Designing a preventative treatment for *Streptococcus mutans* induced dental caries using the antimicrobial properties of *Streptococcus salivarius* M18*

Logan Amos, Lujiong Danzhen, Phalak Dhingra, Erika Frohring, Isabella Haslinger Johnson, Sofia Kovacevic, Danyan Li, Chongming Liu, Yihan Ma, Anya Mathur, Sheilla Muligande, Annie Nguyen, Ruihan Qin, Carmen Reed, Aparajita Shimpi, Lingzhen Wang, Cavin Xue, Siwen Yang, Mira Zamarro, and Yuejiao (Vivian) Zhou Western Reserve Academy, Hudson OH

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*Dental caries is a primary oral health concern, affecting 90% of adults worldwide. Invasive treatments such as dental fillings and root canal therapies stop the degrading effects of caries and accelerate the remineralization process. Although current solutions are effective, they can be painful and expensive for patients. Additionally, these treatments are temporary and may require repetitive maintenance. Streptococcus mutans, a naturally occurring bacterial strain in the human oral cavity, is a principal pathogenic agent of caries and tooth decay. Saliva acidity and accumulation of food debris create an ideal environment for *S. mutans* proliferation. The bacterium becomes part of a biofilm on the surface of teeth, catalyzing plaque production. Studies demonstrate inhibition of the virulence and proliferation of *S. mutans* by *Streptococcus salivarius* A12. Researchers are exploring the possibility of using these bacteria to combat dental caries. Inspired by these findings, our project aims to forestall dental caries by using *S. salivarius* M18, another bacterial strain in the oral cavity that produces two enzymes, urease and dextranase. These enzymes break down dental plaque and decrease saliva acidity. M18 effectively minimizes tooth enamel erosion and impedes dental caries formation by disrupting biofilm formation and reducing salivary pH. We will transform the genes ureA-C and dex, encoding enzymes urease and dextranase, into an early enamel colonizer, Rothia aeria, to suppress caries. Our product will be a *R. aeria* paste applied after dental prophylaxis to forestall dental caries formation. This research demonstrates a biosynthetic method that improves current dental treatments and designs a biocompatible product for oral hygiene.*

Keywords: *Streptococcus mutans*, dental caries, *Streptococcus salivarius* M18, urease, dextranase, caries prevention, probiotics

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*The authors were mentored by Beth Pethel from Western Reserve Academy, Hudson OH. Please direct correspondence to Beth Pethel (pethelb@wra.net). This is an Open Access article, which was copyrighted by the authors and published by BioTreks in 2023. It is distributed under the terms of the Creative Commons Attribution License, which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited.*

Dental caries, also known as cavities or tooth decay, affects approximately 90% of adults ages 20-64. The primary cause of tooth decay is plaque, a sticky film of bacteria that forms on the tooth's surface and remains present for extended periods, as seen in
**Figure 1.** Several factors can contribute to plaque production but the most notable element is the overabundance of sugars in processed foods and drinks (Dental Plaque, n.d.). Plaque, in addition to microorganisms and food debris, contributes to the formation of dental cavities (Forssten et al., 2010, p. 291). Plaque-colonizing bacteria, such as the acid-producing *S. mutans*, damage the tooth structure, especially near carbohydrates such as sucrose or fructose (Forssten et al., 2010, p. 291). Tooth decay may result in tooth loss, tissue damage, or potentially life-threatening issues (Cavities/tooth Decay, 2022).

![Tooth anatomy](image1)

**Figure 1.** A breakdown of tooth anatomy alongside the progression of tooth decay in four stages; enamel decay, advanced enamel decay, dentin decay, and pulp decay. The severity of each stage is indicated by a pain scale that increases from low to high.

Biofilms, layers of bacteria that adhere to a surface (Figure 2), are abundant throughout the mouth. Biofilms protect teeth by maintaining a balance between harmless and destructive bacteria (Brown, 2022). However, they also contribute to plaque production and subsequent tooth decay, as seen in Figure 1. Although proper dental hygiene prevents the continuous growth of biofilms and reduces the risk of cavities, the biofilm still reforms four hours after its removal (Dhir, 2013).

As plaque reforms on clean teeth, certain organisms, known as early colonizers, repopulate the mouth before others. *S. mutans*, one of these early colonizers, is one of the main contributors to tooth decay and plaque formation (Lemos et al., 2019). After binding to the tooth and cultivating a biofilm, *S. mutans* synthesizes dextran (MicrobeWiki, 2016), a common sugar produced by bacteria (Redasani & Bari, 2015).

Another early colonizer of the mouth’s soft tissue, *S. salivarius* M18, produces the enzymes dextranase and urease. Dextranase breaks down dextran in the plaque of decaying teeth (Burton et al., 2013). Bacteria such as *S. mutans* synthesize dextran (MicrobeWiki, 2016). Dextran is found in beets and sugarcane (Infinita Biotech, 2022). Dextranase regulates the mildly toxic polysaccharide dextran levels in the mouth (Infinita Biotech, 2022). Through its breakdown of sugar, dextranase helps neutralize plaques that contribute to tooth decay (Stowik, 2016, p. 16).

The enzyme urease uses the compound urea to generate ammonia, which neutralizes certain acids in the mouth that may cause tooth decay (Chen et al., 2000, p. 4667). Extended exposure to acidic environments contributes to increased growth of harmful bacteria, and a pH value below 5.5 in the oral environment is a critical factor in the progression of cavity development (Matsui & Cvitkovitch, 2010). Urea is a naturally

![Biofilm formation](image2)

**Figure 2.** The three main steps of biofilm formation: adhesion, maturation, and detachment. Adhesion is the initial stage where microorganisms attach to a surface and begin to colonize, followed by maturation where the biofilm develops into a complex structure with different layers. The last step is detachment that perpetuates the cycle of biofilm formation.
occurring compound commonly found in saliva and urine. It is also used in agriculture due to its tendency to form pH-heightening ammonia and decrease acidity (American chemical society, 2021). Urease hydrolyzes the compound, utilizing its high nitrogen content to generate the alkali ammonia (Chen et al., 1996, p. 585). In doing so, it neutralizes acids created by dietary carbohydrates (Morou-bermudez et al., 2011, p. 1283). Urease is made up of three subunits that are encoded by the genes ureA-C. It is also one of the few enzymes that require a nickel ion to function. Consequently, the inclusion of the genes ureDEFG is present for the production of functional urease.

This project aims to insert the dextranase seen in Figure 3 and urease genes from S. salivarius into the early colonizer R. aeria. R. aeria is an effective carrier for the genes, as it colonizes the biofilms of teeth while S. salivarius only inhabits the oral tissue. R. aeria, similar to S. salivarius, possesses a Biosafety Level 1 rating and is gram-positive (Rothia aeria, n.d.).

![Figure 3](image1.png)

**Figure 3.** The process of dextranase breaking down dextran into degraded dextran. When dextranase breaks down dextran, it converts it into degraded dextran, which consists of smaller, non-toxic sugar molecules that can be easily metabolized by the body. By doing so, the dextran becomes non-toxic substances that no longer contribute to the formation of dental caries.

**Systems level**

The goal of this research is to develop a paste that contains enzymes for use in the oral cavity. To achieve this, we are introducing four genes (ureA, ureB, ureC, and dex) that encode for urease and dextranase enzymes into R. aeria. To do this, we will insert each of the target genes into a plasmid called pJPC13, seen in Figure 4, which includes a lac promoter, a lac terminator, and a chloramphenicol resistance marker. We will then transform the plasmid into R. aeria, which will be grown with Medium 92 containing chloramphenicol to isolate a single colony for inoculation. By growing the bacteria on chloramphenicol agar plates, we will ensure that only successfully transformed bacteria form colonies. As we do not need to extract the transformed bacteria, the final product will be the active R. aeria colonies in a paste. While the plasmid we are using is not specifically designed for R. aeria, our team has chosen a plasmid that is designed for a bacterium that genetically mimics it based on limited research.

![Figure 4](image2.png)

**Figure 4:** The pJPC13 plasmid has an origin, a lac promoter and terminator, chloramphenicol resistance, and a multiple cloning site that has the BamHI and EcoRI restriction enzymes that will be utilized. The pJPC13 has 5650 base pairs, which will increase once the genes are transferred into the plasmid.

**Device level**

Our project utilizes Rothia aeria A 1-17B as the chassis, primarily because it is an early colonizer of human teeth and is expected to have a minimal impact on the oral microbiome. We will use the ureA-C gene
cluster, which encodes for the enzyme urease, and the dex gene, which encodes for dextrase. Dextrase is an enzyme that breaks down dextrane sugar, converting it into a non-toxin. Specifically, we only require ureA, B, and C to produce urease. Urease catalyzes the reaction between urea and water, producing ammonia and carbon dioxide, as seen in Figure 5. Ammonia then increases the pH of the oral cavity, thus creating an environment that is not optimal for the growth of S. mutans. In addition, we will use the plasmid backbone pJPC13, which contains a lac promoter, a lac terminator, and a chloramphenicol resistance marker.

![Hydrolysis of urea via urease enzyme](image)

*Figure 5. The process of urea being hydrolyzed by urease to form ammonia and carbon dioxide. The newly produced ammonia helps balance the pH and contributes to preventing the growth of dental plaque.*

**Parts level**

Our project focuses on creating a paste containing dextranase and urease enzymes commonly found in S. salivarius. S. salivarius is a probiotic bacterium that can reduce oral infections by producing bacteriocin-like inhibitory substances. We plan to use the dex gene to produce dextranase and the ureA-C genes to produce urease. To simplify the process, we will use the lac promoter and lac terminator, which are present in the pJPC13 plasmid and are also related to lactose that is found in the oral cavity. The BamH1 and EcoR1 restriction enzymes present in the multiple cloning site of the pJPC13 plasmid will be used to insert the selected genes. As these enzymes occur only once in the multiple cloning site, we can avoid cleavage during insertion. Our selectable marker will be chloramphenicol, as *R. aeria* is not naturally resistant to this antibiotic.

**Safety**

The safety of *R. aeria* must be considered in topical use. The A 1-17B strain of *R. aeria* is classified as a Biosafety Level 1 organism and presents minimal risk to healthy humans (Strain Identifier, 2022; Wei et al., 2020, p 1). In some cases, *R. aeria* has been found at sites of infective endocarditis; however, the oral presence and consumption of the bacteria were not considered a direct cause (Nicodemo et al., 2014). The growth medium used for *R. aeria* cells, Medium 92, is categorized as Biosafety Level 1 (DSMZ, n.d.), meaning the media poses no hazards and does not require further safety measures. The electroporation technique this project will use to transform *R. aeria* poses no safety concerns.

In this project’s *R. aeria* treatment, urease production remains a concern. Urease has been shown to have toxic effects on human cells and can cause possible skin and eye irritation due to its enzymatic activity (Koniczna et al., 2012, p. 789). However, ammonium ions in the growth medium can limit urease transfer across the cell membrane, and glucose-inhibited enzyme synthesis can further regulate its production (Magaña-plaza & Ruiz-herrera, 1967, p. 1294). Dextranase is safe for human consumption, and allergic reactions are rare (Silano et al., 2020, p. 6309).

**Discussion**

This project aims to prevent dental caries using *S. salivarius* M18, another bacterial strain in the oral cavity that produces two enzymes, urease, and dextranase. By disrupting the natural formation of biofilm and reducing salivary pH, M18 should protect the teeth against enamel erosion and the formation of dental caries. By transforming the ureA-C and dex genes into Rothia aeria, a paste of the transformed *R. aeria* could be applied during regular dental
visits to prevent cavities.

However, this method may present potential hazards during practical use. Consultation with professionals and safety experiments would confirm that additional levels of R. aeria in the oral cavity do not cause adverse effects. As outlined in the safety section, its use poses minimal risk. The electroporation technique and use of dextranase are both safe for use in this project. While urease may cause adverse effects, the process of glucose-inhibited enzyme synthesis should neutralize these effects. If this project proves successful, the product could be used as a widespread preventative against plaque and dental caries.

Next steps

Though this project remains theoretical, testing must be considered in the laboratory environment. We will begin with growing Rothia aeria. To prepare for DNA extraction, we will make a stock solution of chloramphenicol and use it to isolate bacterial colonies. Since the bacterium is aerobic, an oxygen-free environment is not required for growth (Taira & Aoki, 2019). Growth will occur on Medium 92 at body temperature (37°C) to simulate the oral environment. Then, we will use electroporation to transform R. aeria. We will then apply our transformed bacteria to lab-grown plaque originating from teeth impacted by caries. This process will be repeated using different concentrations of the bacterial product to gauge the levels of each reaction in relation to the breakdown of plaque samples. Using a previously researched system for in vitro studies of the oral cavity, we will also test adequate quantities of the dextranase and urease enzymes on already formed biofilms that contain S. mutans (Mira et al., 2019). Through this procedure, we will compare the formation of biofilm in an environment that contains urease and dextranase to an environment that does not. Clinical trials can be conducted in small groups of dental providers and patients. If the project shows a reduction in the formation of dental caries, we can seek approval from dental associations to distribute the product to a larger group of dental patients. Ideally, this project will become a prevalent dental treatment for all.

Author contributions


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