Cloning, Sequencing, and Expression of the Cold-Inducible 
\textit{hutU} Gene from the Antarctic Psychrotrophic Bacterium 
\textit{Pseudomonas syringae} 

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A promoter-fusion study with a Tn5-based promoter probe vector had earlier found that the \textit{hutU} gene which encodes the enzyme urocanase for the histidine utilization pathway is upregulated at a lower temperature (4°C) in the Antarctic psychrophilic bacterium \textit{Pseudomonas syringae}. To examine the characteristics of the urocanase gene and its promoter elements from the psychrotroph, the complete \textit{hutU} and its upstream region from \textit{P. syringae} were cloned, sequenced, and analyzed in the present study. Northern blot and primer extension analyses suggested that the \textit{hutU} gene is inducible upon a downshift of temperature (22°C to 4°C) and that there is more than one transcription initiation site. One of the initiation sites was specific to the cells grown at 4°C, which was different from the common initiation sites observed at both 4 and 22°C. Although no typical promoter consensus sequences were observed in the flanking region of the transcription initiation sites, there was a characteristic CAAAA sequence at the −10 position of the promoters. Additionally, the location of the transcription and translation initiation sites suggested that the \textit{hutU} mRNA contains a long 5′-untranslated region, a characteristic feature of many cold-inducible genes of mesophilic bacteria. A comparison of deduced amino acid sequences of urocanase from various bacteria, including the mesophilic and psychrotrophic \textit{Pseudomonas} spp., suggests that there is a high degree of similarity between the enzymes. The enzyme sequence contains a signature motif (GXGX\textsubscript{10}G) of the Rossmann fold for dinucleotide (NAD\textsuperscript{+}) binding and two conserved cysteine residues in and around the active site. The psychrophilic enzyme, however, has an extended N-terminal end.

Antarctic bacteria provide a useful model system for studying cold adaptation (15, 17, 31, 36). These organisms are generally represented by the psychrotrophs and psychrophiles, which have the ability to grow at 0°C. They can transcribe at this lower temperature both in vitro and in vivo (31). However, nothing much is known about the nature of promoter and regulatory elements from these bacteria or about the mechanism of transcription at lower temperatures. Most of the transcriptional studies thus far have been carried out with only mesophilic bacteria, and the RNA polymerase from these bacteria, including \textit{Escherichia coli}, cannot transcribe at 0°C. A recent study from our laboratory has demonstrated that the RNA polymerase of the Antarctic psychrotrophic bacterium \textit{Pseudomonas syringae} can transcribe at 0°C. The polymerase from the bacterium was not only active at the low temperature but also could transcribe in vitro preferentially the cold-inducible gene of \textit{E. coli} cspA from a supercoiled template (43). However, absolutely no information is available with regard to the characteristics of promoter sequence, such as the −10 and −35 elements from the bacterium for such low-temperature-specific transcription. Neither is any information available for the in vivo recognition of promoter sequences by RNA polymerase from the cold-adapted \textit{P. syringae}. Therefore, we initially attempted to identify the genes from the Antarctic \textit{P. syringae} that are upregulated at low temperature, with the help of Tn5-mediated random genomic fusions of a promoter-less reporter gene, \textit{lacZ} (23). One of the fusions that produced at least 10- to 14-fold more β-galactosidase at a low temperature (4°C) was identified by cloning and sequencing of ca. 450 bp of DNA sequence proximal to the Tn5 insertion site. The fusion was in the \textit{hutU} gene, which encodes an enzyme, urocanase, of the histidine utilization pathway of bacteria (13, 23). A direct assay of urocanase activity from the \textit{P. syringae} and a few more Antarctic \textit{Pseudomonas} species and comparison of it with that of the mesophilic \textit{Pseudomonas putida} suggested that the \textit{hutU} gene is upregulated in the psychrophilic but not in the mesophile. Therefore, it appeared to us that the \textit{hutU} gene might be a useful model for investigating the mechanism of gene regulation at low temperatures in the Antarctic bacterium. Accordingly, we cloned and sequenced the DNA encompassing the \textit{hutU} gene and its upstream and downstream regions and identified different open reading frames (ORFs) in the region. We also examined transcripts from bacterial cells grown at low (4°C) and high (22°C) temperatures by Northern and primer extension analyses, and we identified the transcription start sites and other putative regulatory elements of the \textit{hutU} gene. Additionally, we compare here the deduced amino acid sequences of the urocanase from the psychrotrophic \textit{P. syringae} and other bacteria, including the mesophilic \textit{P. putida}, in order to examine the possible amino acid substitutions due to a low-temperature adaptation.
MATERIALS AND METHODS

**Bacterial strains and growth conditions.** The Antarctic psychrotrophic bacterium *P. syringae* Lz4W, which grew optimally at 22°C, was isolated, identified, and maintained as reported earlier (42). Routinely, the culture was grown on Antarctic bacterial medium, which contains 0.5% peptone (wt/vol) and 0.2% (wt/vol) yeast extract, at room temperature (22°C) or at a cold temperature (4°C) when needed. The *E. coli* cells were grown at 37°C in Luria-Bertani (LB) medium and maintained on LB agar plates.

**DNA manipulation techniques and cloning of hutU gene.** The bacterial genomic DNA was prepared as described previously (32). Isolation of plasmid DNA, restriction endonuclease digestion, ligation, transformation, and agarose gel electrophoretic separation of DNA were carried out as described by Sambrook et al. (39).

For cloning of the hutU gene from *P. syringae*, a 522-bp DNA fragment from the plasmid pF43, which contained the hutU proximal region of the promoter fusion clone F43 (23), was used as a probe. Initially, a 2.4-kbp PstI DNA fragment containing the hutU gene was cloned in pUC19 (ph180). Subsequently, three more overlapping fragments containing the upstream 3.5-kbp EcoRI-ApI fragment (phUp) and the downstream 2.54-kbp Sall and 1.2-kbp PstI fragments (ph20 and ph39, respectively) were cloned. A total of 6.578 kb were thus cloned spanning the region (Fig. 1A).

**DNA sequence analysis.** Nucleotide sequence determinations were carried out on an automated DNA sequencer (ABI model 373), by using double-strand tagged plasmid DNAs as a template and the ABI PRISM Dye terminator cycle sequencing method (Perkin-Elmer). The BLAST programs (3, 4) were employed for DNA sequence homology search in the NCBI GenBank sequence database (http://www.ncbi.nlm.nih.gov/). PCGENE programs were also used for various DNA and amino acid sequence analyses. Secondary structure prediction analysis of protein was carried out by using the Predict-Protein PHD mail server at EMBL, Heidelberg (predict-help@EMBL-Heidelberg.DE), Germany, which uses combined evolutionary information and neural networks for structural predictions (34, 35).

The nucleotide sequence reported here has been submitted to GenBank under accession number AF326719.

**Isolation of RNA and Northern analysis.** RNA was isolated by the hot phenol method (1) from the bacterial cells (optical density at 600 nm of ~0.5) grown at 22 and 4°C. For the isolation of RNA from cold-shocked cells, *P. syringae* was initially grown at 22°C and then shifted to 4°C and incubated for various time periods before the isolation. Northern hybridization was carried out as described earlier (32) but with the hutU- and hutP-specific probes. Densitometer scanning (Molecular Dynamics) of the autoradiograms was carried out for estimating the fold induction of RNA.

**Primer extension analysis.** Primer extension analyses were carried out by two methods on the total RNA isolated from 4 and 22°C grown cells. In the first method (39), a 32P-end-labeled primer (5’-GGGATGTTGACGCTGACGGTAC-3’) corresponding to the terminal end of the HutU ORF (–7 to +16 nucleotides with respect to the GTG initiation codon) of *P. syringae* was extended in the presence of Moloney murine leukemia virus reverse transcriptase (Pharmacia). In the second method, the unlabeled primer was extended in the presence of [α-32P]ATP and Moloney murine leukemia virus reverse transcriptase as described earlier (8). Both methods produced similar results, except that the primer-extended products were sharper with less background on the autoradiograms obtained by the second method. DNA sequencing reactions were also carried out with the same primer on the double-stranded DNA template (ph180) with a Sequenase 2.0 Kit (U.S. Biochemicals) and run in parallel with the primer extended products on an 8 M urea-6% polyacrylamide gel for mapping the translational start points as described previously (39).

**Enzyme assays.** Urocanase activity was assayed spectrophotometrically by the method of George and Phillips (14). β-Galactosidase was assayed by the method of Miller (28). Proteins were estimated by the method of Bradford (10).

**RESULTS**

**Cloning and sequence analysis of the hutU region from *P. syringae*.** A 522-bp Tn5-proximal sequence of the hutU promoter fusion clone (F43) of *P. syringae* (23) was used as a probe to clone the whole of the hutU gene and its upstream and downstream sequences, as described in Materials and Methods. The reported region (6.578 kbp) was cloned on three overlapping DNA fragments (Fig. 1A). The DNA sequences of the region were determined (accession no. AF326719) and analyzed by a BLAST search (3, 4). Four complete and one incomplete ORFs were identified which had homology with the hypothetical sdeB gene homologue of *Yersinia pestis* (ac-
This is in contrast to other bacteria, such as Klebsiella and B. subtilis, where a single enzyme encoded by hutG catalyzes the conversion of FIGLU to glutamate and formamide at the final step of the histidine degradation pathway (19, 26).

**Predictive analysis of the amino acid sequence of urocanase from *P. syringae***. In recent times comparative analyses of proteins from psychrophilic, mesophilic, and thermophilic bacteria have shed new light on the structural adaptability of the enzymes for catalysis at low temperatures and have provided some clues as to their thermal stability (11, 15). Therefore, it was interesting to compare the primary sequence of the urocanase from the psychrotrophic and mesophilic bacteria. Accordingly, the deduced amino acid sequences of the enzyme from the psychrotrophic *P. syringae* and mesophilic *P. putida* were compared (Fig. 2). It was observed that the *P. syringae* enzyme contained an overall identity of ca. 90% (similarity of 93%) to the *P. putida* enzyme but had an additional 8-aa extension at the N-terminal end (making it 565 aa long). A BLAST search analysis picked up the urocanase homologues from various other bacteria for which the hutU gene sequences are known. An alignment of the deduced amino acid sequences of the enzyme from these bacteria is shown in Fig. 2. Although the enzyme seems to be quite conserved, it fell into two distinct groups of gram-positive (*B. subtilis*, Streptomyces coelicolor, and *Deinococcus radiodurans*) and gram-negative (*P. syringae*, *P. putida*, *P. aeruginosa*, *Vibrio cholerae*, and *A. rhizogenes*) strains (Fig. 3). The urocanase from the plant (*Trifolium repens*) was found clustered with the enzymes from gram-negative bacteria.

From the compositional analyses of the amino acids of urocanase it was apparent that the contents of isoleucine plus leucine (13.1%), the aromatic amino acids (7.2%), and proline (4.4%), which are generally known to change in the psychrophilic enzymes (15), were similar in *P. syringae* and mesophilic *P. putida* (13.28, 7.5, and 4.4%, respectively). A slight increase in the content of serine and threonine (10.4%) was, however, noticed in *P. syringae* when it was compared with the enzymes from *P. putida* and other organisms (7.98 to 9.5%). The arginine content (5.3%), which is known to stabilize the helices, was also marginally lower in *P. syringae* than in the *P. putida* enzyme (5.9%). The *P. syringae* enzyme also contained eight cysteines compared to the seven cysteines of *P. putida* and *P. aeruginosa*. Six of these cysteines are conserved among them and are located in equivalent positions, including the two important cysteines C-410 and C-354 (C-419 and C-363 in *P. syringae*), which were shown to be involved in catalysis and substrate binding, respectively, in the enzyme from *P. putida* (25). Interestingly, the radiotolerant bacterium *D. radiodurans* contains only these two conserved cysteines in the enzyme.

**Predictive structural analyses of the urocanase sequence from *P. syringae*** exhibited some interesting features (Fig. 2). The enzyme contained a conserved β-α-β structural motif of the dinucleotide binding Rossmann fold (GXGX₃-G-X₁₀G/A) for NAD⁺ binding at the 182- to 198-aa region. A ψ BLAST analysis suggested that this NAD binding region has structural homology with the similar region of the glutamate dehydrogenase, including an acidic amino acid residue (E-205 in the case of *P. syringae* urocanase) at the end of the β-strand (201-SLNEE⁵⁰⁵) of the β-α-β motif (7). The acidic amino acid residue commonly forms the hydrogen bonds to the adenine.
ribose hydroxyls and is generally thought to be an indicator of NAD$^+$ specificity as opposed to the NADP$^+$ specificity (6, 7). An implication of the identification of the Rossman fold in urocanase would be that, although the loop is similar to that of other NAD$^+$ binding proteins, it has to be juxtaposed within a tightly held tertiary or quaternary structural fold of the protein. This is because the exogenously added NAD could not be incorporated into the enzyme in vitro, nor could the coenzyme be dissociated from the protein without irreversible denaturation (33). The present predictive analysis (Fig. 2) also suggests that the *P. syringae* enzyme contains at the N-terminal end a short $\alpha$-helix that is absent in other bacteria.

**Temperature-dependent expression of hutU in *P. syringae***

Northern analyses were carried out to examine the expression of the *hutU* gene in *P. syringae* at low (4°C) and high (22°C) temperatures. It was observed that, during the steady-state growth of the bacterium, the amount of transcripts produced at 4°C from the *hutU* gene and *hut* operon was ca. 20-fold higher than the amount present at 22°C. Three transcripts (ca. 7.1, 2.1, and 1.0 kb) hybridized to the probe of the 2.4-kbp *Pst*I fragment containing both *hutU* and *hutP* genes (Fig. 4a). The *hutP*-specific probe, however, hybridized only to the 7.1-kb transcript (data not shown). Thus, it would appear that the 7.1-kb transcript might represent the polycistronic mRNA, whereas the 2.1- and 1.0-kb transcripts represent the processed and/or degraded product from the operon. The 2.1-kb transcript can potentially encode the full-length *hutU* gene and therefore might be physiologically important.

Interestingly, upon a temperature downshift of the bacterium from 22 to 4°C, a “cold-shock” response was noticed.
repens, is closer to the sequence of gram-negative bacteria. The only sequence of possible plant origin (33), from ph180) are marked by arrowheads.

FIG. 3. Relationship among bacterial urocanases. Similarities between the enzymes have been shown as an unrooted neighbor-joining tree, giving all branch lengths (indicated by numerical values). The tree has been drawn by a neighbor-joining plot of the CLUSTAL X (version 1.8) program. The urocanase sequences of gram-negative bacteria (e.g., P. syringae, P. putida, P. aeruginosa, V. cholerae, and A. rhizogenes) and gram-positive bacteria (e.g., B. subtilis, S. coelicolor, and D. radiodurans) fall into two distinct clusters and may have diverged early in evolution. The only sequence of possible plant origin (33), from T. repens, is closer to the sequence of gram-negative bacteria.

because the amount of mRNAs from the hut operon increased only about a maximum of two- to threefold after the shift (Fig. 4b). The amount of the mRNAs was at a maximum by 2 h after the shift and decreased subsequently. Thus, it appears that the hut operon might have cold-responsive regulatory elements that are known to occur in cold-inducible genes of mesophilic bacteria (12, 29).

Transcription start site and promoter region of the hutU gene of P. syringae. Primer extension analyses were carried out to locate the transcription start sites of the hutU gene, with the RNAs isolated from cells of P. syringae grown at low (4°C) and high (22°C) temperatures (Fig. 5). Among several primer-extended products, the longest product (topmost arrow in Fig. 5) was repeatedly observed only with the RNAs isolated from the cells grown at 4°C. Most of the other extended products were common in cells grown at both both low and high temperatures. Based on the longest primer-extended products, the transcription start sites for the low temperature (4°C) and the common site for both low and high temperatures (4 and 22°C) were located in the 5’ region of the hutU gene (Fig. 6A). The rest of the smaller extended products of the primer extension reactions (Fig. 5) could either represent the true transcription start sites or the processed or degraded ends of the mRNAs.

The low-temperature (4°C) specific transcript starts with a G, which is 219 nucleotides upstream of the putative translation initiation codon GTG of the HutU ORF. The common transcription start site for both low and high temperatures is located 39 nucleotides downstream from the former start site (Fig. 6A). Thus, the transcripts from the hut operon seem to have a long 5’-untranslated region (5’-UTR) sequence that is characteristic of the cold-shock genes in mesophilic bacteria, including E. coli (12). Upon further examination of the DNA sequences around the transcription start sites, it was observed that the −10 region has a characteristic CAAAA sequence at both temperatures. The −35 sequences in the promoter region of the 4°C specific and the 4 and 22°C common transcripts were TGTTTAC and CCTGCG, respectively. Interestingly, the promoter region of the 4°C specific transcript had a second CAAAA sequence at the −15 position; the significance of this, if any, is not yet clear.

The sequence at the upstream of the GTG translational start codon of the hutU gene of P. syringae contained a putative HutC repressor binding motif (CTTGTATGTACAAG), which is slightly different from the sequence observed in P. putida and Klebsiella aerogenes (2, 40). Interestingly, in the latter organisms this sequence overlaps with the −35 element of the promoter region, in contrast to the case in P. syringae.
which contains the sequence within the 5'-UTR of the gene (Fig. 6A). It was also observed that a putative nitrogen assimilatory cofactor-binding sequence [ATA-(N₆)-TAT] is located overlapped with the HutC-binding motif (41). A putative catabolite activator protein (CAP) binding sequence was also noticed in the downstream of transcription initiation site. The sequence, AAGTGTGC(N₆)CCTCTTGT, is located at the 35 nucleotides upstream of the GTG translation initiation codon of the hutU ORF. In K. aerogenes a similar sequence was observed centered around nucleotide -81.5 of the promoter region of the hutUH gene cluster (30). Since the CAP binding sequence occurs downstream of the transcription initiation site of P. syringae, the catabolite repressor in this case could act in theory as a transcriptional roadblock for RNA polymerase. It was also noticed that a putative conserved “cold-box”-like sequence (TTGATGAACAACC), which occurs at the 5’ end of the 5'-UTR of the cold-inducible genes of E. coli, is located 123 nucleotides downstream of the translation initiation codon of HutU in P. syringae (Fig. 6A). Since the location of the cold box is at variance with E. coli, any functional significance of the element in P. syringae remains to be determined.

A putative nitrogen regulatory σ⁸ promoter element [GG-(N₆)-GC] was also noticed 81 nucleotides upstream of the translation start site of the HutU ORF of P. syringae. This is interesting for the observation that the shortest primer-extended product (lowest arrow in Fig. 5) is located very close to the expected position of the transcription start site (at the −13 position rather than at the expected −12 position of the initiation site) of the above σ³ promoter (Fig. 6A). Since histidine is utilized as a source of both carbon and nitrogen in bacteria, the hut operon is likely to be regulated by both vegetative (σ⁷⁰ family) and σ⁸-specific promoters (27), and therefore the occurrence of the σ⁸-promoter might be physiologically important.

Expression of hutU gene in E. coli. Since the hutU gene of P. syringae contains some common features, including a long 5'-UTR of cold-inducible genes observed in E. coli, it was interesting to examine the temperature-dependent expression of the gene in the mesophilic bacterium. The expression of the hutU gene in E. coli was examined by transforming the bacterium with a multicopy plasmid (pH180) that contained a 2.6-kbp DNA fragment with the structural gene for urocanase and its upstream regulatory sequences from P. syringae. The cell extracts prepared from the E. coli grown at 37°C exhibited a modest activity of the enzyme (8.3 μmol min⁻¹ mg of protein⁻¹). However, the extracts prepared from the cells grown at a lower temperature (15°C) did not contain any detectable urocanase activity. Thus, it appears that the hutU gene of P. syringae is not expressed in E. coli at the lower growth temperature. Whether the lack of expression is due to the absence of cognate regulatory sequences and factors for transcription or to the inability of translation of the hutU mRNA at the lower temperature in E. coli remains unknown.

DISCUSSION

The present study was undertaken to determine and analyze the DNA sequences of the hutU gene and its promoter region in order to identify, if possible, the putative regulatory cis elements for the regulation of the hutU operon in the psychrotrophic bacterium P. syringae. Additionally, it was thought that an analysis of the deduced amino acid sequence of the enzyme urocanase (the hutU-encoded product) from P. syringae, in comparison with the enzyme from mesophilic P. putida, might provide the clue to the nature of substitution of amino acids in a cold active enzyme. An earlier study had demonstrated that a urocanase from the psychrophilic P. putida A.3.12 could retain, at 0°C, 30% of its maximum activity seen at 30°C (21). The urocanase in P. putida is a homodimer that contains tightly bound NAD⁺ in each subunit, where an intact NAD⁺ is essential for its catalytic reaction. Mechanistically, the enzyme is unique for the fact that the NAD⁺ in this case does not function as a simple redox reagent but as an electrophile for the catalytic addition of water to the urocanate for its conversion into imidazolone propionic acid (7).
Analysis of the genetic organization and promoter region of the \textit{hutU} in \textit{P. syringae}. The organization of the genes in the \textit{hut} operon is known to be variable among gram-positive and gram-negative bacteria. The present study indicates that the organization of the \textit{hut} operon is also variable within the \textit{Pseudomonas} species (Fig. 1B). The analysis also indicates that the nature of the permease gene within the \textit{hut} operon is variable. For example, Antarctic \textit{P. syringae} has the \textit{hutP} gene encoding permease belonging to transporter class (TC) 2.A.39.1 (family NCS1), which is different from \textit{P. putida} (38). The pathogenic \textit{P. aeruginosa} has both \textit{hutP} and \textit{hutT} within the \textit{hut} gene cluster. The published genetic study of \textit{P. putida} had earlier failed to locate any permease or transporter gene in the operon (19, 20). A recent study of \textit{Sinorhizobium meliloti} has demonstrated that the histidine-degrading \textit{hutH} gene is linked to a histidine transporter of the ATP-binding cassette type (9). Thus, there seems to be a random recruitment of the histidine transporters during the microbial evolution of the histidine degradation pathway.

The regulation of the \textit{hut} operon, which had mainly been studied in three gram-negative bacteria, including \textit{Salmonella} sp., \textit{K. aerogenes}, and \textit{P. putida}, and one gram-positive bacterium (\textit{B. subtilis}), had also been found to be variable (16, 26). For example, in \textit{B. subtilis} histidine was found to be the main inducer of the operon, while in \textit{P. putida} urocanase was found to be the inducer. Similarly, a positive activator, \textit{HutP}, was observed to be the main regulator of the operon in \textit{B. subtilis}, while the repressor \textit{HutC} was found to be the main negative regulator of the operon in \textit{P. putida}. The regulation of the operon is, however, complex in both systems for it is subjected to both carbon catabolite repression and nitrogen metabolite regulation in both of the organisms. The temperature-dependent expression of the operon in the cold-adapted bacterium \textit{P. syringae} might further add to the complexity of the regulation of the operon.
In order to investigate the mechanism of temperature-dependent regulation of the operon, the promoter region of _hutU_, i.e., the first gene of the operon in _P. syringae_, has been identified here by primer extension analysis. It appears that the mRNA for the _hutU_ gene has a long 5'-UTR that is a characteristic feature of many cold-inducible genes of mesophilic bacteria, including _E. coli_. It also appears that the promoter region of the _hutU_ of _P. syringae_ contains various putative cis-acting regulatory elements that have been characterized earlier in mesophilic bacteria. However, the locations of these elements are at variance with the known positions. For example, the repressor HutC-binding site, which is known to overlap either with the −35 sequence (e.g., in _P. putida_) or with the region between the −10 and −35 sequences (e.g., in _K. aerogenes_), of the _hut_ promoters, has been found at the +143 base (with respect to the 4°C specific +1 site) and at the +95 base (with respect to the 4 or 22°C specific common +1 site) region of the 5'-UTR of the _hutU_ gene of _P. syringae_ (Fig. 6A).

Similarly, a putative binding sequence [ATA-(N6)-TAT] for the positive activator NAC (for nitrogen assimilation control) protein (41) is located downstream of the transcription sites (data not shown). Interestingly, the HutC- and NAC binding sites mentioned above overlap each other in _P. syringae_, the significance of which is not yet clear. As pointed out above, the CAP binding sequence is also located downstream of the transcription initiation site of the operon in the cold-adapted bacterium (Fig. 6A).

The identification of a unique sequence, CAAAA, at the −10 site of the _hutU_ promoter of _P. syringae_ is interesting. This sequence is probably important for the initiation of transcription at both low and high temperatures (e.g., 4 and 22°C). An extra CAAAA sequence that is observed a half turn away (5 bp) from the DNA helix in the region of the 4°C specific transcription start site might also be important for increased expression of the gene at lower temperatures. The occurrence of a putative σN (σ348) specific promoter sequence, GG-(N6)-GC, and a corresponding primer-extended product of RNA suggests that the operon might be transcribed by RNA polymerase containing both vegetative sigma and nitrogen regulatory sigma factors, depending upon the environmental signals. The observation of five or six primer-extended products of RNA from the _hutU_ gene might also be a reflection of the complex regulatory process of the operon.

It is also interesting that the putative cold-box sequence observed in the 5'-UTR of the cold-inducible genes of mesophilic _E. coli_ is present in the coding region of _hutU_ mRNA of _P. syringae_ (Fig. 6). Whether this sequence has any role in _P. syringae_ at lower temperatures is a matter of conjecture. Recently, it has been shown in _E. coli_ that the Y-box sequence (5'-ATTTGG-3' or its inverted repeat 5'-CCAT-3') might be important for the regulation of cold-inducible genes by an antitermination mechanism (5). In the psychrotrophic _Yersinia enterocolitica_, the induction of the _ppn_ gene encoding polyphosphate kinase at low temperatures has also been shown to be regulated by the Y-box sequence located 230 bp upstream of the translation start site (18).

The promoter region of the low-temperature-inducible gene _icdH_ for isocitrate dehydrogenase in the psychrophilic bacterium _Vibrio_ sp. strain AE1 also contains a CCAAT sequence 2 bp upstream of the −35 site of the promoter that is essential for low-temperature induction (37). The −10 and −35 sequences of the promoter (TAACTA and TTATAG, respectively) in the bacterium were, however, not novel in any sense compared to other housekeeping genes. The analyses of the present study failed to show any Y-box sequence in the proper context of the promoter, although a 5'-ATTGG-3' sequence is observed 185 bp downstream of the GTG initiation codon of HutU (Fig. 6A).

In order to identify any other putative regulatory elements in the promoter region of _hutU_, the potential secondary structural elements were also examined. One such structure with a dyad symmetry (ΔG = −16.6 kcal) is located 70 nucleotides upstream of the transcription start site. Such a hairpin loop structure can potentially function as a transcription stop signal for upstream _hutC_ or as a regulatory element for transcription of the _hutU_ gene.

**Analysis of the urocanase sequence of _P. syringae_.** The deduced amino acid sequence of the urocanase from _P. syringae_, compared with the sequences of the enzyme homologues from other mesophilic bacteria, including _K. aerogenes_, _A. rhizogenes_, _B. subtilis_, _S. coelicolor_, and _D. radiodurans_, did not show any obvious preference for any specific substitution of amino acids in the protein due to the low-temperature adaptation. The active site of the enzyme in these bacteria (Fig. 2) has an almost identical sequence, FGQGLPARICW, including the essential cysteine of the mesophilic _P. putida_ (25). The two conserved cysteines (C-410 and C-354 of _P. putida_) are present in all of them. All of these enzymes have also a distinct signature motif (GXGX_GX_G) of the Rossmann fold, including an equivalent acidic residue at the end of the β-strand of the α/β-fold (Fig. 2). However, multiple alignment of the amino acid sequence suggests that the urocanase has two distinct diverged branches that might be related to the phylogenetic origin of gram-negative and gram-positive bacteria (Fig. 3).

The cold sensitivity of many cold-labile enzymes, including NAD+-specific glutamate dehydrogenase, from various bacteria has been ascribed to the ready dissociation of the monomeric subunits at a lower temperature as a result of weakening hydrophobic bonds (22). Since the hydrophobic bonds stabilize the quaternary structures of proteins and since the enzymes from cold-adapted bacteria have in general reduced hydrophobic interactions to acquire flexibility for functioning at lower temperatures, it might be suicidal for cold active dimeric and oligomeric proteins to adopt a similar strategy. This could be one of the reasons why there are not many substitutions at the level of primary sequence in the dimeric enzyme urocanase from mesophilic _P. putida_ and psychrotrophic _P. syringae_. In fact, a recent study shows that the hydrophobic character of the homodimeric enzyme malate dehydrogenase of the psychrophilic bacterium _Aquaspirillium arcticum_ is similar to that of the enzyme from the thermophilic bacterium _Thermus flavus_ (24). However, three major differences were noticed in the psychrophilic enzyme that were implicated in the efficient catalysis at lower temperature. The differences include (i) an increased relative flexibility at the active-site region of the enzyme; (ii) favorable surface charge distribution, such as more positive potential around the negatively charged substrate (oxaloacetate) binding site and decreased negative potential around the cofactor NADH binding site; and (iii) reduced intersubunit ion pairs and decreased buried surface area in the dimeric interface of the enzyme (24). A similar struc-
ture-function study on the urocarnase of *P. syringae* would be useful to substantiate the generality of these findings.

In conclusion, the present study shows that the –10 and –35 characteristics of the promoter of the *hutU* gene that is up-regulated at low temperatures in *P. syringae* are unique. The occurrence of multiple A’s in CAAA might be important for low-temperature melting of the promoter. The present study also reflects the possible complexity and uniqueness of regulation in the operon of *P. syringae* due to the presence of many putative regulatory cis elements that are located downstream rather than in the usual location upstream of the transcription start site. The present study also identifies two distinct clusters of urocarnase sequences among the bacteria that might be related to their origin or lineage. The identification of a Rossmann fold for NAD⁺ binding is also important for future modeling of the enzyme since this site is presumed to be different from other NAD-requiring enzymes for its inaccessibility without denaturation of the urocarnase.

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