Genome-wide selective sweeps in natural bacterial populations revealed by time-series metagenomics

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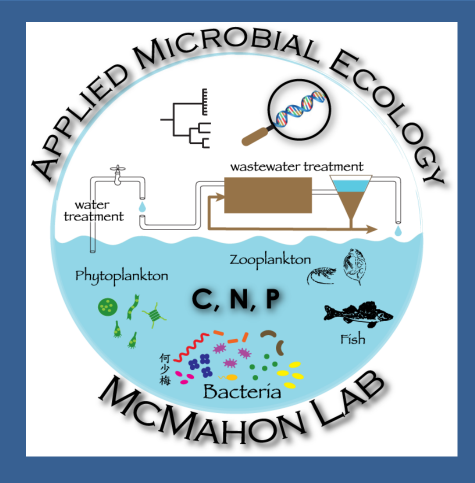
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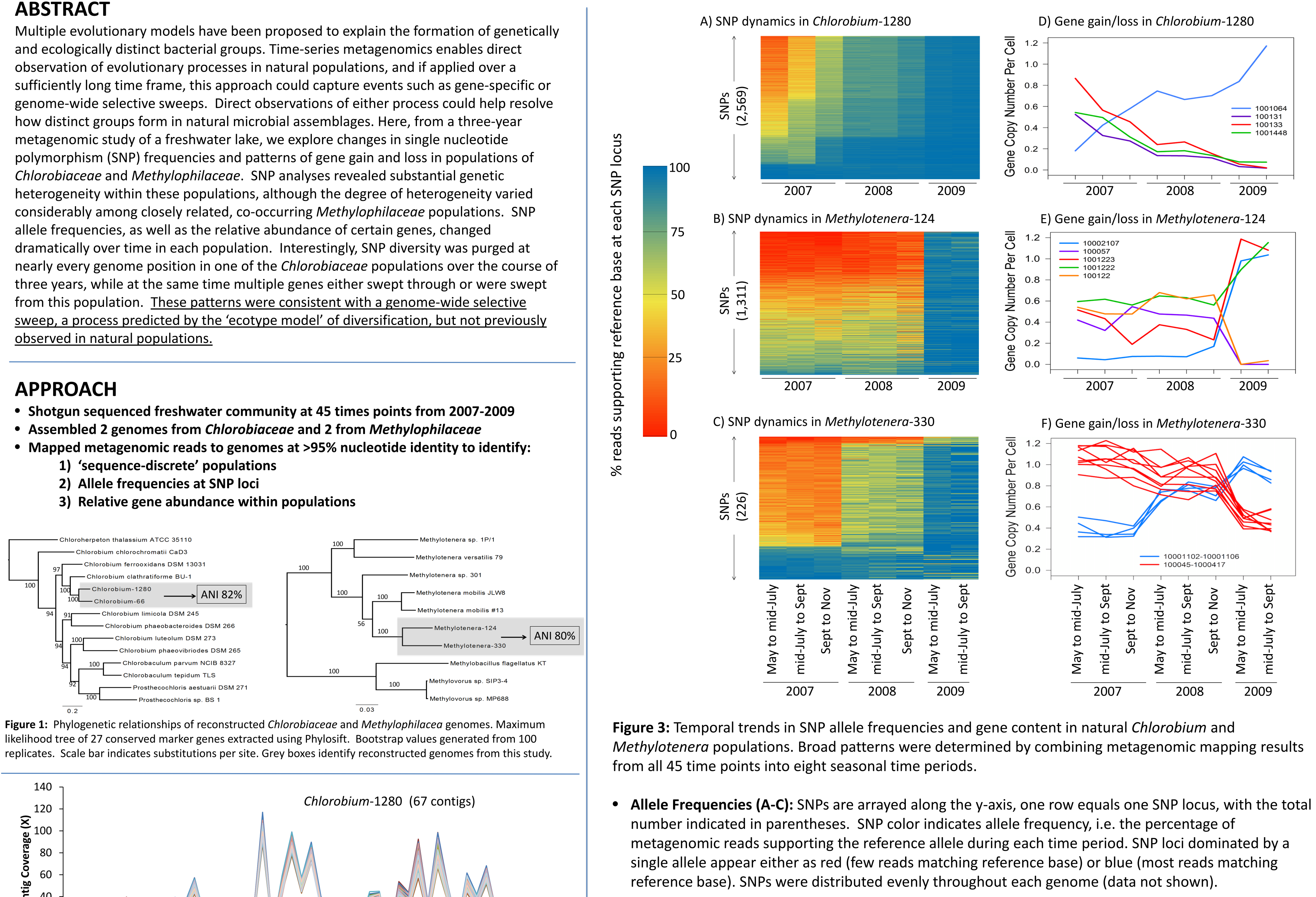
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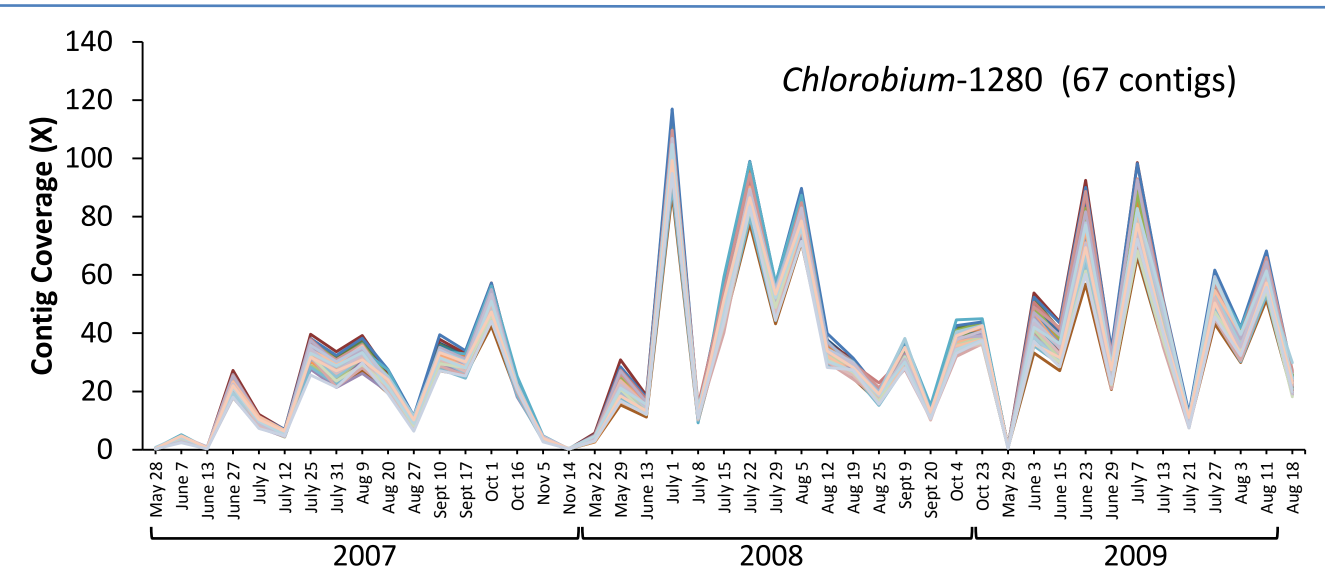
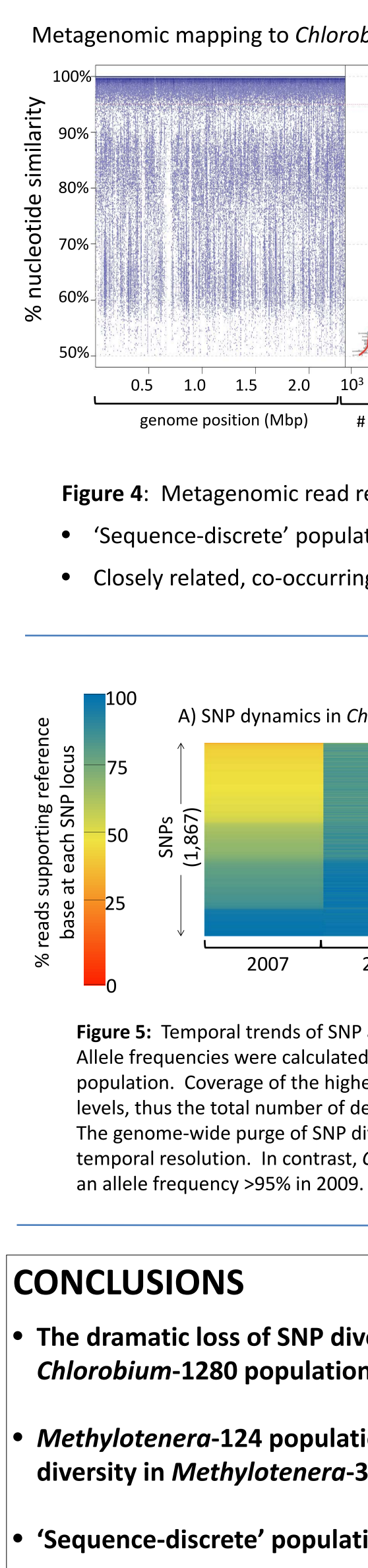


Figure 2: Temporal contig coverage patterns. Each contig is represented by a different colored line. The tight synchronization of contig coverage within each genome bin indicates these contigs were derived from the same organism.

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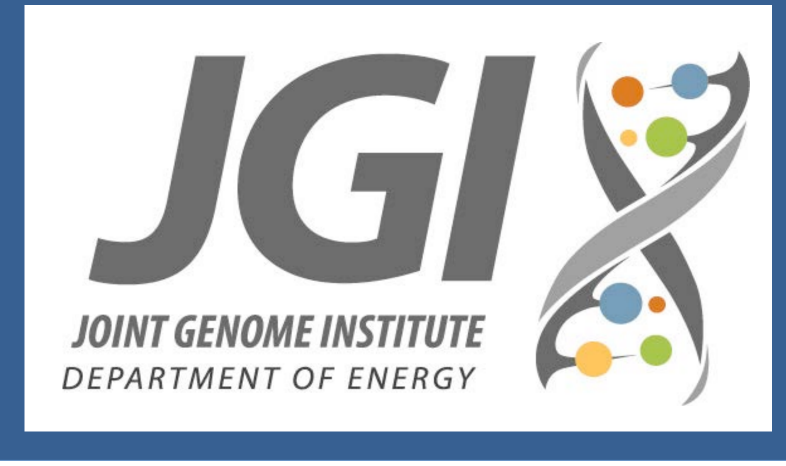
- Gene Gain and Loss (D-F): Relative abundance of genes gained or lost from *Chlorobium* and Methylotenera populations. Copy number per cell determined as coverage of a gene divided by median coverage of all other genes in genome. Gene locus id's are indicated in the legends. Two sets of contiguous genes were gained and lost from *Methylotenera*-330, and genes in each set were plotted with the same line color.



- independently

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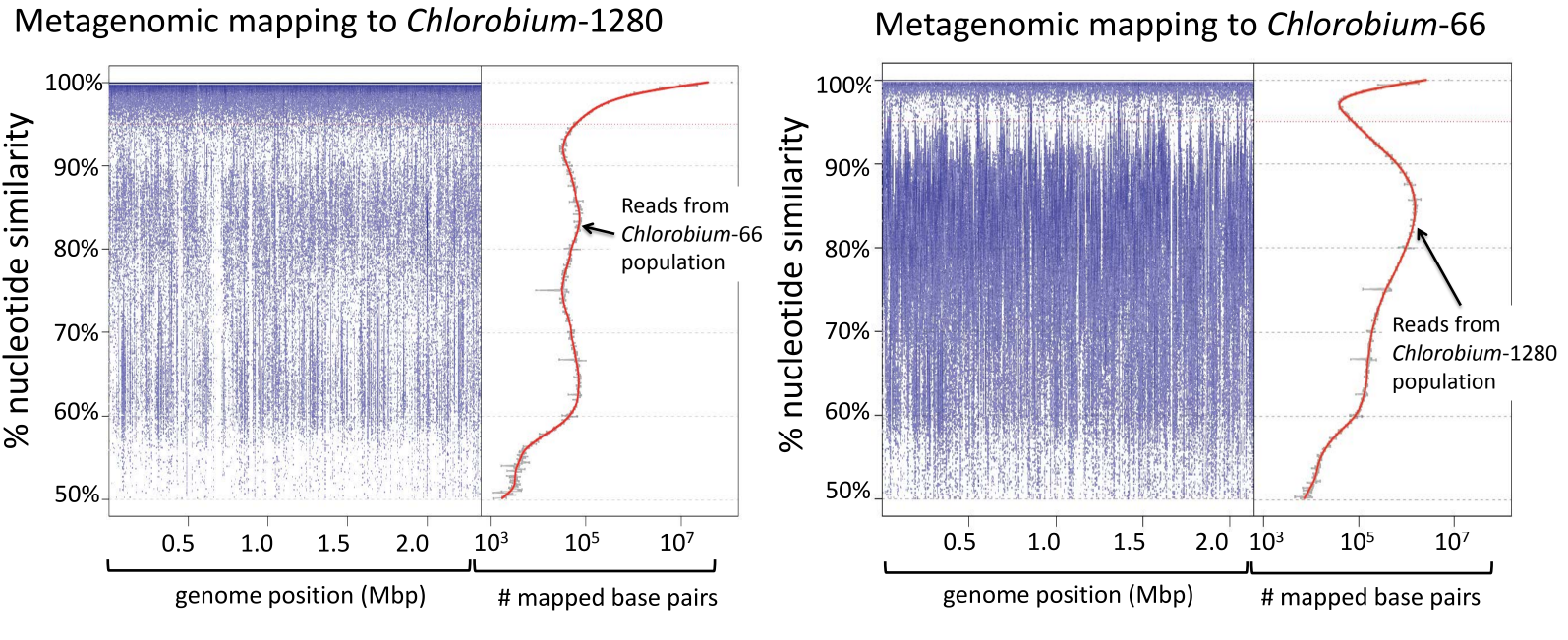


Figure 4: Metagenomic read recruitment to *Chlorobium* genomes.

• 'Sequence-discrete' populations revealed by reads that mapped with >95% nucleotide identity • Closely related, co-occurring populations separated by coverage discontinuity at ~95% identity

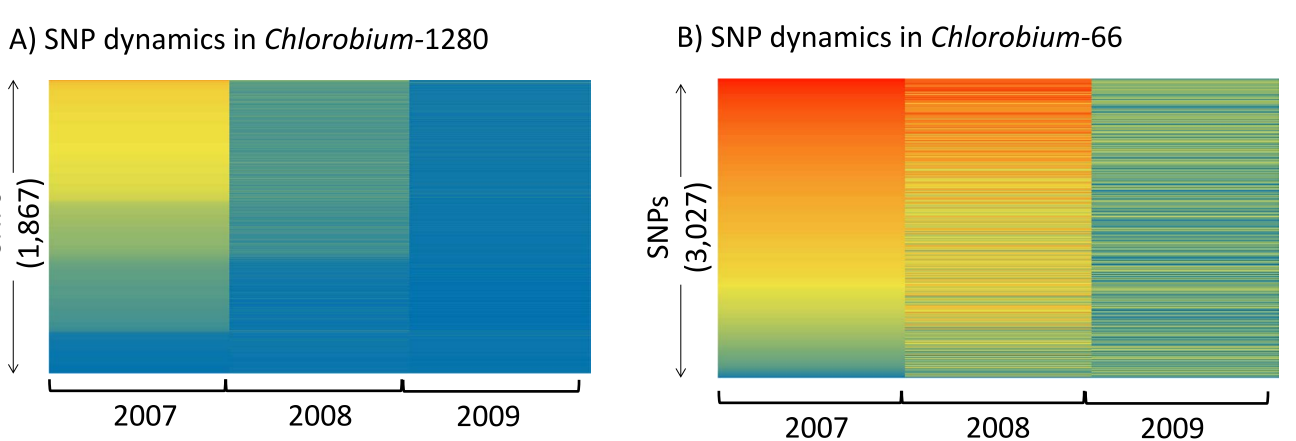


Figure 5: Temporal trends of SNP allele frequencies in *Chlorobium*-1280 (A) and *Chlorobium*-66 (B) populations. Allele frequencies were calculated per year due to coverage limitations in the less abundant *Chlorobium*-66 population. Coverage of the higher abundance *Chlorobium*-1280 was informatically reduced to comparable levels, thus the total number of detected SNPs was lower than reported when all data were used (see Figure 3). The genome-wide purge of SNP diversity in *Chlorobium*-1280 was still apparent with lower coverage and temporal resolution. In contrast, *Chlorobium*-66 did not experience a similar purge; only 414 of the SNP loci had

• The dramatic loss of SNP diversity and the patterns of gene gain and loss in the *Chlorobium*-1280 population were consistent with a genome-wide selective sweep.

• *Methylotenera*-124 population may have experienced a 'soft sweep,' whereas most SNP diversity in *Methylotenera*-330 population was lost prior to the start of this study.

• 'Sequence-discrete' populations behave like theoretically defined 'ecotypes'

• Displacement of many co-existing strains by a single strain/lineage within the same population implies that all population members shared the same ecological niche.

Closely related, co-occurring sequence-discrete populations experience sweeps