (B-Póe, 33.6%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bát, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eco, 53.5%) RP762 fabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rm, 6.48%) RP772 fabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rm, 6.48%) RP783 fabi RP784 fab, 47.3%) RP785 fabi RP784 RP784 fab 3-xxxacyl-lacyl-carrier-protein] synthase (B-Ray A-9xxacyl-lacyl-carrier-protein) synthase (B-Ray A-9xxacyl-lacyl-carrier-protein) synthase (B-Ray A-9xxacyl-lacyl-carrier-protein) synthase (B-Ray A-9xxacyl-lacyl-carrier-protein) synthase (B-Ray 4.79%) RP787 facA acetyl-CoA acetyl-transferase (B-Páe, 54.5%) RP784 gyl-carrier-protein diacyl-gyl-carrier-protein (B-Eco, 30.6%) RP785 phbB acetoacetyl-CoA reductase (B-Zra, 52.9%) RP788 phbCl poly-beta-hydroxybutyrate polymerase (B-Cvi, 22.7%) RP750 pppA phosphatidylglycerophosphates A (B-Hin, 31.9%) RP049 ppsA phosphatidylglycerophosphates ymthase (B-Bsu, 18.9%) RP049 ppsA
RP753 IysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int RP300 nlpD lipoprotein (B-Hin, 22.4%) RP390 rlpD lipoprotein (B-Hin, 22.4%) RP390 rlpD lipoprotein (B-Hin, 23.9%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP048 yidC outer membrane protein (B-Eco, 22.9%) RP049 yidC alanine racemace (B-Hin, 29.5%) RP049 drilb Dentalinie protein precursor (B-Bsu, 33.8%) RP249 drilb Dentalinie ligase (B-Hin, 32.8%) RP454 gimU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.9%) RP595 mra12 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bac, 22.0%) RP807 mra (B-Bac, 81.6%) RP807 mra (DPN-acetylglucosamine 1-carboxyvinyltransferase (B-Aca, 51.6%) RP598 mra UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Aca, 51.6%) RP598 mra UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Aca, 51.6%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protein responsible receptor protein (B-Eco, 49.6%) RP315 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP575 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecB subunit (B-Bsu, 37.3%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP579 secA preprotein translocase SecP subunit (B-Bsu, 37.3%) RP639 secY preprotein translocase SecY subunit (B-Bsu, 37.3%) RP639 secY preprotein translocase SecY subunit (B-Rca, 50.0%) trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion RP803 atpA ATP synthase F o subunit (B-Rru, 66.2%) RP002 atpC ATP synthase F o subunit a (B-Rru, 51.5%) RP800 atpC ATP synthase F i epsilon subunit (B-Rru, 27.0%)	RP633 birá birán biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidyltransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bát, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rm, 6.48%) RP772 fabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rm, 6.48%) RP783 fabi enoyl-lacyl-carrier-protein] synthase (B-Rox, 47.3%) RP365 fabi enoyl-lacyl-carrier-protein] synthase (B-Rox, 47.3%) RP373 facA acetyl-CoA acetyl-transferase (B-Páe, 49.0%) 3R-Hydroxymyristoyl acyl carrier protein dehy dratase (B-Fáe, 47.7%) RP560 fadB latty oxidation complex a subunit (E-Oei, 30.6%) glycerol-3-phosphate dehydrogenase (B-Ecx, 32.1%) RP365 pbbB acetoacetyl-CoA reductase (B-Zra, 52.9%) RP382 phbCl poly-beta-hydroxybutyrate polymerase (B-Wex, 37.4%) RP50 pppA phosphatidylgylcerophosphates A (B-Hin, 31.9%) RP049 plsC 1-acyl-glycerol-3-phosphate acyltransferase (E-Sex, 23.6%) RP242 pssA phosphatidylserine synthase (B-Hpy, 28.8%) RP242 pssA phosphatidylserine synthase (B-Hpy, 28.8%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int application on Inp. lipoprotein (B-Hin, 22.4%) RP369 ripA rare lipoprotein (B-Hin, 22.4%) RP284 toliC RP048 yidC inner membrane protein (B-Eco, 22.9%) RP048 yidC inner membrane protein (B-Eco, 22.9%) RP048 yidC inner membrane protein, 60 kDa (B-Hin, 30.4%) Murein sacculus alanine racemace (B-Hin, 29.5%) RP399 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP454 gifU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP599 mr312 phospho-N-acetylglucoseamine pyrophosphorylase (B-Hin, 49.9%) RP855 mr312 phospho-N-acetylglucosamine penicile-transferase (B-Bac, 22.0%) RP579 mr4 UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-B-Aca, 51.6%) RP248 mr8 UDP-N-acetylquoryuroylglucosamine reductase	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP315 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP576 secA preprotein translocase SecB subunit (B-Ra, 37.3%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP114 secE protein-export membrane protein (B-Hin, 37.7%) RP079 secQ protein-export membrane protein (B-Hin, 37.7%) RP639 secY preprotein translocase SecP subunit (B-Eco, 50.0%) RP630 atpA ATP-synthase Fo subunit (B-Rru, 66.2%) RP003 atpA ATP-synthase Fo subunit a (B-Rru, 51.5%) RP800 atpC ATP-synthase Fo subunit a (B-Rru, 54.5%) RP801 atpA ATP-synthase Fo subunit (B-Ra, 77.0%) RP002 atpE ATP Synthase Fo subunit (B-Ra, 77.0%) RP002 atpE ATP Synthase Fo subunit (B-Ra, 77.0%)	RP533 birá birán biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 có24 phosphatidate oytidylytmas/erase (B-Eo., 31.5%) RP735 tabb malonyl-CoA:Aoyl carrier protein transacylase (B-Bst., 40.3%) RP762 tabb accoacyl-tacyl-carrier-protein] synthase II (B-Eo., 53.5%) RP762 tabb accoacyl-tacyl-carrier-protein] reductase (B-Rme, 54.8%) RP772 tabh accoacyl-tacyl-carrier-protein] reductase (B-Rme, 54.8%) acoacyl-tacyl-carrier-protein] synthase (B-Bac, 47.3%) encyl-tacyl-carrier-protein] synthase (B-Bac, 47.3%) RP737 tadA acetyl-CoA acetyltransferase (B-Pde, 54.5%) RP560 tabb accates (B-Fin, 91.7%) RP73 tadA acetyl-CoA acetyltransferase (B-Pde, 54.5%) RP442 gpsA glycerol-3-phosphate dehydrogenase (B-Eco, 32.1%) RP035 phbC1 poly-beta-hydroxybutyrate polymerase (B-Eci, 32.7%) RP709 phbC2 poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) RP709 pgsA phosphatidylglycerophosphatase A (B-Hin, 31.9%) RP449 plsC 1-acyl-glycerol-3-phosphate acyltransferase (E-Sec, 23.6%) RP449 plsC 1-acyl-glycerol-3-phosphate acyltransferase (E-Sec, 23.6%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP466 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD lipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 nlpD lipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 rlpA rare lipoprotein (B-Hin, 22.4%) RP224 tolic RP048 yldc limer membrane protein (B-Eco, 22.9%) RP495 alr RP399 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP454 glmU UDPN-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mraY1 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bcc, 22.0%) RP807 murA UDPN-acetylglucosamine 1-carboxyvinytltransferase (B-Bca, 23.6%) RP248 murB UDPN-acetylglucosamine reductase (B-Bsu, 35.7%) RP248 murB UDPN-acetylmuryuvoylglucosamine reductase (B-Bsu, 35.7%) RP247 murC UDPN-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP410 murD UDPN-acetylmuramoylalanine-D-glutamate lig-	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 22.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rac, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP575 secA protein-export membrane protein (B-Hin, 37.7%) RP576 secB protein-export membrane protein (B-Hin, 37.7%) RP578 secA protein-export membrane protein (B-Hin, 37.7%) RP579 secG protein translocase SecC subunit (B-Rin, 37.7%) RP579 secG protein translocase SecC subunit (B-Rin, 37.7%) RP579 secG protein translocase SecC subunit (B-Rin, 37.7%)	RP533 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate cytidyltransferase (B-Eoz, 31.5%) RP735 tabb malonyl-CoA:Aoyl carrier protein transacylase (B-Bsu, 40.3%) RP762 tabb accoacyl-tacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 tabb accoacyl-tacyl-carrier-protein] reductase (B-Rme, 54.8%) RP772 tabh accoacyl-tacyl-carrier-protein] synthase (B-Rme, 47.3%) enoyl-tacyl-carrier-protein] synthase (B-Rme, 47.3%) enoyl-tacyl-carrier-protein] synthase (B-Rez, 49.0%) enoyl-tacyl-tacyl-carrier-protein] synthase (B-Rez, 49.0%) enoyl-tacyl-tacyl-tacyl-carrier-protein] synthase (B-Rez, 49.0%) enoyl-tac
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD ilipoprotein N-acyltransferase (B-Hin, 29.1%) RP390 nlpD ilipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 22.9%) rare lipoprotein (B-Hin, 22.9%) rare lipoprotein (B-Hin, 22.9%) rare lipoprotein (B-Hin, 22.9%) rare lipoprotein (B-Hin, 23.9%) rare lipoprotein (B-Hin, 23.9%) rare lipoprotein (B-Hin, 23.9%) rare lipoprotein recursor (B-Bsu, 33.8%) rare lipoprotein recursor (B-Hin, 34.3%) rare lipoprotein recursor (B-Hin, 34.9%) rare lipoprotein recursor (B-Hin, 34.9%) rare lipoprotein recursor (B-Bsu, 34.3%) rare lipoprotein recursor (B-Bsu, 34.3%) rare lipoprotein recursor (B-Rsu, 22.0%) rare lipoprotein recursor (B-Rsu, 22.0%) rare lipoprotein recursor (B-Rsu, 35.7%) rare lipoprotein rare lipoprotein recursor (B-Rsu, 35.7%) rare lipoprotein rare lipo	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP315 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP576 secA preprotein translocase SecB subunit (B-Ra, 37.3%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP114 secE protein-export membrane protein (B-Hin, 37.7%) RP079 secQ protein-export membrane protein (B-Hin, 37.7%) RP639 secY preprotein translocase SecP subunit (B-Eco, 50.0%) RP630 atpA ATP-synthase Fo subunit (B-Rru, 66.2%) RP003 atpA ATP-synthase Fo subunit a (B-Rru, 51.5%) RP800 atpC ATP-synthase Fo subunit a (B-Rru, 54.5%) RP801 atpA ATP-synthase Fo subunit (B-Ra, 77.0%) RP002 atpE ATP Synthase Fo subunit (B-Ra, 77.0%) RP002 atpE ATP Synthase Fo subunit (B-Ra, 77.0%)	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bát, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rme, 64.8%) RP772 fabH 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rme, 64.8%) RP373 fabl and acelyl-CoA acelyl-carrier-protein] synthase (B-Asp, 49.0%) fabZ 3-R-ydcroxymyristoyl acyl carrier protein dehy dratase (B-Rin, 19.1%) RP737 fab acelyl-CoA acelyl-transferase (B-Pde, 54.5%) acelyl-CoA acelyl-transferase (B-Pde, 54.5%) RP442 gpsA glycerol-3-phosphate dehydrogenase (B-Eox, 32.1%) RP373 phbB aceloacetyl-CoA reductase (B-Ze, 39.9%) RP38 phbB aceloacetyl-CoA reductase (B-Ze, 39.9%) RP38 phbB aceloacetyl-CoA reductase (B-Ze, 39.9%) RP38 phbB aceloacetyl-CoA reductase (B-Ze, 39.9%) RP39 phbB phosphatidylglycerophosphates A (B-Hin, 31.9%) RP049 pgsA phosphatidylglycerophosphate synthase (B-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Sex, 23.6%) Phosphatidylserine synthase (B-Hpy, 28.8%) maic enzyme (B-Hin, 45.5%) Vac III poprotein precursor (B-Sil, 33.8%) PURINES, PYRIMIDINES.
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) RP360 lnt RP300 nlpD RP300	Detoxification RP535 sodB Superoxide dismutase (B-Lpn, 53.4%) RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 22.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rac, 51.8%) RP370 secB preprotein translocase SecB subunit (B-Hin, 30.7%) RP578 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP379 secB protein-export membrane protein (B-Hin, 37.7%) RP114 secE preprotein translocase SecB subunit (B-Hin, 37.7%) RP114 secE protein-export membrane protein (B-Hin, 37.7%) RP39 sec7 protein-export membrane protein (B-Fin, 37.7%) RP39 sec7 protein-export membrane protein (B-Fin, 37.7%) RP39 sec7 preprotein translocase Sec7 subunit (B-Eco, 50.0%) trigger factor (B-Eco, 32.0%) ENERGY METABOLISM ATP-proton motive force interconversion RP803 atpA ATP synthase F1 alpha subunit (B-Riu, 24.5%) RP301 atpD ATP synthase F0 subunit (B-Riu, 24.5%) RP302 atpA ATP synthase F1 beta subunit (B-Riu, 24.5%) RP302 atpA ATP synthase F0 subunit (B-Riu, 31.7%) RP303 atpA ATP synthase F1 beta subunit (B-Riu, 31.5%) RP304 atpH ATP synthase F1 beta subunit (B-Riu, 31.5%) RP304 atpH ATP synthase F1 beta subunit (B-Riu, 31.5%) RP304 atpH ATP synthase F1 delfa chain (E-Osi cp 26.4%) RP304 atpH ATP synthase F1 delfa chain (E-Osi cp 26.4%) RP304 atpH ATP synthase F1 delfa chain (E-Osi cp 26.4%)	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bat, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabb 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rme, 64.8%) RP772 fabH 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rme, 64.8%) RP373 fabl anoyl-facyl-carrier-protein] synthase (B-Ra, 47.3%) reductase (B-Ag, 49.0%) fabZ 3R-rydcraymyristoyl acyl carrier protein dehy dratase (B-Fin, 91.7%) reductase (B-Ag, 49.0%) fabZ 3R-rydcraymyristoyl acyl carrier protein dehy dratase (B-Fin, 91.7%) reductase (B-P6, 54.5%) reductase (B-Fin, 51.7%) reductas
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP390 nlpD lipoprotein (B-Hin, 22.4%) RP390 nlpD rlpA rate lipoprotein (B-Hin, 22.4%) RP244 tglC are lipoprotein (B-Hin, 22.4%) RP245 lglC are membrane protein (B-Eco, 22.9%) RP246 lglC are membrane protein (B-Eco, 22.9%) RP399 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP249 ddlB D-Alanine:D-alanine ligase (B-Hin, 32.8%) RP458 mraY2 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Hin, 34.9%) RP857 mra (B-Sac, 22.0%) RP807 mra (B-Sac, 22.0%) RP807 mra (B-Sac, 25.6%) RP248 murB UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Bca, 25.6%) RP247 mur RP48 murB UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP410 murD UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP599 mur UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Hin, 32.9%) RP599 mur UDP-MurNac-pentapeptide synthetase (B-Bsu, 35.2%) RP599 mur UDP-MurNac-pentapeptide synthetase (B-Bsu, 35.9%) RP590 mur UDP-MurNac-pentapeptide transferase (B-Bsu, 35.9%) RP412 murG UDP-MurNac-pentapeptide transferase (B-Bsu, 35.9%) RP412 murG UDP-MurNac-pentapeptide transferase (B-Bsu, 35.9%) RP412 murG UDP-MurNac-pentapeptide transferase (B-Bsu, 35.9%)	Detoxification RP535 sodB Superoxide dismutase (B-Lpn, 53.4%) RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lapA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lapB signal peptidase (B-Sty, 37.3%) RP575 secA preprotein transiocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein transiocase SecA subunit (B-Hin, 30.7%) RP576 secD protein-export membrane protein (B-Eco, 40.4%) RP194 secE preprotein transiocase SecA subunit (B-Hin, 37.7%) RP579 secG protein-export membrane protein (B-Hin, 37.7%) RP580 secD protein-export membrane protein (B-Hin, 37.7%) RP580 secG protein-export membrane protein (B-Rin, 31.5%)	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bat, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP365 fabl enoyl-facyl-carrier-protein] synthase (B-Rme, 64.8%) RP373 fadA aceyl-CoA, aceyl-transferase (B-Pde, 54.5%) RP365 fabl enoyl-facyl-carrier-protein] synthase (B-Rn, 91.7%) radA aceyl-CoA aceyl-transferase (B-Pde, 54.5%) RP442 gpsA glycerol-3-phosphate dehydrogenase (B-Eox, 32.1%) RP373 fadA acetyl-CoA aceyl-transferase (B-Pde, 54.5%) RP469 gpsA glycerol-3-phosphate dehydrogenase (B-Eox, 32.1%) RP373 phbC1 poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) RP580 phbC2 poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) RP580 ppsC1 phosphatidylglycerophosphates A (B-Hin, 31.9%) RP489 pisC 1-2-y-glycerol-3-phosphate acyltransferase (E-Esx, 23.6%) RP499 pisC 1-2-y-glycerol-3-phosphate acyltransferase (E-Exa, 23.6%) RP499 pisC 1-2-y-glycerol-3-phosphate acyltransferase (E-Exa, 23.6%) Phosphatidylsphrene synthase (B-Hpy, 28.8%) maic enzyme (B-Hin, 45.5%) and caryeng (B-Hin, 45.5%) Vaccil ipportein precursor (B-Shi, 33.8%) PURINES, PYRIMIDINES 14
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-30, 19.3%) RP346 lgtD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 lnt pl.3%) RP390 nlpD prolipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 22.4%) rare lipoprotein (B-Hin, 22.9%) RP244 vldC lare lipoprotein (B-Hin, 22.9%) RP399 dacF penicillin binding protein (B-Hin, 30.4%) Murein sacculus Allarine racemace (B-Hin, 29.5%) RP399 dacF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddiB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP249 ddiB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP399 maY1 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Hin, 34.9%) RP399 maY2 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-B-Aca, 51.6%) RP807 mrcA logical penicillin binding protein 11 (B-Eco, 35.6%) RP599 murB UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Aca, 51.6%) RP247 murC lDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP410 murD UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP599 murE UPD-MurNac-pentapeptide synthetase (B-Bsu, 35.2%) RP599 murF UPD-MurNac-pentapeptide synthetase (B-Bsu, 38.8%) RP591 murG UDP-MurNac-pentapeptide transferase (B-Bsu, 28.8%) RP595 bbb41 penicillin binding protein (B-Hin, 34.3%)	Detoxification RP535 sodB Superoxide dismutase (B-Lpn, 53.4%) RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rac, 51.8%) RP370 secB preprotein translocase SecA subunit (B-Hin, 30.7%) RP578 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP379 secB protein-export membrane protein (B-Hin, 37.7%) RP114 secE preprotein translocase SecB subunit (B-Hin, 37.7%) RP114 secE preprotein translocase SecB subunit (B-Hin, 37.7%) RP39 sec7 preprotein translocase SecP subunit (B-Bco, 50.0%) trigger factor (B-Eco, 32.0%) RP842 tig trigger factor (B-Eco, 32.0%) RP842 tig trigger factor (B-Eco, 32.0%) RP858 app A TP synthase F1 alpha subunit (B-Rru, 66.2%) RP803 alph ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP801 alph ATP synthase F1 beta subunit (B-Rru, 24.5%) RP802 alph ATP synthase F1 beta subunit (B-Rru, 24.5%) RP803 alph ATP synthase F0 subunit (B-Rru, 24.5%) RP804 alph ATP synthase F0 subunit (B-Rru, 21.1%) RP805 alph ATP synthase F1 beta subunit (B-Ru, 31.2%) RP806 alph ATP synthase F1 beta subunit (B-Ru, 31.5%) RP807 alph ATP synthase F1 delta chain (E-Dsi op 26.4%) RP808 cmE cytochrome c biogenesis protein (B-Hin, 33.2%) RP406 coxA cytochrome c biogenesis protein (B-Hin, 33.6%) RP405 coxA cytochrome c biogenesis protein (B-Hin, 80.9%)	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate oytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bat, 40.3%) RP764 fabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabb 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabb 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rme, 64.8%) RP772 fabh 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rme, 64.8%) RP365 fabl fabz 3-R-ya(roxymyristoyl acyl carrier protein dehy dratase (B-Fin, 19.1%) Accely-CoA acelyl-transferase (B-Pc/e, 54.5%) acelyl-CoA acelyl-transferase (B-Pc/e, 54.5%) RP404 gry Accelyl-CoA acelyl-transferase (B-Eoz, 52.1%) RP365 fabb acetoacetyl-CoA reductase (B-Zra, 52.9%) RP367 pppA brosphatidyl-glycerophosphates (B-Eox, 52.7%) RP378 phbC1 poly-beta-hydroxybutyrate polymerase (B-Eox, 37.4%) RP50 pppA phosphatidyl-glycerophosphates A (B-Hin, 31.9%) RP404 ppsA phosphatidyl-glycerophosphates (B-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Bsu, 40.1%) Phosphatidyl-glycerophosphates (B-Hin, 31.9%) Phosphatidyl-glycerophosphate (B-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate deaminase (B-Hin, 21.9%) Vac.II glycorol-in procursor (B-Sii, 33.8%) PURINES, PYRIMIDINES. 14
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP446 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP360 nlpD protein (B-Hin, 22.4%) RP300 nlpD protein (B-Hin, 22.4%) RP300 nlpD protein (B-Hin, 22.4%) RP300 nlpD protein (B-Hin, 23.9%) RP244 (gtD) outer membrane protein (B-Eco, 22.9%) RP245 (gtD) outer membrane protein (B-Eco, 22.9%) RP246 gtdD prolipoprotein (B-Hin, 29.5%) RP309 alcr pentillih binding protein precursor (B-Bsu, 33.8%) RP249 ddlB D-alanine:D-alanine ligase (B-Hin, 32.8%) RP309 mr31 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Hin, 49.9%) RP309 mr31 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Rac, 35.6%) RP309 mr42 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Rac, 35.6%) RP309 mr42 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Rac, 35.6%) RP309 mr42 UDP-N-acetylglucosamine 1-carboxyvinyltransferase (B-Rac, 35.6%) RP309 mr42 UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP309 mr42 UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP309 mr42 UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Hin, 32.9%) RP309 mr45 UDP-MurNac-pentapeptide synthetase (B-Bsu, 38.5%) RP309 mr47 UDP-MurNac-pentapeptide synthetase (B-Bsu, 28.8%) RP309 ppoticillih binding protein (B-Hin, 34.3%) RP309 ppoticillih binding protein (B-Hin, 34.3%) RP309 ppoticillih binding protein (B-Bsu, 30.7%) RP309 ppoticillih binding protein (B-Bsu, 30.7%) RP309 ppoticillih binding protein (B-Bsu, 30.7%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) RP769 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP175 tepA GTP-binding membrane protein (B-Bsu, 57.0%) RP575 secA preprotein translocase SecA subunit (B-Ra, 51.8%) RP575 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecB subunit (B-Hin, 37.7%) RP577 secP protein-export membrane protein (B-Bsu, 37.3%) RP114 secE protein-export membrane protein (B-Hin, 37.7%) RP779 sec2 preprotein translocase SecC subunit (B-Bsu, 37.3%) RP39 secY preprotein translocase SecC subunit (B-Bsu, 37.3%) RP679 sec2 preprotein translocase SecY subunit (B-Ra, 51.6%) RP679 sec2 preprotein translocase SecY subunit (B-Ra, 37.7%) RP679 sec2 preprotein translocase SecY subunit (B-Ra, 37.7%) RP679 sec2 preprotein translocase SecY subunit (B-Ra, 37.5%) RP800 app ATP synthase F1 alpha subunit (B-Ra, 56.0%) RP670 app ATP synthase F1 alpha subunit (B-Ra, 56.0%) RP670 app ATP synthase F1 alpha subunit (B-Ra, 71.0%) RP800 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP801 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP802 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP803 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP804 alph ATP synthase F1 beta subunit (B-Ra, 71.0%) RP805 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP806 alph ATP synthase F1 beta subunit (B-Ra, 71.0%) RP807 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP808 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%) RP809 app ATP synthase F1 beta subunit (B-Ra, 71.0%)	RP533 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cd24 phosphatidate cyldylytransferase (B-Púe, 33.6%) RP735 tabb maiony-CoA:Acyl carrier protein transacylase (B-Bút, 40.3%) RP764 tabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eco, 31.5%) RP762 tabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Bút, 47.3%) RP762 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rine, 54.8%) RP768 tabH 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rine, 47.3%) renoyl-lacyl-carrier-protein] synthase (B-Rine, 47.3%) renoyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) RP760 tabB 2 RP404 tabbe (B-Rin, 91.7%) renoyl-dacyl-carrier-protein] reductase (B-Asp. 49.0%) renoyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) renoyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) renoyl-dacyl-carrier-protein] reductase (B-Asp.
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 lgtD 19.3%) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) apolipoprotein (B-Hin, 29.4%) prolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.4%) prolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.5%) RP284 lgtC cuter membrane protein (B-Eco, 29.2%) RP284 lgtC cuter membrane protein (B-Eco, 29.2%) groups dars apoliporotein A (B-Hin, 29.5%) RP289 lgt alanine racemace (B-Hin, 29.5%) RP309 lgt alanine racemace (B-Hin, 39.8%) RP309 lgt alanine racemacemacemacemacemacemacemacemacemacem	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP34 aprE RP373 th signal recognition particle receptor protein (B-Eco, 49.0%) RP173 th signal recognition particle receptor protein (B-Eco, 49.0%) RP175 lepA GTTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP070 secB preprotein translocase SecB subunit (B-Hin, 30.7%) RP586 secD protein-export membrane protein (B-Eco, 40.4%) RP114 secE preprotein translocase SecS subunit (B-Bsu, 37.3%) RP079 secG protein-export membrane protein (B-Hin, 37.7%) RP079 secG protein-export membrane protein (B-Hin, 37.7%) RP079 secG protein-export membrane protein (B-Hin, 37.7%) RP380 apr ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP842 tig translocase SecP subunit (B-Rru, 66.2%) RP803 app ATP synthase F1 signal subunit (B-Rru, 24.5%) RP801 app ATP synthase F1 beta subunit (B-Rru, 24.5%) RP802 app ATP synthase F0 subunit (B-Rru, 21.1%) RP803 app ATP synthase F0 subunit (B-Rru, 21.1%) RP804 app ATP synthase F0 subunit (B-Rru, 21.1%) RP805 app ATP synthase F0 subunit (B-Rru, 21.1%) RP804 app ATP synthase F0 subunit (B-Rru, 21.1%) RP804 app ATP synthase F0 subunit (B-Rru, 21.1%) RP805 app ATP synthase F0 subunit (B-Rru, 21.1%) RP806 app ATP synthase F0 subunit (B-Rru, 21.1%) RP807 app ATP synthase F0 subunit (B-Rru, 21.1%) RP808 come cytochrome c biogenesis protein (B-Hin, 33.2%) RP909 cox A cytochrome c biogenesis protein (B-Hin, 33.2%) RP405 cox A cytochrome c oxidase subunit II (E-Rpm mt, 48.9%) RP207 cox AV cytochrome c oxidase subunit II (E-Rpm mt, 48.9%) RP208 cox AV cytochrome c oxidase subunit II (E-Rpm mt, 48.9%) RP209 cox AV cytochrome c oxidase subunit II (E-Rpm mt, 48.9%) RP209 cox AV cytochrome coxidase subunit II (E-Rpm mt, 48.9%)	RP533 birá birá hotornárca carboxylase synthase (B-Púe, 33.6%) RP424 cdz A phosphatidate cyldylytransferase (B-Púe, 33.6%) RP735 tabb malony-CoA:Acyl carrier protein transacylase (B-Bút, 40.3%) RP764 tabF 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eco, 31.5%) RP762 tabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Bút, 47.3%) RP762 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rine, 54.8%) RP772 tabH 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rine, 47.3%) RP763 tabH 3-xxxacyl-lacyl-carrier-protein] synthase (B-Rine, 47.3%) RP763 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) RP763 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) RP763 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Asp. 49.0%) RP764 tabH 25 reductase (B-Rine, 47.3%) RP764 tabH 25 reductase (B-Rine, 47.3%) RP765 tabH 3-xxxacyl-carrier-protein] reductase (B-Pde, 54.5%) reductase (B-Rine, 47.3%) RP760 pppA phosphatidation complex a subunit (E-Cal, 30.6%) gpt-protein diacylglycerol (B-Eco, 30.1%) acetoacetyl-CoA reductase (B-Zra, 52.9%) RP789 phbC1 poly-beta-hydroxybutyrate polymerase (B-Cvi, 22.7%) RP860 phbC2 poly-beta-hydroxybutyrate polymerase (B-Mex. 37.4%) PP760 pppA phosphatidylglycerophosphate synthase (B-Bsu, 40.1%) RP619 plsC 1-acyl-glycerol-3-phosphate acyltransferase (E-Sea, 23.6%) PpspA phosphatidylserine synthase (B-Bsu, 40.1%) Acyl-glycerol-3-phosphate synthase (B-Bsu, 40.1%) PPRMIDINES PPRMIDINES PRMIDINES PRMIDINES PPRMIDINES
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 lgtD 19.3%) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) apolipoprotein (B-Hin, 29.4%) respectively. Prolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) rare lipoprotein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Hin, 39.8%) respectively. Provided in membrane protein (B-Bsu, 33.8%) respectively. Provided in membrane protein (B-Bsu, 33.8%) respectively. Provided in membrane protein (B-Bsu, 33.8%) respectively. Provided in membrane protein (B-Bsu, 35.7%) respectively. Provided in membrane protein (B-Bsu, 30.7%) report provided protein (B-Bsu, 30.7%) report provided protein (B-Bsu, 30	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 40.0%) RP315 aprB RP375 lepA GTP-binding membrane protein (B-Bae, 32.4%) Signal recognition particle receptor protein (B-Eco, 32.4%) RP375 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sy, 37.3%) RP575 secD protein-export membrane protein (B-Bco, 40.0%) RP575 lepA GTP-binding membrane protein (B-Bsu, 57.0%) RP1680 secD protein-export membrane protein (B-Hin, 30.7%) RP576 secD protein-export membrane protein (B-Hin, 30.7%) RP586 secD protein-export membrane protein (B-Hin, 37.7%) RP114 secE protein-export membrane protein (B-Hin, 37.7%) RP134 secE protein-export membrane protein (B-Hin, 37.7%) RP134 secE protein-export membrane protein (B-Hin, 37.7%) RP680 secY preprotein translocase SecY subunit (B-Eco, 50.0%) trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion RP803 atpA ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP023 atpB ATP synthase F1 solpha subunit (B-Rru, 24.5%) RP801 atpD ATP synthase F1 beta subunit (B-Rru, 24.5%) RP802 atpG ATP synthase F1 beta subunit (B-Rru, 24.5%) RP803 atpA ATP synthase F0 subunit (B-Rru, 21.1%) RP804 atpH ATP synthase F0 subunit (B-Rru, 21.1%) RP805 atpF ATP synthase F0 subunit (B-Rru, 21.1%) RP806 atpF ATP synthase F0 subunit (B-Rru, 33.2%) RP807 atpF ATP synthase F0 subunit (B-Rru, 33.2%) RP808 comE cytochrome c biogenesis protein (B-Hin, 33.2%) RP809 cox A cytochrome c biogenesis protein (B-Hin, 33.2%) RP405 cox A cytochrome c oxidase subunit II (F-Mp mt, 48.0%) RP207 cox A cytochrome c oxidase subunit II (F-Mp mt, 48.0%) RP208 cox Cytochrome c oxidase assembly (E-Sca, 35.2%) RP304 cox Cytochrome c oxidase assembly (E-Rram mt, 49.9%) RP304 cox Cytochrome c oxidase assembly (E-Rram mt, 49.9%) RP304 cox Cytochrome c oxidase assembly (E-Rram mt, 49.9%) RP304 cox Cytochrome	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate cytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bat, 40.3%) RP764 fabF 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabb 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP772 fabr 3-xxxacyl-[acyl-carrier-protein] synthase (B-Rme, 54.8%) RP772 fabr 3-xxxacyl-[acyl-carrier-protein] synthase (B-Rme, 54.8%) RP373 fabr 3-xxxacyl-[acyl-carrier-protein] synthase (B-Rxa, 47.9%) RP305 fabb 4 fabz 3-xxxacyl-[acyl-carrier-protein] synthase (B-Asp, 49.0%) RP737 fadA aceyl-CoA aceyltransferase (B-P6, 54.5%) RP580 fadB fatty oxidation complex a subunit (E-Cel, 30.6%) glycerol-3-phosphate dehydrogenase (B-Ecx, 32.1%) RP738 phbC1 poly-beta-hydroxybutyrate polymerase (B-Cxi, 22.7%) RP30 phbC2 poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) RP580 pph Birch poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) Phosphatidyliglycerophosphates A (B-Hin, 31.9%) phosphatidyliglycerophosphates (B-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate acyltransferase (E-Sex, 23.6%) Phosphatidyliglycerophosphate (B-Bsu, 40.1%) 1-acyl-glycerol-3-phosphate (B-Bsu, 40.1%) 1-acyl-glycer
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 lgtD 19.3%) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) apolipoprotein (B-Hin, 29.4%) respectively. Prolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) rare lipoprotein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Hin, 29.5%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Eco, 29.2%) respectively. Provided in membrane protein (B-Bsu, 39.8%) respectively. Provided in membrane protein (B-Bsu, 39.8%) respectively. Provided in membrane protein (B-Bsu, 39.8%) respectively. Provided in membrane protein (B-Eco, 35.6%) respectively. Provided in membrane protein (B-Eco, 35.6%) respectively. Provided in membrane protein (B-Bsu, 35.7%) respectively. Provided in membrane protein (B-Bsu, 35.2%) respectively. Provided in membrane protein (B-Bsu, 30.5%) respectively. Provided in membrane protein (B-Bsu, 30.5%) report protein protein (B-Bsu, 30.7%) report protein respectively. Protein (B-Bsu, 30	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP313 th signal recognition particle receptor protein (B-Eco, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 32.4%) RP375 lepA GTT-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sty, 37.3%) RP575 secD protein-export membrane protein (B-Eco, 40.4%) RP375 secD protein-export membrane protein (B-Eco, 40.4%) RP576 secD protein-export membrane protein (B-Hin, 30.7%) RP576 secD protein-export membrane protein (B-Hin, 30.7%) RP586 secD protein-export membrane protein (B-Hin, 37.7%) RP586 secV preprotein translocase SecS subunit (B-Eco, 50.0%) RP342 tig trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion RP803 atpA ATP synthase F1 slipha subunit (B-Rru, 66.2%) RP023 atpB ATP synthase F1 slipha subunit (B-Rru, 66.2%) RP023 atpB ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP800 atpD ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP801 atpD ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP802 atpF ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP020 atpF ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP020 atpF ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP021 atpD ATP synthase F1 slipha subunit (B-Rru, 24.5%) RP022 atpF ATP synthase F0 subunit (B-Rru, 21.1%) RP804 atpH ATP synthase F1 slipha subunit (B-Rh, 33.2%) RP025 atpM ATP synthase F1 slipha subunit (B-Rh, 33.2%) RP030 accord cytochrome c biogenesis protein (B-Hin, 33.2%) RP030 accord cytochrome c biogenesis protein (B-Hin, 33.2%) RP0405 coxA cytochrome coxidase subunit II (E-Mp mt, 48.9%) RP261 coxA cytochrome coxidase subunit II (E-Rm mt, 28.6%) RP263 cydA cytochrome coxidase assembly (E-Cox, 35.2%) RP264 cydA cytochrome coxidase assembly (E-Fam mt, 49.9%) RP264 cydA cytochrome coxidase assembly (E-Fam mt, 49.9%) RP264 cydA cytochrome coxidase	RP633 birá RP424 cdz Aphosphatidate cyldylytransylrase (B-Púe, 33.6%) RP735 tabD RP764 tabF RP764 tabF RP762 fabG RP763 fabG RP764 fabC RP764 fabC RP764 fabC RP765 fabG RP765 fabG RP765 fabG RP765 fabG RP766 fabG RP767 fabC RP767 fabC RP767 fabC RP767 fabC RP767 fabC RP768 fabG RP768 fabG RP768 fabG RP768 fabG RP768 fabG RP769 fabG RP769 fabG RP760
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 lgtD 19.3%) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) apolipoprotein (B-Hin, 29.4%) respectively apolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.4%) rare lipoprotein (B-Hin, 29.9%) respectively apolipoprotein (B-Hin, 29.9%) respectively adaptively apolipoprotein (B-Hin, 29.9%) respectively adaptively respectively respecti	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTT-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sty, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP586 secA protein-export membrane protein (B-Eco, 40.4%) RP134 secE preprotein translocase SecA subunit (B-Bsu, 37.3%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP587 secA protein-export membrane protein (B-Hin, 37.7%) RP689 secY preprotein translocase SecY subunit (B-Eco, 50.0%) trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion RP803 atpA ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP022 atpB ATP synthase F1 subunit (B-Rru, 66.2%) RP022 atpB ATP synthase F1 beta subunit (B-Rru, 24.5%) RP801 atpD ATP synthase F1 beta subunit (B-Rru, 24.5%) RP802 atpF ATP synthase F0 subunit (B-Rru, 21.1%) RP802 atpF ATP synthase F0 subunit (B-Rru, 21.1%) RP803 atpA ATP synthase F0 subunit (B-Rru, 31.2%) RP804 atpH ATP synthase F0 subunit (B-Rru, 31.2%) RP805 atpM ATP synthase F0 subunit (B-Rh, 33.2%) RP806 atpH ATP synthase F0 subunit (B-Rh, 33.2%) RP807 atpM ATP synthase F0 subunit (B-Rm mt, 28.6%) RP508 come cytochrome c biogenesis protein (B-Hin, 33.2%) RP509 coxA cytochrome coxidase subunit (II (E-Rim mt, 28.6%) RP518 coxA cytochrome coxidase assembly (E-Fim mt, 48.9%) RP530 cytochrome coxidase assembly (E-Fim mt, 48.9%) RP531 cytochrome coxidase assembly (E-Fim mt, 48.9%) RP525 cytochrome coxidase dsubunit (II (B-Exo, 30.0%) RP527 coxW cytochrome coxidase	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cd24 phosphatidate cyldylytransferase (B-Eo. 31.5%) RP765 tabD malonyl-CoA:Acyl carrier protein transacylase (B-Bát, 40.3%) RP762 tabG 3-xxxacyl-lacyl-carrier-protein] synthase II(B-Eo. 35.5%) RP762 tabG 3-xxxacyl-lacyl-carrier-protein] reductase (B-Bát, 47.3%) RP772 tabH 3-xxxacyl-lacyl-carrier-protein] reductase (B-Rm, 54.8%) RP773 tabH 3-xxxacyl-tacyl-carrier-protein] synthase (B-Ra, 47.3%) RP780 tabE 2 RF-lydroxymyristoyl acyl carrier protein dehy dratase (B-Rn, 91.7%) acelyl-CoA acelytitransferase (B-Pde, 54.5%) acelyl-CoA acelytitransferase (B-Pde, 54.5%) acelyl-CoA acelytitransferase (B-Pde, 54.5%) RP737 tadA acelyl-CoA acelytitransferase (B-Pde, 54.5%) acelyl-CoA acelytitransferase (B-Pde, 54.5%) RP789 phbCl poly-beta-hydroxybutyrate polymerase (B-Eco, 32.1%) aceloacelyl-CoA reductase (B-Zra, 52.9%) RP789 phbCl poly-beta-hydroxybutyrate polymerase (B-Cvi, 22.7%) RP820 phbC2 poly-beta-hydroxybutyrate polymerase (B-Mex, 37.4%) RP750 pgpA phosphatidylglycerophosphate synthase (B-Bsu, 40.1%) RP499 plsC 1-acyl-glycerol-3-phosphate acyltransferase (E-Esc, 23.6%) RP333 tme malic enzyme (B-Hin, 45.5%) Table phosphatidylglycerophosphate synthase (B-Bsu, 40.1%) Punithes, PYRIMIDINES. PYRIMIDINES. 14 Deaxyribonucleetide metabolism deoxycytidine triphosphate tupinosphotydrolase (B-Eco, 30.1%) deoxyguanosine triphosphate tupinosphotydrolase (B-Eco, 30.1%) deoxyguanosine triphosphate nucleotidohydrolase (B-Eco, 30.1%) reductase (B-Bsu, 56.4%) nucleoside diphosphate kinase (B-Hin, 42.9%) thymidylate kinase (B-Blin, 40.9%)
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP466 [att. B. 19.3%) RP390 nlpD lipoprotein (B-Hin, 22.4%) RP390 nlpD lipoprotein (B-Hin, 22.4%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP248 in a lanine racemace (B-Hin, 29.5%) RP048 yidC inner membrane protein (B-Eco, 22.9%) RP390 nlpD lipoprotein Na-evitransierase (B-Hin, 30.4%) Murein sacculus RP095 air alanine racemace (B-Hin, 29.5%) RP390 dis penicillin binding protein precursor (B-Bsu, 33.8%) RP249 dis D-alanine ligase (B-Hin, 32.8%) RP454 gimU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mraYt phospho-N-acetylmuramoyl-pentapeptide-transierase (B-Hin, 39.6%) RP807 mrcA penicillin binding protein 1A (B-Eco, 35.6%) UDP-N-acetylglucosamine 1-carboxyvinyltransierase (B-Aca, 51.6%) RP840 mur DP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP597 mur UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP597 mur UDP-N-acetylmuramoylalanine ligase (B-Hin, 32.9%) RP598 mur UDP-N-acetylmuramoylalanine ligase (B-Hin, 32.9%) RP599 penicillin binding protein (B-Eco, 26.7%) RP599 pbp-penicillin binding protein (B-Bsu, 30.7%) RP590 pbp-penicillin bin	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 22.4%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP173 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 lepA GTR-binding membrane protein (B-Bsu, 57.0%) RP116 lepB signal peptidase (B-Sty, 37.3%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecB subunit (B-Hin, 30.7%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP587 secA preprotein translocase SecB subunit (B-Bca, 40.4%) RP588 secA protein-export membrane protein (B-Hin, 37.7%) RP589 secA protein-export membrane protein (B-Hin, 37.7%) RP693 secA protein-export membrane protein (B-Hin, 37.7%) RP694 tig trigger factor (B-Eco, 32.0%) ENERGY METABOLISM 67 ATP-proton motive force interconversion RP803 atpA ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP022 atpB ATP synthase F1 subunit a (B-Rru, 51.5%) RP800 atpA ATP synthase F1 beta subunit (B-Rru, 24.5%) RP801 atpD ATP synthase F1 beta subunit (B-Rru, 24.5%) RP802 atpB ATP synthase F1 beta subunit (B-Rru, 21.1%) RP803 atpA ATP synthase F0 subunit b (B-Rru, 21.1%) RP804 atpA ATP synthase F0 subunit b (B-Rru, 21.1%) RP804 atpA ATP synthase F0 subunit b (B-Rru, 21.1%) RP805 atpA ATP synthase F1 beta subunit (B-Rb, 30.7%) RP806 atpA ATP synthase F1 beta subunit (B-Rb, 30.7%) RP807 atpA ATP synthase F0 subunit b (B-Rru, 21.1%) RP808 atpA ATP synthase F0 subunit b (B-Rru, 21.1%) RP809 (B-Rru, 24.5%) RP809 (B-Rru, 24.5	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate cytidythransferase (B-Eoz, 31.5%) RP735 fabb maionyl-CoA:Aoyl carrier protein transacylase (B-Bút, 40.3%) RP762 fabb acos (B-Bút, 40.3%) RP772 fabb acos (B-Bút, 40.3%) RP772 fabb acos (B-Bút, 40.3%) RP772 fabb acos (B-Bút, 40.3%) RP773 fab acos (B-Bút, 40.3%) RP773 phbc1 followylasta (B-Bút, 40.3%) PR973 phbc2 polybeta-hydroxybutyrate polymerase (B-Eco, 32.1%) Pr019ph phocy p
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins RP347 asmA outer membrane assembly protein (B-Eco, 19.3%) RP466 (gtD) Prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP366 Int RP300 nlpD lipoprotein (B-Hin, 22.4%) RP390 nlpD lipoprotein (B-Hin, 23.9%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP284 yidC outer membrane protein (B-Eco, 22.9%) RP458 yidC outer membrane protein (B-Eco, 22.9%) RP458 yidC outer membrane protein (B-Eco, 22.9%) RP458 yidC outer membrane protein (B-Eco, 22.9%) RP459 daCF penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddiB outer acamace (B-Hin, 29.5%) RP454 gimU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mra11 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Fin, 34.9%) RP595 mra12 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bca, 25.2%) RP597 murA UDP-N-acetylmuramoyl-pentapeptide-transferase (B-Bca, 25.8%) RP598 mra12 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bca, 25.8%) RP597 murA UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP597 murC UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP597 murC UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Bu, 35.7%) RP598 murF UDP-MurNac-pentapeptide synthetase (B-Bcu, 35.8%) RP599 murG UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Hin, 32.9%) RP599 murF UDP-MurNac-pentapeptide synthetase (B-Bcu, 35.8%) RP599 phpA1 penicillin binding protein (B-Bsu, 30.7%) RP599 phpA1 penicillin binding protein (B-Bsu, 30.7%) RP599 phpA2 penicillin binding protein (B-Bsu, 30.7%) RP599 phpA2 penicillin binding protein (B-Eco, 26.7%) RP599 phpA2 penicillin binding protein (B-Bsu, 25.2%) RP599 phpA2 penicillin binding protein (B-Bsu, 25.2%) RP599 phpE penicillin binding protein (B-Bsu, 30.7%) RP593 capD (B-Sau, 34.9%) RP593 capD (B-Sau, 34.9%) RP594 capMit (B-Sau, 34.9%) RP595 phpA4 capsular polysaccharides biosynthesis protein (25.8u, 24.9%) RP594 capMit (B-Sau, 24.9%) RP594 capMit (B-Sau, 24.9%)	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 22.4%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP315 fight signal recognition particle receptor protein (B-Eco, 49.6%) RP375 fight signal recognition particle receptor protein (B-Eco, 49.6%) RP376 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP578 secA preprotein translocase SecA subunit (B-Bco, 40.4%) RP348 secE preprotein translocase SecE subunit (B-Bco, 40.4%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP579 secG protein-export membrane protein (B-Hin, 37.7%) RP587 secA protein-export membrane protein (B-Hin, 37.7%) RP588 cecF protein-export membrane protein (B-Hin, 37.7%) RP589 atpA ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 66.2%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP581 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP582 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP583 come continue	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate cytidyltransferase (B-Eoz, 31.5%) RP735 fabD malonyl-CoA:Aoyl carrier protein transacylase (B-Bút, 40.3%) RP762 fabF 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-[acyl-carrier-protein] synthase (B-Eoz, 53.5%) RP772 fabH 3-xxxacyl-[acyl-carrier-protein] synthase (B-Eoz, 53.5%) RP365 fabI enoyl-[acyl-carrier-protein] synthase (B-Asp, 49.0%) recomplex fabC enoyl-[acyl-carrier-protein] synthase (B-Asp, 49.0%) RP365 fabI enoyl-[acyl-carrier-protein] synthase (B-Asp, 49.0%) recomplex fabC enoyl-[acyl-carrier-protein] synthase (B-Eco, 30.1%) protein diacylglycerophosphate synthase (B-Eoz, 31.4%) phospatidylglycerophosphate synthase (B-Asu, 40.1%) racyl-glycerol-3-phosphate acyltransferase (E-Soz, 23.6%) phospatidylglycerophosphate synthase (B-Bsu, 40.1%) phospatidylglycerophosphate synthase (B-Bsu, 40.1%) racyl-glycerol-3-phosphate environmental environmental synthase (B-Hin, 45.5%) phospatidylglycerophosphate fabC environmental
RP753 lysC aspartokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 (gtD) 3.3%) RP346 (gtD) prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) RP360 nlpD lipoprotein (B-Hin, 22.4%) RP390 nlpD lipoprotein (B-Hin, 23.9%) RP224 tolC outer membrane protein (B-Eco, 22.9%) RP284 tolC outer membrane protein (B-Eco, 22.9%) RP284 tolC outer membrane protein (B-Eco, 22.9%) RP048 yldC inner membrane protein, 60 kDa (B-Hin, 30.4%) Murein sacculus RP095 air penicillin binding protein precursor (B-Bsu, 33.8%) RP249 ddlB balanine:D-alanine ligase (B-Hin, 32.9%) RP454 glmU UDP-N-acetylglucoseamine pyrophosphorylase (B-Hin, 34.3%) RP595 mra112 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Hin, 49.9%) RP807 mra22 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bac, 22.0%) RP807 mra24 benefibli binding protein 1A (B-Eco, 35.6%) RP807 mra25 phospho-N-acetylmuramoyl-pentapeptide-transferase (B-Bac, 35.7%) RP248 mur B UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP410 mur D UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP596 mur UDP-N-acetylmuramoylalanine ligase (B-Hin, 41.5%) RP597 mur UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Hin, 32.9%) RP598 mur UDP-N-acetylmuramoylalanine-D-glutamate ligase (B-Hin, 32.9%) RP599 popt penicillin binding protein (B-Hin, 34.9%) RP599 popt penicillin binding protein (B-Hin, 34.9%) RP599 popt penicillin binding protein (B-Hin, 34.9%) RP599 popt penicillin binding protein (B-Bsu, 30.7%) RP599 popt penicillin binding protein (B-Bsu, 30.7%) RP599 popt penicillin binding protein (B-Bsu, 30.7%	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Lin, 34.7%) RP769 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP137 th signal recognition particle receptor protein (B-Eco, 49.6%) RP136 the protein secretion ATP-binding protein (B-Bae, 32.4%) RP137 th signal recognition particle receptor protein (B-Eco, 49.6%) RP275 tepA GTP-binding membrane protein (B-Bsu, 57.0%) RP516 the protein secretion translocase SecA subunit (B-Rca, 51.8%) RP517 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP518 secD protein-export membrane protein (B-Bin, 37.3%) RP519 secF protein translocase SecA subunit (B-Bu, 37.3%) RP134 secF protein translocase SecA subunit (B-Bu, 37.3%) RP519 secY protein-export membrane protein (B-Hin, 37.7%) RP639 secY preprotein translocase SecA subunit (B-Bu, 20.4%) RP639 secY preprotein translocase SecA subunit (B-Bu, 37.3%) RP609 aprotein-export membrane protein (B-Hin, 37.7%) RP609 aprotein translocase SecA subunit (B-Bu, 20.4%) RP609 secY protein-export membrane protein (B-Hin, 37.7%) RP609 aprotein translocase SecA subunit (B-Ru, 46.5%) RP601 app ATP synthase F1 alpha subunit (B-Ru, 46.5%) RP602 app ATP synthase F1 beta subunit (B-Ru, 24.5%) RP603 app ATP synthase F1 beta subunit (B-Ru, 21.1%) RP609 app ATP synthase F1 beta subunit (B-Ru, 21.1%) RP609 app ATP synthase F1 beta subunit (B-Ru, 30.0%) RP601 app ATP synthase F1 beta subunit (B-Ru, 30.0%) RP602 app ATP synthase F1 beta subunit (B-Ru, 30.0%) RP603 app ATP synthase F1 beta subunit (B-Ru, 30.0%) RP604 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP605 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP606 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP607 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP608 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP609 app ATP synthase F1 delta chain (E-Osi op 26.4%) RP609 app ATP synthase F1 delta cha	RP633 birá birá biotín Ac-CoA carboxylase synthase (B-Púe, 33.6%) RP424 cóz A phosphatidate cytidyltransferase (B-Eoz, 31.5%) RP735 fabD malonyl-CoA:Aoyl carrier protein transacylase (B-Bút, 40.3%) RP762 fabF 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP762 fabG 3-xxxacyl-[acyl-carrier-protein] synthase II(B-Eoz, 53.5%) RP772 fabH 3-xxxacyl-[acyl-carrier-protein] synthase (B-Eoz, 53.5%) RP772 fabH 3-xxxacyl-[acyl-carrier-protein] synthase (B-Eoz, 53.5%) RP365 fabI enoyl-[acyl-carrier-protein] synthase (B-Eoz, 53.5%) RP365 fabI enoyl-[acyl-carrier-protein] synthase (B-Ag, 49.0%) facal fabC 3-R-yacyl-acyl-carrier-protein] synthase (B-Ag, 49.0%) reacyl-co-X-acyl-facyl-carrier-protein] synthase (B-Ag, 49.0%) reacyl-co-X-acyl-facyl-carrier-protein] synthase (B-Ag, 49.0%) reacyl-co-X-acyl-facyl-carrier-protein] synthase (B-Ag, 49.0%) reacyl-carrier-protein] reductase (B-Ag, 49.0%) reacyl-carrier-protein] synthase (B-Ag, 49.0%) reacyl-carrier-protein] reductase (B-Ag, 49.0%) reductase (B-Ag, 49.0%) reductase (B-Ag, 49.0%) reacyl-carrier-protein] reductase (B-Ag, 49.0%) reductase (B-Ag, 49.0%) reductase (B-A
RP753 lysC asparlokinase (B-Bst, 37.3%) Membranes and lipoproteins (B-Bst, 37.3%) RP346 lytD prolipoprotein diacylglyceryl transferase (B-Vch, 29.1%) apolipoprotein (B-Hin, 29.4%) apolipoprotein (B-Hin, 29.5%) apolipoprotein (B-Hin, 39.8%) apol	Detoxification RP535 sodB RP759 thdF thiophene and furan oxidizer (B-Hin, 34.7%) Protein and peptide secretion RP315 aprD protease secretion ATP-binding protein (B-Pae, 40.0%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 22.4%) RP314 aprE protease secretion ATP-binding protein (B-Pae, 32.4%) RP315 fight signal recognition particle receptor protein (B-Eco, 49.6%) RP375 fight signal recognition particle receptor protein (B-Eco, 49.6%) RP376 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Rca, 51.8%) RP575 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP576 secA preprotein translocase SecA subunit (B-Hin, 30.7%) RP578 secA preprotein translocase SecA subunit (B-Bco, 40.4%) RP348 secE preprotein translocase SecE subunit (B-Bco, 40.4%) RP586 secA protein-export membrane protein (B-Hin, 37.7%) RP579 secG protein-export membrane protein (B-Hin, 37.7%) RP587 secA protein-export membrane protein (B-Hin, 37.7%) RP588 cecF protein-export membrane protein (B-Hin, 37.7%) RP589 atpA ATP synthase F1 alpha subunit (B-Rru, 66.2%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 66.2%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rru, 24.5%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP580 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP581 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP582 atpA ATP synthase F1 sipha subunit (B-Rin, 31.7%) RP583 come continue	RP633 birá RP424 cdz Aphosphatidate cytidylytrans/erase (B-Eco, 31.5%) RP735 tabD RP764 tabF RP762 tabG RP762

RP155 /	pyrH I	uridylate kinase (B- <i>Syn</i> , 53.3%)	RP384	proS	33.9%) proline-tRNA synthetase (B- <i>Zmo</i> , 51.8%)	48.8%) RP214 abc73 ABC transporter, ATP-binding protein (B-Hin,
REGULA	ATORY	FUNCTIONS	RP783 RP221		seryl-tRNA synthetase (B-Cbu, 47.2%) threonyl-tRNA synthetase (B-Hin, 50.6%)	33.6%) RP387 msbA1 ABC transporter, ATP-binding protein (B-Eco,
RP071 d	czcR 1	histidine kinase sensor protein (B- <i>Eco</i> , 23.2%) transcriptional activator protein (B- <i>Aeu</i> , 35.1%)	RP468 RP556	trpS	tryptophanyl-tRNA synthetase (B-Syn, 48.5%) tyrosyl-tRNA synthetase (B-Bca, 38.7%)	26.2%) RP696 msbA2 ABC transporter, ATP-binding protein (B-Eco,
RP426 6		histidine kinase osmolarity sensor protein (B- Sti, 23.65)	RP687		valyI-tRNA synthetase (A-Mja, 38.3%)	28.2%)
RP294 g RP011 /	gppA nifR3 1	pppGpp phosphohydrolase (B- <i>Hpy</i> , 23.3%) transcriptional activator nitrogen assimilation			no acyl-tRNA modification	Amino acids
RP614 /		protein (B- <i>Abr</i> , 59.0%) histidine kinase nitrogen sensor protein (B- <i>Aca</i> ,	RP208 RP209	fmt	methionyl-tRNA deformylase (B-Eco, 49.4%) methinoyl-tRNA formyltransferase (B-Hin, 41.9%)	RP307 atrc1 cationic amino acid transporter (E-Mmu, 29.7%) RP129 glnP glutamine transport system permease (B-Bsu,
RP562 /		30.6%) transcriptional activator nitrogen assimilation	RP152		glutamyľ-tRNA (Gln) amidotransferase subunit A (B- Mca, 48.6%)	48.6%) RP700 glnQ1 glutamine ABC transporter, ATP-binding protein (B-
		protein (B-Aca, 45.2%)	RP151		glutamyl-tRNA (Gln) amidotransferase subunit B (B- Mca, 46.9%)	Eco, 39.1%) RP868 glnQ2 glutamine ABC transporter, ATP-binding protein (B-
		transcriptional activator protein OmpR (B- <i>Rca</i> , 42.9%)	RP153	gatC	glutamyl-tRNA (Gln) amidotransferase subunit C (B- Bsu, 24.7%)	Bst, 51.0%) RP176 gltP glutamate-aspartate transporter (B-Bca, 35.2%)
		histidine kinase phosphatase synthesis sensor protein (B- <i>Bsu</i> , 24.4%)	RP672 RP510		dimethyladenosine transferase (B-Bsu, 35.7%) tRNA delta-2-isopentenylpyrophosphate (IPP) trans	RP483 potE putrescine-ornithine transporter (B-Hin, 26.9%) RP369 potG putrescine ABC transporter, ATP-binding protein (B-
RP312 8	spoT*	(p)ppGpp 3´-pyrophosphohydrolase (B- <i>Eco</i> , 29.9%) (p)ppGpp 3´-pyrophosphohydrolase (B- <i>Mpn</i> , 27.8%)	RP605		ferase (B- <i>Atu</i> , 30.7%) peptidyl-tRNA hydrolase (B- <i>Hin</i> , 40.5%)	Mpn, 29.2%) RP077 proP1 proline/betain transporter (B-Eco, 26.7%)
RP625 s	spoT* I	p)ppGpp 3'-pyrophosphohydrolase (B- <i>Eco</i> , 48.7%) (p)ppGpp 3'-pyrophosphohydrolase (B- <i>Eco</i> , 31.7%)	RP213		S-adenosylmethionine:tRNA ribosyltransferase-iso	RP313 proP2 proline/betain transporter (B-Eco. 24.9%)
RP517	yhbH :	sigma 54 modulation protein (B- <i>Bja</i> , 26.2%)	RP721	tgt	merase (B-Hin, 43.3%) tRNA-guanine transglycosylase (B-Zmo, 61.2%)	RP375 <i>proP</i> 3 proline/betain transporter (B- <i>Eco</i> , 21.2%) RP685 <i>proP</i> 4 proline/betain transporter (B- <i>Eco</i> , 24.0%)
REPLICA	ATION .	46	RP111		tRNA (guanine-N1)-methyltransferase (B-Eco, 44.7%)	RP755 proP5 proline/betain transporter (B-Eco, 27.8%) RP852 proP6 proline/betain transporter (B-Eco, 34.8%)
Degrada			RP857 RP501		pseudouridylate synthase I (B- <i>Eco</i> , 40.1%) tRNA pseudouridine 5S synthase (B- <i>Hin</i> , 37.6%)	RP881 proP7 proline/betain transporter (B-Eco, 28.7%) RP150 yqiX amino acid ABC transporter (B-Bsu, 32.4%)
RP260 >	xthA1 e	ATP-dependent nuclease (B-Bsu, 23.7%) exodeoxyribonuclease III (B-Eco, 30.1%)			proteins, peptides and glycopeptides	Nucleosides and nucleotides
RP675 >	xseA (exodeoxyribonuclease III (B- <i>Eco</i> , 33.2%) exodeoxyribonuclease large subunit (B- <i>Eco</i> , 31.7%)	RP036		ATP-dependent protease, ATP binding subunit (B- Hin, 54.3%)	RP097 mkl ribonucleotide ABC transporter, ATP-binding protein (B-Mie, 36.2%)
RP350	xseB (exodeoxyribonuclease small subunit (B-Eco, 32.5%)	RP520		ATP-dependent Clp protease (B-Yen, 67%)	RP053 tlc1 ATP/ADP translocase (B-Ctr, 43.3%)
DNA rep	dication,	restriction, modification, recombination and repair chromosomal replication initiation protein DnaA (B-	RP692		ATP-dependent protease, ATPase subunit (B-Eco, 62.8%)	RP377 tlc2 ATP/ADP translocase (B-Ctr, 35.2%) RP477 tlc3 ATP/ADP translocase (B-Ctr, 39.6%)
		Eco, 44.1%)	RP228 RP037		tail-specific protease precursor (B-Bba, 42.6%) sialoglycoprotein endopeptidase (B-Hin, 42.2%)	RP500 tlc4 ATP/ADP translocase (B-Ctr, 36.3%) RP739 tlc5 ATP/ADP translocase (B-Ctr, 34.7%)
RP542 (dnaE	DNA helicase (E- <i>Osi</i> cp, 40.9%) DNA polymerase III alpha subunit (B- <i>St</i> y, 37.2%)	RP123 RP122	ĥflC	lambda cll stability-governing protein (B- <i>Eco</i> , 33.9%) lambda cll stability-governing protein (B- <i>Vpa</i> ,	Carbohydrates, organic alcohols and acids
RP859 (dnaG dnaN	DNA primase (B- <i>Sty</i> , 29.0%) DNA polymerase III beta subunit (B- <i>Ppu</i> , 29.9%)			30.3%)	RP054 glpT glycerol-3-phosphate permease (B-Bsu, 37.1%)
RP732	dnaQ	DNA polymerase III epsilon subunit (B-Sty, 46.7%) DNA polymerase III gamma chain (B-Eco, 31.4%)	RP124 RP186	htrA	serine protease (B-Bhe, 37.7%) protease DO (E-Sce, 26.7%)	Cations
RP206 d	qvrA	DNA gyrase A subunit (B-Rsp, 49.4%)	RP450 RP408	ispA	ATP-dependent protease LA (B-Ccr, 53.1%) lipoprotein signal peptidase (B-Bsu, 27.9%)	RP834 afuC iron ABC transporter, ATP-binding protein (B-Eco, 33.9%)
RP227 (gyrB2	DNA gyrase B subunit (B- <i>Sci</i> , 42.0%) DNA gyrase B subunit (B- <i>Ppu</i> , 51.5%)	RP824 RP219	map	methionyl aminopeptidase (B-Sty, 55.3%) mitochondrial protease (B-Bsu, 35.4%)	RP810 kefB glutathionine-regulated potassium-efflux system pro tein (B-Eco, 33.9%)
RP172 /	- 1	DNA polymerase III, delta prime subunit (B- <i>Pae</i> , 22.3%)	RP142 RP174	pepA	aminopeptidase A (B- <i>Pae</i> , 36.6%) peptidase II (B- <i>Rsn</i> , 32.5%)	RP583 mgtE magnesium transporter (B-Syn, 27.0%)
RP171 / RP720 /	hupA lia	DNA binding protein HU (B- <i>Vpr</i> , 47.8%) DNA ligase (B- <i>Zmo</i> , 45.7%)	RP281	ptrB	protease II (B-Eco, 34.2%)	Other
RP777 / RP598 /	metK* :	S-adenosylmethionine synthetase (B- <i>Eco</i> , 66.3%) transcription-repair coupling factor (B- <i>Hin</i> , 33.9%)	RP398 RP525		protease IV (B- <i>Mja</i> , 23.9%) protease IV (B- <i>Hin</i> , 27.6%)	RP205 atm1 mitochondrial ABC transporter, ATP-binding protein (E-Sce, 43.3%)
RP351 /	mpa	DNA-3-methyladenine glycosidase (E-Hsa. 29.7%)	Protein	modific	ation and translation factors	RP794 ccmA haem ABC transporter A, ATP-binding protein (B- Hin, 35.5%)
RP880 / RP298 /	mutS	DNA mismatch repair protein MutL (B- <i>Spn</i> , 35.4%) DNA mismatch repair protein MutS (B- <i>Bsu</i> , 39.0%)	RP238 RP132	efp	elongation factor P (B-Bsu, 39.5%) elongation factor G (B-Atu, 68.7%)	RP268 ccmB haem exporter protein B (E-Ram mt, 20.9%) RP830 ccmC haem exporter protein C (B-Bja, 43.7%)
RP746 / RP067 /	parC	endonuclease III (B- <i>Eco</i> , 50.7%) DNA topoisomerase IV subunit A (B- <i>Hin</i> , 39.0%)	RP814 RP552	infA	initiation factor IF-1 (B-Hin, 67.1%) initiation factor IF-2 (B-Hin, 42.6%)	RP571 panF panthotenate permease (B-Hin, 20.5%) RP630 perM permease PerM homologue (B-Hin, 25.0%)
RP711 /	pin* i	invertase/recombinase (B- <i>Eco</i> , 38.0%) DNA polymerase I (B- <i>Bca</i> , 37.2%)	RP531	infC	initiation factor IF-3 (B-Pvu, 47.7%)	RP374 sec7 transport protein Sec7 (B-Hsa, 29.6%)
RP540 / RP546 /	priA	primosomal protein replication factor (B- <i>Rru</i> , 39.7%) DNA repair (B- <i>Bsu</i> , 46.5%)	RP529 RP274	prfB	peptide chain release factor RF-1 (B-Bsu, 50.1%) peptide chain release factor RF-2 (B-Eco, 50.4%)	RP576 prsA protein export (B-Bsu, 28.9%)
RP761 /	recA I	recombination protein RecA (B-Pde, 71.2%)	RP435 RP693		ribosome binding factor A (B-Bsu, 31.6%) ribosome protein alanine acetyltransferase (B-Eco,	OTHER CATEGORIES
RP029 /		DNA repair protein, ATP binding protein (B- <i>Ccr</i> , 30.4%)	RP154		23.2%) ribosome recycling factor (B-Hin, 43.3%)	Adaptations to atypical conditions RP708 himA integration host factor α (B-Eco, 29.5%)
RP593 / RP528 /		ATP-dependent DNA helicase (B- <i>Eco</i> , 34.1%) single-stranded DNA-specific exonuclease (B- <i>Eco</i> ,	RP397 RP661	tlpA	thiol:disulphide interchange protein (B- <i>Bja</i> , 27.4%) elongation factor Tu (B- <i>Tcu</i> , 81.5%)	RP236 invA invasion protein A (B-Bba, 42.8%)
RP182 /		32.8%) recombination protein RecN (B- <i>Hin</i> , 31.6%)	RP087		elongation factor Ts (B-Sci, 40.7%)	RP717 taxB* conjugative DNA processing (B-Eco, 33.5%)
RP438 /	recR I	recombination protein RecR (B- <i>Bsu</i> , 36.9%) Holliday junction DNA helicase (B- <i>Pae</i> , 35.0%)			teins; synthesis and modification	RP286 trbG conjugal transfer (B-Rsn, 24.7%) RP103 virB4 virulence protein VIRB4 (B-Atu, 30.9%)
RP386 /	ruvB I	Holliday junction DNA helicase (Pae, 51.5%)	RP137 RP656	rplB	ribosomal protein L1 (B-Cgr, 50.2%) ribosomal protein L2 (E-Ram mt, 61.5%)	RP784 virB4 virulence protein VIRB4 (B-Atu, 20.3%) RP287 virB8 virulence protein VIRB8 (B-Atu, 20.4%)
RP119 /		Holliday junction endodeoxyribonuclease (B- <i>Eco</i> , 36.1%)	RP659 RP658		ribosomal protein L3 (E-Sce, 44.1%) ribosomal protein L4 (B-Bst. 39.3%)	RP290 virB9 virulence protein VIRB9 (B-Atu, 24.8%) RP291 virB10 virulence protein VIRB10 (B-Atu, 20.3%)
RP836 s RP326 i	topA	single-stranded binding protein (B- <i>Bab</i> , 52.6%) DNA topoisomerase I (B- <i>Bsu</i> , 44.9%)	RP647 RP644	rpIE	ribosomal protein L5 (B- <i>Aco</i> , 53.6%) ribosomal protein L6 (B- <i>Bst</i> , 45.4%)	RP292 <i>virB</i> 11 virulence protein VIRB11 (B- <i>Atu</i> , 29.6%) RP293 <i>virD</i> 4 virulence protein VIRD4 (B- <i>Atu</i> , 31.3%)
RP835 (repair excision nuclease subunit A (B- <i>Eco</i> , 57.7%) repair excision nuclease subunit B (B- <i>Hin</i> , 56.0%)	RP041	rpll	ribosomal protein L9 (E-Ppu cp, 33.6%)	
RP572 8	uvrC I	repair excision nuclease subunit C (B- <i>Pfl</i> , 36.9%) DNA helicase (B- <i>Sau</i> , 43.5%)	RP138 RP136	rplK	ribosomal protein L10 (B-Laf, 36.7%) ribosomal protein L11 (E-Ram mt, 45.5%)	Drug and analogue sensitivity RP170 acrD acriflavin resistance protein D (B-Eco, 31.3%)
RP817 >	xerC i	integrase/recombinase (B-Bsu, 32.2%) integrase/recombinase (B-Eco, 37.6%)	RP139 RP233		ribosomal protein L7/L12 (B-Bab, 66.9%) ribosomal protein L13 (B-Sca, 52.8%)	RP475 ampG1AMPG protein (B-Eco, 31.4%) RP668 ampG2AMPG prt (B-Eco, 26.3%)
RP361 2			RP649 RP640	rpiN rpiO	ribosomal protein L14 (B- <i>Eco</i> , 69.6%) ribosomal protein L15 (B- <i>Bst</i> , 46.5%)	RP781 ampG3AMPG prt (B-Eco, 27.6%) RP603 bcrl bicyclomycin resistance (B-Eco, 21.7%)
		DN20	RP652 RP634	rpiP	ribosomal protein L16 (B- <i>Aac</i> , 53.3%) ribosomal protein L17 (B- <i>Eco</i> , 57.5%)	RP698 bcr2 bicyclomycin resistance (B-Eco, 18.8%) RP243 emrA multidrug resistance protein A (B-Eco, 26.9%)
Degrada: RP504	ition of F	RNA polyribonucleotide nucleotidyItransferase (B- <i>Eco</i> ,	RP643	rpIR	ribosomal protein L18 (B-Bst, 58.6%)	RP157 emrB multidrug resistance protein B (B-Hin, 29.3%)
BP117 /		48.9%) ribonuclease III (B- <i>Hp</i> y, 40.2%)	RP112 RP609	rpIT	ribosomal protein L19 (B-Eco, 58.8%) ribosomal protein L20 (B-Psy, 61.5%)	RP786 terC tellurite resistance protein (B-Eco, 35.3%)
RP462 /	rnd i	ribonuclease D (B- <i>Eco</i> , 28.5%) ribonuclease E (B- <i>Eco</i> , 35.9%)	RP751 RP654	rpIV	ribosomal protein L21 (E-Sol, 42.3%) ribosomal protein L22 (B-Eco, 50.0%)	Colicin-related functions RP302 tolB colicin tolerance protein (B-Hin, 29.8%)
RP256 /	rnhA i	ribonuclease HI (B-Msm, 43.4%)	RP657 RP648	rpIW	ribosomal protein L23 (B-Bst, 46.3%) ribosomal protein L24 (B-Bst, 55.9%)	RP309 tolQ inner membrane protein (B-Eco, 39.7%) RP310 tolR inner membrane protein (B-Pae, 40.1%)
RP202 /	rnpA i	ribonuclease HII (B- <i>Eco</i> , 44.7%) ribonuclease P (B- <i>Mca</i> , 28.4%)	RP606 RP752	rplY	ribosomal protein L25 (B-Mtu, 26.9%) ribosomal protein L27 (E-Ram mt, 62.9%)	Uncategorized
RP628 /		ribonuclease PH (B- <i>Hin</i> , 55.05%)	RP099	rpmB	ribosomal protein L28 (B-Mge, 43.7%)	RP493 addA adducin alpha subunit (E-Hsa, 32.6%)
RNA syn	nthesis a	and modification transcription elongation factor GreA (B-Hin, 61.4%)	RP651 RP641	rpmD	ribosomal protein L29 (B-Bst, 39.4%) ribosomal protein L30 (B-Mlu, 33.3%)	RP199 adx1 adrenodoxin precursor (E-Spo, 57.1%) RP714 ank2 ankyrin (E-Hsa, 32.7%)
RP553	nusA 1	transcription termination factor NusA (B- <i>Eco</i> , 36.9%)	RP100 RP773	rpmF	ribosomal protein L31 (B-Mtu, 31.6%) ribosomal protein L32 (B-Hin, 49.1%)	RP245 bolA BolA protein (B-Val, 34.2%) RP181 ctaQ* thermostable carboxypeptidase (B-Pho, 29.1%)
RP162 /	nusB 1	transcription termination factor NusB (B- <i>Eco</i> ,	RP879 RP610	rpmH	ribosomal protein L33 (B-Hin, 51.8%) ribosomal protein L34 (B-Ppu, 65.9%)	RP297 cysQ sulphite synthesis pathway protein (B-Eco, 31.2%) RP323 cyaY CyaY protein (B-Ech, 31.1%)
RP135 /	nusG 1	32.9%) transcription antitermination protein NusG (B- <i>Eco</i> ,	RP608 RP456	rpmi rnm.l	ribosomal protein L35 (E- <i>Ppu</i> cp, 38.5%) ribosomal protein L36 (E- <i>Gth</i> cp, 65.8%)	RP118 era GTP binding protein Era (B-Bsu, 33.6%) RP063 hesB1 HesB protein (B-Ava, 37.0%)
RP015 /	pcnB	42.2%) poly (A) polymerase I (B- <i>Bsu</i> , 26.3%)	RP521	rpsA	ribosomal protein S1 (B-Rme, 48.6%)	RP484 hesB2 HesB protein (B-Pbo, 40.2%)
RP669 /	rhIE .	ATP-dependent RNA helicase (B- <i>Eco</i> , 38.3%) transcription termination factor Rho (B- <i>Rsp.</i> 72.0%)	RP086 RP653	rpsC	ribosomal protein S2 (B-Mtu, 41.5%) ribosomal protein S3 (B-Bst, 54.2%)	RP212 n2B N2B, ATPase protein (E-Hir, 27.2%) RP485 nitU nitrogen fixation protein (B-Avi, 43.0%)
RP635 /	rpoA	RNA polymerase alpha subunit (<i>B-Bpe</i> , 47.2%) RNA polymerase beta subunit (<i>B-Rty</i> , 87.4%)	RP345 RP642	rpsE	ribosomal protein S4 (B-Bsu, 43.2%) ribosomal protein S5 (B-Bst, 50.6%)	RP832 <i>p34</i> P34 protein (B- <i>Bri</i> , 91.3%) RP602 <i>pat1</i> patatin B1 precursor protein (E- <i>Stu</i> , 22.9%)
RP141 /	rpoC	RNA polymerase beta subunit (B-Eco, 58.8%)	RP039 RP131	rpsF	ribosomal protein S6 (B-Hin, 30.2%) ribosomal protein S7 (E-Ram mt, 45.2%)	RP317 pkcl protein kinase C inhibitor (B-Abr, 38.6%) RP109 ptb phosphate buturyltransferase (B-Cab, 36.4%)
RP303 /		RNA polymerase sigma-32 factor (B- <i>Atu</i> , 52.0%) RNA polymerase sigma-70 factor (B- <i>Rca</i> , 50.5%)	RP645 RP234	rpsH	ribosomal protein S8 (B-Bsu, 42.0%) ribosomal protein S9 (B-Bst, 48.8%)	RP594 scoB* succinyl-CoA:3-ketoacid-CoA transferase subunit B (B-Mtu, 22.0%)
TRANSL	ATION	118	RP660 RP636	rpsJ	ribosomal protein S10 (B-Hin, 60.8%)	RP031 sco2 Sco yeast precursor protein (E-Sce, 32.6%)
		A synthetases	RP130	rpsL	ribosomal protein S11 (B-Syn, 53.5%) ribosomal protein S12 (B-Tth, 60.5%)	RP587 sco2 Sco yeast precursor protein (E-Sce, 36.6%) RP846 sfhB SfhB protein (B-Zmo, 40.6%)
RP856 a	alaS a	lanyl-tRNA synthetase (B- <i>Bba</i> , 52.7%) arginyl-tRNA synthetase (B- <i>Hp</i> y, 33.0%)	RP637 RP646	rpsN	ribosomal protein S13 (B- <i>Bst</i> , 58.8%) ribosomal protein S14 (B- <i>Syn</i> , 47.0%)	RP430 <i>smpB</i> small protein (B- <i>Syn</i> , 46.7%) RP058 <i>soj</i> SOJ protein (B- <i>Bsu</i> , 50.4%)
RP145 a	aspS a	aspartyl-tRNA synthetase (B-Syn, 43.3%)	RP503 RP878	rpsO	ribosomal protein S15 (B-Bst, 53.4%)	RP486 spl1 tRNA splicing protein (E-Cal, 58.9%) RP487 spl1 tRNA splicing protein (E-Cal, 32.3%)
	gitX1 g	cysteinyl-tRNA synthetase (B- <i>Hin</i> , 46.0%) glutamyl-tRNA synthetase (B- <i>Abr</i> , 45.6%)	RP650 RP040	rpsQ	ribosomal protein S16 (B- <i>Hin</i> , 45.1%) ribosomal protein S17 (B- <i>Tth</i> , 61.8%) ribosomal protein S18 (B- <i>Syn</i> , 50.7%)	RP059 spood sporulation protein (B-Bsu, 40.4%) RP239 suhB supressor protein (B-Eco, 22.8%)
RP085 (gltX2	glutamyl-tRNA synthetase (B- <i>Hpy</i> , 40.3%) glycyl-tRNA synthetase (B- <i>Mca</i> , 60.6%)	RP655	rpsS	ribosomal protein S19 (B-Eco, 58,2%)	RP733 surf1 SurF1 protein (E-Hsa, 23.9%)
RP325 g RP623 g RP850 d	glyQ 4		RP633		ribosomal protein S20 (B- <i>Rmé</i> , 40.9%) ribosomal protein S21 (B- <i>Bbu</i> , 33.3%)	RP710 tra3* transposase (B-Rme, 34.0%)
RP325 (RP623 (RP850 (RP849 (glyS 9	glycyl-tRNA synthetase (B- <i>Bsu</i> , 32.9%)	RP615	. ,	Tibodomai protom GET (B BBB, GG.G76)	
RP325 (RP623 (RP850 (RP849 (RP308) RP620)	glyS (hisS i ileS i	histidyl-tRNA synthetase (B- <i>Eco</i> , 38.3%) isoleucyl-tRNA synthetase (B- <i>Mtu</i> , 48.6%)			AND BINDING PROTEINS38	HYPOTHETICAL PROTEINS
RP325 (RP623 (RP850 (RP849 (RP308) RP620 (RP421) RP371 (glyS 9 hisS 1 ileS 1 leuS 1 lysS 1	histidyI-IRNA synthetase (B- <i>Eco</i> , 38.3%) isoleucyI-IRNA synthetase (B- <i>Bhtu</i> , 48.6%) leucyI-IRNA synthetase (B- <i>Eco</i> , 45.3%) lysyI-IRNA synthetase (B- <i>Bbu</i> , 26.3%)	TRANS	PORT		Integral membrane proteins 19
RP325 (RP623 (RP850 (RP849 (RP308) RP620 (RP421)	glyS (hisS i ileS i leuS i lysS i metS i pheS	histidyl-tRNA synthetase (B- <i>Eco</i> , 38.3%) isoleucyl-tRNA synthetase (B- <i>Mtu</i> , 48.6%) leucyl-tRNA synthetase (B- <i>Eco</i> , 45.3%)	TRANS	PORT		HYPOTHETICAL PROTEINS 110 Integral membrane proteins 19 110 NO SIMILARITY 208 Integral membrane proteins 21 208