Nitric Oxide and Oxygen Regulate Truncated Hemoglobin Gene Expression in *Frankia* Strain CcI3

James Niemann† and Louis S. Tisa*

Department of Microbiology, University of New Hampshire, Durham, New Hampshire 03824-2617

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The *Frankia* genome contains two truncated hemoglobin genes (*hboN* and *hboO*) whose functions remain to be determined. Nitric oxide (NO) generated by the addition of 400 μM SNAP (S-nitroso-N-acetylpenicillamine) caused a 10-fold increase in *hboO* gene expression but had no effect on *hboN* expression. The addition of the NO scavenger, carboxy-PT10, reduced the effect of SNAP. *hboO* gene expression increased under low-oxygen conditions, while *hboN* expression was unaffected. These results suggest that *HboN* may function in protection from nitrosative stress and that *HboO* may act as an oxygen transport molecule for increased respiration in hypoxic environments.

Truncated hemoglobins (trHbs), the newest branch of the hemoglobin superfamily, are typically 20 to 40 amino acids shorter than traditional hemoglobin but retain the classically conserved globin fold (1, 21, 28). Three distinct groups of trHb proteins (trHbN, trHbO, and trHbP) are found distributed among eubacteria, protozoans, and plants, and their functions are currently being elucidated. For example, *Mycobacterium tuberculosis*, which produces two trHbs (trHbN and trHbO), has been hypothesized to use trHbN (glnN) to detoxify nitric oxide produced by macrophages in tubercles (17). *M. tuberculosis* trHbO (glnO) has been proposed to act as an oxygen delivery protein for terminal oxidases to aid in the stationary phase survival of this organism within hypoxic tubercles (10, 18).

*Frankia* is a nitrogen-fixing actinobacterium (gram-positive filamentous bacterium) that forms a symbiotic association with over 200 different species of plants belonging to eight different plant families, which are only distantly related to each other (for reviews, see references 4 and 27). Hemoglobin production seems to be widespread among all of the *Frankia* isolates (3, 26), but its specific function(s) is unknown. Biochemical studies have shown that the presence or absence of a combined nitrogen source does not affect total hemoglobin production (3), but total hemoglobin levels are greater when cells are grown with 2% oxygen than when they are grown with 20% oxygen.

Analysis of the *Frankia* genome elucidated the presence of two trHb genes (*hboN* and *hboO*) (16), and phylogenetic analysis grouped them closest to their respective *Mycobacterium* orthologs (15), suggesting potential analogous functions for the two *Frankia* hemoglobins. Since both microbes are capable of intracellular growth during their life cycle, this hypothesis is not unreasonable. The goal of this work was to evaluate the relative expression levels of the trHbN and trHbO genes in *Frankia* strain CcI3 under various environmental conditions.

*Frankia* strain CcI3 was grown and maintained in propionate basal medium with NH₄Cl as a nitrogen source as described previously (25). Total RNA was isolated from *Frankia* strain CcI3 as described by Sung et al. (23). DNA was removed from RNA samples with a DNase treatment using DNase I (NEB) according to the manufacturer’s recommendations. Reverse transcriptase PCR (RT-PCR) was performed using a Titan One Tube RT-PCR system (Roche) according to the manufacturer’s recommendations. The following primer sets were used: HbNcci92 (5′-CACCCCTCTTGTGCAACC-3′) and HbNcci300 (5′-GGTGTTTTCCGTGGGAC-3′) for *hboN*, HbOcci299 (5′-GGGACGCCTGGCTGAAGA-3′) and HbOcci375 (5′-CCAGAGCTGCGCTGGAAGATC-3′) for *hboO*, and DB41 (5′-TTTCTCATACCGACCCG-3′) and DB44 (5′-GGCTTCGGCGAGTAGGT-3′) for *glnA* (5). The thermocycling parameters were as follows: (i) reverse transcription at 55°C for 30 min; (ii) initial denaturation at 94°C for 2 min; (iii) 9 cycles of denaturation at 94°C for 20 s, primer annealing at 55°C for 30 s, and primer extension at 68°C for 45 s; (iv) 24 cycles of denaturation at 94°C for 20 s, primer annealing at 55°C for 30 s, and primer extension at 68°C for 50 s, with an additional 5 s added to each progressive cycle; and (v) a final extension step at 68°C for 4 min. Amplicons were resolved by gel electrophoresis, and the respective band intensities were quantified using Quantity One software (Bio-Rad).

For quantitative PCR (qPCR), cDNA synthesis was performed using SuperScript III reverse transcriptase (Invitrogen) according to the manufacturer’s directions. qPCR was performed using Power Sybr green PCR master mix (Applied Biosystems) according to the manufacturer’s recommendations. The above-described primer sets were used for real-time PCR, except that for *HboN* expression, primer HbNcci219 (5′-CCTACCGACGCCCCACTT-3′) was used instead of HbNcci92. Amplification was carried out using an ABI GeneAmp 5700 sequence detection system adapted on a 96-well GeneAmp 9600 PCR system (Applied Biosystems). The thermocycling parameters were as follows: activation of the enzyme at 95°C for 10 min, followed by 40 cycles of a two-step denaturation at 95°C for 15 s and primer annealing/extension.
40% oxygen atmospheres. We hypothesized that sphere compared to those in cultures grown under 20% and kinetics (oxygen association and oxygen disassociation rates, with this model, Tjepkema et al. (26) observed rapid oxygen their rates of respiration under hypoxic conditions. Consistent trHbO may deliver oxygen to terminal oxidases to stimulate expression levels of both trHb genes relative to glnA expression were similar (Fig. 1A). Analysis of the intensities of the bands for the three genes confirmed that the expression levels of both trHb genes relative to glnA expression were similar (data not shown). These results indicate that hemoglobin expression was not regulated by cellular nitrogen status.

Effect of oxygen conditions on gene expression. Schwintzer et al. (22) observed an increase in total hemoglobin levels in Frankia strain ArI3 cultures grown under a 1% oxygen atmosphere compared to those in cultures grown under 20% and 40% oxygen atmospheres. We hypothesized that Frankia trHbO may deliver oxygen to terminal oxidases to stimulate their rates of respiration under hypoxic conditions. Consistent with this model, Tjepkema et al. (26) observed rapid oxygen kinetics (oxygen association and oxygen disassociation rates, k'_{oa} and k_{oa}) for Frankia hemoglobin, suggesting the role of oxygen transport over short distances. In M. tuberculosis, trHbO has been shown to function as an oxygen shuttle to respiratory enzymes (10, 12, 18).

Frankia strain CcI3 cultures were grown for 7 days underoxic conditions (cultures aerated with atmospheric oxygen) and hypoxic conditions (static cultures). The relative expression of trHbO was greater in static cultures than in aerated cultures, while trHbN expression levels were similar under both conditions (Fig. 1B). These results suggest that Frankia TrHbO may function under hypoxic conditions to shuttle oxygen to the respiratory chain, similar to mycobacteria.

Effect of NO stress on gene expression. During intracellular pathogenesis, mycobacteria are bombarded with toxic NO species generated by macrophages in tuberculosis granulomas. The trHbN protein functions as a protective molecule, with NO catalytically reacting with oxygen to generate a harmless nitrate molecule (6, 11, 13, 17, 19). Since plants also use NO as a defense mechanism against pathogens (14), we predicted a similar protective function in Frankia. To test our hypothesis, the spontaneous NO donor S-nitroso-N-acetylpenicillamine (SNAP) was added to Frankia strain CcI3 cultures and the relative expression levels were evaluated. RNA samples were taken every hour, and Fig. 2 shows the RT-PCR results from these experiments. The relative expression level of hboO did not change under any of the test conditions. However, the relative expression level of hboN increased through the first 3 h of NO exposure (with SNAP), and the signal decreased after 4 h. The addition of carboxy-PT10 (cPT10), an NO scavenger, to cultures growing in the presence of SNAP decreased the level of hboN expression. Control cultures without SNAP showed no change in their relative expression levels over the same time period. These results indicate that Frankia’s hboN gene expression was stimulated by NO and suggests that Frankia trHbN may be involved in nitric oxide detoxification.

The above-mentioned experiments were also performed with Frankia strain EAN1pec, and similar results were obtained for hboN and hboO gene expression levels in response to nitrogen status, oxygen content, and NO stress (data not shown). These results indicate that these responses were not specific to Frankia strain CcI3.

qPCR results. To support the results from the above-described expression studies, we utilized quantitative RT-PCR (qRT-PCR) to measure gene expression in Frankia strain CcI3. In addition to repeating our previous experiments, H_{2}O_{2} and paraquat were also added to Frankia cultures to evaluate gene expression under oxidative stress conditions. Figure 3 shows the results of these experiments. For each experimental con-
condition, gene expression levels are shown as a ratio to their expression levels in the untreated control cultures.

The semi-qRT-PCR data for *Frankia* strain Cd3 cultured under nitrogen-limiting conditions supported the RT-PCR results described above. There was no substantial difference in gene expression level for either *hboO* or *hboN* under nitrogen-sufficient and -limiting conditions (Fig. 3). *hboN* expression levels were similar under both conditions, while the *hboO* expression level showed a small increase under growth with NH4Cl. These data, in addition to the previous RT-PCR results, suggest that these two hemoglobin genes were not upregulated under nitrogen fixation conditions.

The oxidative burst of reactive oxygen species by plants is another common defense mechanism against invading pathogens (24). One problem that microbial symbionts face is that they may not be initially distinguished by the host as a friend and not a foe. Thus, a defense mechanism must be utilized to help establish this early stage of symbiosis. Both paraquat and \( \text{H}_2\text{O}_2 \) were used to test the effects of oxidative stress on gene expression. The addition of 1.0 mM \( \text{H}_2\text{O}_2 \) caused a small increase in *hboN* (1.5-fold) or *hboO* (2-fold) gene expression compared to that in the control culture (Fig. 3). Interestingly, the addition of 0.1 mM paraquat, which is metabolized to produce endogenous \( \text{H}_2\text{O}_2 \), decreased the expression levels of both *hboN* and *hboO* nearly 10-fold. It is unclear whether or not these decreases in expression were due to the specific downregulation of these genes or possibly to the cytotoxic effects of intracellular concentrations of \( \text{H}_2\text{O}_2 \).

To quantify the effect of nitrosative stress on gene expression, *Frankia* strain Cd3 cultures were exposed to SNAP and/or cPT10 for 3 h before RNA samples were extracted. This time point yielded the strongest band intensity in the previous RT-PCR experiments for *hboN* (Fig. 2). The expression of *hboN* increased nearly 10-fold in cells exposed to 400 \( \mu \text{M} \) SNAP compared to that in the control (untreated) cells (Fig. 3). The addition of 400 \( \mu \text{M} \) cPT10 reduced the effect of 400 \( \mu \text{M} \) SNAP and resulted in only a fourfold increase in *hboN* expression. This result indicates that cPT10 was unable to scavenge all of the NO generated by SNAP under these conditions. *hboO* expression was reduced 0.5-fold under these conditions. Since cPT10 was unable to relieve this effect, this result would suggest that this reduction was not an NO-specific effect.

This NO-specific induction of *hboN* is physiologically relevant for *Frankia*. During initiation of nodule formation, the nitrogen-fixing microsymbiont is initially recognized as an intruder and subjected to host defenses (20). Microsymbionts have developed mechanisms to adapt and regulate these host responses (for a review, see reference 24). NO, the key regulatory molecule of plant pathogen responses (7, 9), has been shown to accumulate in functional legume nodules (2) and also functions as a defense mechanism. Thus, *Frankia* HboN could function to detoxify NO during the initiation stages of the host plant infection process to establish symbiosis or to maintain a functional nodule structure.

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