Genome Update: tRNAs in sequenced microbial genomes

Genomes of the month – microbial genome evolution

Eight microbial genomes have been published in the four weeks since the last Genome Update was written (Ussery et al., 2004). They represent five bacterial and three eukaryotic organisms, and provide several interesting aspects of genome evolution. A very brief overview of the new genomes will be presented below; this is meant merely to wet the appetite of the reader and to provide pointers to the relevant recent literature.

Two spirochaete genomes have been published this month, bringing the total number of genomes from three to five for this phylum. The genome of Treponema denticola strain ATCC 35405 (Seshadri et al., 2004) is more than twice the size of the previously sequenced genome of Treponema pallidum (2.8 Mbp vs 1.1 Mbp), although the number of tRNAs and rRNAs are about the same in both genomes. The difference in genome size appears to be the result of a combination of three types of evolution: genome reduction, lineage-specific recombination and horizontal gene transfer (Seshadri et al., 2004). The other newly sequenced spirochaete genome, of Leptospira interrogans serovar Copenhageni strain Fiocruz L1-130 (Nascimento et al., 2004), has two chromosomes and encodes 3728 genes, two rRNA operons and 37 tRNAs, as shown in Table 1. This genome is nearly identical in size to that of L. interrogans serovar Lai (Ren et al., 2003), which has 4727 annotated genes, or nearly 1000 extra genes. This is perhaps due to the difference in cut-off values for gene-finding from the two different groups.

Members of the Chlamydiae are amongst the most successful bacterial pathogens of humans, and there are currently eight sequenced pathogenic chlamydial genomes, ranging in size from 1.0 to 1.2 Mbp (see table on supplemental web page). Recently, it was discovered that Chlamydia and related species can also exist in free-living amoebae, and the genome of the Acanthamoeba sp. endosymbiont Parachlamydia sp. UWE25 has now been sequenced (Horn et al., 2004); at 2.4 Mbp, it is about twice the size of the other chlamydial genomes. It is estimated that the last common ancestor for the pathogenic and symbiotic chlamydia was about 700 million years ago, and that this bacterium already contained many of the virulence factors found in modern pathogenic chlamydia (Horn et al., 2004).

The thermophilic and halotolerant bacterium Thermus thermophilus has become a model organism for structural biology, as many of its proteins have been crystallized and their structures determined. Examination of the genome of Thermus thermophilus strain HB27, which can grow at temperatures up to 85 °C, has revealed some clues as to what it might take to live in a hot-spring environment (Henne et al., 2004). Based on its genome sequence, it looks like this bacterium is a scavenger which lives on solid surfaces and takes up nutrients as they pass by.

The genome of the parasite Wollbachia pipiensis wMel is unusual in that it is both streamlined and also contains high levels of repeats and mobile DNA elements (Wu et al., 2004). Thus, for this bacterium, natural selection appears to be a bit inefficient, probably due to repeated population bottlenecks (Wu et al., 2004).

Three eukaryotic genomes have also been sequenced this month. As usual, unfortunately the quality of the eukaryotic sequences is not as good as that of the prokaryotic genomes; there are many gaps in the sequences, and also the annotation (when present) is patchy at best (in our opinion). According to Kellis et al. (2004), the genome sequence of the yeast Kluyveromyces waltii strain NCYC 2644 compared to that of Saccharomyces cerevisiae provides ‘the first comparison across an ancient whole genome duplication event and offers the opportunity to study the long-term fate of a genome after duplication’. The intracellular pathogen Cryptosporidium parvum type II isolate has a genome of about 9.1 Mbp in length and encodes a mere 3800 proteins (Abrahamsen et al., 2004). (Note that this is about the size of a medium to small bacterial proteome!) This parasite has undergone massive genome reduction and streamlining, even losing all of its mitochondrial DNA, which has been incorporated into the main chromosome. Finally, the genome of the alga Cyanidioschyzon merolae 10D (Matsuzaki et al., 2004) is 16.5 Mbp long and spread over 20 chromosomes. There are very few introns, and only three rRNA operons (see Table 1). This genome

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Chris Thomas, Editor-in-Chief

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Table 1. Summary of the published genomes discussed in this Update

Note that the accession number for each chromosome is the same for GenBank, EMBL and the DNA DataBase of Japan (DDBJ). chr., Chromosomes.

<table>
<thead>
<tr>
<th>Genome</th>
<th>Size (bp)</th>
<th>AT content (%)</th>
<th>rRNA operons</th>
<th>tRNAs</th>
<th>CDS</th>
<th>Accession nos</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Leptospira interrogans</em> serovar Copenhageni Fiocruz L1-130</td>
<td>5 260 086</td>
<td>64-9</td>
<td>2</td>
<td>37</td>
<td>3728</td>
<td>AE016823</td>
</tr>
<tr>
<td><em>Parachlamydia</em> sp. UWE25</td>
<td>2 414 465</td>
<td>65-3</td>
<td>4</td>
<td>35</td>
<td>2031</td>
<td>BX908798</td>
</tr>
<tr>
<td><em>Thermus thermophilus</em> HB27</td>
<td>1 894 877</td>
<td>31-6</td>
<td>2</td>
<td>47</td>
<td>1988</td>
<td>AE017221</td>
</tr>
<tr>
<td><em>Treponema denticola</em> ATCC 35405</td>
<td>2 843 201</td>
<td>62-1</td>
<td>6</td>
<td>44</td>
<td>2786</td>
<td>AE017226</td>
</tr>
<tr>
<td><em>Wolbachia</em> pipientis wMel</td>
<td>1 267 782</td>
<td>64-8</td>
<td>1</td>
<td>34</td>
<td>1270</td>
<td>AE017196</td>
</tr>
<tr>
<td><em>Cyanidioschyzon merolae</em> type II (8 chr.)</td>
<td>~ 9 100 000</td>
<td>70-0</td>
<td>5</td>
<td>45</td>
<td>3807</td>
<td>AEE01000000</td>
</tr>
<tr>
<td><em>Treponema denticola</em> ATCC 35405</td>
<td>16 520 305</td>
<td>45-0</td>
<td>3</td>
<td>30</td>
<td>5331</td>
<td>AP006483–AP006502</td>
</tr>
<tr>
<td><em>Kluyveromyces</em> waltii NCYC 2644 (8 chr.)</td>
<td>10 613 225</td>
<td>55-6</td>
<td>2</td>
<td>244</td>
<td>5230</td>
<td>AAD01000000</td>
</tr>
</tbody>
</table>

(A) Codon Usage

Chlamydia 7 segments

(B) Nucleotide Bias in triplet position

Chlamydia 7 segments

(C) Amino Acid Usage

Chlamydia 7 segments
provides 'a model system with a simple gene composition for studying the origin, evolution and fundamental mechanisms of [photosynthetic] eukaryotic cells' (Matsuzaki et al., 2004).

**Method of the month – comparison of tRNA genes in sequenced genomes**

The number of tRNA genes in bacterial genomes ranges from 126 in *Vibrio parahaemolyticus* to 29 in *Mycoplasma pulmonis*. Since there are a maximum of 61 possible codons (and hence different tRNA genes), some genomes obviously have missing tRNAs, although all of the genomes can code for the use of all 20 amino acids. The use of base wobble in the third position allows for a given tRNA gene to utilize certain codons which differ only in the third position. Thus, for example, in the case of the *Mycoplasma pulmonis* genome, even though there are only 29 tRNA genes, all 61 codons are found within the protein-coding sequences. However, the frequency of usage within the coding regions varies considerably – for example, of the six possible codons for leucine, UUA is used 13 272 times, whilst CUG is only used 165 times, or nearly 100-fold less.

Codon usage plots for three different phyla are shown in Fig. 1(A). Note that some codons (such as AAA and GAA) are used frequently in all phyla, whilst other codons, such as UAA, UAC and UAU, are used infrequently. A change in the third position in the codon often will code for the same amino acid, and bias in this position is correlated with changes in the AT content of the genome. For example, in the *M. pulmonis* genome mentioned above, the CUN codon usage is strongly biased towards U or A (CUC is only used 399 times, compared to 7523 for CUU and 3932 for CUA). Thus, an AT-rich genome and a GC-rich genome might code for a similar amino acid composition, but each genome would have a different third position bias, as can be seen in Fig. 1(B). Finally, the overall amino acid composition of the genomes from the three different phyla look quite similar, with the amino acids leucine, alanine, glycine and serine being most abundant, and tryptophan, cytosine, histidine and methionine being used infrequently.

A brief word should be mentioned about alternative genetic codes, where 'stop codons' can actually code for an amino acid. First, of course, in the genomes of *Mycoplasma* spp., the stop codon UGA can code for tryptophan (Yamo et al., 1985). Furthermore, selenium is incorporated into some enzymes and has been shown to be incorporated as selenocysteine, again utilizing the UGA stop codon which, with the right enzymic machinery, can code for selenocysteine in other bacterial genomes (Zinoni et al., 1987). About one-quarter of a set of bacterial genomes examined (13/54) contained potential genes incorporating selenocysteine (Wassenaar & Meinersmann, 2003). A 22nd amino acid has also been proposed, which utilizes the stop codon UAG to code for pyrrolysine (Srinivasan et al., 2002). Finally, tRNA editing describes the post-transcriptional modification of a tRNA so that it can only recognize a particular triplet; this has been described for an *Escherichia coli* or a *Bacillus subtilis* tRNA with anticodon CAU (normally encoding Met), which is modified to translate codon AUA exclusively, and is loaded with Ile (Grosjean & Björk, 2004). This may be an explanation as to why a Met tRNA is frequently found duplicated in bacterial genomes, although Met is not a frequently used amino acid.

Next month, the number of genes per genome will be discussed. At the time of writing, the bacterial genome with the fewest genes is that of *Mycoplasma genitalium*, with a mere 480 genes, whilst the largest is that of *Bradyrhizobium japonicum*, with 8317 genes.

**Supplemental web pages**

Web pages containing supplemental material related to this article can be accessed from the following url: http://www.cbs.dtu.dk/services/GenomeAtlas_supp/GenUp005/

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Horn, M., Collingro, A., Schmitz-Esser, S. & 10 other authors (2004). Illuminating the evolutionary history of *Chlamydiae*. *Science* Epub ahead of print, DOI: 10.1126/science.1096330


**Fig. 1.** Codon usage ‘rose-plots’ for genomes of the phyla *Chlamydiae*, *Proteobacteria* and *Spirochaetes*. (A) The first row is of codon usage for all the chromosomes within a given phylum, normalized to the most commonly used codon (outside circle). Note that several of the codons are rarely used, whilst others are quite common. (B) The bias of the third position of the codon. (C) Amino acid usage. Note that in all three phyla, leucine is the most commonly used amino acid.


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