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**Recent Work** 

## Title

Genetic Elucidation of Metabolic Diversity in Dechloromonas aromatica strain RCB

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### Author

Coates, J.D.

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Abstract: Dechloromonas aromatica is a beta class proteobacteria found ubiquitously in soil environments. A facultative anaerobe, capable of aromatic hydrocarbon degradation and perchlorate reduction, it was first isolated from BTEX contaminated Potomac river sediments. Due to its metabolic versatility, strain RCB was selected for complete genome sequencing at the DOE Joint Genome Institute. The completed sequence consists of a single circular chromosomal DNA structure containing a total of 4.5 Mbases of DNA with a G+C content of 60%. Initial draft annotation was conducted at Oak Ridge National Laboratories and the Virtual Institute for Microbial Stress and Survival (http://escalante.lbl.gov) identifying approximately 4000 open reading frames (ORFs). The closest relative with a completed genome sequence is *Ralstonia sola*nacearum, also a gram negative proteobacteria of the Beta sub-class. Dechloromonas aromatica and Ralstonia possess very similar genomic content, duplicative 23S and 16S RNA regions, and identical operon structures for many families of genes. In Ralstonia, the catabolic capabilities leading to degradation of aromatic compounds are thought to be concentrated within plasmid DNA structures. However, D. aromatica appears to lack a large genomic plasmid element (*R eutropha's* pGMI1000MP plasmid is 2.09Mb in size), which indicates its adaptability in degradation of aromatic and toxic compounds is not dependent on a highly transmissible and mutable plasmid structure. Genes have been annotated for InterPro domains, EC assignments, TIGRfams, KEGGS pathways, and GO ontologies. Genes that likely reflect the versatility of *D. aromatica* within the environment include a remarkably high number of two-component sensors and regulators (11% of the annotated ORFs), placing it in the top 5% of microbial genomes in this capability. The D. aromatica genome has been found to contain RuBisCo, indicating the ability to fix carbon dioxide. Conversely, central catabolic pathway enzymes for monoaromatic degradation such as BSS are not present, suggesting the presence of novel aromatic degradation pathways in this organism.

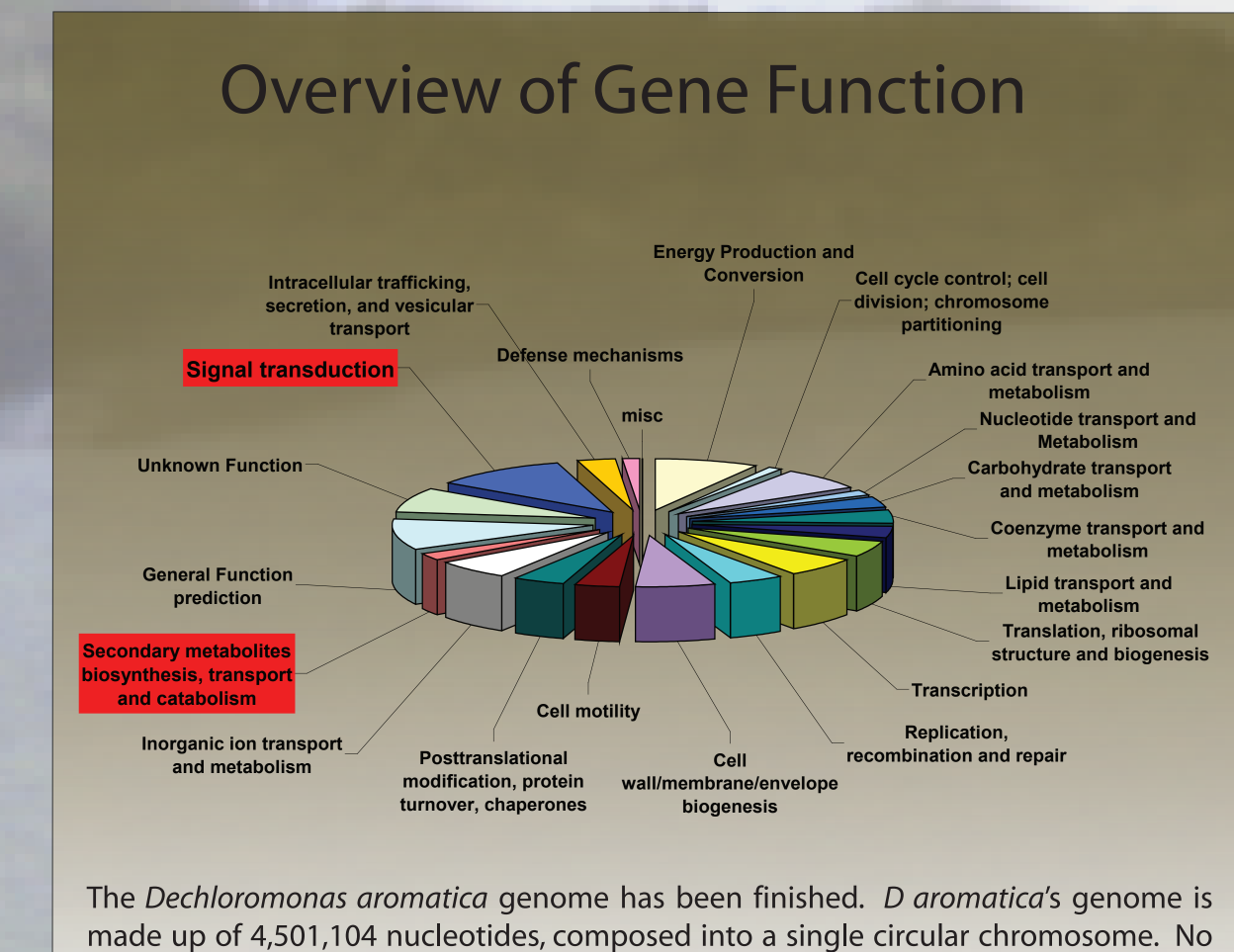
# What's Different about Dechloromonas aromatica?

- A disproportionately high number of two-component sensors and phosphorelay proteins (11% of the genome).

- A highly versatile metabolism, allowing *D. aromatica* to reduce nitrate, oxidize aromatic BTEX compounds, and reduce perchlorate -- all harmful environmental contaminants.

- The genomic sequence indicates *D. aromatica* contains a CO<sub>2</sub> fixation pathway, with the presence of RuBisCo, as well as the ability to fix N<sub>2</sub>. The presence of these capabilities predicts currently unexplored physiological capabilities for this novel organism.

- Though the initial recruitment to established gene families indicates a low number of transport proteins, further analysis indicates a fairly large group of transporters, but they do not belong to the nucleotide permease families expressed in other microbial genomes.

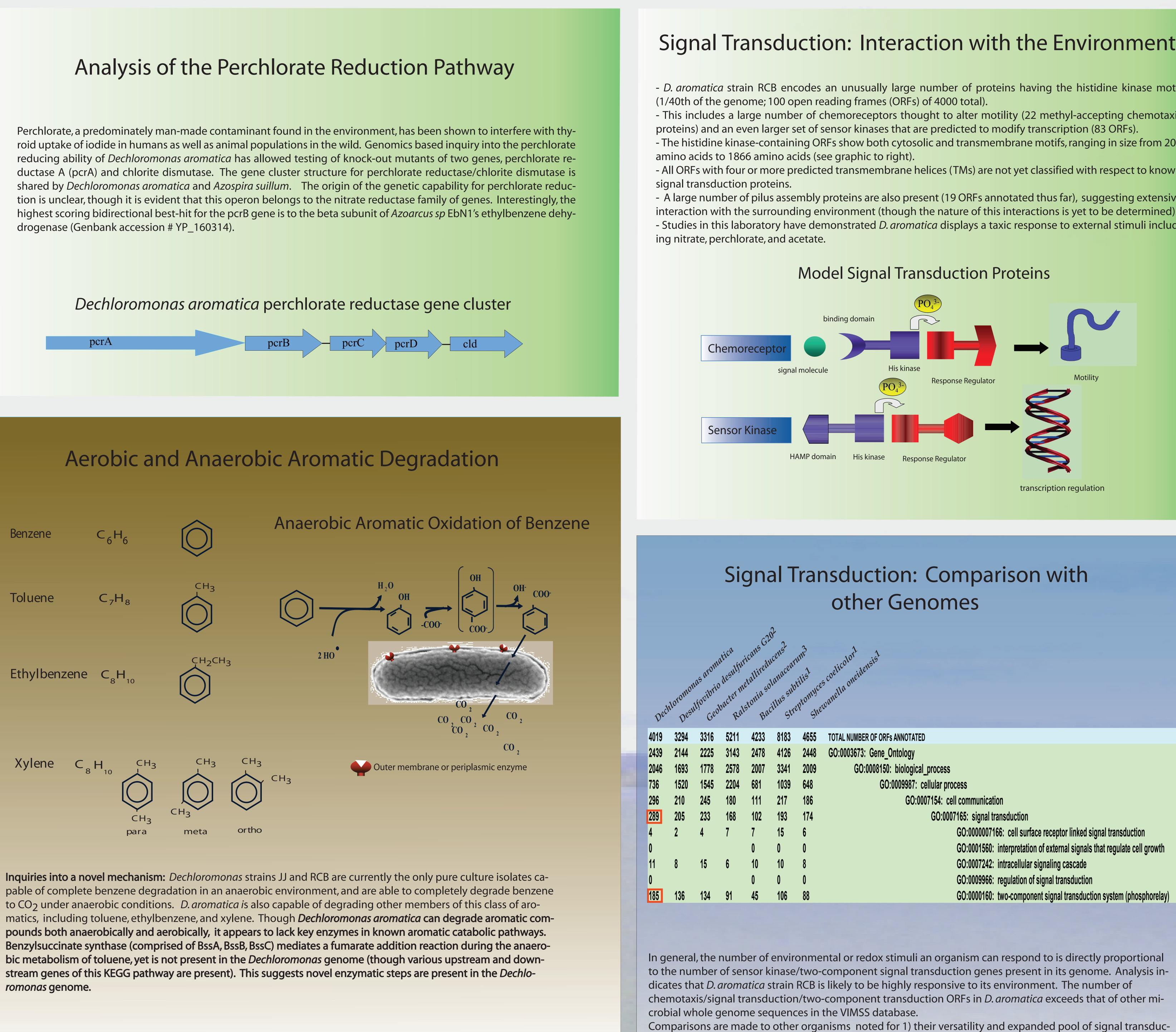


evidence of plasmid DNA was seen.

# Genetic Elucidation of Metabolic Diversity in **Dechloromonas aromatica** strain RCB

K. V. Kellaris<sup>1</sup>, W. S. Feil<sup>1</sup>, H. Feil<sup>1</sup>, R. Chakraborty<sup>1</sup>, E. Alm<sup>1</sup>, K. Huang<sup>1</sup>, M. Price<sup>1</sup>, K. Keller<sup>1</sup>, S. O'Connor<sup>1</sup>, A. Arkin<sup>1</sup>, A. Lapidus<sup>2</sup>, S. Trong<sup>2</sup>, G. Di Bartolo<sup>2</sup>, F. W. Larimer<sup>3</sup>, P. M. Richardson<sup>2</sup>, L.A. Achenbach<sup>4</sup>, J. D. Coates<sup>1</sup> <sup>1</sup>University of California, Berkeley, CA, <sup>2</sup>Joint Genome Institute, Walnut Creek, CA, <sup>3</sup>Oak Ridge National Laboratories, Oak Ridge, TN, <sup>4</sup>Southern Illinois University, Carbondale, IL.

pcrA	pcrB —	pcrC pcrI	D cld	



Reference: Chakraborty, R. and Coates, J.D. (2005). Hydroxylation and carboxylation – two crucial steps of anaerobic benzene degradation by Dechloromonas strain RCB. Applied and Environmental Microbiology (in press).

John D. Coates Email: jcoates@nature.berkeley.edu Phone: (510) 643-8455 Fax: (510)643-4999



	Soluble (no predicted TM): small (<500aa):	6
	medium (500-100aa):	13
	large (1000 - 2000aa): includes two cheA-like proteins and glnL	4
	(involved in nitrogen regulation)	
	One predicted TM:	
	small (<500aa):	1
	medium (500-100aa):	5
	large (1000 - 2000aa): includes phoR and barA	2
	Two predicted TMs (most typical):	
	small (<500aa): $(500, 1000aa)$	11
	medium (500-1000aa): large (1000 - 2000aa):	27 9
	includes 6 copies of the quorum sensor qse	
	envZ, rstB, narX and two barA homologs	
	Three predicted TMs:	
	small (<500aa):	2
	medium (500-1000aa): includes arcB, and kdpD	3
	Four predicted TMs:	
	small (<500aa):	1
	medium (500-1000aa):	3
	Five predicted TMs:	
	small (<500aa):	2
_	medium (500-1000aa): large (1000 - 2000aa):	4
	Seven predicted TMs:	1
	medium (800aa):	I
	Eight predicted TMs:	
	medium (535aa):	1
	Nine predicted TMs:	
	medium (704aa):	1
	Twelve predicted TMs:	1
	medium (903aa):	1
	Fourteen predicted TMs:	
	large (1131aa):	1
	Conclusions	

tion genes, 2) from similar environments with similar capabilities, or 3) close phylogenetic neighbors.

and transport than other microbes sequenced thus far. The reason for this is unknown. - Pathway analysis indicates the ability to fix CO<sub>2</sub> and N<sub>2</sub> exists in this organism. - Several members of currently known catabolic pathways for aromatic degradation have not been found by analysis of the completed genome, suggesting the presence of novel enzymes for aromatic catabolism.

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