A Complex Interplay between Nitric Oxide, Quorum Sensing, and the Unique Secondary Metabolite Tundrenone Constitutes the Hypoxia Response in Methylobacter

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ABSTRACT Methylobacter species, members of the Methylococcales, have recently emerged as some of the globally widespread, cosmopolitan species that play a key role in the environmental consumption of methane across gradients of dioxygen tensions. In this work, we approached the question of how Methylobacter copes with hypoxia, via laboratory manipulation. Through comparative transcriptomics of cultures grown under high dioxygen partial pressure versus cultures exposed to hypoxia, we identified a gene cluster encoding a hybrid cluster protein along with sensing and regulatory functions. Through mutant analysis, we demonstrated that this gene cluster is involved in the hypoxia stress response. Through additional transcriptomic analyses, we uncovered a complex interconnection between the NO-mediated stress response, quorum sensing, the secondary metabolite tundrenone, and methanol dehydrogenase functions. This novel and complex hypoxia stress response system is so far unique to Methylobacter species, and it may play a role in the environmental fitness of these organisms and in their cosmopolitan environmental distribution.

IMPORTANCE Here, we describe a novel and complex hypoxia response system in a methanotrophic bacterium that involves modules of central carbon metabolism, denitrification, quorum sensing, and a secondary metabolite, tundrenone. This intricate stress response system, so far unique to Methylobacter species, may be responsible for the persistence and activity of these species across gradients of dioxygen tensions and for the cosmopolitan distribution of these organisms in freshwater and soil environments in the Northern Hemisphere, including the fast-melting permafrosts.

KEYWORDS hypoxia, Methylobacter tundripaludum, nitric oxide, hybrid cluster protein, quorum sensing, tundrenone

Methylobacter species, members of the Methylococcales that are gammaproteobacterial aerobic methanotrophs, have recently emerged as some of the globally widespread, cosmopolitan species, appearing to play a key role in the consumption of methane in a variety of environments and across gradients of dioxygen tensions (1–3). While multiple species of both gamma- and alphaproteobacterial methanotrophs are detectable in environmental samples using molecular techniques (1–4), and many are also available in pure cultures (5, 6), Methylobacter appears to be the dominant and the most active species in freshwater and soil environments of the Northern Hemisphere (1–4). Some of the environments hosting Methylobacter, such as permafrosts, represent vast areas on this planet, which are also subject to rapid environmental change as part of the global warming effect, manifested by increased methane release resulting from ice melt (7, 8). Importantly, Methylobacter activities have been correlated with methane

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consumption not only in oxic but also in hypoxic and even anoxic niches (9–12), even though the mechanisms for such activity remain poorly understood, given the essential role of dioxygen in methane oxidation catalyzed by methane monoxygenase (MMO). The observations on the dominant nature of Methylobacter have also been supported by results from microcosm incubation experiments in which Methylobacter performed as the most competitive species, especially under low dioxygen tensions (13, 14).

In addition to their proficiency in oxidizing methane, under a variety of conditions, Methylobacter species have also been implicated in shuttling carbon from methane through a bacterial food chain, supporting diverse communities and linking methane consumption to other important biogeochemical processes (15). Methanol-utilizing Methylophilaceae have been noted as some of the most persistent partners of Methylobacter in both natural environments (3) and laboratory simulations (13, 14). Experiments involving synthetic community manipulations have suggested that the methanol-consuming partner species may be capable of modifying carbon metabolism in Methylobacter, by affecting the expression of alternative methanol dehydrogenase (MDH) enzymes, and thus potentially modulating the supply of methanol to the satellite populations of Methylophilaceae (16). Other products that may be released by Methylobacter have been hypothesized to be organic acids and polymeric substances, which would support populations of nonmethylotrophic heterotrophs (15, 17).

What makes Methylobacter such an environmentally successful species? It is unlikely that these are its growth characteristics, as in the laboratory, Methylobacter does not perform robustly and is easily outcompeted by other species such as Methyloptomonas (18, 19). Toward the potential for methane oxidation in niches depleted of dioxygen, the denitrification capability has been proposed to be one metabolic means for coping with hypoxia (20, 21). However, transcriptomics studies applied to natural samples with highly active Methylobacter populations failed to account for the transcription of the denitrification pathway (2).

In this study, we approached the question of how Methylobacter copes with hypoxia that it encounters in natural environmental niches, via laboratory manipulation. We first carried out comparative transcriptomics with cultures grown under high dioxygen partial pressure and cultures starved for dioxygen, identifying genes showing differential transcription. We then mutated several genes of interest and investigated mutant phenotypes. We followed up with additional transcriptomic analyses comparing gene expression in mutant versus wild-type backgrounds and comparing expression patterns between cultures provided with ammonium and those provided with nitrate as nitrogen sources. We uncovered an unexpected connection between the hypoxia stress response and quorum sensing (QS) functions, which, in turn, regulate the expression of a gene cluster responsible for the synthesis of a secondary metabolite, tundrenone (Tun). We also obtained evidence for nitric oxide (NO), generated in the denitrification pathway, playing a role in the stress response, likely as a signaling molecule. We further uncovered that the differential expression of genes for the alternative MDH enzymes was also part of the hypoxia response.

RESULTS

Comparative transcriptomics point to the role of Hcp/Hcr in the hypoxia response by Methylobacter. We exposed cultures of Methylobacter tundripaludum strain 31/32 to hypoxia, sampling cells 24 and 48 h after the last addition of O₂, and compared the transcriptomes of the dioxygen-starved cultures to the transcriptomes of the oxygenated cultures (transcriptomics experiment 1) (see Materials and Methods). Two different experiments (experiments A and B) were run in two replicates each, all involving synthetic community mixtures (see Materials and Methods) (17). For the purposes of this study, we focused solely on the transcriptomes of M. tundripaludum 31/32.

Dramatic changes in transcription patterns were determined for cultures exposed to hypoxia, compared to the oxygenated cultures (Fig. 1; see also Tables S1 and S2 in the supplemental material). In general, among the two independent experiments, the
FIG 1 Heat map depicting differential expression of select genes in *M. tundripaludum* 31/32 under oxic (1 h) and increasingly hypoxic (24 and 48 h) conditions (experiments 1A and B). A single replicate for each experiment is shown. Complete transcriptomics data, including additional replicates, are presented in Tables S1 and S2 in the supplemental material. MFS, major facilitator superfamily.
transcription of over 500 genes was downregulated more than 10-fold, while the transcription of about 150 genes was upregulated more than 10-fold, in response to hypoxia. Between the two experiments, the trends observed were very similar (Fig. 1; Tables S1 and S2). The expression of two genes was dramatically upregulated (up to 4,000-fold) in response to hypoxia, encoding a hybrid cluster protein (Hcp) (gene 2451) and a cognate flavodoxin reductase (Hcr) (gene 2450). A closer look uncovered a long history of misannotation of the gene for Hcp. In most genomes, it is annotated as a hydroxylamine reductase, with this persistent misannotation stemming from the previously reported observation of a secondary, minor side activity of the enzyme (22). In a later study, the respective enzyme was identified as a high-affinity NO reductase (23). Additional studies implicated this protein in nitrosative and oxidative stress responses (24, 25). While the hcp-hcr genes were previously proposed to be involved in the denitrification pathway in a methanotroph (a Crenothrix species [26]), in the experiment presented here, genes for other respiratory and assimilatory nitrate reduction functions were expressed at low levels, and they did not show differential expression in response to the onset of hypoxia (Fig. 1 and Fig. 2B; Tables S1 and S2).

One of the most highly downregulated gene clusters in response to hypoxia was the one for the calcium-dependent MDH, the mxa cluster (genes 4146 to 4157) (Fig. 1; Tables S1 and S2). Conversely, the gene for an alternative MDH, the lanthanide-dependent XoxF enzyme (gene 3928), was upregulated. A pattern of reversed transcriptional regulation of the alternative MDH enzymes in response to hypoxia has been noted previously (19). As for other methylotrophy functions, the transcription of genes for MMO (genes 2846 to 2848) was slightly upregulated, while the transcription of most of the genes for the assimilatory ribulose monophosphate and the (partial) serine cycles (27, 28) as well as of genes for formaldehyde and formate oxidation was downregulated (up to 45-fold) (Fig. 1; Tables S1 and S2). Overall, the transcription of genes for many essential cellular functions, including ribosomal proteins, was decreased, likely indicating a switch to the oxygen starvation mode.

Mutants in the hcp-hcr gene cluster, key denitrification genes, demonstrate altered hypoxia and oxidative and nitrosative stress responses. We focused on the cluster of genes whose transcription was most highly induced by hypoxia, hcp and hcr. Additional genes that were part of the cluster encoded a transcriptional regulator annotated as NorR (anaerobic NO reductase transcriptional regulator), a signal transduction protein possessing PAS, EAL, and GGDEF domains (here named Hst, for hypoxia signal transfer), and a hemerythrin (Hmr), a small diiron protein for the transfer and storage of dioxygen (Fig. 2A). While the Hcp/Hcr pair has been previously demonstrated

![Diagram](https://example.com/diagram.png)
to act as a high-affinity NO reductase (23), the expression pattern for *hcp-hcr*, as reported here, showed a poor correlation with the expression patterns for other genes in the NO-producing denitrification pathway (Fig. 2B), potentially suggesting an additional function for Hcp/Hcr. We hypothesized that this additional function may be in the stress response, as previously reported for other organisms (24, 25). To test our hypothesis, we constructed knockout mutants for each of the genes in the *hcp* gene cluster. To directly test the potential of nitrate in serving as an alternative electron acceptor to allow *Methylobacter* to cope with hypoxia, we mutated the major subunit of the respiratory nitrate reductase (NR) (*NarG*; gene 3186). As a control, we also mutated the assimilatory NR (*NasA*; gene 1731). To select for recombinants, we used both nitrate-supplemented (nitrate minimal salts [NMS]) and ammonium-supplemented (ammonium mineral salts [AMS]) selective media. We noted that in each case, colonies appeared only on the AMS and not on the NMS plates, likely suggesting that all the mutants had decreased survival rates when nitrate served as the nitrogen source.

When inoculated into liquid media, all the mutants were able to grow with both nitrate and ammonium as sources of nitrogen, with one exception. The mutant in *NasA* did not show any appreciable growth in nitrate-supplemented medium, while the remaining mutants showed decreased growth compared to ammonium-supplemented medium and to wild-type controls (Fig. S1). These results suggested that all the functions encoded by the *hcp-hcr* gene cluster and both the assimilatory and the dissimilatory denitrification pathways must be involved in the hypoxia stress response by *Methylobacter*.

We tested the survival rate under hypoxic conditions by incubating cells for 40 days without replenishing dioxygen in the headspace atmosphere, monitoring the fate of cell populations every 10 days using LIVE/DEAD cell stain in medium supplemented with either nitrate or ammonium. Over 40 days, approximately 65% of the initial wild-type cell population died out in both types of media, presumably from dioxygen starvation (Fig. S2A). All the mutants demonstrated reduced survival rates compared to the wild type, with the *NasA* mutant demonstrating a dramatic death rate in nitrate medium while surviving quite well in ammonium medium. Notably, cultures starved for dioxygen do not show significant growth, as reported previously (17, 18). Thus, nitrogen availability should not be essential for building biomass. This suggested that the phenotype of the *NasA* mutant might be reflective of a specific stress response. The pattern of rapid dying out was further tested in a short-term experiment, over 12 h, carried out with the entire group of strains. In this experiment, only the *NasA* mutant revealed a significant decline in live cells and only in nitrate-supplemented medium (Fig. S2B). The most parsimonious explanation for such a phenotype would be the redirection of most of the nitrate into the dissimilatory pathway, via respiratory NR (*NarGHIJ*) and NO-producing nitrite reductase (*NirK*), resulting in a surge in toxic NO (Fig. 2B).

We further employed several stressors that are typically used to test for tolerance to oxidative and nitrosative stresses. When challenged with H$_2$O$_2$, all the mutants ceased growth, and the number of live cells steadily decreased over the course of the experiment (Fig. S3A), in contrast to the nonstressed cultures (Fig. S3D). The *NasA* mutant was one exception, and it revealed a steep decline in live-cell numbers when incubated in nitrate medium, even in cultures with no added stressor, further suggesting that a stressor, likely NO, was produced by this mutant. While wild-type cells showed a diminished growth rate under H$_2$O$_2$ stress, cell death was not observed over the course of the experiment (Fig. S3A). Very similar results were obtained with an alternative oxidative stress agent, paraquat (Fig. S3B).

As a nitrosative stress agent, we employed a diethylenetriamine (DETA)-NO adduct. When tested under oxic conditions, most of the mutants were able to grow at rates similar to wild-type strain rates with both nitrate and ammonium. The *NasA* mutant did not grow in nitrate medium, as expected (Fig. S3C). Two other exceptions were the mutants in Hcp and Hcr, showing severely declining growth. This phenotype was...
consistent with the Hcp/Hcr pair possessing NO reductase activity, with the mutants thus being subject to NO toxicity (Fig. S3C).

**NO and N₂O measurements support the role of NO in the stress response and the role of Hcp/Hcr in NO reduction.** We directly measured the dynamics of instantaneous concentrations of dissolved NO, N₂O, and O₂ in select strain cultures, using microsensors, after the growth medium containing ammonium as the nitrogen source (AMS) was replaced with NMS medium. Upon exposure to nitrate, the NasA mutant immediately accumulated NO. The peak of NO was observed as early as 1 h after exposure to nitrate, while oxygen consumption was very slow (Fig. 3B), corroborating the observations of rapid cell death in the presence of nitrate (Fig. S1 and S2). No short-term NO accumulation was observed when NarG and Hcp mutants or the wild-type strain was exposed to nitrate (Fig. 3A, C, and D). The NarG mutant showed brief dioxygen consumption followed by the loss of metabolic activity. This phenotype is not easy to interpret, but it would be consistent with the involvement of NO in the stress response. Neither NO nor N₂O was produced by this mutant (Fig. 3C). In the case of the Hcp mutant and the wild-type strain, NO accumulated only after a long lag time (>10 h) (Fig. S4A and B). A peak of N₂O was measured only in the wild-type strain, with this peak coinciding with the NO peak (Fig. S4A). Notably, NO and N₂O production by the wild-type strain was observed only after the onset of hypoxia, while the Hcp mutant accumulated NO under oxic conditions (Fig. S4B), suggesting that the denitrification pathway was deregulated in this mutant. The lack of N₂O production by this mutant further supported the role of Hcp in NO reduction.

**Lack of Hcp alters the transcription pattern and reveals connections between the hypoxia stress response, quorum sensing, and the secondary metabolite tundrenone.** With transcriptomics experiment 2, we approached the following questions: does the mutation in Hcp cause a transcriptional response, and is this response hypoxia specific? RNA was isolated from cells exponentially growing under oxygenated conditions with nitrate or with ammonium and from cells subjected to dioxygen starvation for 24 h (see Materials and Methods). A total of 16 RNA samples were sequenced, representing the wild type and the Hcp mutant incubated under the four
conditions, in two replicates each (Table S3). When the transcriptomes of the Hcp mutant and the wild type were compared, under either condition, the most dramatic response to the lack of Hcp was manifested by the recently characterized gene cluster encoding quorum sensing (QS) functions and the novel secondary metabolite tundrenone (Tun) (29) (Fig. 4). The transcription of a total of 27 genes in the cluster was upregulated, including key genes in Tun biosynthesis. The gene for an N-3-hydroxydecanoyl-l-homoserine lactone (HSL) synthase, mbaI, showed the most dramatic response, which was most pronounced in cells subjected to hypoxia in nitrate medium, >1,000-fold upregulation. In comparison, the transcription of mbaI was >100-fold upregulated in cells subjected to hypoxia in ammonium medium and approximately 50-fold upregulated in cells from the oxygenated cultures (Fig. 5). It has been noted previously that in the MbaI mutant not producing native HSL, when exogenous HSL was added, some of the most downregulated genes were the genes for Hcp and Hcr (29). Thus, it appears that the Hcp/Hcr and the QS/Tun systems are inversely regulated and that they potentially regulate each other. It appears that Hcp/Hcr or the products controlled by this enzyme pair (potentially NO) control the transcription of the QS system and, thus, the production of Tun and that a product positively regulated by the QS system, potentially Tun, controls transcription from hcp-hcr. Other notable alterations in transcription included the significantly reduced expression of the mxa operon in the mutant when grown with ammonium but not nitrate. The mxa gene cluster was also downregulated in wild-type cells in response to hypoxia, as has been noted previously (19), but to a much lesser degree than in the Hcp mutant (Fig. 5). This further suggested that an intermediate in the denitrification pathway, likely NO, must be part of the regulatory cascade controlling the expression of the mxa gene operon. The levels of NO should be elevated in the Hcp mutant grown with nitrate, thus positively correlating with mxa expression. The transcription of the nar operon was slightly (~2-fold) elevated in this mutant (Fig. 4).

**NO is involved in transcriptional regulation of quorum sensing, tundrenone synthesis, and methanol oxidation.** With transcriptomics experiment 3, we tested whether NO may be directly involved in gene expression regulation and which, if any, functions would be affected by the elevated levels of NO. We employed the NasA mutant, which revealed the most dramatic phenotype for growth with nitrate, and it emitted NO (Fig. 3B). Samples for RNA extraction were taken after 1 h of exposure to nitrate, based on the evidence for immediate NO release by the mutant (Fig. 3B), and these nitrate (NO)-challenged samples were compared to samples incubated with ammonium under fully oxygenated conditions. Overall, the expression of many genes, including most of the methylotrophy genes, was downregulated (3- to 5-fold) after exposure to nitrate, in agreement with the toxic nature of NO. However, some genes showed upregulation. Among these were genes involved in nitrate metabolism functions, such as ammonium transporters, nitrogen regulatory proteins, nitrate/nitrite transporters, as well as the assimilatory nitrite reductase (Fig. 5; Table S4). In addition, the transcription of genes encoding flagella as well as type IV pilus assembly functions was upregulated. Remarkably, some of the regulated gene clusters included those for QS/Tun, Hcp/Hcr, Mxa, and Xox functions. Of these, the transcription of genes encoding QS/Tun as well as Mxa functions showed rather significant (up to 20-fold) downregulation, while the transcription of genes encoding Hcp/Hcr and XoxF was slightly upregulated, once again suggesting an inverse regulation of QS/Tun/Mxa versus Hcp/Hcr/XoxF genes. These results suggested that the transcriptional response to NO must be dose dependent and that NO can serve as either a stimulator or a toxic agent, depending on its intracellular concentration.

**Quorum sensing and tundrenone are involved in the hypoxia response.** Transcriptomics experiment 4 was designed to further decipher the potentially interlinked roles of Hcp/Hcr and QS/Tun in the hypoxia response. This experiment was carried out with the previously described mutants in QS (MbaI) (29) and Tun biosynthesis (TunJ) (30). The former mutant does not produce the signal molecule HSL, which induces the
FIG 4  Heat map depicting differential expression of select genes in the Hcp mutant in comparison with wild-type M. tundripaludum 31/32 under four different conditions (experiment 2). A single replicate for each strain/condition is shown. Complete transcriptomics data, including additional replicates, are presented in Table S3 in the supplemental material. GNAT, general control nonrepressed-related N-acetyltransferase; HAD, haloadic dehydrogenase; ACP, acyl carrier protein; PGA, poly-beta-1,6-acetyl-D-glucosamine.

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| Maximum | R1 | R2 | R1 | R2 |
|---------|----|----|----|----|
| 962. ammonium transporter, Amt family |    |    |    |    |
| 963. nitrogen regulatory protein P-II family |    |    |    |    |
| 964. ammonium transporter, Amt family |    |    |    |    |
| 965. nitrogen regulatory protein P-II family |    |    |    |    |
| 966. Putative beta-barrel porin-2, OmpL-like bbp2 |    |    |    |    |
| 967. glutamine synthetase |    |    |    |    |
| 1130. nitrogen fixation negative regulator NirL |    |    |    |    |
| 1131. Nif-specific regulatory protein |    |    |    |    |
| 1701. nitrite reductase (NADH) small subunit |    |    |    |    |
| 1702. nitrite reductase (NADH) large subunit |    |    |    |    |
| 1703. response regulator receiver and ANTAR domain protein |    |    |    |    |
| 1715. nitrate/nitrite transport system substrate-binding protein |    |    |    |    |
| 1716. nitrate/nitrite transport system permease protein |    |    |    |    |
| 1717. nitrate/nitrite transport system ATP-binding protein |    |    |    |    |
| 1734. Ntc family, nitrogen regulation sensor Histidine kinase GinH |    |    |    |    |
| 1735. Ntc family, nitrogen regulation response regulator GinG |    |    |    |    |
| 1799. flagellar hook-associated protein 3 FlgI |    |    |    |    |
| 1790. flagellar hook-associated protein 1 FlgK |    |    |    |    |
| 1791. flagellar protein FlgJ |    |    |    |    |
| 1792. flagellar P-ring protein precursor FlgI |    |    |    |    |
| 1793. flagellar L-ring protein precursor FlgH |    |    |    |    |
| 1794. flagellar basal-body rod protein FlgG |    |    |    |    |
| 1795. flagellar basal-body rod protein FlgF |    |    |    |    |
| 1796. flagellar hook protein FlgE |    |    |    |    |
| 1797. flagellar basal-body rod modification protein FlgD |    |    |    |    |
| 1798. flagellar basal-body rod protein FlgC |    |    |    |    |
| 1799. flagellar basal-body rod protein FlgB |    |    |    |    |
| 1800. flagella basal body P-ring formation protein FlgA |    |    |    |    |
| 1801. anti-sigma-28 factor, FigM family |    |    |    |    |
| 1802. flagella synthesis protein FigH |    |    |    |    |
| 1836. type IV pili assembly protein PjaA |    |    |    |    |
| 2014. N-acetyl-L-homoserine lactone synthetase MbaI |    |    |    |    |
| 2016. inositol glycerol phosphate synthase subunit hisH |    |    |    |    |
| 2017. Integrase core domain-containing protein |    |    |    |    |
| 2026. N-acetyl-L-homoserine lactone synthetase |    |    |    |    |
| 2027. HAD superfAMILY subfamily IB hydrolase, TGR01940 TunA |    |    |    |    |
| 2028. Delta-aminolevulinic acid dehydratase, porphobilinogen synthase TunB |    |    |    |    |
| 2029. A-factor bioynthesis holdog domain-containing protein TunC |    |    |    |    |
| 2030. 2-polypropylthio-methylmethanethiol sulfatase TunE |    |    |    |    |
| 2031. Predicted oxidoreductase TunE |    |    |    |    |
| 2032. MFS transporter, DH2 family, multidrug resistance protein TunF |    |    |    |    |
| 2033. membrane fusion protein, multidrug efflux system TunG |    |    |    |    |
| 2034. Acyl carrier protein TunI |    |    |    |    |
| 2035. Condensation domain-containing protein Tunl |    |    |    |    |
| 2036. long-chain acyl-CoA synthetase TunJ |    |    |    |    |
| 2037. Primery-ACP methyl ester carboxyltransferase TunK |    |    |    |    |
| 2038. NADPH:quinone reductase TunM |    |    |    |    |
| 2040. anthranilate synthase, component I (EC 4.1.3.27) TunN |    |    |    |    |
| 2041. anthranilate synthase, component II (EC 4.1.3.27) TunD |    |    |    |    |
| 2042. glutathione S-transferase TunP |    |    |    |    |
| 2450. Ferredoxin-NADP reductase |    |    |    |    |
| 2451. hydroxylamine reductase precursor (EC 1.7.99.1) |    |    |    |    |
| 2452. anaerobic nitric oxide reductase transcription regulator |    |    |    |    |
| 2846. methane monoxygenase subunit C |    |    |    |    |
| 2847. methane monoxygenase subunit A |    |    |    |    |
| 2848. methane monoxygenase subunit B |    |    |    |    |
| 3014. two-component system, sensor histidine kinase FinB |    |    |    |    |
| 3015. two-component system, response regulator FinC |    |    |    |    |
| 3016. flagellar hook-basal body complex protein FinE |    |    |    |    |
| 3017. flagellar M-ring protein FinF |    |    |    |    |
| 3018. flagellar Fil protein |    |    |    |    |
| 3019. flagellar motor switch protein FinM |    |    |    |    |
| 3020. flagellar motor switch protein FIN/FIY |    |    |    |    |
| 3021. flagellar protein FIQ/FIZ |    |    |    |    |
| 3022. flagellar biosynthetic protein FinQ |    |    |    |    |
| 3023. flagellar biosynthetic protein FinR |    |    |    |    |
| 3024. flagellar biosynthetic protein FinB |    |    |    |    |
| 3025. flagellar biosynthesis protein FinA |    |    |    |    |
| 3026. flagellar biosynthesis protein FinA |    |    |    |    |
| 3027. flagellar biosynthesis protein Tune |    |    |    |    |
| 3028. quinoprotein dehydrogenase-associated XoxJ |    |    |    |    |
| 3029. PQQ-dependent dehydrogenase, methanol/nitrophenol family XoxF |    |    |    |    |
| 4144. NIT/TauT family transport system substrate-binding protein |    |    |    |    |
| 4145. hypothetical protein |    |    |    |    |
| 4146. mxaI, protein |    |    |    |    |
| 4147. mxaK protein |    |    |    |    |
| 4148. mxaC protein |    |    |    |    |
| 4149. mxaA protein |    |    |    |    |
| 4150. hypothetical protein |    |    |    |    |
| 4151. MxaR-like ATPase |    |    |    |    |
| 4152. methanol dehydrogenase small subunit MxaI |    |    |    |    |
| 4153. cytochrome cI, apoprotein MxaG |    |    |    |    |
| 4154. mxaI protein |    |    |    |    |
| 4155. methanol dehydrogenase large subunit MxaF |    |    |    |    |
| 4156. two component transcriptional regulator, LuxR family |    |    |    |    |
| 4157. mxaD protein |    |    |    |    |
| 4333. hypothetical protein |    |    |    |    |
| 4336. protein of unknown function (DUF3391) |    |    |    |    |
| 4339. two-component system, chemotaxis family, sensor kinase CheA |    |    |    |    |
| 4356. chemotaxis protein methyltransferase CheR |    |    |    |    |
| 4379. two-component system, chemotaxis family, response regulator CheB |    |    |    |    |
| 4390. purine-binding chemotaxis protein CheW |    |    |    |    |

**FIG 5** Heat map depicting differential expression of select genes in the NasA mutant challenged with nitrate (experiment 3). R1 and R2 are data from biological replicates. Complete transcriptomics data are presented in Table S4 in the supplemental material. PQQ, pyrroloquinoline quinone.
transcription of the genes essential for the biosynthesis of Tun. Thus, this mutant produces neither HSL nor Tun under the conditions used (29). The latter mutant does not produce Tun, while it still produces HSL (30). Note that a different but closely related M. tundripaludum strain, 21/22, was employed in these experiments, due to the availability of validated mutants (29, 30). The mutants and the wild type were grown in either nitrate (NMS)- or ammonium (AMS)-containing medium for 48 h, at which stage cell densities sufficient for the production of HSL were reached (29, 30), and at the same time, dioxygen in the cultures was sufficiently depleted to cause hypoxia (13, 14). Remarkably, neither MbaI nor TunJ mutants produced appreciable amounts of hcp-hcr transcripts with either nitrate or ammonium, revealing phenotypes not responsive to hypoxia. Likewise, the transcriptions of both the mxa gene cluster and xoxF were deregulated, with the mutants not responding to hypoxia (Fig. 6; Table S5). While MbaI and TunJ mutants showed similar transcription patterns for some functions, they demonstrated different patterns for others, suggesting that HSL and Tun may have overlapping but distinct functions in transcriptional regulation. For example, it appears that in the absence of Tun, the expression levels of the pilus function genes (genes 4154 to 4169), exopolysaccharide biosynthesis genes (genes 2050 to 2058), and genes encoding a homolog of MMO (PxmABC; genes 3483 to 3486) were downregulated under both nitrate and ammonium conditions, while the expression patterns between the mbaI mutant and the wild-type strain were reversed (Fig. 6; Table S5).

**DISCUSSION**

In this study, we initiated the quest for untangling the complex mechanistic details of the hypoxia response employed by species of the *Methylobacter* genus that are aerobic methanotrophs. This stress response system uniquely combines modules of central carbon metabolism (i.e., methanol oxidation), denitrification, quorum sensing, and a secondary metabolite, tundrenone. This complex system may be responsible for the persistence and activity of *Methylobacter* species across gradients of dioxygen tensions and for their cosmopolitan distribution in freshwater and soil environments in the Northern Hemisphere, including the fast-melting permafrosts. One of the key elements of this hypoxia response system is the Hcp/Hcr pair of proteins that serve as a NO reductase. While Hcp and Hcr are rather widespread among *Proteobacteria* inhabiting both oxic and anoxic environments (31), to our knowledge, *Methylobacter* presents a unique case in which transcription from these genes appears to be controlled by a homolog of NorR that has been previously demonstrated to directly sense NO (32). Thus, NO appears to act as one of the signaling molecules in the hypoxia response. A hemerythrin (Hmr) encoded by the same gene cluster likely binds O₂, as has been demonstrated for a homolog from a related *Methyllococcales* species, *Methylococcus capsulatus* Bath (33). Thus, O₂ appears to be another signaling molecule that Hmr potentially transfers to the PAS domain of the signal transduction protein named Hst here, while this domain could potentially also sense NO (34). The presence of the EAL and GGDEF domains (35) in the Hst protein provides support for the existence of a downstream regulatory cascade mediated by cyclic diguanylate, thus implying further complexity of the hypoxia response switch. Complex regulatory patterns have been previously proposed for some of the methylotrophy functions, such as the methanol oxidation functions, involving multiple transcriptional regulators (36, 37). From the data presented here, it appears that methanol oxidation is also regulated at the transcriptional level as part of the hypoxia response cascade, potentially responding to NO, O₂, and additional regulators such as NorR.

Importantly, while the traditional respiratory denitrification functions (Nar and Nir) were expressed under both oxic and hypoxic conditions, their transcription patterns did not follow the ones of the hcp-hcr genes. One intriguing scenario that emerges from these data is that the main role of the respiratory denitrification pathway may be in producing the signaling molecule NO in response to hypoxia rather than in serving as a major energy-generating pathway.

The functional interconnection between the Hcp/Hcr and the QS/Tun systems...
FIG 6  Heat map depicting differential expression of select genes in the MbaI and TunJ mutants in comparison with M. tundripaludum 21/22, grown with either ammonium or nitrate (experiment 4). Complete transcriptomics data are presented in Table S5 in the supplemental material. SAM, S-adenosylmethionine.
presents the next level of complexity and suggests a potential role for the recently discovered Tun as a novel signaling molecule whose synthesis, in turn, involves a more traditional signal transducer, HSL (Fig. 7). It is also possible that the QS/Tun system may respond to NO directly, as has been previously described for other QS systems (38, 39).

Why would such complexity be necessary? It is tempting to speculate that the complex interplay between different signals and regulators and their further interconnection with central metabolic pathways constitute a very robust system that allows Methylobacter to better adapt to changing environmental factors such as fluctuations in dioxygen availability and, potentially, to combinations of stressors. This hypothesis would be consistent with the persistence of Methylobacter over other aerobic methanotrophs, including the closely related genera Methylosarcina and Methyloomonas, in methane-fed laboratory microcosms, especially under hypoxic conditions (13, 14).

While carbon metabolism schemes between representatives of Methylococcaceae are very similar (6), they appear to be differentially equipped for stress responses. While Methylosarcina and Methyloomonas may or may not encode the Hcp/Hcr pair and other respiratory denitrification enzymes, none of their known genomes encode the QS/Tun module (6). At the same time, it has been demonstrated that signaling molecules produced by Methylobacter may elicit a response by members of the natural community inhabiting the same niche, such as Methyloomonas (40). On the other hand, it has been demonstrated that Methylobacter may respond to signals from other bacteria by altering its methylotrophy scheme (16). Intriguingly, the QS/Tun module in Methylobacter appears to be encoded by a genomic island (29); thus, it can be potentially spread among the community members under specific selective pressures, such as prolonged hypoxia, or be lost without such selective pressures, for example, as a result of a drought.

The details and the role of the hypoxia response described here in the environmental fitness of Methylobacter need further investigations, via both broader analyses of environmental microbiomes as well as laboratory manipulation of methanotrophs differentially equipped for a hypoxia response. However, the importance of this response likely extends beyond single-organism physiology, as aerobic methanotrophy is increasingly recognized as a community function. Thus, perhaps, further advances into the hypoxia response should involve not only single-species-based experiments but also community-based analyses.

**MATERIALS AND METHODS**

**Strains and growth conditions.** Methylobacter tundripaludum 21/22 and Methylobacter tundripaludum 31/32 (6) were routinely cultivated in nitrate mineral salts (NMS) medium (41) with 25% methane and 75% (vol/vol) air in the headspace, with shaking at 200 rpm at 18°C, or on NMS medium solidified with 1.5% agar (Difco, Thermo Fisher Scientific, Waltham, MA, USA) and placed under a methane-air atmosphere in tight jars. The headspace atmosphere was created essentially as previously described (13, 14, 16–19). For certain experiments, ammonium mineral salts (AMS) medium (41) was used as per the experimental design (see below). All the glassware employed was acid washed for 24 h in 1 M hydrochloric acid before use, to remove trace amounts of any metals that may be adhering to the glass, such as lanthanides, which are known to affect the expression of alternative methanol dehydrogenases (15, 16). For some experiments, the headspace atmosphere was altered by replacing dioxygen with dinitrogen in order to create hypoxic conditions as per the experimental design.
**Genetic manipulations.** All genetic manipulations were achieved through electroporation of the assembled PCR amplification-based constructs into the cells of *M. tundripaludum* 31/32, followed by the selection of chromosomal recombinants, as previously described (42). Briefly, for the gene knockouts, the kanamycin resistance gene cassette (amplified from plasmid pCM433 [43]) and the two regions flanking the gene of interest were assembled by fusion PCR (44). The assembled fragment was further amplified using nested primers at a high annealing temperature (44). The resulting product was purified by using the QiAquick PCR purification kit (Qiagen), and the product was directly electroporated into *M. tundripaludum* 31/32. After electroporation, cells were precultivated overnight in liquid NMS medium without kanamycin, after which cells were harvested and transferred onto either NMS or AMS plates containing 50 µg/ml kanamycin. Single colonies were selected, and the desired mutation was verified by sequencing of the respective PCR-amplified fragment of the chromosomal DNA.

**Measurement of growth rates.** To investigate the growth dynamics of mutant strains in comparison to wild-type *M. tundripaludum* 31/32, cultures were placed into 30-ml tubes sealed with rubber stoppers (Wheaton, Millville, NJ, USA). Each culture (10 ml) was incubated in either NMS or AMS medium with 25% methane in the headspace and various dioxygen concentrations. For “high-oxygen” conditions, a methane/oxygen ratio of 25:75 was used. For “low-oxygen” conditions, a methane/oxygen/dinitrogen ratio of (Wheaton, Millville, NJ, USA). Each culture (10 ml) was incubated in either NMS or AMS medium with 25% methane at 600 nm (OD600) measurements were carried out using a Jenway 7300 spectrophotometer (Bibby Scientific, Burlington, NJ, USA) every 4 h. All values presented with a ± sign represent a minimum of 3 measurements, with ± reporting the standard error.

**Live/dead cell tests.** To test the hypoxia response of the mutants, cells of the seven mutants and the wild-type control were grown in 30-ml tubes, as described above, in both NMS and AMS media, starting with cultures at an OD600 of approximately 0.4. The high-oxygen headspace was created once and not replenished for 40 days. Survival rates were determined using LIVE/DEAD stain. A similar, short-term experiment was run over 12 h, and samples were taken every 4 h. For oxidative and nitrosative stress responses, 0.1 mM H2O2, pararaut, or DETANO, all purchased from Sigma-Aldrich (St. Louis, MO, USA), was added to the cultures under a methane-air (25:75) atmosphere, and samples for dead/live tests were taken every 4 h. To select for the specific concentration of each stressor, series of preliminary experiments were carried out using plate assays with H2O2 and pararaut over 10 days, with concentrations ranging from 0.01 to 2 mM. Concentrations of 0.1 mM and above were sufficient to prevent the growth of each of the mutants, while the wild-type strain was able to grow. As DETANO is oxygen sensitive, only liquid culture experiments were carried out, with 0.02 and 0.1 mM DETANO. The latter concentration produced a stronger negative response by the mutants.

**Instantaneous dissolved gas detection.** Cells were grown under standard conditions in AMS medium to the mid-exponential stage (OD600 of approximately 0.8). Cells were then concentrated by centrifugation (4,500 × g for 10 min) and resuspended in NMS medium to an OD600 of approximately 1.5, and cell suspensions (40 ml) were immediately placed into a 60-ml triple-port microrespirometry (MR) chamber (Unisense, Aarhus, Denmark). All MR experiments were performed at room temperature (approximately 20°C). O2 dynamics were measured using a Stox oxygen microsensor (Unisense, Aarhus, Denmark). The sensor body was partially wrapped with parafilm to fit into the port on the MR chamber. N2O and NO concentrations were measured using an N2O-MR sensor (Unisense) and a NO-MR sensor (Unisense), respectively. For O2 and N2O sensors, calibration was conducted according to the manufacturer’s instructions. For the NO sensor, 100 mM freshly prepared S-nitroso-N-acetyl-D,L-penicillamine (SNAP; Cayman Chemical, MI, USA) in a 0.1 M CuCl2 solution was used for calibration. All sensors were plugged directly into a microsensor multimeter (Unisense) and polarized for >1 day prior to use. All data were logged on a laptop via the microsensor multimeter using SensorTrace Logger software (Unisense).

**Gene expression analysis.** (i) **Experiment 1: time-resolved hypoxia response.** Experiment 1 was described previously (17). Briefly, synthetic communities were assembled by mixing 4 strains (experiment 1A) or 14 strains (experiment 1B). In the 4-strain community, *M. tundripaludum* 31/32 was the only methanotroph species, and in the 14-strain community, *M. tundripaludum* 31/32 was one of the two methanotroph species. The communities were grown in NMS medium with 25%/75% (vol/vol) methane-air in the headspace. After 24 h and at an OD600 of approximately 0.5, samples were flushed with the same atmosphere, and replicate samples were sacrificed after 1, 24, and 48 h, with the first sample beingoxic and the latter samples becoming increasingly hypoxic.

(ii) **Experiment 2: Hcp mutant versus the wild type.** Hcp mutant and wild-type cells were grown under high-oxygen conditions in either NMS or AMS medium to an OD600 of approximately 0.5; these
cells were then used to inoculate replicate cultures, and these cultures were incubated as described above, for another 24 h, to an OD_{600} of approximately 0.5. At this point, half of the samples were given the same methane-air atmosphere for 1 h, after which these (oxic) samples were collected. The second set of samples was given an atmosphere of methane-dinitrogen (25%:75%, vol/vol) to create hypoxic conditions, and the samples were collected after 24 h.

(iii) Experiment 3: NasA mutant NO challenge. The NasA mutant was grown under high-oxygen conditions in AMS medium to an OD_{600} of approximately 0.5. At this point, half of the samples were pelleted by centrifugation (4,500 × g for 10 min) and reinoculated into AMS medium. The second set of samples was pelleted by centrifugation and reinoculated into NMS medium. All the samples were incubated under high-oxygen conditions for 1 h. After incubation, the cells were centrifuged and used for RNA extraction.

(iv) Experiment 4: MbaI and TunJ mutants versus the wild type. MbaI and TunJ mutants and wild-type M. tundripaludum 21/22 were grown in either NMS or AMS medium without replenishing the headspace atmosphere (25%:75% methane-air). Cells were harvested for RNA isolation after 48 h.

RNA was extracted as described previously (17, 19). cDNA library preparation and RNA sequencing were performed by Genewiz (South Plainfield, NJ, USA), using Illumina HiSeq 2-by-150 (paired-end) reads. The raw reads from the sequencing facility were aligned to the corresponding genome assemblies (6), both of which were downloaded from the Joint Genome Institute IMG website (https://img.jgi.doe.gov/). Alignments were performed using BWA version 0.7.12-r1044 using the BWA-MEM algorithm and default parameters (45). The alignments were postprocessed into sorted BAM files with SAMtools version 1.2-232-g87cdc4a (46). Reads were attributed to open reading frames (ORFs) using the htseq-count tool from “HTSeq” framework version 0.6.1p1 in the “intersection-nonempty” mode (47). Differential abundance analysis was performed with DESeq2 1.2.10 (48) using R 3.3.1.

Data availability. Transcriptomics data have been deposited under BioProject accession number PRJNA575263.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, TIF file, 1.9 MB.
FIG S2, TIF file, 0.7 MB.
FIG S3, TIF file, 2.3 MB.
FIG S4, TIF file, 1.1 MB.
TABLE S1, XLSX file, 0.4 MB.
TABLE S2, XLSX file, 0.4 MB.
TABLE S3, XLSX file, 1 MB.
TABLE S4, XLSX file, 0.4 MB.
TABLE S5, XLSX file, 0.8 MB.

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We declare no conflicts of interest.

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