The Whole Genome Sequence of *Sphingobium chlorophenolicum* L-1: Insights into the Evolution of the Pentachlorophenol Degradation Pathway

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Data deposition: GenBank accession numbers for the replicons of *S. chlorophenolicum* are given in table 1.

Abstract

*Sphingobium chlorophenolicum* Strain L-1 can mineralize the toxic pesticide pentachlorophenol (PCP). We have sequenced the genome of *S. chlorophenolicum* Strain L-1. The genome consists of a primary chromosome that encodes most of the genes for core processes, a secondary chromosome that encodes primarily genes that appear to be involved in environmental adaptation, and a small plasmid. The genes responsible for degradation of PCP are found on chromosome 2. We have compared the genomes of *S. chlorophenolicum* Strain L-1 and *Sphingobium japonicum*, a closely related Sphingomonad that degrades lindane. Our analysis suggests that the genes encoding the first three enzymes in the PCP degradation pathway were acquired via two different horizontal gene transfer events, and the genes encoding the final two enzymes in the pathway were acquired from the most recent common ancestor of these two bacteria.

Key words: horizontal gene transfer, biodegradation, enzyme evolution, pentachlorophenol hydroxylase, tetrachlorohydroquinone dehalogenase, tetrachlorobenzoquinone reductase.

Introduction

In the early 20th century, consumers embraced the concept of “better living through chemistry.” Massive quantities of chlorinated anthropogenic chemicals such as dichlorodiphenyltrichloroethane (DDT), lindane, pentachlorophenol (PCP), atrazine, tetrachloroethylene, and polychlorinated biphenyls (PCBs) were introduced into the environment with little concern over their ultimate fate. Many of these compounds persist in the environment and have adverse effects on ecosystems. Use of some of these compounds (e.g., DDT and PCP) is now restricted or prohibited in the United States, although some are still used in developing countries.

The introduction of anthropogenic chemicals into the environment imposes selective pressures that can foster evolution of novel pathways that allow microbes to access new sources of carbon, nitrogen, or phosphorus and/or to detoxify toxic compounds. However, degradation of anthropogenic chemicals is often inefficient because microbes have not yet evolved enzymes that efficiently catalyze the steps needed to convert anthropogenic compounds into metabolites in central carbon metabolism (Copley 2009).

The pesticide PCP is listed as a Priority Pollutant by the US EPA due to its toxicity and persistence in the environment. A number of microbes capable of mineralizing PCP have been isolated from contaminated sites (Saber and Crawford 1985; Apajalahti and Saikinosa-Salonen 1987; Schenk et al. 1990; Uotila et al. 1991; Radehaus 1992; Karn et al. 2010). The best studied of these is *Sphingobium chlorophenolicum* (Saber and Crawford 1985; Steiert and Crawford 1986; Crawford...
Sphingobium chlorophenolicum has patched together a poorly functioning pathway (see fig. 1) that allows mineralization of PCP, although degradation is slow and the bacterium is unable to grow at high concentrations of PCP (Dai and Copley 2004). The first three enzymes in the pathway are unusually ineffective. PCP hydroxylase (PcpB) has a very low $k_{\text{cat}}$ of $0.02 \text{s}^{-1}$ and shows substantial uncoupling; the enzyme frequently fails to hydroxylate the substrate and instead releases H$_2$O$_2$ as the C4a-hydroperoxyflavin intermediate decomposes (Hlouchova K, Rudolph J, Pietari JMH, Behlen LS, Copley SD, unpublished data). Tetrachlorobenzoquinone (TCBQ) reductase (PcpD) is also rather inefficient, with a turnover rate of only $0.7 \text{s}^{-1}$ at 50 $\mu$M TCBQ (Dai et al. 2003). Tetrachlorohydroquinone (TCHQ) dehalogenase (PcpC) is subject to profound substrate inhibition; binding of its aromatic substrates to the active site before completion of the catalytic cycle interferes with the final step required to regenerate the free enzyme (Warner and Copley 2007a). These findings are consistent with the hypothesis that the enzymes in the PCP degradation pathway have not yet evolved to the level of function typical of most metabolic enzymes. Thus, S. chlorophenolicum provides a window on the process of assembly of a new pathway at a relatively early stage in its evolution.

We recently sequenced the genome of S. chlorophenolicum L-1. Analysis of this sequence and comparison with the sequence of the closely related Sphingobium japonicum, which degrades lindane (Nagata et al. 2007), provides insights into the origins of the PCP degradation enzymes. The rDNA genes of S. chlorophenolicum and S. japonicum share 97% identity. The phylogenetic distance between S. chlorophenolicum and S. japonicum is ideal to facilitate identification of genes that were present in the most recent common ancestor of these two Sphingomonads, as proteins involved in core processes show >80% pairwise identities at the amino acid level.

Our analysis suggests that the first three enzymes in the pathway were acquired by S. chlorophenolicum by horizontal gene transfer (HGT) after it diverged from S. japonicum. In contrast, the last two enzymes in the pathway were present in the most recent common ancestor of S. chlorophenolicum and S. japonicum. None of the genes encoding the PCP degradation enzymes arose by recent duplication and divergence of genes within S. chlorophenolicum. The genes occur in two disparate parts of the genome and have not yet been integrated into a compact and consistently regulated operon.

**Materials and Methods**

**Isolation of Genomic DNA from S. chlorophenolicum L-1**

A sample of S. chlorophenolicum L-1 was originally deposited to the American type Culture Collection and designated ATCC 39723. The initial characterization of the enzymes in the PCP degradation pathway was done with the ATCC 39723 strain (Orser and Lange 1994; Copley 2000; Dai et al. 2003; Warner and Copley 2007b). However, the ATCC 39723 strain lost the ability to degrade PCP, so a second sample of S. chlorophenolicum L-1 was deposited and designated ATCC 53874. Genomic DNA was isolated from this strain.

A 5 ml culture of ATCC medium 1687 (Flavobacterium medium) containing 5 $\mu$M PCP was inoculated with a colony from a fresh agar plate of half-strength tryptone soy broth.
containing 50 μM PCP and incubated with shaking at 30 °C overnight. One milliliter aliquots of cells were harvested by centrifugation at 16,100 × g for 1 min. Genomic DNA was isolated using the Aquapure Genomic DNA kit (Bio-Rad, Hercules, CA). Cell pellets from 1 ml of culture were resuspended in 300 μl Genomic DNA Lysis Solution. The solution was incubated at 80 °C for 5 min to lyse the cells. A solution of RNase A (1.5 μl, 4 mg/ml) was added, and tube was inverted 25 times before incubation at 37 °C for 45 min. The temperature of the sample was adjusted to room temperature and 100 μl of Protein Precipitation Solution was added. The sample was vortexed for 20 s and then subjected to centrifugation at 13,000 × g for 3 min to remove cellular debris. The supernatant containing DNA was transferred to a clean 1.5 ml tube, incubated on ice for 5 min, and then subjected to centrifugation again. The supernatant was then transferred to a clean 1.5-ml tube containing 300 μl 100% isopropanol to precipitate the DNA. The tube was inverted 50 times. The DNA was pelleted by centrifugation at 13,000 × g for 1 min. The supernatant was poured off, and the pellet was washed with 300 μl 70% ethanol. The ethanol was poured off and residual ethanol removed by draining the tube onto filter paper for 10–15 min. The pellet was dissolved in 100 μl 10 mM Tris–HCl, pH 7.5, at 65 °C for 5 min and then at room temperature overnight. The quality of the DNA was checked on a 0.8% agarose 1X TAE gel alongside a 1 kb DNA Extension Ladder (Invitrogen, Carlsbad, CA), and the concentration was measured on a NanoDrop spectrophotometer.

Genome Sequencing

The draft genome of *S. chlorophenolicum* L-1 was generated at the DOE Joint genome Institute (JGI) using a combination of illumina (Bennett 2004) and 454 technologies (Margulies et al. 2005). Three libraries were constructed and sequenced: an illumina G4ii shotgun library that generated 40,283,193 reads totaling 3,061 Mb; a 454 Titanium standard library that generated 302,660 reads and a paired end 454 library with an average insert size of 11.26 ± 2.81 kb that generated 174,361 reads. In total, 176.8 Mb of 454 data were obtained. General aspects of library construction and sequencing performed at the JGI can be found at [http://www.jgi.doe.gov/](http://www.jgi.doe.gov/). The initial draft assembly contained 79 contigs in one scaffold. The 454 Titanium standard data and the 454 paired end data were assembled with Newbler, version 2.3. The Newbler consensus sequences were computationally shredded into 2 kb overlapping fake reads (shreds). Illumina sequencing data was assembled with VELVET, version 0.7.63 (Zerbino 2008), and the consensus sequences were computationally shredded into 1.5 kb shreds. The 454 Newbler consensus shreds, the Illumina VELVET consensus shreds, and the read pairs in the 454 paired end library were integrated using parallel phrap, version SPS—4.24 (High Performance Software, LLC). The software Consed (Ewing and Green 1998; Ewing et al. 1998; Gordon et al. 1998) was used in the finishing process. Illumina data were used to correct potential base errors and increase consensus quality using the software Polisher developed at JGI (Lapidus A, unpublished data). Possible misassemblies were corrected using gapResolution (Han C, unpublished data) or Dupfinisher (Han 2006) or by sequencing cloned bridging PCR fragments. Gaps between contigs were closed by editing in Consed, by PCR, and by Bubble PCR (Cheng J-F, unpublished data) primer walks. A total of 87 additional finishing reactions (either sequencing of bridging PCR fragments or primer walking) were necessary to close gaps and to raise the quality of the finished sequence. The total size of the genome is 4,573,221 bp. The final assembly is based on 176.8 Mb of 454 draft data that provides an average 36× coverage of the genome and 3,553 Mb of Illumina draft data that provides an average 790× coverage of the genome.

Genes were identified using Prodigal (Hyatt et al. 2010) as part of the Oak Ridge National Laboratory genome annotation pipeline, followed by a round of manual curation using the JGI GenePRIMP pipeline (Pati et al. 2010). The predicted coding sequences were translated and used to search the National Center for Biotechnology Information nonredundant database, UniProt, TigrFam, Pfam, PRIAM, KEGG, COG, and InterPro databases. These data sources were combined to assert a product description for each predicted protein. Noncoding genes and miscellaneous features were predicted using tRNAscan-SE (Lowe and Eddy 1997), RNAmmer (Lagesen et al. 2007), Rfam (Griffiths-Jones et al. 2003), TMHMM (Krogh et al. 2001), and signalP (Bendtsen et al. 2004).

Results and Discussion

The *S. chlorophenolicum* Genome

The *S. chlorophenolicum* genome consists of two chromosomes and a plasmid (see fig. 2). A summary of the characteristics of each replicon is provided in table 1. Like many bacteria such as *Burkholderia pseudomallei* (Holden et al. 2004) and *Vibrio cholerae* (Heidelberg et al. 2000) that contain multiple replicons, *S. chlorophenolicum* has a dominant chromosome that carries most of the genes for core processes. We generated a list of genes involved in replication, transcription, translation, cell division, peptidoglycan biosynthesis, and core metabolic processes (including glycolysis, the pentose phosphate pathway, the TCA cycle, electron transport, and biosynthesis of amino acids, nucleotides, and cofactors) (see supplementary table 1, Supplementary Material Online). Figure 3 shows the distribution of genes between the two chromosomes. If genes encoding core functions were randomly distributed between the chromosomes, we would expect 354 (72%) to be on chromosome 1.
and 138 (28%) to be on chromosome 2. In fact, 442 core genes (90%) are on chromosome 1 and only 50 (10%) are on chromosome 2. This deviation from the expected distribution is highly significant by chi-square analysis ($P = 2.2 \times 10^{-16}$). An odd collection of essential enzymes, including a few genes for central carbon metabolism, two subunits of RNA polymerase, some components of the electron transfer chain, and a few genes for cofactor biosynthesis are found on chromosome 2 rather than on chromosome 1. However, most of the genes on chromosome 2 appear to be involved in environmental adaptation. Chromosome 2 carries a number of genes predicted to encode transporters for sugars and metal ions and enzymes involved in degradation of various sugars, short-chain fatty acids, and aromatic compounds. All the PCP degradation genes are found on chromosome 2. A secondary chromosome with a low density of essential genes may serve as a convenient storage depot for genes acquired by HGT, as integration of newly acquired DNA is not likely to disrupt an essential function. Chromosome 2 also contains two genes encoding proteins related to ParB, which is involved in plasmid partitioning. These features of chromosome 2 are consistent with the proposal that bacterial secondary chromosomes have arisen by intragenomic transfer of essential genes, including rRNA genes, to a plasmid (Slater et al. 2009).

HGT is rampant among microbes and is known to play a major role in acquisition of resistance to or degradation of toxic compounds, including antibiotics. We examined the *S. chlorophenolicum* genome for features indicative of mobile genetic elements. As mentioned above, *S. chlorophenolicum* contains one plasmid. The genome appears to contain one integrated prophage; a cluster of phage-related

![Fig. 2.—Circle diagram of the replicons of *Sphingobium chlorophenolicum*. The second circle in each replicon indicates the locations of PCP degradation genes in black, phage genes in blue, and transposon-related genes in red. The third circle shows locations of genes that have a close homolog in *Sphingobium japonicum* (>80% identity over >90% of the *S. chlorophenolicum* sequence). The fourth circle shows GC content. Red indicates sequences with GC content <64% and blue indicates sequences with GC content >64%. Light gray lines are placed at intervals of one standard deviation from the mean of 63.8%. (One standard deviation = 0.027%; GC content was calculated by averaging over a 10-kb window and sliding that window in 1-kb increments.) Diagram was generated using Circos (Krzywinski et al. 2009).](https://academic.oup.com/gbe/article-abstract/4/2/184/556989)

### Table 1

Genome Statistics

|              | Length (bp) | GC Content | ORFs | rRNA Operons |
|--------------|-------------|------------|------|--------------|
| Chromosome 1 (NC_015593) | 3080818     | 0.64       | 2,940 | 1            |
| Chromosome 2 (NC_015594) | 1368670     | 0.64       | 1,159 | 2            |
| Plasmid (pSPCH01) (NC_015595) | 123733 | 0.65 | 125 | 0            |
genes (including a major capsid protein, major tail protein, phage portal protein, pro-head peptidase, and some conserved phage proteins of unknown function) is found on chromosome 1 (genes Sc_00004260-00004380). Curiously, seven isolated genes annotated as “phage integrase family” genes are found on both chromosomes 1 and 2 (Chr1: Sc_00026840, Sc_00029370, Sc_00022480, Sc_00028520, Sc_00012030; Chr 2: Sc_00031410, Sc_00030440). These genes have anomalously low GC content (0.48–0.58) and are not closely related to each other. Only one (Sc_00026840) is found in the vicinity of a prophage gene (a CP4-57 regulatory protein). Two (Sc_00017260 and Sc_00026840) are adjacent to transposase genes. The genome also carries 27 sequences annotated as “transposase,” “transposase/integrase core domain,” or “transposase and inactivated derivatives” (16 on chromosome 1, 9 on chromosome 2, and 2 on the plasmid). These proteins are related to transposases found in several different families of insertion elements and in Tn3 family transposons. Thus, like most microbial genomes, the genome of S. chlorophenolicum displays evidence of continual onslaught by mobile genetic elements. However, the PCP degradation genes show no association with any of these elements.

New genes that can help microbes survive and grow in the face of selective pressure from environmental toxins can also arise by gene duplication and divergence (Hughes 1994; Berghthorsson et al. 2007). We carried out a Blast search of the S. chlorophenolicum genome against itself to identify genes that are nearly identical and may have arisen by recent gene duplication. Only 38 proteins have >90% sequence identity to another protein in S. chlorophenolicum. Of these, 13 are related to transposases; three of these genes are found in three identical copies scattered throughout the genome. Although the presence of highly similar transposase genes is not unexpected, there is little rhyme or reason to the identities and locations of the remaining duplicated genes. Seven genes annotated as encoding 2-hydroxychromene-2-carboxylate isomerase, a short-chain alcohol dehydrogenase, a hypothetical protein, a nucleoside-diphosphate-sugar epimerase, an arabinose efflux permease, a Zn-dependent dipeptidase, and an outer membrane receptor protein are present between positions 667462 and 675176 on the —strand of chromosome 2. A second copy of five of these genes are found in a cluster on the + strand of chromosome 2 between positions 1036969 and 1030812. Copies of the first two genes in the cluster are found together in a different location on chromosome 2. Nearly identical copies of the genes encoding the alpha and beta subunits of the E1 component of pyruvate dehydrogenase are found on chromosomes 1 and 2 and nearly identical copies of a gene annotated as an acyl-CoA synthetase/AMP-acid ligase II are found in chromosome 2 but surrounded by completely different genes. The utility of these duplicated genes and the processes leading to their location in distant parts of the genome are not clear. One possibility is that they were not actually duplicated in S. chlorophenolicum but rather acquired by integration into the genome of DNA fragments taken up from lysed cells of S. chlorophenolicum or closely related Sphingomonads. Notably, none of the PCP degradation genes is found among the set of highly similar genes.

Comparison of the Genomes of S. chlorophenolicum and S. japonicum

The genomes of S. chlorophenolicum and S. japonicum are of similar size (4.57 and 4.46 Mbp, respectively) and contain a similar number of open reading frames (ORFs) (4,159 and 4,460, respectively). Both S. chlorophenolicum and S. japonicum (Nagata et al. 2010) have a primary chromosome containing most of the genes for core processes (including glycolysis, the TCA cycle, amino acid and nucleotide biosynthesis, fatty acid oxidation, DNA replication, transcription, and translation) and a secondary chromosome. Sphingobium chlorophenolicum has a single plasmid (pSphCh01), whereas S. japonicum has three (pUT1, pUT2, and pCHQ1).

The third circle in each chromosome map in figure 2 shows in green the positions of genes that encode proteins in S. chlorophenolicum that have close homologs in S. japonicum that exhibit >80% identity over >90% of the length of the S. chlorophenolicum sequence. A total of 2,324 S. chlorophenolicum genes (1,931 on chromosome 1, 446 on chromosome 2, and 108 on pSphCh01) have close homologs in S. japonicum (see supplementary table 2, Supplementary Material Online). In both chromosomes, regions with close homologs in S. japonicum show a typical GC content in a cluster on the —strand of chromosome 2 between positions 667462 and 675176 on the —strand of chromosome 2. A second copy of five of these genes are found in a cluster on the + strand of chromosome 2 between positions 1036969 and 1030812. Copies of the first two genes in the cluster are found together in a different location on chromosome 2. Nearly identical copies of the genes encoding the alpha and beta subunits of the E1 component of pyruvate dehydrogenase are found on chromosomes 1 and 2 and nearly identical copies of a gene annotated as an acyl-CoA synthetase/AMP-acid ligase II are found in chromosome 2 but surrounded by completely different genes. The utility of these duplicated genes and the processes leading to their location in distant parts of the genome are not clear. One possibility is that they were not actually duplicated in S. chlorophenolicum but rather acquired by integration into the genome of DNA fragments taken up from lysed cells of S. chlorophenolicum or closely related Sphingomonads. Notably, none of the PCP degradation genes is found among the set of highly similar genes.

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content of about 64%. Regions without close homologs (Sc islands) might have resulted from either loss of genes in *S. japonicum* or acquisition of genes by HGT in *S. chlorophenolicum*. Some of the Sc islands show a lower GC content, suggesting that these regions may have been acquired by HGT. Most of the genes in the Sc islands are hypothetical proteins, but there are several predicted glycosyltransferases (some predicted to be involved in cell wall biosynthesis), as well as some O-antigen ligases, some ABC transporters, cellobiose phosphorylase, a K⁺-transporting ATPase, and Type IV secretory pathway components. Notably, almost all the sequences associated with transposons and phage genes are found in Sc islands.

A more detailed analysis of the relationships between proteins found in both *S. chlorophenolicum* and *S. japonicum* is shown in figure 4, which shows plots of sequence identity versus coverage for the top hit in the *S. japonicum* genome for each *S. chlorophenolicum* protein. (Coverage is defined as the length of the *S. chlorophenolicum* query sequence that is aligned to a sequence in *S. japonicum* divided by the total length of the query sequence. The data were filtered to remove pairs for which the e value was >0.0001.) On this plot, homologs cluster in three regions: 1) close homologs that share high sequence identity over most of the query sequence; 2) more distant homologs that share moderate sequence identity over most of the query sequence; and 3) homologs that share sequence identity only over part of the query sequence. Notably, chromosome 1 is highly enriched in close homologs, and chromosome 2 is modestly enriched in distant homologs. Together with the observation that most of the genes for core metabolic processes are present on chromosome 1, this observation suggests that chromosome 2 may preferentially collect horizontally transferred genes.

Figure 4a shows that most of the proteins encoded on chromosome 1 share >80% identity with proteins in *S. japonicum*; indeed, 50% share >90% identity. We posit that close homologs with >80% identity were present in the most recent common ancestor of these two species; most are likely to be orthologs. The more distant homologs in region 2 are unlikely to be orthologs derived from the most recent common ancestor, since they are much more divergent than the large number of close homologs in region 1. The genes encoding these proteins may have been acquired by HGT independently from different sources in the two species; they may serve the same or different functions. Alternatively, the *S. chlorophenolicum* protein may indeed have derived from the most recent common ancestor, but the ortholog in *S. japonicum* may have been lost so that the best hit in the *S. japonicum* genome is actually a paralog. Finally, there are 112 proteins for which significant sequence identity is seen over only part of the query sequence (<75%); these include proteins in which conserved domains have been utilized in different structural contexts.

![Figure 4](https://example.com/figure4.png)

**Fig. 4.**—Relationships between proteins encoded by (a) chromosome 1, (b) chromosome 2, and (c) the plasmid of *Sphingobium chlorophenolicum* and the best hits in the *Sphingobium japonicum* genome found by a Blast search using each *S. chlorophenolicum* protein as a query sequence. The number of pairs is plotted as a function of % identity and % coverage of the query sequence for each replicon. (Only pairs for which E < 0.0001 are shown.)
Although the dominant chromosomes of *S. chlorophenolicum* and *S. japonicum* share a common core of genes, there has been considerable rearrangement of genes since the common ancestor of these two bacteria. Figure 5 shows a scatter plot of gene conservation between the chromosomes of *S. chlorophenolicum* and *S. japonicum* made using nucmer in the MUMmer package with the default parameters (Delcher et al. 1999, 2002; Kurtz et al. 2004). This plot shows a pattern known as an “x-alignment” (Eisen et al. 2000) that is often seen in closely related bacteria. Many genes in chromosome 1 of *S. chlorophenolicum* are found in comparable positions in chromosome 1 of *S. japonicum*, as indicated by the red dots along the diagonal. However, a number are found in an inverted orientation (see blue dots) in positions that lie close to a diagonal perpendicular to the red diagonal. This pattern is believed to result from multiple inversions centered around either the origin or terminus of replication. There has evidently been considerable remodeling of the genome via movement of blocks of genes within chromosome 1 since *S. chlorophenolicum* and *S. japonicum* diverged from a common ancestor. The x-alignment pattern is less distinct for chromosome 2,
suggesting greater plasticity in this replicon. This is consistent with the lower number of genes for core processes on chromosome 2 described above. Furthermore, only 12 of the 30 genes encoding core processes found on chromosome 2 of *S. chlorophenolicum* are found on chromosome 2 of *S. japonicum*, suggesting that movement of genes between the two chromosomes has continued to occur since the divergence of *S. chlorophenolicum* and *S. japonicum*. 

Figure 6 depicts the correspondence between the positions of very close homologs (≥90% identity over ≥80% of the query length) in the entire genomes of *S. chlorophenolicum* and *S. japonicum*. Light blue, orange, and dark blue lines connect the positions of genes in chromosome 1, chromosome 2, and the plasmid, respectively, of *S. chlorophenolicum* with the positions of homologs in the two chromosomes and three plasmids of *S. japonicum*. As noted above, chromosome 1 in both species is densely populated with shared genes. A number of genes found on chromosome 2 of *S. chlorophenolicum* have homologs on chromosome 1 of *S. japonicum*, although the converse is not true. Notably, genes found in *S. chlorophenolicum* but not in *S. japonicum* are more heavily represented on chromosome 2 than on Chromosome 1. Since chromosome 2 appears to carry many of the genes for degradation of organic compounds, this difference may be due to the availability of different carbon sources in the environmental niches occupied by the two bacteria. Notably, the *S. chlorophenolicum* plasmid, pSPHCH01, is comprised of a large region that is syntenic with a region of *S. japonicum* chromosome 1 (with the exception of a few small indels) and a smaller region that is syntenic with a region around the origin of *S. japonicum* pCHQ1. This region encodes several proteins, including two chromosome partitioning proteins (ParA and ParB homologs) and the plasmid replication initiation protein (RepA). Thus, the *S. chlorophenolicum* pSPhCh01 and *S. japonicum* pCHQ1 share an origin of replication and the associated genes, but the genes carried on the two plasmids are not closely related. The two smallest plasmids in *S. japonicum* (pUT1 and pUT2) carry genes with no homologs in *S. chlorophenolicum*.

One hundred and eight of 125 genes on pSPHCH01 have >90% identity to genes on plasmid pSLGP in *Sphingobium SYK-6* (a more distantly related Sphingomonad that is the closest relative of *S. chlorophenolicum* and *S. japonicum* for which a whole genome sequence is available), suggesting that these genes may have been present on a plasmid in the ancestor of *S. chlorophenolicum* and *S. japonicum* and may have been incorporated into chromosome 1 of *S. japonicum* after divergence of the two species. Movement of blocks of genes among the plasmids and chromosomes in these organisms may have been facilitated by transposases, as Tn3-family transposase elements are present in both plasmids and in the *S. japonicum* chromosome near one end of the integrated region.

**Insights into the Origin of the PCP Degradation Genes**

The PCP degradation genes are found on chromosome 2. *pcpA, pcpC*, and *pcpE* are found in proximity to each other, although not within an operon, whereas *pcpB* and *pcpD* are found in an operon on the opposite side of the chromosome (see figs. 2 and 7). Expression of all the genes except *pcpC* (Orser et al. 1993) is induced in the presence of PCP, although it is not known whether the genes are regulated by PCP itself or by downstream products. The constitutive expression of *pcpC* suggests that a mechanism for its regulation by PCP has not yet arisen.

In theory, the genes encoding the PCP degradation enzymes might have originated by 1) recruitment of an ancestral enzyme without gene duplication, requiring sharing of the enzyme between the new and original functions; 2) recruitment of a pre-existing enzyme, followed by duplication and divergence of the original gene to provide one copy that is specialized, although possibly not optimized, for its function in PCP degradation; and 3) HGT of genes encoding either enzymes that have already specialized for their functions in PCP degradation or enzymes with inefficient promiscuous activities that became useful when *S. chlorophenolicum* encountered PCP. Comparison of the genomes of *S. chlorophenolicum* and *S. japonicum* allows us to determine which of these possibilities is most likely for each of the PCP degradation genes. If an enzyme originated by recruitment of an enzyme present in the most recent common ancestor of *S. chlorophenolicum* and *S. japonicum*, then a close homolog is likely to be found in *S. japonicum*. If gene duplication and divergence has resulted in an enzyme specialized for its role in PCP degradation, we should find a close homolog in *S. chlorophenolicum*. If a gene has been acquired by HGT, we would likely find no close homologs in either *S. chlorophenolicum* or *S. japonicum* and might find other signatures of HGT, as well.

PCP hydroxylase (PcpB) and PcpD (TCBQ reductase) are encoded by an operon on chromosome 2 that also includes *pcpR*, which encodes a LysR–transcripational regulator that has been reported to be essential for induction of expression of *pcpB, pcpA*, and *pcpE* (Cai and Xun 2002). PCP hydroxylase belongs to a family of flavin monooxygenases that hydroxylate phenols. Such enzymes are common in soil bacteria because phenolic compounds derived from lignin provide an important source of carbon. The lack of a close homolog in *S. japonicum* (see table 2) suggests that *pcpB* was not present in the most recent common ancestor of *S. chlorophenolicum* and *S. japonicum*. Close homologs of *pcpB* are also not found in *S. wittichii*, so it is unlikely that *pcpB* arose from an ancestral gene that was lost in *S. japonicum*. The low pairwise sequence identities between PCP hydroxylase and other flavin monooxygenases in *S. chlorophenolicum* (see table 2) suggest that *pcpB* did not arise by recent duplication and divergence of a previously existing
S. chlorophenolicum gene. These findings, along with the observation that *pcpB* and *pcpD* fall in a region of chromosome 2 with relatively low GC content (see fig. 2), suggest that *pcpB* was acquired by HGT from an unknown source. This hypothesis is supported by the observation that enzymes with 72–98% identity to PcpB are found in *Novosphingobium lentum* (Tiirola, Mannisto, et al. 2002), several polychlorophenol-degrading Sphingomonads from Finland (Tiirola, Wang, et al. 2002), *Sphingomonas* sp. UG30 (Cassidy et al. 1999), and in several uncultured bacteria from environmental samples collected from PCP-contaminated soils (Beaulieu et al. 2000). The high sequence
protein, mostly Fe transport; PcpB (gi21362858) FAD-binding protein; pcpR genes.

ochelin and colicins; LysR-type transcriptional regulator.

PCP hydroxylase might have been before its recruitment into degrading bacteria. The high level of sequence divergence makes it difficult to discern what the original substrate for degrading bacteria. The high level of sequence divergence makes it difficult to discern what the original substrate for PCP hydroxylase might have been before its recruitment into the PCP degradation pathway. (Careless annotation transfers have led to a proliferation of genes that are annotated as encoding PCP hydroxylase. The proteins encoded by these genes are homologous to PcpB, but have only about 35% identity to PcpB, a level at which substrate specificity cannot be confidently assigned. Furthermore, the ability of the corresponding microbes to degrade PCP has not been assessed. These assignments are probably wrong in the majority of cases.)

TCBQ reductase (PcpD) catalyzes the second step in the PCP degradation pathway (Dai et al. 2003). Reduction of a benzoquinone is an uncommon step in aromatic degradation pathways. Benzoquinones are formed as a result of hydroxylation of a phenol only when there is a leaving group, such as a chlorine or nitro group, at the position of hydroxylation. Most commonly, there is a hydrogen at the position of hydroxylation, and the product is a hydroquinone (see fig. 8).

TCBQ reductase is not closely related to any other enzyme in S. chlorophenolicum or S. japonicum (see table 2), suggesting that it, like pcpB, was acquired by HGT. Enzymes that carry out comparable reductions of benzoquinones produced by phenol monooxygenases are found in three other microbes, Pseudomonas sp. Strain WBC-3 (Zhang et al. 2009), Alcaligenes sp. strain Ny2215 (Xiao et al. 2007), and Cupriavidus necator JMP134 (formerly Ralstonia eutropha) (Belchik and Xun 2008), which degrade p-nitrophenol, o-nitrophenol, and trichlorophenol, respectively. Notably, these three benzoquinone reductases and TCBQ reductase belong to three different superfamilies (see fig. 9). The recruitment of enzymes from three different superfamilies to catalyze reduction of quinones in pathways for degradation of four different anthropogenic compounds suggests that, in each case, a bacterium has taken identities between these genes suggest that pcpB has been transferred among a number of bacteria in PCP-contaminated environments, a conclusion previously reached by Tiirila, Wang, et al. (2002). Strikingly, PcpB has <35% identity to most flavin monooxygenases from non-PCP degrading bacteria. The high level of sequence divergence makes it difficult to discern what the original substrate for PCP hydroxylase might have been before its recruitment into

Table 2

| Protein      | Closest Homolog in S. chlorophenolicum | Closest Homolog in S. japonicum | Closest Homolog in Sphingobium sp. SYK-6 |
|--------------|----------------------------------------|---------------------------------|---------------------------------------|
| PcpB (gi21362858) | FAD-binding monooxygenase (gi10740769) | FAD-binding monooxygenase (gi294146540) | p-hydroxy-benzoate hydroxylase (gi347528071) |
| PcpD (gi21362854) | Pthalate 4,5-dioxygenase subunit (gi10723764) | Vanillate monooxygenase oxidoreductase subunit (gi294023794) | Putative flavodoxin reductase (gi347430572) |
| PcpC (gi22417110) | Glutathione S-transferase domain-containing protein (gi10724309) | 2,5-dichlorohydro-quinone dehalogenase (LinD) (gi294023795) | Glutathione S-transferase (gi347527108) |
| PcpA (gi3760223) | None | 2,6-dichlorohydro-quinone 1,2-dioxygenase (LinE) (gi294146914) | None |
| PcpE (gi22417101) | Maleylacetate reductase (gi33432570) | Maleylacetate reductase (LinF) (gi294146911) | Alcohol dehydrogenase (gi347529189) |
advantage of a different pre-existing protein when faced with the need to degrade an aromatic compound that carries a leaving group at the site of the initial hydroxylation reaction.

While PCP hydroxylase catalyzes a reaction similar to that of its homologs, the progenitor of TCBQ reductase may have served a different function. Although there are several families of known quinone reductases, TCBQ reductase is not

![Fig. 8.—Products formed by hydroxylation at a position bearing a hydrogen (a) and a chlorine (b).](image)

![Fig. 9.—Benzoquinone reductases have been recruited from three different superfamilies in microorganisms degrading PCP (Sphingobium chlorophenolicum), 2,4,6-trichlorophenol (Cupriavidus necator JMP134), p-nitrophenol (Pseudomonas sp. Strain WBC-3), and o-nitrophenol (Alcaligenes sp. strain Ny2215). The benzoquinones shown (from top to bottom) result from hydroxylation of a phenol at a position bearing a chlorine atom (for PCP and 2,4,6-trichlorophenol) or a nitro group (for p- and o-nitrophenol).](image)
related to any of these. Rather, TCBQ reductase is most closely related to proteins that serve as the reductase component of two-component dioxygenases that initiate aerobic degradation of aromatic compounds that lack hydroxyl or amine substituents (Dai et al. 2003). These proteins contain a flavin that accepts electrons from NAD(P)H and transfers them one at a time to an iron-sulfur cluster, which then donates electrons to the active site of the oxygenase component of the enzyme. TCBQ reductase is also more distantly related to ferredoxins that restore activity of extradiol dioxygenases that have been inactivated during substrate turnover (Polissi and Harayama 1993; Hugo et al. 1998, 2000; Tropel et al. 2002). (This inactivation occurs after O2 reacts with Fe(II) at the active site of the enzyme-substrate complex to form superoxide and Fe(III). If the superoxide diffuses out of the active site before reacting with the substrate, the iron atom is left in the inactive Fe(III) state.) The reaction catalyzed by TCBQ reductase could certainly take advantage of this protein architecture; donation of hydride to a flavin, followed by transfer of single electrons to the iron-sulfur cluster and thence to bound TCBQ would form first a semiquinone and then TCHQ after transfer of a second electron. Thus, adaptation of the progenitor protein to this new role may have required little more than enhancement of a small molecule-binding site adjacent to the iron-sulfur cluster.

TCHQ dehalogenase (PcpC) catalyzes the third and fourth steps in the PCP degradation pathway. Each reductive dehalogenation step results in oxidation of two molecules of glutathione to glutathione disulfide. Similar reductive dehalogenation steps occur during lindane degradation in S. japonicum (Miyauchi et al. 1998) and during degradation of PCBs in Burkholderia xenovorans LB400 (Batels et al. 1999; Tocheva et al. 2006). All of these dehalogenases are members of the glutathione S-transferase superfamily. Most enzymes in this superfamily catalyze the nucleophilic attack of glutathione upon an electrophilic substrate to form a glutathione conjugate. However, a few, such as TCHO dehalogenase (Warner et al. 2005; Warner and Copley 2007b), maleylacetocetate isomerase (Polekhina et al. 2001), and maleylpyruvate isomerase (Marsh et al. 2008), catalyze more complicated reactions in which additional steps occur before and/or after the canonical attack of glutathione upon an electrophilic intermediate.

Although there are numerous members of the glutathione-S-transferase superfamily in S. chlorophenolicum, TCHQ dehalogenase has no more than 25% identity to any of them, so pcpC likely did not arise by divergence from another gene in S. chlorophenolicum. Close homologs of pcpC are not found in the genome of S. japonicum, so pcpC was probably not present in the ancestor of S. chlorophenolicum and S. japonicum. The GC content of pcpC is 61%, and it falls in a region of chromosome 2 with unusually low GC content. Thus, pcpC was most likely acquired by HGT. Strikingly, S. japonicum LinD, which catalyzes the reductive dehalogenation of 2,5-dichlorohydroquinone to chlorohydroquinone (Miyauchi et al. 1998), has only 25% identity to TCHQ dehalogenase. TCHQ dehalogenase and LinD are the only known reductive dehalogenases in aerobic bacteria that act on aromatic compounds. The high divergence between these enzymes suggests that they were recruited independently from different members of the GST superfamily, possibly due to selective pressure to degrade PCP and lindane, respectively. The function of the progenitors of these enzymes is not clear. Residues in the active sites of TCHQ dehalogenase and LinD resemble those in the active sites of some maleylpyruvate isomerases, and TCHQ dehalogenase has low activity with maleylacetone, an analogue of maleylpyruvate (Anandarajah et al. 2000). Pseudomonic activities can provide clues to the activity of an ancestral enzyme, so TCHQ dehalogenase may have originated from a maleylpyruvate isomerase. However, this cannot be stated with certainty.

2,6-Dichlorohydroquinone dioxygenase (PcpA) catalyzes the cleavage of 2,6-dichlorohydroquinone (DCHQ). DCHQ dioxygenase is 93% identical to LinEb in S. japonicum, which also cleaves DCHQ. (Despite its name, LinEb is not involved in degradation of lindane [Endo et al. 2005].) Thus, this enzyme was likely present in the most recent common ancestor of S. chlorophenolicum and S. japonicum. Whether cleavage of DCHQ is the natural function of this enzyme is not clear, as studies of its substrate range and catalytic efficiency have not been reported beyond an early report that shows that hydroquinone and chlorohydroquinone also serve as substrates (Ohtsubo et al. 1999). Sphingobium japonicum does not degrade PCP, so the role of LinEb in this bacterium is not known. (Sphingobium japonicum contains a homolog of LinEb called LinE that cleaves chlorohydroquinone in the lindane degradation pathway. LinE and LinEb share 53% identity and DCHQ dioxygenase and LinE share 52% identity.)

The final two steps in degradation of PCP are catalyzed by maleylacetate reductase (PcpE), which carries out the successive reductions of 2-chloromaleylacetate to maleylacetate and of maleylacetate to β-keto acid. PcpE has 91% identity to maleylacetate reductase from S. japonicum (LinF), indicating that maleylacetate reductase was found in the progenitor of S. chlorophenolicum and S. japonicum. Thus, PcpE has been probably been recruited to carry out the reduction of 2-chloromaleylacetate in addition to the more typical reduction of maleylacetate. The S. chlorophenolicum genome also encodes two other maleylacetate reductase homologs, Sc_00030750 and Sc_00027830, which have 53% and 55% pairwise sequence identities to PcpE, respectively. These enzymes have not been characterized, but one or both may play a role in reduction of 2-chloromaleylacetate based upon the observation that substantial 2-chloromaleylacetate reductase activity is present in a strain in which pcpE has been knocked out (Cai and Xun 2002).
Conclusions

Comparison of the genome sequences of S. chlorophenolicum and S. japonicum suggests that the first three genes in the PCP degradation pathway, pcpB, pcpD, and pcpC, were acquired by HGT. Since these genes were incorporated into two different places in the genome, at least two different HGT were involved. The documented spread of nearly identical pcpB genes among bacteria in PCP-contaminated soils suggests that transfer of pcpB, and possibly pcpD as well, may have occurred within the last century since the introduction of PCP into the environments.

The last two genes in the PCP degradation pathway, pcpA and pcpE, were inherited from the progenitor of S. chlorophenolicum and S. japonicum. Pathways for degradation of aromatic compounds typically begin with specialized enzymes that convert the initial compounds to catechol or hydroquinone intermediates that are then cleaved by intradiol or extradiol dioxygenases. (Intradiol dioxygenases cleave catechols between the two hydroxyl groups. Extradiol dioxygenases cleave catechols or hydroquinones adjacent to one of the hydroxyl groups.) The ring cleavage products are then processed to C3 and C4 carboxylates that feed into central metabolism. Thus, numerous “upper” pathways funnel into a small number of “lower” pathways. The enzymes encoded by pcpA and pcpE likely existed as part of a standard lower pathway for degradation of naturally occurring compounds. The close proximity of pcpA and pcpE in the genome is consistent with this interpretation.

Duplication and divergence is believed to be the most common mechanism for evolution of enzymes with novel capabilities. This process allows exploration of sequence space while the original activity of the encoded enzyme is provided by a copy that deviates little, if at all, from the original sequence. Ultimately, extraneous copies of the gene are eliminated, leaving one copy encoding the original enzyme and one copy encoding the novel enzyme. Analysis of the S. chlorophenolicum genome sequence does not reveal evidence for recent duplication and divergence of any of the PCP degradation genes within S. chlorophenolicum itself. Duplication and divergence may have occurred in the unknown soil bacterium in which the genes originated, and only the genes encoding enzymes useful for degradation of PCP may have been transferred to S. chlorophenolicum and other PCP-degrading Sphingomonads.

Gene amplification is commonly observed when inefficient enzymatic activity limits the ability of a bacterium to degrade a carbon source. For example, the benzoate degradation genes in a strain of Acinetobacter ADP-1 that lacks the transcriptional regulators that normally induce these genes are amplified up to 20-fold when benzoate was supplied as a sole carbon source (Reams and Neidle 2003). Given the inefficiency of the initial enzymes in the PCP degradation pathway, it is intriguing that there is no evidence for amplification of genomic segments containing the genes encoding these enzymes. In this case, cells carrying duplicated or amplified genes may be at a disadvantage. Amplification of a segment of the genome carrying pcpB and pcpD would indeed increase flux through the initial two steps but would probably not improve fitness. The waste of NADPH and the concomitant production of H₂O₂ by the uncoupled PCP hydroxylase reaction would increase in proportion to the enzyme concentration. Additionally, the increased concentration of TCHQ formed would inhibit TCHQ dehalogenase, leading to a buildup of TCHQ, which would be expected to cause redox cycling and depletion of cellular reductants. On the other hand, duplication of a segment of the genome carrying pcpC would provide no selective advantage because the initial step, conversion of PCP to TCBQ, limits the flux through the pathway (McCarthy et al. 1997). Thus, duplication of this region might simply cause a metabolic burden without providing any benefit. If pcpB, pcpD, and pcpC were colocaled, then duplication of all the genes together might be beneficial. However, given the incorporation of pcpBD and pcpC into two distant regions of chromosome 2, gene duplication does not provide a pathway for further evolution at this point.

Supplementary Material

Supplementary tables 1 and 2 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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