

A Novel CO-Responsive Transcriptional Regulator and Enhanced H₂ Production by an Engineered *Thermococcus onnurineus* NA1 Strain

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Genome analysis revealed the existence of a putative transcriptional regulatory system governing CO metabolism in *Thermococcus onnurineus* NA1, a carboxydophilic hydrogenogenic archaeon. The regulatory system is composed of CorQ with a 4-vinyl reductase domain and CorR with a DNA-binding domain of the LysR-type transcriptional regulator family in close proximity to the CO dehydrogenase (CODH) gene cluster. Homologous genes of the CorQR pair were also found in the genomes of *Thermococcus* species and “*Candidatus* Korarchaeum cryptofilum” OPF8. In-frame deletion of either *corQ* or *corR* caused a severe impairment in CO-dependent growth and H₂ production. When *corQ* and *corR* deletion mutants were complemented by introducing the *corQR* genes under the control of a strong promoter, the mRNA and protein levels of the CODH gene were significantly increased in a Δ CorR strain complemented with integrated *corQR* (Δ CorR/*corQR*⁺) compared with those in the wild-type strain. In addition, the Δ CorR/*corQR*⁺ strain exhibited a much higher H₂ production rate (5.8-fold) than the wild-type strain in a bioreactor culture. The H₂ production rate (191.9 mmol liter⁻¹ h⁻¹) and the specific H₂ production rate (249.6 mmol g⁻¹ h⁻¹) of this strain were extremely high compared with those of CO-dependent H₂-producing prokaryotes reported so far. These results suggest that the *corQR* genes encode a positive regulatory protein pair for the expression of a CODH gene cluster. The study also illustrates that manipulation of the transcriptional regulatory system can improve biological H₂ production.

Carbon monoxide (CO) serves as a central metabolic intermediate in anaerobic metabolism (1), as an enzyme metallo-center ligand (2, 3), as a physiologically significant signal in higher organisms (4), and as a speculative component in an early mode of metabolism and the origin of life (5). CO can be utilized as carbon and energy sources for growth by numerous microorganisms containing carbon monoxide dehydrogenase (CODH), a key enzyme in CO metabolism. CODH oxidizes CO to carbon dioxide (CO₂), and the electrons generated by the process are coupled to diverse reactions, such as oxygen reduction, desulfurification, hydrogenogenesis, acetogenesis, and methanogenesis (6). When CO is aerobically oxidized, as in *Pseudomonas thermocarboxydovorans* and *Oligotropha carboxidovorans*, the reducing equivalents are transferred to oxygen through a CO-insensitive respiratory chain. Under anaerobic conditions, CO oxidation is linked to acetate production through the reductive acetyl coenzyme A (acetyl-CoA) or Wood-Ljungdahl pathway in acetogenic bacteria and to methane production as a substrate of CODH/acetyl-CoA synthase in methanogenic archaea. Carboxydophilic hydrogenogens like *Rhodospirillum rubrum* and *Carboxydotherrmus hydrogenoformans* oxidize CO through a water-gas shift reaction, CO + H₂O → CO₂ + H₂ ($\Delta G^{\circ} = -20$ kJ/mol), which produces hydrogen gas (7–9).

There are several distinct CO regulation systems known for aerobic and anaerobic CO oxidation (1, 10). Aerobic CO oxidation systems are encoded by *cox* genes (*coxMSL*) and transcriptionally regulated by CoxC, CoxH, or RcoM (11, 12). The CoxC and CoxH proteins have a LytTR DNA-binding domain and an MHYT sensor domain of six transmembrane segments with conserved His and Met residues (13). The LytTR DNA-binding domain is unique, in that it is mainly comprised of β strands and predominantly found in pathogenic bacteria (14). RcoM has a LytTR domain and an N-terminal heme-bearing PAS sensor do-

main (12). The PAS domain is a well-known signal transduction module and senses various environmental signals, such as gases (O₂, NO, or CO), light, redox potential, voltage, xenobiotics, and nitrogen availability (15). There is a second type of aerobic CO oxidation system which is found in the Gram-negative carboxydobacteria *P. thermocarboxydovorans* and *Hydrogenophaga pseudoflava* and a Gram-positive bacterium, *Mycobacterium* sp. strain JC1 DSM 3803 (16, 17). The system is encoded by *cut* genes (*cutBCA* or *cutMSL*) and is regulated by the CutR protein, a LysR-type transcriptional regulator (LTTR) with a plausible helix-turn-helix (HTH) motif in the N-terminal domain and a LysR substrate binding domain in the central and C-terminal domains (18). LTTRs, as some of the most common positive regulators in prokaryotes, govern very diverse genes and physiological functions (19).

Anaerobic CO oxidation systems are encoded by *coo* genes,

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which encode CODH (*cooS*), a ferredoxin-like protein (*cooF*), a multisubunit hydrogenase (*cooMKLXUH*), maturation proteins for CODH and hydrogenase, and a transcriptional activator, *CooA* (20). *CooA* is a homodimeric heme-binding CO sensor belonging to the cyclic AMP receptor protein (CRP) family of transcriptional regulators (21). When CO is present, *CooA* binds to CO and undergoes a conformational change to transcriptionally activate the required *coo* genes for CO oxidation (21). *CooA* is also a redox sensor so that *CooA* is not activated in the presence of oxygen (high redox potential) (10).

Two hyperthermophilic archaea, *Thermococcus* sp. strain AM4 and *Thermococcus onnurineus* NA1, can hydrogenogenically grow on CO, and CODH gene clusters are present in their genomes (22–24). The CODH gene is clustered with hydrogenase genes similar to the *coo* gene cluster of *R. rubrum*, but the primary structure and the organization of the genes are considerably different from those of the *coo* gene cluster (24). Additionally, Na^+/H^+ antiporter genes are present in the hyperthermophilic archaea but not in *R. rubrum*. A gene encoding a putative transcriptional regulator is also present in the upstream region of the CODH gene, but it does not encode a *CooA* homolog (25). We previously demonstrated that the CODH catalytic subunit and hydrogenase large subunit encoded by this CODH gene cluster are essential for the carboxydrotrophic hydrogenogenic metabolism in *T. onnurineus* NA1 (26).

In this study, we describe a novel type of CO-responsive regulatory system (CorQR) in *T. onnurineus* NA1 through bioinformatic analysis, characterization of in-frame deletion mutants, and transcriptional analysis. This study also illustrates that the manipulation of the regulatory circuit can improve CO-dependent H_2 production.

MATERIALS AND METHODS

Strain, media, and culture conditions. *T. onnurineus* NA1 (KCTC10859) was isolated from a deep-sea hydrothermal vent area in the Papua New Guinea-Australia-Canada-Manus (PACMANUS) field (27). This strain was routinely cultured in yeast extract-peptone-sulfur (YPS) medium as previously reported (27). Modified medium 1 (MM1) (23, 28) was prepared with 1 g liter⁻¹ yeast extract, 35 g liter⁻¹ NaCl, 0.7 g liter⁻¹ KCl, 3.9 g liter⁻¹ MgSO₄, 0.4 g liter⁻¹ CaCl₂·2H₂O, 0.3 g liter⁻¹ NH₄Cl, 0.15 g liter⁻¹ Na₂HPO₄, 0.03 g liter⁻¹ NaSiO₃, 0.5 g liter⁻¹ NaHCO₃, 0.5 g liter⁻¹ cysteine-HCl, and 0.001 g liter⁻¹ resazurin. One milliliter liter⁻¹ of Holden's trace elements/Fe-EDTA solution (29) and 1 ml liter⁻¹ of Balch's vitamin solution (30) were added as supplements to the medium. After autoclaving, the medium was kept in an anaerobic chamber (Coy Laboratory Products, Grass Lake, MI) filled with an anoxic gas mixture (N₂, H₂, CO₂, 90:5:5) for equilibration, and the final pH of the medium was adjusted to 6.5 with 2 N HCl.

For the cultures in serum bottles, the media were reduced with 0.005% Na₂S·9H₂O, and the headspaces were filled with 100% CO (MM1-CO) or 5 g liter⁻¹ sodium pyruvate was provided to support the growth of the *corQ* and *corR* mutant strains (MM1-pyruvate). The serum bottles were sealed with bromobutyl rubber stoppers and aluminum crimp caps.

For the pH-stat batch culture, *T. onnurineus* NA1 was cultured in a 100-ml serum bottle and then 3-liter bioreactors (Fermentec, Cheongwon, Republic of Korea), and the working volumes were 50 ml and 2 liters, respectively, at 80°C. For the cultures in bioreactors, MM1 was supplemented with 10 g liter⁻¹ yeast extract and a 10 times greater amount of Holden's trace elements/Fe-EDTA solution. Bioreactors were sparged with pure argon gas (99.999%) through a microsparger. The agitation speed was 300 rpm, and the pH was maintained at 6.1 to 6.2 using 0.2 M NaOH in 3.5% NaCl. The inlet gas of 100% CO was supplied by using a

mass flow controller (MKPrecision, Seoul, Republic of Korea) at a feeding rate of 400 ml min⁻¹. The gas outlet was open to let the H₂ and CO₂ gases escape and maintain the total pressure at 10⁵ Pa.

Bioinformatic analysis. The open reading frame (ORF) was predicted using the Glimmer (version 3.02) program (http://www.ncbi.nlm.nih.gov/genomes/MICROBES/glimmer_3.cgi). An homology search was performed using a search with the Basic Local Alignment Search Tool (BLAST; <http://blast.ncbi.nlm.nih.gov/Blast.cgi>) against the nonredundant protein database from the National Center for Biotechnology Information (NCBI). Multiple-sequence alignment was performed using ClustalW2 (<http://www.ebi.ac.uk/Tools/msa/clustalw2>) of the European Bioinformatics Institute (EBI).

Analytical methods. Cell growth was monitored by measuring the optical density at 600 nm (OD₆₀₀) with a BioPhotometer plus UV-visible spectrophotometer (Eppendorf, Hamburg, Germany). Biomass was determined on the basis of the fact that the unit value of the OD₆₀₀ corresponded to 0.361 g (cell dry weight) liter⁻¹. The H₂ production rate was calculated on the basis of the H₂ content in the gases produced from a bioreactor, and the gas flow rate was measured with a wet gas meter (Shinagawa, Tokyo, Japan). The amounts of CO, H₂, and CO₂ were measured by using a YL6100 gas chromatograph (GC; YL Instrument Co., Anyang, Republic of Korea) equipped with a Molsieve 5A column (Supelco, Bellefonte, PA), a Porapak N column (Supelco), a thermal conductivity detector, and a flame ionization detector. Argon was used as the carrier gas at a flow rate of 30 ml min⁻¹.

Reverse transcription-quantitative PCR (RT-qPCR). RNA was prepared with the TRIzol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer's instructions, with some modifications. Genomic DNA was eliminated by DNase I (Thermo Scientific Fermentas, St. Leon-Rot, Germany). One microgram of RNA was incubated with 1 unit of DNase I at 37°C for 30 min and purified through chloroform extraction and ethanol precipitation. RNA was quantified with a spectrophotometer, and cDNA was created using Moloney murine leukemia virus (M-MuLV) reverse transcriptase (Thermo Scientific Fermentas, St. Leon-Rot, Germany). One microgram of RNA was incubated with 40 units of reverse transcriptase, 5 μM random hexamers, and 1 mM deoxynucleoside triphosphate (dNTP) at 37°C for 1 h in reverse transcription buffer (1×, as supplied by the enzyme manufacturer). The reaction products were serially diluted to find an adequate concentration for real-time PCR analysis, and the samples were amplified with SYBR green real-time PCR master mix (Toyobo, Osaka, Japan). Amplified signals were detected using a StepOnePlus system (Applied Biosystems, Foster City, CA), and all primers used are listed in Table S1 in the supplemental material. The relative amount of transcript for each gene was calculated from the cycle threshold (C_T) values using a relative standard curve after normalization to the corresponding 16S rRNA (TON_1979) quantity.

Genetic manipulation. Mutant strains with either complementation of *corQR* or an in-frame deletion of *corQ* or *corR* were constructed by modifying the gene disruption system used for a hyperthermophilic archaeon, *Thermococcus kodakarensis* KOD1 (31) (see Fig. S1 in the supplemental material). Both *corQ* and *corR* deletion mutants were generated by unmarked in-frame deletion through homologous recombination, and deletions were verified by PCR using the primers listed in Table S1 in the supplemental material.

For the complementation of *corQ* and *corR* deletion mutants, the pQRc vector was constructed (see Fig. S2 and Table S1 in the supplemental material). The intergenic region between TON_1126 and TON_1127 was chosen for the gene integration site because the region has a relatively low transcription level, on the basis of deep sequencing data (data not shown). P_{TON_0157}, the promoter of TON_0157 (encoding a glutamate dehydrogenase), which shows a high transcription level in MM1-CO (data not shown), was used for high-level expression of the *Pyrococcus furiosus* *hmg* (*hmg_{Pfu}*) cassette. *corQR* genes were amplified by PCR using the primers listed in Table S1 in the supplemental material, and the am-

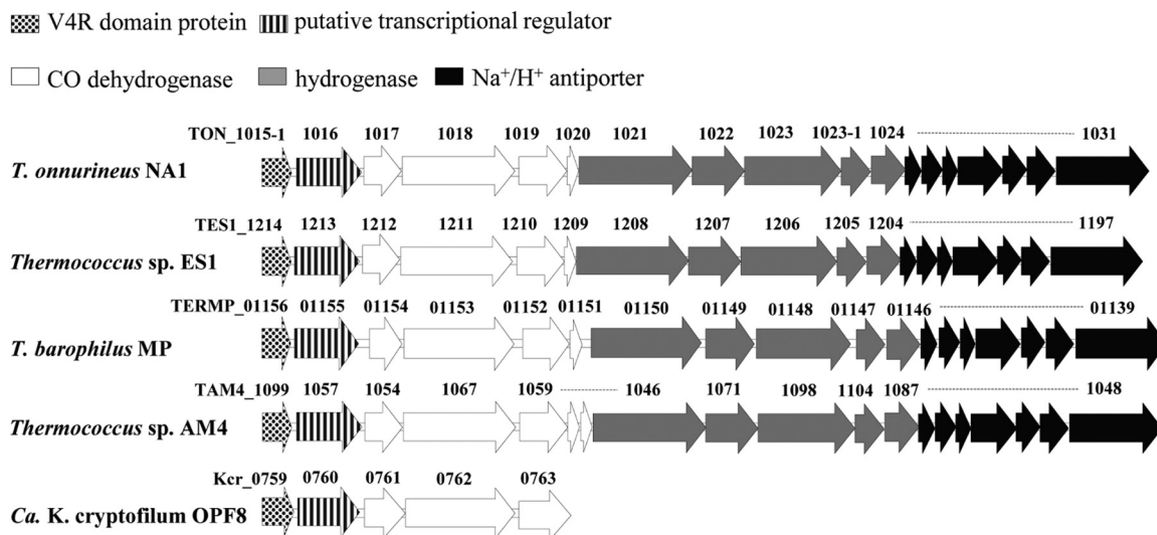


FIG 1 Gene organization of CODH gene clusters in *T. onnurineus* NA1, *Thermococcus* sp. ES1, *T. barophilus* MP, *Thermococcus* sp. AM4, and “*Ca. Korarchaeum cryptofilum*” OPF8. TON_1015-1 was added as an ORF on the basis of the findings of this study. Locus tag information for every gene is indicated as a number above each corresponding arrow.

plified product was inserted in the downstream region of the *hmgP_{flu}* cassette.

Western blotting. Chemiluminescent signals created by Immuno-Star horseradish peroxidase (HRP; Bio-Rad, Hercules, CA) were detected by a ChemiDocMP imaging system (Bio-Rad, Hercules, CA). Western blots were prepared and analyzed using a chemiluminescent dye with an Immuno-Star HRP chemiluminescent kit (Bio-Rad, Hercules, CA). Antibodies against each protein encoded by TON_1016, TON_1018, and TON_1023, which were overexpressed in *Escherichia coli* Rosetta(DE3)pLysS cells (Stratagene, La Jolla, CA), were generated and purified through His-Bind resin (Novagen, Madison, WI) or through excision of the corresponding protein band after sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) separation.

Nucleotide sequence accession number. The DNA and deduced protein sequences have been deposited in GenBank (<http://www.ncbi.nlm.nih.gov/GenBank/>) under accession number KM489057.

RESULTS AND DISCUSSION

Bioinformatic analysis of the putative regulatory genes. In the CODH gene cluster of *T. onnurineus* NA1, an open reading frame (ORF; TON_1016), namely, *corR*, encodes a putative transcriptional regulator belonging to the LTTR family (Fig. 1). A potential helix-turn-helix (HTH) motif, which would represent the DNA-binding domain, is present in the C-terminal part (amino acids 285 to 328) of CorR (see Fig. S3 in the supplemental material). However, a distinctive substrate-binding domain, which plays roles in coinducer recognition and/or response in the LTTR family of transcriptional regulators, was not found in the CorR protein. A BLAST analysis of the CorR protein of *T. onnurineus* NA1 showed that it exhibits a high degree of resemblance (50 to 55% identity and 70 to 74% similarity) to the CorR homologs of *Thermococcus barophilus* MP, *Thermococcus* sp. AM4, and *Thermococcus* sp. strain ES1. The CorR protein also displayed some homology (26% identity and 43% similarity) to a molybdate-binding protein of “*Candidatus* Korarchaeum cryptofilum” OPF8. In all cases, the *corR* genes were associated with CODH gene clusters (Fig. 1), implying an important role of CorR in CO oxidation. The CODH gene clusters were very similar among the strains in

terms of organization and sequence, except in “*Ca. Korarchaeum cryptofilum*” OPF8, which has only a partial CODH gene cluster.

Intriguingly, an ORF annotated as a hypothetical protein was also conserved in the upstream region of the *corR* gene in the CODH gene clusters of the *T. barophilus* MP, *Thermococcus* sp. AM4, and *Thermococcus* sp. ES1 strains. The ORF was not annotated as a protein-coding sequence in the original genome annotation of *T. onnurineus* NA1 (22). Because of the conservation of the ORF in the three genomes mentioned above, a locus tag, TON_1015-1, could be assigned to a missing ORF in *T. onnurineus* NA1. A BLAST analysis of the deduced amino acid sequence of TON_1015-1 (*corQ*) revealed >70% identity (>90% similarity) to the CorQ homologs of *Thermococcus* strains and showed a significant sequence identity (~40%) to the 4-vinyl reductase (V4R) proteins of various methanogens. In contrast, the ClustalW2 pairwise identity between TON_1015-1 and Kcr_0759 from “*Ca. Korarchaeum cryptofilum*” OPF8 was only 20%. All CorQ homologs contained a V4R domain. The V4R domain is present in many bacterial and archaeal proteins either by itself or fused with other domains, such as HTH domains or the AAA⁺ domains, and is primarily involved in transcription regulation and signal transduction (32). The function of the V4R domain is not well understood, but it appears to function in hydrocarbon binding or redox sensing (33–35). The V4R domains of the CorQ proteins contain three conserved cysteines (see Fig. S3 in the supplemental material). The three conserved cysteines of the CorQ proteins can coordinate a metal, as suggested for the V4R domain of MsvR of *Methanothermobacter thermoautotrophicus*, which plays a role in ligand recognition (32).

Importantly, the genomes of these *Thermococcus* strains and “*Ca. Korarchaeum cryptofilum*” OPF8 do not encode any previously known CO-responsive regulator, and therefore, if regulated, the expression of CODH gene clusters must be controlled by another protein(s). We hypothesize that it is the CorQR pair that regulates the CODH gene cluster in these species.

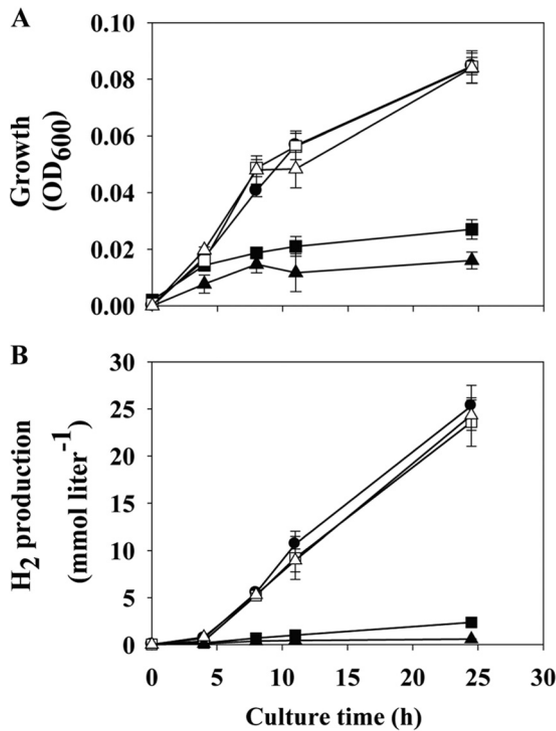


FIG 2 Growth (A) and H₂ production (B) of the wild-type strain *T. onnurineus* NA1 (closed circles) and Δ CorQ (closed squares), Δ CorR (closed triangles), Δ CorQ/*corQR*[↑] (open squares), and Δ CorR/*corQR*[↑] (open triangles) mutants on MM1-CO. Cell growth was monitored by measuring the OD₆₀₀. Average values from triplicate experiments are displayed, and error bars indicate the standard deviations from triplicate experiments.

Inactivation of *corR* and *corQ* by in-frame deletion and complementation of the deletion mutants. To investigate the importance of the *corR* and *corQ* genes in the CO metabolism of *T. onnurineus* NA1, each gene was inactivated by in-frame deletion. The resultant mutant strains, the Δ CorR and Δ CorQ strains, grew normally in YPS medium (heterotrophic condition) and had growth comparable to that of the wild type-strain (data not

shown). However, under the lithotrophic condition where CO was supplied as the main energy source, these mutants grew poorly: the final cell densities of the Δ CorQ and Δ CorR strains were only about 30% and 20% of the final cell density of the wild-type strain, respectively (Fig. 2A). The H₂ production of the mutants was severely impaired as well, with the H₂ production rates being about 10% of the rate of the wild-type strain (Fig. 2B). To confirm that these CO-dependent growth defects in the mutants were caused by the loss of the *corQ* and *corR* genes and not by unexpected genetic variations that could happen during genetic manipulation, each mutant was complemented by inserting intact *corQR* genes at the intergenic region of TON_1126 and TON_1127 (see Fig. S2 in the supplemental material). For this complementation, the *corQR* genes were placed under the control of a strong promoter, P_{TON_0157}, as described in Materials and Methods. In both complemented strains, the transcription levels of *corR* were increased by 5- to 6-fold compared with the level for the wild-type strain (data not shown). The Δ CorQ and Δ CorR strains complemented with integrated *corQR*, designated Δ CorQ/*corQR*[↑] and Δ CorR/*corQR*[↑], respectively, grew normally and produced H₂ gas at a level comparable to that for the wild-type strain (Fig. 2).

These results indicate that the *corQ* and *corR* genes play important roles in CO-dependent growth and the resulting H₂ production in *T. onnurineus* NA1. Also, growth defects were observed only under the CO condition, and it is likely that the functions of the *corQ* and *corR* genes are specific to CO-dependent growth.

Transcriptional analysis of the CODH gene in the wild-type and genetically manipulated strains. To test whether *corR* and *corQ* indeed play a regulatory role in the CO metabolism of *T. onnurineus* NA1, the transcription levels of the catalytic subunit of CODH (TON_1018), a key CO metabolic enzyme, were monitored by RT-qPCR in the wild-type, Δ CorQ, Δ CorR, Δ CorQ/*corQR*[↑], and Δ CorR/*corQR*[↑] strains. While the Δ CorQ and Δ CorR mutant strains barely grew on CO, they could be grown in MM1-pyruvate until exponential phase, harvested, transferred to fresh MM1 with or without 100% CO gas, and incubated for an additional 3 h at 80°C. Total RNAs were extracted from these cells, they were subjected to RT-qPCR as described in Materials and

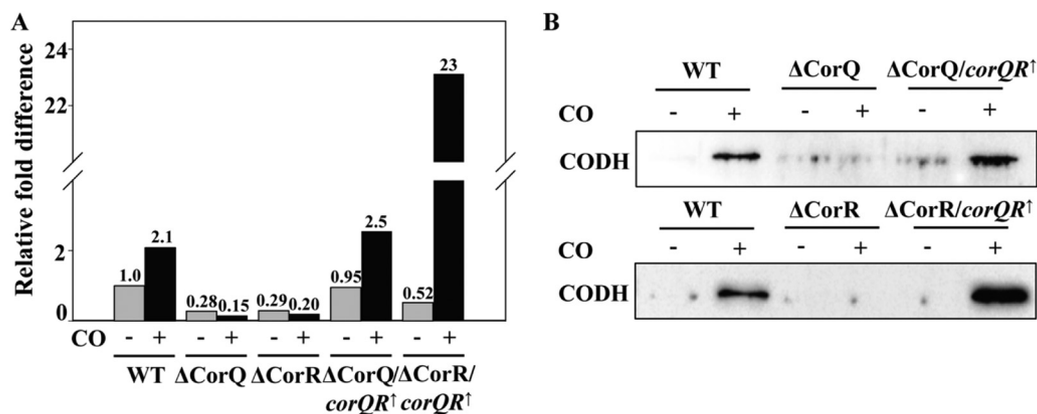


FIG 3 Changes of CODH gene expression at the transcriptional and translational levels after CO addition. (A) mRNA levels of the CODH gene (TON_1018) in the wild-type, Δ CorQ, Δ CorR, Δ CorQ/*corQR*[↑], and Δ CorR/*corQR*[↑] strains. The mRNA quantity was measured by RT-qPCR. The y axis indicates the relative fold difference when the value for the wild-type strain in the absence of CO was set equal to 1.0. (B) Protein levels of CODH in the wild-type, Δ CorQ, Δ CorR, Δ CorQ/*corQR*[↑], and Δ CorR/*corQR*[↑] strains. The amount of the CODH protein was monitored by Western blot analysis. WT, wild-type strain; -, no injection of CO (gray bars); +, injection of CO (black bars).

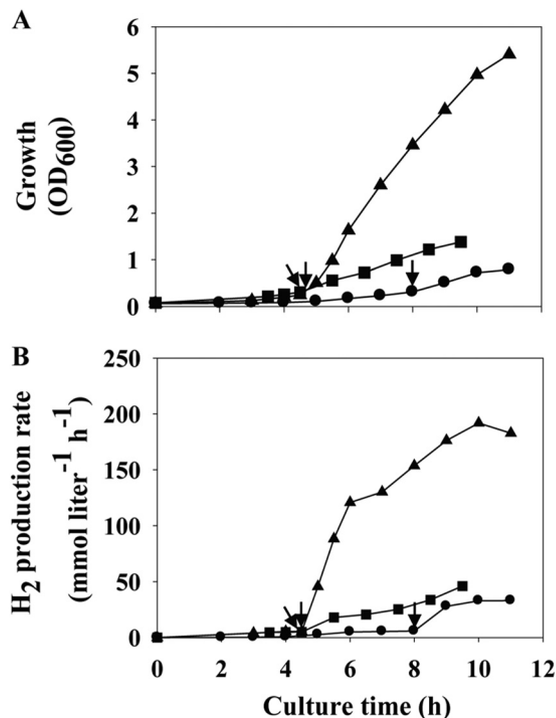


FIG 4 Growth (A) and H₂ production (B) of the wild-type strain *T. onnurineus* NA1 (circles) and the $\Delta\text{CorQ}/\text{corQR}^\uparrow$ (squares) and $\Delta\text{CorR}/\text{corQR}^\uparrow$ (triangles) mutants in a 3-liter bioreactor with CO supplied continuously. Each experiment was performed one time. Cell growth was monitored by measuring the OD₆₀₀. The initial CO flow rate of 20 ml min⁻¹ was raised to 400 ml min⁻¹ when the OD₆₀₀ reached about 0.3, as indicated by arrows.

Methods, and the fold changes in the level of CODH gene expression were calculated. The ΔCorR and ΔCorQ mutants failed to show any CO-dependent changes in the CODH mRNA levels. In contrast, for the wild-type strain and the two complemented mutant strains, there were clear and significant increases in the CODH mRNA level in response to CO (Fig. 3A). However, the two complemented mutants were dissimilar in terms of the fold difference in the level of expression. The fold difference in the level of expression by the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain was about 20-fold higher than the differences for the $\Delta\text{CorQ}/\text{corQR}^\uparrow$ and wild-type strains.

The CODH protein levels were also measured in the strains mentioned above, and the results were generally consistent with the transcript analysis data (Fig. 3B). The CODH protein was de-

tected in the wild-type strain and the two complemented mutants, while the protein was barely visible in the ΔCorR and ΔCorQ mutant strains. Again, the amount of CODH protein was much higher in the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain than in the wild-type and $\Delta\text{CorQ}/\text{corQR}^\uparrow$ strains.

The observation of the difference in the CODH mRNA and protein levels between the two complemented strains is puzzling because we believe that the CorQ and CorR proteins are present at similar saturating levels in both the $\Delta\text{CorQ}/\text{corQR}^\uparrow$ and $\Delta\text{CorR}/\text{corQR}^\uparrow$ strains, where the *corQR* genes are under the control of the same strong promoter, P_{TON_0157}. A mechanistic understanding of this phenomenon awaits further experiments. Nonetheless, our data indicate that the CorQR proteins are required for the CO-dependent transcriptional activation of the CODH gene in response to CO. This may also indicate that the proteins positively regulate the whole CODH gene cluster in *T. onnurineus* NA1.

Enhanced H₂ production in the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain. We previously reported that the overexpression of the CODH gene cluster in the MC01 mutant led to 3.9-fold higher levels of H₂ production (26). The increase in the amount of the CODH protein in the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain, as shown in Fig. 3, may indicate an increase in the level of translation of the whole CODH gene cluster. If this is true, the strain was hypothesized to exhibit a higher H₂ production rate as well. We therefore examined the H₂ production potential of the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain using a bioreactor with 100% CO fed stepwise at a flow rate of 20 to 400 ml min⁻¹. The $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain indeed showed a significantly higher cell density and H₂ production rate than the wild-type and $\Delta\text{CorQ}/\text{corQR}^\uparrow$ strains (Fig. 4). The $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain displayed kinetic parameters 1.6- to 5.8-fold greater than those of the wild-type strain (see Table S2 in the supplemental material). In particular, the maximum H₂ production rate and H₂ productivity were much higher (over 5-fold) in the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain than in the wild-type strain. The $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain showed a slightly higher H₂ production rate (1.6-fold) than even the MC01 mutant, making the $\Delta\text{CorR}/\text{corQR}^\uparrow$ strain the prokaryote with the best CO-dependent H₂ production rate (Table 1).

In summary, we have shown that CorQR is a novel regulatory element of CO metabolism in *T. onnurineus* NA1. The CorQR regulatory system presented here is unique because it consists of two components, while all known CO-sensing transcriptional regulators are single-component proteins where the sensor domain is covalently linked to the response domain. Mechanistic details as

TABLE 1 H₂ production rates of various carboxydophilic hydrogenogenic microbes

| Organism | Cultivation method | H ₂ production rate (mmol liter ⁻¹ h ⁻¹) ^a | Specific H ₂ production rate (mmol g ⁻¹ h ⁻¹) ^a | Reference or source |
|------------------------------------------------------------------------|--------------------|-----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------|---------------------|
| <i>T. onnurineus</i> NA1 ($\Delta\text{CorR}/\text{corQR}^\uparrow$) | Batch | 191.9 | 249.6 | This study |
| <i>T. onnurineus</i> NA1 (MC01) | Batch | 123.5 | 194.7 | 26 |
| <i>T. onnurineus</i> NA1 (wild type) | Batch | 32.9 | 151.3 | 26 |
| <i>Carboxydotherrnus hydrogenoformans</i> | Continuous | 125 | 18.3 | 36 |
| <i>Rhodospseudomonas palustris</i> P4 | Batch | 41 | 41 | 37 |
| <i>Citrobacter</i> sp. strain Y19 | Batch | 5.7 | 27.1 | 38 |
| <i>Rhodospirillum rubrum</i> | Continuous | 4.7 | 11 | 39 |
| <i>Rubrivivax gelatinosus</i> CBS-2 | Continuous | 2.7 | 1.3–33 | 40 |

^a Kinetic parameters were calculated with data from the graphs in Fig. 4. The units of grams represent dry cell weight, except for the volatile suspended solid (VSS) used in *C. hydrogenoformans*.

to how this CorQR system senses CO and transcriptionally activates CODH genes remain to be elucidated. This study also illustrates that transcription regulation circuits can be used to improve biological H₂ production.

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