

# The Nitrogenase Regulatory Enzyme Dinitrogenase Reductase ADP-Ribosyltransferase (DraT) Is Activated by Direct Interaction with the Signal Transduction Protein GlnB

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Fe protein (dinitrogenase reductase) activity is reversibly inactivated by dinitrogenase reductase ADP-ribosyltransferase (DraT) in response to an increase in the ammonium concentration or a decrease in cellular energy in *Azospirillum brasilense*, *Rhodospirillum rubrum*, and *Rhodobacter capsulatus*. The ADP-ribosyl is removed by the dinitrogenase reductase-activating glycohydrolase (DraG), promoting Fe protein reactivation. The signaling pathway leading to DraT activation by ammonium is still not completely understood, but the available evidence shows the involvement of direct interaction between the enzyme and the nitrogen-signaling P<sub>II</sub> proteins. In *A. brasilense*, two P<sub>II</sub> proteins, GlnB and GlnZ, were identified. We used Fe protein from *Azotobacter vinelandii* as the substrate to assess the activity of *A. brasilense* DraT *in vitro* complexed or not with P<sub>II</sub> proteins. Under our conditions, GlnB was necessary for DraT activity in the presence of Mg-ADP. The P<sub>II</sub> effector 2-oxoglutarate, in the presence of Mg-ATP, inhibited DraT-GlnB activity, possibly by inducing complex dissociation. DraT was also activated by GlnZ and by both uridylylated P<sub>II</sub> proteins, but not by a GlnB variant carrying a partial deletion of the T loop. Kinetics studies revealed that the *A. brasilense* DraT-GlnB complex was at least 18-fold more efficient than DraT purified from *R. rubrum*, but with a similar *K<sub>m</sub>* value for NAD<sup>+</sup>. Our results showed that ADP-ribosylation of the Fe protein does not affect the electronic state of its metal cluster and prevents association between the Fe and MoFe proteins, thus inhibiting electron transfer.

**B** iological nitrogen fixation is the reduction of atmospheric N<sub>2</sub> to NH<sub>3</sub>, catalyzed by the enzyme nitrogenase. The most common form of nitrogenase is composed of two component proteins: the molybdenum-iron (MoFe) protein (dinitrogenase, or NifDK), a 240-kDa  $\alpha_2\beta_2$  tetramer, and the iron (Fe) protein (dinitrogenase reductase, or NifH), a 64-kDa  $\gamma_2$  homodimer. The Fe protein is responsible for electron transfer to the MoFe protein in a reaction that requires the hydrolysis of two ATP to two ADP molecules and two P<sub>i</sub> for each electron transferred. The MoFe protein contains the N<sub>2</sub> binding/reduction active site (1). In order to avoid wasteful ATP hydrolysis, several microorganisms have acquired diverse mechanisms to inactivate nitrogenase at the posttranslational level when fixed nitrogen (e.g., ammonium) is available in the environment (2).

The best-described nitrogenase posttranslational regulatory mechanism relies on reversible ADP-ribosylation of the Fe protein. This mechanism has been extensively studied in the alphaproteobacteria Azospirillum brasilense and Rhodospirillum rubrum. Furthermore, bioinformatics analysis supports the idea that a similar mechanism operates in at least 25 bacterial genera (2). Fe protein ADP-ribosylation is catalyzed by dinitrogenase reductase ADP-ribosyltransferase (DraT). This enzyme uses NAD<sup>+</sup> as an ADP-ribose donor to modify a conserved arginine (usually Arg101) in one of the subunits of Fe protein (3). The ADP-ribosylation inactivates the Fe protein, presumably by blocking the docking site between the Fe and the MoFe protein (4, 5), preventing the association of the Fe protein with the MoFe protein, and thus electron transfer. The ADP-ribosyl group attached to the Fe protein is removed by dinitrogenase reductase-activating glycohydrolase (DraG), which thereby promotes nitrogenase complex reactivation (6).

The DraT and DraG activities are subject to opposite regulation *in vivo* (7). In response to a negative stimulus, such as the presence of ammonium or a decrease in the available cell energy, DraT is activated and DraG is inactivated, promoting inactivation of nitrogenase by ADP-ribosylation of the Fe protein. Conversely, when the negative stimulus is removed, DraT is inactivated and DraG is activated, favoring ADP-ribose removal from the Fe protein and nitrogenase activity (2, 7). The signaling pathway linking the cellular nitrogen status to regulated activities of DraT and DraG in *A. brasilense* and *R. rubrum* has been extensively studied and relies on the nitrogen signal transduction proteins of the P<sub>II</sub> protein family. In contrast, the energy-signaling pathway remains unknown (2).

The  $P_{II}$  proteins constitute one of the most widely distributed families of signal transduction proteins. These proteins regulate nitrogen metabolism by direct interaction with enzymes, transcriptional regulators, and transporters (8–10).  $P_{II}$  proteins are homotrimers, and each monomer carries a flexible loop region, the T loop, that is a crucial element for  $P_{II}$  function (11). The signaling status of  $P_{II}$  proteins and their capacity to interact with

Received 30 August 2012 Accepted 24 October 2012

Published ahead of print 9 November 2012

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Supplemental material for this article may be found at http://dx.doi.org/10.1128 /JB.01517-12.

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their target proteins are dictated by the structural conformation of the  $P_{II}$  T loops (9, 12).

In proteobacteria, the  $P_{II}$  T loops are subjected to reversible uridylylation at a conserved residue (Tyr51). Uridylylation reflects the cellular availability of the nitrogen-signaling molecule glutamine, so that under low glutamine,  $P_{II}$  is uridylylated (13).  $P_{II}$  is also modulated through the allosteric binding of ATP, ADP, and 2-oxoglutarate (2-OG). The  $P_{II}$  trimer carries three binding sites for each of these molecules located in the lateral clefts between subunits (11, 14, 15). ATP and ADP bind competitively to the same site on  $P_{II}$  (16), whereas 2-OG interacts cooperatively with the bound Mg-ATP (15, 17, 18). Both the occupancy of  $P_{II}$  by ATP, ADP, and 2-OG and the uridylylation status of  $P_{II}$  alter the T-loop conformation and, hence, the ability of  $P_{II}$  to interact with its target proteins (15).

In A. brasilense, two PII proteins sharing 67% identity in their amino acid sequences were detected, namely, GlnB and GlnZ (19). Under nitrogen-fixing conditions, GlnB and GlnZ are fully uridylylated and nitrogenase is active. Upon an ammonium pulse, nitrogenase is inactivated by ADP-ribosylation of the Fe protein. At the same time, GlnB and GlnZ are deuridylylated (20) and interact with DraT and DraG, respectively (21-24). The DraG-GlnZ complex is directed to the membrane by interaction with the ammonium transporter AmtB, resulting in DraG inactivation (20, 22, 24). Given that the DraT-GlnB complex is detected only under conditions where DraT is active, it was postulated that this protein-protein interaction is responsible for DraT activation; however, this has yet to be confirmed (23). The A. brasilense DraT-GlnB complex can be purified after coexpression in Escherichia coli, and the complex exhibits a stoichiometry of 1 DraT monomer to 1 GlnB trimer, is stable in the presence of Mg-ADP, and is destabilized in the presence of Mg-ATP and 2-OG (23).

Here, we used Fe protein from *Azotobacter vinelandii* as the substrate to analyze the activity of *A. brasilense* DraT *in vitro*. The results show that GlnB activates DraT in the presence of Mg-ADP. The DraT-GlnB complex is at least 18-fold more active than DraT alone in inactivating nitrogenase. A GlnB variant carrying a deletion in the T loop failed both to interact with and to activate DraT *in vitro*. These results suggest a model where DraT can be activated by interaction with the GlnB T loop.

## MATERIALS AND METHODS

**Materials and reagents.** All reagents were obtained from Sigma-Aldrich (St. Louis, MO) and were used as provided unless otherwise specified.

Bacterial strains, plasmids, and protein purification. The *A. brasilense* GlnB $\Delta$ 42-54 and GlnZ variants were obtained using the Quick-Change site-directed mutagenesis kit (Agilent, Santa Clara, CA) with the pLH25PET (25) or pMSA4 (26) plasmid as a template, respectively. The DNA inserts of the resulting plasmids, pLH25PET H42Q, pMSA4 Q22HS52VS54D, pMSA4 S52VS54D, and pLHGlnB  $\Delta$ 42-54, were completely sequenced to confirm their integrity.

A. brasilense GlnB (25), GlnZ (26), GlnZQ42HS52VS54D, GlnZS52VS54D, GlnBH42Q, GlnB $\Delta$ 42-54, His-tagged DraT (27), and His-tagged GlnD (28) and recombinant proteins were overexpressed in *E. coli* strain BL21(DE3) harboring the plasmids listed in Table 1. The P<sub>II</sub> proteins (GlnB, GlnZ, and GlnB $\Delta$ 42-54) were purified using a thermal treatment at 70°C for 15 min prior to loading on the 5-ml HiTrap heparin column (GE Healthcare) (26). The P<sub>II</sub> proteins were eluted in a nonlinear KCl gradient using an Äkta system (GE Healthcare). The samples with purity over 97% as judged by SDS-PAGE were pooled, dialyzed, and stored in aliquots at  $-80^{\circ}$ C until use. His-tagged GlnD was purified essentially as described previously (28) using a 1-ml HiTrap chelating HP

TABLE	1	Plasmids
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Plasmid	Genotype	Reference
pLH25 Km <sup>r</sup> (pET28a)	Expresses the A. brasilense GlnB	25
pMSA4 Km <sup>r</sup> (pET28a)	Expresses the A. brasilense GlnZ	26
pLHPETDraT Km <sup>r</sup> (pET28a)	Expresses the N-terminally His- tagged <i>A. brasilense</i> DraT	27
pLHDK5pII Amp <sup>r</sup> (pDK5)	Expresses the A. brasilense GlnB	25
pMSA4 Q42HS52VS54D Km <sup>r</sup> (pET28a)	Expresses the <i>A. brasilense</i> GlnZ variant Q42HS52VS54D	This work
pMSA4 S52VS54D Km <sup>r</sup> (pET28a)	Expresses the <i>A. brasilense</i> GlnZ variant S52VS54D	This work
pLH25PETGlnB H42Q Km <sup>r</sup> (pET28a)	Expresses the <i>A. brasilense</i> GlnB variant H42Q	This work
pGlnB∆42–54 Km <sup>r</sup> (pET28a)	Expresses the <i>A. brasilense</i> GlnB carrying a T-loop deletion $(\Delta 42-54)$	This work
pGlnDHis Km <sup>r</sup> (pET28a)	Expresses the N-terminally His- tagged <i>A. brasilense</i> GlnD	28

column prepared as recommended by the manufacturer (GE Healthcare). Uridylylated GlnB and GlnZ were obtained as described previously (23). His-tagged DraT was purified using a 1-ml HiTrap chelating HP column preequilibrated with buffer A (50 mM Tris-HCl, pH 8.0). The protein was eluted in a stepwise increase of imidazole (50, 100, 200, 300, 400, and 500 mM) in buffer A, using 5 column volumes at each concentration. Soluble fractions with purity over 95% as judged by SDS-PAGE were pooled, kept in an ice bath, and used immediately. For the purification of the A. brasilense His-DraT-GlnB complex, both proteins were coexpressed, and the complex was purified according to the method of Huergo et al. (23) using a 4-ml immobilized metal ion affinity chromatography (IMAC) column. The complex was dialyzed in buffer containing 10% glycerol, followed by gel filtration chromatography on a Superdex 200 column (320 ml) using 50 mM Tris-HCl, pH 8.0, 10% glycerol, 200 mM NaCl, 1 mM MgCl<sub>2</sub>, 0.5 mM ADP as a buffer. The loaded samples consisted of aggregates of high molecular weight containing both His-tagged DraT and GlnB and a fraction of approximately 64,000 Da, corresponding to the molecular mass of 1 DraT monomer plus 1 GlnB trimer. This low-molecular-weight fraction had higher activity and was used in all subsequent analyses.

The nitrogenase MoFe protein was purified from A. vinelandii strain DJ995 and the Fe protein from strain DJ884 based on the methods of Christiansen and Seefeldt (29, 30), respectively. The cells were grown, and both proteins were expressed. Approximately 240 g of cells (wet weight) was processed for each purification, and the crude extracts were prepared by osmotic shock using glycerol. The DJ995 strain encodes a MoFe protein variant carrying a histidine tag on the α-subunit, which allowed protein purification using metal affinity chromatography (29). Cell extract was loaded onto a Zn(II)-charged IMAC column (60 ml of resin) using a peristaltic pump. After loading the extract, the column was washed with 2 column volumes of buffer A (50 mM Tris-HCl, pH 7.9, 500 mM NaCl, 1 mM sodium dithionite) containing 20 mM imidazole. The protein that remained bound to the column was then eluted using a linear gradient of imidazole in an Äkta system (GE Healthcare). The eluted protein was collected at approximately 250 mM NaCl and diluted 8-fold in a degassed buffer (50 mM Tris-HCl, pH 8.0) containing 1 mM sodium dithionite. The diluted protein was then loaded onto a Q-Sepharose column (30 ml of resin) and eluted using a linear NaCl gradient (100 to 300 mM NaCl over 5 column volumes). DJ884 encodes a wild-type Fe protein, which was purified using standard methods (30). Cell extract was loaded onto a Q-Sepharose column (150 ml of resin) using a peristaltic pump. After loading the extract, the column was washed with 1 column volume of no-salt buffer (50 mM Tris-HCl, pH 8.0) containing 1 mM sodium dithionite. The protein that remained bound to the column was then eluted

using 4 column volumes of a linear gradient of NaCl in an Äkta system (GE Healthcare). The eluted protein was collected at approximately 350 mM NaCl and concentrated in a Q-Sepharose column (40 ml of resin) by loading the protein fraction, washing with 50 ml of no-salt buffer, reversing the flow, and eluting with salt buffer (50 mM Tris-HCl, pH 8.0, 1 M NaCl) containing 2 mM sodium dithionite. The concentrated protein was then loaded onto a Sephacryl S-200 column (600 ml of resin) and eluted using a salt buffer (50 mM Tris-HCl, pH 8.0, 500 mM NaCl) containing 2 mM sodium dithionite. Proteins were concentrated using an Amicon concentrator (Beverly, MA) fitted with a YM100 (for MoFe) and YM30 (for Fe) membrane. The proteins were quantified by the biuret method using bovine serum albumin as the standard, and protein purity was monitored by SDS-PAGE. The final purified proteins were pelleted and stored in liquid nitrogen until use. The Fe protein used in these experiments had purity of over 90% and specific activities for acetylene and proton reduction (30, 31) greater than 2,000 nmol of acetylene reduced or H<sub>2</sub> evolved min<sup>-1</sup> (mg Fe protein)<sup>-1</sup>. All protein manipulations were conducted in septum-sealed serum vials degassed under an argon atmosphere, and all anaerobic liquid and gas transfers were performed using gas-tight syringes.

**DraT-P**<sub>II</sub> **complex formation** *in vitro*. Purified proteins [DraT and GlnB, GlnZ, GlnB-(UMP)<sub>3</sub>, GlnZ-(UMP)<sub>3</sub>, or GlnB $\Delta$ 42-54] were incubated with buffer (50 mM Tris-HCl, pH 7.5, 2 mM MgCl<sub>2</sub>, and 2 mM ADP) at 4°C overnight using the molar ratio indicated, considering the P<sub>II</sub> protein a trimer.

**Protein pulldown assay.** *In vitro* protein complex formation was performed using MagneHis-Ni<sup>2+</sup> beads (Promega Co., Madison, WI) as described previously (23). Reactions were conducted in buffer containing 50 mM Tris-HCl, pH 8.0, 100 mM NaCl, 10% glycerol, 20 mM imidazole, 0.05% (vol/vol) Tween 20, 5 mM MgCl<sub>2</sub>, and 1 mM ADP with 5  $\mu$ g of His-tagged DraT and 10  $\mu$ g of P<sub>II</sub>. After washing, loading buffer was added, and the samples were boiled for 5 min prior to 12.5% SDS-PAGE analysis. The gels were stained with Coomassie blue. The data reported were confirmed in at least two independent experiments.

DraT activity assays. The standard assay for DraT activity was carried out using 20 mM MOPS (morpholinepropanesulfonic acid), pH 7.5, 2 mM NAD<sup>+</sup>, 1 mM ADP, 1 mM MgCl<sub>2</sub>, 100 µg of unmodified oxidized Fe protein, and the indicated amounts of DraT or the DraT-P<sub>II</sub> complex in a 500-µl reaction mixture. The oxidized Fe protein was prepared by passing the reduced Fe protein over a 3-ml G-25 Sephadex column to remove the excess dithionite, which can reduce NAD<sup>+</sup> to NADH (32). The samples were incubated for 5 min at 130 rpm at 30°C. The DraT activity was determined indirectly by measuring the Fe protein activity based on H<sub>2</sub> evolution, as described previously (31) with modifications. The nitrogenase activity buffer contained 200 mM MOPS, 13.4 mM MgCl<sub>2</sub>, 60 mM phosphocreatine, 10 mM ATP, 2.3 mg/ml bovine serum albumin (BSA), 0.4 mg/ml creatine phosphokinase, and 24 mM dithionite. This buffer was added to a 1-ml final volume immediately after incubation for DraT activity. The dithionite reduces the NAD<sup>+</sup>, quenching the DraT reaction. To determine Fe protein activity, MoFe protein was added in a 1:1.4 molar ratio (MoFe/Fe protein), the reaction mixtures were incubated for 8 min at 160 rpm at 30°C, and H<sub>2</sub> evolution was measured by gas chromatography. The pattern of ADP-ribosylation was analyzed by 12% low-crosslinker SDS-PAGE as described previously (33).

The DraT assays were conducted in a small vial inside a bigger septumsealed vial. A dithionite solution (12 mM) was used outside the reaction vial to act as an oxygen scavenger.

**EPR spectroscopy.** Resting state and turnover samples of ADP-ribosylated Fe (Fe-ADPR) protein, prepared as described for the DraT activity assay, were monitored by electron paramagnetic resonance (EPR) spectroscopy essentially as described previously (34) and compared with unmodified Fe protein.

**Stopped-flow spectrophotometry.** Fe protein was modified using NAD<sup>+</sup> in a standard reaction in the presence of the DraT-GlnB complex. To remove the complex, the solution was passed over a 2-ml IMAC col-

umn in the presence of buffer (50 mM Tris-HCl, pH 8.0, 200 mM NaCl, and 0.5 mM ADP). To remove ADP interference, the sample was exchanged with the same buffer in the absence of Mg-ADP and then concentrated in an Amicon cell using a membrane with a 30,000-Da cutoff.

Electron transfer from modified Fe protein to MoFe protein was assessed by absorbance at 430 nm. Observing a change in absorbance at 430 nm is indicative of the redox state change due to one-electron oxidation of the Fe protein metal cluster. This change in absorbance was monitored over time in a Hi-Tech SF61 stopped-flow UV-visible (Vis) spectrophotometer equipped with a data acquisition and curve-fitting system. The reactions were carried out at 25°C under conditions described previously (35).

**Statistical analysis.** Statistical analysis was carried out by one-tailed unpaired t test, using the Graph Pad Prism 5 program. The number of experimental replicates was at least three independent experiments. *P* values of less than 0.05 were considered statistically significant.

## RESULTS

DraT is activated by GlnB in vitro. We have shown previously that A. brasilense DraT interacts with deuridylylated GlnB in vivo and in vitro and that His-tagged DraT has activities comparable to those of the wild-type protein in vivo (20, 21, 23). To assess whether GlnB-DraT complex formation altered DraT activity in vitro, the activity of DraT was compared to that of the DraT-GlnB complex that was obtained either by copurification or by in vitro reconstitution using the purified proteins. DraT activity was determined indirectly by measuring nitrogenase H<sub>2</sub> evolution using the modified Fe protein. For these assays, His-tagged DraT or the DraT-GlnB complex was preincubated with the oxidized A. vinelandii Fe protein in the presence of Mg-ADP and NAD<sup>+</sup> under anaerobic conditions and in the absence of dithionite. After 5 min at 30°C, the ADP-ribosylation reaction was quenched by adding nitrogenase assay buffer containing excess dithionite to completely reduce NAD<sup>+</sup>. The purified A. vinelandii MoFe protein was added to the reaction mixture, and the activity was determined by measuring  $H_2$  production.

Control reactions in the absence of NAD<sup>+</sup> indicated that the Fe protein was fully active and not modified in the presence of the DraT-GlnB complex that was obtained by coexpression and copurification (Fig. 1A and B). Similar activities were observed using DraT alone in the presence of NAD<sup>+</sup> (Fig. 1A). Hence, DraT had no significant activity under these conditions. Preincubation of the Fe protein with the DraT-GlnB complex that was obtained by copurification resulted in complete Fe protein inactivation (Fig. 1A) and ADP-ribosylation (Fig. 1B). Furthermore, preincubation of the Fe protein with DraT in the presence of increasing concentrations of GlnB resulted in Fe protein inhibition in a GlnB-dependent manner, with the highest inhibition when DraT-GlnB was used at a 1:2 molar ratio (Fig. 1A). Under similar conditions, the copurified complex was approximately 3-fold more efficient to ADP-ribosylate the Fe protein than the in vitro-reconstituted complex (260 nmol of Fe protein min<sup>-1</sup> mg DraT<sup>-1</sup> compared to 97 nmol min<sup>-1</sup> mg  $DraT^{-1}$ ) (Fig. 1A). To verify the efficiency of *in vitro* complex formation, DraT and GlnB were mixed at a 1:2 molar ratio and analyzed by pulldown of the His-tagged DraT using Ni<sup>2+</sup> beads. Quantification of the protein bands showed that only 56% of DraT was complexed to GlnB (Fig. 2B). This result may explain the lower efficiency of the in vitro-reconstituted complex in inhibiting the Fe protein in comparison to the DraT-GlnB complex obtained by copurification.



FIG 1 Fe protein activity and modification after DraT-GlnB complex activity assay. (A) Fe protein activity was determined after incubation at different DraT/GlnB (monomer/trimer) ratios or in the presence of the DraT-GlnB complex obtained by copurification. The reaction mixtures for the DraT assay contained 2 mM NAD<sup>+</sup>; 1 mM MgCl<sub>2</sub>; 1 mM ADP; 100 µg of A. vinelandii Fe protein; and DraT, DraT-GlnB complex formed in vitro, or copurified complex and were incubated for 5 min at 130 rpm at 30°C. The reactions were performed using an Fe protein/DraT (dimer/monomer) molar ratio of 42:1. As a control, the reaction was performed in the presence of either copurified complex in the absence of NAD. The DraT-GlnB assay was stopped by reducing NAD<sup>+</sup> using dithionite solution, the Fe protein activity was determined by adding 500 µg of the MoFe protein, and the hydrogen evolution was measured. ns, nonsignificant; \*\*\*, P < 0.001. The error bars represent the calculated standard deviations. (B) NAD<sup>+</sup>-dependent modification of A. vinelandii Fe protein catalyzed by the A. brasilense DraT-GlnB complex. Lane 1, control containing purified, unmodified A. vinelandii Fe protein. Lanes 2 and 3, purified A. vinelandii Fe protein was incubated with the A. brasilense copurified DraT-GlnB complex in the presence (lane 3) or absence (lane 2) of NAD<sup>+</sup>. The unmodified and ADP-ribosylated subunits of the Fe protein were separated using low-cross-link SDS-PAGE, and the gel was stained with Coomassie blue. The percentages of Fe-ADPR and unmodified Fe protein were quantified using Photoshop CS4, resulting in 42 and 58%, respectively.

**DraT activation** *in vitro* requires complex formation with the GlnB T loop. *A. brasilense* encodes two highly similar  $P_{II}$  proteins, GlnB and GlnZ (19), whereas only GlnB seems able to interact with DraT (21, 23). Unexpectedly, the presence of GlnZ in the DraT assay resulted in significant DraT activation in comparison to DraT alone (Fig. 2A), although this GlnZ-induced activation was less than half of that observed with GlnB (Fig. 2A). Given the high similarity between GlnB and GlnZ (see Fig. S1 in the supplemental material), it is possible that GlnZ interacts with DraT to promote its activation. Likewise, the two  $P_{II}$  proteins of *Rhodobacter capsulatus* were able to interact with DraT using yeast two-hybrid assays (36). However, in *A. brasilense*, the lower affinity of DraT for GlnZ precluded detection of the protein complex in pulldown assays (Fig. 2B).

The formation of the DraT-GlnB complex is affected by the uridylylation of GlnB (23). When GlnB-(UMP)<sub>3</sub> or GlnZ-(UMP)<sub>3</sub> was used in the DraT assay, it also resulted in significant DraT activation, though this activation was approximately 2-fold lower



FIG 2 DraT activation and interaction with P<sub>II</sub> proteins. (A) DraT activity was assessed indirectly from Fe protein activity by measuring hydrogen evolution. A DraT assay was performed using A. vinelandii Fe protein as the substrate in the presence of different P<sub>II</sub> proteins to form the *in vitro* complexes. Reaction mixtures containing 2 mM NAD+; 1 mM MgCl<sub>2</sub>; 1 mM ADP; 100 µg Fe protein; and DraT, DraT-P<sub>II</sub> complexes formed in vitro, or copurified DraT-GlnB were incubated for 5 min at 130 rpm at 30°C. The reactions were performed using a DraT/P<sub>II</sub> trimer molar ratio of 1:2 and an Fe protein/DraT molar ratio of 21:1. For a control, the reaction was performed in the presence of either copurified complex, DraT-GlnB (1:2, as shown), or DraT in the absence of NAD<sup>+</sup>. The DraT activity was stopped by reducing NAD<sup>+</sup> using dithionite solution. The Fe protein activity was determined by adding 500 µg of MoFe protein, and the hydrogen production was measured. The activity of the copurified DraT-GlnB complex was 130  $\pm$  2.5 nmol ADP-ribosylated Fe protein min<sup>-1</sup> mg DraT<sup>-1</sup> under the same conditions, corresponding to 100% ADP-ribosylation of Fe protein. ns, nonsignificant; \*, P < 0.05; \*\*, P < 0.01; \*\*\*, *P* < 0.001. The error bars represent the calculated standard deviations. (B) Complex formation between His-tagged DraT and the P11 protein GlnB, GlnZ, or GlnB∆42-54 was assessed by pulldown in the presence of 1 mM ADP. Proteins bound to Ni<sup>2+</sup> beads were analyzed by SDS-PAGE, and the gel was stained with Coomassie blue.

than the activation observed using unmodified GlnB (Fig. 2A). Yeast two-hybrid analysis revealed that in R. rubrum the GlnB T-loop mutants failed to interact with DraT (37). In order to further exploit the role of the GlnB T loop, a GlnB variant carrying a partial deletion of the T loop (GlnB $\Delta$ 42-54) was purified and tested for in vitro interaction with DraT. In contrast to the wildtype GlnB, the GlnB $\Delta$ 42-54 variant did not coprecipitate with His-tagged DraT in pulldown assays (Fig. 2B) and did not significantly activate DraT in vitro (Fig. 2A). The deletion of the GlnB T loop did not affect the quaternary structure of the GlnB $\Delta$ 42-54 variant as judged by migration rates on native PAGE (see Fig. S2 in the supplemental material). Furthermore, the deletion of residues 42 to 54 in the highly similar GlnZ protein did not affect binding of ATP and ADP (24), supporting the conclusion that the GlnB $\Delta$ 42-54 variant retains its overall fold and effector binding properties. Comparison of the GlnB and GlnZ T-loop sequences revealed only 3 nonconserved substitutions at positions 42, 52, and 54 (see Fig. S1 in the supplemental material). When the GlnZ residues at positions 52 and 54 were replaced by the respective residues found in GlnB, interaction between the GlnZ variants (Q42HS52VS54D and S52VS54D) and DraT could be detected by pulldown assay (see Fig. S3 in the supplemental material), reinforcing the involvement of the T loop in the DraT-GlnB interaction. On the other hand, the GlnB T-loop H42 residue seems not to be important for DraT binding, since a GlnB H42Q variant was able to interact with DraT (see Fig. S3 in the supplemental material). Together, these results support a model where the major docking site for DraT is the GlnB T loop and show that this interaction is required for DraT activation.



**FIG 3** Effects of ADP, ATP, and 2-oxoglutarate on the activity of the DraT-GlnB complex. DraT activity was assessed indirectly from Fe protein activity by measuring hydrogen evolution. Reaction mixtures for the DraT assay contained 4 mM MgCl<sub>2</sub>, copurified DraT-GlnB complex, 100 µg *A. vinelandii* Fe protein, and the effectors, as indicated. For a control, the reaction was performed in the presence of copurified DraT-GlnB complex in the absence of NAD<sup>+</sup>. The DraT activity was stopped by reducing NAD<sup>+</sup> using dithionite solution. The Fe protein activity was determined by adding 500 µg of MoFe protein, and the hydrogen production was measured. ns, nonsignificant; \*, *P* < 0.05. The error bars represent the calculated standard deviations.

P<sub>II</sub> effectors ATP, ADP, and 2-oxoglutarate modulate GlnBdependent DraT activation. We have previously shown that the A. brasilense DraT-GlnB complex is stable in the presence of Mg-ADP and is destabilized in the presence of Mg-ATP and 2-OG combined (23). In order to exploit the role of the  $P_{II}$  effectors ATP, ADP, and 2-OG in GlnB-dependent DraT activation, DraT activity was assayed in the presence of added ATP and ADP combined and under different 2-OG levels reflecting the reported physiological range (38). The DraT-GlnB complex (obtained by copurification) was inactive in the presence of ATP without added ADP (see Fig. S4 in the supplemental material). The activities of the DraT-GlnB complex were similar when using ADP alone or using ATP plus ADP at different ATP/ADP ratios (Fig. 3). However, at high ATP/ADP ratios and in the presence of a high 2-OG concentration (2 mM), the activity was inhibited (Fig. 3). These results are in agreement with our previous results reporting on in vitro complex formation between DraT and GlnB (23) and strongly support the idea that only the ADP-bound conformation of the GlnB T loop is able to activate DraT.

**GlnB acts by enhancing the**  $k_{cat}$  and  $k_{cat}/K_m$  of **DraT.** To further investigate the mechanism of DraT activation by GlnB, we determined the  $K_m$  value for NAD<sup>+</sup> of the DraT-GlnB complex (Table 2; see Fig. S5 in the supplemental material). It was not possible to determine the  $K_m$  for NAD<sup>+</sup> using *A. brasilense* DraT alone, given its extremely low activity. The  $K_m$  value determined for NAD<sup>+</sup> using the *A. brasilense* DraT-GlnB complex is within the same order of magnitude as the value reported using DraT purified from the related organism *R. rubrum* (Table 2) (39, 40). Furthermore, the NAD<sup>+</sup>  $K_m$  was well below the reported intracellular concentration of NAD<sup>+</sup> in *E. coli* (about 2 mM) (41). This suggests NAD<sup>+</sup> binding by DraT is not affected by interaction with GlnB. On the other hand, both the  $k_{cat}$  and the  $k_{cat}/NAD^+ K_m$  ratio for the DraT-GlnB complex are well above the reported values for DraT alone; notably, the DraT-GlnB complex was at least

18-fold more efficient than DraT purified from *R. rubrum* (Table 2).

As we did not determine a  $K_m$  value for the Fe protein, we cannot determine whether GlnB acts to enhance the affinity of DraT for the Fe protein and/or simply affects DraT catalytic efficiency. A previous report indicated that NAD<sup>+</sup> binds to the Fe protein-DraT complex but not to DraT alone (42). Thus, given that DraT-GlnB complex formation did not significantly affect the DraT  $K_m$  for NAD<sup>+</sup> (Table 2), it is likely that GlnB acts to enhance the catalytic efficiency of DraT rather than to enhance the affinity for binding the Fe protein.

ADP-ribosylation of Fe protein blocks its ability to transfer electrons to the MoFe protein. The current model for nitrogenase inactivation by Fe protein ADP-ribosylation is based on studies by Murrel and coauthors (4). They provided indirect evidence that ADP-ribosylation of the Fe protein prevents its interaction with the MoFe protein, thus blocking the electron transfer between these proteins. Structural analysis of the nitrogenase complex revealed that, indeed, the site of Fe protein ADP-ribosylation is located in the docking surface for the MoFe protein (5). To evaluate if the ADP-ribosylation of the Fe protein could somehow affect the [4Fe-4S] cluster, we performed EPR experiments, which showed that this modification does not affect the electronic state of the reduced metal cluster ( $[4Fe-4S]^{1+}$ ) bound to Mg-ADP (see Fig. S6 in the supplemental material). This result agrees with the midpoint potential being unaffected by the DraT-catalyzed modification (43). Furthermore, the turnover sample prepared in nitrogen gas demonstrated no association and/or no electron transfer from the Fe to the MoFe protein (data not shown).

Finally, to determine if ADP-ribosylated Fe protein could be weakly associated with the MoFe protein, which could start the first electron transfer based on the deficit spending mechanism recently described (35), we monitored the electron transfer from the Fe protein to the MoFe protein using stopped-flow spectrophotometry. We monitored the absorbance at 430 nm from 3 ms to 50 ms after mixing to observe the change in the redox state of Fe protein due to its oxidation. There was no change in absorbance for ADP-ribosylated Fe protein (Fig. 4), confirming a previous hypothesis (4) that ADP-ribosylation prevents association of the Fe protein with the MoFe protein.

## DISCUSSION

The mechanism of nitrogenase regulation through reversible ADP-ribosylation has been the subject of studies since the late

 TABLE 2 Comparison of kinetic constants for reactions catalyzed by

 DraT from R. rubrum or the DraT-GlnB complex from A. brasilense

 using A. vinelandii

 Fe protein as the acceptor

Fe protein as the acceptor of ADP-ribose	V <sub>max</sub> <sup>a</sup>	$k_{\text{cat}} \ (\text{s}^{-1})^b$	$K_m$ $(\mu M)^c$	$\frac{k_{\text{cat}}/K_m}{(\text{s}^{-1}\text{M}^{-1})}$	Reference
A. brasilense DraT-GlnB	6.72	0.501	166	3,018.1	This work
R. rubrum DraT	0.072 1.48	0.0022 0.044	22 268	100 164.2	39 40

<sup>*a*</sup> The values represent nmol of ADP-ribose transferred  $s^{-1}$  mg DraT<sup>-1</sup> or nmol ADP-ribosylated Fe protein  $s^{-1}$  mg DraT-GlnB complex<sup>-1</sup>.

 $^{c}K_{m}$  for NAD<sup>+</sup> in the presence of Mg-ADP.

<sup>&</sup>lt;sup>b</sup> DraT was considered 30,000 Da and DraT-GlnB complex 74,504 Da (calculated based on the sequence using the Expasy program).



FIG 4 Electron transfer from the Fe protein to the MoFe protein monitored by stopped-flow spectroscopy. Shown is the absorbance at 430 nm plotted against the time after mixing ADP-ribosylated or unmodified Fe protein (68  $\mu$ M) and MoFe protein (18  $\mu$ M) in one syringe with buffer (100 mM MOPS, pH 7.4, and 18 mM Mg-ATP) or in the absence of nucleotide as a control.

1970s. In the 1980s, the enzymes responsible for Fe protein ADPribosylation, DraT and DraG, were isolated and characterized *in vitro* (3, 6). A series of studies revealed that DraT and DraG have opposite regulation *in vivo* in the presence of negative effectors, such as ammonium or anaerobiosis in *A. brasilense* and darkness in *R. rubrum* (33, 44, 45).

In the last decade, it has become clear that proteins from the  $P_{\rm II}$  family are involved in the regulation of DraT in response to ammonium levels (2). The DraT enzyme interacts with GlnB in *A. brasilense* (21, 23), *R. rubrum* (37), and *R. capsulatus* (36). The DraT-GlnB interaction is regulated by the nitrogen status *in vivo*. Under nitrogen fixation conditions, DraT is inactive, whereas upon an ammonium shock, DraT binds to deuridylylated GlnB and is presumably activated (21).

In this work, we used purified proteins to confirm for the first time that GlnB activates DraT *in vitro* (Fig. 1). A GlnB variant carrying a deletion in the T loop (GlnB $\Delta$ 42-54) failed to interact and activate DraT, suggesting that enzyme activation is mediated by interaction between DraT and the GlnB T loop (Fig. 2). Indeed, yeast two-hybrid analysis in *R. rubrum* showed that amino acid substitutions on the GlnB T loop impaired DraT-GlnB interaction (37). Furthermore, when only 2 amino acids of the GlnZ T loop were replaced with those found in GlnB, the GlnZ variants (Q42HS52VS54D and S52VS54D) were able to interact with DraT at detectable levels (see Fig. S3 in the supplemental material).

The hypothesis that the interaction of DraT with GlnB results in enzyme activation is in accordance with *in vivo* analysis of  $P_{II}$ mutants that failed to modify the Fe protein after an ammonium shock (46–48). However, this is in contrast to a previous model that suggested the presence of a negative regulator of DraT *in vivo* (7). The early model was based on the fact that *R. rubrum* DraT was active *in vitro* after purification (3). However, the authors noted that DraT was only stable when Mg-ADP was present during purification. Furthermore, the DraT preparations used in that work had a contaminant with the molecular weight expected for the GlnB monomer, and attempts to remove this contaminant resulted in loss of DraT activity (3). Hence, it is likely the previously described DraT active *in vitro* had copurified GlnB.

Our results showed that the ability of GlnB to activate DraT was positively influenced by ADP and negatively influenced by 2-OG in the presence of high ATP/ADP ratios (Fig. 3), agreeing with our previous protein interaction studies (23). In prokaryotes, the 2-OG levels report the nitrogen status: the molecule accumulates under nitrogen limitation and decreases in response to an ammonium upshift (38, 49, 50). Furthermore, the binding of ATP and ADP to GlnB is influenced by 2-OG so that GlnB is presumably saturated with Mg-ATP and 2-OG under nitrogen-fixing conditions and with Mg-ADP after an ammonium upshift (18, 38, 51). Hence, an ammonium upshift would increase the affinity of DraT for GlnB bound to ADP, resulting in DraT activation. In addition, under nitrogen-fixing conditions, GlnB is fully uridylylated, and this form is a weaker activator of DraT (Fig. 3). The concerted action of GlnB effectors and uridylylation would ensure that GlnB will interact with DraT to activate the enzyme only when ammonium is abundant.

Comparison between the kinetic constants of the *A. brasilense* DraT-GlnB complex and those published for DraT from *R. rubrum* strongly suggests that GlnB acts to enhance the enzyme's catalytic efficiency, although our data cannot precisely define the mechanism of DraT activation by GlnB. Previous *in vitro* analysis showed that some of the properties of His-tagged DraT, including solubility and affinity for Ni<sup>2+</sup>, changed in response to interaction with GlnB, suggesting conformational changes in DraT upon GlnB binding (23). Such conformational changes could result in a DraT structure with improved catalytic efficiency.

Finally, we noted that despite the proximity of the Fe protein ADP-ribosylation site to the [4Fe-4S] cluster ligand Cys-98, the EPR signal of iron in this cluster is not affected by ADP-ribosylation. Furthermore, ADP-ribosylation of the Fe protein completely abolished its ability to transfer electrons to the MoFe protein, as observed by stopped-flow spectrophotometric analysis, confirming that ADP-ribosylation of the Fe protein prevented its association with the MoFe protein, thus inhibiting nitrogenase activity.

#### ACKNOWLEDGMENTS

We are grateful to Luiza M. Araujo for kindly providing the plasmid to express GlnD protein.

V.R.M. was the recipient of a mobility fellowship from CAPES (process number 0388/11-4). This work was supported by CNPq, INCT da Fixação de Nitrogênio/MCT, Fundação Araucária, and CAPES.

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