The *Prevotella ruminicola* 23 genome encodes three different glutamine synthetase (GS) enzymes: glutamine synthetase I (GSI) (ORF02151), GSIII-1 (ORF01459), and GSIII-2 (ORF02034). GSI, GSIII-1, and GSIII-2 have each been heterologously expressed in and purified from *Escherichia coli*. The subunit molecular mass of GS was 56 kDa, while GSIII-1 and GSIII-2 were both 83 kDa. Optimal conditions for γ-glutamyl transferase activity were found to be 35°C at pH 5.6 with 0.25 mM Mn²⁺ ions (GSI) or 37°C at pH 6.0 (GSIII-1 and GSIII-2) with 0.50 to 1.00 mM Mn²⁺ ions. GSIII biosynthetic activity was found to be optimal at 50 to 60°C and pH 6.8 to 7.0 with 10 mM Mn²⁺ ions, while GSI displayed no GS biosynthetic activity. Kinetic analysis revealed *Kₘ* values for glutamate and ammonium as well as for hydrolysis of ATP to be 8.58, 0.48, and 1.91 mM, respectively, for GSIII-1 and 1.72, 0.43, and 2.65 mM, respectively, for GSIII-2. A quantitative reverse transcriptase PCR assay (qRT-PCR) revealed GSIII-2 to be significantly induced by high concentrations of ammonia, and this corresponded with increases in measured GS activity. Collectively, these results show that both GSIII enzymes in *P. ruminicola* 23 are functional and indicate that GSIII-2, flanked by GOGAT (*gltB* and *gltD* genes), plays an important role in the acquisition and metabolism of ammonia, particularly under nonlimiting ammonia growth conditions.

Interestingly, as more genomes are being sequenced, it is clear that many organisms possess multiple enzymes of each type. Based on bioinformatic analysis, three genes encoding glutamine synthetase (one GS with an *Mₜ* of ca. 56,000 and two GSII enzymes with *Mₜ* of ca. 83,000) were identified in the genome of *P. ruminicola* 23. Here we describe the biochemical characterization and transcriptional expression of all three glutamine synthetases and interpret their roles in ammonia assimilation and nitrogen metabolism. Ammonia nitrogen exists in aqueous solution as either NH₄⁺ or NH₃ depending on pH, with a pKₐ of 9.25 (23°C). Thus, at ruminal pH and under buffer conditions for optimal activity, the bulk of ammonia is in the ionized form (NH₄⁺) and in this paper we simply use the term ammonia for the sum of ionized and un-ionized forms unless specified.

**MATERIALS AND METHODS**

**Organism and culture conditions.** *Prevotella ruminicola* strain 23 was kindly provided by M. A. Cotta, USDA-ARS, Peoria, IL, and stored on maintenance slants in the vapor phase of liquid nitrogen in our laboratory.
TABLE 1 Primers used for cloning and qRT-PCR

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>GSI-F</td>
<td>5'-catATGAAACAGCAACGAGCTGATGTTGAAT-3'</td>
</tr>
<tr>
<td>GSI-R</td>
<td>5'-cttcagTACGCCCAATGGAAGGACTTGTG-3'</td>
</tr>
<tr>
<td>GSIII-1 F</td>
<td>5'-catATGCTCAAATCACTTAGTACCGGATCT-3'</td>
</tr>
<tr>
<td>GSIII-1 R</td>
<td>5'-gattttTATTCTGATAAACACACTCAAGG-3'</td>
</tr>
<tr>
<td>GSIII-2 F</td>
<td>5'-catATGGAGCATTAAAGGTTCTCAGTGTT-3'</td>
</tr>
<tr>
<td>GSIII-2 R</td>
<td>5'-ctcggtTACCGGAGGATCAGACCAGTCTCT-3'</td>
</tr>
<tr>
<td>qRT-PCR</td>
<td>PrGSI-F 5'-AGGGCCTTCAATTTGTTTAT-3'</td>
</tr>
<tr>
<td></td>
<td>PrGSI-R 5'-TTCGACGATAGCACATACCC-3'</td>
</tr>
<tr>
<td>PrGSI-1F</td>
<td>5'-CACATCCGACATTCGCCCTTTA-3'</td>
</tr>
<tr>
<td>PrGSI-1R</td>
<td>5'-CACAGCCGATATTGGAAATCCT-3'</td>
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<tr>
<td>PrGSI-2F</td>
<td>5'-CCGATGGGACAAGGATGATC-3'</td>
</tr>
<tr>
<td>PrGSI-2R</td>
<td>5'-GGCACTATGTTCTCCATCAG-3'</td>
</tr>
<tr>
<td>atpD-F</td>
<td>5'-TTGGATCTTATCGGCTGTTG-3'</td>
</tr>
<tr>
<td>atpD-R</td>
<td>5'-TGTCACCGTGCGCGCGAT-3'</td>
</tr>
<tr>
<td>inbB-F</td>
<td>5'-TCAGAATGGCAACACTAGTA-3'</td>
</tr>
<tr>
<td>inbB-R</td>
<td>5'-CACGATCCGACCTGTCGGT-3'</td>
</tr>
<tr>
<td>rpoB-F</td>
<td>5'-GAAGACTGTTCTGAGTGGACTG-3'</td>
</tr>
<tr>
<td>rpoB-R</td>
<td>5'-TAGCAGGCTGGCTGAAAGG-3'</td>
</tr>
</tbody>
</table>

culture collection. The bacterial cultures were grown anaerobiologically at 37°C in the modified medium 2 of Hobson (26) containing 30% rumen fluid and 0.2% glucose, 0.2% cellobiose, and 0.2% maltose under CO₂:H₂ (95.5: vol/vol) gas phase. This maintenance medium contains 6.8 mM (NH₄)₂SO₄.

Continuous culture. Continuous culture of P. ruminicola 23 was carried out with glucose (5 g/liter) and nitrogen-free defined medium (33) with the addition of 10 mM (NH₄)₂SO₄ to provide excess-ammonia growth conditions or 0.7 mM (NH₄)₂SO₄ to provide ammonia-limiting conditions. Residual ammonia concentrations in effluent were 3.66 mM (NH₄)₂SO₄.

PCR amplification. Primers were designed to include a 5' Ndel site in the forward primers and a 5' XhoI (GSI & GSIII-2) or BsaBI (GSIII-1) site in the reverse primers (listed in Table 1). These primers were used to amplify the GSI, GSIII-1, and GSIII-2 genes from genomic DNA isolated from P. ruminicola 23. PCR amplification was performed with the PFU DNA polymerase (Takara Bio Inc., Madison, WI) according to the manufacturer's instructions. Samples were amplified using the following program: hot start at 94°C for 2 min, preheating at 94°C for 30 s, 30 cycles for denaturation at 94°C for 10 s, annealing temperature of 55°C for 30 s, extension at 72°C for 45 s, and final extension at 72°C for 7 min for GSIII-1 and -2. The purity and integrity of the amplified products were determined by visual inspection following separation on a 1% agarose gel. The amplified GSI, GSIII-1, and GSIII-2 were gel purified and isolated using the gel extraction kit (Qiagen).

The purified genes were cloned into pGEM-T Easy Vector System I (Promega, WI). The clones with DNA inserts were identified by colony PCR using primers pUC/M13-F and pUC/M13-R (W. M. Keck Center for Comparative and Functional Genomics at the University of Illinois at Urbana-Champaign) and purified using a QIA spin miniprep kit (Qiagen). The GS genes in purified plasmid DNAs were excised from the pGEM-T vectors using NdeI and XhoI (or BsaBI for GSIII-1) restriction enzymes and ligated into a modified pET28a expression vector.

Protein purification. Epicurian E. coli BL21 CodonPlus (DE3) RIL competent cells (Strategene, La Jolla, CA) were transformed using 100 ng of plasmid DNA by heat shock at 42°C for 30 s. The cells were then spread on LB plates, with ampicillin (100 μg/ml) and chloramphenicol (50 μg/ml) for GSI and GSIII-2 and chloramphenicol (50 μg/ml) and kanamycin (30 μg/ml) for GSIII-1, and incubated overnight at 37°C. Since the competent cells carry the gene resistant to chloramphenicol, a single colony was picked and incubated in 500 μL medium supplemented with ampicillin and chloramphenicol, or chloramphenicol and kanamycin, using the same concentrations as mentioned before. The cells were cultured at 37°C on a rotary shaker until an OD₆₀₀ of 0.3 was reached, at which time isopropyl-β-D-thiogalactopyranoside (IPTG) was added at 1 mM to induce gene expression and protein production. The cells were then further incubated for 16 h at 16°C on a rotary shaker. The cells were harvested and then resuspended in isopropyl (50 mM Na phosphate, pH 7.0, and 300 mM NaCl), and lysed using a French pressure cell (American Instrument Co., Silver Spring, MD). The cell debris was removed by centrifugation. The histidine-tagged recombinant protein was purified using a cobalt-charged affinity resin and eluted into an elution buffer comprising 50 mM Na phosphate, pH 7.0, 300 mM NaCl, and 150 mM imidazole. The GS fractions were then dialyzed against buffer A (50 mM Tris, pH 8.0, 50 mM NaCl, 0.1 mM EDTA, 0.5 mM dithiothreitol [DTT], and 10% glycerol), as previously described (2).

Cytoplasmic proteins for analysis of GS activity from continuous culture were isolated as previously described (15). The cells were harvested by centrifugation, washed, and disrupted by sonication. Briefly, cells were resuspended in 10 ml of lysis buffer (20 mM Tris-HCl, 1 mM DTT, and 1 mM phenylmethylsulfonyl fluoride [PMSF]) and sonicated on ice in three cycles of 30 s at 80 W using a VC130PB (Sonic & Materials Inc., Newtown, CT). Unbroken cells and cell debris were removed by centrifugation at 12,000 × g for 5 min at 4°C. Supernatants were then ultracentrifuged in a fixed angle rotor at 105,000 × g at 4°C, giving a membrane-free cytoplasmic protein fraction.

Size exclusion chromatography. The purified GSI, GSIII-1, and GSIII-2 proteins were dialyzed against a buffer composed of 50 mM Na phosphate (pH 7.0) and 150 mM NaCl and injected into a Superose 12 HR 10/30 gel filtration column (Amersham Biosciences, Piscataway, NJ) already equilibrated with the same buffer at a flow rate of 0.4 ml/min. Fractions were collected with an automated fraction collector, and aliquots were analyzed by SDS-PAGE.

Enzymatic characterization of the GS proteins. Enzyme characterization was carried out by the γ-transferase (3) and biosynthetic assays (9, 20, 21) as previously described. The γ-transferase assay is used to measure the total amount of GS present. The assay mixture and glutamine solutions were prepared immediately prior to use to avoid the breakdown and release of ammonia in the growth media. The assay mixture contained 135 mM imidazole-HCl (pH 6.0), 18 mM hydroxyamine-HCl, 25 mM K-arsenate, 1 mM MnCl₂, 0.36 mM Na-ADP, and 10 μg GS, GSIII-1, or GSIII-2. The mixture was equilibrated at 37°C for 5 min, and the reaction was initiated by adding 50 μl of 20 mM γ-glutamine (final concentration of 20 mM), giving a final assay volume of 500 μl. The reaction was stopped

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following a 10-min incubation with 1 ml of stop mix (5.5% FeCl₃ - 6H₂O [wt/vol], 2% trichloroacetic acid [wt/vol], and 2.1% concentrated HCl) and then centrifuged for 5 min at 10,000 × g to remove precipitates. The product (formation of γ-glutamyl-hydroxamate) was measured spectrophotometrically by absorbance at 540 nm where 1 μmol of γ-glutamyl-hydroxamate had an absorbance of 0.278. A reaction mixture without GS enzyme served as the blank. The reaction is shown as follows.

**Hydroxylamine + γ-glutamyl-hydroxamate → γ-glutamyl-hydroxamate + ammonia + ADP**

The assay mixture for the biosynthetic reaction (measures the ability of the GS to form glutamine through the P, released from ATP) consisted of 100 mM MOPS (morpholinopropanesulfonic acid) (pH 7.5), 50 mM MgCl₂ - 6H₂O, 250 mM γ-glutamate, and 50 mM NH₄Cl to which approximately 10 μg of GSI, GSIII-1, or GSIII-2 was added. The 90 μl of mixture was equilibrated at 37°C for 5 min, and the reaction was initiated by adding 10 μl of 0.1 M ATP (final concentration of 10 mM) in a total volume of 100 μl. The 25 μl of reaction was transferred after 5 min to a microtiter plate, and 75 μl of solution D (2:1 mixture of 12% l-ascorbic acids in 1 N HCl and 2% (NH₄)₂MoO₄ - 4H₂O) was added. The reaction was stopped after 5 min of incubation by the addition of 75 μl of stop color development solution F (2% sodium citrate tribasic dihydrate, 2% acetic acid, and 2% sodium arsinite). The solution was then incubated for 15 min at 37°C to enable the color to fully develop. The inorganic phosphate product was measured spectrophotometrically at 850 nm as for the biosynthetic assay. GS specific activity is expressed as nmol Pi/μg/min. The reaction is shown as follows.

**l-Glutamate + NH₄⁺ + ATP → l-glutamine + ADP + P_i + H⁺**

### Kinetic characterization

The concentration of the substrate for which the affinity was to be calculated was changed while maintaining the concentration of the other substrates in excess. **Kₘ** was determined by γ-transferase, varying the concentration of glutamine from 1.0 to 40 mM, that of ADP from 0.1 to 2.0 mM, and that of hydroxylamine-HCl from 0.5 to 20 mM. **Kₘ** was also analyzed for all the three substrates, glutamate, ATP, and ammonia by biosynthetic assay. Glutamate from 0.5 to 300 mM, ATP from 0.1 to 20 mM, and ammonia from 0.5 to 25 mM were used. A double reciprocal Lineweaver-Burk plot was applied to calculate the **Kₘ** values for the reactions.

### ATPase activity

ATP hydrolysis was assayed using 10 μM nonradioactive ATP and 170 nM γ-[^32]P]ATP. The mixture was equilibrated at 37°C for 5 min, and then the reaction was initiated by the addition of 0.5, 1, and 2.0 μg GSI or 25, 50, and 100 ng GSIII-1 or GSIII-2 protein, respectively. The reaction mixture was incubated at 37°C for 10 min, and then the reaction was terminated by the addition of 2 μl of 0.5 M EDTA (pH 7.5). An aliquot (1 μl) of the reaction mixture was spotted on a polyethyleneimine-cellulose thin-layer plate (Merck, Darmstadt, Germany) and subjected to thin-layer chromatography with 1 M LiCl and 0.5 M formic acid buffer. The plate was exposed to a phosphorimaging plate (DEPC)-treated water. The DNA was purified with a Qiagen RNeasy cleanup kit (Qiagen) by following the manufacturer’s protocol. The RNA was then diluted 1:4 with DNase- and RNase-free water. The qRT-PCR was performed using SYBR green I (Applied Biosystems, Foster City, CA) with an ABI Prism 7900 high-throughput sequence detection system. cDNA (4 μl) was mixed with 5 μl SYBR green master mix (Applied Biosystems, Foster City, CA), 0.4 μl of each 10 μM forward and reverse primer (Table 1), and 0.2 μl of DNase- and RNase-free water. Each sample was run in triplicate along a six-point relative standard curve of internal control or constitutive genes (atpD, ORFB01230; infB, ORFB02450; and rpoB, ORFB02217) determined from cDNA microarray plus nontemplate control (NTC). The qRT-PCRs were performed with the following conditions: 50°C for 2 min, 95°C for 10 min, and 40 cycles of 95°C for 15 s and 60°C for 1 min. In addition, to verify the presence of a single PCR product, a dissociation protocol using incremental temperatures to 95°C for 15 s plus 65°C for 15 s was performed. Data were analyzed using the SDS software version 2.2.1. (Applied Biosystems) using the six-point standard curve. A mixed model with repeated measures (release 9.0; SAS Institute, Cary, NC) using spatial power as the covariate structure was used for all analyses (normalized gene expression). The model included fixed effects of time and treatment. A replicate was considered a random effect. Statistical significance was declared at P ≤ 0.05.

### Phylogenetic analysis

All of the protein sequences were retrieved from GenBank, and GS sequences of *P. ruminicola* 23 were obtained from *P. ruminicola* 23 genome databases (accession number CP002006; http://jctv.jrc.runenomics/). Protein sequences were aligned using ClustalX version 1.83), and the program generated an unrooted neighbor-joining tree (43). The phylogenetic tree was manipulated by using TreeView (version 1.4) (http://taxonomy.zoology.gla.ac.uk/rod/rod.html).

## RESULTS

### Amino acid sequence alignment

Three GS-encoding genes (GSI, 501 amino acids [aa]; GSIII-1, 730 aa; and GSIII-2, 737 aa) were identified bioinformatically in the *P. ruminicola* 23 genome and were aligned with other characterized GSI and GSIII proteins. The alignments among GS proteins showed that GSI is divergent from GSIII proteins while GSIII-1 and GSIII-2 showed high similarity (GSI versus GSIII-1, 6%; GSI versus GSIII-2, 5%; and GSIII-1 versus GSIII-2, 79%). The alignments and phylogenetic tree were consistent with 16S RNA organismal phylogeny, with the *P. ruminicola* 23 GSI being most similar to a GSI type I from *B. fragilis* YCH46 (BF2249, accession no. YP_099530; 77% identity) (Fig. 1 and 2). Likewise, GSIII-1 and GSIII-2 of *P. ruminicola* 23 aligned best with the GSIII from *P. bryantii* B-4 (accession no. AAL87245;
81% and 77% identical, respectively) but less well with those found in *Ruminococcus albus* 8, *Ruminococcus flavefaciens* FD-1, *Synechococcus* sp., and *Psychromonas ingrahamii* 37 (Fig. 2).

Alignment of GSI of *P. ruminicola* 23 allowed for the identification of all five conserved regions common to GS families (1, 44) (motifs I, II, III, IV, and V; see Fig. S1 in the supplemental material). Alignment of GSI with characterized orthologs from *Mycobacterium tuberculosis*, *E. coli*, *Salmonella enterica* serovar Typhimurium, and *Synechocystis* sp. PCC 6803 revealed 10 insertions in motif II and substitutions in the ATP binding site (motif III). In contrast, alignment of GSIII-1 and GSIII-2 with GSIII family enzymes of *R. albus* 8 and *Synechococcus* sp. PCC 7942 revealed strong sequence conservation through all motifs (Fig. 1).

Expression and purification of recombinant proteins. GSI, GSIII-1, and GSIII-2 from *P. ruminicola* 23 were heterologously expressed and purified in *E. coli* (see Fig. S2 in the supplemental material). *P. ruminicola* 23 GS proteins were purified using anion exchange followed by a cobalt affinity column. SDS-PAGE showed expression of protein with a molecular size consistent with that expected for GS proteins of *P. ruminicola* 23 (GSI, 56.0 kDa; GSIII-1 and -2, 83.0 kDa).

Subunit organization of recombinant GS of *P. ruminicola* 23. The subunit organization of the purified GSIII-1 and -2 proteins was estimated using gel filtration and SDS-PAGE analysis. Two peaks were observed for GSIII-1, and one peak was observed for GSIII-2 (Fig. 3). The estimated sizes of the peaks observed for GSIII-1 were 1,339.0 kDa and 133.0 kDa, while that observed for GSIII-2 was estimated to be 230.0 kDa.

Characteristics of GS activities. Optimal GS protein conditions for *P. ruminicola* 23 are shown in Table S1 in the supplemental material. GSI displayed optimal -transferase activity at 35°C, while GSIII-1 and -2 -transferase activities were both optimal at 37°C. In all cases, activity was drastically decreased above 50°C. The optimum pH of GSI was determined to be pH 5.6, with little activity observed below pH 5.2 and above pH 5.6. Conversely, GSIII-1 and -2 were most active between pH 5.6 and 7.2, both having a pH optima of 6.0. -Transferase activity of all three enzymes (GSII-1, GSII-1, and GSII-2) was observed only in the presence of Mn²⁺ ions and was not detected in the presence of Mg²⁺, Cu²⁺, Co²⁺, or Ca²⁺. The optimal
concentrations of Mn\(^{2+}\) ions for GSI, GSIII-1, and GSIII-2 γ-transferase activities were 0.25, 0.5, and 1 mM, respectively, with concentrations below 0.25 mM failing to elicit activity. No biosynthetic activity for GSI was detected with any cation, and optimization of GSI biosynthetic activity for cation concentration, temperature, and pH could not be determined. The biosynthetic activity of GSIII-2 was dependent on the presence of Mn\(^{2+}\), Fe\(^{2+}\), Co\(^{2+}\), or Mg\(^{2+}\), with maximum activity observed in the presence of Mn\(^{2+}\). Biosynthetic activity of GSIII-1 was stimulated by Mn\(^{2+}\) and to lesser extents by Fe\(^{2+}\), Co\(^{2+}\), Mg\(^{2+}\), and Ca\(^{2+}\) (see Table S1 in the supplemental material). The optimal concentrations of Mn\(^{2+}\) for the biosynthetic activities of GSIII-1 and -2 were both 10 mM, with no biosynthetic activity observed below 5 mM (see Table S1). The optimum temperatures for biosynthetic activity of GSIII-1 and -2 were both 50°C, although both enzymes appeared stable up to 70°C. The biosynthetic activity of GSIII-1 had a pH optimum of pH 7.0, while that of GSIII-2 was shown to be pH 6.8. Biosynthetic activity of both GSIII enzymes was stable between pH 5.2 and 8.0 (see Table S1).

Kinetic properties. For the γ-transferase assay of GSI, the calculated apparent \(K_m\) for ADP, hydroxylamine-HCl, and L-glutamine is reported in Table 2. However, apparent \(K_m\) values for the substrates in the biosynthetic assay were not detectable. For the γ-transferase assay of GSIII-1 and GSIII-2, the calculated apparent \(K_m\) values were 0.06 and 0.62 mM for ADP, 2.04 and 0.07 mM for hydroxylamine-HCl, and 1.30 and 1.92 mM for L-glutamine, respectively. The apparent \(K_m\) values of GSIII-1 and GSIII-2 for the substrates in the biosynthetic assay were 8.58 and 1.72 mM for glutamate, 1.91 and 2.65 mM for ATP, and 0.48 and 0.43 mM for ammonia, respectively (Table 2).

ATPase activity. We tested GSI, GSIII-1, and GSIII-2 from \(P.\) ruminicola for their abilities to hydrolyze ATP (Fig. 3; see Fig. S3 in the supplemental material). GSIII-1 and -2 gave the highest ATP hydrolysis activities. Increasing GSI did not increase ATP hydrolysis. However, increasing the GSIII-1 and GSIII-2 concentration increased ATP hydrolysis in a concentration-dependent manner (Fig. 3). GSI had ca. 100-fold lower ATPase activity (7 pmol ATP hydrolyzed/mg/min) (see Fig. S3) compared to ATPase activities of GSIII-1 and -2 (70 to 190 pmol ATP hydrolyzed/mg/min) (Fig. 3).

Transcriptional regulation and GS enzyme activities in continuous culture. Transcription levels of the three different GS-encoding genes were analyzed by qRT-PCR from a continuous culture of \(P.\) ruminicola 23 before and after a shift in ammonia concentration from excess (10 mM) to limiting (0.7 mM). On high concentrations of ammonia, GSIII-2 (ORFB02034) increased 71.3-fold (Table 3). On the other hand, GSI and GSIII-1 were not significantly increased on high concentrations of ammonia (GSI, 1.9-fold; GSIII-1, 1.7-fold).

The biosynthetic activity of GS was around 2-fold higher during growth of \(P.\) ruminicola 23 on a nonlimiting concentration of ammonia than on a limiting ammonia concentration (19.4 ± 2.0 nmol Pi/g/min on nonlimiting versus 8.8 ± 2.6 nmol Pi/g/min on limiting ammonia). No GSI biosynthetic activity was detected during this study; thus, activity likely represents GS type III enzyme activity. Interestingly, this study revealed upregulation when \(P.\) ruminicola 23 was grown on nonlimiting ammonia. Specifically, GSIII-2 showed 71.3-fold upregulation. These results suggest that

FIG 2 Unrooted phylogenetic tree of GS type I, II, and III proteins based on ClustalX. Alignments were constructed using ClustalX, and the phylogenetic tree was built using a neighbor-joining plot (ClustalX version 1.82) (43).
GSIII-2 plays an important role in ammonia assimilation under nonlimiting ammonia growth conditions.

**DISCUSSION**

The well-studied nitrogen metabolic circuit in enteric bacteria consists of three enzymes: glutamine synthetase (GS), glutamate synthase (GOGAT), and glutamate dehydrogenase (GDH). Glutamine synthetase, encoded by \( glnA \), catalyzes the only pathway for the synthesis of glutamine. Glutamate can be synthesized by two pathways: through the combined activity of GS and glutamate synthase, encoded by \( gltBD \), that constitute the GS-GOGAT pathway and through the activity of glutamate dehydrogenase, encoded by \( gdhA \) (34). In enteric bacteria, the GS-GOGAT enzyme system has a high affinity for \( \text{NH}_4^+ \) (\( K_m \), 0.2 mM for GS) whereas GDH has a low affinity (\( K_m \) > 1 mM) (32, 39). The two central intermediates in nitrogen metabolism, glutamine and glu-

**FIG 3** Size exclusion chromatography (A and B) and ATPase assay (C and D) of the \( P.\ ruminicola \) 23 GSIII-1 and -2. (A and B) The chromatograph represents 100 \( \mu l \) (7.5 mg/ml) of purified recombinant \( P.\ ruminicola \) 23 GSIII-1 (A) and -2 (B) dialyzed against 50 mM imidazole-HCl (pH 6.5)–50 mM NaCl–50 mM MgCl\(_2\), injected into a Superdex 200 HR 10/30 gel filtration column (Amersham Biosciences, NJ), and eluted at a rate of 0.5 ml/min. Elution volumes of the standards are represented by arrows 1 to 5: 1, thyroglobulin (670 kDa); 2, \( \gamma \)-globulin (158 kDa); 3, ovalbumin (44 kDa); 4, myoglobin (17 kDa); 5, vitamin B\(_12\) (1.3 kDa). SDS-PAGE analyses of 15-\( \mu l \) aliquots of fractions collected during size exclusion chromatography analysis are presented below the chromatography panel. All bands correspond in size to the \( P.\ ruminicola \) 23 GSIII-1 and GSIII-2 monomeric subunit (83 kDa). (C and D) Histogram showing ATP hydrolysis expressed as pmol ATP hydrolyzed/mg/min determined using the PEI plate assay. Lanes 1, 1 \( \mu g \) positive control (MacHjm of \( \text{Methanosarcina acetivorans} \)); lanes 2, negative control (no protein); lanes 3, 25 ng GS III-1 and -2; lanes 4, 50 ng GS III-1 and -2; lanes 5, 100 ng GS III-1 and -2.

**TABLE 2** Apparent \( K_m \) for different substrates of \( P.\ ruminicola \) 23 GSI and GSIII

<table>
<thead>
<tr>
<th>GS Type</th>
<th>Transferase Assay</th>
<th>Biosynthetic Assay</th>
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<tr>
<td></td>
<td>Glutamine</td>
<td>ADP</td>
</tr>
<tr>
<td>GSI</td>
<td>1.93</td>
<td>0.45</td>
</tr>
<tr>
<td>GSIII-1</td>
<td>1.30</td>
<td>0.06</td>
</tr>
<tr>
<td>GSIII-2</td>
<td>0.62</td>
<td>0.07</td>
</tr>
</tbody>
</table>

\(^a\) NM, no activity for these substrates was obtained using the biosynthetic assay.
motifs II (an active vertical path of descent for these enzymes (Fig. 2). However, and type III proteins in the available databases and support a with those of biochemically validated and putative GS type I 23. The predicted GS amino acid sequences were compared encoding glutamine synthetase in the genome of  coli and Salmonella.

Closed genome sequence for  ruminicola 23 enabled a functional genomics approach to analyze enzymatic mechanisms involved in ammonia assimilation. As a first step, we carried out a bioinformatic analysis and identified three genes encoding glutamine synthetase in the genome of  ruminicola 23. The predicted GS amino acid sequences were compared with those of biochemically validated and putative GS type I and type III proteins in the available databases and support a vertical path of descent for these enzymes (Fig. 2). However, amino acid sequence alignment of GS revealed insertions in motifs II (an active β-barrel site) and III (the ATP-binding site) compared to characterized GS proteins (see Fig. S2 in the supplemental material). In addition, GSIII-1 and GSIII-2 displayed strong conservation of all five GS conserved regions and four GS type III-specific conserved regions (11, 44). Furthermore, key conserved amino acid residues essential for function of the GSIII protein in signature motifs I to IV and motif C in GSIII amino acid sequence (2). These alignments, supported by GS biosynthetic activity of the two GSIII proteins but not GSII, suggest important roles of  ruminicola 23 GSIII proteins in nitrogen metabolism and that GSII is no longer essential to ammonia assimilation and possibly no longer functions as a GS enzyme. Interestingly, despite GSII having no biosynthetic activity, it still has low ATPase activity as well as glutamyl transferase activity and may function as an amino acid transferase, although this has not been tested. It is possible that this indicates a recent loss of GS function for GSII potentially mediated by reduced selective pressure by the duplication of GSII compared to  p. bryantii, a close ruminal and phylogenetic relative, which has a single GSIII and GSII present in the genome. Furthermore, glnA (ORFB02034) encoding GSIII-2 is adjacent to gltD and gltB encoding GOGAT (Fig. 4; see Table S2 in the supplemental material). This gene cluster containing GS adjacent to GOGAT supports the possible function of GSIII-2 in the ammonia assimilation pathway as part of a functional GS-GOGAT-linked enzyme system. The 7-amino acid insertion (or deletion) observed in motif II of the  ruminicola GS is also observed in two homologs found in  B. fragilis and  Bacteroides thetaiotaomicron. This insertion, although potentially contributing to the loss of biosynthetic activity observed in the  ruminicola GS, is unlikely to be the entire contributing factor, since the  albus 8 homolog, which lacks this feature, is also inactive for biosynthetic activity (2). Amino acid residues that may be of significance and are conserved in the functional homologs (  M. tuberculosis,  E. coli K-12, S. Typhimurium, and the  Synechocystis sp. homologs) but have been replaced with nonconservative residues in the two nonfunctional homologs (  albus 8 and  ruminicola 23 homologs) are the aspartate/asparagine in motif III. We predict that the  Bacteroides sp. homologs are also nonfunctional. Mutational analysis will, however, be required to unravel the contributions of the different mutations, insertions, or deletions to the lack of biosynthetic activity observed in some GSII homologs.

Enzymatic characterization of the three GS enzymes of  ruminicola 23 was carried out to define the optimal conditions for the enzymatic activity. The GSII proteins showed a low Km value for ammonia (0.48 mM for GSIII-1 and 0.43 mM for GSIII-2) similar to that of the GS of  e. coli (Km 0.6 mM) (30). These kinetic values suggest that both GSIII enzymes are functional and play a glutamine biosynthetic role in vivo based on their affinity for glutamate and ammonia. Ammonia levels in the rumen vary considerably depending on diet and time post-feeding but are in the general range of 2 to 38 mM (4, 23); however, on poor-quality, low-protein forages they fall below 1 mM. Further support for their role in ammonia assimilation was shown by analysis of transcript level using qRT-PCR and enzyme activities of GSII and two GSIII proteins during chemostat culture of  ruminicola 23 with nonlimiting (residual NH3 3.66 mM) and limiting (residual NH3 0.025 mM) concentrations of ammonia (Table 3). Our results for glnA expression and GS activity in  ruminicola 23 by ammonia concentration are divergent from that described for GSIII proteins in  B. fragilis Bf-1,  Butyrivibrio fibrisolvens,  r.  albus 8,  Pseudanabaena sp. PCC 6903,  Synechocystis sp. PCC 6803, and  Synechococcus sp. PCC 7942, which all showed higher enzymatic activities and high expression levels for cells grown under ammonia limitation (2, 10, 11, 22, 35, 41). The reason for this discrepancy is not obvious, but previously normalization was based on 16S rRNA gene level whereas this comparison is related to constitutively expressed genes. Importantly, cells used for measuring transcript levels and enzyme activity in the present study were prepared in chemostat- and not batch-grown cells and thus steady-state residual concentrations of ammonia could be established for each condition before sampling.

In conclusion, the present study is the first to biochemically characterize multiple GS proteins encoded within a single genome and to establish their kinetic properties. The two GSII genes are paralogs (79% amino acid similarity) resulting from gene duplication that presumably provides functional divergence and the ability to adapt to various environmental conditions and to expand the growth phenotype that  Prevotella ruminicola expresses. Based on its enzyme activity and expression levels, we show that GSIII-2 is responsive to external ammonia concentrations and is likely involved in ammonia assimilation with higher expression levels on nonlimiting than on limiting table 3 relative expression levels of glutamine synthetase enzymes in  ruminicola 23 on high ammonia as determined by qrt-PCR analyses

<table>
<thead>
<tr>
<th>ORF</th>
<th>Gene</th>
<th>Common name</th>
<th>EC no.</th>
<th>Fold change in qRT-PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>ORFB02151</td>
<td>glnA</td>
<td>Glutamine synthetase, type I (GSII)</td>
<td>6.3.1.2</td>
<td>1.9</td>
</tr>
<tr>
<td>ORFB01459</td>
<td>glnA</td>
<td>Glutamine synthetase, type III (GSIII-1)</td>
<td>6.3.1.2</td>
<td>1.7</td>
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<tr>
<td>ORFB02034</td>
<td>glnA</td>
<td>Glutamine synthetase, type III (GSIII-2)</td>
<td>6.3.1.2</td>
<td>71.3</td>
</tr>
</tbody>
</table>

* Fold change in qRT-PCR during growth on high (nonlimiting) ammonia versus low (growth-limiting) ammonia concentrations.
growth concentrations of ammonia. The genomic context of GSIII-2, together with the large and small subunits (gltB and gltD) of GOGAT, also provides additional support for this hypothesis. We propose that GSIII-1 plays a related, rather than entirely new, function and is involved in recycling of ammonia, maintenance of the intracellular glutamate pool, and supply of amine groups for biosynthesis of other N-containing cell components such as amino acids, purines, pyrimidines, and polyamines. Neither GSIII-1 nor GSI have other genes involved in amino acid biosynthesis in contiguous genomic sequence 15 kb upstream or downstream of their respective ORFs. The role for GSI is unknown, since the recombinant enzyme lacked biosynthetic activity although it exhibited both glutamyl transferase and ATPase activity and was functional. We suggest that in vivo this enzyme no longer functions in the synthesis of glutamine from glutamate and ammonia but may play a role in amino-transferase activity and still participate in intracellular nitrogen metabolism, although this remains to be determined.

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