Screening of *Streptococcus mutans* Sortase A Via Myricetin-Like Inhibitors: *In Vitro* Evaluation and Molecular Docking-Based Virtual

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**Abstract**

**Background:** Dental caries is one of the most common causes threatening human health globally. Sortase A (Srt A) as a transpeptidase, mediates the attachment of the *Streptococcus mutans* cell wall to dental surfaces by biofilm formation. Due to the development of multidrug-resistance bacteria, attempting to discover growth inhibitors is logical and promising.

**Objectives:** The current study aimed at the experimental and docking-based virtual screening of myricetin-like inhibitors for the inhibition of Srt A enzyme in *S. mutans* isolates.

**Methods:** Sixty-three *S. mutans* were isolated from pupils based on cultural, morphological, and biochemical characteristics (N = 150). After identifying the *srtA* gene using the polymerase chain reaction (PCR) with specific primers, a broth microdilution test was conducted according to CLSI-2020 criteria to determine the minimum inhibitory concentration (MIC) of myricetin. The *in silico* exploration of Srt A inhibitors was performed using AutoDock 4.2.6.

**Results:** The frequency of *S. mutans* isolates containing the *srtA* gene was 87.3% of which, fifty isolates (79.4%) were categorized as susceptible to myricetin (MIC ≤ 16 μg/mL). Of 20 ligands having a high degree of similarity with myricetin, the best docking results were related to ligand 2.

**Conclusion:** It was concluded that myricetin has an inhibitory effect on oral bacteria *in vitro*, and ligand 2 had the most negative binding energy (−4.66 kcal/mol) and favorably interacts with the key amino acid residues at the active site of Srt A. Accordingly, this ligand can be utilized as a lead compound for further studies to discover novel inhibitors targeting Srt A in *S. mutans*.

**Keywords:** *Streptococcus mutans*, Molecular docking, Myricetin, Sortase A

**Background**

Tooth infection such as dental caries has continuously become the main public health concern globally. Tooth decay contains biological, environmental, and lifestyle-related risk factors. Given biological agents, oral microflora specified in both the root and crown of tooth possess an important situation in dental health (1,2). The main pathogen of the root and coronal caries are viridans-group *Streptococci*, especially *Streptococcus mutans*. *S. mutans* uses the sortase A (Srt A) enzyme for anchoring several surface proteins to the cell walls and forms a biofilm on the tooth surface (3,4). The biofilm production, the so-called dental plaque, is one of the significant virulence features of *S. mutans* that adheres bacteria to the surface of the teeth. The tooth continuously displays enamel decalcification, ultimately resulting in its decay because of the biofilm generation and the acid-manufacturing capacity of carbohydrate fermentation by bacteria. Several studies have reported that the presence of *Streptococcus* spp. reduces in the cell wall after the removal of Srt A; therefore, this enzyme possesses a key function in the interactions between *Streptococcus* spp. and their host and is concerned as a promising target for the treatment of tooth caries (5-7). To date, some Srt A inhibitors have been discovered that are mostly derived from flavonoid compounds, including myricetin (8). However, only a few small molecules are available for inhibiting the transpeptidase by myricetin, which is a naturally benzo-α-pyrene flavonoid derivative. Antibiotics utilized in dentistry were found to raise bacterial resistance, thus naturally occurring ingredients can make a favorable contribution to treating dental caries. Moreover, the computer-based drug design has appeared as a popular procedure for conducting high-throughput virtual screening of potential medicinal compounds,
Experimental Characteristics and Study Population

In this experimental study, 150 samples were taken from the buccal and lingual surfaces of the posterior teeth of elementary school children aged 7-12 years (with the same sex ratio) in Gorgan, northern Iran. The students were excluded from the study if they had used antibiotics within the past three months or had systemic or immunodeficiency disorders. The wooden toothpicks samples were collected and transferred to the laboratory in sterile tubes containing the brain heart infusion broth (Merck, Germany) and buffer. After two hours of incubation at 37°C, the samples were homogenized by vortexing and plated on Mitis Salivarius Bacitracin agar (HiMedia, India) for 24 hours at 37°C in anaerobic conditions. The Streptococcus mutans isolates were identified based on the colony morphology, Gram-staining, and biochemical tests (hemolysis, catalase, bile esculin, optochin susceptibility, methyl red/Voges-Proskauer, and arginine dihydrolase), as well as the fermentation of mannitol, lactose, salicin, and trehalose (13). Then, the broth microdilution test was applied according to CLSI M100 (2020) criteria to determine the minimum inhibitory concentration (MIC) of myricetin (14). First, a suspension (1.5×10^8 CFU/mL) of the 24-hour bacterial cultures in Müller Hinton Broth (Merck, Germany) was prepared spectrophotometrically. After preparing myricetin suspension in 1% of dimethyl sulfoxide, the concentration range of myricetin stock (Sigma Aldrich, USA) was determined at 512 -1 µg/mL. After overnight incubation at 37°C in anaerobic conditions, the growth rate was measured and compared with that of the positive (without the myricetin) and negative (without bacterial suspension) control. The minimum concentration inhibiting bacterial growth up to 50% or 90%, compared to positive controls, is considered MIC_{50} and MIC_{90}, respectively. S. mutans PTCC35668 was used as the control. The DNA of S. mutans isolates was extracted by the glass bead/phenol-chloroform method. DNA was employed as a template for stA gene amplification using specific primers ‘F: 5'- CTGGGATCCAAAACCTCATAT and ‘R: 5'- CTCGTTATTTACATGAC-3 taken from Gene Bank No. AF162657 (15,16). Amplification was programmed for an initial denaturation at 94°C for 1 minute, followed by 31 cycles of denaturation at 92°C for 1 minute, annealing at 55°C for 2 minutes, extension at 72°C for 3 minutes, and a final extension at 75°C for 5 minutes. Polymerase chain reaction (PCR) products were electrophoresed on 1% agarose gel, and the bands were visualized under UV light after staining with ethidium bromide. After verifying the normality of data distribution using the Kolmogorov-Smirnov test, quantitative and qualitative data were analyzed by an independent t test and chi-square test, respectively.

Docking for Potential Inhibitors of S. mutans Sortase A

The appropriate crystallographic structure of the S. mutans Srt A enzyme containing the central catalytic site was selected from the PDB database with the ID 4TQX. The PubChem database and ChemSketch 12.0 molecular modeling program were used to obtain the chemical structure of myricetin-like molecules and create the structure of the molecules, respectively. The HyperChem 8.0 molecular modeling and The PyRx 0.8 virtual screening software were prepared to optimize molecules in terms of energy and evaluation of molecules. After obtaining the results, the binding affinity of all conformers related to each molecule was examined to select the best candidates for docking using AutoDock 4.2.6 molecular modeling simulation software. Electrostatic interactions for each protein atom were investigated by Autogrid 4.0.0, a program, which pre-calculates the grid map of interaction energy for different atom types. The box size 42*42*44 and point spacing 0.375 (equivalent to a quarter of the carbon-carbon bond length) were applied for inhibitor docking. The center of the two copper ions in the active site was designated as the center of the grid box. According to the calculation of the free energy of ligand binding, some ligands were selected for further docking. At this stage, after preparing the protein and its ligands in PDBQT format, the areas where the ligands interact with the protein were identified through the formation of a Grid Box. The best conformer for docking was selected using the relevant clustering diagram and physical parameters such as binding energy, inhibitory constant, and ligand efficiency. In other words, first, the best cluster was obtained according to the number of conformers in each cluster and the binding energy. Then, the best conformer was selected from each cluster by the three above-mentioned physical parameters. If the value of each physical parameter in each cluster was the lowest for a conformer, that conformer was considered as the final docking choice against Srt A. Finally, the interaction of myricetin and its analogs was evaluated and compared in more detail against the Srt A binding site via the Ligplot 4.5.3 computer program, which creates a schematic 2D representation of the protein-ligand complex (12).
Results

Results of In Vitro Assay

Streptococcus mutans (63, 42%) was the predominant streptococcal species. Other isolated species included Streptococcus sanguinis (28.11%) and Streptococcus oralis (8.47%). The presence of 348-bp fragments compared with the DNA marker indicated the presence of the srtA gene and confirmed the accuracy of the reaction. The results revealed that the frequency of S. mutans isolates containing srtA was 87.3% of which 79.4% (MIC ≤ 16 µg/mL) of the isolates were categorized as susceptible to myricetin. The comparative in vitro susceptibilities of the S. mutans isolates to myricetin showed that the MIC₅₀ was 128 µg/mL in comparison with the range of the reference strain (MIC = 32 µg/mL, P = 0.37) and was 4-fold lower than the MIC₅₀ value (512 µg/mL) against the S. mutans isolate (Figure 1). The changes to the MIC of myricetin demonstrated that the difference between the growth and lack of growth of S. mutans isolates containing srtA was significant (P = 0.01).

Data of Docking and Model Validation

In the present study, according to the binding site of the myricetin inhibitor in the Srt A enzyme and its inhibitory mechanism, first, a total of 20 ligands having a high degree of similarity with the myricetin molecule were selected and then analyzed via AutoDock (Figure 2). The matching between the structure of Srt A with myricetin and the ligands represented that the location of the active site is similar. The ligands were docked against Srt A for confirming the validity of the active site. The findings revealed that the key residues phe143 and thr146 were localized in the anticipated active center, and two or three hydrogen bonds between Srt A and its ligands were generated accordingly. This result demonstrated that the determined active site in the Srt A enzyme via AutoDock is comparatively precise and can be used for further virtual screening. The testing sets including 5 positive and 15 negative ligands were applied to measure the accuracy of AutoDock for screening Srt A inhibitors. The 50% of top-ranked ligands were docked via the Hawkins GB/SA score, accounted for 0.95 of the highest area under the receiver operating characteristic curve (data not shown). After studying the pharmacokinetic and quasi-pharmacological properties of the ligands, a total of 4 ligands having a high affinity for Srt A protein were selected as the candidate inhibitors. Each ligand was individually placed in the active site of the enzyme, and the amount of binding affinity, the number of hydrogen bonds with each amino acid, and the inhibitory power were obtained accordingly. The myricetin has a hydrogen bond with the amino acid phe143, which is 2.92 Å in length, and has a hydrophobic bond with 7 amino acids. Ligands 2 and 16 had three hydrogen bonds at the active site of the enzyme, namely, two hydrogen bonds with phe143 and one hydrogen bond with thr146. These ligands were also hydrophobically linked to 7 amino acids. Ligands 12 and 14 had two hydrogen bonds with he143, along with a total of 7 hydrophobic bonds (Figure 3).

The main molecule of myricetin has binding energy and an inhibitory constant of -4.19 kcal mol⁻¹ and 851.28, respectively. Ligand 2 had the lowest inhibitory constant and binding energy among other ligands. Moreover, this ligand had lower binding energy and higher inhibitory efficacy than those of myricetin (Table 1). The most potential ligands displayed more favorable binding energy in contrast to myricetin at -4.19 kcal mol⁻¹. Albeit myricetin belonged to flavonoid compounds, the binding energy was the lowest compared with other ligands.

Discussion

According to the World Health Organization, the problem of tooth decay remains worldwide, especially in socially disadvantaged groups (17). The tooth decay caused by drug-resistant oral bacteria has determined the need to discover new ways to combat these bacteria. Some researchers have confirmed the high role of Srt A in the formation of bacterial biofilms by modulating anchor surface proteins to the cell wall and eventually tooth decay (18, 19). In the present study, more than 80% of the isolates contained the srtA gene. Researchers in Canada (20) and Iran (19) observed that the inactivation of the srtA gene reduces biofilm formation. Therefore, according to widespread reports of drug resistance (13, 17), controlling it with natural compounds and enzymatic inhibitors seems reasonable. In this study, the changes to the MIC of myricetin showed that there was a significant difference between the growth and lack of growth of S. mutans isolates containing srtA. Progress in silico procedures has enabled virtual screening to have a promising effect on drug discovery. Docking-based virtual screening is based on predicting the affinity of each compound in the dataset via docking to an X-ray crystallographic structure. Docking also evaluates the binding mode of compounds with the amino acids present in the active pocket of the intended protein/enzyme (21, 22). The results of the current study indicated the substantial applicability and accuracy of molecular docking. Similarly, this validation method was exerted by Luo et al (11) and Sivaramakrishnan et al (12) for in silico identification of inhibitors targeting sortase. Similar studies have been recorded considering the recognition of the Srt A inhibitors of Streptococcus spp. As well (23-25). However, the exact mechanism behind...
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this inhibition is not yet clear. A variety of plant natural ingredients, especially flavonoid compounds, can hinder sortase. Additionally, the branched chains of flavone can reduce Srt A activity. (26). Some researchers observed that the carbonyl group can have an important function in the mechanism of Srt A suppression (11, 26). Some inhibitors such as morin and curcumin can decrease the activity of Srt A via interacting with key cysteine that takes part in breaking the bond T-G from the motif LPXTG in the enzyme (27-29). Overall, it seems that the ligand targeting Srt A in this study may have occupied the binding site of the motif LPXTG to decrease transpeptidation and thereby affect biofilm production. Considering that myricetin showed a good inhibitory effect at high concentrations *in vitro*, docking-based virtual screening opens a promising window toward the identification of ligands having a higher inhibitory potential than chemical and natural compounds. Likewise, other researchers found that molecular docking can identify ligands having a higher inhibitory potential against *S. mutans* Srt A when compared to natural ingredients (30, 31).

**Conclusion**

According to the results of the present study, myricetin exhibited antibacterial activity and inhibitory effects against *S. mutans* isolates *in vitro*. Docking-based virtual...
screening in this research led to the identification of a promising ligand against S. mutans Srt A. This ligand was the best compound for the selective inhibitor because it favorably interacts with important amino acids located in the active site of Srt A. However, further in vitro and in vivo studies are required to determine the Srt A inhibitory activity of the ligand identified in this research.

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Authors’ Contribution
LF contributed to the study concept and edited the final manuscript. MM performed laboratory examinations and interpreted the data. All authors discussed the results and implications and provided their comments during all stages.

Conflict of Interests
The authors declare that there is no conflict of interests.

Ethical Issues
The study was performed in accordance with the Declaration of Helsinki and approved by the Ethics Committee of I.A.U, Aliabad Branch (code No.: 1398.006), Golestan, Iran.

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Table 1. The Value of Physical Parameters Associated With Each Ligand Targeting Sortase A

| Ligand Name | Myricetin | Ligand 2 | Ligand 12 | Ligand 14 | Ligand 16 |
|-------------|-----------|----------|-----------|-----------|-----------|
| Binding energy (kcal mol⁻¹) | -4.19 | -4.66 | -4.3 | 4.46 | -4.33 |
| Ligand efficiency (kcal mol⁻¹) | 0.18 | 0.23 | 0.2 | 0.2 | 0.2 |
| Inhibition constant (Ki) | 851.28 | 382.64 | 699.87 | 534.23 | 673.22 |

Note: Ligand 2: 2-(3,4-dihydroxyphenyl)-7-hydroxy-4H-chromen-4-one; Ligand 12: Luteolin; Ligand 14: Quercetin; Ligand 16: Robinetin.
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