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Involvement of the adc Operon and Manganese Homeostasis in *Streptococcus gordonii* Biofilm Formation

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Pioneer oral bacteria, including *Streptococcus gordonii*, initiate the formation of oral biofilms on tooth surfaces, which requires differential expression of genes that recognize unique environmental cues. An *S. gordonii*:Tn917-lac biofilm-defective mutant was isolated by using an in vitro biofilm formation assay. Subsequent inverse PCR and sequence analyses identified the transposon insertion to be near the 3' end of an open reading frame (ORF) encoding a protein homologous to a *Streptococcus pneumoniae* repressor, AdcR. The *S. gordonii* adc operon, consisting of the four ORFs adcR, adcC, adcB, and adcA, is homologous to the adc operon of *S. pneumoniae*, which plays a role in zinc and/or manganese transport and genetic competence in *S. pneumoniae*. AdcR is a metal-dependent repressor protein containing a putative metal-binding site, AdcC contains a consensus-binding site for ATP, AdcB is a hydrophobic protein with seven hydrophobic membrane-spanning regions, and AdcA is a lipoprotein permease with a putative metal-binding site. The three proteins (AdcC through -A) are similar to those of the binding-lipoprotein-dependent transport system of gram-positive bacteria. Reverse transcriptase PCR confirmed that adcRCB4 is cotranscribed as an operon in *S. gordonii* and that the transposon insertion in *S. gordonii* adcR:Tn917-lac had resulted in a polar mutation. Expression of adcR, measured by the β-galactosidase activity of the adcR:917-lac mutant, was growth phase dependent and increased when the mutant was grown in media with high levels of manganese (>1 mM) and to a lesser extent in media with zinc, indicating that AdcR may be a regulator at high levels of extracellular manganese. A nonpolar inactivation of adcR generated by allelic replacement resulted in a biofilm- and competence-defective phenotype. The biofilm-defective phenotype observed suggests that AdcR is an active repressor when synthesized and acts at a distant site(s) on the chromosome. Thus, the *adc* operon is involved in manganese acquisition in *S. gordonii* and manganese homeostasis and appears to modulate sessile growth in this bacterium.

Oral streptococci (viridans streptococci) initiate the formation of oral biofilms known as dental plaque and are ubiquitous primary colonizers that are the predominant cultivable bacteria found in supragingival plaque (45). Viridans streptococci are also frequent etiologic agents of bacterial endocarditis (21, 64) and are becoming significant opportunistic pathogens and a major cause of bacteremia in immunocompromised patients, accounting for 40% of infections in neutropenic patients (25, 32). Fully developed, surface-attached oral biofilms are highly structured, with distinct architectural and physiochemical properties commonly observed with other biofilm communities (12). Sessile bacteria found in biofilms represent a distinct mode of growth that differs from planktonic cells, characterized by differences in gene expression and cellular physiology and an increased resistance to antibiotics. The multistep process of biofilm formation has just begun to be understood (12). This complex biofilm developmental process is initiated by the attachment of the colonizing bacteria to a surface. Subsequent accumulation and growth of attached bacteria result in microcolonies that increase in size and eventually appear as towering pillar- and mushroom-shaped biofilms. The growth of this complex, multilayered, cellular matrix-embedded community and its eventual dispersal and spread require the coordinated expression of an array of genes (57, 59). These include general housekeeping genes and some global regulatory genes, which have been identified by using genetic analyses of biofilm-defective phenotypes, DNA microarrays, and proteomic analysis (4, 38, 46, 61, 63, 68, 69). Thus, the transition between planktonic and sessile phenotypes does not appear to arise from the expression of unique genes but probably requires a differential pattern of gene expression as the bacteria enter a distinct physiological state. The same attached bacteria can revert to planktonic cells by switching off this coordinated gene expression (59).

Assays that measure the ability of bacteria to form adherent biofilms on abiotic surfaces by staining attached bacterial biofilm with crystal violet or safranin have been used extensively to screen for biofilm-defective mutants. Specific genes involved in single-species biofilms have been identified in a variety of bacteria, including *Escherichia coli* (24, 52, 54), *Staphylococcus aureus* (13), *Staphylococcus epidermidis* (43), *Streptococcus gordonii* (42), *Streptococcus mutans* (68, 71), *Pseudomonas aeruginosa* (48), *Pseudomonas fluorescens* (49), *Salmonella enterica* (44), and *Vibrio cholerae* (67). Identifying and analyzing biofilm-defective mutants resulted in the identification of a number of biofilm-associated genetic loci. These include genetic competence (39, 40, 42), quorum sensing and signaling (4, 39, 40, 42), exopolysaccharide synthesis (27, 70), and specific bacterial surface proteins (22, 36, 52, 55, 67).

A number of transcriptional regulators that are stress inducible and growth phase dependent have been found to modulate...
bacterial biofilm formation. Biofilm formation of *S. epidermidis* was found to require the expression of RsbU, a positive regulator of α, an alternative sigma factor that modulates biofilm formation during environmental challenges by altering the expression of the polysaccharide intercellular adhesin (33). A transcriptional repressor in *S. epidermidis* encoded by icaR is involved in both the environmental regulation of the ica operon and biofilm formation (10). The *E. coli* OmpR/EnvZ two-component regulatory system modulates biofilm formation in response to changes in osmolality through a complex regulatory network that controls expression of curli (53), whereas the Cpx two-component signal transduction system is involved in the attachment of stationary-phase *E. coli* cells to hydrophobic abiotic surfaces (18, 50). PrvR, a putative sensor regulator homolog, modulates antibiotic resistance and biofilm formation of *P. aeruginosa* during phase variation (19). Crc, the global carbon metabolism regulator in *P. aeruginosa* (47), and CcpA, the catabolite control protein in *S. mutans* (68), are necessary during biofilm formation on abiotic surfaces. CytR, a transcriptional repressor that inhibits polysaccharide synthesis in *V. cholerae*, also modulates biofilm development (26). A ToxR homolog that regulates its own synthesis in *Vibrio anguillarum* serotype 01 has been implicated in biofilm formation on glass surfaces (66). The availability of microbial genome sequences has enabled investigators to mutate global transcriptional regulators identified in *sico* and to identify several biofilm-associated genetic loci among these regulators in the oral pathogen *S. mutans* (4, 38–40, 68). Taken together, these studies demonstrated that certain transcriptional regulators play a crucial role in biofilm formation by regulating the expression of biofilm-related genes and reinforced the concept that coordinated gene expression may modulate the transition between the planktonic and sessile phenotypes.

A mutant generated by transcriptional fusion with a reporter gene would represent an important advantage in the study of genes involved in biofilm formation, as it is an important tool for facilitating studies of gene regulation. In order to investigate factors that may regulate the expression of biofilm-associated genes, the gram-positive transposon Tn917, containing the easily detectable promoterless *E. coli lacZ* reporter gene, was used to generate insertion-mediated transcriptional fusions in *S. gordonii* and the mutants generated were screened to identify genes associated with biofilm formation. An in vitro biofilm formation assay (42) was used to screen for biofilm-defective mutants. The use of Tn917-lac (EryR, Gal') (14) enabled the detection of the levels of gene expression under various environmental conditions. The aims of this study were to identify a biofilm-associated gene(s) in *S. gordonii* and to determine the influence of environmental factors on the expression of the gene(s) identified.

**MATERIALS AND METHODS**

**Bacteria, media, and chemicals.** *S. gordonii* Challis 2, the rifampin-resistant (500 μg/ml) parent strain of *S. gordonii* Challis (42), was used as the wild-type (WT) strain. Unless otherwise indicated, bacteria were subcultured and maintained routinely on brain heart infusion (BHI) agar (BBL; Becton Dickinson, Cockeysville, Md.) or Todd-Hewitt broth (Difco Laboratories, Detroit, Mich.) supplemented with 0.2% yeast extract (THBYE) at 37°C under anaerobic conditions (WVR brand anaerobic chamber; VWR, Plainfield, N.J.). *E. coli* strain XLI-Blue MRF' (Stratagene, Cedar Creek, Tex.) was grown at 37°C in Luria-Bertani broth or on Luria-Bertani agar under aerobic conditions.

All chemicals were purchased from Sigma (St. Louis, Mo.) or Fisher Scientific (Pittsburgh, Pa.). All enzymes for DNA manipulations were purchased from Promega (Madison, Wis.) or Fisher Scientific unless stated otherwise. Oligonucleotide primers were from Invitrogen Life Technologies (Rockville, Md.), and their sequences are available in GenBank.

Tn917-lac mutagenesis of *S. gordonii* Challis. *S. gordonii* Challis 2 was transformed with 2 to 3 μg of pTV32-OK (14) by the method described previously (42). The transformants were then plated on BHI agar containing 10 μg of erythromycin (ERY) ml and 500 μg of kanamycin (KAN) ml at 30°C and were incubated for 3 to 5 days under anaerobic conditions. Single-colony transfor-
mants that were recovered on ERY and KAN containing BHI agar at 30°C were used for marker rescue of interrupted genes of *S. gordonii* by using the methods described by Cvitkovitch et al. (14). Briefly, single colonies were inoculated into 5 ml of THBYE containing ERY and KAN and were grown anaerobically for 48 h at 30°C. Independent pools of Tn917 insertions into host chromosomal DNA were generated by a temperature shift to 40°C. Aliquots (5, 10, 25, and 50 μl) of culture were then inoculated into 5 ml of fresh THBYE with no antibiotics and were incubated overnight at 40°C. Following the temperature shift, overnight aliquots were plated on BHI plates containing ERY alone or both ERY and KAN or no antibiotics to estimate the relative efficiencies of transpositions. *S. gordonii* cells in which the Tn917 had transposed into the chromosome and had lost the plasmid were erythromycin resistant (EryR) and kanamycin sensitive (KanR). An equal volume of sterile glycerol was added to the rest of the culture and mixed, and aliquots were frozen at −80°C.

A Tn917-lac library was made by plating the frozen cultures onto BHI plates containing 5-bromo-4-chloro-3-indolyl-b-D-galactopyranoside (X-Gal; 100 μg/ml) and ERY at the selective concentration (10 μg/ml) and was grown at 37°C. After 36 to 48 h of growth, blue (Lac') and white (Lac') colonies were observed on the plates. Each blue, EryR colony was picked and transferred into a well on a 96-well microtiter plate containing 200 μl of BHI and ERY. After mixing of the inoculum, 100 μl of the BHI was transferred to another well on a new microtiter plate, thus creating a duplicate, and plate contents were incubated anaerobically at 37°C for 24 to 48 h. After glycerol was added to a final concentration of 30%, this library was stored at −80°C.

**Biofilm assay.** The in vitro biofilm formation assay was performed in biofilm medium (BM) as previously described (42). In addition to the microtiter plate assay, biofilm formation on borosilicate glass coverslips was visualized directly by phase-contrast microscopy with a Micromaster phase-contrast microscope (Fisher Scientific). Images were captured with a Nikon Coolpix 950 digital camera.

**Southern hybridization.** Chromosomal DNA was isolated from the *S. gordonii* WT and putative biofilm-defective mutants by the method described by Ganeshe-kanumuri et al. (23). DNA was digested with *Hind*III or *Sca*I, separated by agarose gel electrophoresis, and transferred onto a nitrocellulose membrane for Southern hybridization. The probe used for hybridization was pTV32-OK (14) labeled with digoxigenin (DIG) by using the DIG DNA labeling system (Roche Molecular Biochemicals, Indianapolis, Ind.) according to the manufacturer's instructions. After hybridization, the membrane was developed by enzyme immunoassay with the DIG nucleic acid detection kit (Roche Molecular Biochemicals).

**Localization of transposon insertion site and sequence analyses.** The location of the transposon insertion was determined by sequence analysis of the region flanking the transposon. Initially, pBluescript vector and chromosomal DNA from the mutant were digested with *Hind*III, purified by using the Qiagen Nucleotide Removal kit (Qiagen), and sequenced at the Genetics Core Sequencing Facility at Boston University by using a Model 377 Automated Sequencer (Applied Biosystems, Foster City, Calif.). The sequence obtained was compared with sequences in GenBank by using the BLASTX and TBLASTN programs (1) to identify homologous bacterial sequences. Amino acid sequence alignments and phyloge-
netic analyses were performed by using the AlignX program in Vector NTI (Informax Inc., Bethesda, Md.), which utilizes the neighbor-joining algorithm (56). Transmembrane domains were predicted with TMpred software (http://www.irocr.isb-sib.ch/software/TMPRED_form.html).

**RT-PCR of *S. gordonii* WT strain in order to determine the genes that constitute the adh operon, reverse transcriptase PCR (RT-PCR) was performed with total RNA extracted from an *S. gordonii* WT strain grown to mid-log phase (A600

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of 0.3 to 0.4) by using a Qiagen RNeasy mini kit (Qiagen). Primers used for RT-PCR were adcr 5' for and adca 3' rev (specific for a 1,624-bp region that spans adcr to adca), primers adcr P1 and adcr P2 (specific for an 855-bp region that extends from the open reading frame [ORF] encoding a putative transporter to adcr), and adca P3 and copY (specific for a 770-bp region that spans adca to copY). Locations of the primers are shown in Fig. 5A.

RT-PCR was also performed with RNA isolated from both WT S. gordonii and an adcr::Tn917lac mutant to determine whether the mutant generated was polar or nonpolar. This was carried out by using primers adcr for 1 and adcr P2, specific for a 1,624-bp region spanning adcr to adca; and primers adca for 1 and adca rev 2, specific for a 1,449-bp region within adca.

The reaction mixture (25 μl) contained 0.1 μg of template RNA, a 1.2 μM concentration of each primer, 0.2 mM deoxynucleoside triphosphates, 1 mM MgSO4, 2.5 U of avian myeloblastosis virus RT for first-strand DNA synthesis, and 2.5 U of thermostable Tfi DNA polymerase (from Thermus flavus) for second-strand DNA synthesis and DNA amplification (Access RT-PCR system; Promega). The RT reaction was performed at 37°C for 60 min and was then stopped by raising the temperature to 95°C for 2 min. Subsequent PCR amplification was performed under standard conditions, and the RT-PCR products were visualized after 1% agarose gel electrophoresis.

Expression of adcr in different environmental conditions. The β-galactosidase activity of the biofilm-defective S. gordonii adcr::Tn917lac mutant was determined by a fluorometric assay (28) by using methylumbelliferyl-β-d-glucuronide (MUG). Bacteria were grown in 10 ml of THBYE or BM containing ERY at 37°C (unless otherwise indicated) in the anaerobic chamber for 18 h. The conditions tested were BHI, THB, and THBYE without any supplement; THBYE without any supplement grown under aerobic conditions or at 30 or 42°C; THBYE with 0.8% (wt/vol) sugars (fructose, glucose, lactose, maltose, mannose, and sucrose), or 100 mM amino acid (alanine, arginine, aspartate, cysteine, glutamine, glutamate, glycine, histidine, leucine, lysine, methionine, phenylalanine, proline, serine, threonine, tyrosine, or valine). In a subsequent assay, the conditions tested were BM supplemented with various concentrations of manganese (0.1, 0.5, 1, 5, 10, 50, 100, and 200 μM and 1 and 2 mM).

Cells were centrifuged, resuspended in 10 ml of fresh, prewarmed appropriate liquid medium, and grown to log phase (A∞600 of 0.15 to 0.4). Fresh, prewarmed medium (100 ml) was inoculated with 2 ml of log-phase cultures and grown to a log-phase A600 of 0.15 to 0.4 (~3 h). Bacterial growth was recorded (A600), and cells were washed twice with 5 ml of 10 mM Tris, 0.5 mM EDTA, and 150 mM NaCl (pH 8.0) and resuspended in 5 ml of it and sonicated for 2 min. Cells were plated on ice for 5 min and centrifuged, and the clear sonicate was transferred to a new tube. For each sample, 100 μl of the sonicate was transferred to an opalescent microtiter plate, followed by 100 μl of 0.4 M Na2S2O4 in dimethyl sulfoxide. This was mixed well and incubated for 30 min at room temperature. After addition of 200 μl of 4-methyl-UMB (MU) standards (ranging from 0.1 to 3.2 μM in dimethyl sulfoxide), the microtiter plate was read in a fluorescent plate reader (355-nm excitation and 460-nm emission). A Bradford protein assay (Bio-Rad Laboratories, Richmond, Calif.) was performed according to the manufacturer’s instructions to determine protein concentration in each sonicate preparation. The activity was reported as micromolar concentration of MU per microgram of protein per minute.

Mutagenesis of the adcr gene in S. gordonii. PCR ligation mutagenesis with vectorless intermediates (37) was used to construct adcr deletion mutants. The plasmids pDL151 (containing a KAN resistance gene, kan) and pDL152 (containing a spectinomycin resistance gene, spec, and a 100-bp HindIII fragment) were used as templates for amplifying kan and spec, respectively.

Initially, PCR amplifications of the two flanking regions and the antibiotic marker insert were performed with the appropriate primers, which incorporated MluI and XbaI, to enable directional ligation. After electrophoresis on a 1% agarose gel to confirm amplification, the PCR products were purified with the QIAquick PCR purification kit (Qiagen). The purified PCR products of the 5' and 3' flanking fragments were digested with MluI and XbaI, respectively, while the amplified kan and spec cassettes were digested with both enzymes by overnight double digestion at 37°C. After digestion, purification of the digested fragments was performed with the QIAquick Nucleotide Removal kit (Qiagen). Directional ligation was performed at room temperature for 18 h after mixing together 1 μl of the amplified 5' flanking fragment primers adcrP1 and adcrP2, 2 μl of the amplified 5' flanking fragment, 4 μl of the amplified antibiotic cassette, 1 μl of 10x ligation buffer, and 1 μl of T4 DNA ligase in a total volume of 10 μl.

After ligation, 4 μl of the ligated DNA was used for transformation of the S. gordonii WT by the method described previously (42). Transformants were then plated on BHI agar containing the appropriate antibiotics (either 350 μg of KAN/ml or 1,000 μg of spectinomycin/ml) and were incubated at 37°C anaerobically for 2 to 5 days. The competence of the S. gordonii WT strain and adcr::Tn917lac, adcr::Spe, and adcr::Kan' mutants was assessed by transformation with 4 μg of DNA and by enumerating the CFU on solid media containing the appropriate antibiotics. Experiments were done in triplicate.

RT-PCR was performed with RNA isolated from both adcr::Spe' and adcr::Kan' mutants to determine whether the mutant generated was polar or nonpolar. This was carried out using primers adcr for 1 and adcrP2, specific for a 938-bp region spanning adcr to adca, and primers adca for 1 and adca rev 2, specific for a 1,449-bp region within adca. The RT-PCR was carried out as described above. Locations of the primers are shown (see Fig. 5A).

The growth rates of the S. gordonii WT strain and the adcr::Tn917lac, adcr::Spe', and adcr::Kan' mutants were assessed by inoculating the strains from an overnight THBYE culture into fresh 10 ml of THBYE and growing them at 37°C under anaerobic conditions. Growth was quantified by recording the A600 at regular intervals over 24 h.

Nucleotide sequence accession number. The DNA sequence of the 4,175-bp fragment containing the adcr operon has been assigned the GenBank accession number AY177418.

RESULTS

The transposition of Tn917-lac in S. gordonii Challis 2 occurred at a frequency of approximately 10⁻⁵, and approximately 10% of the transformants were also Kan', indicating that plasmid cointegrants occurred at this frequency. A biofilm-defective Tn917-lac mutant was identified after a preliminary screening of 2,650 isogenic mutants generated by transposon mutagenesis with Tn917-lac (Fig. 1A, lanes 1 and 2).

The biofilm assay showed that biofilm formation of this S. gordonii Tn917-lac mutant (A653 ± standard deviation, 1.03 ± 0.53) was reduced by 67% when compared to the S. gordonii WT strain (A653 ± standard deviation, 3.14 ± 0.33). In addition, biofilm development on borosilicate glass coverslips by both strains was examined microscopically at 1, 3, and 24 h after inoculation. After 1 h of incubation, a similar number of cells of the WT strain and Tn917-lac mutant had attached to the coverslip. Fewer cells of the Tn917-lac mutant were present after 3 h of incubation. After 24 h of incubation, the WT strain formed large chains of cells interspersed with areas of less densely packed cells. In contrast, only a few scattered Tn917-lac mutant cells had attached, with large, empty areas present (Fig. 1A, panels 3 and 4).

The transposition was confirmed by Southern hybridization with Tn917-lac, which has a single KpnI site, and two hybridizing bands were predicted. Results from the Southern hybridization (Fig. 1A, lanes 5 and 6) show two bands that hybridized with the DIG-labeled Tn917-lac probe, at 17 and 19 kb. This confirmed that a single transposon insertion occurred in the biofilm-defective mutant.

Inverse PCR performed with genomic DNA that was digested with HindIII and ligated with pBluescript resulted in a PCR product, which was 2.8 kb in size. Sequence analysis of this PCR product containing the region 5' to the transposon insertion revealed that transposition had occurred within an ORF that was homologous to adcr, which encodes a repressor in the adcr operon of Streptococcus pneumoniae (16) and S. mutans (Fig. 1B to D). This gene was not identified when Tn916 mutagenesis was used in a previous screening of biofilm-defective mutants of S. gordonii Challis (42), suggesting...
FIG. 1. (A) Lanes 1 and 2, biofilm formation assay of two S. gordonii::Tn917-lac mutants. Lane 1, a biofilm-positive mutant (growth = A575 of 1.1); lane 2, biofilm-defective adcR::Tn917-lac mutant (growth = A575 of 1.1). Panels 3 and 4, phase-contrast micrographs of biofilm formation on borosilicate coverslips in BM after 24 h by WT (panel 3) and adcR::Tn917-lac mutant (panel 4). Bar = 10 μm. Images represent what was observed in multiple fields. Lanes 5 and 6, Southern hybridization of KpnI-digested DNA from WT (lane 5) and adcR::Tn917-lac mutant (lane 6). DIG-labeled pTV32-OK containing Tn917-lac (which has one KpnI site) was used as the probe. (B to D) Gene organization in the adc operon in various streptococci. Genes from the adc operon are shown in grey. (B) S. gordonii adc operon, adcR (positions 261 to 704) encodes a putative transcriptional repressor for Mn-responsive expression, adcC (positions 701 to 1411) encodes a putative ATP-binding cassette (ABC) transporter, adcB (positions 1404 to 2210) encodes a putative ABC transporter (membrane protein), and adcA (positions 2220 to 3722) encodes a metal-binding lipoprotein, an ABC transporter. The black vertical arrow indicates the position of Tn917-lac insertion in the biofilm-defective mutant. (C) S. pneumoniae adc operon, consisting of adcR, adcC, adcB, and adcA. (D) S. mutans adc operon, consisting of adcR, adcC, and adcB. The adcA gene is present on a different part of the chromosome.
that utilization of \( Tn917 \) has enabled us to identify an additional biofilm-associated gene.

**Genetic organization of the \( adc \) operon.** The region flanking the transposition insertion in \( S. \) gordonii \( adcR::Tn917-lac \) was amplified by PCR. Appropriate primers were designed based on a homologous sequence found on an unfinished and non-annotated sequence of the \( S. \) gordonii chromosome (http://www.tigr.org), and the amplified products were sequenced.

Sequence similarity BLAST searches using the \( S. \) gordonii nucleotide sequence obtained found that the closest homologs of the deduced amino acid sequences were proteins encoded by genes of the \( S. \) pneumoniae \( adc \) operon, consisting of \( adcR, adcC, adcB, \) and \( adcA \) (Fig. 1B and C). This \( adc \) operon is involved in zinc and/or manganese transport and genetic competence of \( S. \) pneumoniae (9, 16). Four \( S. \) gordonii ORFs were identified and designated \( adcR, adcC, adcB, \) and \( adcA \).

The 444-bp \( adcR \) ORF encodes a 148-amino-acid sequence with a predicted molecular mass of 16.7 kDa. The deduced amino acid sequence showed a high degree of homology to bacterial repressor proteins (Table 1; Fig. 2). Amino acid alignment of the \( AdcR \) of \( S. \) gordonii and all its homologs from \( S. \) pneumoniae, \( Streptococcus \) mitis, \( Streptococcus \) equi, \( Streptococcus \) agalactiae, \( S. \) mutans, and \( Streptococcus \) pyogenes and the \( ScaR \) of \( S. \) gordonii showed that \( ScaR \) was a larger protein that was not homologous to \( AdcR \) and did not contain a putative metal-binding region that is present in \( AdcR \) (Fig. 2).

Analysis of the region 3′ from \( adcR \) revealed the presence of three other ORFs that were highly homologous to \( adcC, adcB, \) and \( adcA \) of \( S. \) pneumoniae. The \( adcC \) ORF was 711 bp, starting 1 bp 5′ of the \( adcR \) stop codon. The \( adcC \) ORF was predicted to encode a 237-amino-acid protein with a predicted molecular mass of 27.1 kDa that exhibited high levels of sim-

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**Table 1. Homology of proteins encoded by the \( S. \) gordonii \( adc \) operon and by those of other oral streptococci**

<table>
<thead>
<tr>
<th>Species</th>
<th>% Identity for ( AdcR ) (% similarity)</th>
<th>% Identity for ( AdcC ) (% similarity)</th>
<th>% Identity for ( AdcB ) (% similarity)</th>
<th>% Identity for ( AdcA ) (% similarity)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( S. ) pneumoniae</td>
<td>69 (79)</td>
<td>89 (91)</td>
<td>87 (96)</td>
<td>81 (86)</td>
</tr>
<tr>
<td>( S. ) mitis</td>
<td>70 (80)</td>
<td>71 (73)</td>
<td>NA</td>
<td>65 (68)</td>
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<tr>
<td>( S. ) equi</td>
<td>60 (72)</td>
<td>76 (83)</td>
<td>71 (86)</td>
<td>60 (71)</td>
</tr>
<tr>
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<td>61 (73)</td>
<td>76 (82)</td>
<td>73 (87)</td>
<td>61 (72)</td>
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<tr>
<td>( S. ) mutans</td>
<td>61 (71)</td>
<td>78 (81)</td>
<td>69 (85)</td>
<td>60 (71)</td>
</tr>
<tr>
<td>( S. ) pyogenes</td>
<td>58 (70)</td>
<td>77 (85)</td>
<td>74 (89)</td>
<td>61 (72)</td>
</tr>
</tbody>
</table>

\(^a\) Incomplete sequence. Homology was calculated over 189 amino acids only.

\(^b\) Incomplete sequence. Homology was calculated over 400 amino acids only.

\(^c\) NA, sequence not found in unfinished \( S. \) mitis genome database.

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**Figure 2.** Multiple alignments of \( S. \) gordonii \( AdcR \) deduced amino acid sequence with \( AdcR \) homologs from various streptococci. \( S. \) gordonii \( AdcR \) was aligned with \( S. \) pneumoniae \( AdcR \) (GenBank accession no. CAA96184), the \( AdcR \) homologs from \( S. \) mitis and \( S. \) equi (identified from their respective unfinished genome sequences), \( S. \) agalactiae \( AdcR \) (NP878189), \( S. \) mutans \( AdcR \) (NP722293), \( S. \) pyogenes \( AdcR \) (AAL96906), and \( S. \) gordonii \( ScaR \) (AAF25184) through use of the AlignX program of Vector NTI (Informax). The black vertical arrow indicates the position of the transposon insertion in the \( S. \) gordonii \( adcR::Tn917-lac \) biofilm-defective mutant (pos 110). MBS, histidine-rich putative metal-binding site. Amino acids that are identical and conserved are highlighted in dark grey and light grey, respectively.
It also contains a consensus site for binding to ATP. The adcB ORF was 807 bp in length, starting 5 bp 5′ of the adcC stop codon. It encodes a deduced 269-amino-acid sequence with a predicted molecular mass of 29.2 kDa, with homology to AdcB of *S. pneumoniae* (Table 1). *S. gordonii* AdcB contains seven hydrophobic transmembrane helices (at amino acid positions 13 to 32, 46 to 78, 89 to 107, 133 to 150, 180 to 210, 220 to 240, and 245 to 260). The adcB ORF was followed by adcA, which was 1,503 bp in length, starting 10 bp 3′ of the adcB stop codon. The adcA ORF has a deduced 501-amino-acid sequence, with a predicted molecular mass of 56.4 kDa, and a high level of homology to AdcA, a zinc-binding lipoprotein found in *S. pneumoniae*, and AdcA homologs in other streptococci (Table 1).

Further 3′ of adcA is another ORF, which starts 148 bp from the stop codon of adcA. This ORF encodes a protein with homology to CopY from *S. mutans* (37% identity; 60% similarity), which is a negative transcriptional repressor of the copYZ operon in *S. mutans* (Fig. 1B). Nine base pairs 3′ of copY is copA, which encodes CopA, a 742-amino-acid protein that shares extensive homology with P-type ATPases (65). In *S. mutans*, the copYZ operon encodes a conserved P-type ATPase, which enables the bacterium to tolerate high concentrations of extracellular copper. The expression of the ATPase appears to be regulated by copper via the activity of copY and copZ of the operon (65). Interestingly, in *S. gordonii*, there is no copZ homolog 3′ of copA. Located 5′ of the adcR is an ORF that is divergently transcribed, starting 567 bp 5′ from the start codon of adcR. This ORF has homology to bacterial transporter and/or permease genes (see Fig. 5A).

**Sequence and phylogenetic analyses of *S. gordonii* adc operon.** The predicted amino acid sequences of the proteins encoded by the four adjacent ORFs in *S. gordonii* showed strong homology to the four proteins encoded by *S. pneumoniae* adcR, adcC, adcB, and adcA. Upstream of the adcR start site, a putative ribosomal binding site, putative −10 and −35 sites, and a 12-bp inverted repeat were found (Fig. 3). The inverted repeat region present in the 3′ region of adcR may be where the active repressor binds. A similar region has been observed in the ScaR repressor-binding domain of *S. gordonii*, which was localized to the 3′ region of scaC and is involved in positively regulating the sca operon in the presence of low levels of extracellular manganese (31). The presence of a histidine-rich putative metal-binding domain (DEHEHHHAH) in AdcR indicates that it may also require the binding of a metal for its function. No such histidine-rich metal-binding domain was found in ScaR (Fig. 2). These data suggest that AdcR may act as a metal-dependent repressor in vivo.

The genetic organization, phylogenetic analysis, and alignment of the deduced amino acid sequences of AdcR through -A were examined by identifying homologous sequences from streptococci deposited in the DNA databases (Fig. 4). Sequences encoding similar proteins were first identified with the TBLASTN algorithm (1) and were retrieved from the microbial genome databases (http://www.ncbi.nlm.nih.gov). The deduced amino acid sequences from these putative adcR through -A genes among the different streptococci were then aligned with the sequences from *S. gordonii* for comparison. Multiple-sequence alignment and the resulting phylogenetic tree of ABC metalloregulators indicated that ScaR belongs to the
DtxR-like family of proteins, while AdcR belongs to the MarR-like family of proteins (Fig. 4A). Phylogenetic analyses demonstrate that *S. gordonii* AdcR is more closely related to repressors that modulate manganese transport and that AdcR is distantly homologous to other metalloregulators. Amino acid sequence alignments of AdcC and AdcB from *S. gordonii*, *S. pneumoniae*, *S. mutans*, *S. equi*, *S. agalactiae*, and *S. pyogenes* indicate that they share high levels of homology (data not shown), but the comparison among forms of AdcA from these streptococci showed distinct differences, even though they share high levels of homology (Fig. 4B).

AdcA in *S. gordonii* contains a consensus sequence for the signal peptide cleavage site (LXXC) of the prolipoprotein signal peptide sequence (60). Notably, a histidine-rich motif (HEHGEHEHHHDDYPH) that is a putative metal-binding domain is present in AdcA but not in ScaA of *S. gordonii* (34). In contrast to ScaA, AdcA has an additional C-terminal region that is also present in AdcA proteins in other streptococci (9).

These data suggested that the two distinct genetic systems, the *adc* and *sca* operons, are involved in metal transport in *S. gordonii*.

Phylogenetic analysis of the operon showed that there are significant differences in the organization of the *adc* operon in different streptococci, as determined by sequences from their completed and unfinished microbial genome databases (http://www.ncbi.nlm.nih.gov). The *adcA* genes in *S. gordonii* and *S. pneumoniae* are contiguous with *adcRCB*, whereas the *adcA* genes in *S. mutans* (Fig. 1B to D), *S. pyogenes*, *S. equi*, and *S. agalactiae* (data not shown) are noncontiguous with *adcRCB*. Also, *S. gordonii* AdcA and *S. pneumoniae* AdcA are lipoproteins and contain a consensus sequence for the signal peptide cleavage site in bacterial lipoproteins, whereas the AdcA proteins in *S. pyogenes*, *S. mutans*, *S. equi*, and *S. agalactiae* do not have the typical cleavage site for lipoproteins. However hydrophobicity

**FIG. 4.** Phylogenetic relatedness (dendrogram) constructed on the basis of amino acid sequence similarities by using the AlignX program in the Vector NTI software (Informax), which utilizes the neighbor-joining algorithm (56). The reliability of the topology was estimated by performing 100 bootstrap trials, and the bootstrap values are expressed in percentages at branch points (http://www.genebee.msu.su). Accession numbers follow the species names, and sequences without accession numbers were obtained from the unfinished microbial genome sequences (http://www.ncbi.nlm.nih.gov). (A) Phylogenetic tree of *S. gordonii* AdcR and other bacterial metalloregulatory proteins. (B) Phylogenetic tree of the metal-binding permease AdcA from *S. gordonii* and AdcA from other streptococcal species.
analyses indicate that they are probably secreted proteins (data not shown). The presence of a lipoprotein signal peptide and phylogenetic analysis of the incomplete deduced AdcA protein sequence of \textit{S. mitis} predict that these bacteria fall within the first group, but this can only be verified when the genome sequence of \textit{S. mitis} is completed (Fig. 4B).

RT-PCR with RNA from the WT strain using primers adcR 5’ for and adcA 3’ rev (Fig. 5A), which are specific for a 1,845-kb region that spans the \textit{adcR} and \textit{adcA} intragenic region, yielded a product of approximately 1,845 bp (Fig. 5B, lane 2). This spans the ORFs \textit{adcR}, \textit{adcC}, \textit{adcB}, and \textit{adcA} and coincides with the predicted amplicon size of 1,845 bp. RT-PCR with primers adcR P1 and adcR P2 (Fig. 5A), which are specific for an 855-bp region that spans the putative transporter and/or permease of \textit{adcR} to the \textit{adcR} gene, and primers adcA P3 and copY rev, specific for a 770-bp region that spans \textit{adcA} to \textit{copY}, yielded no PCR products (Fig. 5B, lanes 3 and 4). All the primer pairs used in the RT-PCR above

FIG. 5. RT-PCR analysis of RNA extracted from \textit{S. gordonii} WT and \textit{adcR} mutants. (A) Organization of \textit{adc} operon and adjacent genes, location of primers used, and predicted size of RT-PCR products. Primers that successfully produced an amplicon and their predicted sizes are in boldface. (B) RT-PCR products were studied by using total RNA extracted from WT (lanes 2 to 7), \textit{adcR::Tn917-lac} (lanes 8 to 10), and \textit{adcR::Spec} (lanes 11 to 13) strains as the template. Lanes: 1, 1-kb DNA marker; 2, primers adcR 5’ for and adcA 3’ rev; 3, primers adcR P1 and adcR P2; 4, primers adcA P3 and copY rev; 5, primers adcR for 1 and adcA P2; 6, primers adcC P3 and adcA P2; 7, primers adcA for 1 and adcA rev 2; 8, primers adcR for 1 and adcA P2; 9, primers adcC P3 and adcA P2; 10, primers adcA for 1 and adcA rev 2; 11, primers adcR for 1 and adcA P2; 12, primers adcC P3 and adcA P2; 13, primers adcA for 1 and adcA rev 2.
were also used in standard PCR amplifications with *S. gordonii* WT DNA, which yielded PCR products with the predicted sizes. These results demonstrate that *adcR*, *adcC*, *adcB*, and *adcA* are cotranscribed as a single operon, while *copY* and *copA* are probably transcribed as a separate operon. Therefore, the *adc* operon consists of the four ORFs *adcR*, *adcC*, *adcB*, and *adcA* only.

No amplified products were obtained from RT-PCR with primers spanning *adcR* to *adcA* or *adcC* to *adcA* or with primers within the *adcA* gene with RNA isolated from the *S. gordonii* *adcR::Tn917-lac* mutant, whereas all three primer pairs yielded an RT-PCR product of the predicted size when RNA from the WT strain was used, indicating that, as expected, the transposition resulted in a polar mutation and transcription did not arise from any alternative downstream promoter (Fig. 5B). All assays were performed in triplicate, and mean values and standard deviations are shown.

**Regulation of *adcR* expression in response to changes in environmental conditions.** When *S. gordonii* *adcR::Tn917-lac* was grown on solid media containing X-Gal, a blue colony phenotype was produced, indicating that the Tn917-lac transposon was inserted as an in-frame fusion within *adcR*. Therefore, it was possible to identify and examine environmental factors that regulate the expression of *adcR* by measuring the β-galactosidase activity of *adcR::Tn917-lac* grown in medium supplemented with a variety of nutrients and under various conditions. Since the *adcR* in *S. pneumoniae* was reported to be involved in zinc and/or manganese transport (9, 15), the role of *adcR* in *S. gordonii* was examined by using the transcriptional fusion present in this *adcR::Tn917-lac* biofilm-defective mutant. Results from β-galactosidase assays show that the expression of *adcR* was increased significantly in the presence of 10 mM manganese and was increased to a lesser extent by the presence of 10 mM zinc (Fig. 6). Initial screening of the *adcR::Tn917-lac* mutant found that growth in BHI induced the expression approximately threefold compared to growth in THBYE. Expression was higher when cells were grown in THBYE at 42°C, under aerobic conditions, and in THBYE supplemented with manganese, zinc, alanine, arginine, glycine, or isoleucine (Fig. 6). These results identified manganese, zinc, arginine, glycine, alanine, and isoleucine and growth at 42°C and aerobic growth at 37°C as potential inducers of *adcR* expression.

There was minimal increase in β-galactosidase activity when cells were grown in media supplemented with 10% saliva, human serum, or THBYE. These conditions may simply reflect the concentration of extracellular manganese in saliva (50 to 176 μg/liter [35]), serum (0.1 to 2.9 μg/liter [35]), and THBYE (not known). The explanation for the enhanced β-galactosidase activity when *adcR::Tn917-lac* was grown in the presence of certain amino acids at 42°C and under aerobic conditions is unclear at this time.

Subsequent β-galactosidase activity assays were performed with *adcR::Tn917-lac* grown in the minimal, chemically defined BM supplemented with increasing concentrations of manganese. There was no increase in activity when the manganese concentrations were low (50 to 200 μM). However, when the concentration of manganese was increased to above 1 mM, *adcR* expression was induced, as indicated by an approximately twofold increase in β-galactosidase activity (data not shown). These results show that the *adc* operon is probably involved in the transport and recognition of high concentrations of extracellular manganese and suggest that *adcR* may be a transcriptional regulator. This is in contrast to the *sca* operon in *S. gordonii*, which is also involved in manganese homeostasis. The expression of *scaR* was induced at an extracellular manganese concentration of 0.1 μM and was repressed when the concent-
Biofilm formation of the adeR mutant became visible only after 48 h. PCR concentration was increased to 50 μM, suggesting that the scaR gene in the sca operon encodes a positive regulator in S. gordonii (31).

Several studies have suggested that the biofilm-associated phenotype is induced by slow growth and under duress (11, 41, 70). When the β-galactosidase activity of adeR::Tn917-lac was assayed at different growth phases, adeR was found to be expressed only during the late exponential phase of growth (Fig. 7).

**Construction and characterization of an adeR insertion mutation.** In order to clarify the role of adeR in producing the biofilm-defective phenotype identified in adeR::Tn917-lac, a nonpolar insertional inactivation of adeR was done by allelic exchange with an antibiotic cassette using PCR ligation mutagenesis (37). The adeR gene (from amino acids 13 to 144) on the S. gordonii chromosome was replaced with either spec or kan, which encodes resistance to spectinomycin or KAN, respectively (see Materials and Methods). In both cases, transformation with the ligation mixture of PCR products resulted in viable mutations. Transformants were visible on BHI agar after 24 h of growth under anaerobic conditions for the adeR::spec' mutant. The adeR::kan' mutation became visible only after 48 h. PCR confirmed that integration of the spec or kan gene in adeR in the chromosome had occurred as predicted (data not shown).

The effects of insertional inactivations on biofilm formation and competence were assessed. Biofilm formation of the adeR::spec' mutant (A_{575} ± standard deviation = 1.07 ± 0.07) and the adeR::kan' mutant (A_{575} ± standard deviation = 0.84 ± 0.10) was similar to that of the adeR::Tn917-lac mutant (A_{575} ± standard deviation = 1.03 ± 0.53), confirming that the adeR is involved in biofilm formation. All three adeR mutants were defective in biofilm formation when compared to the WT strain (A_{575} ± standard deviation = 3.13 ± 0.33). The biofilm formation of both the adeR::spec' and adeR::kan' mutants observed with phase-contrast microscopy was similar to that of the adeR::Tn917-lac mutant (data not shown). Results from transformation of the WT, adeR::Tn917-lac, adeR::spec', and adeR::kan' strains with equal amounts of ligated DNA (amplified adeB flanking regions ligated with either the Spec' or Kan' cassette) demonstrate that competence was defective in the adeR::Tn917-lac, adeR::kan', and adeR::spec' mutants. Therefore, a functional ade operon is probably required for competence in S. gordonii.

In growth experiments, the growth rates and final yields of the adeR::spec' and adeR::kan' mutants were found to be similar to those of the WT strain, indicating that these mutations did not affect the growth rate or yield. In contrast, the adeR::Tn917-lac mutant grew at a lower rate than did the WT strain and the two nonpolar adeR mutants, indicating that a polar mutation in adeR reduced the growth rate. However, the final yield of the adeR::Tn917-lac mutant after 24 h appeared to be the same as that of the WT, adeR::spec', and adeR::kan' strains (data not shown).

When RT-PCRs with primers specific for regions spanning adeR to adeA, adeC to adeA, and within the adeA gene (Fig. 5A) were performed with RNA isolated from the WT or
**DISCUSSION**

In order to understand the environmental cues and the resulting phenotypic responses that influence *S. gordonii* biofilm formation and regulate biofilm-associated genes, the gram-positive transposon Tn917, which contains the promoterless *E. coli* lacZ reporter gene (14), was used to isolate and analyze a biofilm-defective gene. Molecular analyses of a biofilm-defective *S. gordonii* WT DNA, which produced PCR products with the predicted sizes (data not shown). These results indicate that both *S. gordonii* *adcR::Spe* and *adcR::Kan* mutants are nonpolar. These results suggest that the repressor AdcR is probably synthesized as an active repressor de novo, because both polar and nonpolar mutations of *adcR* resulted in a biofilm-defective phenotype. If it was an inactive repressor, then there would not be a change in phenotype. Since the data suggest that AdcR is synthesized as an active repressor, it may also act on a remote region(s) on the *S. gordonii* chromosome and modulate biofilm formation on abiotic surfaces by an unknown mechanism.

Metal ion homeostasis in bacteria is generally controlled during uptake by transcriptional regulators and is mediated by ABC-type permeases (9, 29). When the possible regulation of *adcR* by various environmental factors was examined, two- to fourfold increases in β-galactosidase activity were observed when *adcR::Tn917-lac* was grown in the presence of 1 to 10 mM manganese when compared to growth with 100 μM manganese present in BM. Moderate increases in β-galactosidase activity were also observed when the *adcR::Tn917-lac* mutant was grown in the presence of zinc or some amino acids, in BHI, under aerobic conditions, or at 42°C. These results indicate that the *adcR* gene product plays an adaptive role in *S. gordonii* biofilm formation, probably by modulating *adcR* expression in response to specific environmental conditions. Moreover, the probable specificity for manganese is supported by the observation that, in *adcR::Tn917-lac* mutant, manganese is a more potent activator of *adcR* expression than zinc.

Although the *S. pneumoniae* *adc* operon was reported to be involved in genetic competence and metal ion transport, specifically zinc (9, 16), our studies indicate that the *adc* operon in *S. gordonii* is mainly involved in the transport of manganese and competence, but actual transport experiments are required to support our hypothesis. Conclusions from studies of the *S. pneumoniae* *adc* operon were based on the slow growth of an *adcC* mutant in a chemically defined medium when compared to that of the WT strain and the restoration of growth of the mutant to WT levels by the addition of 0.8 μM ZnSO₄ or ZnCl₂. Interestingly, the growth of the *adcC* mutant was inhibited by the addition of 6 μM MnSO₄, and this inhibition was suppressed by the addition of ZnSO₄ or ZnCl₂ (15). As the disruption of the *adc* operon in *S. pneumoniae* was made in *adcC*, the precise role of metal transport in the *adcC* mutant may have been obscured. On the other hand, the *adc* operon in *S. gordonii* appears to be involved in regulating gene expression primarily in the presence of high extracellular levels of manganese and to a lesser extent of zinc. More importantly, phylogenetic analysis demonstrates that AdcR is more closely related to repressors that modulate manganese transport, and these proteins are only distantly homologous to other metalloregulators.

Phylogenetic analysis, amino acid alignments, and the genetic organization of *adc* operons in a number of streptococci for which DNA sequences are available in the microbial genome databases indicate that there are at least two distinct groups. In one group, all four genes are contiguous and co-transcribed as an *adcRCBA* operon and the AdcA permease is a lipoprotein. *S. gordonii* and *S. pneumoniae*, both naturally transformable bacteria, are in this group. *S. mitis* may also be a member of this group, as AdcA is also a lipoprotein, but whether *adcA* is contiguous with the *adc* operon can only be determined when sequences of this region are determined. In the second group, *adcC* is noncontiguous with the *adc* operon and AdcA is not a lipoprotein but still contains a putative signal peptide. This group contains *S. pyogenes*, *S. agalactiae*, *S. equi*, and *S. mutans*.

Several studies have shown that the expression of biofilm-associated genes is growth phase dependent (11, 70). Similarly, expression of *adcR* was only observed during the later stages of the exponential growth phase in *S. gordonii*. In *E. coli*, RpoS mediates the synthesis of more than 30 proteins during the transition from the exponential to the stationary phase of growth, starvation, and osmotic and oxidative stress (41). Both *adcR* and *rpoS* were expressed during the late exponential phase, but the *adcR* mutant has a biofilm-defective phenotype, while a mutation in *rpoS* in *E. coli* resulted in enhanced biofilm formation (11). The data suggest that AdcR in *S. gordonii* is a positive regulator of biofilm-associated genes, in contrast to RpoS in *E. coli*, which is a negative regulator of biofilm-associated genes (11). This difference may explain the different phenotypes exhibited by these two mutants.

Bacteria have developed sophisticated acquisition systems to scavenge essential metals from the environment, including constitutively expressed or inducible low- and high-affinity transport systems for chelated or free metals (29). Manganese is an essential cofactor for a variety of enzymes, some of which are critical for bacterial growth and survival under oxidative stress, and is an important cofactor for enzymatic antioxidant defenses of bacteria, such as catalase and superoxide dismutase (8, 29, 30, 51). A number of proteins, such as catalases,
oxidoreductases, and transferases have evolved a natural preference for manganese, using this metal for a regulatory or catalytic role. Manganese is also a tightly bound constituent of a few metalloenzymes involved in the metabolism of nitrogen and oxygen and in the modulation of signal transduction pathways (8, 29, 30).

**S. gordonii** has at least three different genes that are involved in manganese homeostasis. The *sca* and *adc* operons appear to be high-affinity manganese transport systems that recognize low and high concentrations of extracellular manganese, respectively, and are necessary for growth, genetic transformation, and biofilm formation (31, 34). Although both *sca* and *adc* operons are ABC solute-binding operons that share a high degree of homology, there are distinct differences between ScaA and AdcA, the lipoproteins responsible for solute binding. ScaA does not contain the histidine-rich metal-binding domain that is present in AdcA. The *adcR* gene is the first gene in the *adc* operon upstream of *adcA*, while the *scaR* gene is distant from the other genes of the *sca* operon elsewhere in the chromosome (31). Phylogenetic analysis revealed that *scaR* and *adcR* belong to two distinct classes of regulatory proteins that regulate manganese transport in *S. gordonii*. While there is clear evidence that Sca-like proteins are implicated in bacterial virulence in animal model studies (6), similar studies have not been done to demonstrate the role of Adc-like proteins in pathogenesis. Clearly these two operons are involved in biofilm formation and genetic competence. Although there is evidence to suggest that a third, unidentified manganese transport system exists in *S. gordonii* and is mediated by proton-dependent ATPase (34), whether this plays a role in biofilm formation, genetic competence, or virulence is yet to be determined. However, mutation in *mntH*, which encodes a P-type ATPase that is involved in manganese transport, does not affect virulence of *Mycobacterium tuberculosis* in a mouse model of tuberculosis (17), suggesting that not all manganese transport systems in bacteria are involved in virulence and perhaps in biofilm formation and competence.

The *adcR* gene is cotranscribed with the rest of the binding-lipoprotein-dependent transport genes *adcCBA*, and both a polar mutation and a nonpolar mutation of *adcR* have been shown to be biofilm defective, indicating that AdcR is an active repressor when synthesized de novo. Also, the biofilm-defective phenotype of the nonpolar mutation of *adcR* strongly suggests that the active *adcR* gene acting on a remote region(s) is essential to the biofilm phenotype observed on an abiotic surface. In addition an active repressor could abolish the transcription of the *adc* operon when a sufficient amount of AdcR is synthesized, thus regulating its own synthesis. Only when a sufficient amount of manganese is acquired by other manganese transport systems can the AdcR may be made inactive and can *adc* transcription be continued. Since the *adc* operon responds to high levels of extracellular manganese, *S. gordonii* can acquire the manganese through other transport genes (ScaA and P-type ATPase). Once AdcR is made inactive, the functional *adc* operon can respond to the high level of extracellular manganese.

Why would *S. gordonii* possess a system to sense a high level of manganese when its natural (saliva-bathed oral cavity) or its opportunistic habitat (human serum) is considered to be low in manganese (29, 30)? It may be that, when a bacterium like *S. gordonii* encounters high level of extracellular manganese, it might be an indication for it to switch off biofilm-associated genes through the inactivation of the AdcR repressor protein. Therefore, the AdcR/adc operon couple could be a sensory mechanism to enable the switch from a sessile to a planktonic phenotype, thus facilitating the dispersal and spread of the bacterium. This type of modulation by manganese may be involved in the pathogenicity of *S. pneumoniae* (3), *Streptococcus parasanguinis* (6), *S. enterica* (5) and *Yersinia pestis* (2), all of which have conserved, ABC-type binding-protein-dependent, manganese acquisition systems that are implicated in virulence.

It is of interest that recent studies have identified metal ions as modulators of bacterial differentiation and growth. In addition to manganese, studies have shown that lactoferrin, a component of the innate immunity of mucosal surfaces in humans, chelates iron, which blocks biofilm development by the opportunistic pathogen *P. aeruginosa* (58). Likewise, the recently discovered autoinducer (AI-2) that is produced by numerous bacterial species allows bacterial populations to coordinate gene expression in a variety of developmental processes, such as bioluminescence, virulence, antibiotic production, and biofilm development, and is a furanosyl borate diester, indicating a potential role for boron in bacterial differentiation (7).

The role of manganese homeostasis in *S. gordonii* Challis biofilm formation provides insight into how metal-binding genes may be involved in biofilm differentiation and is a significant indicator of how changes in the environment can elicit a developmental change in bacteria. Since *S. gordonii* resides in environments such as human saliva or serum, which are low in manganese, it is conceivable that there may be other key extracellular cues, such as the presence of other metal ions that may regulate biofilm formation and virulence in *S. gordonii*. Blocking such transporter functions in bacteria may be an effective therapeutic strategy in controlling bacterial biofilms such as dental plaque and opens new avenues for therapy that do not involve immunological intervention such as vaccination.

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**REFERENCES**


