

# Rex (Encoded by DVU\_0916) in *Desulfovibrio vulgaris* Hildenborough Is a Repressor of Sulfate Adenylyl Transferase and Is Regulated by NADH

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Although the enzymes for dissimilatory sulfate reduction by microbes have been studied, the mechanisms for transcriptional regulation of the encoding genes remain unknown. In a number of bacteria the transcriptional regulator Rex has been shown to play a key role as a repressor of genes producing proteins involved in energy conversion. In the model sulfate-reducing microbe *Desulfovibrio vulgaris* Hildenborough, the gene DVU\_0916 was observed to resemble other known Rex proteins. Therefore, the DVU\_0916 protein has been predicted to be a transcriptional repressor of genes encoding proteins that function in the process of sulfate reduction in *D. vulgaris* Hildenborough. Examination of the deduced DVU\_0916 protein identified two domains, one a winged helix DNA-binding domain common for transcription factors, and the other a Rossman fold that could potentially interact with pyridine nucleotides. A deletion of the putative *rex* gene was made in *D. vulgaris* Hildenborough, and transcript expression studies of *sat*, encoding sulfate adenylyl transferase, showed increased levels in the *D. vulgaris* Hildenborough Rex (Rex<sub>DvH</sub>) mutant relative to the parental strain. The Rex<sub>DvH</sub>-binding site upstream of *sat* was identified, confirming Rex<sub>DvH</sub> to be a repressor of *sat*. We established *in vitro* that the presence of elevated NADH disrupted the interaction between Rex<sub>DvH</sub> and DNA. Examination of the 5' transcriptional start site for the *sat* mRNA revealed two unique start sites, one for respiring cells that correlated with the Rex<sub>DvH</sub>-binding site and a second for fermenting cells. Collectively, these data support the role of Rex<sub>DvH</sub> as a transcription repressor for *sat* that senses the redox status of the cell.

The anaerobic process of microbially influenced corrosion is estimated to account for 15% of the total cost of ferrous metal and concrete/stonework corrosion. For the U.S. energy industry, this amounts to approximately \$100 billion yearly (1). Most notably, the metabolic processes of the sulfate-reducing microbes (SRM) have been implicated in this observed corrosion (2, 3). Clearly, it would be beneficial to understand the mechanism and the regulation of genes involved in the key processes leading to metal dissolution.

The SRM convert energy by dissimilatory sulfate reduction, where sulfate is used as a terminal electron acceptor in respiration. Although SRM are mainly found in sulfate-rich anoxic environments (4), they are also found in anoxic habitats depleted of sulfate because they are able to use electron acceptors other than sulfate (5, 6). *Desulfovibrio vulgaris* Hildenborough, a Gram-negative delta-proteobacterium, is a model organism for examining sulfate reduction as it is genetically accessible and is readily cultured in the lab (7, 8). Also, the genome for *D. vulgaris* Hildenborough has been sequenced (9) and has recently been reannotated (10).

The process of dissimilatory sulfate reduction is carried out by a well-conserved biochemical pathway in heterogeneous SRM (5, 11–13). In brief, sulfate (SO<sub>4</sub><sup>2-</sup>) is taken up by the cell and is activated by sulfate adenylyl transferase (Sat, encoded by *sat*) to adenylyl phosphosulfate (APS) and inorganic pyrophosphate. APS is then reduced to sulfite (SO<sub>3</sub><sup>2-</sup>) and AMP by APS reductase (ApsBA, encoded by *apsBA*). Sulfite is further reduced to hydrogen sulfide (H<sub>2</sub>S) by dissimilatory sulfite reductase (DsrAB, encoded by *dsrAB*). Although enzymes for sulfate reduction have

been well studied (5, 12, 13) and their crystal structures have been solved (14–17), the mechanism for transcriptional regulation of the encoding genes remains unknown.

In *D. vulgaris* Hildenborough, genes involved in sulfate reduction are differentially expressed depending on the available nutrients (<http://www.microbesonline.org/>) (18). For example, *apsBA* expression was decreased in medium containing sulfite compared to expression in sulfate medium (19). From the genome sequence of *D. vulgaris* Hildenborough, more than 150 transcriptional regulators have been predicted, several of which have the potential to be responsible for the observed expression changes (<http://networks.systemsbio.net/dvh/search/advanced>) (20). DVU\_0916 was predicted to encode a Rex protein and was hypothesized to be involved with sulfate

Received 11 July 2014 Accepted 1 October 2014

Accepted manuscript posted online 13 October 2014

Citation Christensen GA, Zane GM, Kazakov AE, Li X, Rodionov DA, Novichkov PS, Dubchak I, Arkin AP, Wall JD. 2015. Rex (encoded by DVU\_0916) in *Desulfovibrio vulgaris* Hildenborough is a repressor of sulfate adenylyl transferase and is regulated by NADH. *J Bacteriol* 197:29–39. doi:10.1128/JB.02083-14.

Editor: W. W. Metcalf

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Supplemental material for this article may be found at <http://dx.doi.org/10.1128/JB.02083-14>.

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reduction by regulating more than 50 genes that are responsible for energy conversion processes (21). From an examination of genome sequences, Rex has been predicted to be present in a wide range of microbes, including Gram-negative and Gram-positive aerobes and anaerobes. To date, Rex proteins have been studied experimentally in several aerobes, including *Thermus aquaticus* (22–24), *Streptomyces coelicolor* (25), and *Bacillus subtilis* (24, 26–28), and the anaerobe *Desulfovibrio alaskensis* G20 (30). The present study will add to our understanding of the function of Rex in anaerobic SRM.

Functional Rex proteins contain an N-terminal DNA-binding domain, a dimerization domain, and a Rossmann fold (22, 24). The latter apparently functions to bind pyridine nucleotides (NADH and NAD<sup>+</sup>). The consensus binding sequence for Rex, TTTGTGAAATATTTTCACAAA, has been compiled from comparisons of more than 100 genomes (21). The sequence contains an inverted repeat (underlined), since Rex functions as a homodimer, and the inverted repeat is the minimum sequence for Rex binding, as observed in *S. coelicolor* (25).

Targets for Rex typically include genes encoding proteins involved in NADH oxidation, and Rex acts as a transcriptional repressor when the NADH/NAD<sup>+</sup> ratio is low (21, 26). Rex from *T. aquaticus*, expressed in *Escherichia coli*, was crystallized in the presence of NADH and NAD<sup>+</sup> separately, and the structures were solved (22, 24). From these *in vitro* assays it was clear that Rex underwent structural changes that modulated the binding activity between Rex and a consensus DNA sequence. Specifically, when Rex bound NADH, the resulting conformation of Rex was no longer able to interact within the major groove of DNA and therefore would no longer repress (24). These results were consistent with *in vitro* protein-DNA interaction assays performed on an upstream DNA sequence of *adhE2* in *Clostridium acetobutylicum*. In these assays, increased expression of *adhE2* was observed in a *rex* deletion strain (31). Examination of expression data in an SRM closely related to *D. vulgaris* Hildenborough, *D. alaskensis* G20, revealed increased transcript expression for *sat* in a *rex* deleted strain; however, minimal differences were observed for *apsBA* and *dsrABD* (30). The increase in expression for *sat*, encoding the first enzyme in the process of sulfate reduction, but not *apsBA* and *dsrABD* provides support for the role of Rex as a transcriptional repressor for early steps in sulfate reduction. Interestingly, *rnfC*, which is predicted to be within the Rex regulon, was decreased in the Rex mutant, which was interpreted to mean that Rex acts not only as a repressor but as an activator as well.

The properties described for Rex (i.e., a regulator that contains genes that encode proteins responsible for energy conversion and a regulator that senses the NADH/NAD<sup>+</sup> ratio) would be those expected for a regulator of genes that encode proteins that function in sulfate reduction in *D. vulgaris* Hildenborough, since sulfate reduction is the primary mechanism for energy conversion in *D. vulgaris* Hildenborough. Therefore, a Rex homolog in *D. vulgaris* Hildenborough (Rex<sub>DvH</sub>) would be a reasonable candidate regulator. By differentially binding NAD<sup>+</sup> and NADH and repressing genes for NADH oxidation when NADH is low, Rex may maintain the NADH/NAD<sup>+</sup> ratio (26). Using this paradigm, we proposed that Rex might be involved in the switch between respiration and fermentation in SRM and be necessary for adaptation to fluctuating electron acceptor concentrations.

We sought to confirm that the protein encoded by DVU\_0916 was a functional Rex homolog in *D. vulgaris* Hildenborough. To

accomplish this, the growth kinetics of a deletion strain for *rex* were determined, and transcript differences for *sat*—encoding the enzyme that activates sulfate for the first step in its reduction—were analyzed. All results were consistent with the conclusion that Rex<sub>DvH</sub> is a redox-responsive transcriptional repressor for *sat*.

## MATERIALS AND METHODS

**Culture, media, and sample collection.** The growth of all cultures was measured in Balch tubes by determining the optical density at 600 nm (OD<sub>600</sub>) with a Spectronic Genesys 20 spectrophotometer (Thermo Fisher Scientific, Waltham, MA). *E. coli* strains were grown at 37°C aerobically in 5 ml of LC medium containing (per liter) 10 g of tryptone, 5 g of sodium chloride, and 5 g of yeast extract (pH 7.0). *D. vulgaris* Hildenborough strains were started from freezer stocks that contained 10% (vol/vol) glycerol in growth medium and frozen at –80 °C. Growth medium was generally MO medium (19) supplemented with 0.1% (wt/vol) yeast extract and 60 mM lactate with 30 mM sulfate (MOYLS4). The medium was reduced with 1.2 mM sodium thioglycolate, and the pH was adjusted to pH 7.2 with 12 M HCl. The inhibitors for *D. vulgaris* Hildenborough strains were as follows: Geneticin (G418; 400 µg/ml), spectinomycin (Sp; 100 µg/ml), kanamycin (Km; 50 µg/ml), ampicillin (100 µg/ml) or 5-fluorouracil (5-FU; 40 µg/ml) obtained from Thermo Fisher Scientific, Sigma-Aldrich (St. Louis, MO), or Gold Biotechnologies (St. Louis, MO). *D. vulgaris* Hildenborough cultures were grown in an anaerobic chamber (Coy Laboratory Product, Inc., Grass Lake, MI) overnight until reaching the stationary phase (OD<sub>600</sub> > 1). The atmosphere of the chamber was ca. 95% N<sub>2</sub> and 5% H<sub>2</sub>. A 2% (vol/vol) inoculum was used to start 5-ml cultures in defined MOLS4 (MOYLS4 lacking yeast extract) or pyruvate fermentation medium (MO medium with 60 mM pyruvate, supplemented with 0.1% [wt/vol] yeast extract and 0.5 mM cysteine [MOY<sub>2</sub>Pyr]). For pyruvate-fermenting cultures, cysteine was provided as a sulfur source and reductant. *D. vulgaris* Hildenborough did not grow on yeast extract alone in the absence of pyruvate (data not shown). For mRNA studies, 4-ml samples were spun down anaerobically at 34°C (10 min at 5,600 × g) and resuspended in 1 ml of TRI reagent (Sigma-Aldrich).

Genomic DNA (Wizard Genomic DNA purification kit; Promega, Madison, WI), plasmid (GeneJET plasmid kit; Thermo Fisher Scientific), and DNA fragments (Wizard SV Gel and PCR Clean-Up System [Promega]) were purified according to the manufacturer's protocol. Sequence analysis was performed by University of Missouri DNA Core Facility. Oligonucleotides necessary for these studies were purchased from Integrated DNA Technologies (Coralville, IA) (see Table S1 in the supplemental material). The DNA and RNA concentrations (and A<sub>260</sub>/A<sub>280</sub> ratio) were calculated from NanoDrop ND-1000 spectrophotometric readings (Thermo Fisher Scientific).

**Strain and plasmid constructions.** All of the strains used in the present study are listed in Table 1. All plasmid constructs were made by the SLIC procedure (32). The protocol used to generate the strain lacking the *sat* promoter (ΔP<sub>sat</sub>), JW9293, conferring kanamycin resistance and 5-FU sensitivity, has previously been described (33). Briefly, 150 bp upstream of *sat* (–150 to –1) was replaced with a cassette conferring resistance to kanamycin and a counterselectable marker (*upp*, the gene encoding uracil phosphoribosyltransferase that confers 5-FU sensitivity to the 5-FU<sup>r</sup> parental strain). To accomplish the cassette insertion, a delivery plasmid (pMO9292) was constructed that contained the selectable cassette flanked by chromosomal regions from either side of the promoter sequence to be deleted. This plasmid was electroporated into the parental strain, JW710, and selected for kanamycin resistance and screened for 5-FU sensitivity. The successful recombinant was designated JW9293. For the construction of the subsequent P<sub>sat</sub> mutants with alterations to the Rex<sub>DvH</sub>-binding site (JW9312, JW9314, JW9316, JW9318, and JW9320), a protocol similar to that described by Parks et al. (34) was followed. JW9293 was transformed with nonreplicating plasmids with individual site-specific mutations in the 150-bp fragment flanked by DNA homologous to that on either side of the integrated cassette. These P<sub>sat</sub> mutant or restored strains were selected

TABLE 1 Strains and plasmids used in this study

Strain or plasmid	Genotype and/or relevant characteristics <sup>a</sup>	Source or reference
<b>Strains</b>		
<i>E. coli</i>		
α-Select (Silver Efficiency) BL21(DE3) competent cells	<i>deoR endA1 recA1 relA1 gyrA96 hsdR17</i> (r <sub>K</sub> <sup>-</sup> m <sub>K</sub> <sup>+</sup> ) <i>supE44 thi-1</i> Δ( <i>lacZYA-argFV169</i> ) φ80 <i>dlacZ</i> ΔM15 F <sup>-</sup>	Bioline Agilent
<i>D. vulgaris</i> Hildenborough		
ATCC 29579	Wild type; 5-FU <sup>s</sup>	ATCC
JW710	WT Δ <i>upp</i> ; 5-FU <sup>r</sup> (used as a WT control for <i>D. vulgaris</i> Hildenborough growth kinetics in this study; parent strain for deletions; retains pDV1 present in WT)	8
JW3311	JW710 ΔDVU_0916::( <i>npt upp</i> ); Km <sup>r</sup> ; 5-FU <sup>s</sup> (Δ <i>rex</i> marker exchange)	33
JW9293	JW710 Δ <sub>-150-1</sub> P <sub><i>sat</i></sub> ::( <i>npt upp</i> ); Km <sup>r</sup> ; 5-FU <sup>s</sup> (P <sub><i>sat</i></sub> disruption)	This study
JW9312	JW710; Km <sup>s</sup> ; 5-FU <sup>r</sup> ( <i>sat</i> promoter restored)	This study
JW9314	JW9293 G <sub>-147</sub> A P <sub><i>sat</i></sub> ; Km <sup>s</sup> ; 5-FU <sup>r</sup> (G -147 A)	This study
JW9316	JW9293 GTA <sub>-147-145</sub> ACG P <sub><i>sat</i></sub> ; Km <sup>s</sup> ; 5-FU <sup>r</sup> (IR1)	This study
JW9318	JW9293 CAC <sub>-136-134</sub> TGT P <sub><i>sat</i></sub> ; Km <sup>s</sup> ; 5-FU <sup>r</sup> (IR2)	This study
JW9320	JW9293 GTA <sub>-147-145</sub> ACG P <sub><i>sat</i></sub> CAC <sub>-136-134</sub> TGT P <sub><i>sat</i></sub> ; Km <sup>s</sup> ; 5-FU <sup>r</sup> (IR1and2)	This study
JW9011	JW710 ΔDVU_2547::( <i>npt upp</i> ); Km <sup>r</sup> ; 5-FU <sup>s</sup> (Δ <i>hcpR</i> marker exchange)	50
GZ0481	Genome position 2680507::Tn5-RL27; insertion 273 bp from predicted AUG start codon within DVU_2567; Km <sup>r</sup> (LysX mutant)	Wall laboratory
<b>Plasmids</b>		
pET14b	6×His tag fusion protein vector with T7 promoter	Novagen
pMO719	pCR8/GW/TOPO containing SRB replicon (pBG1); Sp <sup>r</sup> ; source of Sp <sup>r</sup> and pUC <i>ori</i> fragment; for marker exchange suicide plasmid construction	8
pMO746	Source of <i>upp</i> in artificial operon with <i>npt</i> and Ap <sup>r</sup> -pUC <i>ori</i> ; P <sub><i>npt-npt-upp</i></sub> ; Km <sup>r</sup> ; 5-FU <sup>s</sup> ; for marker exchange suicide plasmid construction	34
pMO3312	pET14b plus <i>rex</i> (without start codon, 642 bp); Ap <sup>r</sup> ; for Rex expression in BL21(DE3) competent cells	This study
pMO3313	pMO9075 with DVU_0916 ( <i>rex</i> ) constitutively expressed from P <sub><i>npt</i></sub>	33
pMO9075	pMO719 containing P <sub><i>npt</i></sub> for constitutive expression of complementation constructs; pBG1 stable SRB replicon; Sp <sup>r</sup>	7
pMO9292	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 383-bp upstream and 319-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150-1) flanking the artificial operon of P <sub><i>npt-npt-upp</i></sub> from pMO746; for marker exchange mutagenesis; Sp <sup>r</sup> and Km <sup>r</sup>	This study
pMO9311	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 403-bp upstream and 467-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150); wild-type sequence; Sp <sup>r</sup> ; for site-directed mutagenesis	This study
pMO9313	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 403-bp upstream and 467-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150); G <sub>-147</sub> A P <sub><i>sat</i></sub> ; Sp <sup>r</sup> ; for site-directed mutagenesis	This study
pMO9315	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 403-bp upstream and 467-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150); GTA <sub>-147-145</sub> ACG P <sub><i>sat</i></sub> ; Sp <sup>r</sup> ; for site-directed mutagenesis	This study
pMO9317	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 403-bp upstream and 467-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150); CAC <sub>-136-134</sub> TGT P <sub><i>sat</i></sub> ; Sp <sup>r</sup> ; for site-directed mutagenesis	This study
pMO9319	Sp <sup>r</sup> and pUC <i>ori</i> from pMO719 plus 403-bp upstream and 467-bp downstream DNA regions from P <sub><i>sat</i></sub> (-150); GTA <sub>-147-145</sub> ACG P <sub><i>sat</i></sub> CAC <sub>-136-134</sub> TGT P <sub><i>sat</i></sub> ; Sp <sup>r</sup> ; for site-directed mutagenesis	This study

<sup>a</sup> Km, kanamycin; Sp, spectinomycin; Ap, ampicillin; 5-FU, 5-fluorouracil; superscript “r” or “s,” resistance or sensitivity, respectively.

as 5-FU<sup>r</sup> and screened for Km sensitivity. The accuracy of the constructs was confirmed by sequencing both strands of a PCR-amplified product across the mutation site. The deletion strain and promoter mutation strains that prevented *sat* transcription were expected not to grow with sulfate as an electron acceptor, and therefore 20 mM sulfite was used in medium to recover these promoter mutations.

**Quantitative reverse transcriptase-PCR (qRT-PCR).** (i) **Initial optimization.** Primers were manually designed to amplify an approximately 100- to 150-bp region at the 3' end of the transcript for each gene (see Table S1 in the supplemental material). qPCRs were set up according to the manufacturer's protocol with SsoFast EvaGreen Supermix (Bio-Rad, Hercules, CA) on a CFX96 and analyzed with CFX Manager (version 3.1; Bio-Rad), with the curve fit to regression (35–37). The amplification protocol was initiated with a 3-min preincubation at 95°C and processed through 40 cycles of denaturation (30 s at 95°C) and annealing/extension (30 s at 65°C). The fluorescent signal was acquired at the end of each

annealing/extension step. The melting curve protocol included annealing at the annealing/extension temperature (65°C) and melting at a ramp rate of 0.5°C/5 s up to 95°C, with the fluorescent signal acquired continuously during the melting curve. The genes to be used as internal (reference) controls were *rplS* (DVU\_0835) and *rpmC* (DVU\_1311) (38) because they are expressed at levels similar to those of the genes to be assessed in the present study and their expression levels were not found to change during exposure to environmental stresses (<http://www.microbesonline.org/>; data not shown). To validate these reference genes, the strategy of Helleman et al. was implemented (35), and the genes were shown to have minimal transcriptional differences among the strains, mutants, and conditions tested, with a mean coefficient of variance (CV) of <0.25 and mean (M) of <0.5.

(ii) **Sample preparation.** Samples were initially resuspended in 1 ml of TRI reagent (Sigma-Aldrich) and kept frozen until RNA isolation by phenol extraction (39, 40). The RNA quality was assessed visually in an



agarose-denaturing gel, and an  $A_{260}/A_{280}$  ratio of  $>1.8$  was required. RNA samples were treated with Turbo DNase (Life Technologies, Carlsbad, CA) according to the manufacturer's protocol and confirmed by PCR to be free of genomic DNA. For each gene to be analyzed, a standard curve (6 logs, serial dilution from 100 ng/ $\mu$ l stock cDNA) was performed to calculate efficiency. For each transcript, the relative abundance was normalized by the reference gene transcripts in the specified sample.

**Transcriptional start site (TSS) determination by 5'-RACE (rapid amplification of cDNA ends).** Samples were initially collected from DNase-treated RNA samples used for qRT-PCR. The procedure used was adapted from that used by Scotto-Lavino et al. (41). In brief, 100 ng of DNase-treated RNA was reverse transcribed to single-stranded cDNA (iScript Select cDNA synthesis kit; Bio-Rad) with primer DVU1295-sat-GSP1, degraded with RNase A/T<sub>1</sub> mix and RNase H (Thermo Fisher Scientific), and the cDNA was purified by column purification. With terminal deoxynucleotidyltransferase (Thermo Fisher Scientific), an adenosine tail was added to the cDNA, followed by the generation of the second DNA strand with iScript and the primer RACE-2nd Strand. The RACE-2nd Strand primer was adapted from primer AUAP (Invitrogen 5' RACE system, v2.0) to include additional T residues and a "V" at the 3' position ("V" as A/G/C but not T) to allow for better anchoring to the modified cDNA. Double-stranded DNA (dsDNA) was purified, diluted 1:1,000, and amplified by PCR (PCR1). Thirty cycles of amplification were carried out (30 s at 94°C, 30 s at 58°C, and 1 min at 72°C), followed by a final extension for 5 min at 72°C with *Taq* DNA polymerase (NEB, Ipswich, MA). An additional PCR (PCR2) followed, which was performed according to the same protocol used for PCR1, but with 1:1,000-diluted PCR1 as the template and nested primers. Fragments were purified, sequenced, and mapped to the genome.

**In vitro protein-DNA interaction assays.** To obtain  $Rex_{DvH}$  for protein-DNA interaction studies, a 6 $\times$ His tag was added to  $Rex_{DvH}$  by cloning *rex* into pET-14b (Novagen, Madison, WI) and transforming this into BL21(DE3) competent cells (Agilent, Santa Clara, CA). After induction with IPTG (isopropyl- $\beta$ -D-thiogalactopyranoside; Gold Biotechnologies), the tagged protein was purified by using a His60 Ni gravity column purification kit (Clontech, Mountain View, CA). Eluted 1-ml fractions were analyzed for protein (42), and fractions were pooled together and passed over a PD-10 desalting column (GE Healthcare Biosciences, Piscataway, NJ). SDS-PAGE was performed to ensure the purity of the  $Rex_{DvH}$  monomer (~25 kDa) (data not shown). The presence of NADH bound to  $Rex_{DvH}$  was checked by measuring the absorbance at 340 nm to ensure the purity of the protein from additional cofactors.

The protocol by Brekasis and Paget (25) was used for DNA template generation of the relevant  $Rex_{DvH}$ -binding site ( $>100$  bp). For smaller DNA fragments (40 bp), reverse complemented primers were annealed (Sigma-Aldrich). These sequences were 5' end labeled with T4 polynucleotide kinase (Promega) and [ $\gamma$ -<sup>32</sup>P]ATP (Perkin-Elmer, Waltham, MA) according to the manufacturer's protocol. The unlabeled nucleotides were removed by using a QIAquick nucleotide removal kit (Qiagen, Valencia, CA).

**EMSA.** Electrophoretic mobility shift assays (EMSAs) were performed according to the method of Ravcheev et al. (21). In brief, dsDNA fragments (0.1 nM) were incubated with  $Rex_{DvH}$  at specified concentrations (0 to 2,000 nM) in a final volume of 30  $\mu$ l. The binding buffer contained 20 mM Tris-HCl (pH 8.0), 10% (vol/vol) glycerol, 1 mM MgCl<sub>2</sub>, and 40 mM KCl. Samples were incubated at 37°C for 25 min, placed on ice for 2 min, and then separated (90 V, 70 min, 4°C) on a 5% (wt/vol) native Tris-borate-EDTA (TBE) polyacrylamide gel that was preincubated (200 V, 30 min at 4°C) in 0.5 $\times$  TBE buffer (Bio-Rad). When pyridine nucleotides were examined, the desired concentrations were added after the initial incubation and then incubated for an additional 10 min at 37°C. The gel was removed from the apparatus, wrapped in plastic wrap, and exposed to a Kodak Imaging Screen K (Bio-Rad) typically for 15 to 60 min

TABLE 2 Transcript analysis of parental,  $\Delta$ *rex*, and complement of *rex* strains grown by sulfate respiration or pyruvate fermentation<sup>a</sup>

Medium and strain	Description or genotype	Mean transcript level $\pm$ SEM for:	
		<i>sat</i>	<i>rex</i>
MOLS4			
JW710	Parental	1.0 $\pm$ 0.2	1.0 $\pm$ 0.0
JW3311	$\Delta$ <i>rex</i>	11.2 $\pm$ 2.3	ND
JW3311(pMO3313)	Complement of <i>rex</i>	5.9 $\pm$ 0.8	10.9 $\pm$ 0.0
MOYPyr			
JW710	Parental	7.3 $\pm$ 1.2	1.7 $\pm$ 0.0
JW3311	$\Delta$ <i>rex</i>	13.6 $\pm$ 3.7	ND
JW3311(pMO3313)	Complement of <i>rex</i>	6.4 $\pm$ 0.6	24.9 $\pm$ 0.2

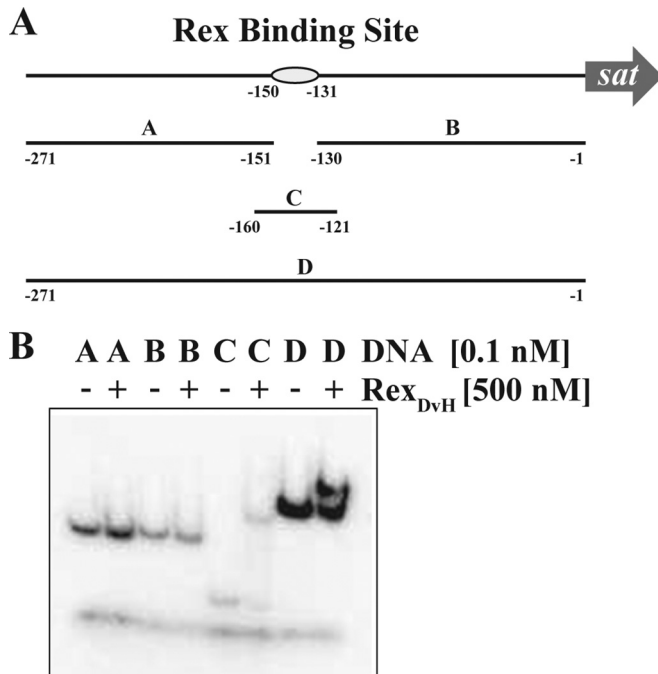
<sup>a</sup> The OD<sub>600</sub> was monitored through growth, and samples were collected for analysis by qRT-PCR at the early exponential phase. Approximately 100 ng of Turbo DNase-treated RNA was converted to cDNA, and 1  $\mu$ l of cDNA (5 ng of RNA) was used per qRT-PCR. Each gene was assessed individually and normalized with respect to the reference genes *rplS* and *rpmC*. The efficiency for each gene was determined as follows: *rplS* = 92.6%, *rpmC* = 95.5%, *sat* = 91.3%, and *rex* = 90.2%. Samples were normalized to JW710 MOLS4. ND, not detected.

in a sealed cassette, followed by imaging with a personal molecular imager (Bio-Rad).

**Fluorescent polarization assay (FPA).** Fluorescent polarization was performed as described previously (43). The binding assay was performed in 96-well black plate (VWR, Radnor PA) with 1 nM fluorescently labeled (6-FAM) oligonucleotides. Different concentrations of protein (10, 25, 50, 100, 250, 500, and 1,000 nM) were incubated with 1 nM labeled oligonucleotides in 100- $\mu$ l reaction mixture in the binding buffer (20 mM Tris-HCl [pH 7.5], 100 mM NaCl, 0.3 mg of bovine serum albumin/ml, 1  $\mu$ g of herring sperm DNA). The fluorescence reading was taken on a Beckman multimode plate reader (DTX 880) with excitation and emission filters at 495 and 520 nm. The background fluorescence from buffer was subtracted and the fluorescence polarization values were defined as follows:  $P_{mp} = [(I_{parallel} - G\text{-factor}) \times I_{perpendicular}] / [(I_{parallel} + G\text{-factor}) \times I_{perpendicular}] \times 1,000$ , where  $I_{parallel}$  and  $I_{perpendicular}$  are the fluorescence intensity in the parallel and perpendicular orientation respective to the orientation of the excitation polarizer. The G-factor is an experimental correction for the polarization bias of the detection system (44).

## RESULTS

**Deletion of *rex*.** To examine the role of  $Rex_{DvH}$ , a marker-exchange deletion of *rex* and a complemented deletion strain were constructed (33). These two strains, in addition to the parental strain, were grown by sulfate respiration in MOLS4 or by pyruvate fermentation in MOYPyr. The former was expected to differ from the latter by changes in the ratio of NADH/NAD<sup>+</sup> proposed to be a signal for Rex regulation. It was assumed that fermenting cultures lacking an inorganic terminal electron acceptor would exhibit an increase in NADH (45). Examination of the growth in either medium revealed no significant differences among the three strains (see Table S2 in the supplemental material). In parental cells, qRT-PCR analysis of *sat* expression (Table 2) showed that *sat* transcription had a 7-fold increase in fermenting cultures compared to cultures respiring sulfate. In the  $Rex_{DvH}$  mutant, *sat* expression was minimally increased when growing fermentatively but was already 11-fold increased from the parental strain while respiring sulfate, a finding consistent with a repressor role for Rex. When this strain was complemented with a plasmid copy of *rex* that was transcribed from a constitutive promoter, partially restored (decreased) levels of *sat* transcription were observed. To



**FIG 1** EMSA demonstrating specific interaction between Rex<sub>DvH</sub> and the predicted Rex<sub>DvH</sub>-binding site within the *sat* promoter. (A) Schematic representation of the *sat* promoter region drawn approximately to scale. The predicted Rex<sub>DvH</sub>-binding site is annotated by an oval. Fragments A, B, C, and D used in EMSA are shown with their positions noted. Fragments A (121 bp), B (130 bp), and D (271 bp) were PCR amplified, while fragment C (40 bp) was generated by annealing two oligonucleotides. (B) Native polyacrylamide gel of individual DNA fragments (A, B, C, and D; 1 nM stock prior to column purification) without (–) or with (+) Rex<sub>DvH</sub>. An equal concentration of DNA was labeled and then passed over a column to separate the fragments from the rest of the components, i.e., unlabeled nucleotides. Fragment C, the smallest fragment, is below the size cutoff for the column (~100 bp), and so only a small amount of this fragment is actually recovered compared to the other three larger fragments. Each fragment was eluted in the same volume of buffer, and so the concentration of this smaller fragment is considerably lower than the rest. Therefore, the band intensity for fragment C is less than the others. The lowest band common in all lanes is the dye front.

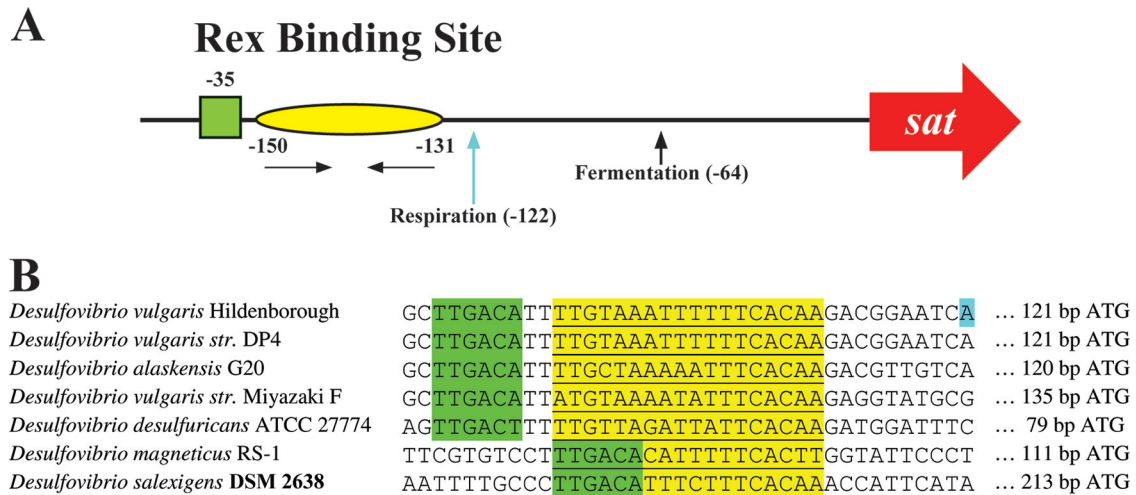
eliminate the possibility that growth modes were affecting the expression of *rex*, transcription of that gene was examined and found to be unchanged in the parental strain and was undetected in the Rex<sub>DvH</sub> mutant as expected. Therefore, these studies provided support that Rex<sub>DvH</sub> is a transcriptional repressor for *sat*.

**Rex binding site in the *sat* promoter.** Since the expression levels for *sat* were increased by the deletion of *rex*, we sought to determine whether the regulation was direct or indirect. There is a putative Rex binding site located at positions –150 to –131 (TTTGTAATTTTTTTCACAAG) relative to the translational start codon for *sat* (21). Therefore, Rex<sub>DvH</sub> was purified for protein-DNA interaction studies. Four dsDNA fragments were examined for interaction with Rex<sub>DvH</sub> (Fig. 1): one upstream and one downstream of the predicted Rex binding site, and two of different sizes that contained the motif. The two fragments that contained the Rex binding site (fragments C and D in Fig. 1) shifted in electrophoretic mobility when the putative Rex<sub>DvH</sub> protein was present, whereas the other two fragments did not. These results confirm a direct interaction between Rex<sub>DvH</sub> protein and the putative Rex<sub>DvH</sub>-binding site upstream of *sat*.

**Transcriptional start sites for *sat*.** With a physical interaction observed *in vitro* between Rex<sub>DvH</sub> and the Rex<sub>DvH</sub>-binding site upstream of *sat*, determination of the relative proximity of this motif to the transcription start site (TSS) for *sat* might provide a logical mechanism for regulation. This analysis could provide evidence that Rex<sub>DvH</sub> repressed by occluding the polymerase from interaction with the promoter region of *sat*. Therefore, RNA samples were isolated from parental and Rex<sub>DvH</sub> strains grown by sulfate respiration or pyruvate fermentation and assessed for the TSS of *sat* (Fig. 2). To determine the 5' end of the transcript, the RACE technique was applied. Examination of the TSS for the parental strain revealed two unique sites: one that was identified from cells growing by sulfate respiration at bp –122 relative to the assumed translational start codon and one from pyruvate-fermenting cells at bp –64. A previous study performed by 5'-RNA-seq analysis also identified bp –122 as the TSS of *sat* for *D. vulgaris* Hildenborough grown by sulfate respiration (10). The Rex<sub>DvH</sub>-binding site (bp –150 to –131) is less than 10 bp upstream of the 5' end of the mRNA and therefore supports the occlusion of RNA polymerase binding for repression by Rex<sub>DvH</sub> during respiration. Furthermore, a potential –35 site of a  $\sigma^{70}$  promoter was identified in close proximity to this region (Fig. 2). Interestingly, examination of the TSS for the Rex<sub>DvH</sub> mutant growing either by respiration or by fermentation revealed the same sites as those identified for the parental strain. Because two sites were observed and deletion of *rex* did not reveal a change, these results suggest that factors in addition to Rex<sub>DvH</sub> are involved in determining the TSS.

**Effect of NADH concentrations on Rex<sub>DvH</sub> function.** Rex proteins contain a pyridine nucleotide binding domain that has been shown in other bacteria to interact with NAD<sup>+</sup> or NADH and influence regulation (22, 46). To determine the role of pyridine nucleotide interaction with Rex<sub>DvH</sub>, DNA-binding assays were performed with purified Rex<sub>DvH</sub> protein in the presence of NAD<sup>+</sup>, NADH, NADP<sup>+</sup>, or NADPH at 0.1 or 1.0 mM (Fig. 3) and a DNA fragment containing the Rex<sub>DvH</sub>-binding site (fragment C, Fig. 1). The addition of NAD<sup>+</sup> appeared to have little effect on Rex<sub>DvH</sub> binding at either concentration compared to purified Rex<sub>DvH</sub> and DNA alone, whereas NADH disrupted the interaction at both concentrations tested. NADP<sup>+</sup> did not appear to have any effect on the binding event, and NADPH disrupted binding only at high, presumably nonphysiological, concentrations. Concentration gradient assays conducted for each of the pyridine nucleotides substantiated these results (data not shown), which are similar to the findings of the Rex homolog in *T. aquaticus* (22).

**Confirmation of essential nucleotides in the Rex binding motif.** To determine the key base pair(s) recognized by Rex<sub>DvH</sub> for binding, a number of mutations within the motif upstream of *sat* were altered and binding studies were performed (Fig. 4A and Table 1). The strategy for the base alterations was to make transitional mutations (i.e., purine to purine [A↔G] or pyrimidine to pyrimidine [C↔T]) for the most conserved bases of the predicted Rex<sub>DvH</sub>-binding sites in *D. vulgaris* Hildenborough (see sequence logo, Fig. 4A). From the JW9293 deletion strain ( $\Delta P_{sat}$ ) lacking the Rex motif, five additional strains were constructed: (i) a restored promoter sequence with the exact promoter region upstream of *sat* as the parental strain (restored P<sub>sat</sub>), (ii) a promoter with the position –147 conserved “G” residue altered to an “A” (G–147A), (iii) a promoter with the distal inverted repeat “GTA” altered to “ACG” (–147 to –145) (IR1), (iv) a promoter with the well-conserved proximal inverted repeat “CAC” altered to “TGT”



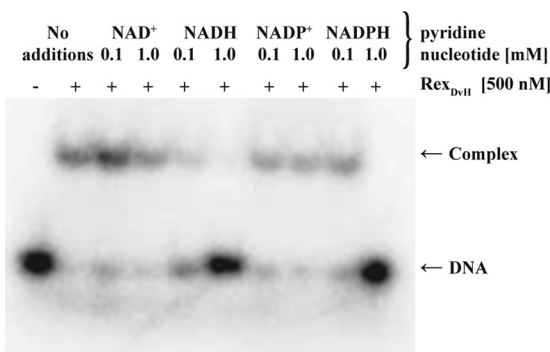
**FIG 2** Two transcription start sites (TSSs) for *sat* identified dependent on growth substrates. Parental and  $\text{Rex}_{\text{DVH}}$  mutant strains were grown in medium that would allow for sulfate respiration or pyruvate fermentation. Samples at the early-exponential-growth phase were analyzed for the TSS of *sat* by 5'-RACE. (A) Schematic representation of *sat* promoter. The predicted  $\text{Rex}_{\text{DVH}}$ -binding site is annotated with a yellow oval with nucleotide positions listed relative to the assumed ATG translational start codon of *sat* in *D. vulgaris* Hildenborough. Horizontal arrows denote the half-sites (inverted repeats) within the  $\text{Rex}_{\text{DVH}}$ -binding site. The predicted -35 site is indicated by a green box. TSSs are identified with vertical arrows (and positions) for each sample tested. (B) The  $\text{Rex}$ -binding site (underlined, highlighted in yellow) and the surrounding region is shown for the promoter sequence of *sat* of *Desulfovibrio* strains, with the predicted -35 site displayed (TTGACA, highlighted in green). A TSS (respiration) for *D. vulgaris* Hildenborough is highlighted in blue.

(-135 to -133) (IR2), and (v) a promoter with mutations to both inverted repeat sequences in a single strain (IR1and2). The mutants along with the restored promoter and parental strains were grown by sulfate respiration (Fig. 4B) and pyruvate fermentation (Fig. 4C). Growth was similar for all strains with the exception of that deleted for the promoter ( $\Delta P_{\text{sat}}$ ), which was unable to grow by sulfate respiration and consistently grew slightly more efficiently by pyruvate fermentation.

It was predicted that the modifications to the  $\text{Rex}_{\text{DVH}}$ -binding site might prevent  $\text{Rex}_{\text{DVH}}$  repression of *sat* and therefore increase expression of *sat*. Therefore, samples early in exponential growth were analyzed for *sat* and *rex* expression (Table 3) (see Table S2 in the supplemental material). All strains with sequence changes in the  $\text{Rex}_{\text{DVH}}$ -binding site had increased *sat* expression levels relative to the parental and restored strains. dsDNA fragments of 40

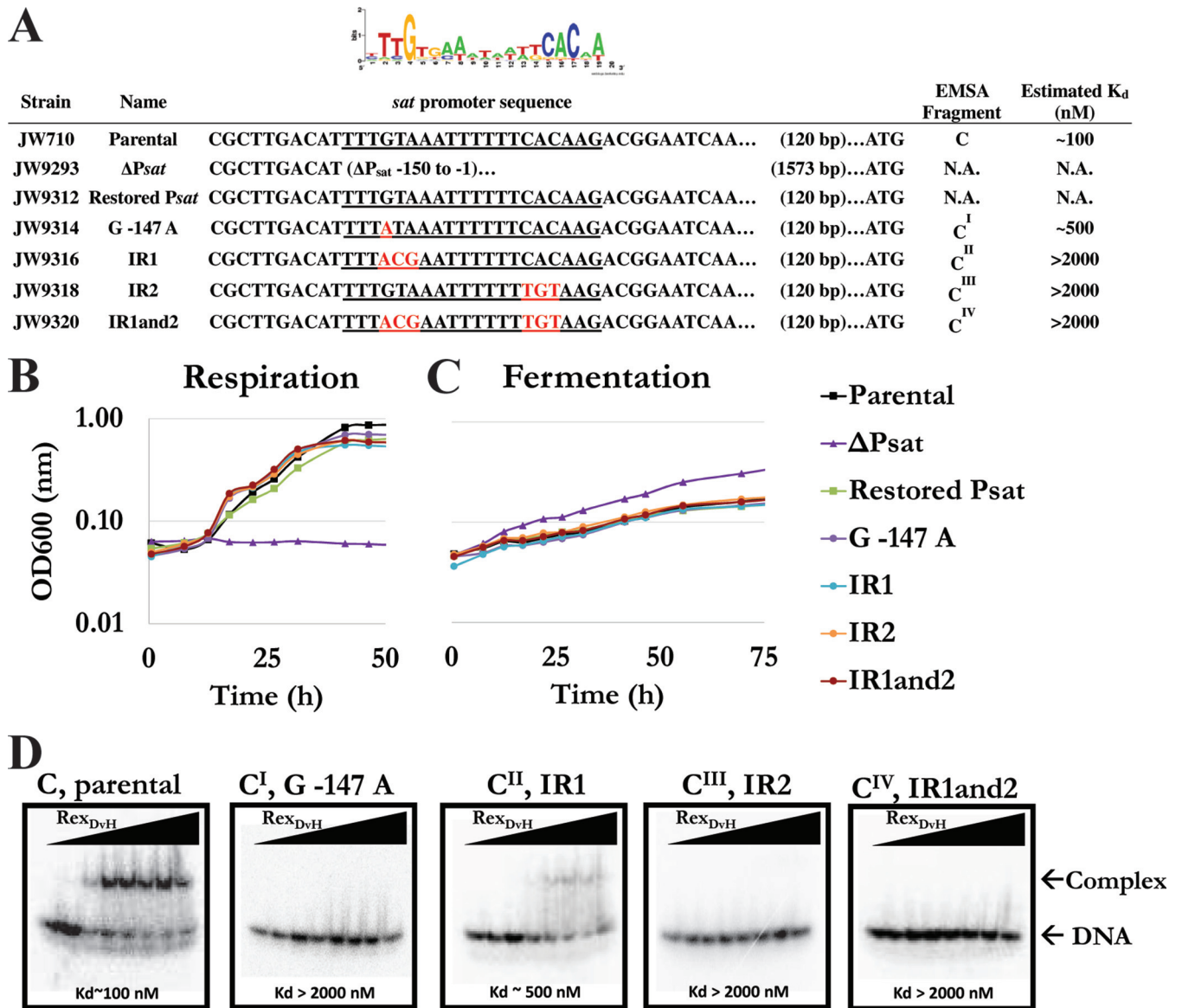
bp containing the same alterations that were introduced into the genome were assayed for interactions with increasing  $\text{Rex}_{\text{DVH}}$  concentrations (0 to 2,000 nM) (Fig. 4D). For three of the altered fragments tested, G-147A, IR2, and IR1and2 ( $C^I$ ,  $C^{III}$ , and  $C^{IV}$ ), no shift was observed (dissociation constant [ $K_d$ ] > 2,000 nM). However, the fragment that contained the three-base alteration to the distal inverted repeat sequence, IR1 ( $C^{II}$ ), did shift ( $K_d \sim 500$  nM), although not to the same extent as the wild-type sequence ( $K_d \sim 100$  nM). This result was rather interesting because this sequence also contained the G-147A alteration that appeared to completely disrupt the binding. As expected, no *sat* transcription was detected for the strain deleted for the *sat* promoter grown by pyruvate fermentation (Table 3). The expression of *rex* was also examined to verify that an unexpected decrease in  $\text{Rex}_{\text{DVH}}$  caused by a transcriptional change was not a factor for the observed differences in *sat* expression. Across the  $P_{\text{sat}}$  strains, *rex* expression was not significantly different. Therefore, alterations to the  $\text{Rex}_{\text{DVH}}$ -binding site within the promoter sequence of *sat* increased *sat* expression confirming the importance of the highly conserved base pairs in the motif.

**$\text{Rex}_{\text{DVH}}$  interacts *in vitro* with all predicted  $\text{Rex}_{\text{DVH}}$ -binding motifs in the *D. vulgaris* Hildenborough genome.** To explore potential  $\text{Rex}_{\text{DVH}}$  regulation of other target genes, a more high-throughput DNA-binding assay was used, fluorescent polarization assay (FPA). Twelve operons with putative Rex motifs in their upstream regions, predicted at the time of the present study (21), were analyzed (see Table S3 in the supplemental material). Exact 20-bp predicted  $\text{Rex}_{\text{DVH}}$ -binding sites were created with five "G's" at the 5' end to improve annealing and with four "G's" with a "T" at the 3' end to which the fluorophore would be attached. This approach takes advantage of the fact that the degree of polarization of a fluorophore is inversely related to its molecular rotation. Thus, the change in fluorescence of a fast-moving small unbound DNA fragment compared to a larger  $\text{Rex}_{\text{DVH}}$ -bound DNA fragment is evidence of protein-DNA interaction. By increasing the



**FIG 3** NADH disrupts interaction between  $\text{Rex}_{\text{DVH}}$  and the  $\text{Rex}_{\text{DVH}}$ -binding site. The results of an electrophoretic assay demonstrate the effect of the pyridine nucleotide on  $\text{Rex}_{\text{DVH}}$  binding. Fragment C (40 bp, including the  $\text{Rex}_{\text{DVH}}$ -binding site upstream of *sat*) at 0.1 nM (1 nM prior to column purification) was incubated with  $\text{Rex}_{\text{DVH}}$  in the presence of a low or high concentration of the specified pyridine nucleotide. The location of the DNA or protein-DNA complex is indicated.





**FIG 4** Characterization of alterations to the  $Rex_{DvH}$ -binding site within the *sat* promoter. (A) List of strains and alterations made to the  $Rex_{DvH}$ -binding site upstream of *sat*, with a sequence logo of the predicted  $Rex_{DvH}$ -binding site shown above. The  $Rex_{DvH}$ -binding site is underlined, with the alterations made shown in red. An alignment of sequences is shown relative to the assumed translational start codon for *sat*. The fragments used for EMSA are displayed, along with the estimated  $K_d$  (nM) for each 40-bp fragment. N.A., not assessed. The strains examined included the parental strain (JW710), the *sat* promoter exchange deletion strain ( $\Delta P_{sat}$ ), the restored promoter strain ( $P_{sat}$ ), the conserved “G-147” altered to “A” strain (G-147A), the distal inverted repeat “GTA” altered to “ACG” strain (IR1), the proximal inverted repeat “CAC” altered to “TGT” strain (IR2), and the strain with alterations to both inverted repeat sites (IR1and2). The relative growth of three replicates of mutants and the parental strain by sulfate respiration (B) or pyruvate fermentation (C) was examined. (D) An electrophoretic assay demonstrated  $Rex_{DvH}$  binding to native (fragment C) and altered (C<sup>I</sup>, C<sup>II</sup>, C<sup>III</sup>, and C<sup>IV</sup>)  $Rex_{DvH}$ -binding sites.  $Rex_{DvH}$  was added with increasing concentrations (0, 10, 100, 250, 500, 750, 1,000, and 2,000 nM) to each DNA fragment (0.1 nM; 1 nM prior to column purification). The estimated  $K_d$  is shown. The location of the DNA or protein-DNA complex is noted. Triangles represent increasing  $Rex_{DvH}$  concentration across the lanes.

protein concentration over a range of values (0 to 1,000 nM), a dissociation constant was determined for  $Rex_{DvH}$  with each 6-FAM-labeled dsDNA fragment. Dissociation constants ( $K_d$ ) of ca. 40 to 105 nM were determined and were similar to previously published values for Rex (i.e.,  $K_d \approx 1$  to 100 nM) (21, 22, 47). The two techniques used in the present study to calculate protein-DNA interaction between  $Rex_{DvH}$  and the  $Rex_{DvH}$ -binding site upstream of *sat* were similar (EMSA,  $K_d \approx 100$  nM; FPA,  $K_d \approx 90$  nM). In conclusion,  $Rex_{DvH}$  protein was determined to interact *in*

*vitro* with all predicted  $Rex_{DvH}$ -binding sites, and the calculated  $K_d$  values were similar.

## DISCUSSION

The sulfate reduction gene *sat* has been shown to be altered in expression depending on the available electron acceptor (18). Bioinformatic predictions for the sulfate reduction pathway, including coexpression studies of the genes, as well as looking for conserved motifs upstream of these genes in multiple *Desulfovib-*

**TABLE 3** Transcript analysis of parental and Rex<sub>DvH</sub>-binding site alteration strains grown by sulfate respiration or pyruvate fermentation<sup>a</sup>

Medium and strain	Description or genotype	Mean transcript level ± SEM for:	
		<i>sat</i>	<i>rex</i>
<b>MOLS4</b>			
JW710	Parental	1.0 ± 0.1	1.0 ± 0.1
JW9293	ΔP <sub><i>sat</i></sub>		
JW9312	Restored P <sub><i>sat</i></sub>	0.9 ± 0.6	0.9 ± 0.5
JW9314	G-147A	2.7 ± 0.2	1.2 ± 0.1
JW9316	IR1	1.8 ± 0.1	0.9 ± 0.1
JW9318	IR2	4.0 ± 0.3	1.0 ± 0.1
JW9320	IR1and2	4.6 ± 0.6	1.4 ± 0.3
<b>MOYPyr</b>			
JW710	Parental	1.0 ± 0.1	1.0 ± 0.1
JW9293	ΔP <sub><i>sat</i></sub>	ND	0.6 ± 0.1
JW9312	Restored P <sub><i>sat</i></sub>	1.0 ± 0.3	1.6 ± 0.7
JW9314	G-147A	2.9 ± 0.7	0.9 ± 0.2
JW9316	IR1	3.7 ± 0.7	0.9 ± 0.1
JW9318	IR2	2.8 ± 0.5	1.1 ± 0.1
JW9320	IR1and2	2.3 ± 0.1	1.5 ± 0.1

<sup>a</sup> The OD<sub>600</sub> was monitored through growth, and samples were collected for analysis by qRT-PCR at the early exponential phase. Approximately 100 ng of Turbo DNase-treated RNA was converted to cDNA, and 1 μl of cDNA (5 ng of RNA) was used per qRT-PCR. Analysis of each gene was conducted separately for each medium tested. For MOLS4, the gene efficiencies were as follows: *rplS* = 91.8%, *rpmC* = 84.2%, *sat* = 92.5%, and *rex* = 83.6%. For MOYPyr, the gene efficiencies were as follows: *rplS* = 90.7%, *rpmC* = 106.1%, *sat* = 97.7%, and *rex* = 89.2%. Each gene was assessed individually and normalized to the reference genes *rplS* and *rpmC*. JW9293 (ΔP<sub>*sat*</sub>) was unable to be grown by sulfate respiration and therefore no data are provided. Transcript values obtained with MOLS4 and MOYPyr were normalized to the corresponding value obtained for JW710. ND, not detected.

*rio* species, proposed regulatory contributions from Rex (encoded by DVU\_0916) (21), HcpR (encoded by DVU\_2547) (48), LysX (encoded by DVU\_2567), and other DNA-binding proteins (encoded by DVU\_0057, DVU\_0744, DVU\_1142, DVU\_2275, DVU\_2690, DVU\_2799, and DVU\_2802) (20). Preliminary transcript studies examining *sat* expression were performed on parental *D. vulgaris* Hildenborough cells, on a marker-exchange deletion of *hcpR*, and on a transposon insertion mutant of *lysX*. No observable differences of >2-fold were found among strains growing on 60 mM lactate with 30 mM sulfate or 20 mM sulfite (see Table S4 in the supplemental material). Therefore, these candidates were no longer pursued as major regulators of the sulfate activation and reduction steps for the present study. However, these proteins may still be regulators under other physiological conditions or regulators of other genes in the complete reduction of sulfate. In particular, annotated transcriptional regulators DVU\_0744, DVU\_2802, and DVU\_2275 have been examined further by others and found to play a role in expression of sulfate reduction targets during respiration (20).

Preliminary transcript studies examining *sat* expression of a Rex<sub>DvH</sub> mutant had shown that the deletion of *rex* increased *sat* expression relative to a parental strain (data not shown). These findings are consistent with the role of Rex<sub>DvH</sub> as a repressor of *sat*. This transcription factor has been proposed to be responsible for the redox poise of the cell through the NADH/NAD<sup>+</sup> ratio and to alter cellular metabolism to reestablish the pyridine nucleotide balance (21). To confirm the role of Rex<sub>DvH</sub>, the Rex<sub>DvH</sub> mutant

was cultured in two media proposed to alter the redox status of the cell, and *sat* transcription was measured (Table 2). Examination of the Rex<sub>DvH</sub> mutant revealed increased *sat* transcripts compared to the parental strain for both growth modes. Furthermore, the differential increase in transcript levels observed in the absence of sulfate for the parental strain was not maintained in the Rex<sub>DvH</sub> mutant. We interpreted these observations to mean that Rex<sub>DvH</sub> is a transcriptional repressor for *sat* responding to redox status in the cell.

The *rex* gene encoded on a plasmid under a *npt* promoter was introduced into the Rex mutant to test restoration of *sat* repression. When this complementation construct was respiring sulfate, *sat* repression was only partially restored. However, *rex* was transcribed >10-fold higher than in the wild type (Table 2). Interestingly, for the fermenting culture the expression of *sat* was restored to wild-type levels. The increased transcription of *rex*, and therefore we assume the protein Rex<sub>DvH</sub>, in the complemented strain might lead to higher-than-normal levels of repression of other targets of Rex<sub>DvH</sub> not yet studied. This aberrant expression of *rex* may adversely affect the overall metabolism of the cell. However, a comparison of growth curves showed little difference in rate or extent of sulfate respiration by the parent, *rex* mutant and complement under the conditions examined in the present study.

Interestingly, there is a conserved hypothetical gene located 33 bp downstream of *rex*, DVU\_0915, that was originally predicted to be in the same operon. However, the expression of DVU\_0915 is quite low and, based on a large battery of transcriptome data publicly available for *D. vulgaris* Hildenborough (<http://www.microbesonline.org/>), these genes do not appear to be coregulated. Examination of the intergenic region revealed several strong hairpins that might function as transcriptional regulators consistent with a separate operonal structure for the adjacent genes. Alternatively, the deletion of the DNA sequence of *rex* may have eliminated regulatory elements responsible for the proper expression of DVU\_0915. Preliminary studies examining DVU\_0915 expression in the three strains revealed that DVU\_0915 expression was indeed elevated in the Rex mutant and complemented strain compared to the parental strain (data not shown). Therefore, the increased abundance of DVU\_0915 may contribute to the inconsistencies in the expression of *sat* in the complemented strain, although no function is known for the protein encoded by this gene.

To date, it has been assumed that Rex<sub>DvH</sub> blocks transcription by preventing the polymerase from binding to promoter DNA, limiting the expression of the downstream gene. Interestingly, based on the identified TSS for respiring cultures the -10 position should be within the Rex<sub>DvH</sub>-binding site, specifically overlapping with the proximal inverted repeat (IR2). However, no conventional -10 site could be identified, but a site similar to a classical -35 consensus (TTCACA) was apparent just upstream of the Rex<sub>DvH</sub>-binding position. This was interpreted to mean that the RNA polymerase binding site of a respiring culture is within the Rex<sub>DvH</sub>-binding site, and therefore the proposed RNA polymerase occlusion mechanism for Rex seems plausible during respiration.

However, this RNA polymerase occlusion mechanism does not explain the second site of transcript initiation apparently functioning when *D. vulgaris* Hildenborough is grown by fermentation and even more *sat* transcript is produced. When sulfate becomes limiting and NADH is plentiful, increased Sat could



possibly scavenge low sulfate as a terminal electron acceptor and/or provide reduced sulfur for cell biosynthesis. Preliminary work suggests that the levels of Sat may be linked to sulfate uptake (G. M. Zane et al., unpublished data). Therefore, it would be reasonable, when the preferred electron acceptor is limiting (e.g., sulfate), that the cell might increase the expression of a gene that encodes a protein that may facilitate sulfate uptake. A second possibility is that there is an alternate regulator that recognizes a specific ligand (e.g., sulfate or sulfite) and that the absence of this ligand is the signal for the selection of the second start site of transcription. It would seem plausible that sulfite, which at high concentrations is toxic to *D. vulgaris* Hildenborough, may be a ligand for controlling the overall expression of *sat* as Sat activates sulfate for reduction to sulfite. This latter interpretation might also explain why *sat* expression is relatively low when sulfite is the electron acceptor and much higher with sulfate.

It was predicted that the deletion of *rex* should resemble the state at which Rex should be removed from the promoter (i.e., fermentation). Therefore, when the Rex<sub>DvH</sub> mutant had the same TSS pattern for *sat* as the parental strain, it was clear that additional regulators are likely involved and that the order of addition of these other factors might be critical for regulation. Therefore, the mechanism proposed above, regarding another potential regulator that responds to sulfite levels, seems reasonable. Sulfite concentrations and not the protein Rex<sub>DvH</sub> could be signaling the selection of the transcriptional start site. Overall, *sat* is still subject to transcriptional control by Rex<sub>DvH</sub> in the fermentative condition since transcription is derepressed to even higher levels when Rex<sub>DvH</sub> is deleted and pyruvate is being fermented.

Because *sat* expression was increased significantly as a result of a deletion of *rex*, we sought to examine the interaction of Rex<sub>DvH</sub> with the promoter of *sat* (Fig. 4). To confirm the specific bases required for interaction, *in vitro* assays were performed on short DNA sequences of the promoter region. All alterations constructed in the consensus binding site disrupted the interaction between Rex<sub>DvH</sub> and the DNA *in vitro*. Interestingly, when a highly conserved base (G-147) was altered to an "A" there was a complete loss of detectable interaction (fragment C<sup>I</sup>, G-147A, "GTA" altered to "ATA"); however, a triple mutation at IR1 (fragment C<sup>II</sup>, "GTA" altered to "ACG"), which included the G-147A mutation, caused only a slight decrease in the interaction and not the complete loss that was observed for the single base mutation. Closer examination of the sequence revealed that the modified sequence for fragment C<sup>II</sup> ("ACG"), for which a "G" is now in the third position, may still resemble the consensus site ("GTG"), but that the single base mutation does not. Therefore, a partially restored binding may occur, but only when the second half site, "CAC," is present as in IR1 (fragment C<sup>II</sup>). However, additional mutation studies would be required to further characterize the specific contribution of each base on Rex<sub>DvH</sub> binding.

Strains containing the altered promoter sequences discussed above were then constructed and grown by sulfate respiration or pyruvate fermentation and compared to the parental strain (Fig. 4). In addition, a strain that was deleted for 150 bp upstream of *sat* ( $\Delta P_{sat}$ ) was also assayed. As expected, *sat* expression increased for any mutation that limited Rex<sub>DvH</sub> binding to the promoter region but was eliminated in the promoter deletion strain. Interestingly,  $\Delta P_{sat}$  grew to a higher cell density while fermenting pyruvate. This phenomenon has been observed for strains with the genes in the sulfate reduction step of the respiratory pathway in *D. vulgaris*

Hildenborough deleted (e.g., quinone-interacting membrane-bound oxidoreductase, *qmoABCD* [data not shown] or a tetraheme cytochrome TplC<sub>3</sub>, *cycA* [49]). The consistent increase in growth on pyruvate may result from a block in sulfate reduction that prevents flux through the activation step, functioning in the parental strain, that requires two ATP equivalents. Although 0.5 mM cysteine is the sulfur source during our growth experiments, it should be noted that nickel (2.3  $\mu$ M) is added as a trace element with sulfate as a counterion in our preparation (NiSO<sub>4</sub>).

In a recent study (21), Rex<sub>DvH</sub> was predicted to interact with the promoter region for 12 operons, including many that encode proteins functioning in other steps of sulfate reduction in *D. vulgaris* Hildenborough (see Table S3 in the supplemental material). Interestingly, most of the targets predicted to be regulated by Rex<sub>DvH</sub> are not involved with NADH oxidation directly (except the Rnf complex) as they are for other bacteria (21).

Since we examined here only a subset of potential electron donors and acceptors for Rex<sub>DvH</sub> regulation of *sat*, the effect of other substrates (e.g., pyruvate, formate, H<sub>2</sub>, sulfite, or thiosulfate) might reveal additional features of Rex regulation. In fact, we recently reported that the Rex<sub>DvH</sub> mutant was inhibited for growth with thiosulfate as the terminal electron (33), an observation that deserves examination. In addition, Rex<sub>DvH</sub> has been predicted to regulate more than 50 genes (21). Recently, one of those genes, *rnfC*, has been shown to decrease in a strain deleted for *rex* in *D. alaskensis* G20 (30), suggesting Rex as an activator as well. Finally, more than 150 potential regulators have been predicted in *D. vulgaris* Hildenborough and should be considered either for their interaction with Rex<sub>DvH</sub> or for their role in regulating genes that encode for proteins responsible for sulfate reduction. Interestingly, the binding site for Rex<sub>DvH</sub> is similar to several known FNR family transcription factors that might compete with Rex<sub>DvH</sub> for binding. Much additional work is needed to identify the transcriptional regulators that signal cellular nutrient and energy status that are integrated at the level of control of sulfate reduction.

## ACKNOWLEDGMENTS

This research, conducted by ENIGMA-Ecosystems and Networks Integrated with Genes and Molecular Assemblies (<http://enigma.lbl.gov/>), a Scientific Focus Area Program at Lawrence Berkeley National Laboratory, was supported by the Office of Science, Office of Biological and Environmental Research, of the U.S. Department of Energy under contract DE-AC02-05CH11231.

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