Acid-resistant genes of oral plaque microbiome from the functional metagenomics

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ABSTRACT
Acid resistance is one of key properties assisting the survival of cariogenic bacteria in a dental caries environment, but only a few genes conferring acid resistance have been identified to date. Functional metagenomics provides a systematic method for investigating commensal DNA to identify genes that encode target functions. Here, the host strain Escherichia coli DH10B and a constructed bidirectional transcription vector pSKII-lacZ contributed to the construction of a metagenomic library, and 46.6 Mb of metagenomic DNA was cloned from carious supragingival plaque of 8 children along with screening for lethal functionality. The screen identified 2 positive clones that exhibited a similar acidic phenotype to that of the positive controls. Bioinformatic analysis revealed that these two genes encoded an ATP/GTP-binding protein and a malate dehydrogenase. Moreover, we also performed functional screening of Streptococcus mutans, since it is one of the predominant cariogenic strains but was not identified in our initial screening. Five positive clones were retrieved. In conclusion, our improved functional metagenomics screening method helped in the identification of important acid resistance genes, thereby providing new insights into the mechanism underlying caries formation as well as in the prevention and treatment of early childhood caries (ECC).

Introduction
Early childhood caries (ECC) is one of the most common infectious diseases in children and affects up to 60–90% of children worldwide [1]. ECC can not only cause tremendous pain [2] but can also result in an enormous social cost [3]. Consequently, the prevention and treatment of ECC are extremely important and mandate intensive study [4]. It is well known that microorganisms living in the dental plaque biofilm play a vital and direct role in the occurrence of tooth decay [2]. Survival and propagation of microorganisms in the caries microenvironment inducing tooth decay require that they exhibit certain properties, such as adhesion, acidogenicity and acid resistance. Among them, acid resistance is a key factor for the survival of cariogenic bacteria [5] in the typical carious microenvironment, in which the pH values fluctuate in the range from pH 7 to 3, primarily due to diet [6]. Hence, the acid resistance of oral microbes is taken into account in the study of ECC prevention and treatment, which offers potential new insights into clinical applications and personalized medicine [7].

Currently, only a minor fraction of acid-resistance genes in oral microorganisms have been discovered and investigated given the limitations of genetic and molecular biology techniques primarily based on culture-dependent methods [8]. In the mid-1980s, Loesche et al. [5] reported that Streptococcus mutans is a principal causative agent of dental caries, and its prevalence was 70–100% of individual cases of caries. Subsequently, a large number of studies on the acid production and acid-resistant properties of Streptococcus were performed [9,10], and some relevant acid-resistant genes, such as fth, uvrA, and dnaK, were discovered and studied [11–13]. These achievements can be partially attributed to the ease in propagating this strain in the laboratory. However, methods in propagating generally difficult-to-grow microbial strains have scarcely been established to date. Recent high-throughput sequencing studies based on culture-independent analysis of childhood caries revealed that S. mutans is not the only dominant pathogenic bacterium in caries. Lactobacillus, Selenomonas and Neisseria can also be detected at high levels in severe carious plaques [14,15], and some of these bacteria cannot be cultured (Human Oral Microbiome Database (HOMD), http://www.homd.org/). In addition to Streptococcus, our knowledge about the acid resistance and related regulatory genes of cariogenic microorganisms, especially those that are not cultivable, is limited; thus, it offers a fertile field for the current study.

To explore this field, we utilized function-based metagenomics [16], which can identify both novel and previously known genes involved in target functions,
even with unknown genetic information. Currently, function-based screening of metagenomic libraries has been widely deployed in different fields, including the screening of environmental libraries [17], the investigation of mouse gut colonization [18], of human faecal samples [19] and others. However, acid-resistance screening studies of oral plaque samples have not yet been reported.

Therefore, this study attempted to identify genes conferring acid resistance from the supragingival plaque of 4–6-year-old children using improved function-based metagenomics to provide a new entry point for the study of the caries mechanism and ECC prevention. A high-copy vector allowing bidirectional transcription was constructed to enhance library expression efficiency. Total microbial DNA of supragingival plaque samples from carious teeth of children was extracted. The DNA was used to construct a metagenomic library, and quality assessment was done before lethal screening was conducted. Bioinformatics analysis of the resulting positive clones identified two genes that were predicted to, respectively, encode an ATP/GTP-binding protein and malate dehydrogenase.

Materials and methods

Ethics statement

Informed consent forms were signed by the parents of all subjects prior to study enrolment. The entire study design, protocol and informed consents were approved by the Medical Ethics Committee at the School & Hospital of Stomatology, Wuhan University, Wuhan, China. Additionally, we confirmed that all experiments were performed in accordance with relevant guidelines and regulations.

Dental plaque samples

Supragingival dental plaques, derived from carious teeth of 4–6 years-old children, were sampled to construct the metagenomic library. The inclusion criteria were as follows: (1) Healthy medical history (2). No current bacterial/viral infections and no antibiotics within the past 3 months (3). At least three cavitated caries teeth among more than 18 present teeth (code >4, according to the criteria of the International Caries Detection and Assessment System (4). No detectable enamel and dentin hypoplasia and (5). No prior treatment (including fluoride) for dental decay.

Only supragingival plaque samples around the cavity were collected using a sterile curette that was immersed into a 1.5 ml Eppendorf tube containing 1 ml of autoclaved saline solution. The kids were asked not to eat, drink and brush within 2 h before sampling. Every individual was repeatedly sampled and all samples were stored at −80°C until use.

Bacterial strains, plasmids, media, and culture conditions

Escherichia coli DH10B, DH5α, and JM110 (TaKaRa Bio Inc. Japan) strains were routinely cultured in Luria–Bertani (LB) medium at 37°C. The shaking speed was set at 200 rpm for liquid cultures. If needed, the growth medium was supplemented with 100 mg/ml ampicillin (Amp) to maintain pUC19, pBluescript SKII(+) (pSKII+) and pSKII−lacZ (pSL, a dual-direction expression vector constructed in this study) vectors. E. coli JM110 was mainly used for demethylation.

The standard S. mutans UA159 strain (ATCC 700,610) was kindly provided by MOST KLOS & KLOBM, School & Hospital of Stomatology, Wuhan University. The strain was amplified in Brain Heart Infusion (BHI) broth in an anaerobic jar at 37°C.

The construction and verification of the vector pSL allowing bidirectional transcription

To enhance the efficiency of library screening, a bidirectional transcription vector pSKII−lacZ was constructed. To verify whether both promoters were functional, a reporter gene encoding green fluorescent protein (GFP) was cloned and ligated into the pSL (for a detailed protocol see Supplementary File). Finally, the strains E. coli DH10B/pSL-GFP (OD = 1.2–1.5) were observed under excitation fluorescence microscopy to detect the fluorescence intensity.

DNA extraction

The plaque samples were mixed in one sterile tube to obtain total DNA. Two independent aliquots of mixed samples were prepared separately for screening and cross validation. The protocol of total DNA extraction was taken from a previous study [20]. The purity and concentration of DNA were measured using agarose gel electrophoresis and NanoDrop 2000 (Thermo Scientific, Wilmington, DE). All the DNA samples were stored at −80°C before further analysis.

Construction of a metagenomic library

For the construction of the supragingival plaque metagenomic library, total DNA was partially digested with Sau3A I (Thermo Scientific) and three DNA fragments of 1–4 kb, 4–7 kb, and 7–10 kb were separately excised from the agarose gel and purified using the OMEGA Gel Extraction Kit (Omega Bio-Tek, Norcross, GA). The DNA
fragments were ligated into a dephosphorylated and BamHI-digested pSL vector at molar ratios of 3:1 to 7:1 (insert/vector) using T4 DNA ligase. Then, 10 μL of ligation mixture was transformed in competent E. coli DH10B, which was plated on LB_Amp agar and incubated for 14 h at 37°C to amplify the libraries. The detection of metagenomics library quality was completed prior to screening. Per plate 1–2% of the colonies were randomly selected to test the efficiency of self-ligation by extracting the plasmids, digesting with EcoRI and XbaI and performing agarose gel electrophoresis. All the ligation and transformation reactions were performed sequentially, and the plates were stored at 4°C for less than 5 days to ensure sufficient time to implement the next step. All primers were designed using Primer Premier 5 (PREMIRE Biosoft, Palo Alto, CA) and are listed in Supplementary Table S1. The protocol for library construction is illustrated in Figure 1(a).

In addition, we used two known acid-resistant genes, orf1 and ffh, to construct two positive controls, E. coli DH10B/pSL-orf1 and pSL-ffh, the former from the environmental microbes [17], the latter from S. mutans [11]. The detailed protocol is described in the Supplementary File.

**Screening of the metagenomic library**

Colonies of the library were directly incubated in 96-well plates containing 0.7 ml of pH 1.9 LB broth supplemented with 100 μg ml⁻¹ Amp for 3 h at 37°C, which is a lethal condition for strain E. coli DH10B/pSL. The LB broth was acidified to pH 1.9 using an aqueous solution of 37% (w/v) HCl and sterilized in an autoclave for 30 min at 115°C. After the acid shock, 3 μL of cells were dropped on LB_Amp agar and incubated at 37°C for 14 h (Figure 1(b)). Meanwhile, negative (E. coli DH10B/pSL) and positive controls were cultured and subjected to acid shock. Colonies that survived like the positive controls were regarded as potential acid resistant clones. Then, the plasmids of these potential acid resistant clones were isolated and transformed into new competent E. coli DH10B,

![Figure 1. Schematic diagram of (a) library construction and (b) functional screening.](image-url)
after which the acid shock was repeated. Those clones that continuously exhibited an acid-resistant phenotype were considered to be positive clones. These plasmids were subsequently sequenced using primer P2 (Sangon Biotech Co.) and stored at −80°C until used.

**In silico analysis**

Based on the sequencing data of the positive clones, potential operon reading frames (ORFs) with start (ATG) and stop (TAA, TAG, TGA) codons were predicted and analysed using Vector NTI Advance11.5.1 and DNAMan v6.0. In addition, alignments of nucleotide sequences and transcribed amino acid sequences were performed against public databases using BlastN and BlastX (basic local alignment search tool) at the NCBI on-line server ([https://blast.ncbi.nlm.nih.gov/Blast.cgi](https://blast.ncbi.nlm.nih.gov/Blast.cgi)). The predicted ORFs similar to known or unknown proteins and longer than 100 bp were obtained and further analysed. The conserved domains of inserts were analysed using the CD search module in NCBI ([https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi](https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi)). Their structure and putative function were annotated based on similarities to the sequences in the Clusters of Orthologous Groups (COG) [21], Protein Families (Pfam) [22] ([http://xfam.org/](http://xfam.org/)), and Blocks [23] ([InterPro http://www.ebi.ac.uk/interpro/](http://www.ebi.ac.uk/interpro/)) databases and based on the results of BLAST searches in UniProt ([http://www.uniprot.org/](http://www.uniprot.org/)). If the similarity of the protein sequence alignment was less than 30%, it was considered to be an unknown sequence. Trans-membrane helices were forecasted with the TMpred programme ([http://www.ch.embnet.org/software/TMPRED_form.html](http://www.ch.embnet.org/software/TMPRED_form.html)) and UniProt.

**Acid-resistance assay**

The acid-resistance assay was based on the library functional screening protocol used in this study. The positive clones and negative/positive controls were activated and inoculated overnight (12 h) in 5 ml LB_Amp broth at 37°C, which contained 2–3 × 10^8 cfu (colony-forming units) per millilitre of culture (OD 1.45–1.55). Then, 10 μL of culture was separately transferred into 1 ml of PBS (phosphate-buffered saline, pH 7.2) and 1 ml of LB broth (pH 1.9). The initial cell population was quantified by plating serial dilutions of PBS buffer on the LB agar and incubating for 14 h at 37°C before counting the cfu. The final cell population was determined as described above after the strains were subjected to acid shock (pH 1.9). Subsequently, the percentage survival value was calculated by dividing the final number of cfu ml⁻¹ after the acid shock by the initial number of cfu ml⁻¹ and the average values of three independent experiments were reported.

**Screening and analysis of S. mutans UA159**

The library construction, functional screening and sequence analysis of S. mutans UA159 were conducted with the above protocol.

**Results**

**Subjects and samples**

Thirty-two caries associated supragingival plaque samples were obtained through four consecutive repeat samplings from eight children aged 4–6 (sample information is shown in Supplementary Table S2). All 32 samples were mixed together and placed into two 1.5 ml Eppendorf tubes. The two resulting samples weighted 28.2 and 29.1 mg, respectively.

**Validation of the dual-direction expression vector pSL**

To improve the efficiency of library screening, a dual-direction expression vector, pSKII−lacZ (pSL), was constructed (the schematic of pSL is shown in Figure 2(a), for a detailed protocol see Supplementary File). We used green fluorescent protein (GFP) to detect whether the two promoters were effective. Strains of E. coli DH10B/pSL-GFP with two different directions of GFP showed green fluorescence (Figure 2(b,c)), indicating that both promoters worked. However, stronger fluorescence was observed with the reverse GFP. Analysis found that this could be attributed to the start codon ATG of the vector-self aiding in the generation of a fusion protein weakening the expression of GFP.

**Preparation of samples and construction of libraries**

The extraction of the total DNA from plaque samples can be found in a previous study [20], named DNA1 and DNA2; the concentrations are 469.8 and 492.9 ng/μl, respectively. The test results of gel electrophoresis and NanoDrop 2000 are shown in Supplementary Figure S1a and Table S3. Approximately 0.08 mg of high-quality DNA per milligram of sample (wet weight) was obtained. Subsequently, we constructed a plaque library using the DNA1 (see Methods). To ensure the library quality, 1–2% colonies per plate were randomly selected to detect the ligation rate of exogenous fragments. We only selected one plate from the same batch of transformation experiments were reported.

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approximately 11,290 colonies were obtained with 46.6 Mb of DNA.

**Acid-resistance screening of the library**

The lethal screening condition was acid shock for 3 h in the medium of pH 1.9, which was confirmed by pre-experimentation to be fatal to the strain *E. coli* DH10B/pSL, but non-toxic to the positive control (E. coli DH10B/pSL-orf1 and pSL-ffh). Both the inserts orf1 and ffh are known aciduric-related genes. The former encodes RNA-binding protein and a separate functional metagenomics study of the environmental sample confirmed its function [17], the latter comes from the *S. mutans* strain [11]. The purpose of construction of the strain *E. coli* DH10B/pSL-ffh and its functional verification was to detect whether the acid-resistant genes from oral bacteria could be heterologously expressed in *E. coli*.

Hence, after the first round of acid shock, seven colonies were identified that could survive in this hostile environment, but only two of the seven clones continuously exhibited acid resistance upon repeated acid shocks (repeated three times). Therefore, these two clones were regarded as positive clones carrying genes conferring acid resistance.

**Molecular analysis of acid-resistant clones**

The sequences of two positive clones are displayed in the Supplementary File. The genetic organization, genetic similarities, potential ORFs, phylogenetic relationship, putative function, and transmembrane domain predictions of the two clones are analysed and summarized in Table 1. As shown in the table, the two acid-resistant clones (pSL-1–31 and 2–41) contained two complete heterologous ORFs. The

| Code | 1–31 | 2–41 |
|------|------|------|
| Seq length (bp) | 836 | 2636 |
| Site* (Nt range)$^b$ | 1–94aa (835–425bp) | 1–643aa (402–2324bp) |
| G + C % | 46.9 | 47.7 |
| Alignment$^c$ | Actinomyces sp. oral taxon 178 str.F0338 | Haemophilus haemolyticus |
| Protein (accession No.)/size (aa)$^c$ | ATP/GTP-binding protein (EFW08722)/137aa | Malate dehydrogenase (WP_005646657.1)/756aa |
| E-value | 2.00E-49 | 0 |
| Identities % | 78/80 (98%) | 372/643 (58%) |
| Accession No. of the similar protein$^d$ | E8JLB4 | H1LMB2 |
| Putative function$^d$ | Play a pivotal role in various distinct biological processes covering protein export, membrane transport, DNA repair etc. | Play a key role in the central oxidative pathway. It assists *E. coli* to survive in the acid environment. |

$^a$Protein range (site) of alignment into the known proteins.

$^b$Nucleotide range (nt range) of the predicted ORF within insert.

$^c$Most similar protein, currently identified with blastp: organism, name, accession no, and protein size.

$^d$Most similar protein, currently identified with UniProt: accession number of the similar protein, conserved domain and putative function.

$^e$Results from the TMpred server.

bp: base pair; aa: amino acids; score: bit score of the alignment using BLAST.
clones exhibited high similarity (98 and 58%) to known functional proteins, indicating that no novel genes were discovered from this plaque library. The G + C contents of the inserts were 46.9 and 47.7%.

The sequences of pSL-1–31 were aligned with sequences from Actinomyces sp. oral taxon 178 str. F0338 (Actinobacteria), which is a gram-positive, anaerobe/facultative anaerobe bacterium that encodes an ATP/GTP-binding protein, belonging to the nucleotide-binding protein family. It is involved in various distinct biological processes. The other positive clone pSL-2–41 was aligned with sequences from Haemophilus haemolyticus (Proteobacteria), which is a gram-negative, facultative anaerobe that encodes malate dehydrogenase, and has a known acid-resistant gene (the details are presented in Table 1).

**Acid-resistance assay**

To quantify the ability to resist acid by the inserts in positive clones, an acid-resistance assay was performed on the two functional clones (pSL-1–31, pSL-2–41) and negative/positive controls (Figure 3). In Figure 3, there were statistically significant differences between all pairs of groups except for group pSL-1–31 and pSL-orf1. The test analysis revealed that the two ORFs in the inserts were responsible for substantial improvement in the acid resistance of host cells but to a different degree. Furthermore, pSL-ffh displayed the highest resistance level \( P < 0.0001 \), which not only further verified the acid-resistance conferred by ffh, but also provided evidence for the heterologous expression of the genes from *S. mutans* in *E. coli*.

**Screening and analysis of S. mutans UA159**

Based on the above results, we found that, in our study, although *S. mutans* was regarded as the primary cariogenic agent, no functional genes were retrieved from the examined strain. This interesting result prompted us to construct and screen another library from *S. mutans* UA159 as outlined above.

Eventually, five positive clones were obtained (the sequences of these clones are presented in the Supplementary File), which encoded molecular chaperone DnaK (U-57), chaperone protein ClpC (U-38), 23S rRNA-methyltransferase (U-41), acyl-phosphatase (U-23), rod-shape-determining protein MreD and glucan-binding protein B (U-8) (Table 2). The DnaK molecular chaperone (U-57) is a ubiquitously distributed class I stress protein, which tolerates the low pH potentially through regulating the biogenesis or stabilization of the F-ATPase compound [13,24]. The functions of Clp protease (U-38) and methyltransferase (U-41) have been corroborated by a previous study [17]. Only SMU_1725 (U-23), mreD and gbpB (U-8) were newly identified functional genes from *S. mutans* UA159. These genes are predominantly involved in catalyzing the hydrolysis of various acyl phosphate carboxyl-phosphate bonds, regulating cell shape and the formation of biofilm on inanimate substrate.

**Discussion**

As one of the most determinant properties of cariogenic bacteria in inducing tooth decay in children, acid resistance deserves extensive research. Currently, various meticulous investigations on the molecular and physiological responses to acid shock are limited to cultivable oral bacteria [25], such as *S. mutans*, the gastric inhabitant *E. coli* [26] and *Helicobacter pylori* [27]. However, knowledge regarding the aciduric features of so far uncultivable microorganisms remains scarce. As a crucial technology used to access and explore the genomes and functional information of microorganisms including uncultivable ones, functional metagenomics [16] exhibit high application value. To the best of our knowledge, this is the first work that applied functional metagenomics to explore the acid-resistant genes of dental plaque microorganisms. The improved function-based screening strategy used here allowed the discovery of two genes involved in acid resistance, which could provide new clues for the development of therapeutic agents targeting the microorganisms to attenuate caries disease progression and prevent the occurrence of ECC.
Table 2. Bioinformatics analyses of five positive clones screened from *S. mutans* UA159.

| Code | U-57 | U-38 | U-41 | U-23 | U-8 |
|------|------|------|------|------|-----|
| **Seq length (bp)** | 1243 | 710  | 1172 | 1050 | 2332 |
| Site^a (Seq No.^b) | 1–239aa (516–1231bp) | 155–368aa (68–709bp) | 1–159aa (494–18bp) | 1–92aa (373–648bp) | 1–168aa (1079–1582bp) / 1–182aa (1695–2330bp) |
| **Gene** | *dnaK* | *dpC* | *rlmH* | *SMU_1725* | *mreD* / *gbpB* |
| **Protein/size (aa)^c** | Molecular chaperone DnaK /612 | Class III stress response-related ATP-dependent Clp protease, ATP-binding subunit/800 | 23S rRNA-methyltransferase RlmH /159 | Acylphosphatase /92 | Rod shape-determining protein MreD/168 / glucan-binding protein B/431 |
| **Putative function^d** | Participates in acid-resistant process | Participates in diverse cellular processes including membrane fusion, proteolysis and DNA replication | Specifically methylates the pseudouridine at position 1,915 in 23S rRNA. | Catalyzes the hydrolysis of various acyl phosphate carboxyl-phosphate bonds | Regulate cell shape / single-species biofilm formation on inanimate substrate |
| **E-value** | 2E-165 | 3E-103 | 2E-107 | 8E-58 | 2E-93 |
| **Identities** | 239/239 (100%) | 214/214 (100%) | 159/159 (100%) | 92/92 (100%) | 168/168 (100%) / 182/182 (100%) |

^a The protein range (site) that can be aligned with the known protein.
^b The nucleotide range (nt range) of predicted ORF within insert.
^c Most similar protein, currently identified with blastp: protein name and protein size.
^d Putative function of protein, currently identified with UniProt.
bp: base pair; aa: amino acids.
Based on successful use of functional metagenomics in various fields, it is well known that the quality of the metagenomic library plays an essential role in successful identification of functional genes. In this study, the following three measures were taken to ensure the quality of the library: (i) Successful construction of a high-copy dual-direction expression vector pSL. Disregarding frameshift mutations, this vector increases the likelihood that some exogenous fragments will be expressed, especially when the direction of the insert is consistent with the direction of the reverse lacZ promoter, which enhances the efficiency of library screening. (ii) Detection of library quality. To ensure that this library contains a variety of appropriately sized DNA fragments for screening, three different sizes of DNA were separately retrieved, the ligation system between them and the vector was optimized, and the ligation and transformation efficiencies were assessed. Finally, we obtained 11, 290 recombinant clones, containing 46.6 Mb DNA (the inserts of approximately 242 clones covered 1 Mb DNA); 85.5% of the colonies contained extraneous DNA fragments. Compared to other studies, 1 Mb DNA covers about 358 [28] or 157 clones [29], and the quality of our library was sufficient to perform the next step. (iii) The setting up of positive controls, pSL-orf1 and pSL-ffh, which can not only assist us to monitor whether there is a deviation in the course of the entire experiment but also allows us to initially evaluate the acidic capability of the positive clones by comparing them with the growth of the positive-control colonies. These improvements guarantee the effectiveness and credibility of acid-resistant genes obtained in this study.

The two acid-resistant genes obtained in this study were aligned with sequences from two common cariogenic genera, Actinomyces and Haemophilus [30], which represent two of seven dominant cariogenic phyla (Actinobacteria and Proteobacteria) [31]. Bioinformatics analysis found that the ORF in pSL-2–41 encodes two domains (the malic and Malic_M domains) that form the N-terminal region of the malate dehydrogenase protein, but the acid-resistance assay indicated that this fragment is sufficient to allow E. coli to survive in this hostile environment. Jain’s group [32] confirmed that the hosts E. coli DH5α and E. coli W3110 can tolerate acidic pH with the help of malate dehydrogenase, which mainly functions to pump out protons and restore the cytoplasmic pH back to neutral. Here, the additional heterologous copy of the malate dehydrogenase from H. haemolyticus (E. coli DH10B/pSL-2–41) can assist the host strain in tolerating the more hostile environment (pH 1.9) compared to the negative control (E. coli DH10B/pSL), which indicated that exogenous insert may enhance the acid resistance of host bacteria. Whether the detailed mechanism is consistent with E. coli requires further study. In addition, this protein can catalyze the interconversion between malate and oxaloacetate with the help of cofactor NAD⁺ or NADP⁺ and plays a critical role in the central oxidative pathway of living organisms [33]. In gram-negative bacteria, this protein generally forms a dimeric molecule and participates in the tricarboxylic acid (TCA) cycle. In addition, this protein can protect against oxidative stress via a reductive TCA cycle [34]. Thus, malate dehydrogenase plays an important role in helping the host resist acid and oxidative stress.

The other positive clone, pSL-1–31, encodes an ATP/GTP binding protein, which is involved in various biological processes; however, its acid-resistant properties and mechanisms remain unknown. Studies showed that this gene plays a pivotal role in a variety of biology processes in both prokaryotes and eukaryotes, including cell signalling [35], cell division, protein export, membrane transport, DNA repair, and multidrug resistance in tumours [36,37]. The functional range of the nucleotide binding protein family is gradually expanding with the accession of some new members. Harris et al. [38] described a gene involved in axon regeneration from the sciatic nerve of mice, named Nma I. Subsequent analysis revealed that it is a novel nucleotide binding protein related to zinc carboxypeptidases. In addition, the proteins encoded by disease-resistant genes in most plants contain ATP/GTP binding motifs, such as kinase-1a, kinase-2, and kinase-3a motifs, which confer resistance on the host tomato against Fusarium oxysporum, root-knot nematodes or potato aphids [39]. However, this study is the first to confirm the acid-resistance of the ATP/GTP binding protein. To preliminarily understand its possible mechanisms, the available mechanisms of acid resistance are summarized in Box1 in the Supplementary File. After acid exposure, the intracellular pH decreases, and DNA is damaged. The cells can resist the acid stress through a variety of defence mechanisms, such as regulating the flow of protons inside and outside cells [11,13] and neutralizing intracellular protons through producing alkaline compounds [40]. Further, the DNA repair enzyme system can also assist cells to resist acid stress through repair of the damaged DNA [12]. The acid-resistance property of the ATP/GTP binding protein could be attributed to the above mechanism. In addition, due to previously established consistency between various mechanisms of antibiotic resistance and acid-resistance [41], we speculate that multidrug resistance was probably involved in influencing acid-resistance. Of course, more in-depth studies are required to confirm this speculation.

Only a low number of functional genes were identified in this study, and no genes were obtained from S. mutans. One possible explanation for this is that addition of a single gene is often not sufficient to detect a
new phenotype, such as acid-resistance. If a single gene encodes only a subunit of a multi-unit enzyme that is critical for an enhanced acid resistance, no phenotype will result. Further, expression of the gene in question and successful translation may not be efficient in the *E. coli* host since some gram-positive proteins are toxic when translated within *E. coli*. An alternative explanation might be the limitations of the samples and method: (i) Studies have confirmed that *S. mutans* is not the only predominant caries-pathogenic bacterium [14,15]; depending on individual patients, it might not be the principal ECC bacterium [42]. Hence, in this study, the abundance of *S. mutans* might only occupy a small fraction of the collected caries samples, limiting the genetic information in our library and reducing the chance of obtaining functional genes. (ii) Some genetic information could be lost during the library construction process. Alternatively, it is known that the process of ligation of the various inserts into a vector is accompanied by a certain amount of randomness and uncontrollability [16], which can diminish or completely lose the genetic information of *S. mutans* and some other strains.

**Conclusion**

This is the first investigation that identifies the acid-resistant genes in the oral microbiota through improved functional metagenomics. Several known and unknown acid-resistant genes are obtained from the library of dental plaque and *S. mutans* in this study, which not only provide evidence for the feasibility of the experimental methods and the credibility of the results, but also promote studies to investigate the aciduric mechanism and the prevention and treatment of ECC. A larger library is needed to be constructed to obtain more functional genes in future work.

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**Disclosure statement**

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