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#### SHORT COMMUNICATION

# Comparative genomic analyses reveal a lack of a substantial signature of host adaptation in *Rhodococcus equi* ("*Prescottella equi*")

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Keywords: *Rhodococcus equi*, "*Prescottella equi*", opportunistic pathogen, genome, metabolic differences

#### Abstract

2

3 Rhodococcus equi ("Prescottella equi") is a pathogenic actinomycete primarily infecting horses but has emerged as an opportunistic human pathogen. We have sequenced the genome 4 of the type strain of this species, R. equi strain  $C7^{T}$ , and compared the genome with that of 5 6 another foal isolate 103S and of a human isolate ATCC 33707. The R. equi strains are closely related to each other and yet distantly related to other rhodococci and Nocardia brasiliensis. 7 8 The comparison of gene contents among R. equi strains revealed minor differences that could be associated with host adaptation from foals to humans, including the presence of a paa 9 operon in the human isolate which is potentially involved in pathogenesis. 10

*Rhodococcus equi*, a Gram-positive coccobacillus that was formerly classified as *Corynebacterium equi*, is primarily a foal pathogen that causes fatal lymphadenitis and ulcerative enteritis in young foals (Prescott, 1991; Giguere, *et al.*, 2011; Vazquez-Boland, *et al.*, 2013). It has recently emerged as an important opportunistic human pathogen causing significant mortality among immunocompromised patients (Kedlaya, *et al.*, 2001; Yamshchikov, *et al.*, 2010) and is also associated with a wide range of other vertebrate hosts and with environmental reservoirs (Prescott, 1991).

In this study, we have sequenced the genome of the *R*. *equi* Type strain  $C7^{T}$  that was isolated 18 from the lung abscess of a foal in 1922 (Goodfellow & Alderson, 1977). Comparative analyses 19 were performed against the available genomes of *R. equi*, namely against those of strains 103S 20 21 (Letek, et al., 2010) and ATCC 33707 (Qin, et al., 2010; GenBank Accession numbers NC\_014659 and NZ\_CM001149, respectively). The *R. equi*  $C7^{T}$  genome was sequenced on a 22 Roche 454 GS-Junior instrument as described previously (Sangal, et al., 2012a; Sangal, et al., 23 2012b). A total of 172,478 reads with an average read length of 464 bp were assembled into 24 48 contigs (>200 bp) by GS de novo assembler v 2.7. The size of assembly was 5.197 MB 25 with an average of 19 fold coverage. The genome was annotated using the NCBI PGAAP 26 pipeline (http://www.ncbi.nlm.nih.gov/genome/annotation\_prok/) and RAST (Aziz, et al., 27 2008). The draft genome of *R. equi* strain  $C7^{T}$  is estimated to have 5,033 features including 28 4,978 coding sequences and 55 tRNA genes. This genome has been submitted to GenBank 29 30 and has the accession number APJC00000000.

The *R. equi* 103S and ATCC 33707 genomes were also annotated using RAST to ensure equivalence of annotation for comparative analyses. The CGView comparative tool analysis (Grant, *et al.*, 2012) revealed a very high conservation of both the nucleotide (Fig. 1A) and the protein sequences (Fig. 1B) among these genomes. Indeed, the genome-wide protein Blast searches, using the SEED Viewer (Aziz, *et al.*, 2012), revealed that only 69, 152 and 187

genes, mostly encoding hypothetical proteins, are specific to strains 103S, C7<sup>T</sup> and ATCC 36 37 33707, respectively that returned no hits in the other *R. equi* genomes (Supplementary Table 1). Some of these strain specific genes belonged to mobile genetic elements (integrase, transposase 38 39 and repeats) and only 10-13 genes are predicted to be involved in cellular activities (Supplementary Table 1). Interestingly, strain ATCC 33707 possessed a set of phenylacetate 40 CoA ligase/oxygenase genes (PaaD, F-I, Z) that are absent in the genomes of strains 103S and 41  $C7^{T}$ . These genes have been implicated in virulence in various pathogens (Law, et al., 2008; 42 Teufel, et al., 2010). Strain ATCC 33707 was isolated from a human abscess and strains 103S 43 and C7<sup>T</sup> from equine hosts. Putative PaaE is common to all three isolates and PaaJ and PaaK 44 subunits of strain ATCC 33707 shared ~30% identities with two proteins (PaaD-like protein 45 DUF59 involved in Fe-S cluster assembly and probable phenylacetic acid degradation NADH 46 oxidoreductase PaaE, respectively; data not shown) in the genomes of strains 103S and  $C7^{T}$ . 47 The genes encoding PaaD and PaaZ have been identified on an horizontally acquired region by 48 the Alien Hunter (Vernikos & Parkhill, 2006) and a lack of these genes in equine isolates 49 50 suggest that paa operon was potentially acquired by strain ATCC 33707 which may be associated with pathogenesis in human hosts. Alternatively, the paa operon may not be 51 essential for pathogenesis in foals and may have been lost in equine isolates through genome 52 reduction. Clearly, a wider analysis of a larger collection of R. equi isolates from humans and 53 foals should be performed to confirm this finding. 54

Strain 103S shares 42 genes with strain C7<sup>T</sup> that are absent from the genome of strain ATCC 33707 and 48 genes with strain ATCC 33707 that are absent from the genome of strain C7<sup>T</sup>. Most of these genes encode hypothetical proteins and only 6-12 of these are potentially involved in cellular activities, including nucleotide/nucleoside (especially RNA) metabolism, carbohydrates and sulphur metabolism, and respiration (Supplementary Table 1). A gene encoding NADH-ubiquinone oxidoreductase chain H (fig|685727.5.peg.1900) and one

encoding a glycosyltransferase (fig|685727.5.peg.4060) are common between the genomes of 61 strains 103S and ATCC 33707 but absent in that of strain C7<sup>T</sup>. Similarly, two genes involved in 62 sulphur metabolism (fig|685727.5.peg.436, fig|685727.5.peg.1276) are common between the 63 genomes of strains C7<sup>T</sup> and 103S but missing from the genome of strain ATCC 33707. Strains 64 C7<sup>T</sup> and ATCC 33707 share 84 genes that are absent from the genome of strain 103S, of which 65 23 are likely involved in a variety of metabolic activities (Supplementary Table 1). In addition, 66 some genes showed poor identity (<50% protein sequence identity) between the different 67 genomes that may also partly contribute to the functional divergence of these strains (data not 68 69 shown). These variations potentially reflect the minor differences in lifestyle of individual strains. Overall, the two equine isolates and the human isolate share most of their gene content 70 71 with 109-258 variable genes between strain pairs (Supplementary Table 1). These results are 72 consistent with a recent report suggesting that R. equi isolates are very closely related to each other (Vazquez-Boland, et al., 2013). 73

Single nucleotide polymorphisms (SNPs) were extracted from the genomic alignment of all 74 three strains using Mauve (Darling, et al., 2004) and were annotated using 103S as the 75 reference genome by TRAMS (Reumerman, et al., 2013), after excluding the SNPs with 76 missing alleles. There were 49,903 SNPs between the genomes of 103S and C7<sup>T</sup>, of which 77 4,976 were intergenic. These SNPs introduced 28,458 synonymous and 12,804 78 nonsynonymous changes in the genome of strain C7<sup>T</sup> including 21 stop codons (Supplementary 79 Table 2). Nine nonsense SNPs were present in genes encoding hypothetical proteins and two 80 nonsense SNPs were present in a gene encoding a MCE family protein, Mce1F (Supplementary 81 Table 2). A third nonsense SNP terminated another MCE family membrane protein. MCE 82 83 family proteins were originally implicated as having an important role in the entry and survival of Mycobacterium tuberculosis inside host cells (Arruda, et al., 1993) although complex 84 phenotypes are associated with mce loci (Clark et al. 2013) and it is likely that the direct 85

function of the MCE proteins is in transport of hydrophobic solutes such as sterols (Mohn *et al.* 2008). Other nonsense SNPs resulted in the inactivation of genes involved in putative ABC transporter ATP binding protein, putative esterase, YndJ, pyruvate oxidase, ribosyl nicotinamide transporter (PnuC-like), polyhydroxyalkanoic acid synthase, epoxide hydrolase, acetate kinase and long-chain fatty acid CoA ligase (Supplementary Table 2). It is not clear whether these changes reflect a difference in ability of this strain to utilize different carbon sources or if they have accumulated over long term culture to adapt to an *in vitro* lifestyle.

There are 43,350 SNPs between the genomes of strains 103S and ATCC 33707 that introduced 93 26,312 synonymous and 10,379 nonsynonymous changes including 19 stop codons 94 (Supplementary Table 2). Seven nonsynonymous changes in genes for an ABC transporter 95 ATP binding protein, a putative esterase, an acetate kinase and four hypothetical proteins are 96 shared with the genome of strain C7<sup>T</sup> (Supplementary Table 2). Stop codons also inactivated 97 eight other hypothetical proteins and genes encoding 3-oxoacyl-[acyl-carrier protein] 98 reductase, secreted peptidase and an oxetanocin A resistance protein (Supplementary Table 2). 99 100 3-oxoacyl-[acyl-carrier protein] reductase is involved in fatty acid biosynthesis by reducing βketoacyl-ACP substrates to beta-hydroxyacyl-ACP products (Lai and Cronan, 2004) but it is 101 unlikely to affect the strain fitness because there are multiple copies of the gene that encode 102 this enzyme in the genome. 103

The draft genome of strain  $C7^{T}$  is ~153 Kb larger than the chromosome of strain 103S but ~59 Kb smaller than the genome of the ATCC 33707 strain. Strains  $C7^{T}$  and 103S were isolated from foals (Goodfellow & Alderson, 1977; de la Pena-Moctezuma & Prescott, 1995) and strain ATCC 33707 from a human skin abscess (Prescott, 1981). The  $C7^{T}$  strain was isolated in 1922 but isolation dates are unknown for the remaining two strains. The smaller chromosome in strain 103S may represent degradation of the accessory genome that may not be essential for host infection. To identify horizontal gene acquisition by these strains, the 111 genomes were analysed by Alien Hunter (Vernikos & Parkhill, 2006) which identified 72 acquired regions in 103S (~662 Kb), 144 in C7<sup>T</sup> (~1.2 Mb) and 115 in ATCC 33707 (~972 Kb; 112 Supplementary Fig. 1). Most of the imports identified in the genome of strain 103S were also 113 identified in those of strains  $C7^{T}$  and ATCC 33707, including two large regions that were 114 previously described (Letek, et al., 2010). The draft status of the C7<sup>T</sup> (48 contigs) and ATCC 115 33707 (35 contigs) genomes might be largely responsible for identification of higher numbers 116 of imports as some gaps may have divided them into more fragments. However, the total length 117 of imports was also variable though most of the genes were conserved among these strains. The 118 119 predicted horizontally acquired regions contributed to 376 of the total 582 variable genes identified by genome-wide protein Blast searches (Supplementary Table 1). The host jumps of 120 Staphylococcus aureus have been characterized by gain or loss of accessory gene pool via 121 122 mobile genetic elements (Lowder, et al., 2009; Resch, et al., 2013) and horizontal gene transfer also appears to have little impact on functional variation among R. equi strains (Supplementary 123 Table 1). Some of the imported genes may be involved in virulence in particular hosts, but their 124 functions remain to be characterized as most of them encode hypothetical proteins. 125

The virulence plasmid, which is essential for virulence in *R. equi* (Takai, *et al.*, 2000; Letek, *et al.*, 2008; Vazquez-Boland, *et al.*, 2013), is missing from strains C7<sup>T</sup> and ATCC 33707,
both of which are avirulent (Takai, et al., 1985; Prescott, 1991). The virulence plasmid has
presumably been lost from these strains at some point during long term culture *in vitro*.

The phylogenetic relatedness of *R. equi* strains was investigated using PhyloPhlAn, a program designed to accurately calculate phylogenies from >400 proteins that can potentially resolve taxonomic groupings (Segata *et al.*, 2013). We also included the genome sequences of *Rhodococcus erythropolis* PR4 (Sekine, *et al.*, 2006; Letek, *et al.*, 2010), *Rhodococcus* strain RHA1 (McLeod, *et al.*, 2006), recently formally classified as *Rhodococcus jostii* (Jones *et al.*, 2013a) and *Nocardia brasiliensis* ATCC 700358 (Vera-Cabrera, *et al.*, 2012) and used 136 Corynebacterium diphtheriae NCTC 03529 (Sangal, et al., 2012b) as the outgroup (Accession numbers NC\_012490, CP000431, NC\_018681 and AJGI00000000, respectively). 137 All of the *R. equi* strains grouped closely together but distantly to the other rhodococci and *N*. 138 139 brasiliensis (Fig. 2), which supports our recent proposal of reclassifying R. equi into a separate genus as "Prescottella equi" (Jones, et al., 2013b; Jones, et al., 2013c). However, the 140 141 name Prescottella has still to be validated, although the taxonomic status of R. equi has been questioned (Kämpfer et al. 2013). Further analyses incorporating a wider range of genomes of 142 rhodococci and closely related bacteria are needed to remove any ambiguity over the proposal 143 144 for the prospective new genus.

In summary, we report the draft genome of the type strain of an important pathogen, R. equi 145 strain C7<sup>T</sup>. R. equi strains are phylogenetically closely related and the nucleotide sequence 146 147 and functional characteristics are highly conserved within this taxon. R. equi has recently emerged as an important opportunistic human pathogen but the comparison of equine and 148 human isolates revealed only small differences in the gene content that may be associated 149 with the host adaptation. The whole Genome Shotgun project of R. equi strain  $C7^{T}$  has been 150 deposited at DDBJ/EMBL/GenBank under the accession no. APJC00000000. The version 151 described in this paper is the first version, APJC01000000. 152

153

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256	Figure Legends
257	
258	Figure 1. Genomic blast maps of <i>R. equi</i> strain 103S against ATCC 33707 and C7 <sup>T</sup> . (A)
259	DNA blast map, (B) CDS blast map. The colour scale (inset) shows the level of sequence
260	identity with the respective sequences from strain $C7^{T}$ (middle circle) and strain ATCC 33707
261	(inner circle).
262	Figure 2. Phylogenetic tree from >400 universal proteins showing the genomic relationship
263	of R. equi to representatives of closely related taxa. Scale bar shows mean nucleotide
264	substitutions per site.







Supplementary Figure 1. Circular diagrams of *R. equi* chromosomes. Outer two rings are the coding sequences on forward and reverse strands. The genomic islands predicted by AlienHunter are shown in red colour followed by GC plot and GC skew, respectively.

Function	Protein Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	41	fig 685727.5.peg.65 (Q)	fig 525370.5.peg.4858 (97.5)		ATCC33707
hypothetical protein	216	fig 685727.5.peg.69 (Q)	fig 525370.5.peg.4854 (100)		ATCC33707
Hydantoin racemase (EC 5.1.99)	252	fig 685727.5.peg.75 (Q)	fig 525370.5.peg.4848 (99.6)		
hypothetical protein	64	fig 685727.5.peg.98 (Q)		fig 1219013.3.peg.93 (98.41)	
hypothetical protein	44	fig 685727.5.peg.101 (Q)	fig 525370.5.peg.4821 (100)		
FIG022958: hypothetical protein	210	fig 685727.5.peg.122 (Q)	fig 525370.5.peg.4977 (100)		
Permease of the drug/metabolite transporter (DMT) superfamily	51	fig 685727.5.peg.242 (Q)		fig 1219013.3.peg.240 (97.73)	
hypothetical protein	69	fig 685727.5.peg.264 (Q)		fig 1219013.3.peg.268 (100)	C7
hypothetical protein	110	fig 685727.5.peg.265 (Q)		fig 1219013.3.peg.269 (98.11)	C7
hypothetical protein	151	fig 685727.5.peg.266 (Q)		fig 1219013.3.peg.270 (98.67)	C7
hypothetical protein	39	fig 685727.5.peg.300 (Q)	fig 525370.5.peg.3529 (100)		
hypothetical protein	56	fig 685727.5.peg.306 (Q)			
hypothetical protein	178	fig 685727.5.peg.313 (Q)			
Nitric-oxide reductase (EC 1.7.99.7), quinol-dependent	778	fig 685727.5.peg.328 (Q)			
hypothetical protein	221	fig 685727.5.peg.329 (Q)			
hypothetical protein	56	fig 685727.5.peg.404 (Q)			
Arylsulfatase (EC 3.1.6.1)	48	fig 685727.5.peg.436 (Q)		fig 1219013.3.peg.462 (100)	
hypothetical protein	42	fig 685727.5.peg.461 (Q)		fig 1219013.3.peg.487 (100)	C7
hypothetical protein	85	fig 685727.5.peg.477 (Q)	fig 525370.5.peg.713 (100)		
hypothetical protein	98	fig 685727.5.peg.484 (Q)	,	fig 1219013.3.peg.514 (98.97)	
hypothetical protein	42	fig 685727.5.peg.509 (Q)	fig 525370.5.peg.2715 (100)		
hypothetical protein	66	fig 685727.5.peg.558 (Q)	on		
hypothetical protein	50	fig 685727.5.peg.563 (Q)			
hypothetical protein	71	fig 685727.5.peg.650 (Q)			
hypothetical protein	149	fig 685727.5.peg.759 (Q)		fig 1219013.3.peg.838 (98.65)	
putative ABC transporter ATP binding protein	612	fig 685727.5.peg.791 (Q)		fig 1219013.3.peg.4071 (28.26)	
hypothetical protein	62	fig 685727.5.peg.892 (Q)		fig 1219013.3.peg.987 (90.16)	
hypothetical protein	71	fig 685727.5.peg.921 (Q)	fig 525370.5.peg.2059 (100)		
hypothetical protein	39	fig 685727.5.peg.947 (Q)			
hypothetical protein	40	fig 685727.5.peg.960 (Q)			103S
hypothetical protein	61	fig 685727.5.peg.961 (Q)			1035
Putative membrane protein	182	fig 685727.5.peg.981 (Q)		fig 1219013.3.peg.1085 (98.34)	
hypothetical protein	51	fig 685727.5.peg.1008 (Q)		fig 1219013.3.peg.1113 (100)	
hypothetical protein	43	fig 685727.5.peg.1059 (Q)		··01	
hypothetical protein	202	fig 685727.5.peg.1103 (Q)	fig 525370.5.peg.3802 (97.5)		
hypothetical protein	45	fig 685727.5.peg.1129 (Q)	01	fig 1219013.3.peg.1237 (100)	
hypothetical protein	53	fig 685727.5.peg.1285 (Q)		01	
hypothetical protein	77	fig 685727.5.peg.1334 (Q)	fig 525370.5.peg.3954 (98.67)		
hypothetical protein	43	fig 685727.5.peg.1461 (Q)	on		
hypothetical protein	60	fig 685727.5.peg.1552 (Q)		fig 1219013.3.peg.1665 (98.31)	C7
hypothetical protein	53	fig 685727.5.peg.1639 (Q)		fig 1219013.3.peg.1757 (100)	C7
hypothetical protein	200	fig 685727.5.peg.1640 (Q)		fig 1219013.3.peg.1757 (98.99)	
hypothetical protein	139	fig 685727.5.peg.1641 (Q)	fig 525370.5.peg.1120 (42.97)	on	
hypothetical protein	152	fig 685727.5.peg.1645 (Q)	fig 525370.5.peg.1116 (84.21)		ATCC33707-1035
hypothetical protein	321	fig 685727.5.peg.1646 (Q)	fig 525370.5.peg.1084 (31.85)		ATCC33707-1035
hypothetical protein	487	fig 685727.5.peg.1647 (Q)			1035
hypothetical protein	255	fig 685727.5.peg.1649 (Q)	fig 525370.5.peg.1109 (98.82)		ATCC33707-103S
DNA helicase related protein	1828	fig 685727.5.peg.1655 (Q)	fig 525370.5.peg.1095 (23.18)		ATCC33707-103S
hypothetical protein	92	fig   685727.5.peg.1656 (Q)	on,,,,,		1035
hypothetical protein	924	fig 685727.5.peg.1658 (O)			1035
Helicase, C-terminal:Type III restriction enzyme, res	2130	fig 685727.5.peg.1659 (Q)			1035
subunit:DEAD/DEAH box helicase, N-terminal					
Type II restriction enzyme, methylase subunits	1530	fig 685727.5.peg.1660 (Q)			103S
putative ATP-dependent helicase	960	fig 685727.5.peg.1661 (Q)	fig 525370.5.peg.2722 (22.42)		
hypothetical protein	1135	fig 685727.5.peg.1662 (Q)			
hypothetical protein	114	fig 685727.5.peg.1664 (Q)			103S
nypotnetical protein	444	tig 685727.5.peg.1665 (Q)			1035
hypothetical protein	63	fig 685727.5.peg.1666 (Q)			103S
hypothetical protein	118	tig 685727.5.peg.1667 (Q)	8 January 1		1035
hypothetical protein	94	fig 685727.5.peg.1669 (Q)	fig 525370.5.peg.1077 (86.02)		ATCC33707-103S
hypothetical protein	173	fig 685727.5.peg.1670 (Q)	fig 525370.5.peg.1076 (88.95)		
hypothetical protein	55	fig 685727.5.peg.1681 (Q)	fig 525370.5.peg.1064 (100)		
cyclase/dehydrase	173	fig 685727.5.peg.1707 (Q)	fig 525370.5.peg.1038 (100)		
hypothetical protein	72	fig 685727.5.peg.1708 (Q)	fig 525370.5.peg.1037 (98.59)		
putative isomerase	122	fig 685727.5.peg.1709 (Q)	fig 525370.5.peg.1036 (100)		
NADH-ubiquinone oxidoreductase chain H (EC 1.6.5.3)	435	fig 685727.5.peg.1900 (Q)	fig 525370.5.peg.4090 (99.08)		
hypothetical protein	38	fig 685727.5.peg.1908 (Q)			

Supplementary Table 1. Differences in the gene content between 103S, ATCC33707 and 103S based on the bidirectional protein blast searches

Function	Protein Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	65	fig 685727.5.peg.1998 (Q)			
COG1359: Uncharacterized conserved protein	79	fig 685727.5.peg.2112 (Q)			103S
hypothetical protein	61	fig 685727.5.peg.2114 (Q)			103S
hypothetical protein	123	fig 685727.5.peg.2116 (Q)			103S
Non-heme chloroperoxidase (EC 1.11.1.10)	52	fig 685727.5.peg.2118 (Q)			
alpha/beta hydrolase fold( EC:1.11.1.10)	64	fig 685727.5.peg.2119 (Q)			
hypothetical protein	38	fig 685727.5.peg.2121 (Q)			
hypothetical protein	82	fig 685727.5.peg.2133 (Q)			
hypothetical protein	58	fig 685727.5.peg.2218 (Q)			
hypothetical protein	38	fig 685727.5.peg.2265 (Q)		fig 1219013.3.peg.2377 (100)	
hypothetical protein	116	fig 685727.5.peg.2347 (Q)	fig 525370.5.peg.4220 (88.89)		
hypothetical protein	57	fig 685727.5.peg.2390 (Q)	fig 525370.5.peg.4177 (100)		
hypothetical protein	444	fig 685727.5.peg.2393 (Q)			
hypothetical protein	43	fig 685727.5.peg.2401 (Q)			
hypothetical protein	82	fig 685727.5.peg.2435 (Q)	fig 525370.5.peg.1805 (98.77)		
hypothetical protein	84	fig 685727.5.peg.2475 (Q)	fig 525370.5.peg.1704 (98.78)		ATCC33707-103S
hypothetical protein	67	fig 685727.5.peg.2533 (Q)		fig 1219013.3.peg.2646 (98.48)	
hypothetical protein	42	fig 685727.5.peg.2566 (Q)			103S
hypothetical protein	56	fig 685727.5.peg.2569 (Q)	fig 525370.5.peg.1612 (100)		ATCC33707-103S
hypothetical protein	86	fig 685727.5.peg.2580 (Q)		fig 1219013.3.peg.2691 (100)	C7-103S
FIG01136303: hypothetical protein	115	fig 685727.5.peg.2585 (Q)		fig 1219013.3.peg.2699 (95.12)	103S
hypothetical protein	86	fig 685727.5.peg.2586 (Q)	fig 525370.5.peg.1595 (95.56)		ATCC33707-103S
hypothetical protein	84	fig 685727.5.peg.2615 (Q)	fig 525370.5.peg.1965 (89.16)	<u> </u>	ATCC33707-103S
hypothetical protein	260	fig 685727.5.peg.2617 (Q)		fig 1219013.3.peg.2734 (96.91)	C7-103S
nypotnetical protein	100	fig 685727.5.peg.2618 (Q)		fig 1219013.3.peg.2735 (94.95)	C7-103S
putative acetyltransferase	143	fig 685727.5.peg.2658 (Q)			1035
hypothetical protein	195	fig 685727.5.peg.2660 (Q)		fig 1210012 2 mag 2802 (100)	1035
hypothetical protein	200	fig 685727.5.peg.2684 (Q)		fig 1219013.3.peg.2803 (100)	C7-1035
hypothetical protein	355	fig 685727.5.peg.2005 (Q)		fig 1219013.3.peg.2804 (99.7)	C7-1035
hypothetical protein	425	fig 665727.5.peg.2667 (Q)		fig 1219013.3.peg.2800 (38.87)	1025
hypothetical protein	425	fig 685727.5.peg.2008 (Q)		fig 1219013.3.peg.4207 (29.87)	1055
hypothetical protein	38	fig 685727.5.pcg.2701(Q)		ng 1213013.3.pcg.2010 (37.00)	0,
hypothetical protein	48	fig 685727.5.peg.3094 (Q)			
hypothetical protein	43	fig   685727.5.peg.3095 (Q)		fig 1219013.3.peg.3228 (92.86)	C7
hypothetical protein	64	fig 685727.5.peg.3272 (Q)		fig 1219013.3.peg.3411 (96.83)	
hypothetical protein	48	fig 685727.5.peg.3310 (Q)	fig 525370.5.peg.3144 (100)		ATCC33707-103S
hypothetical protein	38	fig 685727.5.peg.3318 (Q)			
hypothetical protein	63	fig 685727.5.peg.3447 (Q)			103S
hypothetical protein	65	fig 685727.5.peg.3448 (Q)			103S
hypothetical protein	124	fig 685727.5.peg.3449 (Q)			103S
hypothetical protein	256	fig 685727.5.peg.3450 (Q)			103S
hypothetical protein	281	fig 685727.5.peg.3451 (Q)			103S
hypothetical protein	613	fig 685727.5.peg.3452 (Q)		fig 1219013.3.peg.1777 (32.39)	C7-103S
hypothetical protein	38	fig 685727.5.peg.3480 (Q)		fig 1219013.3.peg.3618 (100)	
hypothetical protein	97	fig 685727.5.peg.3522 (Q)		fig 1219013.3.peg.3660 (97.92)	
YgjD/Kae1/Qri/ family, required for threonylcarbamoyladenosine (t(6)A) formation in tRNA	347	fig 685727.5.peg.3523 (Q)		tig 1219013.3.peg.3661 (99.71)	
hypothetical protein	167	fig 685727.5.peg.3799 (Q)	fig 525370.5.peg.1498 (96.99)		
hypothetical protein	161	fig 685727.5.peg.3853 (Q)			103S
hypothetical protein	45	fig 685727.5.peg.4055 (Q)			103S
hypothetical protein	48	fig 685727.5.peg.4057 (Q)			103S
GumJ protein	504	fig 685727.5.peg.4058 (Q)			103S
hypothetical protein	363	fig 685727.5.peg.4059 (Q)			103S
hypothetical protein	44	fig 685727.5.peg.4063 (Q)	fig 525370.5.peg.673 (95.35)		ATCC33707-103S
hypothetical protein	106	fig 685727.5.peg.4065 (Q)			103S
hypothetical protein	47	fig 685727.5.peg.4066 (Q)	fig 525370.5.peg.671 (89.13)		ATCC33707-103S
hypothetical protein	90	fig 685727.5.peg.4069 (Q)	fig 525370.5.peg.668 (100)		ATCC33707-103S
hypothetical protein	54	fig 685727.5.peg.4140 (Q)	fig 525370.5.peg.4557 (91.07)		
hypothetical protein	53	fig 685727.5.peg.4187 (Q)		fig 1219013.3.peg.4353 (98.08)	
hypothetical protein	38	fig 685727.5.peg.4392 (Q)	fig 525370.5.peg.513 (100)		
hypothetical protein	44	fig 685727.5.peg.4462 (Q)		fig 1219013.3.peg.4643 (97.67)	C7-103S
hypothetical protein	39	fig 685727.5.peg.4465 (Q)		fig 1219013.3.peg.4646 (100)	C7-103S
hypothetical protein	42	fig 685727.5.peg.4466 (Q)		fig 1219013.3.peg.4647 (97.56)	C7-103S
hypothetical protein	39	fig 685727.5.peg.4517 (Q)		fig 1219013.3.peg.4700 (100)	
hypothetical protein	88	fig 685727.5.peg.4520 (Q)		fig 1219013.3.peg.4703 (93.1)	
hypothetical protein	45	fig 685727.5.peg.4524 (Q)	fig 525370.5.peg.382 (97.73)		ATCC33707

Function	Protein Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	48	fig 685727.5.peg.4559 (Q)	fig 525370.5.peg.347 (95.74)		
hypothetical protein	56	fig 685727.5.peg.4616 (Q)			103S
hypothetical protein	48	fig 685727.5.peg.4617 (Q)			103S
hypothetical protein	41	fig 685727.5.peg.4618 (Q)			103S
similar to Ketosteroid isomerase-related protein	136	fig 685727.5.peg.4620 (Q)			103S
hypothetical protein	45	fig 685727.5.peg.4621 (Q)			103S
Transposase	80	fig 685727.5.peg.4622 (Q)			103S
transposase, IS4 family protein	99	fig 685727.5.peg.4623 (Q)			103S
Mobile element protein	71	fig 685727.5.peg.4624 (Q)			103S
hypothetical protein	85	fig 685727.5.peg.4627 (Q)			103S
hypothetical protein	466	fig 685727.5.peg.4628 (Q)			103S
hypothetical protein	245	fig 685727.5.peg.4629 (Q)			103S
Putative DNA-binding protein	277	fig 685727.5.peg.4631 (Q)	fig 525370.5.peg.1035 (31.78)		103S
hypothetical protein	41	fig 685727.5.peg.4632 (Q)	,		103S
hypothetical protein	849	fig 685727.5.peg.4634 (Q)			103S
hypothetical protein	255	fig 685727.5.peg.4635 (Q)			103S
Integrase	279	fig 685727.5.peg.4636 (O)			1035
hypothetical protein	932	fig 685727.5.peg.4637 (Q)			103S
hypothetical protein	138	fig   685727.5.peg.4639 (Q)	fig 525370.5.peg.287 (96.35)		ATCC33707-1035
hypothetical protein	42	fig 685727.5.peg.4734 (Q)	fig 525370.5.peg.4931 (100)		ATCC33707-1035
rRNA small subunit 7-methylguanosine (m7G) methyltransferase GidB	224	fig 685727.5.peg.4735 (Q)	fig 525370.5.peg.4930 (96.41)		ATCC33707-1035
hypothetical protein	40		fig 525370.5.peg.120 (Q)		
hypothetical protein	41		fig 525370.5.peg.134 (Q)		
hypothetical protein	40		fig 525370.5.peg.344 (Q)		
hypothetical protein	38		fig 525370.5.peg.440 (Q)		
hypothetical protein	611		fig 525370.5.peg.464 (Q)		ATCC33707
ATP-dependent DNA helicase UvrD/PcrA	567		fig 525370.5.peg.465 (Q)		ATCC33707
hypothetical protein	39		fig 525370.5.peg.479 (Q)		ATCC33707
hypothetical protein	306		fig 525370.5.peg.621 (Q)		
hypothetical protein	583		fig 525370.5.peg.622 (Q)		
hypothetical protein	45		fig 525370.5.peg.645 (Q)		ATCC33707
Glycosyltransferase	130	fig 685727.5.peg.4060 (32.74)	fig 525370.5.peg.675 (Q)		ATCC33707
FIG00511468: hypothetical protein	535		fig 525370.5.peg.676 (Q)		ATCC33707
hypothetical protein	112		fig 525370.5.peg.681 (Q)		ATCC33707
3-oxoacyl-[acyl-carrier protein] reductase (EC 1.1.1.100)	259		fig 525370.5.peg.751 (Q)		ATCC33707
hypothetical protein	211		fig 525370.5.peg.752 (Q)	fig 1219013.3.peg.4138 (24.56)	ATCC33707
Phenylacetic acid degradation protein PaaD, thioesterase	140		fig 525370.5.peg.754 (Q)		ATCC33707
hypothetical protein	105		fig 525370.5.peg.755 (Q)		ATCC33707
Aldehyde dehydrogenase (EC 1.2.1.3), PaaZ	682		fig 525370.5.peg.756 (Q)		ATCC33707
Transcriptional regulator, TetR family	179		fig 525370.5.peg.757 (Q)		ATCC33707
Phenylacetate-CoA oxygenase, PaaG subunit	317		fig 525370.5.peg.761 (Q)		
Phenylacetate-CoA oxygenase, PaaH subunit	108		fig 525370.5.peg.762 (Q)		
Phenylacetate-CoA oxygenase, Paal subunit	306		fig 525370.5.peg.763 (Q)		
Phenylacetate-coenzyme A ligase (EC 6.2.1.30) PaaF	433		fig 525370.5.peg.766 (Q)		
hypothetical protein	798	fig 685727.5.peg.1668 (82.28)	fig 525370.5.peg.1078 (Q)		ATCC33707
hypothetical protein	253	51	fig 525370.5.peg.1079 (O)		ATCC33707
hypothetical protein	97		fig 525370.5.peg.1080 (Q)		ATCC33707
hypothetical protein	160		fig 525370.5.peg.1081 (O)		ATCC33707
hypothetical protein	104		fig 525370.5.peg.1082 (O)		ATCC33707
hypothetical protein	142		fig 525370.5.neg.1083(0)		ATCC33707
hypothetical protein	318		fig 525370.5.pcg.1085 (Q)		ATCC33707
hypothetical protein	49		fig 525370.5.peg.1086 (Q)		ATCC33707
hypothetical protein	51		fig 525370.5.peg.1087 (Q)		ATCC33707
ATP/GTP hinding protein	627		fig 525370.5.pcg.1087 (Q)		ATCC33707
hypothetical protein	283		fig 525370.5.pcg.1089 (Q)		ATCC33707
hypothetical protein	205		fig 525370 5 nor 1000 (Q)		ATCC22707
hypothetical protein	164		fig 525370.5 peg 1002 (Q)		ATCC22707
hypothetical protein	104		fig 525370.5. peg. 1092 (Q)		ATCC22707
hypothetical protein	749		fig1525570.5.peg.1096 (Q)		ATCC22707
nypoinetical protein	290		iig 525370.5.peg.1097 (Q)		ATCC33707
nypoinetical protein	388		iig 525370.5.peg.1098 (Q)		ATCC33707
nypotnetical protein	340		тіg 525370.5.peg.1099 (Q)		
hypothetical protein	843		tig 525370.5.peg.1100 (Q)		
FIG00995584: hypothetical protein	656		tig 525370.5.peg.1101 (Q)		
hypothetical protein	917		tig 525370.5.peg.1102 (Q)		
FIG00995267: hypothetical protein	1191		fig 525370.5.peg.1103 (Q)		ATCC33707
FIG00998105: hypothetical protein	922		fig 525370.5.peg.1104 (Q)		ATCC33707

Function	Protein Length 103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	97	fig 525370.5.peg.1105 (Q)		ATCC33707
hypothetical protein	78	fig 525370.5.peg.1111 (Q)		ATCC33707
hypothetical protein	120	fig 525370.5.peg.1112 (Q)		ATCC33707
hypothetical protein	201	fig 525370.5.peg.1113 (Q)		ATCC33707
hypothetical protein	98	fig 525370.5.peg.1114 (Q)		ATCC33707
hypothetical protein	45	fig 525370.5.peg.1115 (Q)		ATCC33707
Partial REP13E12 repeat protein	195	fig 525370.5.peg.1224 (Q)		ATCC33707
hypothetical protein	73	fig 525370.5.peg.1425 (Q)		
hypothetical protein	39	fig 525370.5.peg.1441 (Q)		ATCC33707
hypothetical protein	44	fig 525370.5.peg.1444 (Q)		ATCC33707
hypothetical protein	99	fig 525370.5.peg.1599 (Q)		ATCC33707
hypothetical protein	128	fig 525370.5.peg.1600 (Q)		ATCC33707
hypothetical protein	242	fig 525370.5.peg.1721 (Q)		ATCC33707
hypothetical protein	55	fig 525370.5.peg.1722 (Q)		ATCC33707
hypothetical protein	85	fig 525370.5.peg.1723 (Q)		ATCC33707
hypothetical protein	91	fig 525370.5.peg.1724 (Q)		ATCC33707
Phage Gp37Gp68 protein	39	fig 525370.5.peg.1725(0)		ATCC33707
hypothetical protein	240	fig 525370.5.peg.1726 (Q)		ATCC33707
hypothetical protein	515	fig 525370.5.peg 1727 (Q)		ATCC33707
hypothetical protein	206	fig 525370.5 peg 1728 (Q)		ATCC33707
nypometical protein	357	fig 525370.5.pcg.1720 (Q)		ATCC33707
hypothetical protein	124	fig 525370.5.peg.1729 (Q)		Alcession
hypothetical protein	154	fig 525370.5.peg.1730 (Q)		
hypothetical protein	199	fig 525370.5.peg.1732 (Q)		
hypothetical protein	375	fig 525370.5.peg.1733 (Q)	· · · · · · · · · · · · · · · · · · ·	
hypothetical protein	480	fig 525370.5.peg.1734 (Q)	fig 1219013.3.peg.4933 (21.38)	
hypothetical protein	290	fig 525370.5.peg.1735 (Q)		
tail length tape measure protein	1610	fig 525370.5.peg.1736 (Q)	fig 1219013.3.peg.4935 (25.45)	ATCC33707
hypothetical protein	121	fig 525370.5.peg.1737 (Q)		ATCC33707
hypothetical protein	148	fig 525370.5.peg.1738 (Q)		ATCC33707
phage major tail protein	186	fig 525370.5.peg.1739 (Q)		ATCC33707
hypothetical protein	126	fig 525370.5.peg.1740 (Q)		ATCC33707
hypothetical protein	68	fig 525370.5.peg.1741 (Q)		
hypothetical protein	78	fig 525370.5.peg.1742 (Q)		
hypothetical protein	120	fig 525370.5.peg.1743 (Q)		
hypothetical protein	139	fig 525370.5.peg.1744 (Q)		
hypothetical protein	152	fig 525370.5.peg.1745 (Q)		
hypothetical protein	308	fig 525370.5.peg.1746 (Q)		ATCC33707
hypothetical protein	137	fig 525370.5.peg.1747 (Q)		ATCC33707
hypothetical protein	422	fig 525370.5.peg.1749 (Q)		ATCC33707
phage terminase, large subunit, putative	490	fig 525370.5.peg.1750 (Q)		ATCC33707
hypothetical protein	116	fig 525370.5.peg.1751 (Q)		ATCC33707
hypothetical protein	46	fig 525370.5.peg.1752 (Q)		ATCC33707
hypothetical protein	107	fig 525370.5.peg.1753 (Q)		ATCC33707
hypothetical protein	86	fig 525370.5.peg.1754 (Q)		ATCC33707
hypothetical protein	66	fig 525370.5.peg.1755 (Q)		ATCC33707
hypothetical protein	126	fig 525370.5.peg.1756 (Q)		ATCC33707
hypothetical protein	222	fig 525370.5.peg.1757 (Q)	fig 1219013.3.peg.4952 (31.48)	ATCC33707
hypothetical protein	65	fig 525370.5.peg.1758 (Q)		ATCC33707
hypothetical protein	149	fig 525370.5.peg.1759 (Q)		ATCC33707
hypothetical protein	143	fig 525370.5.peg.1760 (O)		ATCC33707
hypothetical protein	92	fig   525370, 5 neg 1761 (O)		ATCC33707
hypothetical protein	59	fig   525370, 5 neg 1762 (O)		ATCC33707
hypothetical protein		fig 525370,5 neg 1763 (0)		ATCC33707
hypothetical protein		fig   525370 5 neg 1764 (0)		
hypothetical protein		fig   525370 5 neg 1766 (0)		
	20	fig 525270 5 peg 1767 (Q)		
	174	fig 525370.3. peg. 1769 (Q)		
Call division protein	1/ <del>4</del> 270	fig = 525570.5. peg. 1770 (Q)	fig 1210012 2 mag 2005 (54.04)	
Cell division protein Ftsk	370	ng 525370.5.peg.1770 (Q)	ng 1219013.3.peg.2065 (54.84)	470020707
nypotnetical protein	332	тіg 525370.5.peg.1771 (Q)		ATCC33707
hypothetical protein	53	tig 525370.5.peg.1772 (Q)		ATCC33707
hypothetical protein	83	tig 525370.5.peg.1773 (Q)		ATCC33707
hypothetical protein	92	tig 525370.5.peg.1774 (Q)		ATCC33707
hypothetical protein	102	fig 525370.5.peg.1775 (Q)		ATCC33707
hypothetical protein	124	fig 525370.5.peg.1776 (Q)		ATCC33707
hypothetical protein	177	fig 525370.5.peg.1777 (Q)		ATCC33707
hypothetical protein	68	fig 525370.5.peg.1778 (Q)		ATCC33707

Function	Protein Length 103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	79	fig 525370.5.peg.1779 (Q)		ATCC33707
hypothetical protein	328	fig 525370.5.peg.1887 (Q)	fig 1219013.3.peg.2816 (96.15)	
hypothetical protein	46	fig 525370.5.peg.1888 (Q)		
hypothetical protein	50	fig 525370.5.peg.1898 (Q)		ATCC33707
hypothetical protein	242	fig 525370.5.peg.2073 (Q)		
hypothetical protein	59	fig 525370.5.peg.2089 (Q)		
hypothetical protein	49	fig 525370.5.peg.2105 (Q)		ATCC33707
hypothetical protein	76	fig 525370.5.peg.2106 (Q)		ATCC33707
hypothetical protein	71	fig 525370.5.peg.2107 (Q)		ATCC33707
hypothetical protein	258	fig 525370.5.peg.2108 (Q)		ATCC33707
hypothetical protein	43	fig 525370.5.peg.2109 (Q)		ATCC33707
hypothetical protein	42	fig 525370.5.peg.2110 (Q)		ATCC33707
hypothetical protein	66	fig 525370.5.peg.2111 (Q)		ATCC33707
hypothetical protein	45	fig 525370.5.peg.2112 (Q)		ATCC33707
hypothetical protein	60	fig 525370.5.peg.2113 (Q)		ATCC33707
hypothetical protein	65	fig 525370.5.peg.2114 (Q)		ATCC33707
hypothetical protein	249	fig 525370.5.peg.2117 (Q)		ATCC33707
hypothetical protein	43	fig 525370.5.peg.2118 (Q)		ATCC33707
hypothetical protein	59	fig 525370.5.peg.2119 (Q)		ATCC33707
hypothetical protein	72	fig 525370.5.peg.2120 (Q)		ATCC33707
hypothetical protein	143	fig 525370.5.peg.2121 (Q)		ATCC33707
hypothetical protein	120	fig 525370.5.peg.2122 (Q)		ATCC33707
hypothetical protein	152	fig 525370.5.peg.2123 (Q)		ATCC33707
hypothetical protein	51	fig 525370.5.peg.2124 (Q)		ATCC33707
hypothetical protein	40	fig 525370.5.peg.2125 (Q)		ATCC33707
hypothetical protein	130	fig 525370.5.peg.2127 (Q)		ATCC33707
hypothetical protein	58	fig 525370.5.peg.2128 (Q)		ATCC33707
hypothetical protein	56	fig 525370.5.peg.2129 (Q)		ATCC33707
hypothetical protein	113	fig 525370.5.peg.2130 (Q)		ATCC33707
Phage terminase	515	fig 525370.5.peg.2131 (Q)		ATCC33707
hypothetical protein	424	fig 525370.5.peg.2132 (Q)		ATCC33707
hypothetical protein	476	fig 525370.5.peg.2133 (Q)		ATCC33707
dehydrogenase / Cytidine deaminase (EC 3.5.4.5)	147	fig 525370.5.peg.2134 (Q)		
hypothetical protein	314	fig 525370.5.peg.2135 (Q)		
hypothetical protein	129	fig 525370.5.peg.2136 (Q)		
hypothetical protein	124	fig 525370.5.peg.2137 (Q)		
hypothetical protein	110	fig 525370.5.peg.2138 (Q)		ATCC33707
Sigma 54-dependent transcriptional activator	94	fig 525370.5.peg.2139 (Q)		ATCC33707
hypothetical protein	133	fig 525370.5.peg.2140 (Q)		ATCC33707
hypothetical protein	268	fig 525370.5.peg.2141 (Q)		ATCC33707
hypothetical protein	88	fig 525370.5.peg.2142 (Q)		ATCC33707
hypothetical protein	38	fig 525370.5.peg.2143 (Q)		ATCC33707
hypothetical protein	98	fig 525370.5.peg.2144 (Q)		ATCC33707
hypothetical protein	149	fig 525370.5.peg.2145 (Q)		ATCC33707
hypothetical protein	1.51	fig 525570.5.peg.2147 (Q)	fig 1210012 2 pog 4025 (22 5)	ATCC22707
hypothetical protein	11/0	fig 525570.5.peg.2147 (Q)	11811111012.3.heg.4932 (33.2)	ATCC32707
hypothetical protein	123	fig   525370 5 nor 2152 (0)		ATCC32707
hypothetical protein	92	fig   525370 5 neg 2152 (Q)		ATCC33707
hypothetical protein	62	fig 525370 5 neg 2154 (0)		ATCC33707
hypothetical protein	118	fig   525370 5 neg 2155 (0)		ATCC33707
hypothetical protein	288	fig 525370 5 neg 2156 (Q)		ATCC33707
hypothetical protein	112	fig 525370.5.peg.2157 (0)		ATCC33707
hypothetical protein	111	fig 525370.5.peg.2158 (O)		ATCC33707
hypothetical protein	115	fig 525370.5.peg.2159 (Q)		ATCC33707
hypothetical protein	329	fig 525370.5.peg.2160 (Q)		ATCC33707
hypothetical protein	73	fig 525370.5.peg.2161 (Q)		ATCC33707
hypothetical protein	72	fig 525370.5.peg.2162 (Q)		ATCC33707
hypothetical protein	53	fig 525370.5.peg.2163 (Q)		ATCC33707
hypothetical protein	40	fig 525370.5.peg.2164 (Q)		ATCC33707
Integrase-like protein	275	fig 525370.5.peg.2165 (Q)		ATCC33707
hypothetical protein	38	fig 525370.5.peg.2166 (Q)		ATCC33707
hypothetical protein	249	fig 525370.5.peg.2167 (Q)		ATCC33707
hypothetical protein	54	fig 525370.5.peg.2168 (Q)		ATCC33707
hypothetical protein	119	fig 525370.5.peg.2169 (Q)		ATCC33707
hypothetical protein	67	fig 525370.5.peg.2298 (Q)		

Function	Protein Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	330		fig 525370.5.peg.2559 (Q)		ATCC33707
hypothetical protein	52		fig 525370.5.peg.2598 (Q)		ATCC33707
hypothetical protein	58		fig 525370.5.peg.2640 (Q)		ATCC33707
Arsenical pump-driving ATPase (EC 3.6.3.16)	380		fig 525370.5.peg.2730 (Q)	fig 1219013.3.peg.532 (100)	
hypothetical protein	82		fig 525370.5.peg.2971 (Q)		
hypothetical protein	79		fig 525370.5.peg.3033 (Q)		
hypothetical protein	61		fig 525370.5.peg.3517 (Q)		
Acyl dehydratase	128		fig 525370.5.peg.3541 (Q)		
FIG00827952: hypothetical protein	420		fig 525370.5.peg.3544 (Q)		
hypothetical protein	52		fig 525370.5.peg.3659 (Q)		
hypothetical protein	38		fig 525370.5.peg.3771 (Q)		ATCC33707
hypothetical protein	38		fig 525370.5.peg.3875 (Q)		
hypothetical protein	164		fig 525370.5.peg.3918 (Q)		
hypothetical protein	80		fig 525370.5.peg.4018 (Q)		ATCC33707
hypothetical protein	619		fig 525370.5.peg.4388 (Q)		
hypothetical protein	44		fig 525370.5.peg.4466 (Q)		
Mobile element protein	285		fig 525370.5.peg.4685 (Q)		
transposase for IS3517	109		fig 525370.5.peg.4686 (Q)		
hypothetical protein	49		fig 525370.5.peg.4845 (Q)		
2,3-dihydroxybiphenyl 1,2-dioxygenase	293	fig 685727.5.peg.72 (100)	fig 525370.5.peg.4851 (Q)		
hypothetical protein rRNA small subunit 7-methylguanosine (m7G) methyltransferase GidB	62 226	fig 685727.5.peg.4735 (95.52)	fig 525370.5.peg.4873 (Q) fig 525370.5.peg.4928 (Q)		ATCC33707-103S
hypothetical protein	54		fig 525370.5.peg.4839 (97.3)	fig 1219013.3.peg.76 (Q)	
hypothetical protein	109		01	fig   1219013.3.peg.177 (Q)	C7
Glycerol-3-phosphate regulon repressor. DeoR family	277		fig 525370.5.peg.2517 (99.64)	fig   1219013.3.peg.258 (Q)	ATCC33707-C7
hypothetical protein	71		fig 525370.5.peg.2514 (98.57)	fig 1219013.3.peg.261 (Q)	ATCC33707-C7
Glycerol-3-phosphate ABC transporter, periplasmic glycerol-3-	204				
phosphate-binding protein (IC 3.A.1.1.3)	394		fig 525370.5.peg.2513 (100)	fig 1219013.3.peg.262 (Q)	ATCC33707-C7
integrase	287		Tig 525370.5.peg.1780 (28.82)	fig 1219013.3.peg.304 (Q)	C7
nypothetical protein	290			fig 1219013.3.peg.305 (Q)	C7
hypothetical protein	540			fig 1219013.3.peg.300 (Q)	67
hypothetical protein	52			fig 1219013.3.peg.307 (Q)	67
hypothetical protein	54			fig 1219013.3.peg.308 (Q)	C7
hypothetical protein	114			fig 1219013.3.peg.309 (Q)	C7
hypothetical protein	205			fig 1219013.3.peg.310 (Q)	C7
hypothetical protein	203			fig 1219013.3.peg.311 (Q)	C7
hypothetical protein	257			fig 1219013.3.peg.312 (Q)	C7
Transcriptional regulator	351			fig 1219013.3.peg.314 (Q)	C7
hypothetical protein	136			fig 1219013.3.peg.315 (Q)	C7
hypothetical protein	222			fig 1219013.3.peg.310 (Q)	C7
hypothetical protein	305			fig 1219013.3.peg.317 (Q)	C7
hypothetical protein	133			fig 1219013.3.peg.319 (Q)	C7
hypothetical protein	80			fig 1219013.3.pcg.319 (Q)	C7
hypothetical protein	66			fig 1219013 3 neg 321 (Q)	C7
hypothetical protein	66			fig 1219013 3 neg 322 (Q)	C7
hypothetical protein	156			fig 1219013.3.peg.334 (O)	
hypothetical protein	101		fig 525370.5.peg.3516 (97)	fig   1219013.3.peg.335 (O)	
hypothetical protein	194		fig 525370.5.peg.794 (96.02)	fig 1219013.3.peg.435 (Q)	ATCC33707-C7
Putative polysaccharide deacetylase	294		fig 525370.5.peg.790 (99.66)	fig   1219013.3.peg.439 (Q)	ATCC33707-C7
L-asparaginase (EC 3.5.1.1)	337		fig 525370.5.peg.789 (100)	fig 1219013.3.peg.440 (Q)	
hypothetical protein	163		01	fig 1219013.3.peg.494 (Q)	
hypothetical protein	65			fig 1219013.3.peg.506 (Q)	
hypothetical protein	91			fig 1219013.3.peg.530 (Q)	
WhiB-type transcription regulator	53			fig 1219013.3.peg.531 (Q)	
Arsenical pump-driving ATPase (EC 3.6.3.16)	347		fig 525370.5.peg.2729 (100)	fig 1219013.3.peg.533 (Q)	
FIG011121: hypothetical protein	44		fig 525370.5.peg.2728 (100)	fig 1219013.3.peg.534 (Q)	
Enoyl-CoA hydratase (EC 4.2.1.17)	269		fig 525370.5.peg.2695 (100)	fig 1219013.3.peg.566 (Q)	
hypothetical protein	369		fig 525370.5.peg.2693 (100)	fig 1219013.3.peg.568 (Q)	
hypothetical protein	350		fig 525370.5.peg.2687 (100)	fig 1219013.3.peg.574 (Q)	ATCC33707-C7
hypothetical protein	131		fig 525370.5.peg.2681 (97.69)	fig 1219013.3.peg.580 (Q)	C7
Transcriptional regulator, IcIR family	273		fig 525370.5.peg.2674 (99.26)	fig 1219013.3.peg.587 (Q)	
hypothetical protein	170		fig 525370.5.peg.2667 (98.82)	fig 1219013.3.peg.594 (Q)	ATCC33707-C7
Beta-mannanase-like protein	329		fig 525370.5.peg.2627 (98.78)	fig 1219013.3.peg.633 (Q)	
Cellulose synthase catalytic subunit	658		fig 525370.5.peg.2625 (99.24)	fig 1219013.3.peg.635 (Q)	
hypothetical protein	146		fig 525370.5.peg.2623 (99.31)	fig 1219013.3.peg.637 (Q)	

Function	Protein Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
FIG00994841: hypothetical protein	90		fig 525370.5.peg.2460 (100)	fig 1219013.3.peg.675 (Q)	
hypothetical protein	51		fig 525370.5.peg.2410 (94)	fig 1219013.3.peg.725 (Q)	C7
hypothetical protein	121			fig 1219013.3.peg.766 (Q)	
hypothetical protein	73		fig 525370.5.peg.2349 (94.44)	fig 1219013.3.peg.787 (Q)	
hypothetical protein	84		fig 525370.5.peg.2301 (94.74)	fig 1219013.3.peg.835 (Q)	
hypothetical protein	147			fig 1219013.3.peg.837 (Q)	
hypothetical protein	64		fig 525370.5.peg.2262 (98.41)	fig 1219013.3.peg.874 (Q)	
hypothetical protein	109		fig 525370.5.peg.2259 (100)	fig 1219013.3.peg.877 (Q)	
hypothetical protein	45		fig 525370.5.peg.2220 (100)	fig 1219013.3.peg.916 (Q)	
hypothetical protein	40		fig 525370.5.peg.2192 (100)	fig 1219013.3.peg.944 (Q)	
hypothetical protein	53			fig 1219013.3.peg.967 (Q)	C7
hypothetical protein	41			fig 1219013.3.peg.968 (Q)	C7
hypothetical protein	40			fig 1219013.3.peg.969 (Q)	C7
hypothetical protein	43			fig 1219013.3.peg.970 (Q)	C7
hypothetical protein	51			fig 1219013.3.peg.999 (Q)	C7
Methyltransferase type 12	252			fig 1219013.3.peg.1001 (Q)	C7
hypothetical protein	38			fig 1219013.3.peg.1097 (Q)	C7
Prophage Lp2 protein 6	374			fig 1219013.3.peg.1189 (Q)	C7
hypothetical protein	73		fig 525370.5.peg.3791 (98.61)	fig 1219013.3.peg.1219 (Q)	
hypothetical protein	38			fig 1219013.3.peg.1227 (Q)	
hypothetical protein	393		fig 525370.5.peg.3707 (98.08)	fig 1219013.3.peg.1306 (Q)	
hypothetical protein	72			fig 1219013.3.peg.1476 (Q)	
hypothetical protein	43			fig 1219013.3.peg.1480 (Q)	
OsmC-like protein	136			fig 1219013.3.peg.1729 (Q)	
hypothetical protein	230			fig 1219013.3.peg.1758 (Q)	C7
hypothetical protein	55			fig 1219013.3.peg.1762 (Q)	C7
hypothetical protein	38			fig 1219013.3.peg.1764 (Q)	C7
hypothetical protein	38			fig 1219013.3.peg.1768 (Q)	C7
hypothetical protein	62			fig 1219013.3.peg.1769 (Q)	C7
hypothetical protein	77			fig 1219013.3.peg.1770 (Q)	C7
YeeC-like protein	402			fig 1219013.3.peg.1771 (Q)	C7
FIG006126: DNA helicase, restriction/modification system component YeeB	657			fig 1219013.3.peg.1772 (Q)	С7
FIG045374: Type II restriction enzyme, methylase subunit YeeA	901			fig 1219013.3.peg.1773 (Q)	C7
hypothetical protein	230			fig 1219013.3.peg.1774 (Q)	C7
hypothetical protein	275			fig 1219013.3.peg.1775 (Q)	C7
hypothetical protein	187			fig 1219013.3.peg.1776 (Q)	C7
Fe-S protein, homolog of lactate dehydrogenase SO1521	107		fig 525370.5.peg.1073 (95.74)	fig 1219013.3.peg.1778 (Q)	C7
hypothetical protein	45			fig 1219013.3.peg.1779 (Q)	C7
hypothetical protein	184			fig 1219013.3.peg.1815 (Q)	C7
hypothetical protein	513			fig 1219013.3.peg.1822 (Q)	C7
hypothetical protein	190			fig 1219013.3.peg.1823 (Q)	C7
hypothetical protein	242			fig 1219013.3.peg.1824 (Q)	C7
hypothetical protein	233			fig 1219013.3.peg.1831 (Q)	
KH domain RNA binding protein YlqC	81		fig 525370.5.peg.946 (100)	fig 1219013.3.peg.1916 (Q)	
hypothetical protein	43			fig 1219013.3.peg.2163 (Q)	
hypothetical protein	40			fig 1219013.3.peg.2285 (Q)	
hypothetical protein	60			fig 1219013.3.peg.2297 (Q)	
hypothetical protein	59			fig 1219013.3.peg.2322 (Q)	
hypothetical protein	58			fig 1219013.3.peg.2417 (Q)	
hypothetical protein	69			fig 1219013.3.peg.2511 (Q)	C7
hypothetical protein	377		fig 525370.5.peg.4061 (37.2)	fig 1219013.3.peg.2531 (Q)	
hypothetical protein	157			fig 1219013.3.peg.2544 (Q)	C7
Protein of unknown function DUF262	606			fig 1219013.3.peg.2545 (Q)	C7
Uncharacterized protein SCO1141	138			fig 1219013.3.peg.2547 (Q)	C7
hypothetical protein	78		fig 525370.5.peg.1796 (98.57)	fig 1219013.3.peg.2556 (Q)	
hypothetical protein	40			fig 1219013.3.peg.2693 (Q)	C7
hypothetical protein	46			fig 1219013.3.peg.2694 (Q)	C7
Glycosyltransferase	334		fig 525370.5.peg.1985 (98.21)	fig 1219013.3.peg.2712 (Q)	
hypothetical protein	71			fig 1219013.3.peg.2722 (Q)	C7
hypothetical protein	74			fig 1219013.3.peg.2732 (Q)	C7
hypothetical protein	120			fig 1219013.3.peg.2733 (Q)	C7
hypothetical protein	133		fig 525370.5.peg.1925 (100)	fig 1219013.3.peg.2776 (Q)	C7
hypothetical protein	38		fig 525370.5.peg.1924 (100)	fig 1219013.3.peg.2777 (Q)	ATCC33707-C7
hypothetical protein	60		fig 525370.5.peg.1907 (98.31)	fig 1219013.3.peg.2794 (Q)	
hypothetical protein	67			fig 1219013.3.peg.2846 (Q)	C7
hypothetical protein	57			fig 1219013.3.peg.2868 (Q)	

	Protein				
Function	Length	103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	141	fig 685727.5.peg.266 (32)		fig 1219013.3.peg.2873 (Q)	
hypothetical protein	203		fig 525370.5.peg.4357 (99.5)	fig 1219013.3.peg.2924 (Q)	
Oxidoreductase, short-chain dehydrogenase/reductase family (EC	211			fig 1219013.3.peg.2961 (Q)	
1.1.1)					
hypothetical protein	150			fig 1219013.3.peg.3004 (Q)	
hypothetical protein	85		fig 525370.5.peg.4395 (100)	fig 1219013.3.peg.3026 (Q)	
hypothetical protein	148			fig 1219013.3.peg.3074 (Q)	
hypothetical protein	39			fig 1219013.3.peg.3084 (Q)	C7
hypothetical protein	106		fig 525370.5.peg.3360 (97.12)	fig 1219013.3.peg.3227 (Q)	C7
hypothetical protein	43		fig 525370.5.peg.3348 (100)	fig 1219013.3.peg.3240 (Q)	
hypothetical protein	38			fig 1219013.3.peg.3280 (Q)	
Aspartyl-tRNA(Asn) amidotransferase subunit A (EC 6.3.5.6) /	67		fig 525370.5.peg.3297 (98.48)	fig 1219013.3.peg.3292 (Q)	
Glutamyi-tRNA(GIn) amidotransferase subunit A (EC 6.3.5.7)					
hypothetical protein	47		fig 525370.5.peg.3287 (95.65)	fig 1219013.3.peg.3302 (Q)	C7
hypothetical protein	70			fig 1219013.3.peg.3342 (Q)	
hypothetical protein	56			fig 1219013.3.peg.3346 (Q)	
hypothetical protein	63			fig 1219013.3.peg.3393 (Q)	C7
possible sugar phosphate isomerase/ epimerase	274		fig 525370.5.peg.3096 (99.63)	fig 1219013.3.peg.3494 (Q)	
FIG00995642: hypothetical protein	365		fig 525370.5.peg.3094 (99.18)	fig 1219013.3.peg.3496 (Q)	
hypothetical protein	38			fig 1219013.3.peg.3626 (Q)	
nutative transnosase	89		fig 525370 5 neg 2750 (80 23)	fig 1219013 3 neg 3841 (O)	
nutative acultransferace domain protein	66		16192937019196612790 (00129)	fig 1219013.3.peg.3812 (Q)	
bur ath atian anatain	00			fie   1219013.3.peg.3642 (Q)	
nypotnetical protein	39			fig 1219013.3.peg.3914 (Q)	
Ferrichrome ABC transporter	322		fig 525370.5.peg.1527 (99.38)	fig 1219013.3.peg.3919 (Q)	
hypothetical protein	48		fig 525370.5.peg.1526 (95.12)	fig 1219013.3.peg.3920 (Q)	
hypothetical protein	50		fig 525370.5.peg.1518 (100)	fig 1219013.3.peg.3928 (Q)	ATCC33707-C7
Adenylate kinase (EC 2.7.4.3)	173			fig 1219013.3.peg.3948 (Q)	
hypothetical protein	248		fig 525370.5.peg.1440 (77.42)	fig 1219013.3.peg.4001 (Q)	ATCC33707-C7
hypothetical protein	38			fig 1219013.3.peg.4007 (Q)	
hypothetical protein	40			fig 1219013.3.peg.4017 (Q)	C7
hypothetical protein	40			fig 1219013 3 peg 4090 (O)	
hypothetical protein	679		fig 525270 5 peg 1248 (99 56)	fig 1219013.3.pcg.4090 (Q)	ATCC33707-C7
hypothetical protein	722		ng 525570.5.peg.1248 (55.50)	fie   1219013.3.peg.4197 (Q)	ATCCSSTOT-CT
nypothetical protein	/23			fig 1219013.3.peg.4209 (Q)	
hypothetical protein	76			fig 1219013.3.peg.4212 (Q)	C7
hypothetical protein	457			fig 1219013.3.peg.4213 (Q)	C7
hypothetical protein	75			fig 1219013.3.peg.4214 (Q)	C7
hypothetical protein	380			fig 1219013.3.peg.4215 (Q)	C7
putative glycosyl transferase	336			fig 1219013.3.peg.4216 (Q)	C7
hypothetical protein	73			fig 1219013.3.peg.4219 (Q)	C7
hypothetical protein	222			fig 1219013.3.peg.4221 (Q)	C7
hypothetical protein	39			fig 1219013.3.peg.4246 (O)	
Believed to be an alternative form of N-formylglutamate deformylase (EC 3.5.1.68), but experimentally invalidated	435		fig 525370.5.peg.619 (97.15)	fig 1219013.3.peg.4270 (Q)	
hypothetical protein	860		fig 525370.5.peg.616 (99.28)	fig 1219013.3.peg.4273 (Q)	
hypothetical protein ABC transporter, periplasmic spermidine putrescine-binding protein PotD (TC 3.4.1.11.1)	229 336		fig 525370.5.peg.614 (98.68) fig 525370.5.peg.610 (100)	fig 1219013.3.peg.4275 (Q) fig 1219013.3.peg.4279 (Q)	ATCC33707-C7
Transcriptional regulator IcIR family	248		fig 525370 5 neg 609 (99 19)	fig 1219013 3 neg 4280 (O)	ATCC33707-07
Gentisate 1 2-diovygenase (EC 1 13 11 4)	257		18192997019196B1009 (99129)	fig 1219013.3.peg.1260 (Q)	
Chables destance (50.4.5.4.20)	100	fiel (05727 5 and 427( (27.27)		fie   1219013.3.peg.4309 (Q)	
FMN reductase (EC 1.5.1.29)	199	Tig   685727.5.peg.1276 (27.27)	· · · · · · · · · · · · · · · · · · ·	fig 1219013.3.peg.4371 (Q)	
Transcriptional regulator, ICIR family	278		fig 525370.5.peg.2082 (27.54)	fig 1219013.3.peg.4374 (Q)	
hypothetical protein	60			fig 1219013.3.peg.4377 (Q)	
hypothetical protein	48		fig 525370.5.peg.4657 (97.87)	fig 1219013.3.peg.4390 (Q)	
hypothetical protein	102		fig 525370.5.peg.540 (92.08)	fig 1219013.3.peg.4546 (Q)	
hypothetical protein	233		fig 525370.5.peg.510 (97.5)	fig 1219013.3.peg.4573 (Q)	ATCC33707-C7
hypothetical protein	48			fig 1219013.3.peg.4606 (Q)	C7
hypothetical protein	318		fig 525370.5.peg.401 (99.37)	fig 1219013.3.peg.4686 (Q)	
regulatory protein, TetR	208		fig 525370.5.peg.400 (98.55)	fig 1219013.3.peg.4687 (Q)	
hypothetical protein	39			fig 1219013 3 neg 4711 (O)	
hypothetical protein	401		fig = 25270 5 pog 245 (09 75)	fig 1210012.2 pog 4742 (Q)	
	401		fig [ 223270.3. peg. 345 (98.75)	fig 1215015.5.peg.4/43 (Q)	ATCC22707 07
nypotnetical protein	222		тів   525370.5.peg.328 (98.19)	пg 1219013.3.peg.4759 (Q)	ATCC33707-C7
hypothetical protein	93			tig 1219013.3.peg.4810 (Q)	
Enoyl-CoA hydratase (EC 4.2.1.17)	213		fig 525370.5.peg.3654 (31.58)	fig 1219013.3.peg.4824 (Q)	C7
phage integrase family protein	398		fig 525370.5.peg.1780 (28.89)	fig 1219013.3.peg.4900 (Q)	C7
hypothetical protein	56			fig 1219013.3.peg.4901 (Q)	C7
hypothetical protein	72			fig 1219013.3.peg.4902 (Q)	C7
hypothetical protein	201			fig 1219013.3.peg.4903 (Q)	C7
hypothetical protein	129			fig 1219013.3.peg.4905 (O)	C7
hypothetical protein	117			fig 1219013.3.peg.4906 (O)	C7

Function	Protein Length 103S Feature (blast hit)	ATCC33707 Feature (blast hit)	C7 Feature (blast hit)	HGT
hypothetical protein	110		fig 1219013.3.peg.4907 (Q)	C7
hypothetical protein	134		fig 1219013.3.peg.4908 (Q)	C7
hypothetical protein	332		fig 1219013.3.peg.4909 (Q)	C7
hypothetical protein	123		fig 1219013.3.peg.4910 (Q)	C7
hypothetical protein	97		fig 1219013.3.peg.4911 (Q)	C7
hypothetical protein	47		fig 1219013.3.peg.4924 (Q)	C7
hypothetical protein	364	fig 525370.5.peg.2599 (24.66)	fig 1219013.3.peg.4925 (Q)	ATCC33707
hypothetical protein	146		fig 1219013.3.peg.4926 (Q)	C7
hypothetical protein	138		fig 1219013.3.peg.4927 (Q)	C7
hypothetical protein	220		fig   1219013.3.peg.4928 (Q)	C7
hypothetical protein	107	fig 525370.5.peg.1731 (30.09)	fig 1219013.3.peg.4929 (O)	C7
hypothetical protein	143	··81 ()	fig 1219013 3 peg 4930 (O)	C7
hypothetical protein	251	fig 525370 5 neg 2151 (69 96)	fig 1219013 3 neg 4931 (O)	C7
hypothetical protein	123	hgj525570.5.pcg.2151(05.50)	fig 1219013.3.pcg.4932 (Q)	C7
Phage minor tail protein	123	fig 525270 5 pog 2140 (24.42)	fig 1219013.3.peg.4932 (Q)	ATCC22707 C7
Phage minor tail protein # Cn27	304	fig 525370.5.peg.2149 (34.43)	fig 1219013.3.peg.4933 (Q)	ATCC33707-C7
hunsthatisel protein	304	lig 525370.5.peg.2146 (29.34)	fig 1219013.3.peg.4934 (Q)	ATCC33707-C7
hypothetical protein	100		fig 1219013.3.peg.4936 (Q)	67
	246		fig 1219013.3.peg.4937 (Q)	07
hypothetical protein	112		fig 1219013.3.peg.4938 (Q)	C/
FIG00545832: hypothetical protein	109		fig 1219013.3.peg.4939 (Q)	C7
hypothetical protein	99		fig 1219013.3.peg.4940 (Q)	C7
hypothetical protein	137		fig 1219013.3.peg.4941 (Q)	C7
hypothetical protein	71		fig 1219013.3.peg.4942 (Q)	C7
hypothetical protein	128		fig 1219013.3.peg.4943 (Q)	C7
hypothetical protein	335		fig 1219013.3.peg.4944 (Q)	C7
hypothetical protein	200		fig 1219013.3.peg.4945 (Q)	C7
hypothetical protein	593		fig 1219013.3.peg.4946 (Q)	C7
hypothetical protein	534		fig 1219013.3.peg.4947 (Q)	C7
hypothetical protein	433		fig 1219013.3.peg.4948 (Q)	C7
hypothetical protein	166		fig 1219013.3.peg.4949 (Q)	C7
hypothetical protein	238		fig 1219013.3.peg.4950 (Q)	C7
hypothetical protein	100		fig 1219013.3.peg.4951 (Q)	C7
hypothetical protein	225	fig 525370.5.peg.2126 (38)	fig 1219013.3.peg.4952 (Q)	ATCC33707-C7
hypothetical protein	111		fig 1219013.3.peg.4953 (Q)	C7
hypothetical protein	71		fig 1219013.3.peg.4954 (Q)	C7
hypothetical protein	124		fig 1219013.3.peg.4955 (Q)	C7
hypothetical protein	205		fig 1219013.3.peg.4957 (Q)	C7
hypothetical protein	43		fig 1219013.3.peg.4958 (Q)	C7
hypothetical protein	116		fig 1219013.3.peg.4959 (Q)	C7
hypothetical protein	107		fig 1219013.3.peg.4960 (Q)	C7
hypothetical protein	63		fig 1219013.3.peg.4961 (Q)	C7
hypothetical protein	121		fig 1219013.3.peg.4962 (Q)	C7
hypothetical protein	41		fig 1219013.3.peg.4963 (Q)	C7
hypothetical protein	117		fig 1219013.3.peg.4964 (Q)	C7
hypothetical protein	90		fig 1219013.3.peg.4965 (O)	C7
hypothetical protein	165		fig 1219013 3 peg 4966 (O)	C7
hypothetical protein	122		fig 1219013 3 peg 4967 (Q)	C7
hypothetical protein	92		fig 1219013 3 peg 4968 (O)	C7
hypothetical protein			fig 1219013 3 peg 4969 (0)	с, С7
hypothetical protein	281	fig 525370 5 neg 2116 /20 27	fig 1219012 2 pog 4070 (Q)	ATCC 22707. C7
Dhage-related protein predicted endopuelosso	212	fig 525370.5.peg.2110 (23.37)	fig 1219013 3 pog 4071 (Q)	ATCC22707 C7
hypothetical protein	01	1151929310.3.peg.2113 (23.30)	fig 1210012 2 pog 4072 (Q)	C7
hypothetical protein	20		fig 1210012 2 mor 4072 (Q)	C7
hypothetical protein	00		fig 1219012.3.peg.4973 (Q)	C7
nypotneucal protein	90 102		fig 1219013.3.peg.4974 (Q)	C7
nypoinetical protein	102	8-1525270 5 4511 (ST S)	iig 1219013.3.peg.4975 (Q)	L/
Integrase	352	тіg   525370.5.peg.1714 (24.2)	тіg   1219013.3.peg.4976 (Q)	ATCC33707

Note: 1. Genes used as the query for blast searches are marked with (Q) whereas the blast score is mentioned with the target features.

2. Two genes of strain 103S, one of ATCC33707 and four C7 genes showed similarities with more than one features in another genome. These genes and their target genes are labelled in green, blue and red, respectively.

3. The last column HGT represents "Horizontal Gene Transfer" and we mentioned the strain IDs where the genes are borne by the genomic islands that are predicted by the program Alien Hunter.

1035		C7		ATCC 33707			
Position	Ref. allele	Locus/gene	Product	SNP allele	SNP type	SNP allele	SNP type
717240	С	fig 685727.5.peg.690	CONSERVED 13E12 REPEAT FAMILY PROTEIN			а	nonsense
777219	С	fig 685727.5.peg.749	MCE-family protein Mce1F	а	nonsense		
777362	С	fig 685727.5.peg.749	MCE-family protein Mce1F	t	nonsense		
778256	G	fig 685727.5.peg.751	MCE associated membrane protein	а	nonsense		
832375	С	fig 685727.5.peg.791	putative ABC transporter ATP binding protein	t	nonsense	t	nonsense
953696	С	fig 685727.5.peg.895	3-oxoacyl-[acyl-carrier protein] reductase (EC 1.1.1.100)			t	nonsense
1018281	А	fig 685727.5.peg.965	hypothetical protein	t	nonsense		
1046258	С	fig 685727.5.peg.994	putative esterase	t	nonsense	t	nonsense
1060941	С	fig 685727.5.peg.1008	hypothetical protein			а	nonsense
1086868	G	fig 685727.5.peg.1037	hypothetical protein			а	nonsense
1420888	G	fig 685727.5.peg.1369	YndJ	а	nonsense		
1441851	G	fig 685727.5.peg.1390	hypothetical protein			а	nonsense
1474013	G	fig 685727.5.peg.1422	Pyruvate oxidase (EC 1.2.3.3)	а	nonsense		
1719095	С	fig 685727.5.peg.1649	hypothetical protein	t	nonsense		
1766963	А	fig 685727.5.peg.1671	Ribosyl nicotinamide transporter, PnuC-like	g	nonsense	С	nonsynonymous
1806315	С	fig 685727.5.peg.1711	secreted peptidase			а	nonsense
1939596	G	fig 685727.5.peg.1836	hypothetical protein			t	nonsense
2159015	G	fig 685727.5.peg.2048	oxetanocin A resistance protein			а	nonsense
2593202	С	fig 685727.5.peg.2435	hypothetical protein	t	nonsense		
2646430	С	fig 685727.5.peg.2481	Polyhydroxyalkanoic acid synthase	t	nonsense		
2754494	С	fig 685727.5.peg.2584	Epoxide hydrolase (EC 3.3.2.9)	t	nonsense		
2755296	А	fig 685727.5.peg.2585	FIG01136303: hypothetical protein	g	nonsense	g	nonsense
2777171	С	fig 685727.5.peg.2615	hypothetical protein	t	nonsense		
3092582	С	fig 685727.5.peg.2906	hypothetical protein			а	nonsense
3296283	С	fig 685727.5.peg.3095	hypothetical protein			t	nonsense
3395747	G	fig 685727.5.peg.3187	hypothetical protein			а	nonsense
3735874	G	fig 685727.5.peg.3515	hypothetical protein	а	nonsense	а	nonsense
4133706	С	fig 685727.5.peg.3890	hypothetical protein	t	nonsense	t	nonsense
4160134	С	fig 685727.5.peg.3917	Acetate kinase (EC 2.7.2.1)	t	nonsense	t	nonsense
4300318	G	fig 685727.5.peg.4054	Long-chain-fatty-acidCoA ligase (EC 6.2.1.3)	а	nonsense		
4350321	С	fig 685727.5.peg.4088	FIG00863843: hypothetical protein	t	nonsense	t	nonsense
4379112	С	fig 685727.5.peg.4112	hypothetical protein			а	nonsense
4959414	С	fig 685727.5.peg.4661	hypothetical protein	g	nonsense		

Supplementary Table 2. A list of nonsense SNPs in *R. equi* strains C7 and ATCC33707 using 103S as the reference