

Porphyromonas gingivalis galE Is Involved in Lipopolysaccharide O-Antigen Synthesis and Biofilm Formation[∇]

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Porphyromonas gingivalis is a crucial component of complex plaque biofilms that form in the oral cavity, resulting in the progression of periodontal disease. To elucidate the mechanism of periodontal biofilm formation, we analyzed the involvement of several genes related to the synthesis of polysaccharides in *P. gingivalis*. Gene knockout *P. gingivalis* mutants were constructed by insertion of an *ermF-ermAM* cassette; among these mutants, the *galE* mutant showed some characteristic phenotypes involved in the loss of GalE activity. As expected, the *galE* mutant accumulated intracellular carbohydrates in the presence of 0.1% galactose and did not grow in the presence of galactose at a concentration greater than 1%, in contrast to the parental strain. Lipopolysaccharide (LPS) analysis indicated that the length of the O-antigen chain of the *galE* mutant was shorter than that of the wild type. It was also demonstrated that biofilms generated by the *galE* mutant had an intensity 4.5-fold greater than those of the wild type. Further, the *galE* mutant was found to be significantly susceptible to some antibiotics in comparison with the wild type. In addition, complementation of the *galE* mutation led to a partial recovery of the parental phenotypes. We concluded that the *galE* gene plays a pivotal role in the modification of LPS O antigen and biofilm formation in *P. gingivalis* and considered that our findings of a relationship between the function of the *P. gingivalis* *galE* gene and virulence phenotypes such as biofilm formation may provide clues for understanding the mechanism of pathogenicity in periodontal disease.

Porphyromonas gingivalis has been implicated in the causation and pathogenesis of periodontal disease. Although this gram-negative anaerobe possesses a diverse repertoire of virulence factors, including fimbriae, gingipains, hemagglutinins, lipopolysaccharide (LPS), and others, such as outer membrane vesicles (6, 9, 13, 14, 18), the mechanism by which it initiates periodontal disease is not fully understood. On the other hand, it has been suggested that *P. gingivalis* is usually a late colonizer in the oral cavity and plays a central role in subgingival pockets (18). Thus, subgingival plaque biofilms containing *P. gingivalis* appear to be important for the progression of periodontal disease. In *P. gingivalis* strain W50, the bacterial surface polysaccharide thought to contribute to biofilm formation was found to contain mannose, galactose, rhamnose, glucose, and 2-acetamido-2-deoxy-D-glucose in a relative molar ratio of 13.5:2.0:1.4:1.0:1.0 (7).

Among the virulence factors of *P. gingivalis*, LPS, the major integral component of the outer membrane, which exhibits immunostimulatory and inflammatory activities, has three general components: O-antigen polysaccharide, core oligosaccharide, and lipid A. Most of the biological effects of LPS result from the lipid A part; however, there is an increasing body of evidence indicating that O antigen plays an important role in its effective colonization of host tissue (5, 11, 26–28) as well as in resistance to some bactericidal effects (2, 29, 37). The location of O antigen places it at the interface between the bacterium and its environment. Hence, a critical density of the long O-antigen chain not only prevents access of detrimental mol-

ecules into the outer membrane but also plays an important role in the initial attachment of the bacterium to other organisms, such as other bacteria and mammals, or inorganic surfaces. However, it is not clear whether changes in O antigen of *P. gingivalis* have effects on the properties of attachment and biofilm formation.

To elucidate the virulence mechanisms associated with periodontal biofilms, we analyzed several genes of *P. gingivalis* related to the synthesis of polysaccharides, which are contained as building components of the bacterial surface and biofilms. We selected three genes involved in the synthesis of LPS O antigen: *wecA*, encoding GlcNAc-1-phosphate transferase; *wbaP*, encoding galactose-1-phosphate transferase; and *wzt*, encoding an ATP-binding protein subunit. In addition, we analyzed the *galE* gene, which is involved in the synthesis of sugar nucleotides in the Leloir pathway. The GalE enzyme (UDP-galactose 4-epimerase) catalyzes the interconversion between UDP-glucose and UDP-galactose and is universally conserved among all organisms from bacteria to *Homo sapiens*. Further, *galE* mutants of *Salmonella enterica* serovar Typhimurium, *Neisseria gonorrhoeae*, and *Haemophilus influenzae* have been shown to be relatively avirulent compared with their parental strains (15, 22, 33).

In the present study, inactivation of the *galE* gene of *P. gingivalis* resulted in a shortened O antigen and a significant increase in biofilm formation, which demonstrates the relationship between the *galE* gene and biofilm formation in *P. gingivalis*.

MATERIALS AND METHODS

Bacterial strains, culture conditions, and plasmids. The bacterial strains and plasmids used in this study are listed in Table 1. *Escherichia coli* DH5 α was grown in LB broth or on LB agar plates under aerobic conditions. *P. gingivalis* was grown in brain heart infusion (BHI) broth supplemented with hemin and

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TABLE 1. Bacterial strains and plasmids used in this study

Bacterial strain/plasmid	Relevant phenotype, description, and/or selective marker ^a	Source or reference
Strains		
<i>Escherichia coli</i> DH5 α	F' ϕ 80 <i>lacZ</i> Δ M15 Δ (<i>lacZYA</i> · <i>argF</i>)U169 <i>deoR recA1 endA1</i> <i>hsdR17</i> (r _K ⁻ m _K ⁺) <i>phoA supE44</i> λ ⁻ <i>thi-1 gyrA96 relA1</i>	Takara Bio, Otsu, Japan
<i>Porphyromonas gingivalis</i> ATCC 33277	Wild type	American Type Culture Collection
ATCC 33277 <i>htpG</i> strain	Erm ^r	This study
ATCC 33277 <i>wecA</i> strain	Erm ^r	This study
ATCC 33277 <i>wbaP</i> strain	Erm ^r	This study
ATCC 33277 <i>galE</i> strain	Erm ^r	This study
ATCC 33277 <i>galE</i> -c strain	Tet ^r Erm ^s	This study
ATCC 33277 <i>wzt</i> strain	Erm ^r	This study
Plasmids		
pUC19	Cloning vector; Amp ^r	Takara Bio, Otsu, Japan
pBR322	Cloning vector; Amp ^r	Takara Bio, Otsu, Japan
pHS19	Contains the <i>ermF-ermAM</i> cassette between EcoRI and BamHI sites of pBR322 without <i>bla</i> ; Erm ^r	10
pBR322-erm	Contains the <i>ermF-ermAM</i> cassette between EcoRI and BamHI sites of pBR322; Amp ^r Erm ^r	This study
pRN2	Contains PG0045 between KpnI and AvaI sites of pUC19; Amp ^r	This study
pRN2-erm	Contains the <i>ermF-ermAM</i> cassette in PmaCI site within PG0045 of pRN2; Amp ^r Erm ^r	This study
pRN3	Contains PG0106 between XbaI and BamHI sites of pUC19; Amp ^r	This study
pRN3-erm	Contains the <i>ermF-ermAM</i> cassette in BglII site within PG0106 of pRN3; Amp ^r Erm ^r	This study
pRN4	Contains PG1964 between EcoRI and XbaI sites of pUC19; Amp ^r	This study
pRN4-erm	Contains the <i>ermF-ermAM</i> cassette in Bsp1407I site within PG1964 of pRN4; Amp ^r Erm ^r	This study
pRN6	Contains PG0347 between EcoRI and BamHI sites of pUC19; Amp ^r	This study
pRN6-erm	Contains the <i>ermF-ermAM</i> cassette in BseRI site within PG0347 of pRN6; Amp ^r Erm ^r	This study
pRN7	Contains PG1225 between XbaI and BamHI sites of pUC19; Amp ^r	This study
pRN7-erm	Contains the <i>ermF-ermAM</i> cassette in ApaI site within PG1225 of pRN7; Amp ^r Erm ^r	This study
pKD375	Contains <i>tetQ</i> cassette in pUC19; Amp ^r Tet ^r	25
pUC19Q	Contains <i>tetQ</i> cassette between SmaI and BamHI sites in pUC19; Amp ^r Tet ^r	This study
pQG	Contains an entire region of <i>galE</i> and the 0.3-kb downstream region between BamHI and XbaI sites of pUC19Q; Amp ^r Tet ^r	This study
pQGQ	Contains a region of 0.8 kb upstream of <i>galE</i> between KpnI and SmaI sites of pQG; Amp ^r Tet ^r	This study

^a Amp^r, ampicillin resistant; Erm^r, erythromycin resistant; Erm^s, erythromycin sensitive; Tet^r, tetracycline resistant.

menadione (HM) or on BHI-HM blood agar plates in an anaerobic chamber (miniMACS anaerobic workstation; Don Whitley Scientific Ltd., Shipley, United Kingdom) in 80% N₂, 10% H₂, and 10% CO₂. For *E. coli*, ampicillin and erythromycin were supplemented at 100 μ g/ml and 300 μ g/ml, respectively, when required. For *P. gingivalis*, erythromycin and tetracycline were supplemented at 5 μ g/ml and 0.7 μ g/ml, respectively, when required. Recombinant plasmids derived from pUC19 or pBR322 were transformed into *E. coli* DH5 α .

Sequence analysis. The publicly available sequences from the *P. gingivalis* genome project (<http://www.tigr.org>) were examined for the presence of the target genes; *wecA* (glycosyl transferase group 4 family protein), *wbaP* (bacterial sugar transferase), *wzt* (an ATP-binding protein subunit), and *galE* (UDP-galactose 4-epimerase). The amino acid sequences encoded by these genes of *Salmonella enterica* serovar Typhi or *Bacteroides thetaiotaomicron*, which were obtained from the GenBank database, were used as query sequences. Homology analysis was performed with BLAST.

Construction of mutants. All DNA modifications and manipulations were carried out using standard methods (25, 34). An overview of the construction of the *galE* mutant is shown in Fig. 1A. The oligonucleotides used in this study are listed in Table 2. A 2.2-kb *ermF-ermAM* cassette obtained from pHS19 (10) was cloned between the EcoRI and BamHI sites of pBR322, resulting in pBR322-erm. The *ermF-ermAM* cassette of pBR322-erm was used as a selection marker. A 1.0-kb *galE* region was PCR amplified from the chromosomal DNA extracted from *P. gingivalis* ATCC 33277 by using the high-fidelity DNA polymerase Pyrobest (Takara Bio Inc., Shiga, Japan). The amplified region was cloned

between the BamHI and XbaI sites of pUC19, resulting in pRN6. The *ermF-ermAM* cassette of pBR322-erm was subcloned into the BseRI site of *galE* of pRN6 to yield pRN6-erm. The *ermF-ermAM* cassette and the flanking *galE* regions were retrieved by BamHI and XbaI digestions and then electrotransformed into *P. gingivalis* ATCC 33277 to generate an insertion mutation in the *galE* gene. Construction of the other mutant strains was carried out in a similar manner: the BglII, NaeI, and ApaI sites in the *wecA*, *wbaP*, and *wzt* regions, respectively, were utilized for insertion of the *ermF-ermAM* cassette. Construction of the *htpG* mutant as a control strain was also performed, since HtpG, which is a homolog of the human heat shock protein Hsp90, was reported to have no effect on the growth of *P. gingivalis* or on its adherence to other bacterial and human cells (21, 36). The PmaCI site in the *htpG* region was utilized for insertion of the *ermF-ermAM* cassette.

Complementation of *galE* mutation. The construction of the vectors used for complementation of the *galE* mutation is shown in Fig. 1B, while an overview of the complementation of *galE* mutation is shown in Fig. 1C. A 2.7-kb *tetQ* cassette was used as a selection marker for the complementation of the *galE* mutant. First, the *tetQ* cassette, which was tagged with additional sequences for restriction enzyme SmaI and BamHI sites, was amplified from pKD375, kindly gifted by K. Nakayama of Nagasaki University (25). The amplified DNA fragment was digested by the enzymes and cloned between the SmaI and BamHI sites of pUC19, producing pUC19Q. Next, a 1.0-kb cassette of *galE* gene with the 0.3-kb downstream region was coamplified from the chromosomal DNA of *P. gingivalis* 33277 and cloned between the BamHI and XbaI sites of pUC19Q, producing

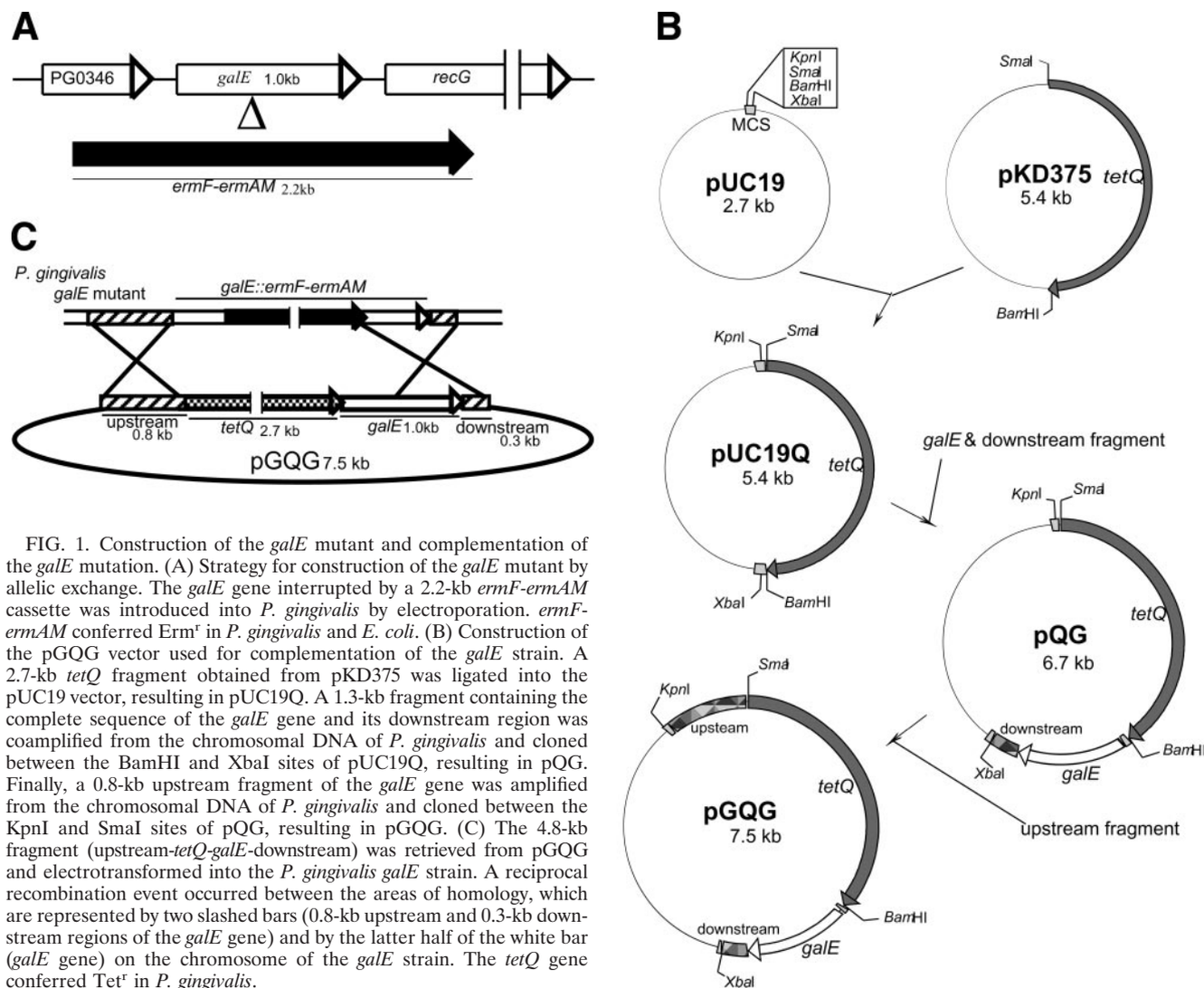


FIG. 1. Construction of the *galE* mutant and complementation of the *galE* mutation. (A) Strategy for construction of the *galE* mutant by allelic exchange. The *galE* gene interrupted by a 2.2-kb *ermF-ermAM* cassette was introduced into *P. gingivalis* by electroporation. *ermF-ermAM* conferred *Erm^r* in *P. gingivalis* and *E. coli*. (B) Construction of the pQG vector used for complementation of the *galE* strain. A 2.7-kb *tetQ* fragment obtained from pKD375 was ligated into the pUC19 vector, resulting in pUC19Q. A 1.3-kb fragment containing the complete sequence of the *galE* gene and its downstream region was coamplified from the chromosomal DNA of *P. gingivalis* and cloned between the BamHI and XbaI sites of pUC19Q, resulting in pQG. Finally, a 0.8-kb upstream fragment of the *galE* gene was amplified from the chromosomal DNA of *P. gingivalis* and cloned between the KpnI and SmaI sites of pQG, resulting in pQGQ. (C) The 4.8-kb fragment (upstream-*tetQ*-*galE*-downstream) was retrieved from pQGQ and electrotransformed into the *P. gingivalis galE* strain. A reciprocal recombination event occurred between the areas of homology, which are represented by two slashed bars (0.8-kb upstream and 0.3-kb downstream regions of the *galE* gene) and by the latter half of the white bar (*galE* gene) on the chromosome of the *galE* strain. The *tetQ* gene conferred *Tet^r* in *P. gingivalis*.

pQG. Finally, a 0.8-kb cassette of the upstream region of the *galE* gene was amplified from the chromosomal DNA of *P. gingivalis* and cloned between the KpnI and SmaI sites of pQG, producing pQGQ. The 4.8-kb cassette (upstream-*tetQ*-*galE*-downstream) of pQGQ was retrieved by KpnI and XbaI digestions and then electrotransformed into the *P. gingivalis galE* strain. Transformants that grew on tetracycline plates and not on erythromycin plates were considered to be potential target colonies. One of the many transformants that had lost erythromycin resistance and retained tetracycline resistance was named the *galE-c* strain. To confirm the integration of the target region into the chromosome of the *galE* strain, generated following the predicted recombination event, PCR analysis and partial sequencing of DNA from *P. gingivalis* ATCC 33277 and the *galE* and *galE-c* strains were performed (ABI Prism 310 genetic analyzer; Applied Biosystems, California).

Preparation of cell extracts and UDP-galactose 4-epimerase assay. One-milliliter portions of fresh *P. gingivalis* wild-type, *galE*, and *galE-c* strains at an optical density at 660 nm (OD_{660}) of 1.0 were harvested by centrifugation ($10,000 \times g$ for 2 min at 4°C) and then washed twice and suspended with 20 mM of phosphate buffer (pH 6.5) containing 50 mM NaCl, 10 mM $MgCl_2$, and 1 mM dithiothreitol (1). The cells were disrupted ultrasonically in an ice bath for 60 cycles of 1 second each at 100 W (IKASONIC U50 control; Janke & Kunkel GmbH & Co. KG, Staufen, Germany). To obtain the cell extract, debris was removed by centrifugation ($10,000 \times g$ for 2 min at 4°C). The protein concentration of the cell extract was determined using a Bio-Rad protein assay based on the method of Bradford (3). UDP-galactose 4-epimerase activity was assayed as described by Maxwell et

al. (23). The procedure was executed as follows. The reaction mixture contained 50 μ l of glycine-NaOH buffer (1 M, pH 8.7), 20 μ l of β -NAD⁺ (25 μ mol/ml; ORIENTAL YEAST Co., Osaka, Japan), 25 μ l of UDP-glucose dehydrogenase (2 U/ml; EMD Biosciences, Inc., San Diego, CA), 475 μ l of distilled water, and 20 μ l of cell extract. The reaction was started by the addition of 30 μ l of UDP-galactose (2.2 μ mol/ml; EMD Biosciences, Inc.). After 30 min, the formation of NADH was determined by measuring the increase in absorbance at 340 nm. Enzyme activity was expressed as nmol per minute per 1 mg of total cell extract protein. The blank consisted of the reaction mixture without the cell extract.

Quantification of intracellular carbohydrates. *P. gingivalis* wild type or the *galE* strain was cultured in 0.5-liter portions of BHI-HM broth supplemented with and without galactose (0%, 0.01%, 0.05%, and 0.1%). The cells were collected and washed with phosphate-buffered saline (PBS) (pH 7.4) three times ($3,000 \times g$ for 20 min at 4°C) and then lyophilized. Forty milligrams of lyophilized cells was suspended in 5 ml of phosphate buffer (pH 7.2) with 1% Triton X-100 and then disrupted ultrasonically in an ice bath for five cycles of 30 seconds each at 100 W (IKASONIC U50 control). Cell debris was removed by centrifugation ($10,000 \times g$ for 2 min at 4°C). The supernatant was centrifuged at $100,000 \times g$ for 2 h at 10°C. The soluble fraction was collected as intracellular components. The quantity of carbohydrates was determined using a phenol-sulfuric acid colorimetric method (12) with glucose as the control sugar.

LPS analysis. *P. gingivalis* wild-type and mutant strains were cultured in 0.5 liter BHI-HM broth. The cells were collected and washed with PBS (pH 7.4)

TABLE 2. Primer pairs used for gene cloning

Target region	Name	Sequence (5'-3') ^a
<i>htpG</i>	htpGp2-f	GGGGTACCTGAGGAGAATAGCGAGTT
	htpGp2-r2	CCCCCGGGTTGGGTTTTCAGCACAAAC
<i>wecA</i>	PG0106-D	GCTCTAGATGCGACCTCAATCCTTC
	PG0106-E	CGGGATCCCAACCAGTGTTCCTCT
<i>wbaP</i>	PG1964-A	GGAATTCGGGATACTACAACAAACC
	PG1964-C	GCTCTAGAAGGCTCATATTCTCGTAG
<i>galE</i>	PG0347-A	GGAATTCGGATCCCATACGACTGTG
	PG0347-C	GCTCTAGAACGACTCCAAAGCTTTC
<i>wzt</i>	PG1225-A	GCTCTAGAGATCGGACTGCTTCTGGT
	PG1225-B	CGGGATCCTTTTCGACGGTCTGCCTTTC
<i>tetQ</i>	SmaI-tetQ-F	TCCCCCGGGCGTTCATTGGCCCTCAAAC
	BamHI-tetQ-R	CGGGATCCCTCTGCCATTATAGAGGC
<i>galE</i> and its downstream	BamHI-down-F	CGGGATCCGAAACCGAAATGAAACAAAAG
	XbaI-down-R	GCTCTAGAAGGACACCGCGCAGCTGGAT
Upstream of <i>galE</i>	KpnI-up-F	GGGGTACCCTCTCCAATGCCAGACTTTG
	SmaI-up-R	TCCCCCGGGCGGTTTCTATTTCGTAGC

^a Restriction enzyme sites incorporated into oligonucleotides for subcloning are italicized.

three times (3,000 × g for 20 min at 4°C) and then lyophilized. LPS was extracted from the same weight (40 mg) of the lyophilized cells by using a hot phenol extraction method (39). The aqueous phase was separated by centrifugation (9,000 × g for 30 min at 4°C) and then collected and dialyzed at 4°C for 12 h against distilled water to remove phenol contamination. To aid in the elimination of nucleic acids, the dialyzed solution was brought to 0.15 M NaCl-NaOH (pH 6.5) and treated with RNase A (0.02 mg/ml; QIAGEN, GmbH, Hilden, Germany) for 2 h at room temperature, followed by the addition of MgCl₂ to 4 mM Mg²⁺, and then treated with DNase I (5 µg/ml; Sigma, St. Louis, MO) for 6 h at room temperature. Next, the solution was dialyzed against distilled water overnight at 4°C. To aid in the elimination of protein, the solution was brought to 30 mM Tris-Cl (pH 8.0) and treated with proteinase K (0.4 mg/ml; QIAGEN) for 1 h at 60°C and dialyzed with distilled water. Finally, the dialyzed solution was centrifuged twice at 100,000 × g for 12 h at 10°C, and the pellet was lyophilized as purified LPS. An equivalent weight of the lyophilized LPS (5 µg) was electrophoresed on a 15% sodium dodecyl sulfate-polyacrylamide gel (17) and subjected to silver staining (38). The band densities of the O-antigen ladder in the wild-type, *galE*, and *galE-c* strains were evaluated using densitometric scanning (CS analyzer 1.0; ATTO Co., Tokyo, Japan).

Biofilm formation assay. Biofilm formation by the *P. gingivalis* wild-type and mutant strains was assayed using a method described previously (20) with some modifications. To produce biofilms, 2 × 10⁷ CFU of *P. gingivalis* in 200 µl of BHI-HM broth (1 × 10⁸ CFU/ml) was added to the wells of 96-well flat-bottom polystyrene microtiter plates (Corning, New York, NY). After the plates were anaerobically incubated at 37°C for 34 and 48 h, planktonic cells in liquid medium were discarded and the plates were washed twice with distilled water. The plates were then air dried, and attached biofilms were stained with 0.25% safranin for 30 min. Then, the plates were rinsed twice with distilled water to remove excess dye and air dried. All dye associated with the attached biofilms was dissolved with 200 µl of 100% ethanol, and then OD_{492/620} absorbance was measured by use of a microplate reader (Multiskan Ascent; Thermo Electron Oy, Vantaa, Finland) to determine the amount of biofilm formation.

Scanning electron microscopy. For scanning electron microscopy (SEM) examinations, *P. gingivalis* wild type or the *galE* strain was developed on nontreated plastic sheets (Wako Chemical Ltd., Osaka, Japan) placed in six-well polystyrene cell culture plates (Corning). *P. gingivalis* (2 × 10⁸ CFU) in 2 ml of BHI-HM broth per well was added and incubated at 37°C under an anaerobic condition. After 12 h, the sheets were rinsed twice with PBS to remove any planktonic cells. Attached cells on the sheets were fixed with 2.5% glutaraldehyde and 2% paraformaldehyde in PBS for 30 min at room temperature and subsequently washed three times in PBS. Then, secondary fixation with 1% osmium tetroxide in PBS was performed. The samples were washed in PBS, dehydrated in 50% ethanol to absolute ethanol, immersed in isoamyl acetate, dried by critical point drying, coated with osmium vapor by an osmium plasma coater, and observed by SEM (S5200; HITACHI Corporation, Hitachi, Japan).

Determination of MIC. Etest benzylpenicillin, oxacillin, ampicillin, cefotaxime, ceftriaxone, imipenem, tetracycline, doxycycline, erythromycin, clindamycin, tobramycin, kanamycin, and vancomycin strips (AB Biodisk, Solna, Sweden) were used to determine MICs. An inoculum of each *P. gingivalis* strain adjusted

to a standard turbidity of McFarland 0.5 was applied to BHI-HM blood agar plates. The Etest strips with continuous gradients of the antibiotics were placed on the surface of the agar. After a 5-day incubation anaerobically, the MICs were determined by the regions where the zones of inhibition intersected the MIC scale on the strips.

Statistical analysis. Statistical analysis was performed using the Mann-Whitney U test. *P* values of 0.05 or less were considered to indicate statistical significance.

Nucleotide sequence accession numbers. The nucleotide sequences of the PG0106 (*wecA*), PG1964 (*wbaP*), PG0347 (*galE*), PG1225 (*wzt*), and PG0045 (*htpG*) regions of *P. gingivalis* W83 strain are listed under accession nos. AAQ65351, AAQ66941, AAQ65558, AAQ66314, and AAQ65296, respectively.

RESULTS

Construction of *P. gingivalis* mutants. To determine whether *P. gingivalis* possesses homologs of *wecA*, *wbaP*, *galE*, and *wzt* genes, we performed BLAST search analysis using the genome sequence entries of *S. enterica* serovar Typhi or *B. thetaiotaomicron* as query sequences. All of those homologs were found in the genome of *P. gingivalis*, and the amino acid sequences corresponding to the *wecA*, *wbaP*, *galE*, and *wzt* genes in *P. gingivalis* showed 25%, 44%, 40%, and 38% identity, respectively, to those of the corresponding genes of *S. enterica* serovar Typhi, and 35%, 43%, 66%, and 52% identity, respectively, to those of *B. thetaiotaomicron*. In order to analyze the biological roles of *wecA*, *wbaP*, *galE*, and *wzt* genes, we constructed each target gene mutant by use of *P. gingivalis* ATCC 33277. Each gene disrupted by insertion of the *ermF-ermAM* cassette was introduced into *P. gingivalis* by electroporation (Fig. 1A). We also generated a *galE*-complemented mutant, the *galE-c* strain, by *cis* replacement of the *ermF-ermAM* cassette-inserted *galE* mutation to the intact *galE* gene with the *tetQ* gene (Fig. 1B and C) in order to confirm that inactivation of the *galE* gene results in the phenotypes of the *galE* strain.

Loss of UDP-galactose 4-epimerase activity and accumulation of intracellular carbohydrates by *galE* mutation. First, we examined the growth of the wild-type and mutant strains. All mutant strains grew to same extent as the parental strain in BHI-HM broth (Fig. 2A). Since it has been reported that *galE* mutants of a number of species accumulated intracellular Gal-

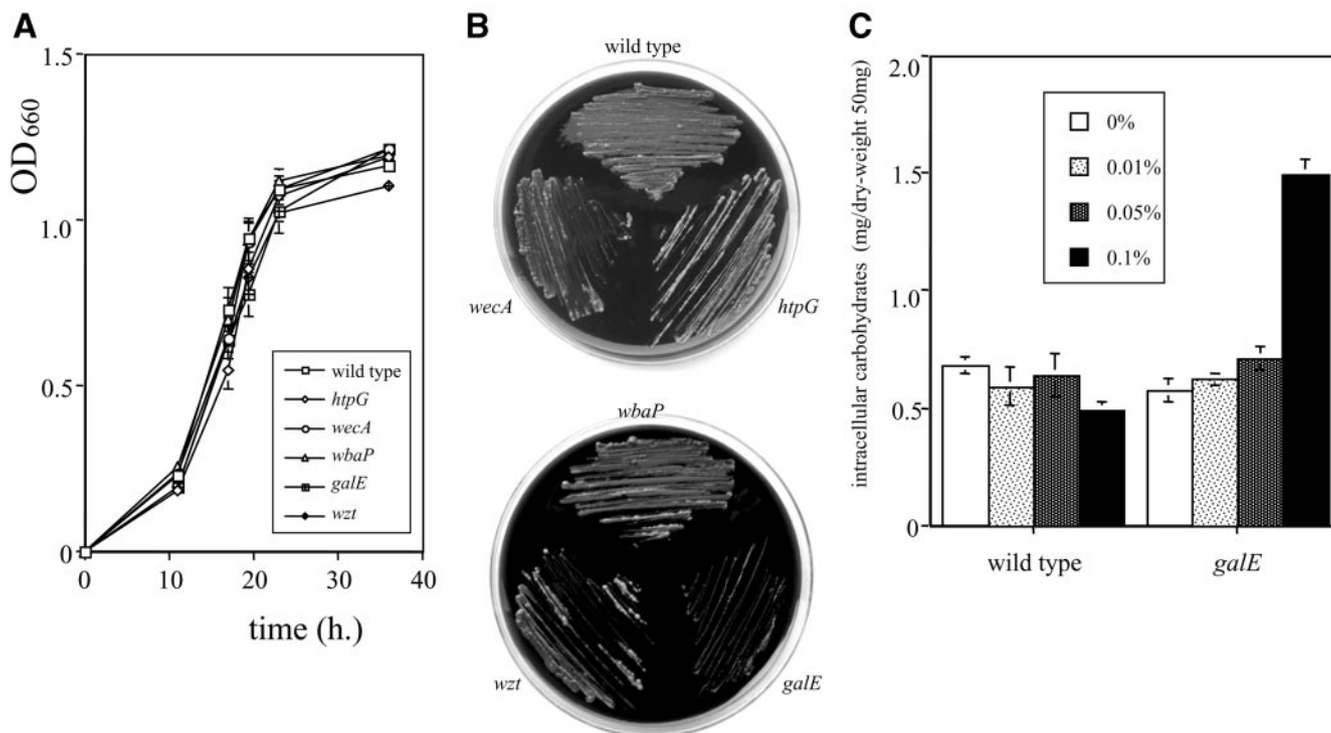


FIG. 2. Growth assays and quantitation of intracellular polysaccharide in the absence or presence of galactose. (A) *P. gingivalis* wild-type and mutant strains were grown in BHI-HM broth, and OD₆₆₀ absorbance was measured at different time points. Data shown are representative of three independent experiments. The results are expressed as the mean \pm standard deviation (SD) from a triplicate assay. Similar results were obtained in three independent experiments. (B) The strains were also grown on BHI-HM agar plates supplemented with 1% galactose. (C) *P. gingivalis* wild-type and *galE* strains were grown in BHI-HM broth supplemented with or without galactose (0%, 0.01%, 0.05%, and 0.1%). Quantities of intracellular polysaccharides, which were extracted from every 50 mg of lyophilized cells, were determined by a phenol-sulfuric acid colorimetric method with glucose as the control. Total intracellular carbohydrates are expressed as mg per 50 mg of dry weight. The results are expressed as the mean \pm SD of a triplicate assay. Similar results were obtained in two independent experiments.

1-P and UDP-Gal in the presence of galactose, leading to bacteriostasis and bacteriolysis (8, 26, 30), we examined their growth in BHI-HM broth supplemented with galactose. Among the mutant strains, only the growth of the *galE* strain was delayed in the presence of more than 0.1% galactose (data not shown). The *galE* strain did not grow in the presence of more than 1% galactose (Fig. 2B). In contrast, the complemented strain, the *galE-c* strain, was able to grow in the presence of 1% galactose (data not shown). Moreover, the *galE* strain accumulated intracellular carbohydrates in large quantities in the presence of 0.1% galactose (Fig. 2C). In order to confirm the defect of GalE activity in the *galE* strain, a UDP-galactose 4-epimerase assay was carried out. As expected, the *galE* strain lost the enzyme activity, while that of the parental strain was active (0 versus 111.8 ± 2.3 nmol/mg protein \cdot min). The *galE-c* strain demonstrated that activity at an intermediate level compared to the wild-type and *galE* strains (6.7 ± 0.6 nmol/mg protein \cdot min).

Modification of LPS O antigen in *wecA* and *galE* strains. We hypothesized that the defect of the proposed genes would have an effect on the synthesis of LPS in this organism. To confirm this hypothesis, we analyzed the LPS profiles of each strain by using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (PAGE) and silver staining. Although the distribution of

the bands of LPS in the *htpG*, *wbaP*, and *wzt* strains scarcely changed relative to those of the parental strain, LPS in both the *wecA* and *galE* strains showed an altered profile (Fig. 3A). A minor shift of the O-antigen ladder was observed for the *wecA* strain; this shift was denoted by *1. For the *galE* strain, though the lowest three bands of O antigen scarcely change, the densities of the O-antigen bands at high molecular mass, from 25 to 35 kDa, nearly disappeared (*2), while those at low molecular mass, from 16 to 19 kDa, became more intense (*3). The *galE-c* strain recovered production of LPS at an intermediate level compared to the wild-type and *galE* strains, as shown in the inset in Fig. 3B. When the densities of the O-antigen ladder bands were quantitated using gel scanning, those of the *galE* strain were higher at a lower molecular mass (Fig. 3B), indicating that the O-antigen length of the *galE* strain was shorter than that of the wild type.

Biofilm formation and SEM analysis. To confirm the involvement of the proposed genes in biofilms organized from *P. gingivalis*, biofilm formation assays were carried out using 96-well microplates. No major change in biofilm mass was observed in any of the mutants, except for the *galE* strain. After a 34-hour incubation, biofilms organized from the *galE* strain showed an intensity more than threefold greater than those from the wild-type strain (Fig. 4A). After a 48-hour incubation,

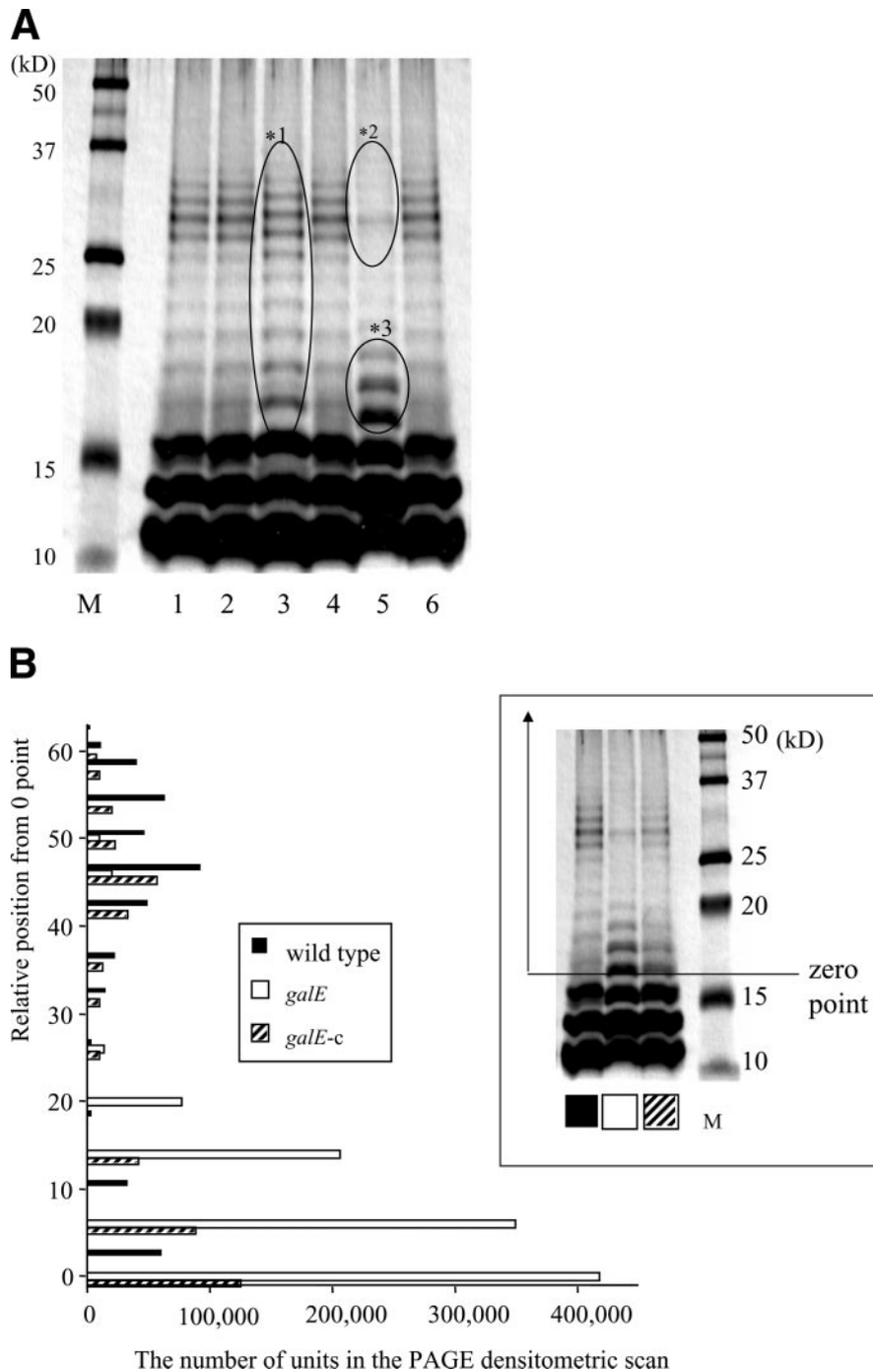


FIG. 3. LPS profiles of *P. gingivalis* wild-type and mutant strains. LPS samples extracted from the *P. gingivalis* wild-type and mutant strains were analyzed by sodium dodecyl sulfate-PAGE and silver staining. Each lane contains 5 μ g of LPS. (A) Lanes: 1, wild type; 2, the *hipG* strain; 3, the *wecA* strain; 4, the *wbaP* strain; 5, the *gale* strain; and 6, the *wzt* strain. M: molecular mass marker. The areas shown with ovals represent a minor shift of the O-antigen ladder (*1) in the *wecA* strain and a decrease of the high molecular band of O antigen (*2) and an increase of the low molecular band of O antigen (*3) in the *gale* strain. (B) The band densities of the O-antigen ladders after silver staining in the wild-type, *gale*, and *gale-c* strains were evaluated using densitometric scanning. The y axis of the histogram indicates the relative position from 0 point, whose position, denoted in the inset, was determined to be approximately 16 kDa.

biofilms related to the *gale* strain showed greater than 4.5 times the mass of the wild-type strain (Fig. 4B). Further, the *gale-c* strain formed biofilms of an intermediate level compared to those of the wild-type and *gale* strains (Fig. 4B). We

also examined the biofilm structure of the *gale* strain by SEM. After a 12-hour incubation on plastic surfaces, there was no difference between the wild-type and *gale* strains in terms of the number of attached cells. However, the *gale* strain was

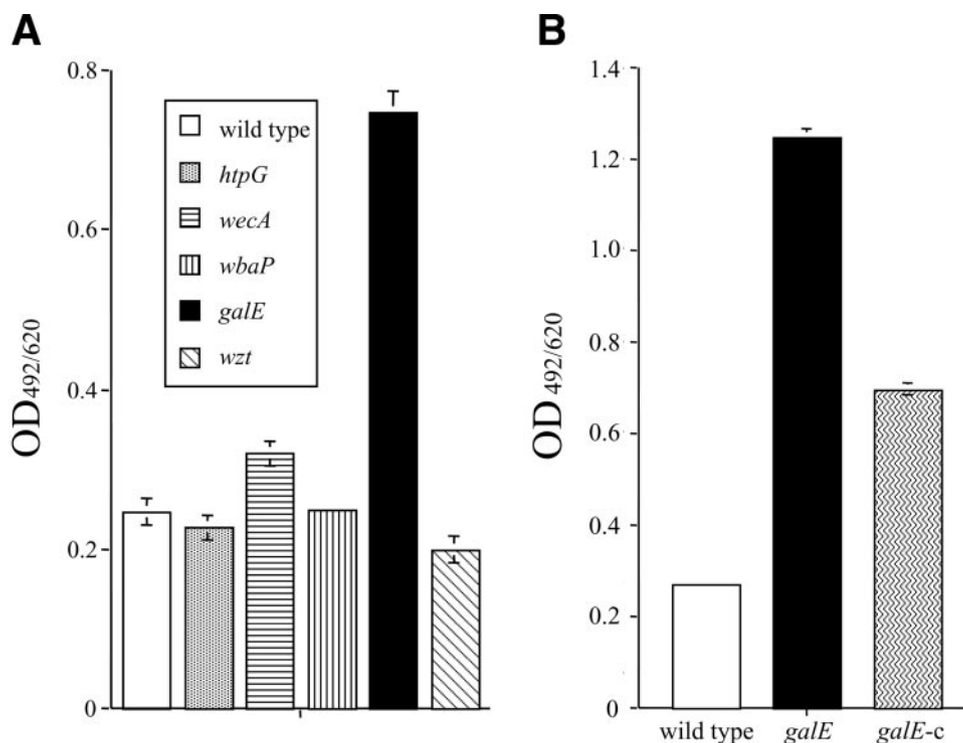


FIG. 4. Biofilm formation by *P. gingivalis* wild-type and mutant strains. Biofilm formation was examined after 34 (A) and 48 (B) hours of culturing in BHI-HM broth. OD_{492/620} absorbance was measured to determine the biofilm mass. Data shown are representative of three independent assays. The results are expressed as the mean \pm SD of a triplicate assay. Similar results were obtained in three independent experiments.

found to have cell-aggregative and long-chain phenotypes, in contrast to the wild type (Fig. 5A and B). After 24 h, the number of attached cells for the *galE* strain was greatly increased compared with that for the parental strain (data not shown).

High sensitivity to antibiotics by *galE* mutation. Etest strips were used to examine whether the *galE* strain had an altered sensitivity to antibiotics. The *galE* strain was found to be significantly more susceptible to benzylpenicillin, oxacillin, cefotaxime, imipenem, and vancomycin than the wild type (Table 3).

DISCUSSION

In the present study, the essential role of the *galE* gene of *P. gingivalis* in LPS biosynthesis was established following PAGE analysis of its LPS (Fig. 3). The O-antigen ladder bands of *P. gingivalis galE* strain were found to be distributed at molecular masses lower than those seen for the parental strain, indicating that the O polysaccharide length of the *galE* strain was decreased. Further, the *galE-c* strain partially restored the synthesis of the O antigen (Fig. 3B). The incomplete recovery observed in our complementary trial might have been dependent on the low level of activity of the *tetQ* promoter in comparison with that of the native promoter, because the GalE activity of the *galE-c* strain was also incomplete in comparison with that of the wild type.

GalE protein catalyzes the interconversion of UDP-glucose to UDP-galactose. In previous studies, the *galE* mutant of

Vibrio cholerae, whose O antigen does not include galactose, did not have an altered LPS profile (26), while *galE* mutants of other bacteria, such as *Neisseria meningitidis* and *Helicobacter pylori*, were found to produce truncated LPS molecules that lacked galactose (16, 19). Bramanti et al. reported that galactose accounted for 25.3% of the total monosaccharides of LPS in *P. gingivalis* strain 33277 (4). Further, Paramonov et al. also showed that the O antigen of *P. gingivalis* strain W50 consists of the tetrasaccharide repeating unit (6)- α -D-Glcp-(1-4)- α -L-Rhap-(1-3)- β -D-GalNAc-(1-3)- α -D-Galp-(1) (31). Therefore, we speculated that the truncated O antigen in the *P. gingivalis galE* strain is a result of the inability of this mutant to utilize UDP-galactose, which is one of the essential building blocks of O antigen in *P. gingivalis*. Further, the *galE* strain was found to exhibit the long-chain phenotype, in contrast to the wild-type strain. This phenotype may be due to a failure of cellular segregation resulting from alteration of the cell surface, which contains a variety of glycoconjugates, such as LPS and peptidoglycan. However, a more precise analysis is required to fully understand the relationship between the *galE* mutation and these effects.

Inactivation of the *galE* gene in *P. gingivalis* also had an effect on biofilm formation (Fig. 4), which may have been due to the fact that galactose is one of the components of the carbohydrate polymer on the surface of *P. gingivalis* (35). Therefore, we considered that the diminished levels of galactose in biofilms may have an effect on the total amount of biofilms and/or chemical behavior, such as hydrophobicity, re-

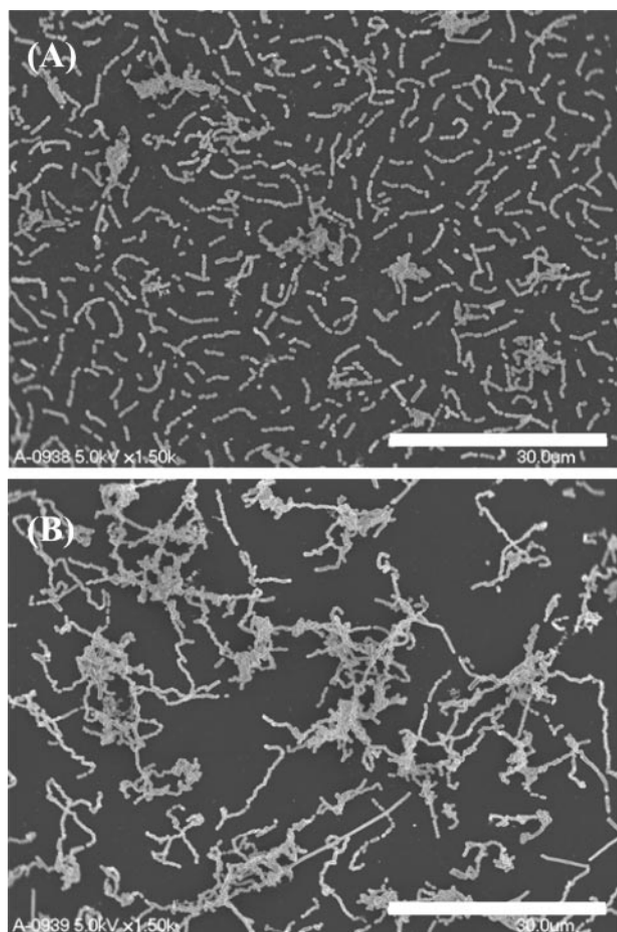


FIG. 5. SEM images of *P. gingivalis* wild-type and *galE* strains. *P. gingivalis* wild-type (A) and *galE* (B) strains were grown on plastic sheets for 12 h, after which attached cells were investigated using SEM. Scale bars are shown at the lower right of each electron micrograph.

sulting in aggregation of the cells (Fig. 5D) and overproduction of biofilms (Fig. 4). Alternatively, the shortened O antigen of the *galE* strain might promote the property of initial attachment to the materials, followed by an increase in biofilm formation. The O antigen on the bacterial surface is located at the interface between the bacterium and its environment. When the length of the O polysaccharide chain as a physical obstacle becomes short on its surface, an essential component of the outer membrane involved in the affinity of the organism to solid surfaces or other organisms may become easily exposed to the environment. This may be true in the case of the *galE* strain of *P. gingivalis*, which was shown to promote biofilm formation. It has also been reported that O-antigen chain length has an effect on the sensitivity of bacteria to bactericidal molecules. For example, a truncated O-antigen mutant of *V. cholerae* raised its sensitivity to complement and cationic peptides (26). Likewise, the increased sensitivity to some antibiotics shown for the *galE* strain (Table 3) might be dependent on the shortened O antigen on its surface.

We also examined the LPS profiles of two glycosyltransferase mutants, the *wecA* and *wbaP* strains, and of an ABC transporter mutant, the *wzt* strain (Fig. 3A). For the *wecA*

strain, a minor shift in the O-antigen ladder was observed; this might have been because *WecA* functions in the first step of O polymerization involved in the formation of lipid-linked GlcNAc by transfer of GlcNAc-P residue to the lipid anchor (24). In contrast, the distribution of the O-antigen ladder in the *wbaP* strain did not change in relation to that of the parental strain. It is possible that there is an alternative gene product that compensates for the loss of function by *WbaP*. For example, PG1135 is paralogous to the *wbaP* gene (PG1964) (42% amino acid identity) and may fulfill the role of *WbaP* in the *wbaP* strain. Further, the defect of the *wzt* gene, which encodes the ABC transporter subunit, also did not have an effect on the LPS profile. In the same way, *P. gingivalis* possesses two paralogs of the *wzt* gene (PG1225), PG2199 and PG2206 (31% and 30% amino acid identities, respectively). Since both PG2199 and PG2206 encode the ABC transporter, they may fulfill the role of *Wzt* in the *wzt* strain. Otherwise, another pathway for O-antigen transport in the *wzt* strain might compensate for the ability. The three currently known pathways for O-antigen transport are distinguished by their respective export mechanisms. The pathways are called ABC transporter dependent, Wzy dependent, and synthase dependent (32). Thus, it is possible that the Wzy-dependent and/or synthase-dependent pathway might be active in *P. gingivalis*.

In conclusion, our results suggest that *P. gingivalis galE* plays an important role in both the synthesis of O antigen and the formation of biofilms, which were found to be closely related with its virulence. Although the effects following the *galE* mutation are insufficient to explain the detailed mechanism, we suggest that the phenotypes related to the *galE* mutation are associated with polysaccharide changes that occur at the bacterial surface. Further studies to elucidate the precise mechanism of these phenomena are in progress. Finally, our conclusive evidence for the relationship between the *P. gingivalis galE* mutation and these virulence factors may provide important clues for understanding the mechanism involved in the progress of periodontal diseases.

TABLE 3. MICs for *P. gingivalis* wild-type, *galE*, and *galE-c* strains

Antibiotic	MIC ($\mu\text{g/ml}$) ^a for:		
	33277 (wild type)	<i>galE</i> strain	<i>galE-c</i> strain
Benzylpenicillin	0.011 \pm 0.0023	0.0047 \pm 0.0012 ^b	0.0073 \pm 0.0012
Oxacillin	0.10 \pm 0.020	0.053 \pm 0.010 ^b	0.084 \pm 0.017
Ampicillin	<0.016	<0.016	<0.016
Cefotaxime	0.0067 \pm 0.0012	0.002 \pm 0.00 ^b	0.006 \pm 0.002 ^c
Ceftriaxone	<0.002	<0.002	<0.002
Imipenem	0.084 \pm 0.035	0.024 \pm 0.0080 ^b	0.043 \pm 0.018
Tetracycline	0.042 \pm 0.0087	0.037 \pm 0.0087	1.7 \pm 0.29 ^d
Erythromycin	<0.016	>256 ^e	<0.016
Clindamycin	<0.016	>256 ^e	<0.016
Kanamycin	>256	>256	>256
Tobramycin	>256	>256	>256
Vancomycin	12 \pm 4	1.8 \pm 0.29 ^b	4 \pm 0 ^e

^a Values are shown as means \pm standard deviations.

^b The *galE* strain had significant difference in sensitivity to the indicated antibiotic compared to the wild-type 33277.

^c The *galE-c* strain had significant difference in sensitivity to the indicated antibiotic compared to the $\Delta galE$ strain.

^d Resistance was dependent on insertion of *tetQ*.

^e Resistance was dependent on insertion of *ermF*.

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