

DNA methyltransferases of the cyanobacterium *Anabaena* PCC 7120

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ABSTRACT

From the characterization of enzyme activities and the analysis of genomic sequences, the complement of DNA methyltransferases (MTases) possessed by the cyanobacterium *Anabaena* PCC 7120 has been deduced. *Anabaena* has nine DNA MTases. Four are associated with Type II restriction enzymes (*AvaI*, *AvaII*, *AvaIII*, and the newly recognized inactive *AvaIV*), and five are not. Of the latter, four may be classified as solitary MTases, those whose function lies outside of a restriction/modification system. The group is defined here based on biochemical and genetic characteristics. The four solitary MTases, DmtA/M.*AvaVI*, DmtB/M.*AvaVII*, DmtC/M.*AvaVIII*, and DmtD/M.*AvaIX*, methylate at GATC, GGCC, CGATCG, and rCCGGy, respectively. DmtB methylates cytosines at the N4 position, but its sequence is more similar to N6-adenine MTases than to cytosine-specific enzymes, indicating that it may have evolved from the former. The solitary MTases, appear to be of ancient origin within cyanobacteria, while the restriction MTases appear to have arrived by recent horizontal transfer as did five now inactive Type I restriction systems. One MTase M.*AvaV*, cannot reliably be classified as either a solitary or restriction MTase. It is structurally unusual and along with a few proteins of prokaryotic and eukaryotic origin defines a structural class of MTases distinct from all previously described.

INTRODUCTION

DNA methyltransferases (MTases) in bacteria are most often associated with cognate restriction endonucleases (1), serving to protect the host organism from self-digestion. However, some MTases modify DNA sequences for purposes distinct from restriction. These enzymes, sometimes called orphans, will be termed solitary MTases to emphasize their functional role rather than their possible origins. The best studied enzyme in this group is Dam of *Escherichiacoli* (2). Methylation of DNA by Dam at GATC permits the cell to distinguish recently synthesized (hemimethylated) DNA from older (fully methylated) DNA. The cell uses this information to regulate the timing of DNA synthesis and to choose the proper strand as template in order to correct mismatched bases. MTases nearly identical to Dam in sequence and function appear to be widespread amongst enteric bacteria (see **Discussion**), and MTases of similar function have been observed in the α division proteobacteria (3,4). Little is known, however, about a second MTase in *E. coli*, the CCwGG-specific enzyme

Dcm (2), nor about the range of functions served in bacteria by solitary MTases.

Modification of GATC sequences is prevalent also amongst the cyanobacteria (5), an ancient class of photosynthetic eubacteria. Characterization of GATC modification in cyanobacteria has extended little past demonstrating the general inability of adenine-specific, GATC-recognizing restriction endonucleases to cut genomic DNA (5,6). One gene, *mbpA*, in the fully sequenced genome of the cyanobacterium *Synechocystis* PCC 6803 has been identified whose putative product bears a striking resemblance to Dam of *E. coli* and to other MTases that modify the adenine within GATC (<http://www.kazusa.or.jp/cyano/>). The insensitivity of genomes of the filamentous cyanobacteria within the genera *Anabaena* and *Nostoc* to digestion by the GGCC-specific endonuclease *HaeIII* (6,7) coupled with the apparent absence of a corresponding restriction enzyme (8), led us to speculate that these cyanobacteria carry at least two solitary MTases.

Type II DNA MTases may be grouped according to the positions of the modifications they catalyze. Enzymes methylating cytosine at the 5-carbon of the pyrimidine ring form a coherent group, according to a common ordering of ten well conserved blocks within their amino acid sequences (9). Enzymes that methylate the exocyclic N4 amine of cytosine or N6 amine of adenine form a looser group whose members share some sequence similarities. The latter group has been subdivided into three types, α , β , and γ , according to their different orderings of common motifs and sequence similarities within these motifs (10). Crystal structures have recently been deduced for two C5-methylcytosine (5mC) MTases (11,12) and for an N4-methylcytosine (N4mC) or N6-methyladenine (N6mA) MTase from each of the three types (13-15). These have led to the view that all MTases share a common architecture (16).

Type I DNA MTases differ markedly from Type II in both structure and mechanism of action (17). Most notably, the former consist of two distinct subunits, a catalytic subunit (M, encoded by *hsdM*) and a subunit (S, encoded by *hsdS*) responsible for specific binding to DNA. The addition of a third subunit (R, encoded by *hsdR*) confers upon the complex the additional ability to cleave DNA unmethylated at the target recognition site. Type I DNA MTases have been divided into four families, IA through ID, originally on the basis of antigenic crossreactivity.

The possibility that DNA replication, modulated by a solitary MTase, might regulate differentiation within filaments of *Anabaena* PCC 7120 (henceforth referred to as *Anabaena*) motivated us to screen the strain for genes encoding such enzymes. Unusual features in the deduced amino acid sequences of protein encoded by two of the genes found led us to examine them more closely within the context of current understanding of structure/function relationships.

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MATERIALS AND METHODS

DNA isolation and cloning

DNA was isolated from *Anabaena* grown on BG11, modified as previously described (18), by vortexing with glass beads in phenol (19). Plasmids bearing expressed DNA MTase genes *avaMV*, *dmtB/avaMVII*, and *dmtD/avaMIX* were isolated essentially according to the protocol of Kiss *et al.* (20). In brief, *Anabaena* DNA partially digested with *Sau3AI* to an apparent average size of 2 to 3 kb, was ligated with pBluescript II KS+ (Stratagene). The ligated DNA was electroporated into *E. coli* strain GM4715 (*dam⁻ mcrB⁻*) (2), for cloning of *avaMV*, or K803 (*hds3 mcrB*) (21), for cloning of *dmtB* and *dmtD*. Approximately 400,000 colony forming units were incubated in 4 ml LB, 1 hr, then 100 ml LB + 50 µg/ml ampicillin, 6 hrs. Plasmid DNA was isolated through a Qiaprep spin column. Approximately 1 µg plasmid DNA was digested for at least 6 hrs with excess restriction enzyme (*DpnII* for *avaMV*, *HaeIII* for *dmtB*, and *BsrFI* for *dmtD*). The DNA was precipitated, and 20% of it was used to electroporate the same strain of *E. coli* to ampicillin resistance. For *dmtB* and *dmtD*, this was sufficient to isolate several colonies carrying the gene. For *avaMV*, it was necessary to go through a second round of amplification, starting with several hundred colonies scraped off of the transformation plate.

The gene encoding DmtA was isolated by PCR amplification using the primers 5'-ATATCATCAGGTGATCGCGC-3' and 5'-TTTGGCGCTGGGATAGTACC-3'. The gene encoding M.AvaIII and its downstream open reading frame (ORF) were isolated on the same PCR-amplified fragment, using the primers 5'-CAGTATGCTTCAGGGGAAA-3' and 5'-GTTGTTGATGCTTTGAGCGA-3'. Both sets of primers were derived from available genomic sequence (<http://www.kazusa.or.jp/cyano/>) of the regions identified as described below and in the **Results**.

Routine plasmid isolation and manipulations followed standard procedures. The presence of small DNA digestion fragments was assessed by electrophoresis on 3% gels made with Metaphor (FMC). DNA was prepared for pulsed-field gel electrophoresis as previously described (22)

Analysis of DNA and protein sequences

Plasmid DNA purified through a Qiaprep minispin column was sequenced by the University of Chicago DNA Sequencing Facility. Most of the DNA, and all regions where there were any ambiguities, were sequenced from both strands. ORFs were identified and translated using EditBase, provided by Niels Nielsen (Purdue University) and OrfFinder (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>). Protein sequences were aligned by ClustalX (23), which uses distance matrices and nearest neighbor joining, and displayed by ClustalInColor, a locally written program that facilitates the coloring of amino acids to accentuate relationships between groups of sequences. Tree structures were drawn using TreeView (24) and colored by hand.

In sequence comparisons, an amino acid was judged to be similar to the consensus amino acid if the entry of the pair in BLOSUM62, a matrix of amino acid similarities (25), gives a positive value. Groups of amino acids were judged to be similar to each other if they fell into one of the following sets defined by positive values in BLOSUM62 with respect to the first listed amino acid: DEN, EDKQ, HNY, LIMV, SANT, QEKR, YFHW. Since BLOSUM62 groups amino acids by the frequency with which they occur at the same position in

aligned positions, the sets do not necessarily correspond to obvious biochemical characteristics.

DNA MTases and restriction endonucleases were found by searching GenBank with Gapped BLAST (26) and searching other databases listed below (last searched June 2000). DNA sequences from *Synechocystis* PCC 6803 and preliminary DNA sequences from *Anabaena* were obtained from Cyano-Base (www.kazusa.or.jp/cyano/). Preliminary DNA sequences from *Nostoc punctiforme* ATCC 29133 (henceforth called *N. punctiforme*), and *Enterococcus faecalis* V583 were obtained from the DOE Joint Genome Institute (http://spider.jgi-sf.org/JGI_microbial/html/nostoc_homepage.html), and from The Institute for Genomic Research (<http://www.tigr.org>), respectively. DNA MTase sequences were identified in genomic sequences by three means: annotation of the sequence, annotation in REBASE (reference 27 and <http://rebase.neb.com>), a depository of information regarding restriction endonucleases and DNA MTases, and BLAST searches using the following archetypical sequences, with each followed by its class (10,17) and accession/enzyme number: M.EcoBI (IA; rebase|3381), M.EcoEI (IB; rebase|3386), M.EcoR124I (IC; rebase|3392), M.StySBLI (ID; rebase|3571), M.Hae III (II-5mC; sp|P20589), M.Dpn IIA (II-N6mAα; sp|P04043), M.Nla III (II-N4mCα; pir|XYNHAL), M.HinFI (II-N6mAβ; rebase|3429), M.Pvu II (II-N4mCβ; gb|AAA96336), M.EcoPI (III-N6mAβ; sp|P08763), M.Pst I (II-N6mAγ; rebase|3483). An ORF was judged to encode a DNA MTase if it matched a positively identified DNA MTase with a BLAST score better than 10⁻⁴. Unidentified ORFs were searched for known motifs using EMOTIF (reference 28 and <http://motif.stanford.edu/emotif/>). Possible membrane-spanning regions were found in protein sequences by the Dense Alignment Surface method (DAS) (reference 29 and <http://www.sbc.su.se/~miklos/DAS/maindas.html>).

Chemical analysis of base composition

Plasmid DNA isolated from the *dam⁻ dcm⁻* strain *E. coli* GM48 (2) was hydrolyzed by a procedure modified from that described by Gehrke *et al.* (30). About 50 µg DNA in 100 µl deionized water was denatured by boiling for 5 minutes, and plunging the microfuge tube into ice water. 200 µl 30 mM sodium acetate pH 5.2 and 10 µl 20 mM zinc sulfate was added, followed by 6 units of nuclease P1 (Roche). After 6 hrs of incubation at 37°, 40 µl of 0.5 M Tris (not adjusted for pH) and 5 units shrimp alkaline phosphatase (Roche) was added prior to an overnight incubation at 37°. The resulting nucleosides were purified through a microcon YM-3 filter (Millipore). The purified nucleosides were analyzed on a 3.9x150mm Nova-Pak reverse phase C18 column (Waters) using a previously described step gradient (30).

RESULTS

Cloning of three genes from *Anabaena* encoding DNA modification enzymes

As part of a general screen, DNA from *Anabaena* was digested with a number of restriction enzymes and run out on conventional and sometimes pulse field gels. We confirmed previously published results (6,31) that DNA from *Anabaena* could not be cut by *Hae* III, which recognizes GGCC, nor by *DpnII*, which recognizes GATC. It is, however, cut by *DpnI*, which cuts only at methylated GATCs. In addition, *Anabaena*

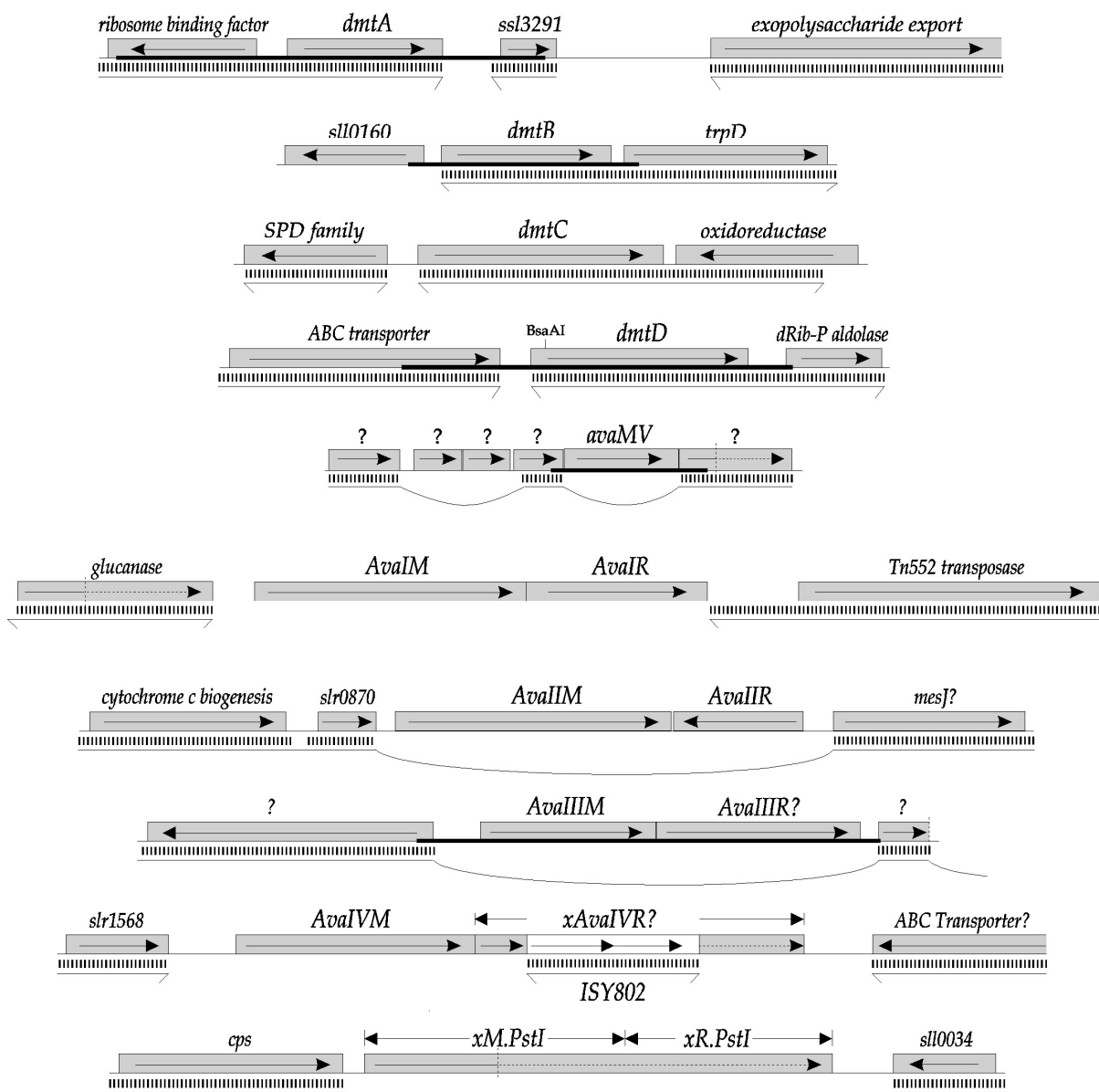


Figure 1. Genetic context surrounding genes encoding or potentially encoding DNA MTases of *Anabaena*. Gray boxes represent ORFs, with the direction of translation shown. The white box indicates an insertion sequence. Vertical dotted lines indicate the point at which a frameshift occurs. Thick horizontal lines indicate the region that was cloned. Thin horizontal lines separated from the *Anabaena* genes by vertical bars indicate that a contiguous DNA sequence from *N. punctiforme* exhibits sequence similarity in that region. Horizontal lines with bent ends represent DNA sequences that continue without similarity to the *Anabaena* sequence. Loops indicate interpolations of sequences without similarity to the *Anabaena* sequence. Genes with names preceded by “x” are defective by reason of a frameshift, nonsense codon, or insertion sequence.

DNA could not be cut by *BsrFI*, which recognizes the degenerate sequence rCCGGy (data not shown). The three genes responsible for the protection of *Anabaena* from digestion by *DpnII*, *HaeIII*, and *BsrFI* were cloned by their ability to extend protection in *E. coli* to plasmid DNA bearing the genes and characterized as described below.

Analysis of *avaMV*, encoding a GATC-specific DNA MTase

A plasmid isolated as resistant to cutting by *DpnII* (which recognizes GATC) proved to be sensitive to digestion by *DpnI* (which recognizes G^{me}ATC) and *Bsp143I* (which recognizes GATC regardless of adenine methylation). A 1.3-Kb fragment subcloned from that plasmid (forming pUR147) retained these

characteristics. Analysis of the original plasmid from a *dam*⁻ *E. coli* by HPLC showed a minor base with the same retention time as N6mA. These results are consistent with pUR147 encoding an enzyme that modifies the adenine residue within GATC.

The sequence of the insert within pUR147 revealed a single large ORF (Figure 1), named *avaMV* (see **Discussion**), potentially encoding a protein of 210 amino acids. The region of the *Anabaena* genome surrounding *avaMV* was carefully examined to determine whether there may be any genes nearby that could conceivably encode a restriction endonuclease (Figure 1). Such genes are often not recognizable by sequence, but of 227 sequenced Type II

restriction endonucleases found in GenBank, only 8 have fewer than 200 amino acids, and the smallest, *R.PvuII*, has 157 amino acids. Thus it is reasonable to expect a gene of at least that size. Furthermore of 110 cases where the relative positions of the genes encoding the restriction enzyme and corresponding MTase(s) were readily available, only one was found where the gap between the two genes exceeded 1000 bp (1451 bp for *Eco47I*), and the average gap size was only 82 bp. Furthermore, only one pair of genes (*R.CviAI* and *M.CviAI* from a chlorella virus) was found where a gene not part of the RM system intervened between the two. Thus we deemed it unnecessary to look beyond 2 Kb and more than two large ORFs from a gene encoding a MTase to seek a gene encoding a corresponding endonuclease.

The available 1968 base sequence 5' to the *avaMV* gene lacks any ORF capable of encoding a protein greater than 119 amino acids. Immediately 3' to *avaMV* lies an ORF capable of encoding a protein of 200 amino acids similar through most of its length to hypothetical proteins from the plant *Arabidopsis thaliana* (AAD23616) and from *Streptomyces coelicolor* (gb|CAB93740), but it is disrupted by a frameshift in the 5' end of the gene. Other ORFs within the 2500 bp region 3' to *avaMV* are either very small (the largest is 106 amino acids) or show great similarity to proteins of known function.

It was of interest to see whether *avaMV* is found generally in cyanobacteria. The sequence surrounding *avaMV* is found in the closely related strain *N. punctiforme*, but *avaMV* itself is not (Figure 1). Instead it is replaced almost precisely by a different ORF that is not significantly similar to any in GenBank. No protein in the region shows any significant similarity to proteins of the distantly related cyanobacterium *Synechocystis* PCC 6803.

All previously characterized MTases with GATC specificity and one cyanobacterial ORF (*mbpA*) highly similar to genes encoding such enzymes fall into either the α class (nine genes) or β class (two genes) of MTases, and we anticipated that *M.AvaV* would do the same. In fact, a search through known protein sequences brought up three bacterial and several eukaryotic proteins (Table 1 and Figure 2) that do not readily fit into any described class. One, *M.MunI*, is the MTase of a restriction/modification (RM) system, and another, MT-A70, is the S-adenosylmethionine (AdoMet)-binding subunit of a human RNA MTase that acts on adenine residues (32). The functions of the other proteins are unknown.

These proteins all have the region, Motif IV, most highly conserved in MTases and regions that could possibly correspond to Motifs X and I, as previously noted for *M.MunI* (33). All three regions participate in the binding of AdoMet

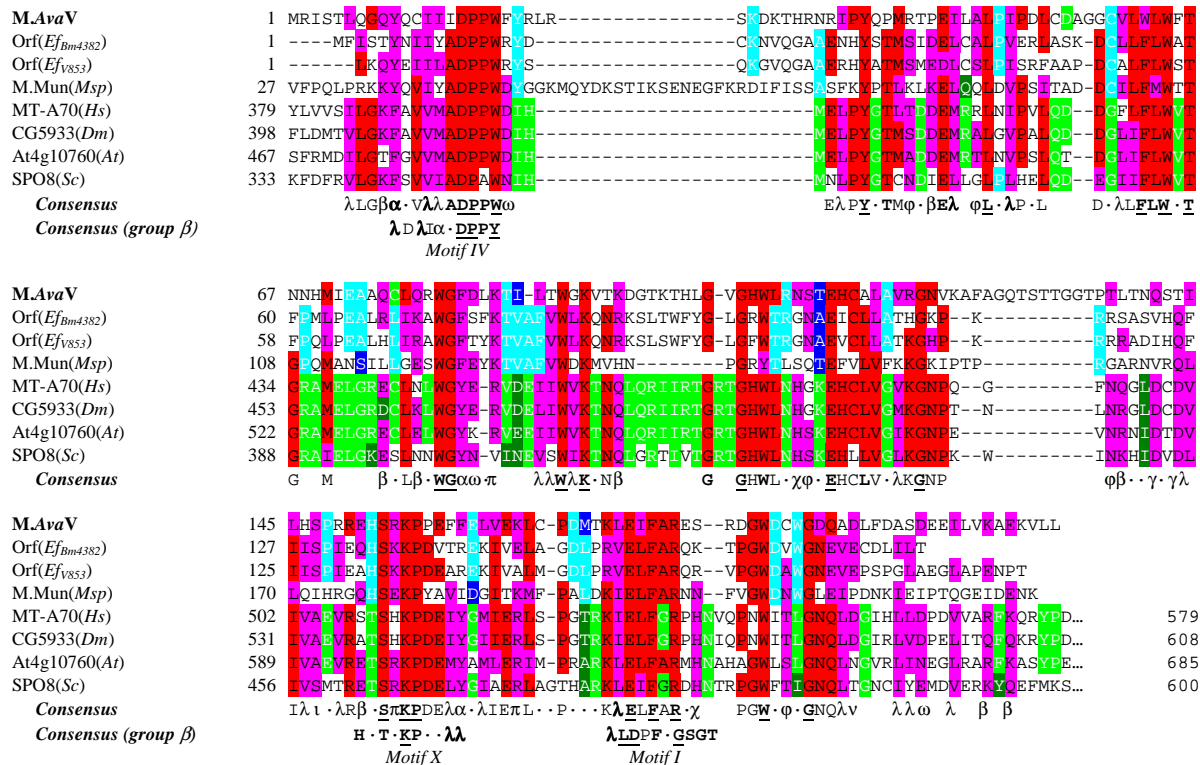


Figure 2. Alignment of *M.AvaV* with known and putative DNA and RNA methyltransferases. Sequences listed in Table 1 (except the mouse sequence, which is substantially the same as the human sequence) were aligned as described in **Materials and Methods**. For each position, amino acids substantially conserved over the entire group are colored red, and similar amino acids colored magenta. Identical amino acids within the bacterial subgroup (first four sequences) are colored cyan and similar amino acids dark blue. Identical amino acids within the eukaryotic subgroup (last four sequences) are colored bright green and similar amino acids dark green. A consensus sequence over the entire group is shown below the alignment and for the corresponding region of Group β N6mA DNA MTases (10). A letter is shown if greater than 50% of the sequences have the same amino acid residue or class of amino acids at the given position. The letter is in bold type if the frequency exceeds 75% and underscored if the position is invariant. Greek letters represent classes of amino acids: α , aromatic (FHXY); β , big hydrophilic (EKQR); λ , leucine family (ILMV); v, negative (DE); π , positive (KR); γ , glutamate family (EQDK); χ , (NHY); ϕ , (ANST); ι , one carbon (AS); ω ,

Table 1. Proteins similar to putative solitary Type II DNA methyltransferases of *Anabaena*

Protein ^a	Recognition sequence ^b	Source	Score ^c	Accession no. ^d
Protein sequences similar to M.AvaV				
1. Ef-Orf18	?	<i>Enterococcus</i>	4·10 ⁻²¹	gb AAF72345.1
2. Ef10277.13628	?	<i>Enterococcus</i>	1·10 ⁻²⁰	genome site ^e
3. DmCG5933	?	<i>Drosophila</i>	6·10 ⁻¹⁷	gb AAF56221.1
4. MT-A70	?	<i>Homo</i>	2·10 ⁻¹⁶	gb AAB71850.1
5. AT4g10760	?	<i>Arabidopsis</i>	2·10 ⁻¹⁶	gb CAB81177.1
6. Mouse ORF	?	<i>Mus</i>	2·10 ⁻¹⁶	gb AAD33673.1
7. Spo8	?	<i>Saccharomyces</i>	3·10 ⁻¹⁵	ref NP_011323.1
8. M.MunI	CAATTG	<i>Mycoplasma</i>	4·10 ⁻¹²	sp P43641
Protein sequences similar to DmtA/M.AvaVI				
1. DmtA(<i>Np</i>)	GATC?	<i>Nostoc</i>	1·10 ⁻¹¹⁸	genome site ^f
2. M.PgiI	GA ⁶ TC	<i>Porphyromonas</i>	3·10 ⁻⁶⁶	pir S34414
3. Ph-orf	GATC?	<i>Pyrococcus</i>	8·10 ⁻⁶⁰	pir C71096
4. St8-orf3	GATC?	<i>Streptococcus</i>	3·10 ⁻⁵⁹	gb CAB46541.1
5. M.DpnIIA	GA ⁶ TC	<i>Diplococcus</i>	9·10 ⁻⁵⁹	sp P04043
6. M.LlaDCHIA	GA ⁶ TC	<i>Lactococcus</i>	8·10 ⁻⁵⁸	sp P50179
7. M.MjaIII	GATC	<i>Methanococcus</i>	7·10 ⁻⁵³	sp Q58015
8. MbpA	GATC?	<i>Synechocystis</i>	6·10 ⁻⁵²	pir S77170
22. DmtB	GGCC	<i>Anabaena</i>	7·10 ⁻¹⁷	gb AAF75229
Protein sequences similar to DmtB/M.AvaVII				
1. DmtB(<i>Np</i>)	GGCC?	<i>Nostoc</i>	1·10 ⁻¹²³	genome site ^f
2. Sli0729	GGCC?	<i>Synechocystis</i>	1·10 ⁻⁸⁸	pir 76841
3. St8-orf3	GATC?	<i>Streptococcus</i>	6·10 ⁻²⁰	gb CAB46541.1
4. M.PgiI	GA ⁶ TC	<i>Porphyromonas</i>	4·10 ⁻¹⁸	pir S34414
5. M.MjaIII	GATC	<i>Methanococcus</i>	2·10 ⁻¹⁶	sp Q58015
6. M.LlaDCHIA	GA ⁶ TC	<i>Lactococcus</i>	5·10 ⁻¹⁴	sp P50179
7. M.CviAI	GA ⁶ TC	<i>Chlorella virus</i>	5·10 ⁻¹⁴	pir T18083
8. M.DpnIIA	GA ⁶ TC	<i>Diplococcus</i>	2·10 ⁻¹³	sp P04043
9. M.CviQVI	GA ⁿ TC	<i>Chlorella virus</i>	5·10 ⁻¹³	gb AAC03126.1
10. MbpA	GATC?	<i>Synechocystis</i>	9·10 ⁻¹³	pir S77170
11. M.StsI	GGG ⁶ TC	<i>Streptococcus</i>	1·10 ⁻¹¹	sp P29347
16. M.BalI	TGGCCA	<i>Brevibacterium</i>	8·10 ⁻⁷	pir S71506
Protein sequences similar to DmtC/M.AvaVIII				
1. DmtC(<i>Np</i>)	CGATCG?	<i>Nostoc</i>	1·10 ⁻¹⁸⁸	genome site ^f
2. M.XorII	CGATCG	<i>Xanthomonas</i>	1·10 ⁻¹²⁸	sp P52311
3. SynMI	CGATCG	<i>Synechocystis</i>	1·10 ⁻⁹⁷	pir S76359
4. M.NspHI	rC ⁵ ATGy	<i>Nostoc</i>	6·10 ⁻⁴³	gb AAC97192.1
5. M.NspI	rC ⁵ ATGy	<i>Nostoc</i>	1·10 ⁻⁴²	gb AAC97190.1
6. M.DdeI	C ⁵ TnAG	<i>Desulfovibrio</i>	5·10 ⁻⁴²	sp P05302
Protein sequences similar to DmtD/M.AvaIX				
1. DmtD(<i>Np</i>)	rCCGGy?	<i>Nostoc</i>	1·10 ⁻¹⁷⁶	genome site ^f
2. M.NmeDI	rCCGGy?	<i>Neisseria</i>	1·10 ⁻¹¹⁹	gb CAB59897.1
3. M.Cfr10I	rC ⁵ CGGy	<i>Citrobacter</i>	1·10 ⁻¹⁰⁶	pers. commun. ^g
4. M.Bse634I	rCCGGy	<i>Bacillus</i>	4·10 ⁻⁵⁸	pers. commun. ^g
5. M.HaeIII	GGC ⁵ C	<i>Haemophilus</i>	9·10 ⁻²⁴	sp P20589
6. M.MthTI	GGCC	<i>Methanobacter</i>	3·10 ⁻²³	sp P29567
7. M.Phi3TII	TC ⁵ GA	<i>Bacillus phage</i>	6·10 ⁻²³	pir S47248
25. SynMI	CGATCG	<i>Synechocystis</i>	1·10 ⁻¹⁴	pir S76359

^aProtein sequences obtained by BLAST search (26) of the combined GenBank, PDB, SwissProt, PIR, and PRF databases (unless otherwise noted), last accessed June 2000, ranked by score.

^bRecognition sequences of MTases are given along with the position, if known, where they methylate: A⁶, N6-methyladenine; C⁵, C5-methylcytosine.

^cScore represents expected number of proteins at least as similar to target within the combined databases.

^dgb, GenBank; sp, SwissProt; pir, Protein Information Resource; ref, NCBI Reference Sequence project.

^eSequence obtained from the *E.faecalis* web site (<http://www.tigr.org>), last accessed June 2000.

^fSequences obtained from the *N.punctiforme* web site (http://spider.jgi-psf.org/JGI_microbial/html/nostoc_homepage.html), last accessed June 2000.

^gSequences graciously provided by Virgis Siksnys (Fermentas).

(10). The Motif I assignment is particularly dissatisfying because the aligned sequences all lack a hydrophobic residue four amino acids prior to F and a G residue two amino acids after F, both found in almost all known adenine MTase sequences (34).

Analysis of *dmtB*, encoding a GGCC-specific DNA MTase

A second plasmid, pUR109, was isolated repeatedly from the library of *Anabaena* inserts, based on its ability to withstand digestion by *HaeIII*, which recognizes and cuts at GGCC. We

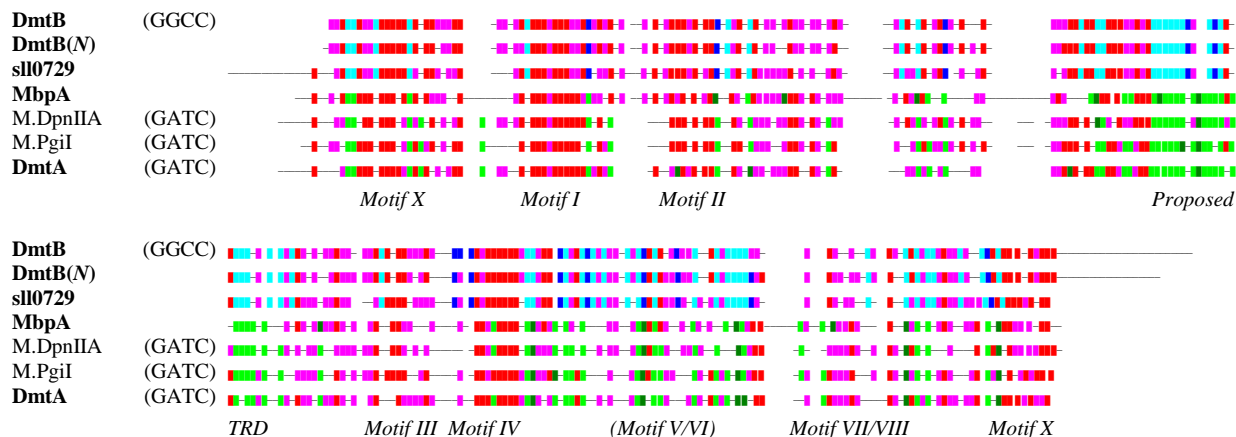


Figure 3. Alignment of DmtB and DmtA with representative Group α Nm6A/Nm4C DNA methyltransferases. Names of cyanobacterial sequences are shown in bold. Coloring conventions are as describe in the legend to Figure 2, with the cyano/dark blue and green/dark green groups representing DNA MTases of proven and putative GGCC-specificity and GATC-specificity, respectively. The motifs (including the TRD, target recognition domain) are those described by Tran *et al.* (15).

expected that the enzyme encoded by pUR109 would determine a GGCC-specific MTase, which, like all others known, methylated at 5mC. To test this, pUR109 DNA was isolated from the *dam⁻ dcm⁻* strain Gm48 and hydrolyzed. Surprisingly, gas chromatography showed a minor base that eluted with the same retention time as N4mC, not 5mC. *Hae*III is sensitive to N4 methylation at the outer cytosine (35), but nothing has been reported regarding its sensitivity to N4 methylation at the inner cytosine.

To identify the precise site of N4mC methylation, pUR109 was digested with *Msp*I. pUR109 contains one site (GGCCCGG) where a GGCC sequence overlaps by one base with the recognition site (CCGG) of *Msp* I. A derivative of pUR109 was made that contains an additional site (GGCCGG) that overlaps by two bases. The ability of *Msp* I to cut at the first site but not the second is consistent with known sensitivity of *Msp*I to N4mC methylation at the outer cytosine of CCGG (36) only if the enzyme encoded by pUR109 methylates at the inner cytosine of GGCC.

The sequence of the 1211-bp insert within pUR109 contains one large ORF (Figure 1), denoted *dmtB* (for DNA MTase; see **Discussion**), capable of encoding a protein of 293 amino acids. There are at least 11 distinct GGCC-specific MTases from diverse sources that have been characterized sufficiently to distinguish the MTase type, and all methylate at the C5 position (7 of the MTases are from a *Bacillus* or its phage and the remaining are from *Fusobacterium*, *Haemophilus*, *Methanobacterium*, and *Neisseria*) (27). The deduced protein sequence of DmtB does not fall into this class but instead shows significant similarity to N6mA-specific MTases of the α class, particularly those that recognize GATC (Table 1). The first similar sequence with known specificity that does not methylate at GATC is that of *M.BalI*, an N4mC MTase. DmtB is most similar to two predicted proteins from cyanobacteria: Orf352.2360R from *N. punctiforme* and SII0729 from *Synechocystis* PCC 6803. They share 73% and 55% amino acid identity over the full length of the protein.

In order to assess the relationship of a GGCC-modifying enzyme to those that modify at GATC, eight such MTases of the α class were compared to the predicted amino acid

sequence of DmtB, and a representative part of the comparison is shown in Figure 3. The regions of similarity to these MTases are focused on conserved motifs spread throughout the protein. The target recognition domain (TRD), the region through which MTases make contact with DNA, is highly conserved amongst the adenine-modifying, GATC-specific MTases, but in this region, there is little similarity with DmtB. The corresponding region in DmtB is highly similar to that in SII0729 but bears no resemblance to the TRDs of known GGCC-specific MTases. The motifs that are particularly well conserved (Motifs X, I-IV) are those that participate in the binding of AdoMet and in the case of Motif IV serves as the catalytic site. Those motifs that are least conserved (Motifs V-IX) are required for the structural integrity of the protein (15).

It seemed peculiar that a MTase that presumably methylates a cytosine within GGCC should appear so similar to MTases that methylate an adenine within GATC, so a dendrogram was constructed from all known MTase sequences of the α class, including those that methylate cytosine at the N4 position (Figure 4). The tree uses sequences in which the TRD region has been removed, so as not to bias the relationships by regions that are known to be constrained by target-specificity, but the resulting tree was not appreciably different from that constructed with full sequences (not shown). In both cases, the sequences of the N6-adenine-methylating enzymes clustered together, apart from sequences of the N4-cytosine-methylating enzymes. DmtB fell into the N6mA group, as did a second N4mC MTase, *M.BalI*.

If DmtB serves to protect GGCC sequences from a corresponding restriction enzyme of *Anabaena*, then one would expect the genes encoding the RM system to lie nearby one another. Immediately downstream lies a gene very similar to that encoding anthranilate synthetase (TrpD) from diverse bacteria. All other ORFs capable of determining a protein greater than 100 amino acids and lying within 2 kb of either side of *dmtB* encode membrane-spanning proteins (data not shown) or a protein that exist in *Nostoc* in four copies. These were judged unlikely candidates to encode a restriction endonuclease.

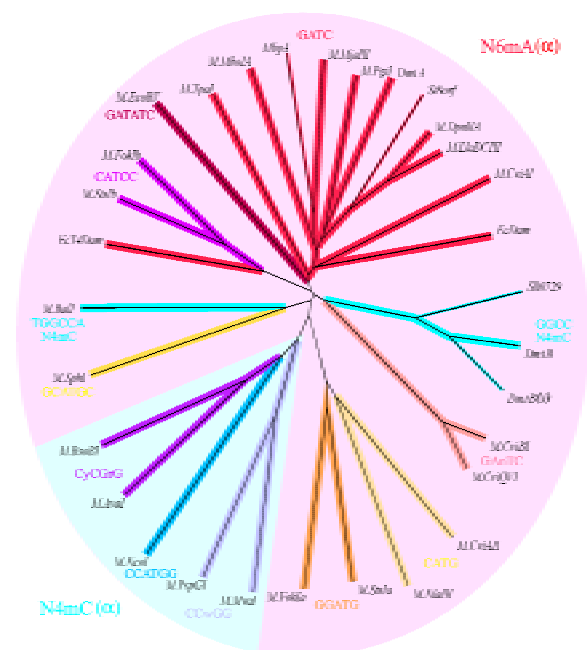


Figure 4. Dendrogram of all available sequences of N4 and N6 methyltransferases of the α group. The sequences compared consisted of the N-terminus and C-terminus, with the central TRD removed. The TRD was liberally defined as the block corresponding to residues 77 through 172 according to *M.DpnII* coordinates (15). The branches of all N6mA MTases are some hue of red and of all N4mC MTase some hue of blue. Thin lines indicate that the target sequence of the enzyme has not been established by biochemical criteria. Besides those sequences listed in Table 1, the following were found through the indicated accession numbers: *M.AvaI* (CAA66984), *M.BsoBI* (CAA66933), *M.CviAII* (S27901), *M.CviBI* (Q01511), *M.EcDam* (P00475), *M.EcoRV* (P04393), *M.EcoT4Dam* (X17641), *M.FokI* (P14871), *M.MboIA* (P34720), *M.MvaI* (P14244), *M.NcoI* (AAC23514), *M.NlaIII*, *M.PspGI* (AAC67523), *M.SphI* (AAB40377), and *M.TpaI* (AAB82782). *M.FokI* and *M.StsI* consist of two separable MTases that are fused as a single polypeptide. The N terminal portion (a) recognizes one asymmetric target and the C-terminus (b) recognizes the other.

Analysis of *dmtD*, encoding an rCCGGy-specific DNA MTase

One element (pUR115) of a library of *Anabaena* DNA was consistently isolated after selection for resistance to *BsrFI*. Plasmid DNA was fully protected only when isolated from cells in the stationary phase of growth. The rCCGGy sites within pUR115 were resistant not only to *BsrFI* but also to *MspI*, which cuts at the internal CCGG tetranucleotide. *MspI* recognizes CCGG and is sensitive to methylation at the C5 position of the outer cytosine but not to methylation at the inner cytosine. The resistance of pUR115 to *MspI*, coupled with the sequence analysis later in this section, suggests that the enzyme encoded by the plasmid modifies at least the 5' cytosine of rCCGGy at the C5 position.

From the inability of *BsrFI* to cut genomic DNA from *Anabaena* and its inability to cut pUR115 (which contains GCCGGC, GCCGGT, and ACCGGC sites), it follows that the modification enzyme protects all sequences within rCCGGy. However, it is still conceivable that the enzyme modifies other sequences as well. To assess this possibility, the *MspI* digest of pUR115 was examined more closely. pUR115 contains 11

of the 12 possible nCCGGn sequences not described by rCCGGy (only TCCGGA is lacking). Since *MspI* cut at all of these sites (data not shown), the sequence recognized by the encoded enzyme cannot be more redundant at its outer bases than rCCGGy.

Only one intact ORF was found within the sequenced 2071-bp fragment (Figure 1). When that fragment was interrupted at the *BsaAI* site by insertion of a kanamycin resistance cassette, protection against *BsrFI* was lost. The intact ORF evidently encodes a DNA modifying enzyme and was named *dmtD*. It is surrounded by two genes of readily identifiable function. Further downstream from *dmtD* lies a gene encoding a protein of unknown function with strong similarity to a protein in *Synechocystis*, which does not possess a DmtD-like MTase. No other significant ORFs lie within 1700 bp of *dmtD*.

The 385 amino acid protein sequence of DmtD, deduced using the first potential start codon of the ORF, was compared to known protein sequences, and more than a hundred sequences with e-values of less than 10^{-5} were found. Most of these had been identified as 5mC DNA MTases. Particularly prominent were the DNA MTases *M.Cfr10I* and *M.Bse634I*, both of which recognize and modify rCCGGy, and many DNA MTases recognizing the sequence GGCC (Table 1).

The sequence of DmtD contains all six highly conserved motifs common to 5mC MTases and matches all 21 of the most conserved residues (data not shown), those residues common to almost all 5mC MTases. Alignments with 20 5mC MTases revealed lesser but significant similarity between DmtD and the less well conserved motifs II, III, V, and VII. DmtD shows no significant similarity to most 5mC MTases in the TRD region, where the enzyme makes contact with its DNA substrate. However, it shares in this region (residues 257 to 313) 43 of 67 amino acid residues with *M.Cfr10I* and 35 of 67 with *M.Bse634I*.

Analysis of *dmtA* and *dmtC*, encoding GATC- and CGATCG-specific DNA MTases

The release of the partial genomic sequence of *Anabaena* made it possible to screen the strain for genes encoding MTases on the basis of sequence similarity rather than function. The proteins predicted from the available sequence were compared to several representative 5mC, N4mC, and N6mA MTases covering all established groups, and significant hits were further analyzed.

One predicted protein showed considerable sequence similarity to class α N6mA MTases, particularly those that recognize GATC. Since the GATC-recognition ability of *M.AvaV* had been established, we were keen on determining whether *Anabaena* truly possesses a second such enzyme. The gene was cloned by PCR on a fragment containing no other significant ORFs and placed parallel to the *lac* promoter of pBluescript. The resulting plasmid was partially digested by both *DpnI* (cutting at G^{me}ATC) and *DpnII* (cutting at GATC). Activity, while evident, was not sufficient either to protect the plasmid from digestion by *DpnII* or to permit full digestion by *DpnI*. On the basis of this activity, the cloned gene was named *dmtA*. The limited protection afforded by *dmtA* in *E. coli* may explain why the gene was not picked up by the strategy that led to the cloning of *avaMV*. *dmtA* is flanked by two large open reading frames of recognizable function. Besides these, there are no ORFs capable of encoding protein greater than 95 amino acids within 2000 bp of *dmtA*.

Table 2. Proteins similar to proven and putative Type II DNA R/M methyltransferases of *Anabaena*

Target	Protein ^a	Recognition sequence ^b	Source	Score ^c	Accession ^d
Protein sequences similar to <i>AvaI</i> R/M proteins					
M. <i>AvaI</i>	1. M. <i>NspIII</i>	CyCGrG	<i>Nostoc PCC 7524</i>	4·10 ⁻¹⁴¹	gb AAC97192
	2. M. <i>BsoBI</i>	CyCGrG	<i>Bacillus</i>	6·10 ⁻⁸³	gb CAA66933
R. <i>AvaI</i>	1. R. <i>BsoBI</i>	CyCGrG	<i>Bacillus</i>	1·10 ⁻¹⁰⁴	gb CAA66932
	2. R. <i>NspIII</i>	CyCGrG	<i>Nostoc PCC 7524</i>	1·10 ⁻¹⁰³	gb AAC97193
	Next match			> 5	
Protein sequences similar to <i>AvaII</i> R/M proteins					
M. <i>AvaII</i>	1. M. <i>SinI</i>	GGwC ⁵ C	<i>Salmonella</i>	1·10 ⁻¹⁴⁹	sp P09795
	2. M. <i>EcoO109I</i>	rGGnCCy	<i>Escherichia</i>	4·10 ⁻⁶⁴	gb AAF06965
R. <i>AvaII</i>	1. M. <i>SinI</i>	GGwCC	<i>Salmonella</i>	4·10 ⁻⁶⁶	sp P09796
	2. M. <i>Eco47I</i>	GGwCC	<i>Escherichia</i>	7·10 ⁻⁶⁶	sp P50194
	Next match			>0.2	
Protein sequences similar to putative <i>AvaIII</i> R/M proteins					
M. <i>AvaIII</i>	1. ?	?	<i>Escherichia</i>	1·10 ⁻⁷⁶	sp P28638
	2. M. <i>HpaI</i>	GTAA ⁶ C	<i>Haemophilus</i>	3·10 ⁻³¹	sp P29538
	3. M. <i>BglIII</i>	AGATC ⁴ T	<i>Bacillus</i>	1·10 ⁻²⁴	gb AAC45061
R. <i>AvaIII</i> ?	No match			> 3	
Protein sequences similar to <i>AvaIV</i> R/M proteins					
M. <i>AvaIV</i>	1. M. <i>BlpI</i>	GCTnAGC	<i>Bacillus</i>	1·10 ⁻¹⁰²	NEB database ^e
	2. M. <i>DdeI</i>	C ⁵ TnAG	<i>Desulfovibrio</i>	1·10 ⁻⁸⁴	sp P05302
	3. M. <i>Bpu10IC1</i>	GC ⁵ TnAGG	<i>Bacillus</i>	1·10 ⁻⁷⁹	gb CAA74996
R. <i>AvaIV</i> (reconstructed) ^f	1. R. <i>BlpI</i>	GCTnAGC	<i>Bacillus</i>	2·10 ⁻⁹⁸	NEB database ^e
	2. R. <i>Bpu10Iβ</i>	GCTnAGG	<i>Bacillus</i>	6·10 ⁻¹⁹	gb CAA74999
	3. R. <i>Bpu10Iα</i>	GCTnAGG	<i>Bacillus</i>	9·10 ⁻⁷	gb CAA74998
	Next match			> 1	
Protein sequences similar to <i>PstI</i> R/M-like proteins					
Orf362.31394R (M-portion)	1. M. <i>PstI</i>	CTGCA ⁶ G	<i>Providencia</i>	5·10 ⁻⁵³	sp P00474
	2. Orf-F60	?	<i>Aeromonas</i>	2·10 ⁻⁵²	gb AAF45040
	3. M. <i>BsuBI</i>	CTGCA ⁶ G	<i>Bacillus</i>	7·10 ⁻⁵²	sp P33563
Orf362.31394R (M-portion)	1. R. <i>BsuBI</i>	CTGCAG	<i>Bacillus</i>	6·10 ⁻⁸⁹	sp P33562
	2. R. <i>PstI</i>	CTGCAG	<i>Providencia</i>	2·10 ⁻⁷²	sp P00640
	3. R. <i>Rle39BI</i>	CTGCAG	<i>Rhizobium</i>	9·10 ⁻²⁰	gb CAA67875
	4. R. <i>XphI</i>	CTGCAG	<i>Xanthomonas</i>	5·10 ⁻¹⁴	gb AAF22367
Next match				> 2	

^aProtein sequences obtained by BLAST search (26) of the combined GenBank, PDB, SwissProt, PIR, and PRF databases (unless otherwise noted), last accessed June 2000, ranked by score.

^bRecognition sequences of MTases are given along with the position, if known, where they methylate: A⁶, N6-methyladenine; C⁵, C5-methylcytosine.

^cScore represents expected number of proteins at least as similar to target within the combined databases.

^dgb, GenBank; sp, SwissProt

^eSequence graciously provided by Geoffrey Wilson (New England Biolabs).

^fReconstructed by removing insertion sequence from Orf323B.1084'R and Orf339B.706 and combining the results.

In addition, a predicted 5mC MTase was found that is very similar to M.*XorII*, which methylates CGATCG, and to SynMI, a cyanobacterial enzyme (37) with the same specificity. The similarity is particularly striking in the variable TRD region: 56% and 46% amino acid identity, respectively. Since *PvuI*, a CGATCG-specific endonuclease sensitive to cytosine methylation, fails to cut *Anabaena* DNA (38), even though recognition sites can be found in the genome with a frequency of one in about 1100 bp (J. Elhai, unpublished data), we concluded that the predicted protein actually functions in the cyanobacterium and on this basis named the gene that encodes it *dmtC*.

The gene is preceded by an ORF capable of encoding a 188-amino acid protein. This upstream gene, and close variants of it, occurs at least 11 times in the *Anabaena* genome, 7 times in the genome of *N. punctiforme* and 6 times in the genome of *Synechocystis* PCC 6803 (data not shown). Its putative product shows no obvious similarity to noncyanobacterial protein. The gene family has been termed SPD, after a three-amino acid sequence found in almost all variants. Apart from this ORF, there is no possible gene 1117 bp upstream or 2000 bp downstream of *dmtC* capable of encoding a protein greater than 95 amino acids that does not have a function readily recognizable from the sequence.

Analysis of genes capable of encoding Type II RM systems

The presence of three restriction activities in *Anabaena* (8) implies the existence of corresponding MTases, and three likely genes were found. One gene shows 100% nucleotide identity with the previously described gene encoding M.AvaI (accession X98339) cloned from *Anabaena* PCC 7118, and the adjacent gene is identical to R.AvaI (which also recognizes CyCGrG). Likewise, a second gene pair initially identified for the similarity of their predicted products to the 5mC MTase M.SinI (P09795) and its corresponding restriction enzyme R.SinI (an isoschizomer of R.AvaII) have 100% amino acid identity to the proteins predicted from the cloned M.AvaII and R.AvaII genes from *Anabaena* PCC 7118 (G.G. Wilson and K.D. Lunnen, personal communication). The enzymes from *Anabaena* PCC 7120 were originally given names (e.g. M.Asp(7120)I) to distinguish them from the corresponding enzymes from *Anabaena* PCC 7118 (39,40). Since the proteins are evidently identical, those names may now be abandoned. In fact, the near identity between the two strains over all known nucleotide sequences (4 differences in 4783 bases; data not shown) compels us to accept the suggestion of Rippka and Herdman (41) that the two strains are derived from the same natural isolate.

Detection of the gene encoding M.Ava III (which recognizes ATGCAT) was more problematic, since no sequence has been reported of a MTase with the same target specificity. It seemed likely that the MTase was encoded by Orf337.35147, evidently an N6mA or N4mC MTase (2), since that is the only ORF in the available *Anabaena* genome sequence predicted to encode a MTase that remained unaccounted for. This ORF and the one downstream from it was cloned (Figure 1), and the resulting plasmid gained resistance to digestion by *Nsi* I, which shares the same recognition sequence as *Ava* III. On this basis, we judged that Orf337.35147 encodes M.Ava III.

The adjacent ORF, Orf337.36048, most likely encodes R.Ava III. The failure to find a match between that ORF and known restriction enzymes, while not very informative, is expected, since restriction enzymes seldom show similarity to proteins that recognize different sequences (1; also compare with R.Ava I and R.Ava II in Table 2).

Two other gene pairs are similar to those encoding Type II RM enzymes. Orf323B.3213 is capable of encoding a 5mC DNA MTase highly similar to M.Blp I, while the adjacent gene is highly similar to R.Blp I (Table 2), except for an intervening insertion sequence near the 5' end of the gene (Figure 1). Since neither *Blp*I (data not shown) nor its isoschizomer *Esp*I (38) cut *Anabaena* DNA, despite the presence of at least 16 sites (J. Elhai unpublished data), it is likely that such sites are methylated owing to the action of the M.*Blp*I-like gene product. On that basis, the putative MTase has been named M.Ava IV, though the cognate restriction enzyme is surely inactive.

Orf362.31394R shows strong similarity (Table 2) to M.*Pst* I and R.*Pst* I fused together (also noted independently by Bill Buikema, University of Chicago). The gene is interrupted by a stop codon appearing between a region encoding Motif VIII of the MTase and its TRD, so it is not surprising that *Anabaena* does not methylate at *Pst* I sites (38).

None of the five predicted RM proteins show significant sequence similarity to proteins potentially encoded by *N. punctiforme* genes, beyond the general similarity shown by all members of their particular classes; nor do the genes that

encode the proteins show similarity to *N. punctiforme* DNA (Figure 1). In two cases, M/R.AvaII and M/R.AvaIII, the genes encoding the RM system replace a significant amount of *N. punctiforme* DNA: about 740 bp and 5720 bp, respectively. The novel *Anabaena* sequences are not flanked by discernible direct or indirect repeats that might be indicative of the ends of insertion sequences.

Analysis of genes capable of encoding Type I MTases

Five regions of the *Anabaena* genome were found containing genes similar to those encoding elements of Type I RM systems (Table 3). In all cases, one or more genes were defective, as judge by frameshifts or insertion sequences within the gene. Two regions bear genes related to Type IB RM systems. The two sets of genes are nearly identical to each other, except for the specificity subunit (HsdS) and in both cases defects in that subunit should render restriction and modification inactive. The two modification (HsdM) and restriction (HsdR) proteins show striking resemblance to those of the *EcoA* and *EcoE* systems of *E. coli* strains T15 and A58. In fact, these *E. coli* systems are more similar to the Type IB systems from *Anabaena* than to those found thus far in any other organism, including the closely related enteric bacterium *Salmonella typhimurium* (data not shown).

The three other Type I systems of *Anabaena*, two of Type IC and one of Type ID, follow the same pattern: the HsdM and HsdR proteins are very similar to known Type I enzymes, while the HsdS protein (when present) is less similar. In contrast, the *Anabaena* proteins show much less resemblance to Type I RM proteins from the cyanobacterium *N. punctiforme* (Table 3). None of the systems have restriction capability, by reason of a defective HsdR subunit, defective HsdS subunit, or both. There are no obvious defects in the genes encoding the Type ID MTase and specificity subunits, so that system may be functional.

DISCUSSION

Nine genes that encode MTases with demonstrated or deduced activity have been identified from the cyanobacterium *Anabaena* (Table 4). Of these, four have characteristics associated with solitary MTases, and a fifth may also fall into that category. While this number is considerably higher than that established thus far for any other bacterium, solitary MTases may be common amongst bacteria.

Definition of solitary MTases

Solitary MTases are defined primarily by a cellular role distinct from RM. The demonstration of such a function is no trivial task, but in the absence of proven function, it is possible to recognize solitary MTases by other characteristics. The failure to detect a corresponding restriction endonuclease is supportive evidence but by no means definitive, only in part because of the impossibility of proving that an activity is not present in a strain. It is more telling that a gene encoding a MTase has no nearby gene that can conceivably encode an active endonuclease. Here too, however, caution must be exercised. A MTase not long ago associated with a restriction endonuclease may momentarily exist alone, before selective pressures swallow it as well.

What sets solitary MTases apart from transiently orphaned MTases is stability over evolutionary time, conferred by the selective advantage its function confers on their hosts. This is seen in the two cases, Dam and CcrM, where independent

Table 3. Proteins similar to those of putative Type I R/M systems of *Anabaena*

Target ^a	Protein ^b	Source	Score ^c	Accession no. ^d
Sequences similar to orf308A Type IB R/M proteins				
HsdM (271 amino acid,C-terminus)	- Orf311A	<i>Anabaena</i>	1·10 ⁻¹⁵³	genome site ^e
	1. EcoE HsdM	<i>Escherichia</i>	2·10 ⁻⁹⁴	pir I41293
	2. EcoA HsdM	<i>Escherichia</i>	2·10 ⁻⁹⁴	pir A47200
	3. StySKI HsdM	<i>Salmonella</i>	9·10 ⁻⁹³	gb CAA71895
	- Orf658	<i>Nostoc</i>	6·10 ⁻²³	genome site ^f
	- Orf588	<i>Nostoc</i>	1·10 ⁻⁴	genome site ^f
HsdS (550 amino acid)	- Orf311A	<i>Anabaena</i>	3·10 ⁻⁹⁰	genome site ^e
	1. CfrA	<i>Citrobacter</i>	3·10 ⁻⁵⁸	pir S06097
	2. EcoE	<i>Escherichia</i>	2·10 ⁻⁴⁷	sp P19705
	3. EcoA	<i>Escherichia</i>	2·10 ⁻⁴⁶	sp P19704
	- Orf658	<i>Nostoc</i>	4·10 ⁻⁸	genome site ^f
HsdR (430 amino acid,C-terminus)	- Orf308B	<i>Anabaena</i>	<10 ⁻²⁰⁰	genome site ^e
	1. EcoA	<i>Escherichia</i>	1·10 ⁻¹⁴¹	pir I41291
	2. EcoE	<i>Escherichia</i>	1·10 ⁻¹⁴⁰	pir I41292
	3. StySKI	<i>Salmonella</i>	2·10 ⁻⁵³	gb CAA71894
	- Orf658	<i>Nostoc</i>	7·10 ⁻⁸	genome site ^f
Sequences similar to Orf308B/311A Type IB R/M proteins				
HsdM (296 amino acid, C-terminus)	1. EcoE HsdM	<i>Escherichia</i>	1·10 ⁻¹⁰¹	pir I41293
	2. EcoA HsdM	<i>Escherichia</i>	1·10 ⁻¹⁰¹	pir A47200
	3. StySKI HsdM	<i>Salmonella</i>	1·10 ⁻¹⁰⁰	gb CAA71895
HsdS (542 amino acid)	1. EcoA	<i>Escherichia</i>	4·10 ⁻⁸¹	sp P19704
	2. StySKI	<i>Salmonella</i>	1·10 ⁻⁴⁰	gb CAA71896
	3. CfrA	<i>Citrobacter</i>	7·10 ⁻³⁵	gb CAA35604
	4. EcoE	<i>Escherichia</i>	8·10 ⁻³⁰	sp P19705
HsdR (776 amino acid)	1. EcoA	<i>Escherichia</i>	<10 ⁻²⁰⁰	pir I41291
	2. EcoE	<i>Escherichia</i>	<10 ⁻²⁰⁰	pir I41292
	3. StySKI	<i>Salmonella</i>	3·10 ⁻⁵³	gb CAA71894
Protein sequences similar to Orf260 Type IC R/M protein				
HsdM (537 amino acid)	1. <i>LldI</i>	<i>Lactococcus</i>	1·10 ⁻¹²⁰	pir T09460
	7. HsdM	<i>Helicobacter</i>	1·10 ⁻⁹³	pir E71886
	- Orf339A	<i>Anabaena</i>	1·10 ⁻⁶⁷	genome site ^e
	- Orf376B	<i>Anabaena</i>	4·10 ⁻⁴⁷	genome site ^e
	- Orf641	<i>Nostoc</i>	4·10 ⁻²¹	genome site ^f
HsdR (933 amino acid)	1. HsdR	<i>Helicobacter</i>	1·10 ⁻¹²⁷	pir B71890
	- Orf376B	<i>Anabaena</i>	1·10 ⁻²⁵	genome site ^e
	- Orf641	<i>Nostoc</i>	3·10 ⁻¹⁰	genome site ^f
Sequences similar to Orf339A Type IC R/M protein				
HsdM (803 amino acid)	1. HsdM	<i>Helicobacter</i>	1·10 ⁻¹⁸⁰	pir C71810
	- Orf376B	<i>Anabaena</i>	6·10 ⁻⁴³	genome site ^e
	- Orf641	<i>Nostoc</i>	8·10 ⁻¹⁹	genome site ^f
HsdS (425 amino acid)	1. HsdS	<i>Methanococcus</i>	8·10 ⁻²²	pir B64316
	- Orf641	<i>Nostoc</i>	3·10 ⁻³	genome site ^f
HsdR (179 amino acid, N-terminus)	1. HsdR(179)	<i>Klebsiella</i>	6·10 ⁻⁶⁵	pir T30818
	- Orf376B	<i>Anabaena</i>	2·10 ⁻⁴	genome site ^e
Sequences similar to Orf376B Type ID R/M protein				
HsdM (527 amino acid)	1. Cj1553c	<i>Campylobacter</i>	1·10 ⁻¹⁷¹	gb CAB73544
	2. HsdM	<i>Pasteurella</i>	1·10 ⁻¹³⁵	gb AAC44666
	- Orf641	<i>Nostoc</i>	4·10 ⁻²⁰	genome site ^f
HsdS (390 amino acid)	1. Cj1551c	<i>Campylobacter</i>	6·10 ⁻²⁹	gb CAB73967
	2. MJEL41	<i>Methanobacter</i>	5·10 ⁻¹⁵	pir H64514
HsdR (1011 amino acid)	1. Cj1549c	<i>Campylobacter</i>	<10 ⁻²⁰⁰	gb CAB73965
	2. HI1285	<i>Haemophilus</i>	<10 ⁻²⁰⁰	pir F64114
	3. HsdR	<i>Pasteurella</i>	1·10 ⁻¹⁹⁸	pir JC5216
	- Orf641	<i>Nostoc</i>	6·10 ⁻⁸	genome site ^f

function of an MTase has been demonstrated (2,3,42-44). Modification of GATC sites is found generally in enteric bacteria and closely related gamma proteobacteria (Figure 5A), and the lineage of the responsible MTase closely follows that of 16S rRNA (Figure 5B). In contrast, the ability to restrict GATC sites is sporadic, and the lineage of restriction-associated GATC MTases shows a heavy influence of lateral DNA transfer (45). The same phenomenon is seen with CcrM and other GAnTC-specific MTases (Figs. 5C and 5D). Just as the lineage of GATC-specific MTases is coherent within a subset of the gamma proteobacteria, so is the lineage of GAnTC-specific MTases within the alpha proteobacteria.

One might have expected that the presence of a MTase would make it easier for the host to acquire a RM system that recognizes the same target sequence, but just the opposite is the case with Dam and CcrM. It is striking that while methylation at GATC is perhaps universal amongst enteric bacteria, there is no known instance of a restriction enzyme carried by a member of this group that cuts at GATC (Figure 5A). Similarly, GAnTC-specific restriction enzymes are absent from the alpha proteobacteria (Figure 5C), though the rarity of such enzymes in general makes the observation not particularly informative. Perhaps the functional constraints placed on Dam- and CcrM-methylation do not permit the total modification of DNA required by the presence of a restriction system. It should be noted that restriction enzymes that target the recognition sequence (CCwGG) of Dcm are very common in enteric bacteria.

In recognition of the needs of two scientific communities, those that study the structures of MTases and restriction enzymes and those that study bacterial physiology, we have given two names to solitary MTases. Like other MTases, they have been given names of the form *M.AvaX* (encoded by *avaMX*). In addition, to emphasize their cellular roles distinct from RM, they have been given names of the form *DmtX* (encoded by *dmtX*). A new name was chosen for its generality. It seemed inappropriate to use CcrM for any enzyme except that from *Caulobacter crescentus*, and Dam fits only deoxyadenine-specific MTases.

Multiple solitary MTases in *Anabaena*

On the basis of the criteria set forth above, *Anabaena* appears to have four solitary MTases, denoted as DmtA, DmtB, DmtC, and DmtD. None of the genes encoding these MTases lies near any plausible candidate to encode a corresponding restriction endonuclease. Restriction activities have not been found in *Anabaena* that recognize any of the four target sequences (8,46). Indeed, in two cases, *dmtB* and *dmtD*, null mutants are viable (A.V.Matveyev, K.T.Young, and J.Elhai, unpublished observations), indicating that their activities are not required to protect the host against self-digestion by a cognate restriction enzyme. Supportive of the absence of such restriction enzymes in *Anabaena* is the observation that the efficiency of transfer from *E. coli* to *Anabaena* of plasmids bearing recognition sites for unmethylated DmtB, DmtC, and DmtD is high. In contrast,

the presence of unmethylated sites recognized by R.AvaI, R.AvaII, or R.AvaIII drastically reduces the efficiency of conjugal transfer (8).

Another important criterion is evolutionary stability. GATC methylation (as specified by DmtA) is widely distributed and perhaps universal amongst cyanobacteria (5,6; Figure 5A), and yet none of the 162 characterized cyanobacterial restriction enzymes (27) have GATC specificity. The other cases are less clear. GGCC methylation (as specified by DmtB) is not universal (6,7), though it is widespread, and GGCC-specific restriction enzymes have been found in six cyanobacterial genera, including *Anabaena* (27). Little is known about the prevalence of CGATCG (DmtC) or rCCGGY (DmtD) methylation amongst cyanobacteria. Restriction enzymes recognizing the former sequence have been described in two cyanobacterial genera (27), but those recognizing the latter have not yet been found.

The GATC-specific *M.AvaV* poses an unusual case in that it, like RM systems (see below), appears to be a relatively recent arrival to *Anabaena*. On the other hand, there is no plausible ORF in the vicinity that might encode a corresponding endonuclease. Conceivably, an intact RM system was recently transferred to *Anabaena*, and the gene encoding the restriction enzyme has deteriorated. However, we have found that mutation of *avaMV* appears to alter the frequency at which cells differentiate (A.V. Matveyev and J.Elhai, unpublished results), so the MTase has evidently acquired a physiological role during its brief stay in *Anabaena*, even though the cyanobacterium already possesses DmtA, another GATC-specific MTase.

Unusual structure of Type II MTase *M.AvaV*

The 5mC MTases of *Anabaena* (DmtD, DmtC, *M.AvaII*, and *M.AvaIV*) fall readily within the definitions of established groups, as do four of the N6mA/N4mC MTases (DmtB, DmtA, *M.AvaI*, and *M.AvaIII*). One MTase, *M.AvaV*, does not and deserves special mention.

From the order of the few motifs that can be identified and the limited sequence similarity (Figure 2), *M.AvaV* appears to be a distant relative of group β N6mA MTases. Presuming the identifications of Motif X (and possibly I) to be valid, the order of motifs corresponds to that found in group β MTases (IV-V-VI-VII-VIII-TRD-X-I-II-III; 10). The consensus sequences for the three identified regions are closer to the corresponding sequences for motifs from group β than to those of any other group. However, the differences between *M.AvaV* (and similar proteins) and group β MTases are also striking. The two groups show no significant similarity, according to BLAST comparisons, and *M.AvaV* and two of the three similar bacterial proteins are smaller than any of the 43 DNA MTases used to establish the conserved motifs amongst N4mC/N6mA MTases (10). We propose the creation of a new subgroup, β_2 , to accommodate *M.AvaV*-like MTases, but only a three-dimensional structure determination can resolve whether these proteins share the common architecture

Table 3. *continued*

^aProtein sequence from *Anabaena* used in BLAST search (26) of the combined GenBank, PDB, SwissProt, PIR, and PRF databases, followed by length of sequence in amino acid residues. Partial sequences were used when full sequences were unavailable.

^bProteins found by BLAST search, ranked by score. Scores of proteins not found in combined databases were obtained by a pairwise BLAST comparison. Proteins in bold are from cyanobacteria.

^cScore represents expected number of proteins at least as similar to target within the combined databases.

^dgb, GenBank; sp, SwissProt; pir, Protein Information Resource

^eSequence obtained from the *Anabaena* web site (<http://www.kazusa.or.jp/cyano/anabaena/>), last accessed June 2000.

^fSequences obtained from the *N. punctiforme* web site (http://spider.jgi-psf.org/JGI_microbial/html/nostoc_homepage.html), last accessed June 2000.

Table 4. DNA Methyltransferases in *Anabaena* PCC 7120

Methyltransferase	Locus ^a	Specificity ^b	Type ^c	Found in ^d	Restriction Enzyme ^e	Comments
DNA methyltransferases of demonstrated functionality						
M.Ava I	Orf308a.5063	CyCGrG (<i>AvaI</i>)	N4mC(α)	—	Orf308A.6511	= RM.Ava I
M.Ava II	Orf333.17488	GGwCC (<i>AvaII</i>)	5mC	—	Orf333.19647R	= RM.Ava II
M.Ava III	Orf337.35114	ATGCAT (<i>NsiIII</i>)	N6mA(β)	—	Orf337.36048?	= RM.Ava III?
M.Ava IV	Orf323b.2328R	GCTnAGC (<i>BlpI</i>)?	5mC	—	(<i>Orf323B.1084'R/339B.706</i>)	~M. <i>BlpI</i> ; R.Ava IV interrupted by IS
M.AvaV	AF220506	GATC (<i>Dam</i>)	N6mA(β_2)	—	none	Solitary?
DmtA/M.AvaVI	Orf304.22874	GATC (<i>Dam</i>)	N6mA(α)	NS	none	Solitary
DmtB/M.AvaVII	AF220507	GGCC (<i>HaeIII</i>)	N4mC(α)	NS	none	Solitary
DmtC/M.AvaVIII	Orf350.1117	CGATCG (<i>PvuI</i>)	5mC	NS	none	Solitary
DmtD/M.AvaIX	AF220508	rCCGGy (<i>Cfr10I</i>)	5mC	N	none	Solitary
Putative or defective DNA methyltransferases						
Defective?	<i>Orf362.31394R</i>	CTGCAG (<i>PstI</i>)?	N6mA(γ)	—	<i>Orf362.31394R</i>	Stop before TRD; fused to R. <i>PstI</i>
Defective?	<i>Orf275.9656R</i>	?	N4mC(β)	—	none	Frame shift in TRD region
Defective?	<i>Orf335.3576</i>	?	5mC	—	none	Motifs I, IV, VI, and VIII only
Defective?	<i>Orf351a.9898</i>	?	5mC	—	none	Motifs I, IV, VI, and VIII (=Orf335.3576)
Defective?	Orf308a.17412R	?	Type IB	—	<i>Orf308A.15244R</i> (HsdS)	IS in <i>hsdS</i>
					Orf308A.19733R (HsdR)	
Defective?	<i>Orf308b.11683'/311a.30178</i>	?	Type IB	—	<i>Orf311A.29298</i> (HsdS)	Stop at 5' end of <i>hsdM</i> ; frameshift in <i>hsdS</i>
Defective?	Orf260.11347R	?	Type IC	—	Orf308B.8327 (HsdR)	
					<i>Orf260.4629R</i> (HsdR)	Frameshift, no start in <i>hsdR</i> ; no nearby <i>hsdS</i>
Defective?	<i>Orf339a.24489R</i>	?	Type IC	—	Orf339A.22077R (HsdS)	Frameshift in <i>hsdM</i> ;
					<i>Orf339A.20008R'</i> (HsdR)	IS in <i>hsdR</i>
M.Ava V?	Orf376b.55687	?	Type ID	—	Orf376B.58994 (HsdS)	frameshift in <i>hsdR</i> at C-terminus
					<i>Orf376B.60879</i> (HsdR)	

^aLocus given is either the accession number or the contig deposited in the *Anabaena* PCC 7120 web site (<http://www.kazusa.or.jp/cyano/anabaena/>). The number after "Orf" refers to the number of the contig and the number after the period refers to the position within the contig of the start codon. Numbers followed by "R" indicate that the ORF lies on the strand opposite to that given in the contig. Numbers followed by an apostrophe indicate that the contig stops in the middle of the gene. Numbers preceded by an apostrophe indicate that the contig begins in the middle of the gene. Italicized names refer to ORFs with obvious defects (stop codons, frameshifts, or insertion sequences within the gene).

^bSpecificities were determined either as described in the text (M.AvaV, DmtB, and DmtD) or by comparison of the TRD region with MTases of known specificity. A well-known R/M system or solitary MTase with the same specificity is given in parentheses.

^cThe subtype of MTase is according to Malone *et al.* (10) for Type II MTases (except for M.AvaV) or Murray *et al.* (17) for Type I MTases.

^d"N" or "S" is given if the MTase is substantially similar to one found in *Nostoc punctiforme* ATCC 29133 or *Synechocystis* PCC 6803, respectively. A MTase is judged to be substantially similar if it exhibits greater similarity than most MTases of the same type or subtype.

^eNearby gene encoding putative restriction enzyme or (in the case of Type I enzymes) specificity subunit (HsdR) or restriction subunit (HsdR). Naming conventions are the same as described in note a.

of previously characterized MTases despite unusual primary sequences or instead represent a second mechanism found by nature to methylate DNA.

There are other MTases besides M.AvaV that fit poorly into the conventional classification scheme. GATC-specific adenine MTases from *Vibrio cholerae*, *E. coli* phage T1 and *H. influenzae* phage P1 exhibit no obvious similarity to other MTases motif IV (47,48).

Evolution

We have attempted to discern the evolutionary history of MTases of *Anabaena* by comparing their sequences to those of other bacteria and in particular to those of the closely related cyanobacterium *N. punctiforme* and the distantly related cyanobacterium *Synechocystis* PCC 6803 (49). From this analysis, the generality emerges that solitary MTases have ancient antecedents in the cyanobacterial lineage. DmtD is common to both *Anabaena* and *N. punctiforme*. DmtB, DmtA, and DmtC homologues are found in *Synechocystis* as well, and probably throughout the cyanobacterial radiation (the DmtB-like protein *sll0729* of *Synechocystis* is no doubt the GGCC-specific MTase postulated by Scharnagl *et al.*; 37).

In contrast, RM systems are transient visitors to *Anabaena*. The active RM systems of *Anabaena* are absent even in

N. punctiforme, and most of the genes show striking resemblances to those that exist in organisms outside the cyanobacteria (Tables 2 and 3). The systems are not long lived, however: the *Anabaena* genome is a graveyard of R/M genes in various states of decomposition, as is the genome of *Helicobacter pylori* J99 (50). A survey of such enzymes must therefore represent a snapshot that would differ markedly from one taken at a different moment in evolutionary time.

These results are consistent with a genome that is showered with DNA derived from diverse organisms. Most foreign genes presumably disappear without a trace, but there remain a few well documented examples of horizontal transfer into the cyanobacterial lineage (51-53). The great divergence between cyanobacteria and other eubacteria (54) makes it unlikely that genes whose products interact with host machinery would function in their new host. Restriction modification systems, which can operate independently, seem well suited to proliferate by horizontal gene transfer (55,56) and either confer on their new hosts a selective advantage, e.g. defense against phage attack (57), or act as molecular parasites (56).

The alternative hypothesis that inactive RM systems represent an internal pool to be drawn upon for defense against foreign DNA (50) is less likely. None of the genes

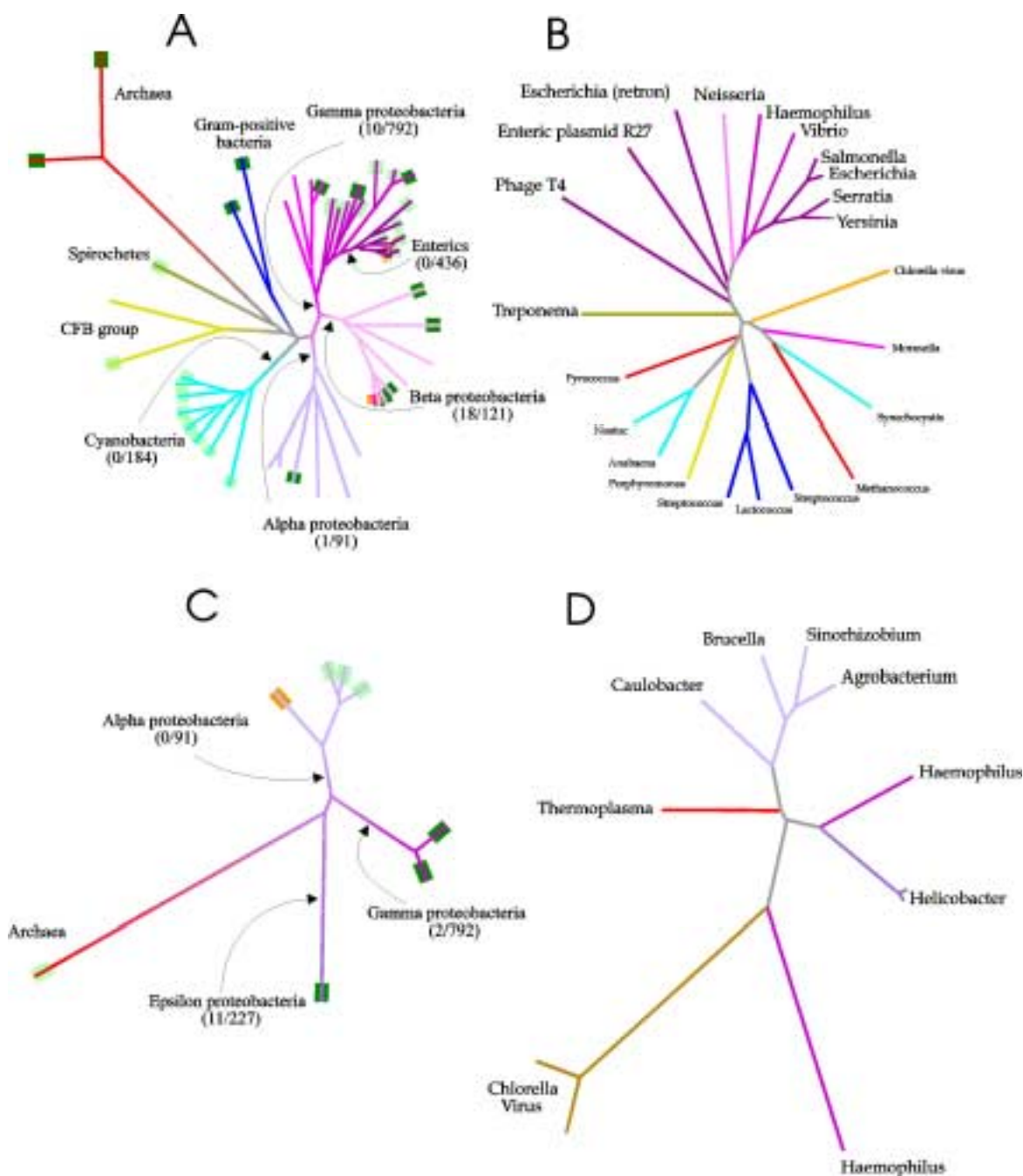


Figure 5. Relationship amongst enzymes functionally similar to known solitary methyltransferases, Dam and CcrM. Aligned 16s rRNA sequences were obtained from the Ribosomal Database Project (<http://www.cme.msu.edu/RDP/html/index.html>), and only the part of the alignment common to all was used in constructing the tree. (A) Dendrogram of 16s rRNA sequences from organisms that have been tested for GATC-specific adenine methylation (5) or that possess sequenced GATC-specific MTases of type N6mA(α). The number of restriction endonucleases known to recognize GATC is given followed by the total number of known endonucleases, for all bacteria beyond the indicated branch point. Lines are colored according to taxonomic groups. Tags at the end of branches are green if GATC-specific methylation has been demonstrated, dark green if the strain possesses a GATC-specific MTase, and orange if the strain possesses a proven solitary GATC-specific MTase. (B) Dendrogram of all GATC-specific MTases of type N6mA(α) for which sequences are available. The lines are colored as in (A). The source of the sequences are given in the legend to Figure 4, except for Dam from *Yersinia pseudotuberculosis* (AAG23175), Dam from *Serratia marcescens* (P45454), Dam from *Salmonella typhimurium* (P55893), gene from *Vibrio cholerae* O395 (AAG23174), gene from *Haemophilus influenzae* Rf (P44431), Dam from *Neisseria meningitidis* BF13 (AAD34292), gene from *Escherichia coli* retron Ec67 (P21311), and gene from enteric plasmid R27 (AAF69879). (C) Dendrogram of 16s rRNA sequences from organisms that possess sequenced GAnTC-specific MTases. The conventions are analogous to those described in (A). (D) Dendrogram of all GAnTC-specific MTases for which sequences are available. The lines are colored as in (C). The source of the sequences are (reading counterclockwise from Agrobacterium): gene from *Agrobacterium tumefaciens* C58 (Brad Goodner, personal communication), gene from *Sinorhizobium meliloti* 1021 (AAB71350), *M.BabI* (O30570), *M.BabI* (Q45971), CcrM (Q45971), gene from *Thermoplasma acidophilum* DSM1728 (CAC12293), *M.CviBI* (Q01511), *M.CviQVI* (AAC03126), *M.HhaII* (P00473), *M.HpyAIV* (H64688), *M.Hpy99IX* (F71827), and *M.HinI* (P20590).

encoding the four active RM systems in *H. pylori* J99 are found in the closely related strain *Helicobacter pylori* 26695, although the two strains share several inactive systems (<http://www.tigr.org/> and unpublished observations). The life history for RM systems seems more consistent with a one way path from acquisition to decay than with internal genetic recycling.

The commonality of certain R/M systems in cyanobacteria related to *Anabaena* supports the idea that DNA exchange occurs more frequently among themselves than with the bacterial world at large. Of the 40 strains of *Anabaena* and other heterocystous cyanobacteria (Sections IV and V per Rippka et al, ref.49), 11 have *Ava* I activity, 14 have *Ava* II activity, and 6 have both (27). In contrast, *Ava* I and *Ava* II are relatively rare amongst other cyanobacteria, and the one characterized *Ava* I-like system, *Aqu* I (58), found in a distantly related cyanobacterium is part of a 5mC system, with no similarity to the enzymes of *Anabaena*. There are several other restriction enzymes that recur amongst heterocystous cyanobacteria. A self-consistent phylogeny accounting for these occurrences without recourse to multiple horizontal transfers requires an ancestor with more than six R/M systems and bears little resemblance to the currently accepted taxonomy (J.Elhai, unpublished observations). We conclude, then, that horizontal transfer amongst heterocystous cyanobacteria has been responsible on several occasions for the spread of R/M systems, perhaps by transduction by phage with a broad host range amongst heterocystous cyanobacteria (59) or by conjugation (60). A perusal of known RM systems organized by their hosts indicates that the spread of certain systems within taxonomic groups is common. One may view this either as infections of taxa by selfish DNA or as sharing of valuable resources amongst relatives.

The transient presence of R/M systems commonly exchanged among related cyanobacteria explains the observation that *Anabaena* strains (61; J.Elhai, unpublished observations) and *E. coli* K-12 (62) are deficient in sites recognized by restriction enzymes of related strains. This would be the expected result if organisms selected against recognition sites for their own restriction enzymes and if there were rampant genetic exchange of RM systems amongst similar strains (62-64).

The evidence suggests an unusual origin for DmtB (recognizing GGCC). In general, N4mC MTases form a coherent phylogenetic group, distinct from N6mA MTases (65). DmtB, however, lies within the class N6mA radiance, particularly close to MTases with GATC-specificity. In general, DNA MTases that modify the same target sequence at the same site share broad sequence similarity (66), despite the taxonomic diversity of their hosts (e.g. the archaeal *M.Mja*III and eukaryotic virus-derived *M.Cvi*AI mixed amongst the eubacterial GATC-specific MTases in Figure 4). The phylogenetic position of DmtB suggests that it evolved from the line leading to DmtA and other cyanobacterial GATC-dependent N6mA MTases. DmtB adds credence to the hypothesis of Bujnicki and Radlinska (65) that N4mC MTases may have occasionally originated from N6mA precursors.

Function

The present work was inspired by a desire to understand the function of solitary MTases in *Anabaena*. Pertinent results along those lines will be published elsewhere, but in brief, DmtA appears to be essential for the viability of *Anabaena* under standard laboratory conditions, DmtB lies in a region

intimately involved in the cellular differentiation evoked by nitrogen starvation, and loss of DmtD causes a slight defect in growth under laboratory conditions (A.V.Matveyev, J.Rumble, J.Elhai, unpublished results). The functional importance of the GATC sequences modified by M.AvaV and DmtA MTases is further indicated by the absence of such sites in some phages that infect *Anabaena* (38)

The target sequence (CGATCG) of DmtC is provocative, because it is contained within a highly iterated sequence (GCGATCG; HIP1) found in many cyanobacteria (67) and also envelops the target sequence (GATC) of DmtA and M.AvaV. Loss of the DmtC homolog SynMI of *Synechocystis* leads to poor growth, especially under conditions most favorable to rapid doubling (37).

It is interesting that two DNA MTases, M.AvaV and DmtA, recognize the same sequence. There are several instances known of restriction enzymes accompanied by two MTases with the same target sequence. In one such case, *Streptococcus pneumoniae* possesses two GATC-specific MTases, whose genes are adjacent to that encoding the GATC-specific restriction endonuclease *Dpn* II (68). As with M.AvaV and DmtA, the two MTases of *Streptococcus* are of different groups (N6mA and N6mA). The presence of the two MTases may be explained by their different biochemical properties. Furthermore, a strain of *E. coli* has been described that carries a second functional GATC-specific MTase besides Dam (69), and *Vibrio cholerae* may also carry an unusual GATC-specific enzyme (48) in addition to the conventional Dam-like MTase.

In short, very little is yet known about the function of cyanobacterial solitary MTases. In pursuing their functions, we hope to extend our understanding of the roles played by solitary MTases beyond what is known from Dam and CcrM.

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