Chromosomal clustering of nuclear genes encoding mitochondrial and chloroplast proteins in *Arabidopsis*

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We present a statistical analysis of chromosomal clustering among nuclear genes encoding mitochondrial or chloroplast proteins in *Arabidopsis*. For both organelles, the clustering was significantly increased above the expectation, but the clustering effect was weak, and most clusters were small and dispersed. Clustered genes showed coexpression but not more than expected, and no substantial synteny was detected in other eukaryotic genomes. We propose that the unexpected clustering results from continuous selection favoring chromosomal proximity of genes acting in the same organelle.

Introduction

Mitochondria and chloroplasts both originated from endosymbiotic events. During the course of evolutionary history, most of the essential genes required for mitochondrial and chloroplast function were transferred to the nuclear genome [1]. The organelle protein sets encoded in the nucleus today are not simply the genes originally transferred from the ancient endosymbiont but have a much more complicated genetic history. Evidence for proteins derived from the original endosymbiont, from the host genome, and even originating from other endosymbionts is found in both mitochondrial and chloroplast protein sets [2,3,4]. Due to the discrete metabolic roles of the two plant energy organelles, their origins, the mechanisms of gene transfer and the need for coordination of nuclear and organelle function in plants, it is possible that chromosomal organization of nuclear genes according to function could be an important aspect of regulation. Genes with various kinds of functional relationships are known to be in clusters on the chromosomes of a number of organisms [5–10]. Combining the experimental organelle sets [11–13] with clustering of their location and their expression in *Arabidopsis*, we have attempted to determine if any of the complex factors noted above link physical location with function or origin in this model plant.

Results

Chromosomal clustering of genes encoding targeted organelle proteins

Clusters of neighboring genes were built by finding stretches of organelle genes closer than 10 kb to another organelle gene along the five chromosomes of *Arabidopsis*. To resolve the problem of tandemly duplicated genes, clustered homologs (those with a unidirectional BLAST similarity score > 100 bits) were counted as one gene. To calculate the statistical significance of the number of clustered genes in the experimental sets of 473 mitochondrial and 664 chloroplast genes, we picked the same number of genes randomly from the genome 5000 times to generate a probability distribution (Figure 1 and Table 1). The P-values for the observed clustered genes in the mitochondrial and chloroplast sets were 0.0034 and 0.0004, which are greatly significant.

No large clusters were found using a 10-kbp cutoff. Most chloroplast and mitochondrial clusters contained two genes. Five clusters had three organelle genes, and one contained four genes. Variable numbers of other genes were found in these clusters—the largest total cluster size was observed with three mitochondrial genes and six other genes. The importance of the cutoff distance was assessed by using a number of cutoffs between 5 and 80 kbp, as summarized in Figure 2. Approximately 50% of all *Arabidopsis* genes have a neighbor within 80 kbp, hence it was not meaningful to investigate higher cutoffs. For the chloroplast genes, the observed clustering was significant (P < 0.05) at all cutoffs except 80 kbp, whereas the mitochondrial genes are generally less significantly clustered, particularly at the 5-kbp cutoff.

The clustering analysis was also made with a gene-based distance cutoff, that is, counting the number of intermediate genes rather than the number of base pairs. The cutoffs 0, 1, 2, 4 and 9 in-between genes were used. In this case, we found the clustering tendency much weaker, and, again, less significant in the mitochondrial set (only one P-value was < 0.05). The optimum was seen at two genes in between for both groups. This indicates that the genes clustered by absolute distance tend to reside...
in gene-rich regions. We confirmed this by comparing the neighbor distance distributions of the two organelle sets with genes from none of the sets. Both the chloroplast and mitochondrial sets were significantly ($P < 0.00001$ by Fisher’s exact test on four bins) enriched in genes having a close neighbor (data not shown).

For comparison, the same procedure was applied to the set of 750 proteins found in yeast (*Saccharomyces cerevisiae*) mitochondria [14]. The base-pair cutoffs were reduced in accordance with the more compact (2.15 times shorter intergenic distances) yeast genome. This displayed a similar, but weaker, pattern of gene clustering, with a minimum $P$-value of 0.0066 at 4 kbp. The clustering tendency practically disappeared when using the gene-based distance cutoffs (minimal $P$-value = 0.065 at two genes in between). The enrichment of genes with a close neighbor was also significant ($P < 0.05$).

It is striking that the yeast results were so similar to *Arabidopsis* despite the fact that a much larger set of the proteome encodes experimentally confirmed mitochondrial proteins (12% compared with 1.7% and 2.4% in the *Arabidopsis* organelle sets).

**Function and coexpression of the genes in organelle groups and clusters**

The clustered genes were found to encode proteins with a wide set of biological functions, and covered a broad range of expression levels based on the available EST information (see Supplementary Material Table 1). We found no evidence for greater enrichment of targeting presequences in the clustered genes compared with the organelle sets as a whole. Also, no increased coexpression among genes in the same cluster compared with the average coexpression among all genes in the same organelle sets was observed. The lack of elevated coexpression in chromosomal clusters might be caused by the fact that the nuclear organelle genes are often coexpressed [mean Pearson correlation coefficient ($r$) = 0.35 and 0.11 for chloroplast and mitochondria, compared with 0.01 for random gene pairs and 0.03 for close (<10 kbp) chromosomal neighbors].

**Table 1. Gene and clustering statistics using a 10-kbp cutoff, and $P$-values of the observed number of clustered genes relative to the random distribution for a range of distance cutoffs**

<table>
<thead>
<tr>
<th>Organelle set</th>
<th>Total number of genes</th>
<th>Observed number of clusters (10 kbp)</th>
<th>Number of clustered genes (10 kbp)</th>
<th>$P$-value at different cutoffs (kbp)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Observed</td>
<td>Above expected</td>
<td>5</td>
</tr>
<tr>
<td>Mitochondrial</td>
<td>473</td>
<td>29</td>
<td>62</td>
<td>22.7</td>
</tr>
<tr>
<td>Chloroplast</td>
<td>664</td>
<td>55</td>
<td>111</td>
<td>37.2</td>
</tr>
</tbody>
</table>

Figure 1. Chromosomal clustering of genes for mitochondrial and chloroplast proteins. Distribution of cluster sizes in 5000 random samples of (a) chloroplast and (b) mitochondrial gene groups using a 10-kbp cutoff. The observed real number of clustered genes is shown with black arrows.
only greatly coexpressed ($r > 0.77$) clustered gene pairs gave fractions also statistically indistinguishable from the complete organelle sets. The occurrence of elevated coexpression in the clusters thus seems to correspond to the general degree of functional coupling, but does not exceed it.

**Clustering of orthologous genes between Arabidopsis and rice**

We analyzed the degree of clustering among orthologs to the mitochondrial and chloroplast genes in *Oryza sativa* (rice), using the InParanoid program [15]. The fraction of *Arabidopsis* nuclear organelle genes with orthologs was great: 52% for mitochondrial genes and 69% for chloroplast ones, whereas the whole-genome average was 26%. However, conservation of the gene clusters was scant – we extracted all possible gene pairs in each cluster in each species, and looked for such pairs that were found clustered in both species. Only 1 of the 131 clustered *Arabidopsis* gene pairs had conserved their nearby genomic position in *O. sativa* (Table 2). Even when adopting a much looser limit (*Arabidopsis* pairs at <60 kbp and *O. sativa* orthologs at <1 Mbp) we found conservation of only two (of 179) mitochondrial and five (of 287) chloroplast pairs (Table 2). This is approximately the same as the fraction of such neighbors in the whole genome (~1.0%); hence we found no evidence of increased conservation of organelle clusters between rice and *Arabidopsis*. No clustered mitochondrial pairs defined in *Arabidopsis* were found conserved in four other eukaryotes (*S. cerevisiae, Caenorhabditis elegans, Drosophila melanogaster* and *Homo sapiens*).

**Discussion**

If a biological mechanism is driving the statistically significant coupling of genes in the two organelle groups

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**Table 2. Pairs of genes that have intergenic distance < 60 kbp in Arabidopsis and orthologs in rice (O. sativa) at a distance < 1 Mbp**

<table>
<thead>
<tr>
<th>Arabidopsis</th>
<th>Rice</th>
<th>Genes in between</th>
<th>Genes in between*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mitochondrion</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At2g43780</td>
<td>At2g43750</td>
<td>4122</td>
<td>13</td>
</tr>
<tr>
<td>At5g50850</td>
<td>At5g50810</td>
<td>13 166</td>
<td>3</td>
</tr>
<tr>
<td><strong>Chloroplast</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At1g21750</td>
<td>At1g21650</td>
<td>41 615</td>
<td>9</td>
</tr>
<tr>
<td>At3g56940</td>
<td>At3g56910</td>
<td>6227</td>
<td>2</td>
</tr>
<tr>
<td>At4g25100</td>
<td>At4g25090</td>
<td>6521</td>
<td>1</td>
</tr>
<tr>
<td>At5g35170</td>
<td>At5g35100</td>
<td>58283</td>
<td>6</td>
</tr>
<tr>
<td>At5g42765</td>
<td>At5g42650</td>
<td>51 146</td>
<td>11</td>
</tr>
</tbody>
</table>

*aMinimal value of alternative orthologs.*

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Figure 2. Number of clustered genes in chloroplasts and mitochondria as a function of the clustering cutoff. The chloroplast groups were clustered under the same cutoffs as the mitochondrial ones, but their points were shifted to the left for visibility. Observed real data points for the groups are denoted with arrows. The bars denote 95% confidence intervals.
to be chromosomal neighbors in the Arabidopsis nuclear genome, it seems to be operating mostly at the intergenic distance of ~10 000–20 000 base pairs (Figure 2). Open chromatin domains extending >2–5 genes in C. elegans chromosomes have been identified that contain coexpressed genes responsible for tissue-specific functions in the worm muscle [7]. Also, genes that are strongly expressed in a variety of human tissues or that are coexpressed in yeast are known to be clustered along the chromosomes across similarly sized intergenic regions [6,10,16].

However, we show here that coexpression can be largely ruled out as the driving force for the observed clustering. There have been many other reports that physical clustering is not required for coexpression or regulation of genes for organelle proteins. Although insertion into the vicinity of other genes encoding organelle proteins could initially provide signals required for expression or targeting (or both), coexpression of genes encoding subunits of multisubunit complexes can be readily achieved without physical clustering [17,18]. A study analyzing expression of 3292 genes enriched for nuclear-encoded plastid genes defined coexpression of only three gene pairs, when strongly homologous genes were excluded [19].

Could the insertion sites of the original gene transfer be the cause of some of the clusters? In a range of studied cases of gene transfers the genes are located adjacent to or inserted into a gene encoding a mitochondrial protein, for example RPS11 (a gene encoding a mitochondrial protein of the small ribosomal subunit) in rice [20], RPS10 in carrot and fuchsia [21], succinate dehydrogenase RPS10 for example inserted into a gene encoding succinate dehydrogenase subunit 2 – the cause of some of the clusters? In a range of studied genes pairs, when strongly homologous genes were excluded [19].

As study encoding subunits of multisubunit complexes can be read-into the vicinity of other genes encoding organelle proteins could initially provide signals required for expression or targeting (or both), coexpression of genes encoding subunits of multisubunit complexes can be readily achieved without physical clustering [17,18]. A study analyzing expression of 3292 genes enriched for nuclear-encoded plastid genes defined coexpression of only three gene pairs, when strongly homologous genes were excluded [19].

The preferential location of the clustered genes in gene-dense regions serves as additional evidence of a post-insertion rearrangement.

A mixture of other factors might also be responsible for the observed chromosomal clustering of genes for organelle proteins in the Arabidopsis genome. Further functional genomics studies will be necessary to identify specific patterns of functional coupling in individual species. The discovered clustering could be important in our understanding of the evolutionary history of gene transfer and activation, and the co-inheritance of this material in particular lineages.

Supplementary material
The data and methods for our analysis are described in the supplementary material associated with this article, which can be found online at doi:10.1016/j.tig.2006.09.002.

References
Overlapping genes as rare genomic markers: the phylogeny of γ-Proteobacteria as a case study

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Phylogenies can be constructed in many ways, including using shared complex characters known as rare genomic changes (RGCs), such as insertions and deletions (indels), retroposon integrations and intron positions. Here, we demonstrate that distance-based phylogenies, which were determined by shared overlapping genes from 13 completely sequenced γ-Proteobacteria genomes, are consistent with phylogenies based on 16S rRNAs and other robust markers. These findings suggest that overlapping genes could provide interesting additional insights into the phylogenomics of completely sequenced microbial genomes.

Introduction

With more and more completely sequenced genomes available, phylogenies based on whole genomes capture more phylogenetic signatures and are less influenced by anomalous events than those based on single genes. Along with sequence-based methods, which mainly involve the comparison of primary sequences, gene content and gene order are currently used in phylogenetic analyses of completely sequenced genomes; see Delsuc et al. [1] for a detailed review. However, gene content might prove to change too little, and gene order might change too much, for adequate analyses to be performed [2]. Genomes can also be compared by looking for shared complex characters known as rare genomic changes (RGCs), such as indels, retroposon integrations and intron positions. Until recently, only a few types of RGC character have been used for inferring phylogenetic relationships among completely sequenced genomes; for example, Gupta et al. [3,4] deduced the branching order for bacterial groups using conserved indels in protein sequences, and Yang et al. [5] determined prokaryotic phylogeny based on protein domain content (see also Rokas and Holland [6] for a more detailed review).

Orthologous overlapping genes among bacterial genomes

Overlapping genes in bacterial genomes are adjacent genes whose coding sequences partially or entirely overlap. Intuitively, we would assume that this character does not evolve as slowly as gene content because the formation of overlapping genes is more frequently observed than the variation of gene repertoires among completely sequenced genomes, especially in closely related genomes [7,8]. By contrast, overlapping genes might be more conserved than gene order during the course of evolution, because functional constraints might prevent breaking of the linkage of two overlapping genes [9–11]. The rate of evolution is also expected to be slower for stretches of DNA encoding overlapping genes than for similar DNA sequences that encode different proteins.

The complete sequences of 13 γ-Proteobacteria genomes were downloaded from GenBank in August 2004. Open reading frames (ORFs) annotated as ‘hypothetical’ or ‘putative’, or with products annotated as ‘unknown

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