Transcriptional Profiling Reveals the Importance of RcrR in the Regulation of Multiple Sugar Transportation and Biofilm Formation in *Streptococcus mutans*

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ABSTRACT  The ability of *Streptococcus mutans* to survive and cause dental caries is dependent on its ability to metabolize various carbohydrates, accompanied by extracellular polysaccharide synthesis and biofilm formation. Here, the role of an *rel* competence-related regulator (RcrR) in the regulation of multiple sugar transportation and biofilm formation is reported. The deletion of the *rcrR* gene in *S. mutans* caused delayed growth, decreased biofilm formation ability, and affected the expression level of its multiple sugar transportation-related genes. Transcriptional profiling revealed 17 differentially expressed genes in the *rcrR* mutant. Five were downregulated and clustered with the sugar phosphotransferase (PTS) systems (mannitol- and trehalose-specific PTS systems). The conserved sites bound by the *rcrR* promoter were then determined by electrophoretic mobility shift assays (EMSAs) and DNase I footprinting assays. Furthermore, a potential binding motif in the promoters of the two PTS operons was predicted using MEME Suite 5.1.1. RcrR could bind to the promoter regions of the two operons in vitro, and the sugar transporter-related genes of the two operons were upregulated in an *rcrR*-overexpressing strain. In addition, when RcrR-binding sites were deleted, the growth rates and final yield of *S. mutans* were significantly decreased in tryptone-vitamin (TV) medium supplemented with different sugars, but not in absolute TV medium. These results revealed that RcrR acted as a transcription activator to regulate the two PTS systems, accompanied by multiple sugar transportation and biofilm formation. Collectively, these results indicate that RcrR is a critical transcription factor in *S. mutans* that regulates bacterial growth, biofilm formation, and multiple sugar transportation.

IMPORTANCE  The human oral cavity is a constantly changing environment. Tooth decay is a commonly prevalent chronic disease mainly caused by the cariogenic bacterium *Streptococcus mutans*. *S. mutans* is an oral pathogen that metabolizes various carbohydrates into extracellular polysaccharides (EPSs), biofilm, and tooth-destroying lactic acid. The host diet strongly influences the availability of multiple carbohydrates. Here, we showed that the RcrR transcription regulator plays a significant role in the regulation of biofilm formation and multiple sugar transportation. Further systematic evaluation of how RcrR regulates the transportation of various sugars and biofilm formation was also conducted. Notably, this study decrypts the physiological functions of RcrR as a potential target for the better prevention of dental caries.

KEYWORDS  *Streptococcus mutans*, transcription regulator, sugar transportation, biofilm formation, dental caries
Dental caries is a prevalent and consequential oral disease worldwide (1). In 2016, caries of permanent teeth was among the top 10 causes of the most pervasive disorders (2). Tooth decay arises from biofilms containing cariogenic microorganisms that break down fermentable carbohydrates into organic acids. These acids subsequently cause demineralization and destruction of the tooth tissue (3). Streptococcus mutans is a primary cariogenic pathogen that utilizes glucosyltransferases (GtfB, GtfC, and GtfD) to synthesize extracellular polysaccharides (EPSs) from sucrose (4), thereby providing adhesive sites for the colonization of other microorganisms. This occurrence leads to the formation of cariogenic biofilms (5). Keeping this in mind, EPS synthesis and biofilm formation by S. mutans are crucial virulence factors in the pathogenesis of dental caries.

Sugar transportation and metabolism play significant roles, such as EPS synthesis, biofilm formation, and lactic acid production, in the physiology and virulence of S. mutans (6). There are two primary sugar incorporation systems reported in S. mutans, phosphotransferase (PTS) systems and non-PTS systems (6). These systems are responsible for binding, transmembrane transportation, and phosphorylation of various sugar substrates. Meanwhile, they also regulate various metabolic and transcriptional processes and virulence factors, such as growth, energy metabolism, competence, EPS production, and biofilm formation (6–10). An increasing number of studies have shown that genes coding for sugar transportation and metabolism in S. mutans are regulated by transcription factors, sigma factors, and two-component regulatory systems (11, 12). Recently, some transcription factors involved in sugar metabolism have been most studied in S. mutans. For example, TreR, CelR, and StsR act as transcriptional activators that regulate the trehalose operon, cellobiose operon, and multiple transporter genes, respectively (12–14). However, there are also negative regulators. For instance, FruR and NigR negatively regulate the transcription of the fruRki operon and the PTSbio operon, respectively (9, 15).

Here, a new function of rel competence-related regulator (RcrR; SMU_921) was discovered. RcrR is a multiple antibiotic resistance regulator (MarR) family transcription factor in S. mutans UA159. Previously, RcrR has been demonstrated to play critical roles in environmental stresses, genetic competence, and (p)ppGpp metabolism in S. mutans (16). In this study, the loss of rcrR influenced bacterial cell growth, multiple sugar transportation, and biofilm formation. Further investigations via mRNA sequencing of an rcrR mutant revealed that five genes in two PTS systems (mannitol- and trehalose-specific PTS systems) that were involved in sugar transportation were significantly downregulated. Thus, our findings indicate that RcrR is the transcription activator of two PTS systems that control multiple sugar utilization and biofilm formation.

**RESULTS**

**Deletion of rcrR causes growth delay in S. mutans.** SMU_921 (rcrR) can be cotranscribed with SMU_922 (rcrP) and SMU_923 (rcrQ). As such, they are annotated as transmembrane ATP-binding cassette (ABC) transporters in a polycistronic operon (16). Studies have previously focused on RcrR regulation of rcrP and rcrQ expression and physiological functions (16). Here, we found that RcrR possesses other important functions. Growth delay of the rcrR mutant (ΔrcrR) was observed regardless of the nutritional condition (brain heart infusion [BHI], 1/2 BHI, and 1/4 BHI media) (Fig. 1; $P < 0.05$). Meanwhile, the ΔrcrR mutant showed an extended lag phase in the 1/2 BHI medium compared with those of the wild-type and complement strains (Fig. 1B). Nonetheless, the cells of all strains reached the same optical density at the stationary phase. These results indicated that rcrR deficiency affected the growth of planktonic bacteria.

**Effect of rcrR deficiency on the growth of S. mutans with different sugar supplementations.** Based on the findings of growth delays caused by rcrR deficiency in different nutritional conditions (Fig. 1), it was intriguing to speculate on a specific relationship between rcrR and multiple sugar utilization. The growth curves of the wild-type, the ΔrcrR, and the complement strains were thus measured when incubated in TV medium supplemented with common sugars, including glucose, galactose,
lactose, fructose, maltose, and trehalose (1% [wt/vol]), to verify this phenomenon (12, 17). All ΔrcrR mutants had a significantly extended lag phase and decreased final yield compared with those of the wild-type and the complement strains (Fig. 2; \( P < 0.05 \)). The different growth rates and final yield showed that the ability to utilize diverse sugars was also dramatically different between \( S. mutans \) strains. The growth rate and final yield of complement strains incubated in TV medium plus galactose/lactose also gradually decreased (Fig. 2B and C). These results also illustrated that \( rcrR \) deficiency had an important effect on the bacterial growth of \( S. mutans \) with different sugar supplementations. \( S. mutans \) does not grow in TV medium without carbohydrate supplementation (18). It is possible that \( rcrR \) deletion inhibited the transportation or metabolism of multiple sugars of \( S. mutans \), indirectly affecting bacterial growth.

**Biofilm formation of \( S. mutans \) is susceptible to \( rcrR \) deletion.** The ability to form a sucrose-dependent biofilm is regarded as one of the most important virulence
factors in *S. mutans*. Biofilms of the wild-type, the ΔrcrR, and the complement strains were further quantified by crystal violet staining assays (Fig. 3A and C). The ΔrcrR biofilms were significantly reduced compared to the biofilms of the wild-type and complement strains when cultured in BHI plus 1% sucrose (BHIS) or TV plus 1% sucrose (TVS) medium (Fig. 3A and C; *P* < 0.05). The total CFU from biofilms were then counted, and we found that the CFU from the ΔrcrR mutant were also decreased compared to those from the wild-type and complement strains (Fig. 3B and D; *P* < 0.05). These results demonstrated that the deletion of the rcrR gene led to the reduction of bacterial growth and biofilm formation, suggesting the decline in biofilm formation ability was due to a decrease in the number of bacteria.

The 24-h time point was selected to visualize the biofilm using a confocal laser scanning microscope (CLSM) after culturing in BHIS or TVS medium (Fig. 4A or D). The representative three-dimensional images of bacteria (green) and EPS (red) in *S. mutans* biofilms are shown in Fig. 4. In the BHIS medium, the ΔrcrR biofilm contained fewer bacteria (Fig. 4A, upper, green) than those observed in wild-type and complement strain biofilms. However, EPS (Fig. 4A, middle, red) showed no apparent differences, which is consistent with fluorescence quantitation of total bacterial and EPS biomass (Fig. 4B; *P* < 0.05). Meanwhile, the ratio of bacteria to EPS in ΔrcrR biofilm was significantly lower than that in wild-type and complement strain biofilms (Fig. 4C; *P* < 0.05). In the TVS medium, the biofilm formed by the ΔrcrR mutant contained fewer bacteria (Fig. 4D, upper, green) and less EPS matrix (Fig. 4D, middle, red) than those of wild-type and complement strains, similar to the quantitative analysis of total bacteria and EPS.
EPS biomass (Fig. 4E; $P < 0.05$). However, there were no significant differences in the ratio of bacteria to EPS among these strains (Fig. 4F). In line with the results shown in Fig. 3, these data also indicated that bacterial growth (CFU) and biofilm formation of *S. mutans* were susceptible to the deletion of *rcrR*, regardless of the nutritional conditions (BHIS or TVS).

**Transcriptomic analysis of the *S. mutans ΔrcrR* mutant.** Transcriptome analysis was performed to examine the gene expression in the *rcrR* mutant strain. There were 17 differentially expressed genes (DEGs) between UA159 and the *rcrR* mutant (Fig. 5; see also Table S3 in the supplemental material; $P < 0.05$). Of these, 12 genes were significantly upregulated, and 5 genes were downregulated in the *rcrR* mutant compared to the wide-type strain (Fig. 5 and Fig. 6A and B). These genes were mainly associated with posttranslational modifications, protein turnover, chaperones, translation, ribosomal structure, and biogenesis (Fig. 6B). They were also associated with amino acid transport and metabolism and carbohydrate transport and metabolism (Fig. 6B). The DEGs were found to be enriched in three Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and 73 gene ontology (GO) terms by bioinformatics analyses (Majorbio Co., Ltd., China) (Fig. 6C and D and Tables S4 to S7). Notably, the downregulated genes during KEGG pathway analysis included those involved in the PTS system, starch and sucrose metabolism, and fructose and mannose metabolism (Fig. 6C; $P < 0.05$). Similarly, they primarily belonged to various transport systems (sorting $P$ values corrected from low to high), including kinase activity and the phosphoenolpyruvate-dependent sugar phosphotransferase system (Fig. 6D). They were also associated with carbohydrate transport, transferase activity, and organic substance transport (Fig. 6D and Table S6; $P < 0.05$). These findings revealed that the deletion of *rcrR* downregulated the sugar transportation-related genes, leading to a decrease in

**FIG 4** Biofilm formation of *S. mutans* is susceptible to the deletion of *rcrR*. Double labeling of *S. mutans* biofilms formed in BHIS (A) and TVS (D) media. Green denotes bacteria (SYTO 9), and red denotes exopolysaccharide (EPS) (Alexa Fluor 647). Confocal laser scanning microscopy (CLSM)-captured images ($\times 60$ magnification). Three-dimensional reconstruction of the biofilms was performed with Imaris 9.0.1. Representative images were displayed from at least three randomly selected positions in each sample. Quantification of bacteria and EPS biomass (B and E) was conducted, and the ratio of bacteria to EPS (C and F) was calculated using Comstat 2.1. Results are the average of three randomly selected positions in each sample. Data are presented as mean ± SD (*, $P < 0.05$).
transportation of multiple sugars, further influencing the bacterial growth and biofilm formation of S. mutans.

**RcrR directly binds to its own operon promoter.** The 551-bp spacer region between SMU_919 and SMU_921 (rcrR) was divided into three fragments of equal lengths (P1 and P2 = 184 bp each; P3 = 183 bp) to study the DNA binding activity and specificity of RcrR. The three DNA fragments incubated with purified recombinant His-RcrR protein were then subjected to an electrophoretic mobility shift assay (EMSA). There were mobility shifts when RcrR was incubated with the P3 fragment. However, there were no shifts when it was incubated with the P1 and P2 fragments (Fig. 7A). When the P3 fragment-RcrR protein complex was exposed to DNase I, three regions, namely, 5'-ATTAAATTAGTTCTCA-3' (oligonucleotide 1), 5'-ATTATAATAGTTTTCA-3' (oligonucleotide 21), and 5'-TGAGAACTATTATAAT-3' (oligonucleotide 22), were protected by RcrR (Fig. 7B). Interestingly, oligonucleotide 21 plus oligonucleotide 22 was a palindromic sequence with high AT content. These results indicated that RcrR is specifically bound to its promoter region.

**Positive regulation of two PTS systems by RcrR.** There were only 5 downregulated genes based on transcriptome sequencing (RNA-seq) results (Fig. 5), which were classified by two sugar transporter operons, the mannitol-specific PTS system (SMU_1183, SMU_1184c, and SMU_1185) and the trehalose-specific PTS system (SMU_2037 and SMU_2038) (Fig. 8A). The promoter regions (PSMU_1185 and PSMU_2038) were used to search for a generally conserved motif for RcrR binding using MEME Suite 5.1.1 (Fig. 8B), generating a 14-nucleotide (nt) sequence logo (Fig. 8C; P = 0.05). The deletion of rcrR induced the downregulation of two PTS operon genes, suggesting that RcrR can function as an activator to regulate the expression of downstream genes. An rcrR-overexpressing UA159 strain, containing plasmid pDL278-rcrR, was constructed and was used to verify whether RcrR could upregulate two PTS operons. The expression levels of SMU_1183, SMU_1184c, SMU_1185, SMU_2037, and SMU_2038 in the rcrR-overexpressing strain were found to be upregulated 2- to 7-fold compared with the wild-type strain by quantitative reverse transcription-PCR (qRT-PCR) (Fig. 9B). The predicted binding sequences were then subjected to EMSA with RcrR to further investigate this phenomenon in vitro. The mobility shift result generated many DNA-protein complex strips and showed that RcrR could more specifically bind to PSMU_1185 and PSMU_2038 compared with the corresponding controls of P1 and P2 (Fig. 9A).
To further verify whether RcrR binds with PSMU_1185 and PSMU_2038 to activate the expression of the two PTS operons responsible for transporting multiple sugars in vivo, we constructed two mutants (Δ1185p and Δ2038p) in which the RcrR binding sites were deleted (Fig. 10A). The growth curves of UA159, Δ1185p, and Δ2038p were then measured in the presence of different sugars (Fig. 10B to H). We found that the growth rates and final yield of Δ1185p and Δ2038p decreased compared to the those of the wild-type strain when cultured in TV plus glucose/galactose/mannitol medium (Fig. 10B, C, and G). Simultaneously, the growth rate and final yield of Δ2038p significantly decreased compared to those of the wild-type and Δ1185p strains when cultured in TV plus trehalose medium (Fig. 10H). However, the growth rates and final yield of the three strains showed unstable levels when cultured in TV plus lactose/fructose/maltose medium (Fig. 10D to F). These results demonstrated that the deletion of RcrR binding sites decreased the capacity for multiple sugar transportation and the bacterial growth of S. mutans, but not completely. In brief, RcrR binding with the predicted sites was likely to function as a transcription activator to positively regulate the expression of the two PTS operons in S. mutans.

FIG 6 Transcriptomic analysis of the S. mutans ΔrcrR mutant. (A) A volcano plot revealing the differences in gene expression between the ΔrcrR strain and the wild type. (B) Classification and percentage of the differently expressed genes (DEGs) were performed based on the COG type. (C) Functional categories and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis of DEGs using the Majorbio cloud platform (P < 0.01). (D) Functional categories and gene Ontology (GO) enrichment analysis of DEGs using the Majorbio cloud platform (P < 0.05). Red and green denote the upregulated and downregulated genes, respectively.
DISCUSSION

Biofilm formation of *S. mutans* plays an extremely important role in the development of dental caries. Sugar transport and metabolism participate in the regulation of EPS synthesis and biofilm formation, via the PTS systems and ABC transporters (6, 11). The PTS system can also be regulated by various regulatory factors (11, 12). Here, we found that the expression of the two PTS systems was susceptible to *rcrR* deletion. This phenomenon resulted in decreased growth, biofilm formation, and multiple sugar transportation. This study provides further insights into the mechanisms of RcrR as a
transcription activator that manipulates two PTS systems by directly binding to their promoter regions.

Previous studies have found that the rcrRPQ operon plays a critical role in physiological homeostasis and stress tolerance by linking acid and oxidative stress tolerance, (p)ppGpp metabolism, and genetic competence in S. mutans (16). It is well established in the literature that different mutant constructs of the rcrR gene (including \(D_{rcrR}-P\) for a polar mutation and \(D_{rcrR}-NP\) for a nonpolar mutation) can cause different gene expression changes (16, 19). Our method for constructing an \(D_{rcrR}\) mutant is based on the markerless mutagenesis protocol using the IFDC2 cassette that absolutely deletes the open reading frame (ORF) (from ATG to TAA) (20). Interestingly, there are extreme differences in gene expression profiling between the RNA-seq results of Kaspar et al. for \(D_{rcrR}-P\) and \(D_{rcrR}-NP\) strains (19) and our RNA-seq results for the \(D_{rcrR}\) strain. Only the expression levels of a few genes were consistent. For example, the gene expression levels of the rcrRPQ operon and SMU_1185 were upregulated and downregulated in both strains (\(D_{rcrR}-NP\) and our \(D_{rcrR}\) mutants). We further verified the expression levels of rcrRPQ operon in the wild-type, mutant, complement, and rcrR-overexpressing strains by qRT-PCR (see Fig. S1 in the supplemental material), consistent with our RNA-seq results. The above-mentioned results suggested that the phenotype of our mutant was partially, but not absolutely, similar to that of the \(D_{rcrR}-NP\) mutant. It is possible that S. mutans has developed various strategies (including changing the expression levels of different genes) in response to different rcrR mutations caused by complex environmental stresses in the oral cavity.

One more difference between this RNA-seq study of an rcrR mutant and the Kaspar et al. RNA-seq study is that our study was completed in an anaerobic environment and that of Kaspar et al. was completed in 5% CO\(_2\) atmosphere (oxygen) (19). A previous study demonstrated that the biochemical and phenotypic properties of S. mutans were changed during growth with aeration compared with growth under an anaerobic condition (21, 22). In particular, S. mutans bacteria that initially colonize an oral surface are exposed to higher levels of oxygen than those in mature biofilms, where diffusion of oxygen appears to be limited (22). Thus, S. mutans may regulate the expression level of the corresponding genes in response to exposure to oxygen. Seaton et al. showed that \(\Delta_{crp-NP}\) and \(\Delta_{crq-NP}\) mutants were sensitive to oxidative stressors (16), suggesting that the rcrRPQ operon may play important roles in response to

![Predicting the potential binding sites of RcrR. (A) The operon promoters of mannitol-specific (PSMU_1185) and trehalose-specific (PSMU_2038) phosphotransferase (PTS) systems were used to search for a generally conserved motif (B) for RcrR binding using MEME Suite 5.1.1. (C) A predicted sequence logo for the RcrR binding motif was generated using MEME Suite 5.1.1 (\(P < 0.05\)).](msystems.asm.org/FIG_8.png)
environmental changes. In our present study, S. mutans strains were cultured in an anaerobic environment, eliminating the interference of oxygen. When exposed to oxygen, the expression level of the \textit{rcrRPQ} operon and the observed phenotype(s) of our strains, including biofilm formation and sugar utilization, might be different from those under anaerobic conditions. These speculations remain to be further verified in future studies.

The MarR family transcriptional regulators are widely distributed in bacteria. They regulate various functions, such as resistance to antibiotics and oxidative stressors, degradation of harmful chemicals, and the expression of virulence genes (23). These proteins mainly bind to recognizable palindromic sequences within the intergenic region of divergent genes. Binding leads to reduced gene expression, as the RNA polymerase is hindered from binding to the promoter (23). Previous studies have shown that RcrR, as one of the MarR family members, functions as a repressor to regulate \textit{rcrP} and \textit{rcrQ} expression in the \textit{rcrRPQ} operon by binding to its promoter region in \textit{S. mutans} (16, 24). In this study, the expression levels of the two PTS operons were downregulated when the \textit{rcrR} gene was knocked out, suggesting that RcrR directly functioned as an activator to regulate the gene expression of the two PTS operons. The results of the EMSA (Fig. 9A) showed that RcrR was able to bind to promoter regions of the two PTS operons. Moreover, keeping in mind the results of qRT-PCR (Fig. 9B) \textit{in vivo} (Fig. 9D), it is probable that RcrR functions as an activator binding to the promoter regions to recruit or stabilize RNA polymerase, or alter promoter conformation for activating transcription, via a mechanism similar to that of the CueR regulator (25).
Nonetheless, the actual mechanism underlying activation after RcrR binding with the two promoter regions should be further explored.

Our DNase I footprinting assays results showed that there were three fragments (oligonucleotides 1, 21, and 22; 16 nt) protected by RcrR and that their sequences were highly identical, which further verified and supplemented the conserved DNA sequence (24). Meanwhile, oligonucleotide 21 plus oligonucleotide 22 was a palindromic sequence with high AT content, suggesting that RcrR prefers binding to DNA regions with high AT content. This preference was also similar to those of the MarR family regulators CouR (26) and FabT (27).

*S. mutans* appears to have developed diverse strategies for sugar utilization in response to complex environmental changes in the human oral cavity. A series of carbohydrate transporter systems have been found in *S. mutans* UA159, including 14 PTS systems and 4 sugar ABC transport systems (11, 28). The PTS system consists of three components, namely enzyme I (EI), EII (EIIA-D), and a histidine-containing phospho-carrier protein (HPr). Glucose binds to EI and is transported into the cell membrane, where HPr phosphorylates it to form glucose 6-phosphate (Glc-6P), which then immediately enters the glycolysis pathway (6, 29). The well-characterized mannitol operon contains *SMU_1185* (encoding a mannitol-specific EIIBC PTS component), *SMU_1184c* (a transcriptional regulator, antterminator), *SMU_1183* (encoding a mannitol-specific EIIA PTS component), and *SMU_1182* (encoding a mannitol-1-phosphate dehydrogenase) (28). The trehalose-specific PTS system contains *SMU_2037* (encoding a trehalose-6-phosphate hydrolase), *SMU_2038* (encoding a trehalose-specific EIIBC PTS component), and *SMU_2040* (encoding a GntR family regulator, TreR) (13, 28). These components of the two operons play a critical role in sugar transportation. In this study, deletion of *rcrR* resulted in decreased expression of these components. Our data also showed that the deletion of binding sites could result in a low growth rate and low final yield of *S. mutans* when cultured in TV plus sugar medium (Fig. 10B, C, G, and H). Given that RcrR binds to the promoter regions of the two operons, it is easy to think that the deletion of binding sites abolished the activation ability of RcrR upon binding with the promoter regions, consequently leading to decreased expression of the two operons and a decreased ability to transport multiple sugars. In addition, we also speculate that the deletion of *rcrR* mainly influenced sugar uptake, resulting in a decrease in...
substance sources, energy for bacterial growth, and sugar metabolism in S. mutans, as genes related to sugar metabolism showed no significant changes based on RNA-seq results. These results possibly explain the reason for the delay in bacterial growth and decreased biofilm formation ability. However, this hypothesis should be further tested.

Sugar transport and metabolism are regulated by different mechanisms (15). Organisms can express and coordinate the regulation of multiple sugar uptake systems in response to complex environments. Previous studies have postulated that PTS systems are widely regulated by various factors, such as FruR (9), StsR (12), TreR (13), CelR (14), and NigR (15). For example, deletion of stS changes the expression levels of PTS and ABC transporter genes, including mannitol- and trehalose-specific PTS systems (12). Other studies have also reported that the mannitol- and trehalose-specific PTS systems are regulated by MtrR and TreR, respectively (13, 30). Here, RcrR, as an activator, also regulated these two operons. As such, these results suggest that RcrR, probably interacting with proteins (StsR or TreR), can coregulate the transcription of SMU_2037 and SMU_2038.

In summary, S. mutans has multiple pathways to coordinate sugar transport and metabolism in response to complex environmental conditions. The findings demonstrate that RcrR is a transcription activator with pleiotropic effect on bacterial growth, biofilm formation, and multiple sugar transportation, which are all important virulence traits of S. mutans. Based on this, it is of paramount interest to determine the specific signals that can induce the RcrR-related regulatory pathway. Notably, this study also decrypts the physiological functions of RcrR as a potential target for controlling biofilm formation and multiple sugar uptake for the better prevention of dental caries.

MATERIALS AND METHODS

Bacterial strains and their growth conditions. All bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. S. mutans UA159 and its derivatives were routinely grown in BHI broth (BD, USA) or BHI agar at 37°C in an anaerobic incubator (80% N2, 10% H2, and 10% CO2). Escherichia coli strains were grown in Luria-Bertani medium at 37°C under aerobic conditions.

Construction of in-frame mutants. An in-frame rcr deletion (ΔrcrR) mutant strain was constructed using S. mutans UA159. Construction was performed via a two-step transformation procedure (5, 20). PCR and DNA sequencing methods were used to confirm whether a correct rcr deletion mutant strain was obtained. Mutants of Δ1185p and Δ2038p were obtained following the same method. All primers used here are listed in Table S1.

Construction of rcrR complement and overexpression strains. The rcrR coding sequence, including its promoter region (upstream 490 bp) was cloned into a SacI/Sall-digested E. coli-Streptococcus shuttle vector, pDL278 (31). The recombinant plasmid pDL278-rcrR was then transformed into an rcrR mutant strain to generate the complement strain ΔrcrR/pDL278-rcrR. The transformation efficiency (or genetic competence) is reduced because of the deletion of rcrR (16); therefore, we constructed the complement strain (ΔrcrR/pDL278-rcrR) using electroporation. pDL278-rcrR was also transformed into UA159 to generate the overexpression strain UA159/pDL278-rcrR, as previously described (5). PCR and DNA sequencing were performed to confirm whether the correct strains were obtained. The primers used in this study are listed in Table S1.

Growth curves. Cultures grown overnight in BHI medium were diluted to 1:10 in fresh BHI, 1/2 BHI, and 1/4 BHI media. Cell growth was monitored using Multiskan GO (Thermo Fisher Scientific, USA) for 12 h under anaerobic conditions (80% N2, 10% H2, and 10% CO2). OD600 was measured at 1-h intervals. The growth of the strains was further assessed in the presence of different carbohydrates. Cultures grown overnight in BHI medium were diluted to 1:10 in fresh BHI and grown to the mid-exponential phase (OD600 ≤ 0.5). They were then diluted to 1:100 into fresh BHI, 1/2 BHI, and 1/4 BHI media. Cell growth was monitored using Multiskan GO (Thermo Fisher Scientific, USA) for 12 h under anaerobic conditions (80% N2, 10% H2, and 10% CO2). OD600 was measured at 1-h intervals. The growth of the strains was further assessed in the presence of different carbohydrates. Cultures grown overnight in BHI medium were diluted to 1:10 in fresh BHI and grown to the mid-exponential phase (OD600 ≤ 0.5). They were then diluted to 1:100 in a tryptone-vitamin (TV) base medium supplemented with 1% (wt/vol) glucose, galactose, lactose, fructose, maltose, or trehalose (17). The base TV medium contained 3.5 g of tryptone/100 ml, 0.04 μg of p-aminobenzoic acid/ml, 0.2 μg of thiamine-HCl/ml, 1 μg of nicotinamide/ml, and 0.2 μg of riboflavin/ml. Cell growth was monitored using Multiskan GO for 24 h under anaerobic conditions. OD600 was then measured at 2-h intervals. Overnight cultures of strains were diluted to 1:10 in fresh BHI and grown to the mid-exponential phase (OD600 ≤ 0.5). Subsequently, the cultures were diluted to 1:10 and transferred to fresh TV medium supplemented with glucose, galactose, lactose, fructose, maltose, or trehalose (1%, wt/vol) (Fig. 10). Cell growth was monitored using Multiskan GO for 12 h under anaerobic conditions. OD600 was measured at 2-h intervals. Each analysis was performed in triplicate, and the representative growth curves were plotted.

Biofilm detection by crystal violet staining assay. As described previously (32), cultures grown overnight were diluted to 1:10 in fresh BHI and grown to an OD600 of approximately 0.5 to 0.6. The log-phase cells were then diluted to 1:100 in fresh BHI plus 1% sucrose (BHI5; wt/vol) or TV plus 1% sucrose (BHI5; wt/vol) for 24 h.

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(TVS) and cultured in sterile 48-well microtiter plates for 24 h. The cells were then dumped out after incubation by turning the plate over, followed by gently submerging the plate in a small tub of water. The water was then drained out, and 0.01% crystal violet solution was added to stain the biofilm for 15 min. The plate was then rinsed 3 or 4 times by submerging in water. Ethanolic acid (30%) was then added to solubilize the dye for 15 min. Finally, 100 µl of ethanolic acid solution was transferred to a new plate, and its absorbance at 570 nm was measured.

Confocal laser scanning microscopy analysis of biofilm. Biofilms were grown on a glass-bottomed petri dish (NEST, China) in BHI or TVS medium for 24 h. Bacterial cells and EPS from biofilms were then labeled with 5 µM SYTO 9 and 1 µM Alexa Fluor 647 (Invitrogen, USA), respectively (33). Alexa Fluor 647-labeled dextran conjugate was then added to the culture medium before inoculation. The medium was subsequently decanted after incubation, and the petri dishes were washed twice with 0.85% NaCl. The bacteria were then stained with SYTO 9 for 15 min, and then the residual dye was washed twice with 0.85% NaCl. Confocal imaging was captured using an FV3000 instrument (Olympus, Japan) with a 60× oil immersion objective. The image collection gates were set at 650 to 750 nm for Alexa Fluor 647 and 500 to 540 nm for SYTO 9. During imaging, Gain (1.0×), HV (365 V for SYTO 9; 410 V for Alexa Fluor 647), and offset (4%) were kept constant. Z sections were used to determine the biofilm thickness. Three-dimensional reconstruction of biofilms and calculation of bacteria and EPS biomass were performed using Imaris 9.0.1 and ImageJ contained in Comstat 2.1, respectively (34).

Transcriptome sequencing and data analysis. For transcriptome analysis, the S. mutans UA159 and S. mutans ΔrcrR strains were routinely grown at 37°C under anaerobic conditions in BHI medium until an OD₆₀₀ of 0.5 was reached. They were then centrifuged (4,000 rpm for 10 min) and resuspended in liquid nitrogen for 15 min. They were subsequently submitted to Majorbio Co., Ltd. (Majorbio, China) for genome-wide RNA sequencing. Three independent total RNA extracts of each sample were extracted using TRIzol reagent according to manufacturer’s instructions (Invitrogen, USA), and genomic DNA was removed using DNase I (TaKaRa, Japan). Then, RNA quality was determined using a 2100 Bioanalyzer (Agilent, USA) and quantified using an ND-2000 instrument (NanoDrop Technologies, USA). cDNA libraries were constructed from enriched mRNA samples using the TruSeq RNA sample prep kit (Illumina, USA). RNA depletion from total RNA was performed using the Ribo-Zero magnetic kit (Epicentre, USA), and the mRNA was chemically fragmented to short pieces (200 nt) using a 1× fragmentation solution (Ambion, USA) for 2.5 min at 94°C. Double-stranded cDNA was produced using the SuperScript double-stranded cDNA synthesis kit (Invitrogen, USA) with random hexamer primers (Illumina, USA). Then, the synthesized cDNA was subjected to end repair, phosphorylation, and “A” base addition according to Illumina’s library construction protocol. Libraries were selected for cDNA target fragments of 200 bp on 2% low-range agarose gels. PCR amplification was performed using Phusion DNA polymerase (NEB, USA) for 15 PCR cycles. After quantification by TBS380, the paired-end RNA-seq sequencing library was sequenced with the Illumina HiSeq X Ten instrument (2 × 150-bp read length). Genes with a fold change of >2.0 and a P value of <0.05 were selected for further gene expression pattern discovery. All analyses were performed using the free online Majorbio cloud platform (www.majorbio.com).

Expression and purification of the recombinant RcrR protein. The rcrR gene was cloned into pET28a to produce recombinant vectors (pET28a-rcrR). The EcoRI and XbaI restriction sites were used. The RcrR protein was then purified using the His tag protein purification kit (Beyotime, China), following the manufacturer’s protocol. The concentration of purified His-RcrR protein was determined by measuring its spectrophotometric absorbance at 280 nm. The primers used are listed in Table S1.

DNA substrate preparation and EMSA. The rcrR promoter (551 bp) was divided into three DNA fragments, P1 (184 bp), P2 (184 bp), and P3 (183 bp). P1, P2, and P3, the mannotol-specific promoter (184 bp), and the trehalose-specific promoter (173 bp) were acquired by PCR. The DNA fragments (20 pmol) were then incubated with various amounts of RcrR proteins in a total volume of 20 µl EMSA buffer for 30 min on ice. The buffer contained 20 mM Tris-HCl (pH 7.0), 100 mM NaCl, and 5% (vol/vol) glycerol. The DNA-protein complexes were then resolved by electrophoresis on a 5% (wt/vol) nondenaturing polyacrylamide gel in 0.5× Tris-borate-EDTA (TBE) buffer at 110 V for 60 min. The gels were dyed with ethidium bromide for 20 min, and images were captured using a ChemiDoc touch imaging system (Bio-Rad, USA). The primers used in the study are listed in Table S1.

DNase I footprinting assays. The P3 fragment was amplified using 6-carboxyfluorescein (FAM)-labeled primers. Amplified fragments were then subjected to a binding reaction similar to that in EMSA. DNase I footprinting was then performed (35). The reaction mixtures, composed of 20 pmol of gene products and either 0, 80, or 160 pmol of RcrR, were treated with DNase I (0.015 U) at 37°C for 5 min. The samples were then extracted with phenol, precipitated with ethanol, and eluted in 15 µl of distilled water. The fragments were analyzed using a 3730XL DNA analyzer (Applied Biosystems, Tsingeke, China). The two operon promoters were used to search for a generally conserved motif for RcrR binding using MEME Suite 5.1.1.

RNA extraction and quantitative reverse transcription-PCR. Bacterial strains were grown overnight in BHI broth and diluted to 1:10 in fresh BHI. They were allowed to grow to an OD₆₀₀ of 0.5 and then harvested for total RNA isolation. Extraction was performed using the MasterPure complete DNA and RNA purification kit (Lucigen, USA) following the manufacturer’s protocol. cDNA was synthesized using the PrimeScript RT reagent kit with the gDNA Eraser kit (TaKaRa, Japan) according to the manufacturer’s instructions. qRT-PCR was performed on a QuantStudio 6 Flex real-time PCR system (ABI, USA) using TB Green Premix Ex Taq II (TaKaRa, Japan). The QuantStudio 6 Flex software determined threshold cycles and dissociation curves. The determination was conducted to ensure that only one PCR product was amplified and detected. The differential expression fold was quantified following the comparative
threshold cycle (2–ΔΔCt) method with 16S rRNA as an internal standard. The primers used in this study are listed in Table S1.

Statistical analysis. All experiments were performed in triplicate and separately reproduced three times. Data were analyzed using SPSS statistical software (version 20 for Windows; SPSS, Inc., Chicago, IL). Student’s t test was used to compare data between two groups. Analysis of variance (ANOVA) and Tukey’s test were performed to compare data between multiple groups. A P value of <0.05 indicated that there were significant differences between groups.

Data availability. The raw data from the transcriptomic sequencing analysis of S. mutans UA159 and the S. mutans ΔcrR mutant have been deposited in the NCBI Sequence Read Archive (SRA) database under BioProject accession number PRJNA749547.

Supplemental material

Supplemental material is available only.

FIG S1, TIF file. 1 MB.

TABLE S1, XLSX file. 0.01 MB.

TABLE S2, XLSX file. 0.01 MB.

TABLE S3, XLSX file. 0.01 MB.

TABLE S4, XLSX file. 0.01 MB.

TABLE S5, XLSX file. 0.01 MB.

TABLE S6, XLSX file. 0.01 MB.

TABLE S7, XLSX file. 0.01 MB.

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We declare no conflicts of interest.

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