

Complete genome sequence of the metabolically versatile photosynthetic bacterium *Rhodospseudomonas palustris*

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Rhodospseudomonas palustris is among the most metabolically versatile bacteria known. It uses light, inorganic compounds, or organic compounds, for energy. It acquires carbon from many types of green plant-derived compounds or by carbon dioxide fixation, and it fixes nitrogen. Here we describe the genome sequence of *R. palustris*, which consists of a 5,459,213-base-pair (bp) circular chromosome with 4,836 predicted genes and a plasmid of 8,427 bp. The sequence reveals genes that confer a remarkably large number of options within a given type of metabolism, including three nitrogenases, five benzene ring cleavage pathways and four light harvesting 2 systems. *R. palustris* encodes 63 signal transduction histidine kinases and 79 response regulator receiver domains. Almost 15% of the genome is devoted to transport. This genome sequence is a starting point to use *R. palustris* as a model to explore how organisms integrate metabolic modules in response to environmental perturbations.

R. palustris is a purple photosynthetic bacterium that belongs to the alpha proteobacteria and is widely distributed in nature as indicated by its isolation from sources as diverse as swine waste lagoons, earth-worm droppings, marine coastal sediments and pond water. It has extraordinary metabolic versatility and grows by any one of the four modes of metabolism that support life: photoautotrophic or photosynthetic (energy from light and carbon from carbon dioxide), photoheterotrophic (energy from light and carbon from organic compounds), chemoheterotrophic (carbon and energy from organic compounds) and chemoautotrophic (energy from inorganic compounds and carbon from carbon dioxide) (Fig. 1). *R. palustris* enjoys exceptional flexibility within each of these modes of metabolism. It grows with or without oxygen and uses many alternative forms of inorganic electron donors, carbon and nitrogen. It degrades plant biomass and chlorinated pollutants and it generates hydrogen as a product of nitrogen fixation^{1,2}. Thus *R. palustris* is a model organism to probe how the web of metabolic reactions that operates within the confines of a single cell adjusts and reweaves itself in response to changes in light, carbon, nitrogen and electron sources that are easily manipulated experimentally. As a critical step in the further development of this model we have sequenced and annotated the *R. palustris* genome. The genome comprises one circular chromosome that is

5.46 Mb in size. The sequenced strain also harbors a 8.4-kilobase (kb) circular plasmid.

RESULTS

Major features of the genome

The *R. palustris* genome has very few repeat nucleotide sequences, insertion sequence elements or transposons. It has just 16 insertion sequence elements including representatives of the 'phage' integrase family, four ISR1-like elements and two *xerD* type elements. No horizontally transferred islands of DNA are apparent based on anomalous G + C content. *R. palustris* has 4,836 predicted protein-encoding genes (Table 1 and <http://genome.ornl.gov/microbial/rpal/>). These include genes required for the biosynthesis of all its cellular components from carbon dioxide in keeping with its robust growth in media lacking organic carbon sources. *R. palustris* has many genes associated with energy metabolism, reflecting its metabolic versatility (Fig. 2). The chromosomal positions and numbered designations of these genes can be found in Supplementary Table 1 online. There are genes allowing oxidation of hydrogen, thiosulfate and carbon monoxide as energy and reductant sources. Two homologous NADH dehydrogenase complexes that are encoded in the genome likely broker the catabolism of a wide variety of organic compounds, including fatty acids, dicarboxylic

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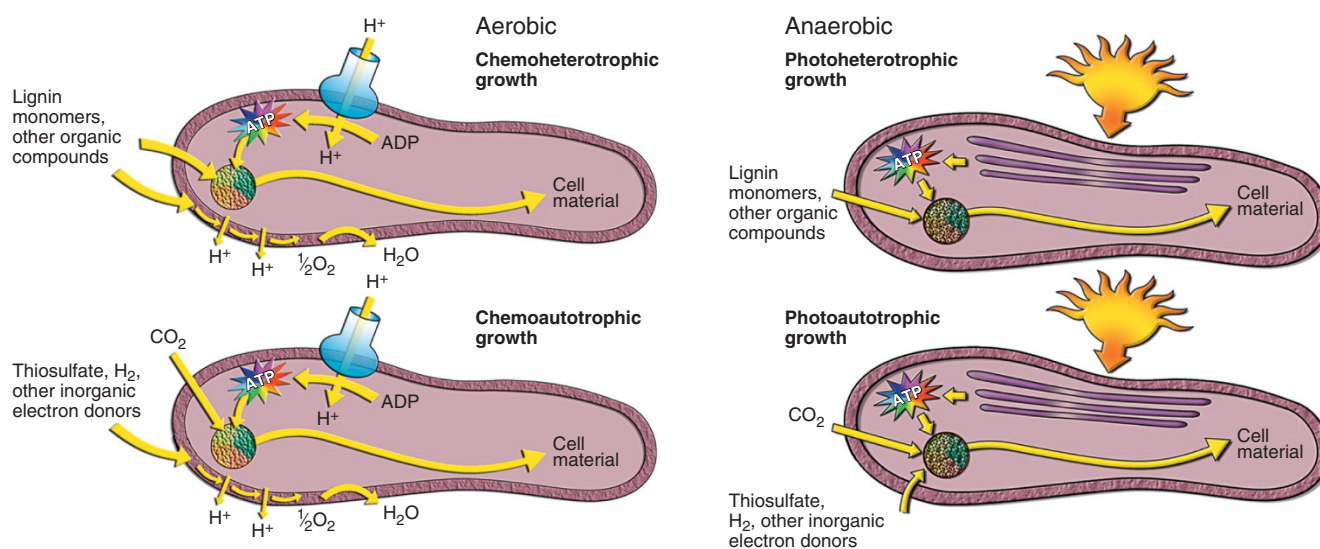


Figure 1 Overview of the physiology of *R. palustris*. Schematic representations of the four types of metabolism that support its growth are shown. The multicolored circle in each cell represents the enzymatic reactions of central metabolism.

acids and lignin monomers. The conditions under which these two seemingly redundant enzyme systems are expressed have not been defined. Terminal oxidase genes should enable *R. palustris* to use nitrite, nitric oxide and nitrous oxide as electron acceptors during anaerobic respiration³. There are four sets of genes for terminal oxidases that can function with oxygen: a cytochrome *aa*₃ oxidase, a cytochrome *cbb*₃ oxidase, a cytochrome *d* quinol oxidase and a quinol *bd* oxidase. Photosynthesis genes enable the use of light as an energy source by cyclic photophosphorylation under anaerobic conditions.

Phototrophy

Genes *rpa1505*–*rpa1554* required for the generation of energy by photophosphorylation reside in a 55-kb region of the *R. palustris* chromosome. These include genes for bacteriochlorophyll and carotenoid biosynthesis as well as genes encoding the L, M and H polypeptides that form the membrane-bound reaction center complex, where light energy is absorbed to initiate electron transfer reactions. The reaction center genes *rpa1527*, *rpa1528* and *rpa1548* are the most highly conserved aspect of this region, sharing from 45 to 60% predicted amino acid identity with the corresponding genes from *Rhodobacter sphaeroides*, a model organism for the study of anoxygenic photosynthesis⁴. However the *R. palustris* reaction center proteins are most similar (on the order of 75% amino acid identity) to homologs in the unusual photosynthetic *Bradyrhizobium* sp. strain ORS278 (ref. 5). This strain forms nitrogen-fixing nodules on the stems of the plant *Aeschynomene sensitiva*, a tropical legume that grows in water logged soils⁶. In addition to a conserved arrangement of photosynthesis genes, the *A. sensitiva* symbiont and *R. palustris* each contain a bacteriophytochrome regulatory gene that is absent in other purple phototrophs. The symbiont's bacteriophytochrome absorbs far-red light and is required for expression of photosynthesis in response to illumination at 740 nm⁷. In our strain the homologous bacteriophytochrome gene *rpa1537* contains a frameshift mutation and is probably inactive. Analysis of rRNA sequences indicates that *R. palustris* is closely related to the *A. sensitiva* symbiont as well as to the soybean symbiont *B. japonicum*⁸. However, *R. palustris* has never been found in symbiotic association with plants, and its genome lacks nodulation genes.

R. palustris, like other purple phototrophic bacteria, responds to lowered light intensity by increasing the amount of light harvesting (LH) complexes. These consist of α and β polypeptides bound to bacteriochlorophyll and a carotenoid, to form a unit that oligomerizes to produce complexes that transfer light energy to the reaction center⁹. The pathway of light energy transfer is LH2 \diamond LH1 \diamond reaction center. *R. palustris* differs from other phototrophs in that it has multiple LH2 complexes that differ slightly in the wavelengths of light absorbed. It tunes its complement of LH2 complexes to harvest light of differing qualities and intensities¹⁰. The genome sequence reveals four complete sets of LH2 genes (*pucBA*) and one incomplete set (Fig. 2 and Supplementary Table 1 online). Two of the four complete sets of *pucBA* genes are located near bacteriophytochrome genes *rpa3015*, *rpa3016* and *rpa1490* that may function in the regulation of LH2 complex gene expression.

R. palustris has genes (*rpa0008* and *rpa0009*) that are similar to the circadian clock genes, *kaiB* and *kaiC* previously identified only in oxygenic photosynthetic bacteria¹¹. *R. palustris* cells present in anoxic environments generate ample energy by photophosphorylation during daylight hours, but may be energy limited at night. Circadian regulation of energy consuming reactions such as nitrogen fixation would make sense, but has yet to be shown in *R. palustris*.

Carbon dioxide fixation

The *R. palustris* genome encodes two active forms of RubisCO, the key enzyme of the Calvin-Benson-Bassham (CBB) pathway of CO₂ fixation¹². The form I (*cbbLS*, *rpa1559* and *rpa1560*) and form II (*cbbM*, *rpa4641*) RubisCO genes are located on almost opposite sides of the chromosome. The *cbbM* gene is linked to other CBB pathway genes in an arrangement that is similar, but not identical to form II *cbb* operons from other purple phototrophs. The *R. palustris* RubisCO form I gene cluster includes an expected divergently transcribed LysR type regulatory gene *cbbR*, but it differs from form I gene clusters in other species in that it includes three additional regulatory genes situated between *cbbR* and the *cbbLS* structural genes. These encode two predicted response regulators (Rpa1556 and Rpa1557) and a hybrid sensor kinase/response regulator (Rpa1558) that contains two PAS domains.

Table 1 General features of the *R. palustris* genome

Total bases	5,459,213	
Gene density	0.881 genes per kb	
Average gene length	987 bases per gene	
Protein coding features	4,836	
Protein coding bases	4,757,178	
Protein coding percentage	87.1	
tRNA	49	
rRNA operons	2, leading strand	
tmRNA	1	
Pseudogenes	17	
GC percentage	65.05	
IS elements	16	
Circular plasmid	1 of 8,427 base pairs	
Gene categories	Number of genes	% of genome
Energy metabolism, biosynthesis, carbon and nitrogen metabolism and cellular processes	1,514	31.0
Transport	700	14.5
Signal transduction	225	4.7
Transcription	288	6.0
Replication and repair	129	2.7
Translation	170	3.5
General function prediction only	404	8.4
Unknown function ^a	432	9.0
Conserved hypothetical	545	11.3
Hypothetical	429	8.9

^aConsists of members of COG group 'S' (function unknown) and also hypothetical and conserved hypothetical genes, not belonging to COG group S, that have been confirmed by proteomics. COG, Clusters of Orthologous Groups of proteins.

Inorganic compounds as a source of reducing power

R. palustris oxidizes inorganic compounds such as thiosulfate and hydrogen gas as energy sources for respiratory growth and as sources of reducing power for carbon dioxide and nitrogen fixation. *R. palustris* has a large cluster of genes (*rpa0959–rpa0979*) for the synthesis and assembly of a nickel-containing uptake hydrogenase. Its periplasmic thiosulfate:cytochrome *c* oxidoreductase complex is encoded by genes *rpa4459–rpa4467* that are very similar to *sox* genes that are found in many other sulfur oxidizing organisms¹³. Its use of reduced sulfur compounds as electron donors sets *R. palustris* apart from closely related phototrophic bacteria¹⁴. The genome also encodes carbon monoxide dehydrogenases and a formate dehydrogenase (Fig. 2 and Supplementary Table 1 online). These can potentially function to supply reductant and substrate for carbon dioxide fixation during anaerobic phototrophic growth or to supply reductant for both energy generation and carbon dioxide fixation under aerobic chemoautotrophic growth conditions.

RubisCO-like proteins

R. palustris is the only organism known to date that encodes two RubisCO-like proteins (RLPs)^{12,15}. RLPs contain varying numbers of substitutions in conserved active site residues. The single RLP from the green sulfur bacterium *Chlorobium tepidum* contains nine active site substitutions and cannot function as a RubisCO¹⁵. One of the *R. palustris* RLPs (RLP2, Rpa0262) is 66% identical to the *C. tepidum* RLP protein and contains the same pattern of active site substitutions. *R. palustris* RLP1 (Rpa2169) has seven active site substitutions distinct from those in its RLP2. A *C. tepidum rlp* mutant is defective in its

ability to oxidize reduced sulfur compounds and from this we infer that the *R. palustris* RLPs are probably involved in sulfur metabolism¹⁶.

Biodegradation

Purple photosynthetic bacteria are a major component of microbial populations found in wastewater treatment facilities exposed to sunlight^{17,18}. *R. palustris* thrives in such environments because it metabolizes structurally diverse compounds found as components of degrading plant and animal wastes. These include lignin monomers, fatty acids and dicarboxylic acids of the types derived from green plants, animal fats and seed oils. *R. palustris* also degrades nitrogen-containing compounds including amino acids and heterocyclic aromatic compounds², and it dehalogenates and degrades chlorinated benzoates and chlorinated fatty acids^{19,20}, compounds that are sometimes found in industrial wastes.

Although *R. palustris* has been studied for its biodegradation abilities and is a model for molecular studies of aromatic ring degradation in the absence of molecular oxygen²¹, its genome has revealed a much larger inventory of degradation genes than expected. It encodes four distinct oxygenase-dependent ring cleavage pathways for the aerobic degradation of the aromatic compounds protocatechuate, homoprotocatechuate, homogentisate and phenylacetate (Fig. 2 and Supplementary Table 1 online). *R. palustris* has the potential to combine oxygen-sensitive and oxygen-requiring enzyme reaction sequences to accomplish complete degradation. An example is the anaerobic transformation of phenol to 4-hydroxyphenylacetate, which is then degraded aerobically via either the homogentisate or homoprotocatechuate pathways²². These types of transformations would be expected to occur in populations straddling oxic to anoxic transition zones. The genome contains 19 mono- or dioxygenase and four cytochrome P450 genes. Additional genes that may be useful in bioremediation or biocatalysis include nitrile hydratase (*rpa2805* and *rpa2806*) and amidase (*rpa2415*) genes, phosphonate utilization genes (*rpa0687–rpa0700*) and carboxylesterase genes (*rpa1568*, *rpa2627*, *rpa3893* and *rpa4646*). The *R. palustris* genome has 16 glutathione *S*-transferase genes, some of which may catalyze the cleavage of β -aryl ether bonds²³.

R. palustris encodes a complete tricarboxylic acid cycle, an Embden-Meyerhof pathway and a pentose phosphate pathway. A predicted glyoxylate shunt permits use of acetate as a sole carbon source, and the genome sequence indicates the synthesis of glycogen and poly β -hydroxyalkanoates as carbon storage polymers. Other genes encode enzymes to mobilize and degrade these polymers during times of carbon starvation. *R. palustris* has a limited ability to grow on sugars and this is reflected by the absence in its genome sequence of glucose or fructose transporters or a hexokinase gene. Genes of the Entner-Doudoroff pathway are absent.

Nitrogen fixation and nitrogen assimilation

We were surprised to find that *R. palustris* has structural genes for three different nitrogenases as well as the related cofactor and assembly genes for these nitrogenases (Fig. 2 and Supplementary Table 1 online). Previously, only *Azotobacter* sp., a heterotrophic obligate aerobe, had been found to encode three nitrogenases. *R. palustris* encodes a molybdenum-dependent nitrogenase, found in all nitrogen-fixing bacteria, and also a vanadium-dependent and an alternative iron nitrogenase. *R. palustris* encodes dinitrogenase reductase ADP-ribosyltransferase (DraT) (Rpa1431 and Rpa2405) and dinitrogenase reductase activating glycohydrolase (DraG) (Rpa2406) enzymes that likely modulate the activity of dinitrogenase reductase by reversible ADP ribosylation. Homologs of NifA (Rpa4632), VnfA (Rpa1374) and

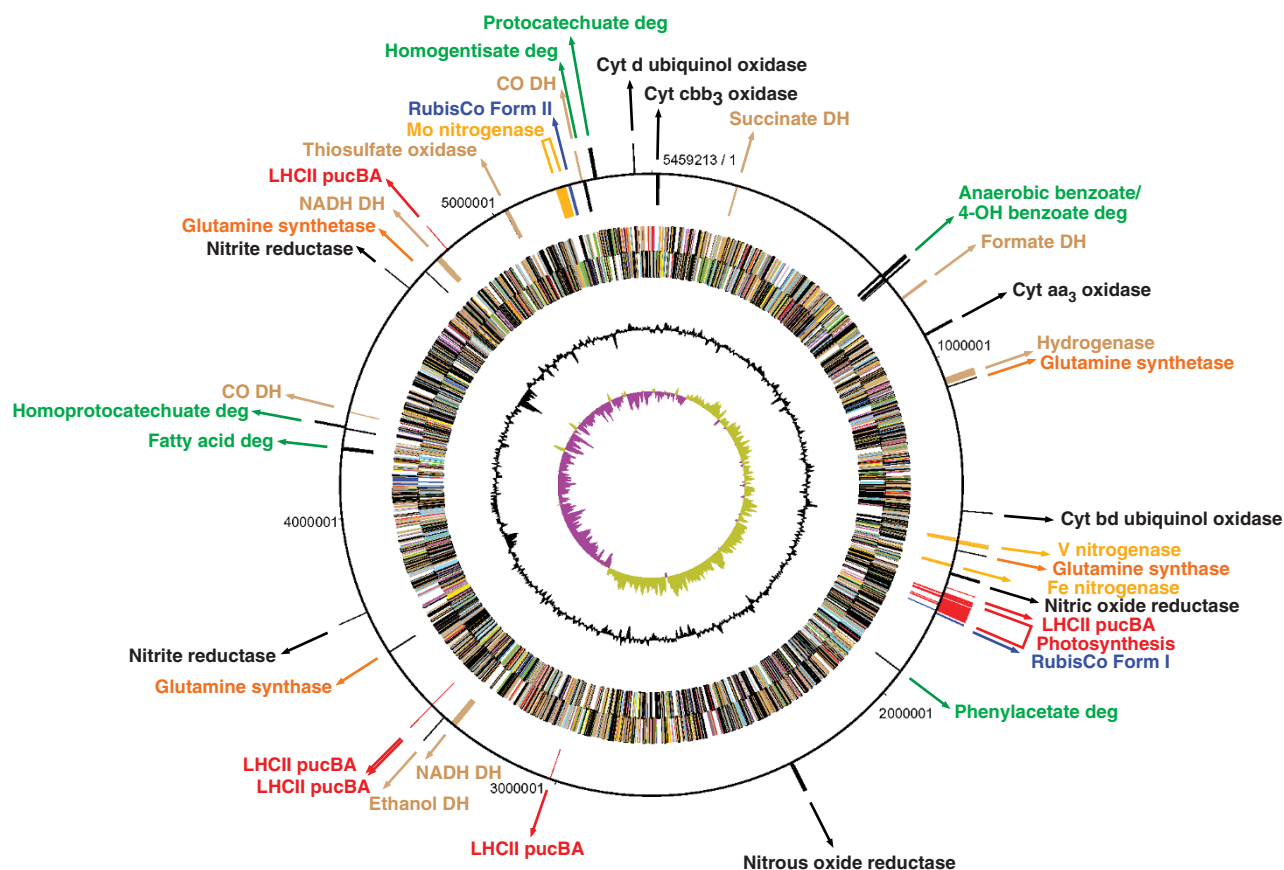


Figure 2 The chromosome of *R. palustris* strain CGA009. Major metabolic features and the locations of the genes that encode them are indicated on the outer circle. Progressing inward, the second circle depicts predicted coding regions on the plus strand colored by functional category: white, hypothetical; dark gray, unknown function; red, replication and repair; green, energy metabolism; blue, carbon and carbohydrate metabolism; cyan, lipid metabolism; magenta, transcription; yellow, translation; pale green, structural RNAs; sky blue, cellular processes; orange, amino acid metabolism; brown, general function prediction; pink, metabolism of cofactors and vitamins; light gray, conserved hypothetical; dark green, transport; lavender, signal transduction; light red, purine and pyrimidine metabolism. Third circle, predicted coding regions on minus strand (same color scheme as the second circle). Fourth circle, G + C content (deviation from average); fifth circle, G + C skew in purple and olive. Scale (in bp) is indicated along the outside of the circle.

AnfA (Rpa1439) regulators are present to potentially activate their cognate clusters of nitrogenase genes in conjunction with the single RNA polymerase sigma factor, RpoN (Rpa0050).

Its genome sequence indicates that *R. palustris* incorporates ammonia exclusively through glutamine synthetase and glutamine:oxoglutarate aminotransferase reactions. It encodes four glutamine synthetases and genes for post-translational control of glutamine synthetase activity by reversible adenylation are present. *R. palustris* has contiguous duplicated, although not identical, *amtB* genes *rpa0273* and *rpa0275* encoding ammonium transporters. Additional transport and metabolic capacity exists to use cyanate (*rpa2115*), urea (*rpa3658–rpa3664*) and ethanolamine (*rpa3747–rpa3749*) as potential nitrogen sources.

Regulation and signal transduction

Because it is a successful metabolic opportunist, *R. palustris* should be able to sense diverse environmental conditions to appropriately regulate gene expression for survival and growth. It also needs to integrate its metabolism and distribute limited pools of ATP and reductant to competing processes such as nitrogen fixation and carbon dioxide fixation. *R. palustris* has 451 potential regulatory and signaling genes, many of which encode multiple domain motifs (Table 2; see

Supplementary Table 2 online for a complete list)²⁴. It devotes about the same proportion of its genes (9.3%) to regulation as do the soil bacteria *Pseudomonas putida*, *Streptomyces coelicolor* and *Streptomyces avermitilis* (<http://www.tigr.org/>). Regulatory genes comprise 5–6% of the genomes of most free-living bacteria. The great variety in the domain architecture of *R. palustris*' 63 signal transduction histidine kinases points to their involvement in regulating many different processes. Half of these genes encode from one to ten predicted transmembrane regions, 20 have PAS domains, 9 have GAF domains (which are characteristic of phytochromes) and 2 have very large, novel cytoplasmic domains. The genome has genes for 19 different RNA polymerase sigma factors, 16 of which are classified as extracytoplasmic function (ECF) sigma factors²⁵. Two of the ECF sigma factor genes (*rpa0639* and *rpa1635*) are located near flagella biosynthesis genes and another (*rpa0550*) is translationally coupled to a gene resembling the cytochrome *c*₂ anti-sigma factor gene *chrR*²⁶, suggesting specific functions.

R. palustris has an acylhomoserine lactone (HSL) synthase gene (*rpa0320*) that is adjacent to the HSL-responsive regulator gene *rpa0321*. HSLs produced by gram-negative bacteria serve as intercellular signals that allow cells to monitor their population density. Generally, HSLs activate expression of genes that are advantageous to

Table 2 *R. palustris* regulatory and signaling proteins

Type of protein	Number encoded in genome	Type of protein	Number encoded in genome
AraC type helix-turn-helix	23	Signal transduction histidine kinase	63
Bacterial regulatory protein, DeoR family	1	Response regulator receiver domain	79
Bacterial regulatory protein, LuxR family	11	Bacterial chemotaxis sensory transducer	30
Bacterial regulatory protein, LysR family	27	CheB methyltransferase	2
Bacterial regulatory protein, MarR family	17	CheR-type MCP methyltransferase	4
Bacterial regulatory protein, ArsR family	9	CheW	4
Bacterial regulatory protein, AsnC family	5	CheY	3
Bacterial regulatory protein, Crp family	15	CheA(STHK)	3
Bacterial regulatory protein, GntR family	13	Cyclic nucleotide-binding domain	4
Bacterial regulatory protein, IclR family	7	EAL domain, DUF2	3
Bacterial regulatory protein, MerR family	3	GGDEF domain, DUF1	22
Bacterial regulatory protein, TetR family	39	EAL/GGDEF domain	14
Transcriptional regulatory protein, C terminal	11	GAF domain	14
Helix-turn-helix Fis type	13	PAC motif	23
Helix-turn-helix protein, CopG family	2	PAS domain	45
Sigma 70 (RpoD)	1	Nitrogen regulatory protein PII (1 GlnB, 2 GlnK)	3
Sigma 54 (RpoN)	1	Serine/threonine protein kinase	3
Sigma 32 (RpoH)	1		
ECF Sigma-24	16	ECF, extracytoplasmic function; MCP, methyl-accepting chemotaxis protein.	

a species when cells of that species are at a population density perceived as a quorum. *R. palustris* genes that might be controlled by quorum sensing include genes *rpa1885–rpa1906* for a phage-like particle called a gene transfer agent²⁷, polyketide synthase gene *rpa3339*, and genes *rpa3342–rpa3357* for the production and export of exopolysaccharides^{28,29}.

R. palustris has genes for three complete chemotaxis signal transduction complexes and it has 30 chemotaxis sensory transducer genes. All but five of the transducers are predicted to be membrane-bound proteins. Four of the transducer genes (*rpa4202*, *rpa4311*, *rpa4481* and *rpa4483*) are translationally coupled to or located just a few base pairs away from a sensor gene with a PAS domain. These gene pairs may have originally existed as single genes but have been translationally frameshifted. The existence of the same split genes in *Magnetospirillum magnetotacticum* and *Rhodospirillum rubrum* suggests that this arrangement may have been present in an ancestor common to these three organisms.

Transport

The genome of *R. palustris* encodes about 325 transport systems comprising at least 700 genes, adding up to almost 15% of the genome. Transport genes account for 5–6% of most bacterial genomes³⁰. A complete listing, classified using the TC Number system³¹ can be found as **Supplementary Table 3** online. There are 102 primary transport systems, defined as systems powered directly by ATP hydrolysis. These include 86 ATP-binding cassette (ABC) systems and 7 P-type ATPases and type II, III and IV secretion systems. The P-type ATPases likely confer resistance to heavy metals³². Separate *R. palustris* Type II secretion systems are likely used for the biogenesis of type IV pili and general protein secretion (the Sec system), with a type III secretion system for flagella biosynthesis. *R. palustris* has two sets of type IV secretion genes (*rpa2224–rpa2233* and *rpa4115–rpa4124*) similar to the *Trb* genes from *Agrobacterium tumefaciens* for conjugal transfer of DNA³³.

R. palustris encodes 137 secondary transport systems including 36 major facilitator superfamily (MFS) members, 22 resistance-nodulation-cell division (RND) pumps, 15 divalent metal transport (DMT) members and 8 tripartite ATP-independent periplasmic (TRAP) transporters^{34,35}. All but two of the RND systems are classified as heavy

metal and drug efflux pumps. This is the largest number of RND pumps observed in any bacterium to date and may explain the high intrinsic resistance of *R. palustris* to antibiotics. *R. palustris* has been isolated in high numbers from polluted environments³⁶. Heavy metal efflux transporters should allow *R. palustris* to live in a variety of environments and still acquire the necessary nutrients while resisting heavy metal toxicity.

Of the 86 ABC systems, 20 are related to the branched chain amino acid uptake (*ilvFGHKL*) system of *E. coli*. Isoleucine, leucine and valine are hydrophobic amino acids and we speculate that other members of this amplified family are specific for other sorts of hydrophobic compounds such as lignin monomers, fatty acids and dicarboxylic acids derived from oils and fats. One system of this *ilv* ABC family (Rpa0665–Rpa0668) has tentatively been identified as a 4-hydroxybenzoate transport system²¹. Another (*rpa1789* and *rpa1791–1793*) lies adjacent to a feruloyl

CoA ligase gene implying that it catalyzes the uptake of the lignin monomer ferulate. A third example is an *ilv* family ABC system (*rpa3719–3725*) that is next to genes for the degradation of the dicarboxylic acid pimelate. An analysis of 73 other microbial genomes shows that 34 of them have no *ilv*-like transport systems. Another 25 microbes have between one and five of these systems and 11 microbes have between six and ten *ilv* family ABC transporters. Only three other species, *Burkholderia fungorum* LB400 and *Ralstonia eutropha*, both β -proteobacteria, and *B. japonicum*, have 19 or more versions of the *ilv*-like ABC transport operon.

Iron acquisition appears to be particularly important for *R. palustris*. It encodes 24 outer membrane ferric iron siderophore receptors, and 7 TonB systems for powering these and other outer membrane receptors (**Supplementary Table 3** online). This implies that *R. palustris* uses a large number of different types of siderophores for iron acquisition. However, genes *rpa2388–rpa2390* to synthesize only one siderophore, rhizobactin³⁷, were detected suggesting that *R. palustris* may transport iron-loaded siderophores produced by other soil bacteria. As many as seven of the ECF sigma factors encoded by *R. palustris* are either translationally coupled to ferrisiderophore-like receptor genes or are located very close to genes involved in iron acquisition; in one case siderophore biosynthesis genes and in another, a predicted heme uptake system. This suggests a role for multiple alternative sigma factors in activating gene expression in response to iron starvation³⁸.

DISCUSSION

R. palustris owes much of its metabolic versatility to known genes encoding metabolic modules of carbon dioxide fixation and photophosphorylation that act in concert with dehydrogenases, oxidoreductases and carbon degradation pathways to support its four modes of growth (Fig. 1). The number of options that *R. palustris* has within the major metabolic modes to take advantage of fluctuating supplies of carbon, nitrogen, light and oxygen is unusually large. The existence of genes for three nitrogenases, multiple aromatic degradation pathways and multiple oxidoreductases was not known before the genome sequence. Its large inventory of transport and chemotaxis genes implies that *R. palustris* is adept at sensing and acquiring diverse compounds from its environment. The groundwork has now been laid to

explore regulatory strategies used by *R. palustris* to appropriately select and integrate its large number of metabolic choices.

R. palustris is ideally suited for use as a biocatalyst because it generates ample supplies of ATP from light thus catalyzing reactions that are thermodynamically unfavorable and beyond the potential of chemotrophic organisms. The metabolic group of purple phototrophic bacteria to which it belongs have been evaluated as sources of single cell protein, for the synthesis of polyhydroxyalkanoate 'bioplastics' and for the production of hydrogen, which they generate as a product of nitrogen fixation³⁹. Its genome sequence reveals that *R. palustris* has additional capabilities, not shared by other purple bacteria, that enhance its potential for use in biotechnological applications. These include modulating photosynthesis according to light quality and degrading aromatic compounds that are typically found in agricultural and industrial wastes. That the genome encodes oxygen-requiring, as well as anaerobic reductive pathways, for the degradation of aromatic rings, suggests the possibility of designing hybrid degradation pathways of broader substrate specificity than those that occur naturally. *R. palustris* has physical attributes that are well suited for process development. It undergoes asymmetric cell division and produces a cell surface adhesin at one end of the cell that causes cells to stick to solid substrates. *R. palustris* has especially good potential for use as a biocatalyst for hydrogen production. It is unique among purple phototrophic bacteria in encoding a vanadium-containing nitrogenase that catalyzes the production of approximately three times as much hydrogen as do molybdenum-containing nitrogenases⁴⁰. *R. palustris* derives reductant for hydrogen generation from plant biomass, and energy captured from sunlight drives the process. Manipulating *R. palustris* to produce hydrogen efficiently will require a detailed knowledge of how each of its three nitrogenases is regulated. It will also be important to know in detail how the metabolic modules of photophosphorylation, biodegradation, carbon dioxide fixation and hydrogen uptake are regulated and how their activities are integrated.

METHODS

Construction, isolation and sequencing of small-insert and large-insert libraries. Genomic DNA, isolated from the *R. palustris* CGA009, was sequenced using a conventional whole genome shotgun strategy⁴¹. Briefly, random 2–3 kb-DNA fragments were isolated after mechanical shearing. These gel-extracted fragments were concentrated, end-repaired and cloned into pUC18. Double-ended plasmid sequencing reactions were carried out using PE BigDye Terminator chemistry (Perkin Elmer) and sequencing ladders were resolved on PE 3700 Automated DNA Sequencers. One round (117,510 reads) of small-insert library sequencing was done, generating roughly 9.6-fold redundancy.

A large insert (~30 kb) fosmid library was also constructed by *Sau3AI* partial digestion of genomic DNA and cloning into the pFos1 cloning vector⁴². End sequencing of ~300 fosmid clones (0.02-fold redundancy) generated roughly 2-fold genome scaffold coverage. The fosmids were fingerprinted with *EcoRI* to aid in assembly verification and determination of gap sizes and provided a minimal scaffold used for order and orientation across assembly gaps. The 8.4-kb plasmid was assembled from a total of 107 reads.

Sequence assembly and gap closure. Sequence traces were processed with Phred^{43,44} for base calling and assessment of data quality before assembly with Phrap (P. Green, University of Washington, Seattle, Washington, USA) and visualization with Consed⁴⁵. Gaps were closed by primer walking on gap-spanning library clones (identified using linking information from forward and reverse reads). Alternatively, some of the larger gaps, including the larger regions covered only by fosmid clones, were closed by primer walking on PCR products. Remaining physical (uncaptured) gaps were closed by combinatorial (multiplex) PCR. Sequence finishing and polishing added a total of 300 reads and assessment of final assembly quality was done as previously described⁴⁶.

Sequence analysis and annotation. Gene modeling was done using the Critica⁴⁷, Glimmer⁴⁸ and Generation (<http://compbio.ornl.gov/generation/index.shtml>) modeling packages, the results were combined and a basic local alignment search tool (BLAST) for proteins (P) search of the translations versus GenBank's nonredundant database (NR) was conducted. The alignment of the N terminus of each gene model versus the best NR match was used to pick a preferred gene model. If no BLAST match was returned, the Critica model was retained. Gene models that overlapped by greater than 10% of their length were flagged, giving preference to genes with a BLAST match. The revised gene/protein set was searched against the KEGG GENES, InterPro (incorporating Pfam, TIGRFams, SmartHMM, PROSITE, PRINTS and ProDom) and Clusters of Orthologous Groups of proteins (COGs) databases, in addition to BLASTP versus NR. From these results, categorizations were developed using the KEGG and COGs hierarchies. Initial criteria for automated functional assignment required a minimum 50% residue identity over 80% of the length of the match for BLASTP alignments, plus concurring evidence from pattern or profile methods. Putative assignments were made for identities down to 30%, over 80% of the length. Automated assignments were reviewed and curated manually using a web-based editing environment.

Nucleotide sequence accession number. The sequence of the complete genome of *R. palustris* CGA009 is available under GenBank/EMBL/DDJB accession numbers BX571963 (chromosome) and BX571964 (plasmid).

Note: Supplementary information is available on the Nature Biotechnology website.

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COMPETING INTERESTS STATEMENT

The authors declare that they have no competing financial interests.

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