Burkholderia Xenovorans LB400 Harbors a Multi-Replicon, 9.73-Mbp Genome Shaped for Versatility

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Burkholderia xenovorans LB400 harbors a multi-replicon, 9.73-Mbp genome shaped for versatility


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Contributed by James M. Tiedje, August 10, 2006

Burkholderia xenovorans LB400 (LB400), a well studied, effective polychlorinated biphenyl-degrader, has one of the two largest known bacterial genomes and is the first nonpathogenic Burkholderia isolate sequenced. From an evolutionary perspective, we find significant differences in functional specialization between the three replicons of LB400, as well as a more relaxed selective pressure for genes located on the two smaller vs. the largest replicon. High genomic plasticity, diversity, and specialization within the Burkholderia genus are exemplified by the conservation of only 44% of the genes between LB400 and Burkholderia cepacia complex strain 383. Even among four B. xenovorans strains, genome size varies from 7.4 to 9.73 Mbp. The latter is largely explained by our findings that >20% of the LB400 sequence was recently acquired by means of lateral gene transfer. Although a range of genetic factors associated with niche adaptation and intercellular interactions are present, these genetic factors are likely related to niche breadth rather than determinants of pathogenicity. The presence of at least eleven “central aromatic” and twenty “peripheral aromatic” pathways in LB400, among the highest in any sequenced bacterial genome, supports this hypothesis. Finally, in addition to the experimentally observed redundancy in benzoate degradation and formaldehyde oxidation pathways, the fact that 17.6% of proteins have a better LB400 paralog than an ortholog in a related species, the presence of at least ten closely related species, the Burkholderia cepacia complex (Bcc), are secondary colonizers of the lungs of cystic fibrosis patients, often causing rapid decline in lung function (12). Similar to most other Burkholderia species, Bcc species are commonly found in the environment, often associated with the plant rhizosphere (18).

The diversity of the Burkholderia genus is exemplified by the high diversity of ecological niches occupied by the different species, ranging from soil to aqueous environments, associated with plants, fungi, amoeba, animals, and humans, from saprophytes to endosymbionts, from biocontrol agents to pathogens (10–15). Historically, most of the attention toward this genus was directed toward species with malign properties, including plant pathogens and species causing animal and human disease, such as Burkholderia mallei and Burkholderia pseudomallei (16, 17). A subgroup of at least ten closely related species, the Burkholderia cepacia complex (Bcc), are secondary colonizers of the lungs of cystic fibrosis patients, often causing rapid decline in lung function (12). Similar to most other Burkholderia species, Bcc species are commonly found in the environment, often associated with the plant rhizosphere (18).

Common to most Burkholderia species are a large, multireplicon genome and the presence of multiple insertion sequences that confer high genome plasticity, which could help explain the versatility of the genus (19). Twenty-two Burkholderia strains are being, or have been recently, sequenced (Joint Genome Institute [JGI]/Integrated Microbial Genomes [IMG]; http://img.jgi.doe.gov). The complete genomes of the closely related B. pseudomallei K96243 (20) and B. mallei ATCC23344 (21) became available in 2004, whereas Burkholderia cenocepacia J2315 (http://www.sanger.ac.uk), B. cenocepacia H12424 (JGI), and the environmental isolate...
\textit{Ralstonia solanacearum} Group at the Sanger Institute (http://www.sanger.ac.uk/Projects/B.cenocepacia/); \textit{B. pseudomallei} (blue, rings 10 and 11); \textit{Bcc} strain 383 (green, rings 12 and 13); and \textit{Ralstonia solanacearum} (magenta, rings 14 and 15). Ring 16 (black) represents \(G + C\) content, and ring 17 represents the \(G + C\) skew.

Results and Discussion

General Genome Description. General overview. The LB400 genome has a size of 9.73 Mbp, harboring \(\approx 9,000\) coding sequences distributed over three circular replicons that have been designated chromosome 1 (4.90 Mbp), chromosome 2 (3.36 Mbp), and megaplasmid (1.47 Mbp) (Table 1, which is published as supporting information on the PNAS web site). Origins of replication were determined based on \(G + C\) skew analysis (Fig. 1) and the presence of genes involved in DNA replication close to the predicted origin for each of the replicons. Homologs of \textit{dnaA} and \textit{dnAN} are present on chromosome 1, whereas \textit{parA} and \textit{parB}, which are responsible for plasmid partitioning and replication, are present on chromosome 2. Similarly, genes encoding ParA and ParB are also found on the small chromosome of other \textit{Burkholderia} species (20, 21), and these proteins are \(\approx 80\%\) identical to those of LB400. We defined the 3.36-Mbp replicon as chromosome 2 and not a megaplasmid based on the presence of three ribosomal operons and several tRNAs (although none are unique), as well as the unique occurrence of core cellular functions involved in DNA replication [DNA primase (\textit{dnaG}), DNA polymerase I (\textit{polA})] and associated elongation factor (\textit{polB}) and amino acid metabolism. Moreover, there are several functions crucial to this organism’s adaptation to its niche uniquely located on this replicon (see Genetic Factors Indicating the Ecological Niche of \textit{B. xenovorans}). The 1.42-Mbp megaplasmid, on the other hand, contains neither RNA genes nor any essential functions and is mostly absent in other strains of the same species (see Comparison with other \textit{Burkholderia} species). The overall \(G + C\%\) of the megaplasmid is \(1\%\) lower than the two chromosomes, and, together with the \(G + C\%\) patterns and the high fraction of inserted sequences, this replicon appears to be a mosaic of foreign genomic material. Nevertheless, the megaplasmid’s \(G + C\) skew is well defined, although asymmetrical (Fig. 1C). ParAB share only \(\approx 20\%\) amino acid sequence identity with ParAB on chromosome 2 whereas they are \(> 80\%\) identical to ParAB on the 0.64-Mbp replicon of \textit{Cupriavidus necator} (\textit{Ralstonia eutropha}) JMP134.

Gross functional content. The genome’s functional content and its distribution over the replicons were analyzed based on the replicons’ bias toward particular clusters of orthologous groups of proteins (COG) (Fig. 2A). The large chromosome carries the core cellular function (e.g., translational machinery, DNA replication, cell division, and nucleotide metabolism) and can be considered the “core” chromosome. The small chromosome and the megaplasmid have a functional bias toward energy metabolism, secondary metabolism, inorganic ion transport and metabolism, and amino acid metabolism and transport (COG E). The bias toward COG E can be explained by the large number of amino acid transporters on these replicons: 58% of the megaplasmid-located and 42% of small chromosome-located COG E proteins compared with only 32% of the large chromosome-located COG E proteins. On the megaplasmid, the high number of proteins in the subclass of branched amino acid transporters is remarkable: 49 (28% of its COG E proteins) vs. 25 (8%) on the small and 40 (12%) on the large chromosome. The role of this subclass of transporters is unclear but is most likely not limited to amino acid transport. The bias toward inorganic ion...
transport/metabolism is caused by an uneven presence of oxygenase enzymes on these replicons, which are classified in this COG because they contain metal ions. Compared with the other replicons, the protein content of chromosome 2 is biased toward transcription, carbohydrate transport and metabolism, and signal transduction mechanisms, whereas the megaplasmid has a bias toward lipid metabolism. The latter observation is explained by the presence of a large number of CoA ligases as well as other enzymes similar to fatty acid metabolic pathways. This distribution of functions leads us to define the small chromosome as the “lifestyle-similar to fatty acid metabolic pathways. This distribution of functional distribution over the three replicons based on the COG classifications. Presented is the percent of genes on each replicon belonging to each COG.

**Paralogs and functional redundancy.** Gene and functional redundancy seems to be an important theme in large bacterial genomes. Based on our analysis, 1,581 coding sequences (17.6% of all genes) were found to have a better match within the LB400 genome than in the database of 260 completed genomes. To date, this percentage represents the highest extent of gene and potential functional redundancy among the genomes of free-living bacteria, where the average is 7.6% (±4.0%) of their genes. The list of paralogs in the LB400 genome is enriched in genes related to transport (230 paralogs), signal transduction (164 paralogs), mobile elements (112 paralogs), membrane proteins (66 paralogs), and secondary metabolism, including 120 dehydrogenases, 32 di-oxygenases, and 13 mono-oxygenases. Although the aim of our analysis was to uncover potential functional redundancy, it is important to recognize that, for most paralogs, divergent evolution can lead to changes in substrate range, kinetic properties, or even function, thus further extending LB400's great metabolic versatility. Interestingly, these 1,581 paralogs are numerically more or less equally distributed among the three replicons (~500 genes per replicon), and, consequently, the relative contribution of the smaller replicons to gene redundancy is substantially greater than chromosome 1.

The high number of paralogs among transport proteins correlates with the large fraction of LB400 genes (~1,400, encoding ~610 transport systems; Table 2, which is published as supporting information on the PNAS web site) dedicated to this function. There are >180 efflux systems, including 89 drug efflux pumps, 18 protein secretion systems, 21 heavy metal efflux pumps, and 18 amino acid/amino acid lactone efflux pumps (LivE, RhtB, and LysE). The latter suggests a role beyond amino acid transport. The other 430 systems are involved in uptake of organic compounds as well as all necessary inorganic cations and several anions. The most extensive redundancy is observed for the abundant branched and polar amino acid transporter families, which can indicate functional diversification to transport-related compounds.

A second class of proteins for which a high number of paralogs is observed is signal transduction. In light of this trait and the fact that it has been previously shown that the signal transduction network size increases disproportionately with genome size (28), we analyzed in more detail this class of proteins (Table 3, which is published as supporting information on the PNAS web site). Among currently published bacterial genomes, LB400 dedicates the second-largest share (9%) of its genome to signal transduction (second to Streptomyces coelicolor) and utilizes almost the entire spectrum of input (sensory) and output (regulatory) domains found in prokaryotic signal transduction (29). Whereas regulation of gene expression (vs. enzymatic domains) on average constitutes ~75% of the output activity in prokaryotic signal transduction (29), it constitutes almost 90% in LB400. One-component signal transduction (1CST) proteins (29) that combine a sensory and a regulatory module are the dominant signal transduction mechanism in LB400 (629 proteins). The LysR and AraC families (transcription activation in response to a small ligand) constitute the two largest classes (176 and 74 proteins, respectively) among 1CST systems. Most are encoded within metabolic gene clusters. In contrast, we found fewer GntR-type (71 proteins) and TetR-type transcriptional repressors (57 proteins), indicating that most metabolic pathways in LB400 require induction. The majority of 1CST systems in LB400 are predicted to be soluble, cytoplasmic sensors, indicating that it detects most environmental signals intracellularly after transport into the cell. Two-component signal transduction (2CST) systems are less abundant. Apart from the chemotaxis system (see Metabolic traits and Chemotaxis), the genome contains 76 sensor histidine kinases and 73 response regulators. Thirty histidine kinases are predicted to be soluble cytoplasmic sensors whereas many membrane-bound sensor histidine kinases contain intracellular sensory domains, such as PAS (30). The extraordinary sensing capacity of LB400 is further emphasized by the fact that Rhodococcus sp. RHA1, an organism with similar niche and genome size, has 25% fewer transporter systems and less than half the number of 2CST systems (31).

What is the added value of gene and functional redundancy for organisms with such large genomes? In LB400, functional redundancy of formaldehyde oxidation (23) and benzoate catabolism (32), for each process which LB400 has three potential pathways, has been studied, and, in both cases, the functionality of each of the three pathways present in the genome was confirmed. Redundancy is not entirely unexpected because these organisms presumably have to contend with varying conditions in nutrient source types and concentrations. The latter is exemplified by the detailed study of benzoate catabolism, which suggests that each of these pathways is conditionally regulated depending on benzoate and oxygen levels presented to the cell (33). Further studies of LB400 metabolic and other functional redundancies, such as the presence of multiple Type IV secretion systems (see Burkholderia virulence-associated genes), will be important to gain a better understanding of their evolutionary origin, their role, and whether or not there is a selective benefit of functional redundancy in large-genome bacteria.

**Genome Evolution. Comparison with other Burkholderia species.** We compared the gene complement of LB400 with that of *B. pseudomallei* and *Bcc* strain 383, two strains about equally distant to LB400 (77.5% and 76.8% average nucleotide identity, respectively). The comparisons gave very similar results, and we summarize the findings for the comparison with strain 383 only for simplicity. Less than half of the genes in the LB400 genome (3,961) are conserved between LB400 and strain 383, revealing the great genetic diversity within the *Burkholderia* genus. These 3,961 conserved genes are unevenly distributed in the three replicons of LB400: 2,602 conserved genes on chromosome 1 (57.7% of the
replicon’s genes), 1,150 on chromosome 2 (37.8%), and 209 on the megaplasmid (15%). Therefore, the smaller replicons seem to contribute more to the phenotypic uniqueness of the species and strains, whereas the largest replicon possesses most of the core functions of the Burkholderia genus. This uneven distribution of conserved genes over the replicons is consistent with comparisons of B. pseudomallei and B. mallei genomes (20, 21) and reveals an important, universal property of the Burkholderia genus. Furthermore, extending our analysis to other fully sequenced Burkholderia strains confirms the large chromosome as the core and most conserved chromosome (Fig. 1). Similar to the comparison with Bcc strain 383 alone, the small chromosome displays a significantly larger variation between the various species as compared with the large chromosome (Fig. 1B), whereas the megaplasmid is virtually unique to LB400 (Fig. 1C). The large number of shared genes on the small chromosomes of the different Bcc strains (data not shown) and the high sequence similarity of the small chromosome’s parAB genes between all Burkholderia species (see General Genome Description) suggest a shared ancestry.

Interestingly, our best reciprocal BLAST comparison of LB400 with Bcc strain 383 showed that the orthologous genes between the LB400 and strain 383 genomes are generally found in the same replicon for the genes in the largest replicon whereas this is not always the case for the smaller replicons. For instance, 40, 125, and 44 of the orthologs on the LB400 megaplasmid are located on the largest, medium, and smallest replicon of strain 383, respectively. The sequence identity of the orthologs is also noticeably different between the replicons: the orthologs share 74.9% average amino acid identity for chromosome 1, 66.3% for chromosome 2, and 55.9% for the megaplasmid. Together, these results reveal that the smaller replicons are more “plastic” and show higher sequence evolution rates. Although we cannot exclude that some “false” orthologs may confound some of our findings, when we repeated the analysis with genomes of more closely related Burkholderia strains, or more stringent cut-offs (which are less error-prone), we found a very similar pattern.

To further test for differences in selective pressure acting on the replicons, we used the ratio of nonsynonymous vs. synonymous substitutions (Ka/Ks) between orthologous pairs in LB400 and Bcc strain 383, with smaller Ka/Ks ratios representing higher negative, purifying selection. The Ka/Ks ratio differs substantially between replicons: the megaplasmid’s genes have an average Ka/Ks ratio of 0.32 (±0.14), vs. 0.28 (±0.13) for chromosome 2 and 0.22 (±0.11) for chromosome 1 (Fig. 3). Also, a significantly larger share of the Ka/Ks values was significantly <1 (Z tests, P < 0.05) on the large chromosome as compared with the two smaller replicons (Fig. 5), which is published as supporting information on the PNAS web site). These results further support the notion that the smaller replicons may be under more relaxed selective pressure, which probably accommodates and drives their plasticity. It is interesting to note that, in most of the previous comparisons, chromosome 2 behaves more like the megaplasmid than like chromosome 1. Consistent with these interpretations, whole genome alignments (34) of LB400 with B. pseudomallei and Bcc strain 383 indicate that the degree of recombination/rearrangement is significantly higher for the small chromosome (Fig. 6, which is published as supporting information on the PNAS web site). Interestingly however, and similar to B. pseudomallei (20), chromosome 2 carries a smaller fraction of recently acquired functions (6 genomic islands, containing ~7% of the small chromosome) compared with chromosome 1 (15 genomic islands containing ~17% of the large chromosome) (Table 1 and Table 4, which is published as supporting information on the PNAS web site). Considering that 7 of 15 genomic islands on the large chromosome (8/21 in total) are adjacent to a tRNA gene and contain integrases or transposases, this preferential insertion into the large chromosome is probably due to the much larger number of tRNA genes on the large chromosome (57 of 64). The lower number of insertions into the small chromosome could also indicate that the specific functions encoded by genes located on the small chromosome, which presumably help differentiate the various Burkholderia species, were acquired long ago and have been maintained by selection.

Comparison with other B. xenovorans strains. By using the LB400 genomic microarray, comparative genome hybridizations (CGHs) were performed for B. xenovorans LMG 21720 (plant rhizosphere origin) and LMG 16224 (blood sample origin). Sixty-nine percent of LB400’s genes are conserved in both strains. Pairwise comparisons correlate well with previously determined DNA:DNA hybridization ratios (1): 78% are conserved in LMG 21720 and 73% in LMG 16224 as compared with DNA:DNA hybridization ratios of 83% and 73%, respectively. Based on pulsed-field gel electrophoresis (PFGE), LMG 21720 and LMG 16224 were estimated to be 7.83 and 8.87 Mbp, respectively. A fourth B. xenovorans strain, LMG 22943 (plant rhizosphere origin), was sized at 7.42 Mbp. These three strains harbor two chromosomes, whereas only LMG 22943 and LMG 16224 contain a plasmid, roughly half the size of the one in LB400 (Fig. 7, which is published as supporting information on the PNAS web site). These data support the CGH results, indicating that most of the megaplasmid is indeed missing from the other members of the species. Based on these data ~1.75 Mbp of additional genomic material is found in LMG 16224 compared with only 0.30 Mbp in LMG 21720.

The most evident CGH result was that 70% of the genes located on the megaplasmid were uniquely present in LB400. Consequently, the functional distribution of the genes unique to LB400 correlates well with the functional bias of the megaplasmid (Fig. 8, which is published as supporting information on the PNAS web site). Apart from five islands on the megaplasmid (~12%) that are conserved in the other B. xenovorans strains (Fig. 1C), conserved genes are scattered throughout the megaplasmid and are mostly involved in transport and regulation. The conserved islands include genes encoding carbon monoxide dehydrogenase (cox), diterpenoid metabolism (dit), anthranilate dioxygenase (andA), and formaldehyde oxidation. We also verified the presence of dit genes experimentally by growth of both LMG strains on abietic and dehydroabietic acid. Important megaplasmid-derived functions, unique to LB400, include the periplasmic nitrate reductase (nap), the biphe-
We have also found in LB400, but not in the other *B. xenovorans* strains used for CGH (confirmed by PCR, data not shown), the presence of IS*pu*12, a 3,372-bp insertion sequence first described on the toluene-xylene TOL catabolic plasmid pWWO of *Pseudomonas putida* mt-2 (35). In LB400, IS*pu*12 was found in its entirety in seven different locations (three copies on the large chromosome, two on the small chromosome, and two on the megaplasmid). Furthermore, because its sequence was found in a portion of the sequencing reads at four additional locations (one on chromosome 1, two on chromosome 2, and one on the megaplasmid), we believe this IS element is actively copying itself within the genome, which resulted in distinct genotypic (and potentially phenotypic) subpopulations within the original culture. Similar observations of strain variation within clonal populations grown for sequencing have been previously reported for large-scale genomic rearrangements in *Yersinia pestis* (36) and for phase variable regions in the *Bacteroides fragilis* genome (37). It is interesting to note that IS*pu*12 occurs in several other bacteria, generally associated with plasmids and xenobiotic degradation genes, and that it has been suggested to be involved in gene silencing and activation (38).

**Genetic Factors Indicating the Ecological Niche of *B. xenovorans***

**Metabolic traits.** Based on the origin of its isolates, *B. xenovorans* presumably has a versatile environmental niche (soil and plant rhizosphere), which could imply an ability to degrade aromatic compounds that originate from root exudates and root turnover. CGH with other *B. xenovorans* strains showed that LB400 is enriched in aromatic catabolic pathways contained in genomic islands, providing biphenyl, 3-chlorocatechol, 2-aminophenol, and other catabolic capacities (Table 4) to this strain. Based on in silico analysis, we found eleven “central aromatic” catabolic pathways in the LB400 genome (Fig. 4 and Table 5, which is published as supporting information on the PNAS web site), indicating an unusually high metabolic versatility. Other sequenced aromatic compound-degrading bacteria, such as the aerobic *P. putida* KT2440 (39), *Rhodococcus* RHA1 (31), and anaerobic strain EbN1 (40), have six, eight, and five central aromatic pathways, respectively. Nine such pathways were found in the combined genomes of all fully or partially sequenced *Pseudomonas* strains (41). In addition, the LB400 genome contains twenty “peripheral aromatic” catabolic pathways (Fig. 4). Seven and 13 such pathways were described in *P. putida* KT2440 (39) and strain EbN1, respectively (40), whereas *Rhodococcus* RHA1 (31) contains at least 25. LB400 has >170 genes encoding these aromatic catabolic pathways, spread over the three replicons (Fig. 9, which is published as supporting information on the PNAS web site, and Table 4). The genes of the central aromatic pathways are generally organized in operon-like structures, and genes encoding transcriptional regulators are adjacent to 10 of 11 central aromatic pathway operons, which suggests integration into the signal transduction network. Several operons contain transporter genes as well (Table 4). There are at least 23 aromatic acid transporters, and some of the ∼100 (undefined) major facilitator superfamily and amino acid family transporters could be involved as well, e.g., branched amino acid transporters, two systems of which are within aromatic degradation operons (Table 4). The high abundance of transporters in LB400 suggests...
that its diverse metabolism is complemented by the same diversity in transport capacity.

**Primary reactions.** To be able to metabolize aromatic, non-hydroxylated carbon sources, initial ring activation is generally required. This can occur by di- or monohydroxylation through the activity of a primary oxygenase. In other aerobic aromatic degradation pathways, CoA ligation is the initial step mediated by an AMP-dependent CoA ligase. In the latter case, ring hydroxylation is still required, but the initial substrate selectivity is determined by the CoA ligase. The LB400 genome contains 12 Rieske-type primary dioxygenases (Fig. 10.4, which is published as supporting information on the PNAS web site), 6 cytochrome P450 monooxygenases, and 45 AMP-dependent synthetases/ligases (Fig. 10B). The latter group are mainly studied for their role in fatty acid metabolism, but the same mechanism is involved in degradation of benzoate, phenylacetate, and diterpenoids (see Aerobic hybrid pathways). The functions of most of these divergent CoA ligases and associated β-oxidation-like pathways are unclear and require experimental verification.

**The β-keto adipate pathway and peripheral aromatic pathways.** Degradation pathways for many aromatic compounds eventually funnel into the β-keto adipate pathway via the catechol branch or the protocatechuic acid branch, which converge at β-keto adipate enol-lactone. The cat genes, which encode the three enzymes of the catechol pathway and participate in the transformation of catechol to β-keto adipate enol-lactone, are arranged as a catRBAC cluster located on chromosome 1. The genes of the complete protocatechuic acid pathway, arranged as pcaIJBDC and pcaHG clusters and a single pcaF gene, are located on chromosome 2. The cat and pca gene arrangement is similar to other β-proteobacteria such as *B. pseudomallei* and *Ralstonia metridii clusrans* (39). Fifteen peripheral aromatic pathways that lead to the β-keto adipate pathway were identified. Nine substrates are degraded via catechol: (i) (chboroxyethyl glycol (bpg, megaplasmid); (ii) benzoate, (ben, chromosome 2); (iii) benzoic acid (nit, chromosome 1), which is metabolized to benzoic acid via (iv) benzamide (ami, megaplasmid); (v) mandelate (mdl, chromosome 2), which is transformed to benzaldehyde via (vi) benzaldehyde (gh, megaplasmid); (vii) tryptomphan (kyn, chromosome 1 and 2); which is converted to benzoate via (viii) anthranilate, (and, two paralogous copies on chromosome 2, megaplasmid); (ix) salicylate, which is transformed to catechol by salicylate hydroxylase (nahW, megaplasmid). Five substrates are degraded via protocatechuate: (i) vanillin (van, chromosome 1); (ii) 4-hydroxybenzoate (poh, chromosome 1) (41); (iii) phtalate (oph, chromosome 1 and 2); (iv) terephtalate (poh, two copies on chromosome 2); and (v) 4-carboxy diphenyl ether (poh, copies on chromosome 1 and 2). Finally, 3-chlorocatechol is degraded to β-keto adipate by the so-called modified or tho pathway (clc, chromosome 1), organized similar to *Pseudomonas* sp. B13 (42).

**Aerobic hybrid pathways.** Recently, a new aerobic hybrid pathway for benzoate degradation has been described in various bacteria including LB400, which has two homologous and functional copies of this pathway (chromosome 1, megaplasmid) (32, 33, 43, 44). Phenylacetate is transformed by a similar hybrid pathway via phenylacetate-CoA (45) because of activity of the paa pathway (chromosome 1, megaplasmid). Also, a particular cluster of genes, which includes the characterized *Pseudomonas abienanphila* BKME-9 dit cluster, encodes proteins required for abietane diterpenoid catabolism. Abietane diterpenoids are tricyclic compounds commonly found in the oleoresin of coniferous trees. The dit cluster includes genes coding for two cytochromes P450 (ditQ, ditU), a Rieske-type dioxygenase (ditA) (10.4A), an extradiol ring-cleavage dioxygenase (ditC), a CoA ligase (ditL) (Fig. 10B), and associated β-oxidation homologs (D.S., J. Park, J.M.T., and W. Mohn, unpublished data). Based on the abundance of CoA ligase/β-oxidation homologs, more of such auxiliary, aerobic hybrid pathways are probably present in LB400 but are yet to be recognized.

**Other metabolic traits.** Similar to some other *Burkholderia* species, *B. xenovorans* contains the *nif* nitrogen fixation genes (chromosome 2). Regarding nitrogen metabolism, the presence of *nap* genes (megaplasmid) for aromatic nitrite reduction is interesting as well. Furthermore, *C* oxidation and assimilation may play an important role in this species because we observed in the LB400 genome (i) three formaldehyde oxidation pathways, (ii) a methanol dehydrogenase-like enzyme (XoxF) and associated POQ cofactor synthesis genes, (iii) three carbon monoxide dehydrogenases (ace togenesis), and (iv) the entire Calvin–Benson–Bassham cycle for CO₂-fixation in autotrophic organisms. Importantly, several of these pathways also seem to be present in the *B. xenovorans* strains tested by CGH, but not in other sequenced *Burkholderia* species (23). There is no evidence of a recent acquisition event for most *C* genes, indicating that these genes were presumably acquired early on in the evolution of the *B. xenovorans* lineage. No growth of LB400 on *C* compounds has yet been reported; however, growth was observed on the methylated-glycine derivatives betaine and choline (23). Because many plant aromatic compounds are methylated or methoxylated, it is possible that these *C* pathways are involved in plant aromatics metabolism, which is a possible explanation for their coexpression in biphenyl-grown LB400 cells at transition to stationary phase (33, 44).

**Chemotaxis.** The signal transduction network of LB400 contains several chemotaxis genes, some of which might be involved in taxis toward aromatic compounds. Only one chemoreceptor (BxeA0121, methyl-accepting chemotaxis protein, MCP) is located within the chemotaxis operon, whereas 31 other chemoreceptors are scattered throughout the genome. Twenty-one chemoreceptors belong to Class I MCPs (46), i.e., are transmembrane proteins with a periplasmic sensory domain and are therefore predicted to detect extracellular signals. One MCP (BxeA2405) is an ortholog of the *Escherichia coli* Acr aerotaxis transducer, and all amino acid residues implicated in the FAD binding and the aerotactic response (47) are conserved in this protein, suggesting that it governs energy taxis (48) in LB400.

**Presence/absence of pathogenicity traits.** *Burkholderia* virulence-associated genes. Although the genetic factors required for animal and human pathogenicity by *Burkholderia* are largely unknown, multiple traits have been experimentally implicated in the virulence of *B. pseudomallei* (29) and the *Bcc* (12) (Table 6, which is published as supporting information on the PNAS web site). Resistance to cationic antimicrobial agents such as polymixin B, presence of a flagellum (49), and iron acquisition by siderophores (50) (pyoverdin, pyochelin biosynthesis, chromosome 1) are virulence traits but also important in their soil habitat. Also, several genes are present that may mediate resistance to reactive oxygen species and enable LB400 to survive within intracellular environments. Although LB400 encodes three phospholipase homologs and an LasA-type protease, its repertoire of phospholipases, proteases, and collagenses, which have been implicated in *Burkholderia* virulence (20, 51), is limited. LB400 also lacks homologs of the *B. cenocepacia* cable pilus, a defining pathogenic feature of one of the most problematic strains in cystic fibrosis infection (12). However, in common with other *Burkholderia* genomes (20), LB400 possesses multiple hemagglutinin-like adhesin genes (54); however, neither of the LB400 systems shares significant sequence similarity with the latter plant pathogenic gene cluster.

The *Cenocepacia* island (12) is not present in LB400. However,
components of the amino acid metabolism gene cluster within the *B. cenocepacia* pathogenicity-related island show remarkable similarity with LB400 megaplasmid segments (Fig. 11, which is published as supporting information on the PNAS website). Six of the eight branched-chain amino acid metabolism genes, including the porin implicated in inflammation during infection (55) (BxeC0438), are highly conserved between the two species. The LB400 megaplasmid also has a cluster of type IV secretion genes immediately adjacent to the latter amino acid gene cluster, which shares considerable sequence similarity with a *B. cenocepacia* type IV secretion system island encoded ~46 kb downstream of the *Cenocepacia* island on the J2315 second chromosome (54). These corresponding LB400 megaplasmid regions do not possess the hallmarks of genomic islands, suggesting that the megaplasmid may represent an ancestral replicon from which the *B. cenocepacia* laterally acquired the *Cenocepacia* and type IV secretion islands. The fact that the quorum-sensing system on the *Cenocepacia* island is most closely related to that encoded on the LB400 megaplasmid supports this hypothesis. Quorum-sensing systems regulate the expression of a number of traits involved in *Burkholderia* pathogenicity (20, 56, 57). LB400 possesses two gene sets with homology to autoinducer synthesis and response regulator systems, one on the megaplasmid (BxeC0415, BxeC0416) and a classical luxIR-like system (BxeB0608, BxeB0610 on chromosome 2), which is phylogenetically most closely related to the MuplIR, PpuIR, and LaslIR systems of *Pseudomonas* species (57). Their regulatory roles are uncharacterized.

**Genes associated with in vivo survival.** Hunt et al. (58) reported the isolation of 102 transposon mutants of *B. cenocepacia* strain K56-2 that were unable to survive in a rat model of chronic lung infection. Of the 69 mutants that were mapped to orthologs within the finished *B. cenocepacia* J2315 genome (58), direct orthologs were found for 46 genes in LB400 (Table 7, which is published as supporting information on the PNAS website). Although these findings suggest that LB400 encodes a large proportion of the *Burkholderia* genes required for in vivo survival, the lack of LB400 homologs for several of the virulence factors for pathogenic *Burkholderia* species indicates that LB400 has little potential to cause infection.

**Concluding Remarks.** The LB400 genome and its comparison to closely related *B. xenovorans* strains, as well as to more distantly related *Burkholderia* species, underscore major themes in prokaryotic niche adaptation and genome evolution. Our analysis highlights that the niche and phenotypic diversity and versatility of the *Burkholderia* genus are reflected in the genomic composition of this group. This genus exemplifies the importance to niche versatility of lateral gene transfer and plasmid acquisition, as well as extensive gene and functional redundancy, because of both gene duplication and independent acquisition events. Based on (i) the distribution of gene function among the replicons, (ii) the dominance of intra- over interreplicon rearrangements when compared with other *Burkholderia*, and (iii) the differential gene conservation and evolutionary rates observed for the replicons, we (i) define the large chromosome as the core chromosome representing the major phenotypic characteristics of the *Burkholderia* genus, (ii) define the small chromosome as the lifestyle-determining replicon representing the adaptation of the species to its niche, and (iii) assert that the megaplasmid, the individuality replicon, provides the highly specialized, unique metabolic capabilities to LB400. The ecologic, phenotypic, and genomic features of *B. xenovorans* suggest that it and probably many environmental *Burkholderia* are versatiles, i.e., adapted to complex or diverse niches.

**Materials and Methods**

**Bacterial Strains.** The sequenced strain *B. xenovorans* LB400 was isolated from PCB-contaminated landfill soil in Moreau, New York (6). The sequenced clone is the original strain, which is deposited in the U.S. Department of Agriculture–Agricultural Research Service (USDA-ARS) Culture Collection (Peoria, IL) (NRRL B-18064). Three additional *B. xenovorans* strains were used in this study: LMG 16224 was retrieved from a blood culture specimen from a 31-year-old woman in Göteborg, Sweden; LMG 21720 is a coffee plant rhizosphere isolate from Costa Rica, Veracruz, State, Mexico (8); and LMG 22943 was isolated from a grass rhizosphere in Wageningen, The Netherlands (9).

**Genome Sequencing.** Whole-genome shotgun sequencing was performed on 3-kb, 8-kb, and 40-kb insert DNA libraries by the Production Genomics Facility of the Department of Energy (DOE)-Joint Genome Institute (DOE-JGI, Walnut Creek, CA). Paired-end sequencing generated >270,000 reads and resulted in approximately 16× depth of coverage. The reads were assembled to produce the primary high-quality draft assembly of 715 contigs, which were linked into larger scaffolds (55 total) by paired-end information. Genome finishing, including gap closure, repeat resolution, and polishing, was performed as described (59).

**Genome Annotation and Analysis.** Automated gene prediction was performed by using the output of Critica (60) complemented with the output of Generation and Glimmer (61), and is available at http://genome.ornl.gov/microbial/bfun/. The tRNAscanSE tool (62) was used to find tRNA genes, whereas ribosomal RNAs were found by using BLASTn vs. the 16S and 23S ribosomal RNA databases. Other “standard” structural RNAs (e.g., 5S rRNA, rnpB, tRNA, SRP RNA) were found by using covariance models with the Infernal search tool (63). The assignment of product descriptions was made by using search results of the following curated databases in this order: TIGRfam; PRIAM (e−30 cutoff); Pfam; Smart; COGs; Swissprot/TrEMBL (SPTR); and KEGG. If there was no significant similarity to any protein in another organism, it was described as “hypothetical protein.” “Conserved hypothetical protein” was used if at least one match was found to a hypothetical protein in another organism. EC numbering was based on searches in PRIAM at an e−10 cutoff; COG and KEGG functional classifications were based on homology searches in the respective databases.

The automatic annotation was manually curated. The start codons were refined, and pseudogenes were identified. The product description was then curated for all genes. Those genes that had a >70% identity match in *B. pseudomallei* or *B. mallei* or a >70% identity match to a protein with an experimentally verified function were designated with the same product description unless other evidence overruled this annotation. All other product descriptions, except (conserved) hypothetical proteins, were preceded by “putative.” Next, previously studied functions in LB400 were manually curated (taurine metabolism; C1 metabolism; diterpenoid metabolism; and biphenyl and benzoate metabolism). Additionally, a detailed annotation of all aromatic catabolic pathways in LB400 was performed.

Functional redundancy in the LB400 genome was evaluated in the following way: every protein-coding gene in the LB400 genome was searched against the protein database of 260 fully sequenced genomes, including the LB400 genome itself, by using the BLASTp algorithm and a minimum cut-off for a match of at least 30% amino acid identity and an alignable region at least 70% of the length of the query sequence. The cut-off is above the twilight zone of similarity searches where inference of homology is error-prone because of low similarity between aligned sequences (64, 65). Therefore, query proteins were presumably homologous to their match and share at least the same general biochemical function. The genes that had the best match within the LB400 genome (with the exclusion of the self-match) presumably represent duplicated genes (paralogs) and/or genes of similar function that have been laterally acquired (these two processes cannot easily be differentiated) after the emergence of the LB400 lineage and, thus, represent a measure of the functional redundancy within the genome.
We also compared the gene complement of strain LB400 to that of the published B. pseudomallei and the environmental Bce strain S83, which has been fully sequenced by our group, using a reciprocal best BLASTP approach and the same cut-off for a match as above.

**Comparative DNA Microarray Hybridizations.** LB400, LMG 16224, and LMG 21720 DNA were fragmented to 1–5 kb by sonication. Aminoallyl labeling was performed by using the BioPrime labeling kit, based on the manufacturer’s protocol (Invitrogen, Carlsbad, CA), except for the dNTP mixture (final concentrations of 0.5 mM dATP, dGTP, and dCTP; 0.1 mM dTTP; and 0.4 mM aa-dUTP). Sheared DNA (250 ng) was mixed with random hexamers, denatured, snap-cooled, mixed with the dNTP mix and Klenow polymerase, and incubated for 2 h at 37°C. Reaction cleanup, coupling of monoreactive fluorescent Cy dye, and hybridization to the LB400 microarray were performed as described (32, 44). Data normalization was performed in Genespring (Agilent Technologies, Palo Alto, CA) by using “division by control channel,” and data of two biological replicates with dye-swapping were averaged. If for a particular gene the signal produced by the labeled test strain DNA was less than half the signal produced by the LB400 control DNA (threshold) and displayed less than a difference of 2 between the ratios of the biological replicates (data quality filter), that gene was considered absent in the test genome. The current LB400 microarray lacks probes for 356 genes (6.2%). Data for 8,049 genes passed the used data quality filter for both strains.

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