

Genome Sequence of *Azotobacter vinelandii*, an Obligate Aerobe Specialized To Support Diverse Anaerobic Metabolic Processes^{∇†}

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Azotobacter vinelandii is a soil bacterium related to the *Pseudomonas* genus that fixes nitrogen under aerobic conditions while simultaneously protecting nitrogenase from oxygen damage. In response to carbon availability, this organism undergoes a simple differentiation process to form cysts that are resistant to drought and other physical and chemical agents. Here we report the complete genome sequence of *A. vinelandii* DJ, which has a single circular genome of 5,365,318 bp. In order to reconcile an obligate aerobic lifestyle with exquisitely oxygen-sensitive processes, *A. vinelandii* is specialized in terms of its complement of respiratory proteins. It is able to produce alginate, a polymer that further protects the organism from excess exogenous oxygen, and it has multiple duplications of alginate modification genes, which may alter alginate composition in response to oxygen availability. The genome analysis identified the chromosomal locations of the genes coding for the three known oxygen-sensitive nitrogenases, as well as genes coding for other oxygen-sensitive enzymes, such as carbon monoxide dehydrogenase and formate dehydrogenase. These findings offer new prospects for the wider application of *A. vinelandii* as a host for the production and characterization of oxygen-sensitive proteins.

Azotobacter vinelandii is a free-living nitrogen-fixing bacterium of the gammaproteobacteria. It is found in soils world-

wide, with features of nitrogen and energy metabolism relevant to agriculture (41, 42). This organism has been studied for more than 100 years by numerous scientists throughout the world. Prior to Joshua Lederberg's discovery of sexuality in *Escherichia coli* (47), *A. vinelandii* was the experimental organism of choice for many investigators during the emergence of biochemistry as a dominant discipline within the life sciences. Examples include the classical Lineweaver-Burk kinetic parameters, developed using enzymes from *A. vinelandii* (51), and the isolation by Severo Ochoa of polynucleotide phosphor-

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ylase from *A. vinelandii*, which was used in studies that contributed to the elucidation of the genetic code (62).

A. vinelandii is able to adapt its metabolism to diverse sources of nutrients. If no carbon source is present, *A. vinelandii* will undergo a differentiation process to form cysts that are resistant to desiccation and other chemical and physical challenges (74). While the process of encystment has been known for many years and studied at the physiological and morphological levels, there is little knowledge about the unique biosynthetic pathways that are involved and how they are regulated. Previous work has implicated the alternative sigma factors AlgU and RpoS in the differentiation process (13, 57, 64). Alginate polymers with different monomer compositions are an important structural component of the cyst, and at the end of exponential growth, *A. vinelandii* cells accumulate poly-beta-hydroxybutyrate (PHB) as a reserve carbon and energy source (81). The physiology of PHB formation has been well studied in a variety of different systems, and the PHB biosynthetic operon has been described (67, 77). *A. vinelandii* can also produce copolymers of hydroxybutyrate and hydroxyvalerate, known to improve the flexibility and stretch of bioplastics (63).

A. vinelandii has long served as a model for biochemical and genetic studies of biological nitrogen fixation, the conversion of N₂ into NH₃ by a nitrogenase enzyme. The best-studied nitrogenase consists of two oxygen-sensitive metalloproteins that, in the case of the molybdenum nitrogenase, are denominated the Fe protein and the MoFe protein. *A. vinelandii* is unusual in that it is one of the few bacteria that contain three nitrogenases with different subunit and metal cofactor compositions, namely, the molybdenum nitrogenase, the vanadium nitrogenase, and the iron-only nitrogenase. Expression of these nitrogenases is differentially regulated by metal availability from the medium (27).

Here we present the complete genome sequence of *A. vinelandii* DJ and discuss what the genome has revealed about the organism's ability to protect oxygen-sensitive processes. *A. vinelandii* has been cited as having one of the highest respiratory rates of any known bacterium (10). Diazotrophic growth under aerobic conditions is possible because *A. vinelandii* can adjust oxygen consumption rates to help maintain low levels of cytoplasmic oxygen, which is otherwise detrimental not only to nitrogenase but also to other oxygen-sensitive enzymes expressed by *A. vinelandii*. This phenomenon is called respiratory protection. In this work, we identify unique features of the *A. vinelandii* genome that help to explain the coexistence of oxygen-sensitive reactions and strict aerobic metabolism. The genome sequence and annotation allowed identification of the genes involved in respiration, including key players in respiratory protection. In addition, we have identified unexpected gene clusters encoding a carbon monoxide dehydrogenase (CODH), a formate dehydrogenase (FDH), and a second hydrogenase, all of which are also oxygen-sensitive enzymes.

MATERIALS AND METHODS

Strain description. *A. vinelandii* O is a strain that forms gummy, slimy colonies of pale color. The earliest report of this strain dates from 1952 (92). *Azotobacter agilis* (later named *Azotobacter vinelandii* strain O) was part of the bacterial collection at the University of Wisconsin, Madison. In 1959, Bush and Wilson (11) reported the isolation of a nongummy chromogenic isolate of *A. vinelandii* O, which they named *A. vinelandii* OP. This new strain has a fluorescent color

and well-defined colony shape. *A. vinelandii* strain OP has also been referred to as UW and CA. *A. vinelandii* strain DJ is a high-frequency transforming variant of *A. vinelandii* UW generated in 1984 through multiple rounds of transformation, using chromosomal DNAs from rifampin (rifampicin)-resistant and -sensitive strains (Dennis Dean, personal communication), and is available from the American Type Culture Collection under number ATCC BAA-1303.

Genome sequencing and assembly. A total of 87,000 reads were generated by the Joint Genome Institute for a draft sequence released prior to this project. These reads came from two DNA libraries (insert sizes of 2 to 4 kb and 4 to 8 kb) obtained using mechanical shearing of DNA and cloning into pUC18, followed by a shotgun sequencing approach. The reads and clones were sent to Monsanto, where finishing occurred. The genome was then assembled and edited using Phred, Phrap, and Consed (23, 24, 32). Finishing was completed by generating an optical map (46) cut with the restriction enzymes BamHI and BsiWI and aligning the assembled sequences to the map. Gaps were closed by sequencing specific products. All rRNA operons were amplified with specific flanking primers, sequenced, and assembled individually. All positions with Phred scores of <40 were resequenced using an independent PCR fragment as a template. The error rate is estimated to be less than 1:10,000 bp.

Annotation and comparative genomics. Genome annotation was done by supervised teams of undergraduate students using a web-based system over a preliminary automated annotation, both developed by J. C. Setubal. Comparison to other genomes was done using MUMmer (45) for alignments and orthoMCL (50) for protein families.

Phylogenetic analysis. A total of 18 genomes (see Table S1 in the supplemental material) were used to build the phylogenetic tree. *Chromohalobacter selexigens*, *Hahella chejuensis*, and *Marinomonas* sp. strain MWYL1 were used as outgroup species. OrthoMCL (50) provided 1,445 protein families containing only one representative member of each ingroup genome. These protein sequences were aligned with Muscle (20), and the noninformative columns of the resulting (concatenated) alignment were removed by Gblocks (14). The tree was built using RAxML (79) with the PROTGAMMAWAGF model. Individual gene phylogenies were obtained by aligning protein sequences from *A. vinelandii* and *Pseudomonas* species, using the same method as that described above, but using MrBayes (35) for tree building.

Prediction of σ^{54} - and NifA-binding sites. Multiple alignments of known σ^{54} - and NifA-binding-site DNA sequences were constructed from the literature (3). Using HMMER 1.8.5 (25), we generated profile hidden Markov models representing the binding sequence motifs from the alignments and used HMMER to search the genome for matches.

CydR binding site prediction. We used the sequence proposed by Wu et al. (93) as a query against the genome in a BLASTN (1) search.

Nucleotide sequence accession number. The *A. vinelandii* DJ sequence and its annotation are available at GenBank under accession number CP001157 and at www.azotobacter.org.

RESULTS

General features and phylogeny. The general features of the *A. vinelandii* genome are shown in Table 1. Based on the sequence and its annotation, we have obtained a phylogeny for *A. vinelandii* (Fig. 1) that indicates that it groups most closely with the pseudomonads (family *Pseudomonadaceae*). Among the pseudomonads, the closest relative is the nitrogen-fixing strain *Pseudomonas stutzeri* A1501 (96). Table 1 shows that *A. vinelandii* and *P. stutzeri* A1501 share 46% and 56% of their respective protein coding gene complements. Figure S1 in the supplemental material presents a whole-chromosome alignment between *A. vinelandii* DJ and *P. stutzeri* A1501 in which large-scale conservation of gene order can readily be observed. The "X" pattern seen in this comparison is likely due to inversions around the origin of replication (22) and suggests that many such inversions have taken place since these two species diverged.

Respiration and respiratory protection genes. *A. vinelandii* is an obligately aerobic bacterium that uses the electron transport chain, with molecular oxygen as the final electron acceptor. This strict aerobic metabolism is supported by our analysis

TABLE 1. General features of *Azotobacter vinelandii* DJ and *Pseudomonas stutzeri* A1501 genomes^a

Parameter	Value	
	<i>Azotobacter vinelandii</i> DJ	<i>Pseudomonas stutzeri</i> A1501
Chromosome features		
Size (bp)	5,365,318	4,567,418
% GC	65.7	63.9
Protein-encoding genes		
Total no.	5,051	4,128
No. (%) with functional assignment	3,561 (70.5)	3,180 (77.0)
No. (%) of conserved hypothetical protein genes	739 (14.6)	0 (0.0)
No. (%) of hypothetical protein genes	751 (14.9)	948 (23.0)
No. of pseudogenes	66	11
No. (%) shared protein-encoding genes	2,332 (44.1)	2,298 (55.7)
RNAs		
No. of rRNA operons	6	4
No. of tRNAs	64	61
No. of other RNAs	18	0

^a Data for *P. stutzeri* were taken from reference 96. The number of shared protein-encoding genes was computed using orthoMCL (50).

of the genome, which lacks genes for complete systems involved in anaerobic respiration using alternative terminal electron acceptors or fermentation processes. At the interface of catabolism and respiration, NADH and succinate are the major electron carriers that feed the electron transport chain, via complex I and complex II, respectively. The genome encodes four NADH-ubiquinone oxidoreductases, including one ATP-coupled NADH oxidoreductase of the Nuo type (Avin28440 to Avin28560) and three ATP-uncoupled NADH-ubiquinone oxidoreductases. The latter are Ndh (Avin12000) and two membrane-associated enzymes involved in the transport of cations across the membrane, i.e., the Nqr type (Avin14590 to Avin14640) and the Sha type (Avin19530 to Avin19580). While there is no association of Nuo, Nqr, and Sha with oxygen protection, the *ndh* gene product has been shown to be important for aerobic nitrogen fixation (5). It is possible that NADH-driven protection of nitrogenase is dependent on Ndh and CydAB oxidase via a ubiquinol-dependent electron transfer pathway. Despite its apparent involvement in nitrogen fixation (5), a copy of *ndh* is found in several *Pseudomonas* species, some of which are not known to fix nitrogen. In addition, *A. vinelandii* carries two copies of the Rnf system (Rnf1 [Avin50930 to Avin50980] and Rnf2 [Avin19220 to Avin19270]), which shows sequence similarity to a sodium-dependent NADH-ubiquinone oxidoreductase (18). Although there is no change in respiration upon inactivation of either one or both Rnf systems, Rnf1 is required for accumulation of nitrogenase Fe protein (18). An ortholog of Rnf1 exists in the diazotroph *P. stutzeri* A1501 (YP_001171857 to YP_00117852) but not in any other members of the *Pseudomonas* genus.

Ubiquinol-oxygen oxido-reduction can occur either through a two-step pathway, via cytochrome *c* reductase (complex III; Avin13060 to Avin13080) followed by a cytochrome terminal ox-

idase (complex IV), or in a single-step process, via a ubiquinol-dependent cytochrome terminal oxidase. The genome annotation identified the catalytic and biosynthetic genes of the following five terminal oxidases: cytochrome *c* oxidase (Cdt oxidase; Avin00950 to Avin01020), cytochrome *o* oxidase (Cox; Avin11170 to Avin11180), cytochrome *bd* copy I (CydAB I; Avin19880 to Avin19890), cytochrome *bd* copy II (CydAB II; Avin11040 to Avin11050), and cytochrome *ccb*₃ (Cco; Avin19940 to Avin20010). While the Cox, CydAB II, and Cco oxidases are encoded in all *Pseudomonas* species analyzed to date, the Cdt oxidase gene is found only in *P. stutzeri* A1501 and *Pseudomonas mendocina* ymp.

The presence of two CydAB oxidases in *A. vinelandii* was unexpected. Phylogenetic analysis revealed that CydAB oxidase I (encoded by *cydAB* gene copy I) has its closest orthologs in *P. stutzeri* A1501 but groups with similar genes from members of the *Acinetobacter* and *Shewanella* genera rather than with those from other members of the *Pseudomonas* genus. Copy II, on the other hand, groups with similar genes from several members of the *Pseudomonas* genus (see Fig. S2 to S5 in the supplemental material). CydAB oxidase I has been characterized extensively (21, 39, 40, 94). The precise function of CydAB oxidase II remains to be elucidated.

Oxygen consumption by the aforementioned terminal oxidases is not the only factor responsible for oxygen protection of nitrogenase. The FeSII protein, known as the Shethna protein and encoded by Avin01520, forms a protective complex with nitrogenase when the enzyme is exposed to oxygen (58). In addition, it has been determined that cellular levels of ATP also contribute to protection of nitrogenase against oxygen damage (53). High concentrations of ATP directly correlate with high electron flux to nitrogenase, which influences the dissociation rate constant of the nitrogenase components (84) and, consequently, the susceptibility of the Fe protein to oxygen damage (83). The *A. vinelandii* genome encodes two sets of ATP synthase machineries. The first complex (Avin52150 to Avin52230) is the ortholog of the *Pseudomonas* ATP synthase and is located close to the putative origin of replication. The second copy (Avin19670 to Avin19750) is located 8 genes upstream of the *sha* operon (complex I-like) and 10 genes downstream of *cydAB* I (complex IV). This copy does not seem to be related phylogenetically to ATP synthases from the *Pseudomonas* genus, and the only orthologs are found in *Burkholderia* species, most of which are obligate aerobes, although very few can fix nitrogen.

Genomic analyses of *A. vinelandii* and microaerobic diazotrophs such as *P. stutzeri* show that they have similar respiratory complexes, suggesting that the regulation of these machineries, especially at the transcriptional level, is very important for adjusting rates of oxygen consumption in order to protect oxygen-sensitive processes. The transcriptional regulator CydR (Avin19910) seems to regulate various physiological processes associated with respiratory protection. Wu et al. (94) showed that CydR can coordinate an oxygen-labile [Fe-S] cluster, which provides a mechanism for CydR to sense subtle changes in the intercellular oxygen concentration and to regulate the expression of respiratory genes. During nitrogen fixation, CydR repression is presumably relieved, resulting in increased expression of uncoupled NADH dehydrogenase (Ndh) and CydAB I. A series of reports have suggested a role

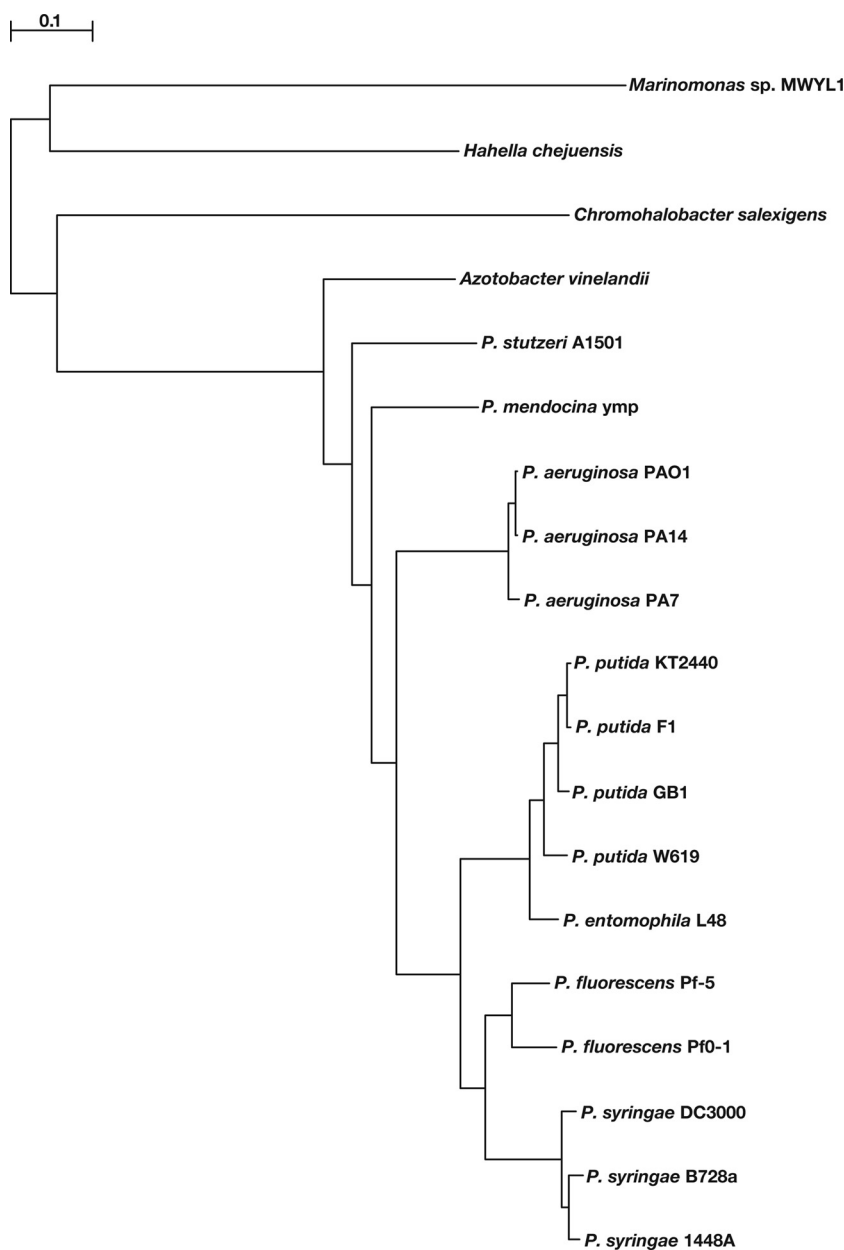


FIG. 1. Phylogenetic tree for *A. vinelandii*, all fully sequenced members of the *Pseudomonas* genus, and three outgroups. Accession numbers for the genomes used are given in Table S1 in the supplemental material. Bootstrap support for all branches was 100%.

for CydR in several other metabolic processes, including synthesis of PHB (95) and flagellar motility (48). Although the role of CydR in respiration is not completely defined, its direct or indirect participation in other cellular processes is even more obscure.

The genomic location of *cydR* (Avin19910) downstream of the *cco* region (containing the *cco* genes [Avin19920 to Avin20010]) suggests that it is involved in the regulation of this system. In support of this, a putative CydR binding site was identified upstream of the *cco* genes, and expression of a *ccoN::lacZ* fusion is CydR dependent (data not shown). Similar regulation of Cco by FNR (a CydR ortholog) is seen in other organisms (16). CydR also binds upstream of the CydAB

oxidase I promoter and is known to repress transcription of this operon (93). While *cydR* in *A. vinelandii* is adjacent to the *cydAB* operon (Avin19890 to Avin19880) encoding CydAB oxidase I, this organization is not seen in other organisms. Our promoter analysis has further identified a CydR binding site immediately upstream of Avin11170, suggesting that CydR also regulates the *cox* genes in *A. vinelandii*.

A putative CydR binding site is also located upstream of the iron-sulfur cluster biosynthetic genes (*isc* genes; Avin40380 to Avin40410). Cross regulation of Fe-S cluster biosynthesis and respiration makes biochemical sense for at least the following two reasons: (i) a large number of Fe-S proteins are involved in respiration, and (ii) elevated oxygen concentrations result in

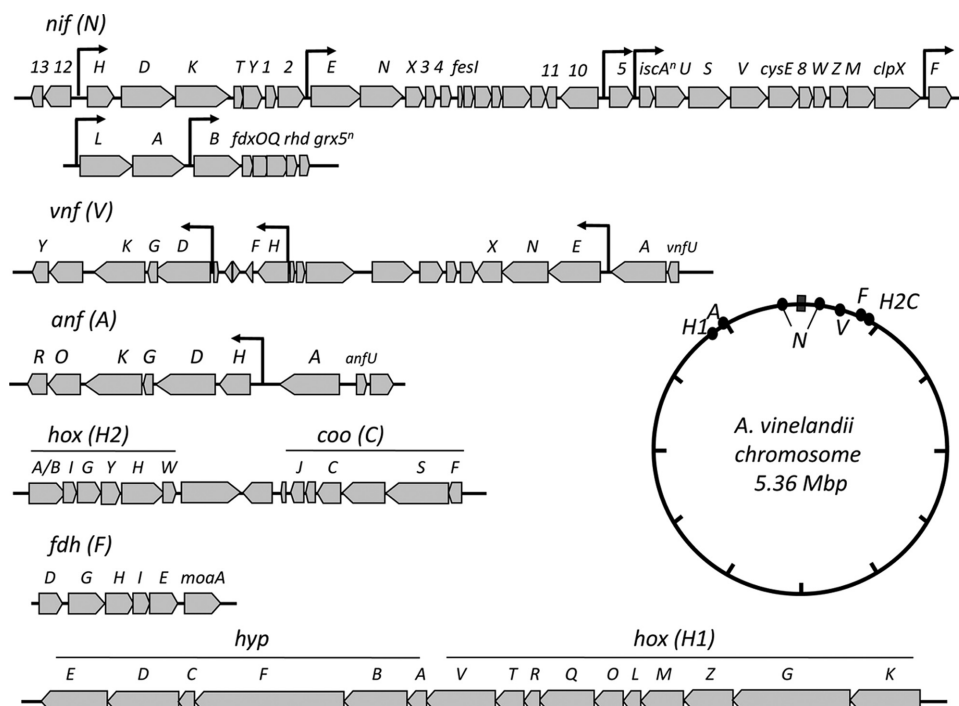


FIG. 2. Oxygen-sensitive protein-encoding genes in *A. vinelandii* and their genomic locations.

increased damage to Fe-S proteins under conditions in which the respiratory proteins are more highly expressed. Therefore, under conditions of high oxygen concentrations, CydR regulation would increase the expression of respiratory components and increase the capacity of the Fe-S cluster biosynthetic apparatus to supply the temporary higher demand for Fe-S clusters.

Nitrogen fixation. *A. vinelandii* expresses three oxygen-sensitive nitrogenase enzymes with different structural subunits and metal cofactor dependencies. Genes encoding the well-studied molybdenum-dependent nitrogenase and its assembly machinery and regulation (*nif*) are located in two regions of the chromosome, adjacent to and equidistant from the origin of replication. Proximity to the origin might result in a higher gene dosage during active growth, which can contribute to the high expression levels of the Nif components. The major *nif* region (comprising genes Avin01360 to Avin01710, oriented away from the putative origin, on the plus strand) encodes the structural subunits and the majority of the assembly machinery. The minor *nif* region (genes Avin50990 to Avin51060, also on the plus strand) contains the regulatory genes *nifL* and *nifA* and genes required for Mo trafficking and nitrogenase cofactor biosynthesis. Adjacent to the minor *nif* region are the *mfI* region (upstream) and genes for a putative rhodanese (*rdh_{nif}*; Avin51050) and a monothiol glutaredoxin (*grx_{nif}*; Avin51060) (downstream). The *mfI* gene products are associated with the accumulation of active nitrogenase component 2, also referred to as Fe protein (18). Likewise, the inactivation of the *nif*-associated glutaredoxin resulted in a 50% loss of Fe protein activity (data not shown). The dramatic effect on Fe protein activity and the less pronounced effect on nitrogenase component 1, the MoFe protein, could be attributed to an inherent

susceptibility to oxygen damage and/or deficiency in repair of the [Fe-S] cofactor in strains lacking these genes.

The genome sequence reveals that the region containing the structural genes for the vanadium nitrogenase (Avin02650 to Avin02660) is in close proximity to the gene cluster carrying genes (Avin02740 to Avin02780) involved in its assembly and regulation (Fig. 2). However, these two *vnf* clusters are separated by three genes predicted to participate in molybdopterin biosynthesis (Avin02700 to Avin02720) and a gene, *pcaK*, encoding a hydroxybenzoate transporter (Avin02690). In contrast, the *anf* genes, encoding the iron-only nitrogenase, are located in a single gene region (Avin48950 to Avin49000), which is regulated by the AnfA transcriptional activator (Avin49020) (Fig. 2).

Although no additional nitrogenase genes were identified in the genome sequence, genes with sequence similarity to the *nif*, *vnf*, and *anf* genes were found scattered throughout the genome. Whether or not these other genes are involved in nitrogen fixation remains to be determined. Interestingly, a gene with sequence similarity to *nifD* was identified (Avin39870), but its inactivation did not affect growth under standard diazotrophic or nondiazotrophic conditions (data not shown). Phylogenetic analysis of the NifD-like sequence in comparison with its paralogs in *A. vinelandii* and *P. stutzeri* reinforced the idea that the three structural and accessory nitrogenase genes were derived from a common ancestor and that the *nifD*-like sequence resembles the ancestral gene (see Fig. S6 in the supplemental material).

A paralog of *nifA*, Avin26490, was identified as encoding a protein that contains an almost identical DNA recognition helix in the C-terminal DNA-binding domain, implying that this protein binds similar enhancer sequences to those recog-

nized by NifA (Avin51000). A search for σ^{54} promoters and NifA upstream activator sequences detected the experimentally determined NifA-dependent promoters upstream of the *nif* and *mif* genes and additional putative NifA-dependent σ^{54} promoters upstream of genes encoding ModE (Avin50680), a putative molybdate binding protein, ModA3 (Avin50730), and a putative MbcC-like oxidoreductase (Avin48680). In view of the similarity in DNA recognition motifs, it is possible that these promoters are activated by either NifA or its paralog, Avin26490.

The genome sequence revealed two genes (Avin33440 and Avin47100) with 78% and 71% identity to *vnfA* (Avin02780), respectively. These VnfA paralogs have two of the three cysteines present in the proposed metal-binding cluster in the amino-terminal GAF domain of VnfA, similar to that of AnfA, which also has two cysteine residues implicated in metal or redox sensing (37, 69).

The *nfuA* gene (Avin28760) encodes a protein with a NifU-like C-terminal domain that shares sequence similarity with the products of *vnfU* (Avin02790) and *anfU* (Avin49030), which are located adjacent to *vnfA* (Avin02780) and *anfA* (Avin49020), respectively. Biochemical experiments suggest that NfuA represents an intermediate [Fe-S] cluster carrier involved in [Fe-S] protein maturation (2). It is likely that VnfU and AnfU play a role in the maturation of clusters present in vanadium- and iron-only nitrogenases.

In addition to paralogs of nitrogen fixation genes, other systems critical to nitrogen fixation appear to have been duplicated. Positioned 35 kb upstream of the minor *nif* gene cluster is the high-affinity molybdate transport system, encoded by *modG* and *modEA1B1C1* (Avin50650 to Avin50690), which supports the expression of Mo-dependent nitrogenase under molybdenum-limiting conditions (65). The second known molybdate transport locus, *modA2B2C2* (Avin01280 to Avin01300), is located 10 kb upstream of the major *nif* gene cluster. Interestingly, we identified a third putative molybdate transport system (Avin50700 to Avin50730), located directly next to the *modEA1B1C1* operon.

Other oxygen-sensitive processes. (i) CODH. The genome of *A. vinelandii* DJ contains genes with sequence similarity to membrane-bound Ni-dependent anaerobic CODH genes. CODH is an α_2 homodimer of the *cooS* product (Avin04490) that carries out the reversible oxidation of CO to CO₂ (71). Electrons extracted from CO are transferred to CooF (Avin04500), a hydrophobic FeS protein similar to the electron transfer subunits of oxidoreductases, such as the FDH beta subunit FdhH (Avin03820).

CODH contains a NiFe₄S₄ center, known as the C cluster, at its active site. The assembly of the CODH active site, which has been well studied in *Rhodospirillum rubrum*, requires the activities of three proteins, encoded by the *cooJ*, *cooC*, and *cooT* genes (43), involved in Ni storage and insertion (36, 91). Whereas one gene similar to *cooC* is present in the *A. vinelandii* genome (Avin04470), the genome does not code for proteins similar to CooT or CooJ. Avin04460, located downstream of *cooC*, encodes a 73-amino-acid histidine-rich protein with sequence similarity to the N-terminal portion (first 20%) of HypB proteins in *R. rubrum* (E value, 10⁻⁷) and *Ralstonia eutropha* (10⁻⁵), which are involved in Ni processing for hydrogenase maturation. Thus, it is likely that the product of

Avin04460 substitutes for the role of CooJ in the maturation of CODH. There is a gene similar to *hupE/ureJ* genes (Avin04450) downstream of Avin04460. HupE/UreJ proteins are secondary Ni transporters, and their coding genes are widespread among bacteria, normally clustered with urease or hydrogenase genes. There is another gene in the *hupE/ureJ* family (Avin50400), located near the *hypEDCFB* gene cluster, that probably encodes a HupE protein. We hypothesize that Avin04450 and Avin50400 may encode two different Ni transport systems, specific for CODH and hydrogenase maturation, respectively.

The *A. vinelandii* *coo* gene cluster includes a gene for a flavin adenine dinucleotide-dependent pyridine nucleotide-disulfide oxidoreductase (Avin04480) inserted between the *cooS* and *cooC* genes. A similar protein is also encoded downstream of *cooFS* in *Carboxydotherrmus hydrogenoformans*. Proteins of this family utilize flavin adenine dinucleotide to shuttle electrons from NADH to a redox-active disulfide bridge. The product of Avin04480 might be involved in CODH maturation. It has been proposed that Ni insertion into the C cluster involves reversible Ni binding to an Fe₃S₄ center followed by coordination to a specific cysteinyl residue (Cys531 in *R. rubrum* CooS) (36). It is also noteworthy that in vitro Ni insertion into CODH requires dithionite as a reductant. Thus, the Avin04480 gene product could be involved in reduction of a cysteine residue in *A. vinelandii* apo-CODH to facilitate Ni insertion.

The genome sequence reveals the presence of a *cooA* gene (Avin47010), as previously reported (98). The *cooA* gene encodes a transcriptional activator distantly related to the CRP family. CooA is a heme-containing homodimeric protein that functions as a CO sensor and controls the expression of the *coo* genes (72). In *R. rubrum*, CooA activates transcription of two contiguous gene clusters, namely, one that encodes CODH and accessory proteins (*cooFSCTJ*) and another encoding a membrane-bound CO-tolerant NiFe hydrogenase and its accessory proteins (*cooMKLXUH*). Although *A. vinelandii* lacks the genes encoding the CO-tolerant hydrogenase, it contains a gene cluster encoding a chimerical soluble NiFe hydrogenase in the corresponding locus (see below).

(ii) FDH. FDHs combine heterogeneous groups of enzymes found in both prokaryotes and eukaryotes that catalyze the oxidation of formate to CO₂ and H⁺. In aerobic organisms, the FDHs are mostly NAD⁺-dependent FDHs. Many prokaryotes, however, thrive in anoxic environments, where FDHs are NAD⁺-independent enzymes containing a variety of redox centers with oxygen-sensitive active sites composed of molybdenum or tungsten cofactors. These organisms utilize formate (produced from pyruvate) as a main electron donor for a variety of inducible anaerobic respiratory pathways (49).

The *A. vinelandii* genome contains a gene cluster that encodes a NAD⁺-independent FDH (*fdhDGHIE*; genes Avin03800 to Avin03840). While best BLAST (1) hits of *fdhD* are against *Pseudomonas* spp., including *P. stutzeri* A1501 (E value, 10⁻⁸⁴), the catalytic subunit of FDH encoded by *fdhGHI* shows significant similarity to the alpha, beta, and gamma subunits of the well-studied nitrate-inducible Fdh-N enzyme from enterobacteria, including *E. coli* (85). Similarity E values against *E. coli* O157 were as follows: for *fdhG*, 0.0; for *fdhH*, 10⁻¹²⁹; and for *fdhI*, 10⁻⁴⁸. In the catalytic site of the alpha subunit (FdhG), the molybdopterin-guanine dinucleotide co-

factor extracts electrons from formate. These electrons are transferred first to the [FeS]-containing beta subunit (FdhH) and finally to the heme-containing gamma subunit (FdhI). Although in *E. coli* Fdh-N function is associated with the activity of a respiratory nitrate reductase complex (NarGHI) located in the inner membrane, *narGHI* genes were not found in the genome of *A. vinelandii*. Interestingly, FdnG from *Klebsiella pneumoniae* was suggested to participate in relieving NifL inhibition of NifA (33), since *K. pneumoniae* strains carrying null mutations of *fdnG* (or the NADH-ubiquinone oxidoreductase genes *nuoCD*) showed significantly reduced *nif* induction under nitrogen-fixing conditions. The *moaA* gene (Avin03850), which encodes a SAM-dependent radical enzyme involved in the first step of the biosynthesis of the molybdenum cofactor (34), is located directly downstream of the *fdhDGHIE* gene cluster. Avin30330 encodes another MoaA homolog.

(iii) Hydrogenases. Prokaryotes, mostly from anaerobic ecosystems, have the ability to use H₂ by employing uptake hydrogenases or to produce H₂ by the activity of H₂-evolving hydrogenases. About one thousand hydrogenase sequences have been identified, many by genome sequencing, and more than 100 have been characterized genetically and/or biochemically (89). Three phylogenetically distinct classes of hydrogenases have been described, namely, [NiFe] hydrogenases, [FeFe] hydrogenases, and [Fe] hydrogenases ([Fe-S] cluster-free hydrogenases) (89, 90). Hydrogenases are usually sensitive to oxygen. For example, while [FeFe] hydrogenases are irreversibly destroyed by oxygen, the catalytic function of [NiFe] hydrogenases is reversibly inactivated by oxygen, but their structural integrity remains stable (8). The *A. vinelandii* genome contains genes encoding two [NiFe] hydrogenases, in membrane-bound and water-soluble forms (see below). Genes predicted to encode [FeFe] or [Fe] hydrogenases were not found.

The membrane-bound [NiFe] hydrogenase of *A. vinelandii* has been characterized extensively at both the biochemical (44, 75) and genetic (15, 31, 56) levels. The structural genes for this enzyme (*hoxKG*) (56) are clustered together with genes coding for hydrogenase accessory proteins, *hoxKGZMLQORTV* (Avin50500 to Avin50590) and *hypABFCDE* (Avin50440 to Avin50480), encoding the components of the hydrogenase electron transport chain and proteins for the biosynthesis of the hydrogenase cofactors and hydrogenase maturation (15, 31, 56). Mutational analysis has shown the requirement of these genes for H₂ oxidation activity (15, 31, 56).

It has generally been assumed that the presence of an uptake hydrogenase in nitrogen-fixing bacteria provides the advantage of recycling H₂ produced by nitrogenase, thereby yielding extra reductant and/or ATP for N₂ reduction and also providing respiratory protection for nitrogenase. Although competitive fitness experiments under carbon-limited conditions supported that notion for the related strain *Azotobacter chroococcum* (97), no significant difference was observed in ATP accumulation or in H₂-dependent O₂ uptake in a *hoxKG*-deficient mutant of *A. vinelandii* (52). Thus, the physiological role of the uptake hydrogenase in N₂ fixation is not completely understood.

Close to *hypE* (Avin50440) there is a gene, Avin50400, which codes for a protein that belongs to the HupE/UreJ family of proteins involved in Ni transport or processing. Two adjacent

open reading frames (Avin50410 and Avin50420), encoding putative membrane proteins, are unique to the *A. vinelandii* genome. Most pseudomonads possess a single copy of this membrane protein-encoding gene, but 12 copies are predicted for the *A. vinelandii* genome (Avin00560, Avin00570, Avin01720, Avin04580, Avin28060, Avin31780, Avin32050, Avin33410, Avin35450, Avin38580, Avin50420, and Avin50410). Whether these genes encode previously unknown accessory proteins for the *A. vinelandii* uptake hydrogenase and/or display an *A. vinelandii*-specific trait needs to be addressed.

A soluble NAD⁺-reducing [NiFe] hydrogenase is predicted to be encoded by Avin04360 to Avin04410. This putative multisubunit enzyme shares characteristics with both the soluble sensing hydrogenase HoxFUYHWI from *Ralstonia eutropha* (9) and the soluble H₂-evolving, S⁰-reducing sulfhydrogenase ShyBCDA from *Pyrococcus furiosus* (54). There is weak but still significant similarity (E value, 10⁻⁴) between Avin04410 and *R. eutropha* *hoxW* (CAA63575), encoding the hydrogenase peptidase, and between Avin04370 and *hoxI* (NP_942732), encoding the NADPH-binding subunit (E value, 10⁻⁶) (9). These two *A. vinelandii* genes appear to be interspersed among the *P. furiosus* *shyBCDA*-like genes (Avin04360 and Avin04370 to Avin04390) for the hydrogenase structural genes and the accessory subunits encoded by *shyBC*. Thus, *A. vinelandii* would encode a putative soluble [NiFe] hydrogenase of chimerical nature among other previously characterized hydrogenases. A similar arrangement of genes to those of *A. vinelandii* is present in the genomes of other bacteria, such as the aerobic, nitrogen-fixing alphaproteobacterium *Beijerinckia indica*, the aerobic ammonia-oxidizing gammaproteobacterium *Nitrosococcus mobilis* Nb-231, and others, which have been uncovered by genome sequencing projects and represent the most similar relatives to the putative *A. vinelandii* soluble hydrogenase. The corresponding amino acid sequence identities to *A. vinelandii* *shyB-hoxI-shyCDA-hoxW*-like genes of these putative hydrogenase subunits are 66%, 75%, 63%, 73%, 76%, and 52% for *B. indica* and 58%, 60%, 65%, 71%, 65%, and 45% for *N. mobilis*, respectively. Interestingly, this gene cluster appears to have undergone extensive rearrangements during the evolution of hydrogenases. While this cluster is adjacent to the putative CODH gene cluster in *A. vinelandii* (Avin04450 to Avin04490), in *N. mobilis* it is adjacent to the *hypBAEDCF*-like genes for the maturation of the [NiFe] hydrogenase, and in *B. indica* they do not appear to be linked physically to hydrogen metabolism genes. Mutagenesis studies of these genes would shed light on hydrogen metabolism in *A. vinelandii* and other bacteria.

Alginate. There are two possible strategies for keeping the cytoplasm anaerobic: either the bacterium can remove the oxygen after it has entered the cell, or it can create a barrier to impede O₂ transfer into the cell. Sabra et al. showed that the alginate capsule of *A. vinelandii* was affected by the oxygen tension (73). Alginate is a linear copolymer of 1→4-linked β-D-mannuronic acid and α-L-guluronic acid, where some of the mannuronic acid residues may be acetylated. The alginate biosynthetic gene set consists of 12 genes (Avin10860 to Avin10970). The physical organization of this cluster is highly conserved in *A. vinelandii* and *Pseudomonas* species able to produce alginate, including *Pseudomonas aeruginosa*, *Pseudomonas fluorescens*, *Pseudomonas syringae*, *Pseudomonas putida*, *Pseudomonas entomophila*, and *P. mendocina* (see Fig. S7 in

the supplemental material). In addition, *A. vinelandii* encodes a set of seven secreted mannuronan C-5 epimerases (AlgE1-7) that modify the polymer outside the cells (*algE1*, Avin51190; *algE2*, Avin51180; *algE3*, Avin51170; *algE4*, Avin51200; *algE5*, Avin33710; *algE6*, Avin51230; and *algE7*, Avin51250). Some of these epimerases are responsible for introducing consecutive guluronic acid residues into the polymer, and this structural feature is necessary for forming a gel with divalent cations. Close homologs of the epimerase genes *algE1-7* are not described for other species except for *P. syringae*, which encodes a single bifunctional alginate epimerase/deacetylase, PsmE (7).

Wild-type strains of *Pseudomonas* usually do not produce alginate under laboratory conditions, while most strains of *A. vinelandii* produce alginate constitutively and use the polymer both for making the vegetative state capsule and for the cyst coat (see below). This probably explains why the regulation of alginate synthesis in *A. vinelandii* differs from that in *Pseudomonas*. Alginate production in *P. aeruginosa* is controlled by a complex regulatory system that includes AlgR, AlgB, AlgQ, AmrZ, and AlgP (30, 59). Orthologs of the genes encoding these regulators are present in the genome of *A. vinelandii* DJ, as follows: *algR* (Avin47610), *algB* (Avin11120), *algP* (Avin47540), *algQ* (Avin47550), and *amrZ* (Avin34410). While in *Pseudomonas* AlgR is required for alginate production, *algR* mutants of *A. vinelandii* produce alginate but are unable to form mature cysts (60). The participation of AlgB, AlgP, AlgQ, and AmrZ in the regulation of alginate synthesis has not been studied in *A. vinelandii*. Orthologs of these genes are present, however, in *P. stutzeri* A1501, which does not produce alginate.

Other regulators of alginate synthesis, including the *algU-mucABCD* operon (Avin13660 to Avin13730), the protease genes *algW* (Avin12950), *mucP* (Avin38920), and *prc* (Avin35170), and the diguanylate cyclase gene *mucR* (Avin49140), are also conserved in *A. vinelandii* and *Pseudomonas* species. *A. vinelandii* does not carry any close homolog to *mucE*, which in *P. aeruginosa* is necessary for the proteolytic events that degrade the anti-sigma factor MucA enabling alginate production (70). Strain DJ does not produce alginate due to the presence of an insertion sequence that splits *algU* (Avin13660) in two (55).

Polymer production and encystment. A distinctive characteristic of *A. vinelandii* is the formation of a desiccation-resistant cell, described as a cyst, upon encountering adverse growth conditions or upon induction of vegetative cells with specific reagents. Only a few bacterial species have been observed to make cysts. Initial studies of cyst formation in *Rhodospirillum centenum* and *Azospirillum brasiliense* identified a few genes (4, 88) that are involved in the regulation of encystment in these bacteria. Genes similar to these regulators, however, have not been found in the *A. vinelandii* genome. Alginate, in addition to its role in respiratory protection, is a major component of the cyst capsule, and alginate with consecutive guluronic residues is essential for the formation of mature cysts (12, 80). During encystment, *A. vinelandii* was shown to incorporate phenolic lipids (alkylresorcinols and alkylpyrones) into the membrane. The proteins involved in the biosynthesis of these lipids have been characterized in *A. vinelandii* (29), and it has been shown that these lipids play a structural role in the cyst capsule, although they are not essential for desiccation resistance (78).

Even though *A. vinelandii* seems unable to utilize alginate as

an energy source, it carries at least six enzymes with alginate lyase activities, including AlyA1-3 (Avin31810, Avin23960, and Avin13810), AlgE7 (Avin51250), Avin46500 (82; our unpublished data), and AlgL (Avin10900), the alginate lyase encoded in the alginate biosynthetic gene cluster that is also present in *Pseudomonas* species. One of these, AlyA3, is required for cyst germination, probably helping the germinating cells to escape from the cyst coat by depolymerizing the alginate (H. Ertesvåg, personal communication). In contrast, AlgL is not required for cyst germination (86).

Abundant PHB granules accumulate during encystment and are a major component of the central body of the cyst. Conditions or strains favoring greater polymer accumulation also produce more mature cysts (81), although PHB synthesis is not essential for encystment (76). The PHB biosynthetic genes *phbB* (Avin23650), *phbA* (Avin23640), and *phaC* (Avin23630), the regulatory genes *phbR* (Avin23660) and *phbF* (Avin23680), and the phasin gene *phbP* (Avin23670) were previously identified (66, 76, 77). The biosynthetic genes seem to be under *CydR* control, since the PhbB and PhbA proteins are increased in a *cydR* mutant (94) and a putative *CydR* binding site is present upstream of *phbB* (66). Genome analysis revealed additional genes probably involved in PHB metabolism, including genes similar to *phbB* and *phbA* and genes encoding putative phasins, PHB synthases, PHB depolymerases, and PHB oligomer hydrolases (see Table S6 in the supplemental material).

PHB is the polyhydroxyalkanoate (PHA) usually produced by *A. vinelandii*. However, the addition of valerate, heptanoate, or nonanoate to a culture of *A. vinelandii* UWD grown in glucose allows the synthesis of a copolymer, poly-(hydroxybutyrate-cohydroxyvalerate) (PHB-co-HV) (63). In the pseudomonads belonging to rRNA homology group I, an (*R*)-specific enoyl-coenzyme A hydratase, the product of the *phaJ* gene, is responsible for the channeling of β -oxidation intermediates to PHA synthesis (28). In accordance with the PHB-co-HV synthetic capacity of *A. vinelandii*, a *phaJ* gene (Avin30160) is present in its genome.

DISCUSSION

While *P. stutzeri* is the closest relative to *A. vinelandii* among fully sequenced prokaryotes, there are marked physiological and metabolic differences between these organisms. Unlike *A. vinelandii*, *P. stutzeri* can fix nitrogen only under microaerobic conditions, but in contrast to *A. vinelandii*, this organism can grow under anaerobic conditions by utilizing nitrate as a terminal electron acceptor. The classical high rates of respiration exhibited by *A. vinelandii* are apparently bestowed by the provision of five terminal oxidases (Fig. 3), supported by a number of NADH- and other substrate-driven respiratory chains. Unexpectedly, the genome encodes two cytochrome *bd*-type terminal oxidases. One of these, *CydAB I*, is the well-studied low-affinity *bd* oxidase known to be involved in respiratory protection of nitrogenase (68). This terminal oxidase appears to be essential for aerotolerant nitrogen fixation in *A. vinelandii*. The second *bd*-type terminal oxidase, *CydAB II*, resembles the cyanide-tolerant *Cio* terminal oxidases, which differ in their heme complement from the canonical cytochrome *bd* (17).

The distinctive ability of *A. vinelandii* to carry out nitrogen fixation under aerobic conditions is subject to regulation by the

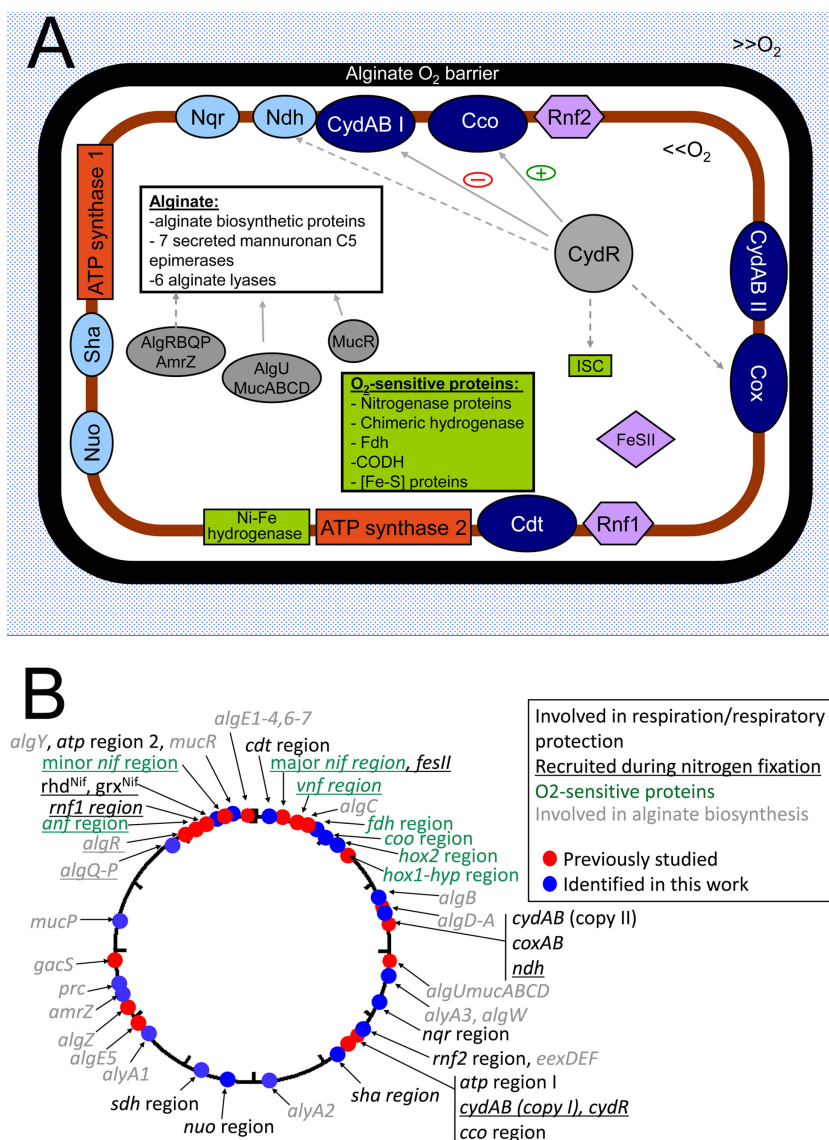


FIG. 3. Respiratory protection in *Azotobacter vinelandii*. (A) Illustration of *A. vinelandii* cell and the key proteins involved in respiratory protection. A number of alginate biosynthetic proteins are produced that direct the formation of an alginate coating around the cell, acting as a physical O₂ barrier (black lining). Respiratory proteins, including five terminal oxidases (dark blue), four NADH-ubiquinone oxidoreductases (light blue), two ATP synthases (orange), and other proteins involved in respiratory protection (lavender), provide the second line of defense against O₂. Proteins sensitive to O₂ exposure are shown in green boxes. Solid line arrows represent regulatory events for which experimental evidence exists. Dashed line arrows represent putative regulation on the basis of promoter binding regions. Regulatory proteins are depicted in gray. (B) Locations of respiratory and respiratory protection genes in the *A. vinelandii* genome. Proteins are color coded according to their role in respiration, respiratory protection, and nitrogen fixation. Genes whose protein products have been identified previously are represented by red dots. The genes identified in this work (represented by blue dots) encode the following proteins: cytochrome *o* oxidase (Avin11160 to Avin11180), Rnf2 complex (Avin19220 to Avin19270), Sdh complex (Avin29780 to Avin29810), Sha complex (Avin19530 to Avin19580), Nuo complex (Avin28440 to Avin28560), Nqr complex (Avin14590 to Avin14640), Rhd^{Nif} (Avin51050), Grx^{Nif} (Avin51060), cytochrome *c* oxidase (Avin00940 to Avin01020), chimeric soluble Ni-Fe hydrogenase (Avin04360 to Avin04410), Fdh (Avin03820), CODH (Avin04450 to Avin04490), ATP synthase 2 complex (Avin52150 to Avin52230), AlgB (Avin11120), AlgQ (Avin47550), AmrZ (Avin34410), AlgP (Avin47540), AlgW (Avin12950), MucP (Avin38920), Prc (Avin35170), AlyA1-3 (Avin13810, Avin23960, and Avin31810), AlgY (Avin49400), and the alginate lyase (Avin 46500).

oxygen-responsive transcriptional regulator CydR, which like Fnr contains a [4Fe-4S]²⁺ cluster and negatively regulates expression of the *cydAB* genes that encode the low-affinity CydAB I enzyme (93). CydR also appears to control expression of the uncoupled NADH-ubiquinol dehydrogenase (Ndh) (6), which is thought to supply electrons to CydAB I oxidase and is also essential for aerotolerant nitrogen fixation (5). Our stud-

ies suggest that CydR activates expression of Cco, which may explain why CydR is required for growth under microaerobic conditions (94). Overall, CydR may have a role analogous to that of Anr in controlling the expression of terminal oxidases in *Pseudomonas* (87), but its involvement in the regulation of PHB metabolism (95) implies a wider role in the integration of carbon source and oxygen availability (Fig. 3).

Mechanisms for respiratory protection are complemented by the barrier to oxygen diffusion into the cell provided by the alginate capsule, which appears to play a major role in protecting nitrogenase from oxygen damage, particularly under phosphate-limiting conditions (73). It is noteworthy that alginate biosynthesis is constitutive in *A. vinelandii* and that a large number of secreted mannuronan C-5 epimerases are encoded in the genome by *algE1-7*. These enzymes may be required to increase the L-guluronic acid content of the alginate when the organism is grown at high oxygen tensions (73). Some of the six alginate lyase enzymes encoded by the genome may be responsible for altering the composition of alginate in response to oxygen availability (Fig. 3). *P. stutzeri* does not possess genes for alginate biosynthesis and therefore is not able to take advantage of this mechanism for reducing oxygen diffusion into the cell.

In addition to the oxygen diffusion barrier and respiratory removal of oxygen at the cell surface, a number of other strategies may be used by *A. vinelandii* to protect oxygen-sensitive enzymes in the cytoplasm. These include maintenance of a low redox state and efficient energy metabolism (61). In the latter context, it is interesting that *A. vinelandii* contains a second operon encoding an F-type ATP synthase (Avin19670 to Avin19750) that is not closely related to the ancestral operon, with its closest relative being present in the genus *Burkholderia*. The redundancy of oxygen protection mechanisms may explain why *A. vinelandii* DJ is able to fix nitrogen under standard atmospheric conditions in spite of its inability to produce alginate.

The *A. vinelandii* genome encodes several unexpected anaerobic enzymes in addition to the three well-characterized oxygen-sensitive nitrogenases and the membrane-bound [NiFe] hydrogenase (Fig. 3). The presence of NAD⁺-independent FDH is unusual for a strict aerobe that does not possess a respiratory nitrate reductase. Even more intriguing is the presence of genes predicted to encode a Ni-dependent CODH, although a homolog of the heme-containing CO sensor, CooA, has been found previously (98). These proteins are characteristic of anaerobic CO metabolism and contrast with the molybdopterin-containing CODHs found in aerobes (71). CO oxidation by *A. vinelandii* has not been reported previously, although it may be an important detoxification process given the CO sensitivity of hydrogenase and respiratory enzymes. The electron acceptor for CO oxidation could potentially be provided by the gene cluster encoding the soluble chimeric NiFe hydrogenase (Avin04360 to Avin04410), which intriguingly is positioned in the corresponding locus to that of the membrane-bound CO-tolerant NiFe hydrogenase in *R. rubrum*. It has been suggested that this energy-conserving hydrogenase produces a proton gradient to drive ATP synthesis by coupling CO-dependent H₂ production and proton translocation in *R. rubrum* (26). Clearly, the role of CO in *A. vinelandii* metabolism requires further investigation.

The ability of *A. vinelandii* to reconcile an obligate aerobic lifestyle with the maintenance of fundamental oxygen-sensitive processes, such as nitrogen fixation, is a remarkable metabolic accomplishment that has important implications for biotechnological exploitation. *A. vinelandii* is a model organism for biochemical studies on the basis of the high yield and quality of enzymes that can be prepared from it. Moreover, powerful

genetic approaches facilitated by homologous recombination are readily available, as are a stringent system for controlled expression of proteins encoded within the genome (38) and a means to achieve high-level protein expression (19). Given the high quality of proteins purified from *A. vinelandii*, these combined genetic and biochemical tools make this an ideal organism for the production of enzymes, particularly those that are oxygen sensitive.

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This report is dedicated to the memory of Christina Kennedy (1945–2009). Christina made numerous contributions to our understanding of nitrogen fixation and *Azotobacter* biology over her nearly 4-decade-long career, culminating in the publication of its genome.

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