Identification and Cloning of an Erwinia carotovora subsp. carotovora Bacteriocin Regulator Gene by Insertional Mutagenesis

DUEN-YAU CHUANG, AMPAABENG G. KYEREMEH, YUICHI GUNJI,† YOSHIYUKI TAKAHARA, YOSHIO EHARA, and TOSHIK KIKUMOTO

Institute of Genetic Ecology, Tohoku University, 2-1-1 Katahira, Aoba-ku, Sendai 980-8577, Central Glass Co. Ltd., Chemical Research Center, 2805 Imafuji-nakadai, Kawagoe, Saitama 350-1151, and Faculty of Agriculture, Tohoku University, 1-1 Tsuetsumidori-Amamiyamachi, Aoba-ku, Sendai 981-8555, Japan

Received 6 August 1998/Accepted 8 January 1999

Avirulent Erwinia carotovora subsp. carotovora CGE234-M403 produces two types of bacteriocin. For the purpose of cloning the bacteriocin genes of strain CGE234-M403, a spontaneous rifampin-resistant mutant of this strain, M-rif-11-2, was isolated. By Tn5 insertion mutagenesis using M-rif-11-2, a mutant, TM01A01, which produces the high-molecular-weight bacteriocin but not the low-molecular-weight bacteriocin was obtained. By thermal asymmetric interlaced PCR, the DNA sequence from the Tn5 insertion site and the DNA sequence of a contiguous 1,280-bp region were determined. One complete open reading frame (ORF), designated ORF2, was identified within the sequenced fragment. The 3' end of another ORF, ORF1, was located upstream of ORF2. A noncoding region and a putative promoter were located between ORF1 and ORF2. Downstream from ORF2, the 5' end of another ORF (ORF3) was found. Deduction from the nucleotide sequence indicated that ORF2 encodes a protein of 99 amino acids, which showed high homology with Yersinia enterocolitica Yrp, a regulator of enterotoxin (Y-ST) production; Escherichia coli host factor 1, required for Qf-replicase; and Azorhizobium caulinodans NrfA, required for the expression of nifA. ORF2 was designated brg, bacteriocin regulator gene. A fragment containing ORF2 and its promoter was amplified and cloned into pBR322 and pHSG415, and the recombinant plasmids, pBYL1 and pHYL1, were transferred into E. coli DH5. Plasmid pBYL1 was reisolated and transferred into the insertion mutant TM01A01. Transformants carrying the plasmid, which was reisolated and designated pBYL1, re-produced the low-molecular-weight bacteriocin.

Erwinia carotovora subsp. carotovora is a phytopathogenic bacterium responsible for the soft-rot disease of many plant species. Despite its economic importance, no efficient method, either chemical or otherwise, has been found to control this worldwide disease. Agrochemicals are generally used for the control of this disease, but in a quest for a more environmentally friendly control methods, biological control is under investigation.

Some bacterial species produce one or more antibacterial substances called bacteriocins, which enhance their competitiveness with other related bacterial species (27). According to the report of Kikumoto et al. (13), the antibacterial activity of two types of bacteriocin produced by biocontrol agents (avirulent E. carotovora subsp. carotovora) may contribute to the suppression of soft-rot disease (18). There is also strong evidence of the effectiveness of bacterial control of the soft-rot disease of Chinese cabbage (14, 22). A biological control agent with the trade name Biokeeper has therefore been developed for the control of this disease in Japan. In view of these reports, identification and cloning of the gene(s) controlling bacteriocin production may facilitate its use in further control methods, biological control is under investigation.

avirulent E. carotovora subsp. carotovora may contribute to the suppression of soft-rot disease (18). There is also strong evidence of the effectiveness of bacterial control of the soft-rot disease of Chinese cabbage (14, 22). A biological control agent with the trade name Biokeeper has therefore been developed for the control of this disease in Japan. In view of these reports, identification and cloning of the gene(s) controlling bacteriocin production may facilitate its use in further control methods, biological control is under investigation.

Bacterial strains, plasmids, and growth media. The bacterial strains and plasmids used in this study are listed in Table 1. The putative biocontrol agent produces two types of bacteriocin, low- and high-molecular-weight bacteriocins. E. carotovora subsp. carotovora strains were propagated at 28°C in nutrient agar (NA) containing 1.4% agar or with shaking in Luria-Bertani (LB) medium with 5 g of NaCl per liter substituted for 10 g. Escherichia coli strains were propagated at 37°C in LB medium with shaking. Rifampin, kanamycin, and ampicillin (all at 50 μg per ml) were added to NA and LB agar when necessary.

Bacterial mating. Bacterial mating was carried out on NA by the membrane-filter mating method (8), by using 0.22-μm-pore-size membrane filters (Millipore, Inc. Bedford, Mass.). The filters were placed on NA and incubated overnight at 28°C. Appropriate dilutions of the suspension of the progeny of the mating were spread on modified Drigalski's agar plates (26) containing 50 ppm each of rifampin and kanamycin and were incubated at 28°C for 24 to 48 h before the colonies were counted.

Bacteriocin assays. Bacteriocin production was examined by the double-layer method of Fredericq (7), but hard and soft IFO-802 media containing, respectively, 1.4 and 0.65% agar were used. Growth inhibition zones around the colonies were considered an indication of bacteriocin production.
Genetic engineering technique. Plasmids of *E. carotovora* subsp. *carotovora* were isolated according to the method of Kado and Liu (11), and *E. coli* plasmids were isolated by the method of Sambrook et al. (20). Total DNA was isolated as previously described (16).

Oligonucleotide DNA primers were synthesized by the Gibco BRL Co. and Takara Co. (Tokyo, Japan). Reagents were purchased from the Takara Co. The general PCR procedure has been described by Sambrook et al. (20). Thermal asymmetric interlaced PCR (TAIL-PCR) was performed according to the method of Liu and Whittier (15), but the annealing temperature for specific primers was increased from 63°C to 65°C for this study. For TAIL-PCR, specific primers that are complementary to the respective sequences of Tn\textsuperscript{5} were synthesized: AGAGAACACAGATTTAGCCCAGTCGG (PF-1), CCGCACGATGAAGAGCAGAAGTTAT (PF-2), GATCCTGGAAAACGGGAAAGGTTC (PF-3), GCCGAAGAGAACACAGATTTAGCCCA (PR-1), CCGCACGATGAAGAGCAGAAGTT (PR-2), and CAGATCTCTGGAAAACGGGAAAGG (PR-3). In addition, two arbitrary degenerate primers, GTNCGA(C/G)(A/T)CANA(A/T)GTT (N-2) and (A/T)GTG(N)ANCANAGA (N-3), were used.

For sequencing of TAIL-PCR products, the ABI PRISM Dye Terminator Cycle Sequencing Ready Reactions Kit was used. Cycle sequencing was carried out on a GeneAmp System 9600 thermocycler. Sequencing using an automated DNA sequencer 373A (ABI) was carried out according to the manufacturer's protocol.

Southern and colony hybridizations, probe labeling, and detection were performed by using the DIG DNA Labeling and Detection Kit (Boehringer GmbH, Mannheim, Germany) as described by the manufacturer. Hybridization was performed overnight, and the membrane was washed according to the recommendation of the manufacturer.

DNA electrophoresis, restriction digestion, ligation, and transformation for *E. coli* were carried out as described by Sambrook et al. (20). Plasmid DNA transformation for *E. carotovora* subsp. *carotovora* was performed by the methods of Hinton et al. (10) and Hanahan (9).

Computer analysis of sequence data. The nucleotide sequence and deduced amino acid sequence of Brg were compared by the BLAST and FASTA programs of the National Center for Biotechnology Information server. Sequence data were compiled with DNASIS-Mac software (Hitachi, Tokyo, Japan).

Isolation of transposon insertion mutants. For the Tn\textsuperscript{5} mutagenesis, the transmissible plasmid pJB4J1 was used. By mating *E. coli* 1830 with *E. carotovora* subsp. *carotovora* M-rif-11-2, 5,500 insertion mutants that could grow on a selective medium containing 50 ppm each of rifampin and kanamycin were isolated. In order to ascertain their antibiotic resistance, their growth on the selective medium was rechecked and found to be a stable property of the isolates. Of these 5,500 isolated mutants, only 1 was defective in the production of low-molecular-weight bacteriocin. This defective mutant was further purified on modified Drigalski's medium containing 50 ppm each of rifampin and kanamycin. All the mutants defective in low-molecular-weight bacteriocin production were therefore siblings isolated from the same purification plates.

**Bacteriocin assay of mutants.** The bacteriocin activity of the test isolates was examined. The parental strain produces two
After TAIL-PCR was performed three times, two to three fragments of more than 1,280 bp were obtained. All the insertions characterized were at the same position in the brg gene, as they are all siblings.

Sequence analysis and homology. The gene structure of the 1,280 bp was determined (GenBank accession no. AF039142). In the direction of transcription indicated by the complementation studies, one complete ORF (ORF2) was present, and Tn5 was located in the same ORF between bp 878 and 879.

The predicted amino acid sequence of ORF2 was compared with those of the high-molecular-weight bacteriocin, which is restricted to the immediate surroundings of the colony, and the low-molecular-weight bacteriocin, which diffuses relatively further away from the colony. It was therefore expected that a mutant deficient in the low-molecular-weight bacteriocin would produce a restricted inhibition zone. The inhibition zones of the putative isolates (insertion mutants), typical of the high-molecular-weight bacteriocin, were restricted compared to those of the parent strain (Fig. 1). This indicates the possibility that transposon Tn5 has been successfully inserted into the genes of the low-molecular-weight bacteriocin. The mutants therefore produced only the high-molecular-weight bacteriocin. It was also observed that the insertion mutants were sensitive to UV light, and the duration of the UV light induction (irradiation) in the course of the bacteriocin assay had to be shortened in order to avoid complete killing of the cells. This is also an indication that the gene that is defective in the insertion mutants may not only control low-molecular-weight bacteriocin but also influence the degree of tolerance of this bacterium to UV light.

Detection of Tn5 in the mutants. To ascertain whether Tn5 had actually been inserted into the putative isolates, nested PCR was used to amplify the nptII gene (25), and two oligonucleotides, CTCGACGGTGCATAGGTTGG, and AAGCAGAGGAAACCGGTAGCATGCCCCTCATG (P-4), were synthesized. Almost all the test isolates except M-rif-11-2, which does not harbor the Tn5 gene, produced a short DNA fragment of 500 bp, indicating the presence of the Tn5 insert in the mutants. Southern blot hybridization also confirmed the above results (data not shown).

Amplification of Tn5 insertion junction DNA and sequencing. After TAIL-PCR was performed three times, two to three bands of different sizes were obtained for each sample. All the fragment products were isolated by electrophoresis and purified, and the sequences of the recovered products were analyzed. Analysis of the respective bands showed a high homology of about 95% or more, indicating a possible similarity in origin. A nucleotide sequence of 1,280 bp was obtained. All the Tn5 insertions characterized were at the same position in the brg gene, as they are all siblings.

Sequence analysis and homology. The gene structure of the 1,280 bp was determined (GenBank accession no. AF039142). In the direction of transcription indicated by the complementation studies, one complete ORF (ORF2) was present, and Tn5 was located in the same ORF between bp 878 and 879.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Low-mol-wt bacteriocin</th>
<th>High-mol-wt bacteriocin</th>
</tr>
</thead>
<tbody>
<tr>
<td>M-rif-11-2</td>
<td>7 mm</td>
<td>+</td>
</tr>
<tr>
<td>TM01A01</td>
<td>None</td>
<td>+</td>
</tr>
<tr>
<td>TM01A01/pBYL1</td>
<td>7 mm</td>
<td>+</td>
</tr>
</tbody>
</table>

* M-rif-11-2 (parent strain), TM01A01 (Tn5 insertion mutant), and pBYL1- TM01A01/pBYL1 (transformed mutant) were used as bacteriocin producers, and strain T-29 was used as an indicator.
Bacteria were cultured in LB medium with shaking at 28°C. by colony hybridization using brg into E. coli. The presence of the brg DNA was detected by colony hybridization using brg DNA probes and by electrophoresis after digestion with BamHI and HindIII. The brg band size was certified to be 400 bp (data not shown). The DNA of plasmid pBYL1 was isolated from DH5/pBYL1 and transferred into the insertion mutant TM01A01. One hundred colonies were isolated by selection on modified Drigalski's medium containing 50 ppm each of kanamycin, rifampin, and ampicillin, and the brg DNA was detected as previously described. The new plasmid was designated pBYL1 after reisolation from the transformed colonies.

Recovery of bacteriocin production and characteristics of the transformed mutants. The bacteriocin assay for the insertion mutants after transformation indicates a successful recovery of their ability to produce the low-molecular-weight bacteriocin. Their larger inhibition zones were therefore comparable to those of the parent strain, M·rif-11-2 (Table 2). This proves that production of the low-molecular-weight bacteriocin was actually regulated by brg DNA.

The cloning of the brg DNA and its subsequent transfer into the insertion mutant TM01A01 resulted in the recovery of production of the low-molecular-weight bacteriocin. The homology search and computer analysis for the DNA sequence of the bands obtained by TAIL-PCR also confirmed the above results indicating that brg is required for the production of the low-molecular-weight bacteriocin. The homology search also indicates that Brg may also regulate the expression of genes other than those for bacteriocin production, as observed in the roles of Yrp in Y. enterocolitica (17) and hfq in E. coli K-12 and other bacteria (3, 5, 6, 21, 23, 24).

It has been reported that an hfq-defective mutant (hfq::Ω) exhibited defects in osmosensitivity, cell growth, cell shape and size, plasmid supercoiling, and sensitivity to UV light (23). We also observed that on modified Drigalski's medium, colonies transformed with pBYL1, containing the brg gene, were green and were larger in diameter (about 13 mm) than the parents and the insertion mutants, which were only 5 mm in diameter and yellow. Further incubation of the brg-defective mutant under this condition results in the death of the cells. The transformed colonies were also found to have a very strong smell compared to those of the parents and the insertion mutants. A possible explanation of this observation is that the utilization of the lactose present in modified Drigalski's medium results in the formation of lactic acid (lowering the pH), indicated by the yellowish color, which is further utilized as a sole carbon source, leading to an increase in the pH around the colony and the green color of the colony. The differences in color and colony size are therefore due to the differences in the efficiency with which the respective colonies utilize lactic acid and tolerate a low pH. The relative tolerance of the reduced pH and efficiency of lactic-acid utilization by the brg transformant points to a possible control by this gene. The observed sensitivity of the brg-defective mutant to UV light is similar to that previously observed for the E. coli hfq::Ω mutant (23).

It was also observed that among the insertion mutants, most of the cells, formed a precipitate at the bottom of the test tube when cultured in LB medium with shaking for 24 or 48 h. Further examination showed that, as observed for the hfq::Ω mutant (23), the brg-defective mutants were larger and more elongated than the parent and the transformed cells (Fig. 3). However, among the 160 cells measured for each strain (parent, mutant, and transformant), it was observed that the data for the mutant strain were the least uniform. The nonuniformity seems to stem from impaired or delayed cell division. This could be supported by larger cell size during the 4th and 11th h, followed by a marked reduction in cell size at the 24th h, with a subsequent reduction in standard deviation. It therefore stands to reason that the brg gene affects not only the size of the cells but also, either directly or indirectly, the process of cell division. This may be the reason for the increase in the doubling time observed for the hfq-defective mutant (23).

All these results show that Brg may be an analogue of Hfq which operates in this strain of E. carotovora subsp. carotovora. Therefore further study of this gene is needed, as it may have pleiotropic effects on other aspects of cell physiology which have not yet been studied.

Nucleotide sequence accession number. The GenBank accession no. of the sequence of the brg genes is AF039142.
iversity, Sendai, Japan, for donating *E. coli* DH5 and for insightful discussion and guidance.

**REFERENCES**


9. Hanahan, D. 1983. Studies on transformation of *Escherichia coli* with plas-


