

The H₂ Sensor of *Ralstonia eutropha* Is a Member of the Subclass of Regulatory [NiFe] Hydrogenases

LAURA KLEIHUES,[†] OLIVER LENZ, MICHAEL BERNHARD, THORSTEN BUHRKE,
AND BÄRBEL FRIEDRICH*

Institut für Biologie, Humboldt-Universität zu Berlin, 10115 Berlin, Germany

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Two energy-generating hydrogenases enable the aerobic hydrogen bacterium *Ralstonia eutropha* (formerly *Alcaligenes eutrophus*) to use molecular hydrogen as the sole energy source. The complex synthesis of the nickel-iron-containing enzymes has to be efficiently regulated in response to H₂, which is available in low amounts in aerobic environments. H₂ sensing in *R. eutropha* is achieved by a hydrogenase-like protein which controls the hydrogenase gene expression in concert with a two-component regulatory system. In this study we show that the H₂ sensor of *R. eutropha* is a cytoplasmic protein. Although capable of H₂ oxidation with redox dyes as electron acceptors, the protein did not support lithoautotrophic growth in the absence of the energy-generating hydrogenases. A specifically designed overexpression system for *R. eutropha* provided the basis for identifying the H₂ sensor as a nickel-containing regulatory protein. The data support previous results which showed that the sensor has an active site similar to that of prototypic [NiFe] hydrogenases (A. J. Pierik, M. Schmelz, O. Lenz, B. Friedrich, and S. P. J. Albracht, *FEBS Lett.* 438:231–235, 1998). It is demonstrated that in addition to the enzymatic activity the regulatory function of the H₂ sensor is nickel dependent. The results suggest that H₂ sensing requires an active [NiFe] hydrogenase, leaving the question open whether only H₂ binding or subsequent H₂ oxidation and electron transfer processes are necessary for signaling. The regulatory role of the H₂-sensing hydrogenase of *R. eutropha*, which has also been investigated in other hydrogen-oxidizing bacteria, is intimately correlated with a set of typical structural features. Thus, the family of H₂ sensors represents a novel subclass of [NiFe] hydrogenases denoted as the “regulatory hydrogenases.”

Molecular hydrogen is frequently used as an energy source by diverse prokaryotic organisms. Many of these bacterial and archaeal species harbor multiple hydrogenases which mediate heterolytic cleavage of H₂ into 2 H⁺ and 2 e⁻. [NiFe] hydrogenases are the most dominant enzymes, representing a fairly conserved family of proteins, composed of at least a large active site-containing subunit and a small electron-transferring subunit which bears one to three FeS clusters (1, 2, 18).

The facultative chemolithoautotrophic proteobacterium *Ralstonia eutropha* H16 (formerly *Alcaligenes eutrophus* [7]) harbors two energy-generating [NiFe] hydrogenases, a membrane-bound enzyme (MBH) and a cytoplasmic enzyme (SH). The MBH is primarily involved in electron transport-coupled phosphorylation, whereas the SH is able to reduce NAD and thus provides the cell with reducing equivalents (38, 40). The composition of the MBH resembles the prototype of [NiFe] hydrogenases whose atomic structure has been resolved by X-ray analysis (50). The two subunits of the *R. eutropha* MBH, encoded by *hoxK* and *hoxG*, are anchored to the outer face of the cytoplasmic membrane via a *b*-type cytochrome (4). The SH, encoded by *hoxF*, *hoxU*, *hoxY*, and *hoxH*, contains an FeS-flavoprotein in addition to the hydrogenase moiety (30). Mutants disrupted in either one of the two hydrogenases maintain their ability to grow on H₂, which indicates that the two enzymes can replace each other physiologically (23).

The hydrogenase-related genes of *R. eutropha* are organized in the MBH and the SH operons, which are regulated coordi-

nately (42). The MBH operon comprises 10 MBH-specific genes in addition to a set of accessory genes whose products are involved in the complex posttranslational maturation of the hydrogenases and the regulation of both the MBH and the SH operon (5, 11, 24, 41). The SH operon harbors the structural genes of the NAD-reducing hydrogenase together with a set of accessory genes which code for maturation proteins (45, 47, 52).

Hydrogenase gene expression is controlled by the major transcription factor HoxA, a member of the NtrC family of response regulators (12). HoxA binds specifically at the upstream regions of the MBH and SH operons and activates transcription in concert with the σ^{54} -containing RNA polymerase (42, 54). Transcription activation by HoxA is stimulated by at least two environmental signals: the presence of molecular hydrogen and/or limitation of organic carbon and energy sources (25, 42). Recognition of molecular hydrogen by cells of *R. eutropha* is mediated by a complex signal transduction system consisting of the proteins HoxB and HoxC which share features of [NiFe] hydrogenases, and HoxJ, a histidine protein kinase which has autophosphorylation capacity with ATP as the phosphoryl donor (25, 26). Deletions in *hoxB* or *hoxC* of *R. eutropha* prevent hydrogenase from being synthesized, whereas a knockout of *hoxJ* leads to H₂-independent high-level hydrogenase gene expression. The data suggest a model in which HoxBC functions as a hydrogen receptor which interacts either directly or indirectly with the sensor kinase HoxJ. Furthermore, unlike in most other two-component regulatory systems, the autophosphorylation-active kinase acts negatively on hydrogenase gene expression. This observation indicates that the HoxJ-mediated phosphorylation of the response regulator HoxA blocks hydrogenase gene transcription. The negative effect of HoxJ is released by HoxBC, provided H₂ is present (25). Proteins similar to HoxBC, designated HupUV, have

* Corresponding author. Mailing address: Institut für Biologie, Humboldt-Universität zu Berlin, Chausseestr. 117, 10115 Berlin, Germany. Phone: 49-30-2093-8100. Fax: 49-30-2093-8102. E-mail: baerbel.friedrich@rz.hu-berlin.de.

[†] Present address: Max-Planck-Institut für molekulare Genetik, 14195 Berlin, Germany.

TABLE 1. Strains, plasmids, and oligonucleotides used in this study

Strain, plasmid, or oligonucleotide	Relevant characteristic(s) or sequence	Source or reference
Strains		
<i>A. eutrophus</i>		
H16	MBH ⁺ SH ⁺ RH ⁺ , HoxJ ^{G422S}	DSM428, ATCC 17699
HF371	<i>hoxGΔ hoxHΔ</i> ; MBH ⁻ SH ⁻ RH ⁺ , HoxJ ^{G422S}	31
HF375	<i>hoxNΔ</i> ; HoxN ⁻ HoxJ ^{G422S}	This study
HF433	MBH ⁺ SH ⁺ RH ⁺	25
HF459	<i>hoxNΔ</i> ; HoxN ⁻	This study
HF500	<i>hoxGΔ hoxHΔ hoxCΔ</i> ; MBH ⁻ SH ⁻ RH ⁻ , HoxJ ^{G422S}	This study
<i>E. coli</i>		
JM109	F' <i>traD36 lacI^q, Δ(lacZ)M15 proA⁺B⁺/e14⁻ (McrA⁻) Δ(lac-proAB) thi gyrA96 (Nal^r) endA1 hsdR17(r_K⁻ m_K⁺) relA1 supE44 recA1</i>	53
S17-1	Tra ⁺ <i>recA pro thi hsdR, chr:RP4-2</i>	43
Plasmids		
LITMUS 29	Ap ^r <i>lacZ'</i> , ColE1 <i>ori</i>	New England Biolabs
pLO2	Km ^r , <i>sacB</i> , RP4 <i>oriT</i> , ColE1 <i>ori</i>	27
pNEB193	Ap ^r <i>lacZ'</i> , ColE1 <i>ori</i>	New England Biolabs
pQE-30	Amp ^r , T5 promoter, ColE1 <i>ori</i>	Qiagen Inc.
pCH128	Derivative of pSUP202 carrying <i>hoxF</i>	13
pCH231	pBluescript KS(+) with a 2.2-kb <i>HindIII-XhoI</i> fragment containing <i>hoxN</i>	14
pCH394	pACYC177 with a 2.38-kb <i>HindIII-SmaI</i> fragment containing <i>hoxB</i>	O. Lenz and B. Friedrich
pCH352	pBluescript KS(+) with a 9.6-kb <i>HindIII/BamHI</i> fragment containing ' <i>hoxA, hoxB, hoxC, hoxJ, and hoxN</i>	K. Marin and B. Friedrich
pCH591	LITMUS 29 with a 0.26-kb <i>HindIII-NdeI</i> fragment containing the SH promoter (P _{SH})	This study
pCH592	pNEB193 with a 0.68-kb <i>NdeI-EcoRI</i> fragment containing <i>hoxB'</i>	This study
pCH593	pCH592 with a 3.3-kb <i>PstI</i> fragment containing ' <i>hoxB, hoxC, hoxJ</i>	This study
pCH594	pCH591 with a 2.5-kb <i>NdeI</i> fragment from pCH593	This study
pCH615	0.8-kb <i>Eco47III</i> fragment containing ' <i>hoxJ</i> into <i>PmeI</i> -digested pLO3	25
pCH644	pLO1 with a 2.45-kb <i>PstI</i> fragment containing <i>hoxCΔ</i>	25
pCH655	1.5-kb <i>PspI</i> (Klenow-treated) fragment containing <i>hoxC</i> into <i>Ecl136II</i> -digested pQE-30	This study
pCH658	0.45-kb <i>FspI-SmaI</i> fragment from pCH231 into <i>EcoRI</i> -digested (Klenow-treated) pCH231	This study
pCH659	pLO2 with a 1.5-kb <i>SalI</i> fragment containing <i>hoxNΔ</i>	This study
pEDY305	RK2 <i>ori</i> , Tc ^r , Mob ⁺ , promoterless <i>lacZ</i> gene	E. Schwartz and B. Friedrich
pEDY309	Derivative of pEDY305 with a modified multiple cloning site	This study
pGE151	Derivative of pRK404	24
pGE301	Φ(<i>hoxK'-lacZ</i>), Tc ^r	27
pGE377	pEDY309 with a 2.8-kb <i>HindIII-XbaI</i> fragment containing P _{SH} - <i>hoxB-hoxC</i>	This study
pGE378	2.2-kb <i>PvuII-Ecl136II</i> containing P _{lac} - <i>hoxA</i> into <i>SwaI</i> -digested pGE377	This study
pGE400	1.95-kb <i>SmaI-Ecl136II</i> fragment containing <i>hoxA</i> into <i>Ecl136II</i> -digested pGE151; P _{lac} - <i>hoxA</i>	This study
Oligonucleotides		
189	GGTGGAAAGGGGTGGCCG	This study
340	CGCCATGGCATATGGTCTCCTCCTACTAATGTTCCG ^a	This study
341	CGCTGGCACAAGCTTGC	This study
342	TGGAGGACATATGAACGCGCCTGTATGTACC ^a	This study
372	CTCTAGAGGAGATCTCACAAGCTTCGAAGCTTAGTCTAGAT	This study
373	CGATCTAGCTAAGTTCGAAGCTTGTGAGATCTCCTCTAGAGGTAC	This study

^a Mutagenic oligonucleotide (underlined residues represent altered nucleotides)

been identified in *Rhodobacter capsulatus* and *Bradyrhizobium japonicum*. Mutant analysis revealed that these proteins play a pivotal role in the H₂-dependent regulation in these organisms. These results led to the conclusion that the HupUV proteins act as a hydrogen sensor (6, 15).

To study the mechanism of H₂-signal transduction in more depth, the interacting partners of the system have to be isolated and characterized in vitro. Because previous attempts to overproduce active [NiFe] hydrogenases heterologously in *Escherichia coli* had not been successful, we constructed a novel expression vector for the native host *R. eutropha*. With the aid of this vector, we achieved an efficient expression of *hoxBC* and show that the resulting protein catalyzes H₂ oxidation. The inspection of mutants revealed that the physiological role of HoxBC is H₂ sensing and not the generation of energy for growth on H₂. Thus, the third hydrogenase of *R. eutropha*

is denoted as the “regulatory hydrogenase” (RH). Both the enzymatic activity and the regulatory function of the RH protein are strictly dependent on the availability of nickel in the medium, showing that nickel is essential for H₂ sensing.

MATERIALS AND METHODS

Strains and plasmids. The strains and plasmids used in this study are listed in Table 1. Strains with the initials HF were derived from *R. eutropha* H16 (wild type). *R. eutropha* HF433 harbors an active H₂-sensing signal transduction chain, including the sensor kinase HoxJ. *R. eutropha* H16 is a natural variant in which the H₂-dependent signal transduction is interrupted by a glycine-to-serine exchange at position 422 in HoxJ (HoxJ^{G422S} [25]). The newly isolated strains *R. eutropha* HF375, a derivative of strain H16, and HF459, a derivative of strain HF433, carry in-frame deletions in the nickel permease gene *hoxN*. *R. eutropha* HF500 bears in-frame deletions in *hoxG, hoxH, and hoxC*, resulting in an MBH⁻ SH⁻ RH⁻ phenotype. Strain HF371 (31) harbors the inactive sensor kinase HoxJ^{G422S} in addition to in-frame deletions in *hoxG* and *hoxH* and was used as host for plasmid-based overexpression of *hoxB* and *hoxC*. *E. coli* JM109 was used

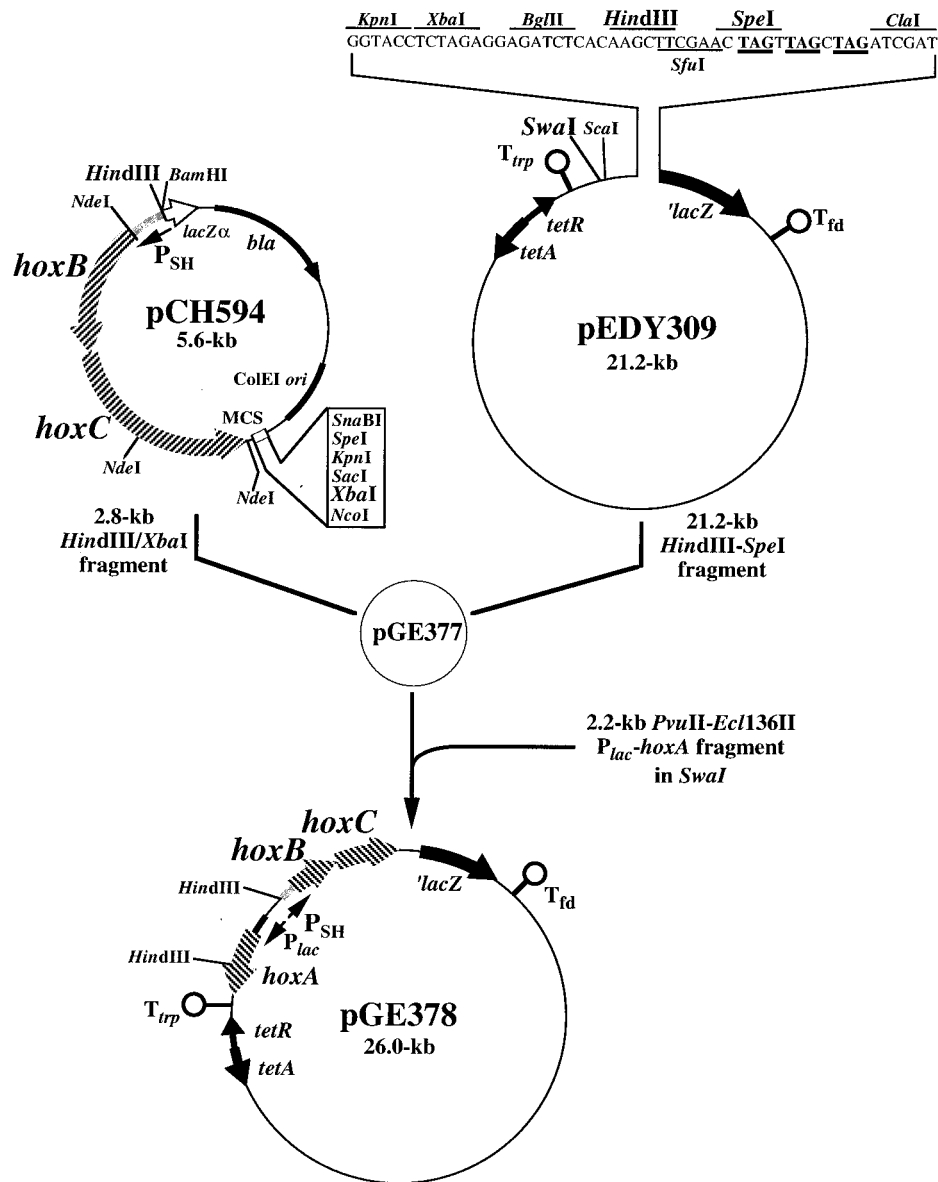


FIG. 1. Construction of the *hoxBC* overexpression vector pGE378. Genes are marked by bold arrows. *R. eutropha*-derived genes are marked by hatched arrows. The orientations of P_{SH} and P_{lac} are given by arrows. Stop codons in all three reading frames downstream of the multiple cloning site (MCS) from pEDY309 are underlined. T_{trp} , *trp* terminator; T_{fd} , phage *fd* terminator.

as a host in standard cloning procedures (53). *E. coli* S17-1 (43) served as a donor in conjugative transfers.

Plasmid pCH591 contains the modified SH promoter region and an *NdeI* site at the ATG start codon of the first gene *hoxF* of the SH operon (Table 1). pCH591 was constructed by amplification of the *hoxF* upstream region from pCH128 using the mutagenic primer 340 and the primer 341. The amplification product was cut with *HindIII* and *NcoI*, and the resulting 0.26-kb fragment was inserted into *HindIII-NcoI*-digested LITMUS 29, yielding pCH591. An *NdeI* site at the ATG start codon of *hoxB* was introduced as follows. pCH394 was used as the template for amplification of the 5' region of *hoxB* with primer 189 and the mutagenic oligonucleotide 342. A 0.68-kb *NdeI-EcoI* fragment of the amplified product was inserted into pNEB193, resulting in pCH592. Subsequently, a 3.3-kb *PstI* fragment from pCH352 was inserted into *PstI*-digested pCH592, resulting in plasmid pCH593 containing the tandemly arranged genes *hoxB* and *hoxC*. A 2.5-kb *NdeI* fragment of pCH593 was cloned into pCH591 yielding pCH594 which harbors the *hoxB* and *hoxC* genes under control of the SH promoter (Fig. 1).

For overexpression of the *hoxB* and *hoxC* genes in the native host *R. eutropha*, we constructed the broad-host-range vector pEDY309 by modification of the multiple cloning region of pEDY305 (Table 1). pEDY305 was digested with *KpnI-ClaI* and used to introduce the polynucleotide kinase-treated hybridization

product of oligonucleotides 372 and 373. Subsequently, the 2.8-kb *HindIII-XbaI* fragment of pCH594 was inserted into pEDY309, yielding plasmid pGE377. Finally, a *hoxA*-containing 2.2-kb *PvuII-Ecl136II* fragment, originating from pGE400, was ligated into *SwaI*-cut pGE377 to yield the *hoxB-hoxC* overexpression vector pGE378 (Fig. 1).

The *hoxN* in-frame deletion allele was constructed as follows. A pCH231-derived 0.45-kb *FspI-SmaI* fragment containing the 3' region of *hoxN* was inserted into *EcoRI*-digested, end-polished pCH231. The resulting plasmid, designated pCH658, contains a *hoxN* allele in which 768 of 903 bp (85%) were deleted. For recombination into *R. eutropha* the *hoxNΔ* allele was subcloned as a 1.5-kb *SalI*-fragment into pLO2, yielding pCH659. The fusion sites in the *hoxNΔ* allele and in the PCR amplification products were verified by sequencing.

Plasmid pCH655, which was used for overexpression of *hoxC* in *E. coli*, was constructed by insertion of a 1.5-kb *PspI* (Klenow-treated) fragment harboring the *hoxC* sequence without the first two codons into the *Ecl136II* site of pQE-30.

Media and growth conditions. Strains of *R. eutropha* were grown in nutrient broth (NB), in a modified Luria broth (LB) with 0.25% sodium chloride (LSLB), or in mineral salts medium as described previously (12). Synthetic media for heterotrophic growth contained 0.4% (wt/vol) fructose (FN) or 0.2% (wt/vol) fructose and 0.2% (vol/vol) glycerol (FGN). Cultivation under lithoautotrophic

conditions was done in mineral salts medium under an atmosphere of hydrogen, carbon dioxide, and oxygen (8:1:1, vol/vol/vol). Sucrose-resistant segregants of *sacB*-harboring strains were selected on LSLB plates containing 15% (wt/vol) sucrose (27).

E. coli strains were grown in LB medium. Solid medium contained 1.2% (wt/vol) agar. Antibiotics were supplemented with the following: kanamycin (400 µg/ml) and tetracycline (15 µg/ml) for *R. eutropha* and kanamycin (25 µg/ml), tetracycline, (15 µg/ml), and ampicillin (100 µg/ml) for *E. coli*.

Gene replacement. The *hoxNΔ* allele was reintroduced into *R. eutropha* H16 via conjugation using the suicide vector pCH659. The allelic exchange procedure was based on the conditionally lethal *sacB* gene (27). The resultant sucrose-resistant isolates were screened for the presence of the desired mutation by amplification of the respective target sites as previously described (5). Deletion-carrying isolates were identified on the basis of the altered electrophoretic mobility of the amplification products. The resulting *hoxNΔ* strain HF375 served as the recipient for the *hoxJ*-containing suicide vector pCH615 to generate the isogenic *hoxNΔ hoxJ* strain HF459. Strain HF500, a derivative of HF371 (31), was isolated by the same recombination technique using pCH644 which contains the *hoxCΔ* allele (25).

Cell fractionation. Cells were disrupted in a French pressure cell, and the resulting crude extract was separated into soluble and membrane fractions as described earlier (19). Cytoplasmic and periplasmic fractions were separated by a modified version (4) of the procedure of Probst and Schlegel (36).

Immunogenic techniques. *E. coli* JM109 cells harboring the *hoxC* expression plasmid pCH655 were grown in LB medium at 30°C to an optical density at 600 nm (OD₆₀₀) of 0.8. Expression of His₆-HoxC was induced by adding IPTG (isopropyl-β-D-thiogalactopyranoside) to a final concentration of 1 mM. Cells were harvested after 3 h of induction and disrupted by two passages through a French pressure cell. Inclusion bodies which contained most of the His₆-HoxC protein were treated with 6 M guanidinium-HCl. Subsequently, the His₆-HoxC protein was purified using the Ni-nitrilotriacetic acid Spin Kit (Qiagen, Inc.) according to the manufacturer's instructions. Purified His₆-HoxC was used as antigen for immunization of rabbits (BioGenes GmbH, Berlin, Germany). Proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), transferred to Protran BA85 nitrocellulose membranes (Schleicher and Schuell), and identified immunologically according to a standard protocol (46). HoxC was detected with anti-HoxC serum, diluted 1:1,000, and an alkaline phosphatase-labeled goat anti-rabbit immunoglobulin G (Dianova, Hamburg, Germany).

Labeling with ⁶³NiCl₂. Labeling of hydrogenases with ⁶³NiCl₂ was essentially performed as previously described (5). Cells were grown in FGN medium in the presence of 120 nM of ⁶³NiCl₂ (6.38 mCi/ml; Amersham-Buchler). Soluble extracts were prepared and subjected to native PAGE. Gels were run in a continuous buffer system consisting of 90 mM Tris, 80 mM borate, and 2.5 mM EDTA (pH 8.3) at 200 V and 4°C for 2,500 V-h. After electrophoresis the gels were dried under vacuum and subjected to autoradiography using a SI 550 storage PhosphorImager (Molecular Dynamics).

Assays. Hydrogenase assays were performed with cells grown heterotrophically in FGN medium. SH (hydrogen-NAD⁺ oxidoreductase; EC 1.1.2.1.2) activity was determined by photometric recording of the H₂-dependent NAD reduction in the soluble fraction (39). MBH (hydrogen-acceptor oxidoreductase EC 1.18.99.1) and RH activities were photometrically measured in the membrane fraction using methylene blue as an electron acceptor (38). Amperometric H₂ uptake measurements using an H₂ electrode and methylene blue as an electron acceptor were done as previously described (35). For in-gel chromogenic detection of hydrogenase activity (5), soluble extracts were resolved on native PAGE gels as described above. The gels were subsequently incubated in H₂-saturated 50 mM potassium phosphate buffer (pH 5.5) containing 0.09 mM phenazine methosulfate and 0.06 mM nitroblue tetrazolium under an atmosphere of H₂ at 30°C. O₂ uptake assays were performed with whole cells using a Clark electrode (Rank Brothers Model 10). O₂ consumption was recorded amperometrically in 2.7 ml of H₂-saturated potassium phosphate buffer (50 mM, pH 7.0) at 30°C. Then, 200 µl of O₂-saturated water was added, and the reaction was started by the addition of 100 µl of cell suspension which was previously adjusted to an OD₄₃₆ of 11. H₂-independent O₂ consumption was monitored in N₂-saturated potassium phosphate buffer. β-Galactosidase assays were performed as described previously (54), and the activities (in units) were calculated according to the Miller method (33) except that cell density was measured at 436 nm. The level of protein in extracts was determined by the method of Lowry et al. (28).

RESULTS AND DISCUSSION

The RH and homologous H₂-sensing proteins form a subclass of [NiFe] hydrogenases. The regulatory region of the megaplasmid-borne hydrogenase gene complex in *R. eutropha* has previously been extended by three additional open reading frames (ORFs), designated *hoxB*, *hoxC*, and *hoxJ* (25, 26). The ORFs fill a gap between *hoxA*, the response regulator gene, and *hoxN*, the nickel permease gene. Database searches revealed similarity of the *hoxJ* product to histidine protein ki-

nases (26) and of HoxB and HoxC to [NiFe] hydrogenases, in particular to a small group of proteins which are present in aerobic H₂-oxidizing bacteria (Fig. 2). The closest relatives are the HoxB and HoxC proteins of *Alcaligenes hydrogenophilus* (26) and the HupU and HupV proteins of *R. capsulatus* (15) and *B. japonicum* (6), with sequence identities ranging from 53 to 79%.

HoxB and HoxC and their close relatives show typical signatures of standard [NiFe] hydrogenases (Fig. 2), represented by the prototypic periplasmic [NiFe] hydrogenase from *Desulfovibrio gigas* (51). The product of *hoxC* displays the conserved amino acid motifs which are considered as essential elements for the coordination of the NiFe cofactor (Fig. 2A) (2). The N-terminal RGxE motif (element 1) shows the regular spacing of 16 residues to the metal binding motif RxCGxCxxxH (element 2). It is worth noting, however, that the highly conserved histidine residue of element 2 is replaced by a glutamine residue in all HoxC-like proteins known so far (Fig. 2A). The conserved signature of the more variable element 3 is restricted to only two of five histidine residues. In element 4, GxxxPRGxxxxH, which is oriented to the C-terminal end of the polypeptide, an alanine substitutes for the highly conserved proline residue, whereas the C-terminal NiFe coordination site DPCxxCxxH (element 5) shows the perfect consensus of standard hydrogenases. Moreover, HoxC-like proteins terminate at a histidine residue and are devoid of a C-terminal extension (Fig. 2A). This observation suggests that, unlike the situation in most [NiFe] hydrogenases (29), the regulatory proteins do not undergo a proteolytic cleavage prior to metalcenter insertion and subunit oligomerization. The lack of a C-terminal tail has also been reported for the CO-induced hydrogenase from *Rhodospirillum rubrum* and for the Ech hydrogenase from *Methanosarcina barkeri*. However, these enzymes are related to hydrogenase 3 of *E. coli* and are considered to be involved in H₂ evolution rather than in H₂ sensing (17, 32).

HoxB and its homologues have potential coordination sites for three FeS clusters similar to the small HynB subunit of *D. gigas* (Fig. 2B). Four cysteines (Cys27, -30, -130, and -178) correlating with the ligands of the proximal [4Fe-4S] cluster (P), are present in the N-terminal region of HoxB. Three cysteine residues (Cys220, -240, and -247) and one histidine (His217) coincide with the ligands of the distal [4Fe-4S] cluster (D) in HynB. Interestingly, HoxB and its close relatives reveal four instead of three conserved cysteines (Cys256, -267, -274, and -277) for binding the putative medial FeS cluster (M), suggesting that the common [3Fe-4S] center might be replaced by a [4Fe-4S] cluster in this group of proteins. Most notably, HoxB and its homologues lack the N-terminal leader sequence which directs the export of periplasmic and membrane-bound hydrogenases (3, 34). This observation points to a cytoplasmic location of these proteins. Another interesting structural feature of HoxB-like proteins is a C-terminal peptide of 54 to 55 amino acids which is not present in the periplasmic HynB protein. Although the small subunits of membrane-bound hydrogenases also carry an extension of about 50 amino acids, including a stretch of 22 hydrophobic amino acids and a highly conserved histidine residue, the primary structure is clearly distinct from the tail of the HoxB-type proteins (Fig. 2B). It has been shown that the C-terminal domain plays a pivotal role in anchoring the membrane-bound hydrogenases to the membrane and in coupling the proteins to the primary electron acceptors, the *b*-type cytochromes (4, 21). In analogy, it seems likely that the C-terminal domain of HoxB-like proteins links the regulatory hydrogenases to their specific partners, namely, the histidine protein kinases HoxJ and HupT, respectively (16,

B

<i>R. eu.</i>	HoxB	-----MNA P VCTGLASAK P G	15
<i>A. hy.</i>	HoxB	-----MNPNDV K R P GR P	12
<i>R. ca.</i>	HupU	-----	0
<i>B. ja.</i>	HupU	-----MSRSD G	6
<i>D. gi.</i>	HynB	MKCY I GRG K DQVEER L ERRG V SR R DFMK F CTAVAVAMGM G PA F AP K V A EAL T AK K	55
<i>R. eu.</i>	HoxK	MVETFYEV M RR Q GISRR S FL K Y C SL T AT S L G L G P S L P Q T A H A M E T K P	48
▲			
<i>R. eu.</i>	HoxB	VLN V L W I Q S G C G C S M S L L C A D T D F T G M L K S A G I H M L W H P S L S L E S G V E Q L Q I	70
<i>A. hy.</i>	HoxB	HF N V L W L Q S G C C G C S M S L L C A D S A D F F G S L Q D A G I N M L W H P S L S L E T G A D L Q I	67
<i>R. ca.</i>	HupU	-MK V L W L Q A S C G C T M S A L C A E A P D L I D T L A T A G V E F L W H P A L S L A T G G E V R Q L	54
<i>B. ja.</i>	HupU	T T N V L W L Q A S C G C T M S I L E S G A S G W F D E L R Q F G I N L L W H P S V S E E T G E A V E V	61
<i>D. gi.</i>	HynB	R P S V V Y L H N A C T C S E S L L R T V D P-Y V D E L I L D V I S M D Y H E T L M A G A G H A V E A	109
<i>R. eu.</i>	HoxK	R T P V L W L H G L E C T C S E S F I R S A H P -L A K D V V L S M I S L D Y D D T L M A A A G H Q A E A	102
* * * * * *			
<i>R. eu.</i>	HoxB	LE D C L Q R V A L H A L C V E G A M L R G P H G T G R F H L L A G T G V P M L E W S R L A A V A D Y T L	125
<i>A. hy.</i>	HoxB	L R A C A E G E V P L H A L C I E G S M L R G P N G S G R F H M L A G T G K P L D W V E A L A G M A D Y T I	122
<i>R. ca.</i>	HupU	L Q A L E A G E I A L D C L A V E G A I A R G P M G T G R F Q M L S G T G R S M L D W R A L A P L A G H V	109
<i>B. ja.</i>	HupU	L Q S V L D G K V Q L D L L L L E G S V A R G P N D S G R F N M L A G T N R S V Y R W M L D L A P L A D Y V I	116
<i>D. gi.</i>	HynB	L H E A I K G D F V C---V I E G G I ---P M G D G G Y W G K V G R-R N M Y D I C A E V A P A K A K A I	157
<i>R. eu.</i>	HoxK	L E E I M T K Y K G N Y I L A V E G N P ---P L N Q D G M S C I I G -G R P F I E Q L K Y V A K D A K A I	153
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<i>R. eu.</i>	HoxB	AV G T C A A Y G G I T A G G N P T D A C G L Q Y E G D Q P G L L G L N Y R S R A G L P V I N V A G C P T	180
<i>A. hy.</i>	HoxB	AV G T C A A Y G G I T A A G N P T D A C G L Q Y D G D Q V G L L G A D Y R S R S G L P V I N S C P T	177
<i>R. ca.</i>	HupU	AV G S C A A Y G G V T S A G G N P S D A V G L A F E G A H P G G V L A E F R A R S G L P V V N I A C P T	164
<i>B. ja.</i>	HupU	AV G S C A A Y G G V P A A G S N P T D A V G L Q F E G S D S G G A L G A G F R S R L G L P V I N V A G C A P	171
<i>D. gi.</i>	HynB	A I G T C A T Y G G V Q A A K N P T G T V G V N-----E A L G -----K L G V K A I N I A C P P	200
<i>R. eu.</i>	HoxK	SW G S C A S W G C V Q A A K N P T Q A T P V H K V I T D -----K P I K V P G P P	194
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<i>R. eu.</i>	HoxB	H P G W V T D A L A L S A R L L T A S D L D T L G R P R F Y A D Q L V H G C T R N E Y E ---F K A S -	231
<i>A. hy.</i>	HoxB	H P N W V T D T L M A L A A G M P D A S Q I D P L G R P R F Y A D Q L V H G C T R N E F Y E ---F K A S -	228
<i>R. ca.</i>	HupU	H P G W V T E T L M L A R G H L A A D L D A L G R L F Y A Q H L V H G C P R N E F Y E---Y K A S -	215
<i>B. ja.</i>	HupU	H P G W M M E T I L A L T S K D L A A T D L D G Y G R P K F I A N H L A H G C S R N E F Y E ---F K A S -	222
<i>D. gi.</i>	HynB	N P M N F V G T V V H L L T K G M--P E L D R Q G R P V M F F G E T V H D N C P R L K H F E A G E F A T S F	253
<i>R. eu.</i>	HoxK	T A E V M T G V I T Y M L T F D R I -P E L D R Q G R P K M F Y S Q R H D K V Y R R P H F D A G Q F V E W	248
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<i>R. eu.</i>	HoxB	A E K P S D L G -C M M E N M G C K G T Q A H A D C N T R L W N G E G S-C T R G G Y A C I S C T E P G F E E	284
<i>A. hy.</i>	HoxB	A E K P S D L G -C M M E H M G C K G T Q A H G D C N T R L W N G D G S-C T R G G Y A C I S C T E P G F E E	281
<i>R. ca.</i>	HupU	A L Q L S D L G -C M M E H L G C V G T Q A V G D C N I R P W N G E G S-C T R G G Y F C I A C T A P E F E E	268
<i>B. ja.</i>	HupU	A E T M S E R G -C L M E H L G C K A T Q A V G D C N Q R S W N G E G S-C T R G G Y A C I A C T S P G F E G	275
<i>D. gi.</i>	HynB	G S P E A K K G Y C L Y E-L G C K G P D T Y N N C P K Q L F N Q-V N W P V Q A G H F C I A C S E P N F W D	306
<i>R. eu.</i>	HoxK	D D E S A R K G F C L Y K-M G C K G P T T Y N A C S T R W N E G T S F P I Q S G H C I G C S E D G F W D	302
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<i>R. eu.</i>	HoxB	P G H P F H Q T P K V A G I P I G L P T D M P K A W F V A L A S L S K S A T P K R V R L N A T A D H P L I A P A	339
<i>A. hy.</i>	HoxB	P G H P F A L T P K I A G I P I G L P T D M P K A W F V A L A S L S K S A T P K R V R Q N A T A D H R V V V P A	336
<i>R. ca.</i>	HupU	P R H P F T E T P K V A G I P V G L P A D M P K A W F M A L A S L S K A A T P E R I A K N A V A P R L T V P P T	323
<i>B. ja.</i>	HupU	A Q N-F L E T A K L A G I P V G L P T D M P K A W F V A L A L S K S A T P R R V R L N A T A D H V V V P P G	329
<i>D. gi.</i>	HynB	L Y S P F Y S A -----	314
<i>R. eu.</i>	HoxK	K G S-F Y --D R L T G I S Q F G V E A N A D K I G G T A S V V G A V T A H A A S A I K R A S K K N E T	354
*			
<i>R. eu.</i>	HoxB	A I R K T R L K -	347
<i>A. hy.</i>	HoxB	A I R K T R L K -	344
<i>R. ca.</i>	HupU	T I R K P G G R R	332
<i>B. ja.</i>	HupU	G R S T A K G K P	338
<i>D. gi.</i>	HynB	-----	314
<i>R. eu.</i>	HoxK	S S G S E H ---	360

FIG. 2—Continued.

signal transduction (44). Since PAS domains monitor changes in light, redox potential, oxygen, and the general energy status of the cell, the HoxJ sensor kinase is the favorite candidate to receive and convert the signal from the RH.

The outstanding biological role of H₂-sensing hydrogenases (6, 15, 25), obviously based on a series of individual structural features, places this group of regulatory proteins into a distinct subclass of [NiFe] hydrogenases. Public databases provided evidence for the occurrence of this group of hydrogenases in *Rhodobacter sphaeroides* (20) and *Oligotropha carboxydovorans* (37). Moreover, one of the three putative hydrogenases of *Aquifex aeolicus* shows characteristics of HoxBC-like proteins, pointing to a possible H₂-sensing process in this phylogenetically ancient bacterium (10). Thus, regulatory hydrogenases may be more common than originally envisaged.

The RH of *R. eutropha* is a soluble H₂-oxidizing protein located in the cytoplasm. The structural similarity to hydrogenases and its role in H₂-sensing implied the fundamental questions whether the RH catalyzes H₂ oxidation and if so, whether the hydrogenase activity is necessary for its regulatory function. It was not possible to discriminate the RH activity in wild-type cells clearly from the MBH- and SH-derived hydrogenase activities. To exclude assay interferences the MBH⁻ SH⁻ RH⁺ strain HF371, which carries in-frame deletions in the large subunit genes of the MBH (*hoxGΔ*) and the SH (*hoxHΔ*), was grown under heterotrophic, hydrogenase-derepressing conditions and then tested for H₂-oxidizing activity. Soluble extracts of this mutant showed a low level of H₂ uptake activity (1.5 nmol H₂/min/mg of protein) measured amperometrically with methylene blue as the electron acceptor. However, the activity

was clearly above the background level (<0.5 nmol of H_2 /min/mg of protein) of extracts derived from strain HF500, which is disrupted in all three hydrogenase proteins. Notably, the $MBH^- SH^- RH^+$ strain HF371 was not able to grow autotrophically with H_2 as the energy source. These results indicate that the RH is either formed at an extremely low level and/or that the protein exhibits only poor hydrogenase activity, a finding which is in agreement with its regulatory role.

To determine the cellular localization of the RH protein, cells of the $MBH^- SH^- RH^+$ strain HF371 were separated into the cytoplasmic, periplasmic, and membrane fractions. In order to detect the RH immunologically a hexahistidine-tagged variant of the HoxC protein was purified from *E. coli* to raise a polyclonal antiserum. Immunoblots developed with this antiserum gave a faint band corresponding to a 52-kDa protein which was exclusively present in the cytoplasmic fraction (data not shown). The size of this protein was in good agreement with the molecular mass of 52.4 kDa predicted for HoxC. A HoxC signal was absent in extracts of the control strain HF500, which lacks all three hydrogenases, showing that the antiserum is specific for HoxC (data not shown). These immunological data are completely in line with the prediction for a cytoplasmic location of the RH deduced from the primary sequence. Since dihydrogen is a freely diffusible molecule, there is no need for the cell to anchor the H_2 -sensing protein to the membrane.

Homologous overproduction of the RH protein. To get further insight into the biochemical properties of the H_2 sensor, the intracellular level of the protein had to be increased. This was achieved by overexpressing the native *hoxB* and *hoxC* genes in *R. eutropha* under the control of the SH promoter (P_{SH}), which directs transcription of the SH operon (42, 54). The HoxA-controlled, homologous system has the advantage that the Hyp proteins, which are required for metallocenter assembly (11), are potentially available for RH maturation. Moreover, P_{SH} is a well-characterized, relatively strong promoter, and the putative ribosome-binding site of the first gene *hoxF* of the SH operon is in perfect agreement with the consensus in *E. coli* (42, 54).

The construction of the expression vector is based on three steps, which are described in detail in Materials and Methods. (i) The native *hoxB* and *hoxC* genes were tandemly fused to a modified SH promoter region yielding plasmid pCH594 (Fig. 1). (ii) A fragment containing the P_{SH} -*hoxBC* fusion was transferred to the broad-host-range vector pEDY309 that replicates stably in *R. eutropha*. (iii) To enhance transcription from P_{SH} , a copy of the *hoxA* activator gene, governed by the *lac* promoter, was inserted into pGE377, resulting in the expression vector pGE378 (Fig. 1). Ongoing research in our laboratory showed that the vector system is also suitable for a general application (3, 8, 10). A moderate expression of the cloned genes in slowly growing cells obviously prevents the occurrence of toxic effects and the formation of inclusion bodies.

To estimate the effectiveness of the overexpression system, pGE378 was introduced into strain HF371. The resulting transconjugant was grown under hydrogenase-derepressing conditions, and the cells were fractionated into membrane, cytoplasmic, and periplasmic extracts. Immunological analysis showed that the level of HoxC was enhanced significantly and that the protein was located in the cytoplasm (data not shown). A 40-fold increase in hydrogenase activity (58.6 nmol H_2 /min/mg of protein) was obtained with HF371(pGE378) in comparison with the control strain HF371(pEDY309) (1.5 nmol of H_2 /min/mg of protein). Nevertheless, even the enhanced RH activity did not support autotrophic growth of the strain with H_2 as the sole energy source, again indicating that the RH is not coupled to an energy-generating electron trans-

port process. This conclusion is consistent with the observation that the O_2 uptake rates of the strains HF500 ($MBH^- SH^- RH^-$, HF371 ($MBH^- SH^- RH^+$), and HF371(pGE378) ($MBH^- SH^- RH^{++}$) remained constant at a basal level (30 nmol of O_2 /min/mg of protein) upon addition of H_2 , whereas the O_2 uptake rate of the MBH^- and SH^- -harboring wild-type cells increased significantly under these conditions from 30 to 120 nmol of O_2 /min/mg of protein. The results do not unambiguously show that the RH has no potential for providing energy for growth, since the experiment did not exclude the possibility that the RH is linked to an unknown electron transport component that was not overexpressed by the vector system used.

The function of the RH is nickel dependent. The structural similarity with [NiFe] hydrogenases and the ability to react with H_2 raised the question as to whether the enzymatic and the regulatory functions of the RH depend on nickel. The RH-overproducing strain HF371(pGE378) was grown under hydrogenase-derepressing conditions in the presence of various concentrations of $NiCl_2$. Soluble extracts were prepared, and the proteins were separated by native PAGE and subjected to a hydrogenase-specific activity staining assay with phenazine methosulfate as the electron acceptor. The RH activity strictly correlated with the addition of nickel to the growth medium (Fig. 3A). In the presence of 1 μ M $NiCl_2$ the dye reaction was very intense and did not occur with cells cultivated under nickel starvation. The quantitative RH data obtained from extracts with methylene blue as the electron acceptor (Fig. 3A) supported the conclusions drawn from the staining assay. The occurrence of double bands in native PAGE gels indicates that the RH displays different conformations. We observed that the ratio of the two bands varied with respect to the preparation. From gel filtration experiments we obtained evidence that the slowly migrating band correlates with a tetramer consisting of two RH moieties and that the rapidly migrating band correlates with the sole RH dimer (M. Bernhard and B. Friedrich, unpublished results).

Preliminary Fourier transform infrared (FTIR) spectroscopy analyses have indicated that the active site of the RH is similar to that of standard [NiFe] hydrogenases (35). To determine whether the RH is a nickel-containing protein, we tested the accumulation of ^{63}Ni by autoradiography (5, 11). In fact, the soluble extract of the overproducing strain HF371(pGE378) developed a strong ^{63}Ni signal, and a faint signal occurred below the strong one (Fig. 3B, lane 4). The migration behavior and, to a lesser extent, the relative intensity of the two signals correlated with the two bands obtained in the activity staining assay (Fig. 3A, lane 3). Only traces of a ^{63}Ni signal were recognized with cells which produced the RH at a normal level (Fig. 3B, lane 3). Interestingly, the RH-specific signal decreased in extracts of cells containing intact MBH^- and SH^- proteins, indicating competition of the three hydrogenases for ^{63}Ni or for the pleiotropically acting Hyp proteins which are required for metallocenter assembly (11, 29).

Experiments to test whether the H_2 -sensing function is also nickel dependent are not trivial since only a low intracellular level of the RH is probably instrumental in the regulatory process. Therefore, traces of nickel might be sufficient for the regulatory function of the RH. In order to demonstrate a correlation between nickel supply and the level of hydrogenase induction, we had to expose the cells to conditions of severe nickel limitation. This was achieved by using *R. eutropha* HF459, a derivative of strain HF433, which carries a deletion in the high-affinity nickel permease gene *hoxN*, and by addition of the chelating agent nitrilotriacetic acid. Since the MBH^- and SH^- promoters are regulated coordinately (42), β -galactosidase

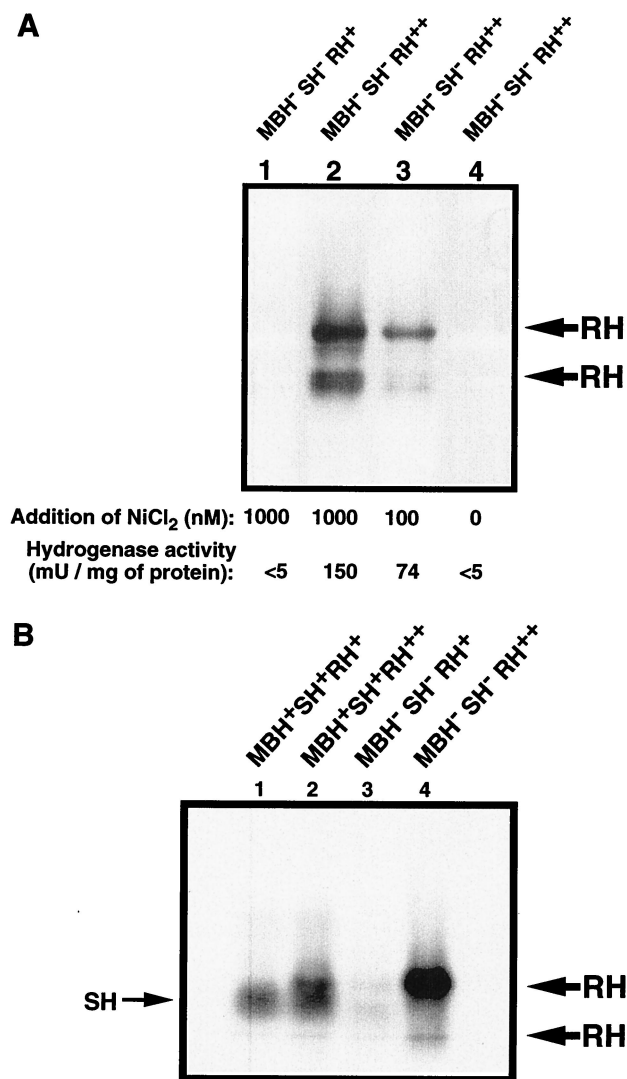


FIG. 3. (A) In-gel RH activity staining. Soluble extracts prepared from cells grown in FGN medium in the presence of various NiCl₂ concentrations were separated on 4 to 15% native PAGE gels (40 μg of protein in each lane). The gel was soaked for 3 h in H₂-saturated 50 mM potassium phosphate buffer (pH 7.0) containing 0.09 mM phenazine methosulfate and 0.06 mM nitroblue tetrazolium under a hydrogen atmosphere. Methylene blue reducing activity (lower part of the figure) was measured photometrically in soluble extracts at pH 7.0 using methylene blue as an electron acceptor. Lane 1, HF371(pEDY309); lanes 2 to 4, HF371(pGE378). (B) ⁶³Ni incorporation into the RH in vivo. Cells were grown in FGN medium in the presence of 120 nM ⁶³NiCl₂. Soluble extracts were separated in a 4 to 15% native PAGE gel (200 μg of protein on each lane). Lane 1, H16(pEDY309); lane 2, H16(pGE378); lane 3, HF371(pEDY309); lane 4, HF371(pGE378). RH⁺⁺, RH overproduction.

activity was monitored representatively with a plasmid-based Φ (*hoxK'*-*lacZ*) fusion. The result is illustrated in Fig. 4 and shows a clear correlation between β -galactosidase activity and supplementation of NiCl₂ to the medium. As expected, the expression level was low when H₂ was omitted, even with nickel excess. The conclusion that nickel is essential for H₂ recognition by the RH protein is confirmed by the behavior of strain HF375, in which the H₂-dependent signal transduction is interrupted due to a lesion in the sensor kinase (Table 1). This strain displayed high-level hydrogenase gene expression independently of H₂ and hence independently of nickel (Fig. 4).

It is obvious from these results that the regulatory function

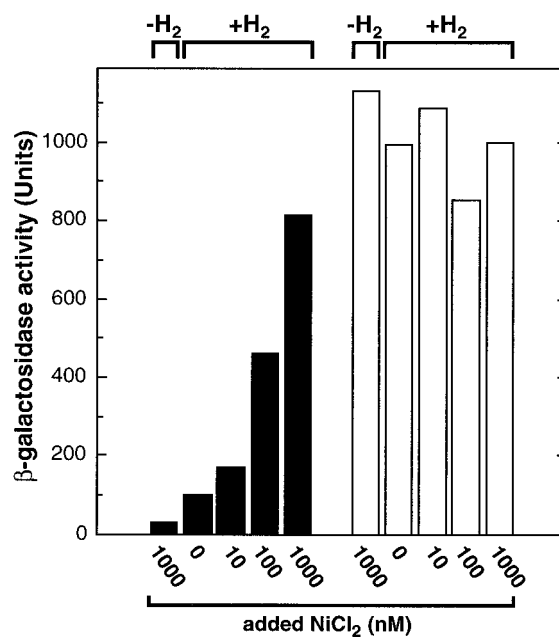


FIG. 4. Hydrogenase gene expression in the presence of various NiCl₂ concentrations. The *R. eutropha* strains HF459 (HoxN⁻; black bars) and HF375 (HoxN⁻ HoxJ⁻; white bars), each containing the plasmid-based Φ (*hoxK'*-*lacZ*) fusion pGE301, were grown in FGN medium containing 10 μM nitrilotriacetic acid at the nonpermissive temperature of 37°C until they reached an OD₄₃₆ of 7. At time zero the cells were shifted to 30°C and 10% (vol/vol) H₂ (+H₂) or 10% (vol/vol) N₂ (-H₂) was added. After further incubation for 5 h the cells were collected, and the β -galactosidase activity was determined.

of the RH relies on the availability of nickel in the medium. For HupUV of *B. japonicum* it was proposed that the protein acts as a sensor of nickel and/or of oxygen and hydrogen (6). The content of hydrogenase-specific mRNA in *B. japonicum* correlated with the addition of nickel to the medium showing that nickel acts as an effector of transcriptional regulation (22). Our experiments revealed that only trace amounts of the metal were required for the H₂-sensing function of the RH in *R. eutropha*, suggesting that nickel is a tightly bound component of the active site rather than an effector of regulation.

The results of this study, together with previous findings regarding the electron paramagnetic resonance and FTIR properties of the RH (35) and the H₂-binding capacity of the HupUV protein from *R. capsulatus* (49), indicate that H₂ recognition is based on a nickel-iron-containing active site similar to that of standard [NiFe] hydrogenases, including two CN⁻ groups and one CO molecule at the iron site (35). The capability of the sensors to oxidize H₂ does not yet allow the conclusion that H₂ binding is intimately connected with a redox reaction. Thus, elucidation of the underlying mechanism of H₂ signal transduction is a fascinating subject of current research in our laboratory.

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