

# *Geobacter sulfurreducens* Contains Separate C- and A-Adding tRNA Nucleotidyltransferases and a Poly(A) Polymerase<sup>∇†</sup>

Patricia Bralley, Madeline Cozad, and George H. Jones\*

Department of Biology, Emory University, Atlanta, Georgia 30319

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**The genome of *Geobacter sulfurreducens* contains three genes whose sequences are quite similar to sequences encoding known members of an RNA nucleotidyltransferase superfamily that includes tRNA nucleotidyltransferases and poly(A) polymerases. Reverse transcription-PCR using *G. sulfurreducens* total RNA demonstrated that the genes encoding these three proteins are transcribed. These genes, encoding proteins designated NTSFI, NTSFII, and NTSFIII, were cloned and overexpressed in *Escherichia coli*. The corresponding enzymes were purified and assayed biochemically, resulting in identification of NTSFI as a poly(A) polymerase, NTSFII as a C-adding tRNA nucleotidyltransferase, and NTSFIII as an A-adding tRNA nucleotidyltransferase. Analysis of *G. sulfurreducens* rRNAs and mRNAs revealed the presence of heteropolymeric RNA 3' tails. This is the first characterization of a bacterial system that expresses separate C- and A-adding tRNA nucleotidyltransferases and a poly(A) polymerase.**

RNA nucleotidyltransferases play an important role in the maturation, processing, and degradation of RNAs. tRNA nucleotidyltransferases (TNTs) and poly(A) polymerases (PAPs) are both members of a superfamily of nucleotidyltransferases (13, 27). TNTs add C and A residues to the 3' ends of immature or damaged tRNA molecules and are found in bacteria, archaea, and eukaryotes (24). PAPs, which are present in both eukaryotes and bacteria, polyadenylate RNA 3' ends (27). With regard to the distribution of RNA nucleotidyltransferases in bacteria, with only a few exceptions, the genomes of the alpha- and epsilonproteobacteria and of the low- and high-G+C-content gram-positive bacteria appear to encode only one member of this superfamily, presumably a CCA-adding TNT (1). Several of the so-called deeply branching bacterial species, including *Aquifex aeolicus* and *Deinococcus radiodurans*, and some cyanobacteria contain two RNA nucleotidyltransferases, which have been shown to function as separate C- and A-adding TNTs (22, 23). Interestingly, at least one low-G+C-content gram-positive species, *Bacillus halodurans*, has also been shown to contain separate C- and A-adding TNTs (1). The beta- and gamma-proteobacteria, such as *Escherichia coli*, also contain two RNA nucleotidyltransferases, a CCA-adding TNT and a PAP, PAP I, that adds 3' tails to the ends of RNA molecules in cells of these organisms (8, 11, 18, 19). mRNAs and 16S and 23S rRNAs are polyadenylated in *E. coli* and presumably in other beta- and gamma-proteobacteria, and polyadenylation facilitates the exonucleolytic degradation of these RNAs by RNase II and polynucleotide phosphorylase (9, 16, 18, 19).

The genomes of several deltaproteobacteria also encode two members of the RNA nucleotidyltransferase superfamily

(NTSF), and the genomes of a few deltaproteobacteria appear to encode three NTSF enzymes (1). In particular, the genomes of *Geobacter sulfurreducens*, *Geobacter metallireducens*, *Pelobacter carbinolicus*, and *Desulfotalea psychrophila* all contain three open reading frames with significant homology to open reading frames encoding other members of the superfamily of nucleotidyltransferases. We describe here biochemical characterization of the three NTSF enzymes of the dissimilatory metal- and sulfur-reducing bacterium *G. sulfurreducens* (6). This report is the first characterization of NTSF enzymes from a bacterial species with three RNA nucleotidyltransferases.

## MATERIALS AND METHODS

**Bacterial strains and genomic DNA.** *G. sulfurreducens* ATCC 51573 genomic DNA was obtained from the American Type Culture Collection. Frozen late-log-phase cells of *G. sulfurreducens* PCA were generously provided by Lori Hayes (laboratory of Derek Lovley) of the University of Massachusetts.

**Sequences, alignment, and phylogenetic analysis.** Sequences were retrieved from the microbial genomes database at www.ncbi.nlm.nih.gov. Sequences were aligned using ClustalX, and neighbor-joining phylogenetic trees were constructed using PHYLIP 3.6b (12) as described previously (1) and in the supplemental material.

**RT-PCR analysis.** Reverse transcription (RT)-PCR was carried out essentially as described by Bralley et al. (1). For studies of expression of the enzymes designated NTSFI, NTSFII, and NTSFIII, 5 µg of total RNA was reverse transcribed with random hexamer primers. PCR was then performed using primers specific for each of the NTSF enzymes using the Expand high-fidelity PCR system (Roche). PCR primers were designed so that they were complementary to nonhomologous regions of the three proteins to avoid detecting the transcript of one gene with more than one set of primers. The gene-specific primers were GeoBac I F1, GeoBac I R1, GeoBac II F1, GeoBac II R1, GeoBac III F1, and GeoBac III R1 (see Table S1 in the supplemental material).

**Cloning and overexpression of the NTSF genes.** PCR primers corresponding to the 5' and 3' ends of the three genes were designed and used in the PCR as described previously (1). Primers were designed to incorporate BamHI sites at the 5' and 3' ends of each of the NTSF genes, and the PCR products were cloned into pET11A for overexpression in *E. coli* BL21(DE3)/pLysS Rosetta cells. Overexpressed proteins were purified as described in the supplemental material.

**Preparation of tRNA<sup>Asp</sup>-C and tRNA<sup>Asp</sup>-CC.** Plasmid pmBsDCCA encodes *E. coli* tRNA<sup>Asp</sup> (15). Plasmid DNA (27 µg) was digested with either BbsI or FokI. The digested DNAs were isolated by phenol extraction and ethanol precipitation and used as templates for transcription with T7 RNA polymerase. The reaction

\* Corresponding author. Mailing address: Department of Biology, Emory University, Atlanta, GA 30319. Phone: (404) 727-0712. Fax: (404) 727-2880. E-mail: george.h.jones@emory.edu.

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mixtures (300  $\mu$ l) contained 5 $\times$  reaction buffer, 9  $\mu$ g DNA template, 10 mM dithiothreitol, 300 U 5 PRIME Stop RNase inhibitor (Fisher), 2 mM ATP, 2 mM CTP, 2 mM GTP, 2 mM UTP, and 270 U T7 RNA polymerase (Promega). The reaction mixtures were incubated for 4 h at 37°C, brought to 1.5 mM with CaCl<sub>2</sub>, and RNase-free DNase (15 U; Promega) and additional RNase inhibitor (45 U) were added. Then the mixtures were incubated for 30 min at 37°C, DNase and RNase inhibitor were added again as described above, and the mixtures were incubated for 30 min. RNAs were collected by phenol extraction and two ethanol precipitations. Transcription of the BbsI-digested template produced tRNA<sup>Asp</sup>-CC, while transcription of the FokI-digested template produced tRNA<sup>Asp</sup>-C.

**Enzyme assays.** Generally, PAP and TNT assays were performed using 60- $\mu$ l reaction mixtures as described previously (1, 2, 5). The PAP assay reaction mixtures contained [ $\alpha$ -<sup>32</sup>P]ATP, while the TNT assay mixtures contained [ $\alpha$ -<sup>32</sup>P]ATP or [ $\alpha$ -<sup>32</sup>P]CTP (3,000 Ci/mmol; GE Health Sciences). To examine the products of the TNT and PAP reactions, large-scale mixtures (60 to 240  $\mu$ l) were incubated as described above and extracted with phenol-chloroform. Labeled RNAs were precipitated with ethanol. RNAs were examined by electrophoresis on 5 or 10% denaturing polyacrylamide gels along with Ambion Century-Plus size markers which had been end labeled with [<sup>32</sup>P]pCp and T4 RNA ligase. The amounts of enzyme (per 60- $\mu$ l reaction mixture) used to generate the data shown in Fig. 3 were as follows: PAP I (New England Biolabs), 5 U; NTSFI, 5.9  $\mu$ g; NTSFII, 7.5  $\mu$ g; NTSFIII, 7.1  $\mu$ g; and SCO3896, 8.6  $\mu$ g.

**Determination of poly(A) tail lengths, cDNA cloning, and sequencing.** Poly(A) tails were isolated and characterized as described previously by end labeling total RNA with [<sup>32</sup>P]pCp and then digesting the RNA with RNases A and T<sub>1</sub> (1, 4, 5). Products of this digestion were displayed on a 12% denaturing polyacrylamide gel. RT of *G. sulfurreducens* total RNA was carried out using primer AD20 containing an oligo(dT<sub>17</sub>) sequence that could anneal to poly(A) tails. For cDNA amplification by PCR, gene-specific forward primers and a reverse primer complementary to the primer used for RT were used. The sequences of the AD20 primers have been described previously (3). The gene-specific primers used in these experiments were 23SF, 16S1, NTSFI F3, and NTSFII F2 (see Table S1 in the supplemental material).

In an alternative method for cDNA preparation, a DNA adaptor, GSUENDR2 (see Table S1 in the supplemental material) was attached to the 3' ends of *G. sulfurreducens* total RNA using RNA ligase. The reaction mixtures (20  $\mu$ l) contained 25  $\mu$ g of RNA, 0.4 nmol of adaptor, 10% dimethyl sulfoxide, 30 U of 5 PRIME Stop RNase inhibitor, 10 U of T4 RNA ligase (Ambion), and 5 $\times$  RNA ligase buffer (Ambion). The reaction mixtures were incubated overnight at 4°C, and the products were purified by phenol extraction and ethanol precipitation. The RNA products were then reverse transcribed, and cDNA amplification was performed as described previously using GSURT (see Table S1 in the supplemental material) and the gene-specific primers indicated above (3). Amplification products were cloned using a TOPO-TA cloning kit (Invitrogen) and then sequenced.

## RESULTS AND DISCUSSION

**Phylogenetic analysis of the *Geobacter* NTSF protein sequences.** We initially subjected the three *Geobacter* NTSF protein sequences, arbitrarily designated NTSFI (annotated GSU3250 in the *G. sulfurreducens* genome database), NTSFII (GSU2184), and NTSFIII (GSU1581), to phylogenetic analysis and compared them with RNA nucleotidyltransferases from other bacteria to which functions have been assigned. Figure 1 shows an unrooted neighbor-joining phylogenetic tree constructed using PHYLIP (12). Representative NTSF enzymes from species with separate C- and A-adding TNTs belonging to the beta- and gammaproteobacteria and from other deltaproteobacteria with three NTSF genes were compared in this analysis. Protein sequences were aligned using ClustalX, and the alignments were refined using TuneClustal as described previously (1). The results of this analysis show that *G. sulfurreducens* NTSFI, along with the corresponding NTSF enzymes from the other deltaproteobacteria that were examined, occurs in a clade that contains the PAPs from the beta- and gammaproteobacteria. *G. sulfurreducens* NTSFII and the cor-

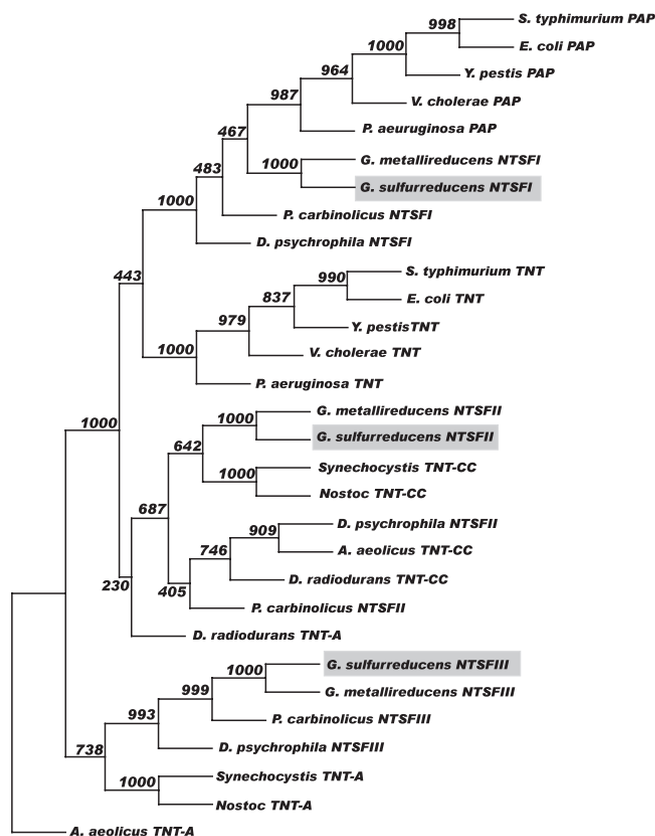


FIG. 1. Neighbor-joining phylogenetic tree for the bacterial NTSF enzymes. The tree was constructed using PHYLIP. Bootstrap scores are shown at the nodes. The branch lengths do not represent evolutionary distances. The species represented are *E. coli*, *Salmonella enterica* serovar Typhimurium, *Vibrio cholerae*, *Yersinia pestis*, *Pseudomonas aeruginosa*, *D. radiodurans*, *A. aeolicus*, *Nostoc* spp., *Synechocystis* spp., *G. sulfurreducens*, *G. metallireducens*, *P. carbinolicus*, and *D. psychrophila*.

responding proteins from the other deltaproteobacteria occur in the clade that contains the C-adding enzymes from the species with separate C- and A-adding TNTs, while NTSFIII occurs in the clade containing the A-adding TNTs (Fig. 1).

***G. sulfurreducens* NTSF genes are transcribed.** We next determined whether the three NTSF genes were transcribed in *G. sulfurreducens*. RNA was isolated from frozen cells, and RT-PCR was performed using random primers for RT and primers specific for each NTSF enzyme for PCR. The primers utilized for PCR were expected to yield 1,160-, 1,183-, and 1,250-bp amplification products for NTSFI, NTSFII, and NTSFIII, respectively. An agarose gel of the PCR products (Fig. 2A) revealed bands of the expected size in each case and also showed that the PCR templates were generated by RT rather than by contaminating DNA in the RNA preparations. Thus, the genes encoding NTSFI, NTSFII, and NTSFIII are transcribed in *G. sulfurreducens*.

**Cloning and overexpression of the *G. sulfurreducens* NTSF enzymes.** PCR products corresponding to the coding sequences of NTSFI, NTSFII, and NTSFIII were prepared using primers that incorporated BamHI sites at the ends of each gene (NTSFI, NTSFII, and NTSFIII forward and reverse primers [see Table S1 in the supplemental material]) and *G.*

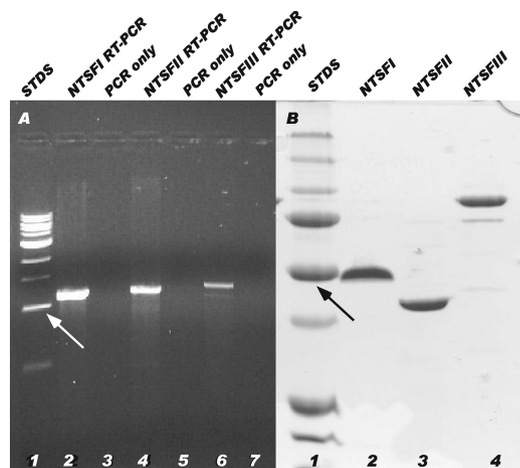


FIG. 2. (A) RT-PCR analysis showing the expression of the genes encoding *G. sulfurreducens* NTSF enzymes. Lane 1, size standards (the arrow indicates the position of the 1-kb standard); lanes 2, 4, and 6, RT-PCR products obtained with primers designed to identify NTSFI, NTSFII, and NTSFIII, respectively; lanes 3, 5, and 7, PCR products obtained with the same primers but without RT. (B) Sodium dodecyl sulfate-polyacrylamide gel electrophoresis gel of purified NTSFI (1  $\mu$ g) (lane 2), NTSFII (2.5  $\mu$ g) (lane 3), and NTSFIII (2.5  $\mu$ g) (lane 4). Lane 1 contained size standards having molecular weights ranging from 20,000 to 106,000. The arrow indicates the position of the 50,000- $M_r$  standard.

*sulfurreducens* chromosomal DNA as the template. The products were then cloned into the BamHI site of the overexpression vector pET11A, and the proteins were overexpressed in *E. coli* BL21(DE3)/pLysS Rosetta cells. We note that the overexpressed proteins contained a 13-amino-acid N-terminal tag upstream of each coding sequence, contributed by the cloning vector. These amino acids were derived from the sequence between the NdeI and BamHI sites in pET11A. Inclusion of this tag was necessary in order to express the proteins in *E. coli*. Attempts to express the NTSF enzymes from PCR products cloned into pET11A as NdeI/BamHI fragments lacking the upstream tag were unsuccessful. We speculate that the additional amino acids facilitated the translation of the mRNAs for the *G. sulfurreducens* NTSF enzymes by *E. coli* ribosomes. Overexpressed proteins were purified by a combination of anion-exchange chromatography and hydrophobic interaction chromatography (NTSFI and NTSFIII) or by ammonium sulfate fractionation, size exclusion chromatography, and hydrophobic interaction chromatography (NTSFII) as described in the supplemental material. Figure 2B shows that nearly homogeneous protein was obtained for each NTSF enzyme. The predicted molecular weights of the NTSF enzymes, based on the genomic sequences, are 50,633 for NTSFI, 45,586 for NTSFII, and 98,394 for NTSFIII.

**Functional analysis of the *G. sulfurreducens* NTSF enzymes.** The phylogenetic analysis suggested that *G. sulfurreducens* NTSFI functioned as a PAP, NTSFII functioned as a C-adding TNT, and NTSFIII functioned as an A-adding TNT. To confirm these assignments, the activity of NTSFI was tested using reaction mixtures optimized for RNA polyadenylation with a model mRNA, the 5650 transcript derived from the *rpsO-pnp* operon of *Streptomyces coelicolor*, as a substrate (10). Positive

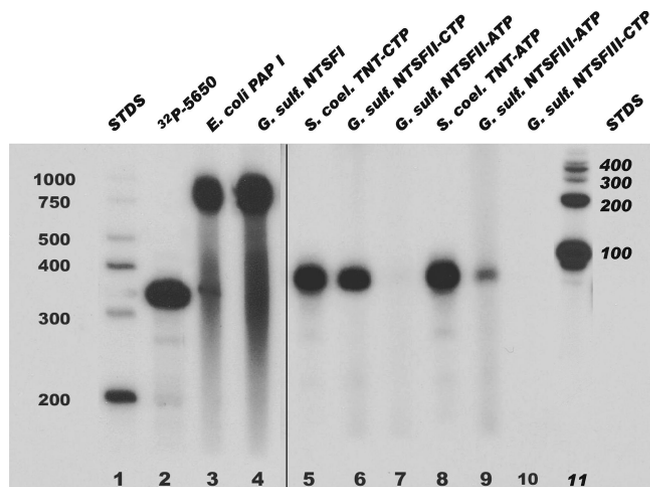


FIG. 3. Autoradiogram of denaturing polyacrylamide gels of the reaction products from assays of *E. coli* PAP I (lane 3), *G. sulfurreducens* (*G. sulf.*) NTSFI (lane 4), *S. coelicolor* (*S. coel.*) SCO3896 with CTP and tRNA<sup>ASP</sup>-C as substrates (lane 5), SCO3896 with ATP and tRNA<sup>ASP</sup>-CC as substrates (lane 6), NTSFII with CTP and tRNA<sup>ASP</sup>-C as substrates (lane 7), NTSFII with ATP and tRNA<sup>ASP</sup>-CC as substrates (lane 8), NTSFIII with CTP and tRNA<sup>ASP</sup>-C as substrates (lane 9), and NTSFIII with ATP and tRNA<sup>ASP</sup>-CC as substrates (lane 10). Lanes 1 to 4 are lanes from a 5% polyacrylamide gel, and lanes 5 to 11 are lanes from a 10% polyacrylamide gel. Lanes 1 and 11 contained end-labeled size markers (STDS). For comparison, lane 2 shows the migration position of the end-labeled *rpsO-pnp* transcript, but it should be noted that unlabeled transcript was used in the polyadenylation assays. The *rpsO-pnp* transcript migrated more rapidly than expected, presumably due to the presence of some secondary structure in the transcript even in the presence of urea (20).

control reaction mixtures for polyadenylation contained authentic *E. coli* PAP I. For the TNT reactions, an *E. coli* tRNA<sup>ASP</sup> lacking either the 3'-terminal CA residues or the terminal A residue and prepared by transcription of plasmid pmBsDCCA (15) as described in Materials and Methods was used as the substrate. NTSFII and NTSFIII were examined for C- and A-adding TNT activity, and the *S. coelicolor* SCO3896 gene product, which has been shown to be a TNT (2, 21), was used as a positive control for C and A addition to tRNA 3' ends. Products from these reactions were separated by polyacrylamide gel electrophoresis on 5 or 10% denaturing gels, followed by autoradiography of the gels, and the results are shown in Fig. 3. Lane 2 shows the migration position of the *rpsO-pnp* transcript used in these experiments (note that the unlabeled transcript was used as a substrate in the PAP assays), lane 3 shows the products obtained by using the polyadenylation reaction mixture containing authentic *E. coli* PAP I, and lane 4 shows the products obtained by using the reaction mixture containing *G. sulfurreducens* NTSFI. Essentially identical patterns were observed in each case, showing the higher-molecular-weight products expected from addition of A residues to the 3' end of the mRNA substrate. Lanes 5 and 8 show the products obtained by using TNT reaction mixtures containing SCO3896 with CTP and tRNA<sup>ASP</sup>-C and ATP and tRNA<sup>ASP</sup>-CC as substrates, respectively. A single band reflecting the addition of C or A to the tRNA<sup>ASP</sup> substrate was observed. Lanes 6 and 7 show the products obtained by using similar reaction mixtures containing *G. sulfurreducens* NTSFII.

It is apparent that the product obtained by using the reaction mixture containing CTP and tRNA<sup>Asp</sup>-C as substrates (lane 6) migrated in the same fashion as the corresponding product obtained by using the reaction mixture containing SCO3896 (lane 5), while only a faint band was observed with ATP and tRNA<sup>Asp</sup>-CC (lane 7). Similarly, lane 9 shows that the product of the reaction catalyzed by NTSFIII with ATP as the substrate and tRNA<sup>Asp</sup>-CC had a mobility identical to that observed for the product obtained by using the reaction mixture containing SCO3896 (lane 8), while NTSFIII was inactive with CTP and tRNA<sup>Asp</sup>-C as substrates (lane 10). These results confirmed the identity of *G. sulfurreducens* NTSFII as a C-adding TNT and the identity of NTSFIII as an A-adding TNT. NTSFII and NTSFIII were inactive with the 5650 transcript as the substrate (data not shown).

#### Characterization of the *G. sulfurreducens* RNA 3' tails.

Given that *G. sulfurreducens* contains a functional PAP, it was of interest to confirm that *G. sulfurreducens* RNAs possess poly(A) tails. Total RNA from *G. sulfurreducens* cells was end labeled with [<sup>32</sup>P]pCp and digested with a combination of RNases A and T<sub>1</sub> as described previously (1, 4, 5). This digestion procedure cleaves all phosphodiester bonds except those following A residues, leaving the poly(A) tails from the 3' ends of the labeled RNA intact. Products of this digestion were displayed on a 12% denaturing polyacrylamide gel, as shown in Fig. 4. Lane 1 shows the migration of a size marker, oligo(dT<sub>18</sub>). Lane 2 shows the migration of tails associated with RNAs from *E. coli*, lane 3 shows the migration of tails from RNAs of *Bacillus subtilis*, lane 4 shows the migration of tails from *B. halodurans* RNA, and lane 5 shows the migration of tails from *G. sulfurreducens* RNA. Lane 5 contains prominent bands for tails that are ca. 20 and 30 residues long, but most of the poly(A) tails associated with the *G. sulfurreducens* RNAs are relatively short (15 residues or less). The pattern obtained for *G. sulfurreducens* is similar to that observed for organisms with heteropolymeric 3' tails, including *Streptomyces* species (2, 4, 5).

To characterize the *G. sulfurreducens* 3' tails further, cDNA cloning was performed as described previously (1, 3, 7) and in Materials and Methods, using total RNA as the template for RT. PCR was performed using gene-specific primers designed from the sequences of the 16S and 23S rRNA genes and the genes encoding NTSFI and NTSFII (see Table S1 in the supplemental material). Results of this analysis are shown in Fig. 5. cDNA cloning revealed the presence of 3' tails associated with decay intermediates (or premature transcription termination products) of 16S and 23S rRNAs and with intermediates from the mRNA populations corresponding to NTSFI and NTSFII. Tails of various lengths were observed for the rRNA and mRNA populations and ranged from a single residue attached to putative decay intermediates of 16S and 23S rRNAs to a 253-nucleotide tail attached at position 2618 of 23S rRNA. It is noteworthy that two 23S cDNAs and one 16S cDNA representing putative decay intermediates that did not possess a 3' tail (designated "adaptor attachment site" [Fig. 5]) were isolated. Rather, these products terminated at presumed cleavage or processing sites in the transcripts in question and may have been isolated prior to posttranscriptional tail addition. Of particular interest is the observation that, as suggested by the results shown in Fig. 4, the 3' tails associated with the

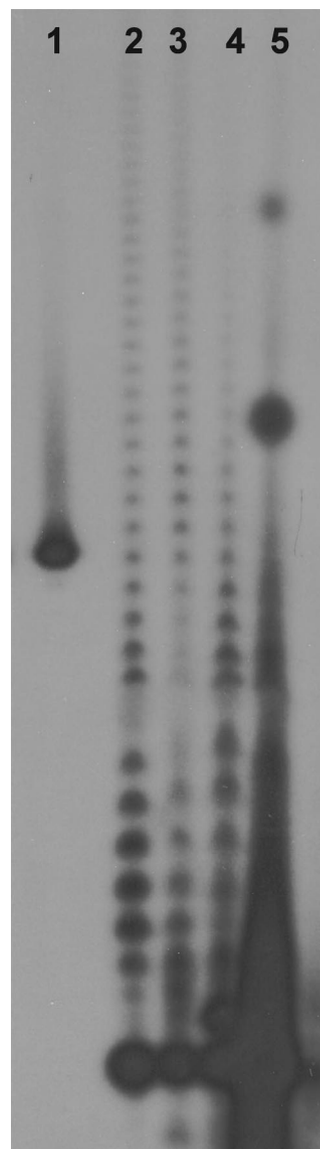


FIG. 4. Measurement of the lengths of oligonucleotide and poly(A) stretches found in the 3' tails of RNAs from *E. coli* DH5 $\alpha$  (lane 2), *B. subtilis* (lane 3), *B. halodurans* C-125 (lane 4), and *G. sulfurreducens* (lane 5). Lane 1 contained an oligo(dT<sub>18</sub>) size standard. Labeled 3' tails were fractionated on a 12% denaturing polyacrylamide gel.

*Geobacter* RNAs are heteropolymeric, containing G, C, and U residues as well as A residues. In *E. coli*, *Streptomyces*, some cyanobacteria, and spinach chloroplasts, polynucleotide phosphorylase is the enzyme responsible for the addition of G, C, and U residues to RNA 3' tails (2, 14, 17, 21, 25). This may also be the case in *G. sulfurreducens*, which, like all other bacteria except mycoplasmas, contains a gene for polynucleotide phosphorylase. It should be noted that, at least in vitro, *G. sulfurreducens* NTSFI utilized CTP and UTP (but not GTP) as substrates in reactions performed under PAP assay conditions (data not shown). *E. coli* PAP I also utilizes other nucleoside triphosphates as substrates in vitro, although only A residues appear to be added to RNA 3' ends in vivo (26). It is possible

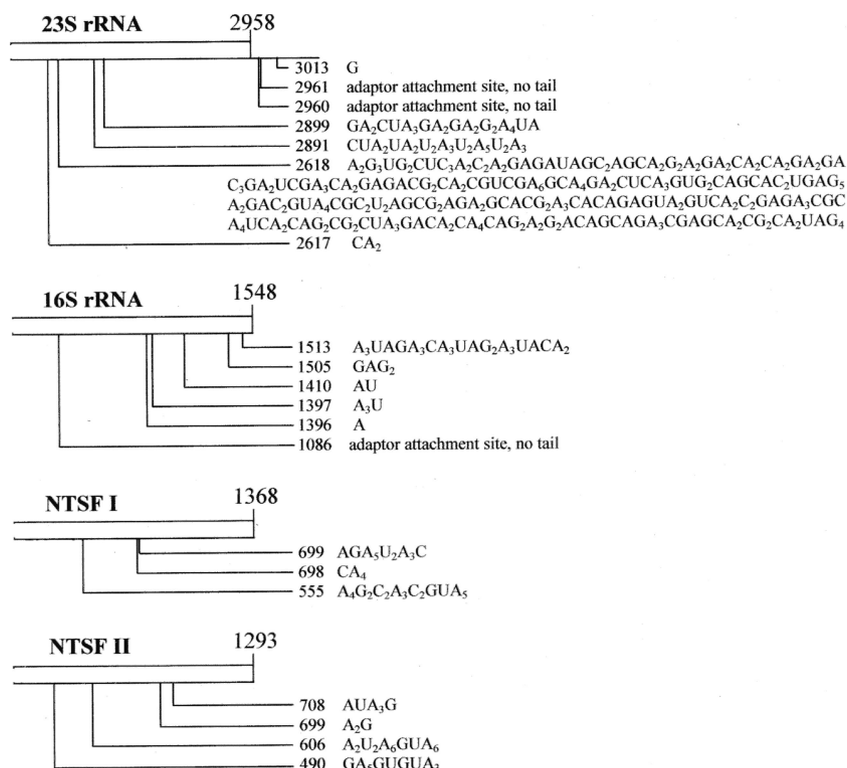


FIG. 5. Positions and compositions of *G. sulfurreducens* RNA 3' tails. The position of each tail was determined from the mature 5' end of rRNA gene or the translation start site of the NTSF enzyme. The number of the last nucleotide of the mature rRNA or translation stop codon is indicated at the top right. The figure is not drawn to scale.

that, unlike the situation in *E. coli*, *G. sulfurreducens* NTSFI utilizes all three nucleoside triphosphates *in vivo*. This would explain our failure to observe tails composed of only A residues in the cDNA cloning experiments whose results are shown in Fig. 5. Alternatively, a combination of NTSFI and polynucleotide phosphorylase activities may be responsible for the synthesis of 3' tails in *G. sulfurreducens*. It is also possible that examination of other *G. sulfurreducens* transcripts will reveal the presence of tails composed of only A residues.

In summary, we showed that unlike the alpha-, beta-, gamma- and epsilonproteobacteria, the low- and high-G+C-content gram-positive bacteria, and many other bacterial genera, at least some deltaproteobacteria encode separate C- and A-adding TNTs and a PAP. We demonstrated that the genes encoding all three of these enzymes are transcribed in *G. sulfurreducens*, and although we did not determine whether the transcripts were translated, we were able to detect all three enzymatic activities in crude *G. sulfurreducens* extracts (data not shown). We speculate that the CCA-adding TNT activity observed in the beta- and gammaproteobacteria evolved via mutations in the genes encoding either the C- or A-adding TNTs of an ancestral species and that subsequent mutations resulted in the conversion of the CCA-adding TNT to a PAP. The apparent absence of a PAP I-like activity in the so-called deeply branching bacterial species, such as *Aquifex*, *Thermotoga*, and *Deinococcus* species, and the cyanobacteria (1, 22, 23) is consistent with the hypothesis that polyadenylation of RNA 3' tails by a PAP I-like enzyme evolved subsequent to the evolution of TNTs. The *G. sulfurreducens* PAP gene and the

PAP I genes of the beta- and gammaproteobacteria may have arisen via independent evolutionary events. The *G. sulfurreducens* PAP gene and the *E. coli* *pcnB* (PAP I) gene are ca. 53% identical. It is also possible, therefore, that the *G. sulfurreducens* gene arose by horizontal gene transfer from a beta- or gammaproteobacterium.

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