Identification of Potential Glucosyltransferase Inhibitors from Cinnamic Acid Derivatives Using Molecular Docking Analysis: A Bioinformatics Study

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Abstract

Background: Dental caries is one of the most common oral chronic diseases. Streptococcus mutans is the main pathogenic bacteria playing a role in degrading the mineral texture of the teeth. Glucosyltransferase (GTFase) of S. mutans is responsible for producing glucan, which is the main exopolysaccharide found in the cariogenic biofilms. Further, previous studies have reported that cinnamic acid diminished biofilm formation of S. mutans. Therefore, we hypothesized that cinnamic acid and its derivatives might act as GTFase inhibitors.

Methods: The binding affinity of a total of 12 plant-based compounds including cinnamic acid and its 11 derivatives to the GTFase active site were examined by utilizing the AutoDock tool. The possible interactions between top-ranked cinnamic acid derivatives and the residues within the GTFase catalytic site were also taken into consideration.

Results: Five of the cinnamic acid derivatives including rosmarinic acid (RA), cynarine, chlorogenic acid (CGA), caffeic acid 3-glucoside, and N-p-coumaroyllyramine demonstrated inhibitory effects on GTFase at nanomolar concentration. Stabilizing interactions such as π–π stack pairing and pi-charge interactions were detected between top-ranked GTFase inhibitors and residues within the enzyme active site.

Conclusions: The present study suggests that RA, cynarine, CGA, caffeic acid 3-glucoside, and N-p-coumaroyllyramine might have protective effects on dental caries, and therefore, may be considered as anti-tooth caries compounds.

Keywords: Cinnamic acid, Dental caries, Glucosyltransferase, Inhibitor, Molecular docking

Background

Dental caries (tooth decay) is the most frequent oral disorder worldwide that has destructive effects on the quality of life. According to the World Health Organization (WHO) report, it affects 60-90% of children, especially in developing countries, although it can affect people at any age. Several factors have been reported to be involved in the dental caries occurrence. While matrix metalloproteinases (MMPs) contribute to degrading the organic tissue within the teeth, tooth-adherent bacteria are involved in metabolizing sucrose, leading to acid production and demineralization of the mineral structure of teeth (1-6). Streptococcus mutans is the most prevalent bacteria extracted from human cariogenic dental cavities (7,8). It mediates the synthesis of exopolysaccharides, the main texture of cariogenic biofilms, resulting in more bacterial adhesion (9,10). Glucan is found to be the most common exopolysaccharides synthesized by glucosyltransferase (GTFase) of S. mutans. Therefore, GTFase inhibition has been considered as an effective strategy to diminish dental biofilm formation and to prevent dental caries occurrence (11-13).

Cinnamic acid is an aromatic carboxylic acid compound that can be synthesized by deamination of phenylalanine and is primarily found in Cinnamomum cassia, Panax ginseng, vegetables, grains, and honey (14,15). Figure 1 illustrates the chemical structure of cinnamic acid achieved by the ACD/ChemSketch version 12.01. Cinnamic acid derivatives are naturally produced by modifying their aromatic ring and the acrylic acid group (16). Several pharmaceutical features (i.e., antimicrobial, anticancer, and anti-inflammatory) have been reported for cinnamic acid and its derivatives (17,18). In addition to antibacterial activities of cinnamic acid derivatives, Mojtabavi et al (19) demonstrated that the combination of cinnamic acid and laccase resulted in approximately 90% reduction in S. mutans biofilm formation.

In the present study, we hypothesized that cinnamic acid and its derivatives might act as GTFase inhibitors in S. mutans. The binding affinity of cinnamic acid and its 11 derivatives to the GTFase active site were estimated by molecular docking analysis. Five of the tested compounds were revealed to block the GTFase catalytic site at the nanomolar scale. Two-dimensional structures of the tested...
compounds in this study are presented in Table 1.

Methods
Structural Preparation
The three-dimensional structure of GTFase was collected from the Protein Data Bank (PDB ID: 3AIE) with a criteria of X-ray resolution = 2.10 Å, which is available at https://www.rcsb.org. A total of eight subunits were included within the 3AIE file. Chain A, with a total of 844 residues, was chosen for computational dockings. It should be noted that the molecular energy of GTFase was optimized prior to molecular docking analysis by using the Swiss-PdbViewer version 4.1.0, which is available at http://www.expasy.org/spdbv (20).

The binding affinity of 12 compounds including cinnamic acid and its derivatives to the GTFase catalytic site was examined by using the AutoDock software (version 4.0), which is available at http://autodock.scripps.edu (21). The components included rosmarinic acid (RA), cynarine, chlorogenic acid (CGA), caffeic acid 3-glucoside, N-p-Coumaroyltyramine, caffeic acid phenethyl ester (CAPE), o-Coumaric acid, caffeic acid (CA), ferulic acid, sinapinic acid, p-coumaric acid, and cinnamic acid. In addition, Acarbose (PubChem ID: 41774), Maltose (PubChem ID: 6255), and WP1066 (PubChem ID: 11210478) were considered as standard inhibitors of GTFase (22, 23). All ligand structures were firstly achieved as SDF files from the public repository for information on chemical substances and their biological activities (PubChem database), which is available at https://pubchem.ncbi.nlm.nih.gov (24-26). Thereafter, the SDF files were converted to PDB formats using the web-server of the Computer-Aided Drug Design (CADD) Group of the Chemical Biology Laboratory (CBL), NCI, and NIH located at the Frederick National Laboratory for Cancer Research (FNLCR), formerly NCI-Frederick (http://cactus.nci.nih.gov/chemical/structure). The energy minimization of small molecules was also executed before binding energy predictions using the HyperChem software (version 8.0.10) (27).

Molecular Docking and Post-docking Analyses
A windows-based computer (with the criteria of installed memory: 32 GB, processor: Intel Core i7, and system type: 64-bit) was used for in silico simulations. The AutoDock tool imposes limited flexibility on the protein. It uses an accurate free energy force field based on a Lamarckian genetic algorithm, leading to a rapid ligand conformation prediction within the binding site and estimating the Gibbs free binding energy from the following algorithm (28-30):

\[
\Delta G_{\text{binding}} = \text{Intermolecular Energy} + \text{Total Internal Energy} + \text{Torsional Free Energy} - \text{Unbound System's Energy}
\]

The active site of GTFase was considered as a receptor for the ligands. The grid box settings in the AutoDock tool (included spacing, 0.375 Å; X-dimension, 58; Y-dimension, 74; Z-dimension, 52; X-center, 190.161; Y-center, 46.104; and Z-center, 191.584. A total of 14 amino acids were identified to be located within the GTFase catalytic site from the Ito and colleagues’ study (22), including Tyr430, Leu433, Leu434, Arg475, Asp477, Asn481, Glu515, Trp517, Arg540, His587, Asp588, Asp909, Tyr916, and Gln960. It is worth mentioning that a total of 50 runs were set for each ligand.

For each ligand, the lowest \(\Delta G_{\text{binding}}\) within the largest cluster of results was considered for post-docking analyses including protein-ligand complex imaging and interaction mode study. The BIOVIA Discovery Studio Visualizer version 19.1.0.18287 (https://discover.3ds.com/discovery-studio-visualizer-download) was used for visualizing the two-dimensional images of interactions between top-ranked inhibitors and residues within the GTFase active site as well as demonstrating the three-dimensional docked pose of the top-ranked CA derivatives.

Results
Binding Affinity and Interaction Modes Between GTFase and Small Molecules
The Gibbs free energy changes of interactions between GTFase and the studied compounds were estimated using the AutoDock tool to identity potential GTFase inhibitors for combating dental caries. According to the results, a total of five cinnamic acid derivatives including RA, cynarine, CGA, caffeic acid 3-glucoside, and N-p-coumaroyltyramine were predicted to bind to the GTFase catalytic site at the nanomolar scale (nM); therefore, these cinnamic acid derivatives were considered as top-ranked GTFase inhibitors in the present study. It was also estimated that CAPE, o-Coumaric acid, and CA could inhibit the GTFase activity at the micromolar scale (uM). Moreover, ferulic acid, sinapinic acid, p-coumaric acid, and cinnamic acid revealed a dismal affinity to the GTFase active site, based on the inhibition constant values (Ki) calculated for these molecules that were predicted to be at the millimolar (mM) scale. In addition, acarbose demonstrated the highest binding affinity to the GTFase active site among control inhibitors followed by maltose and WP1066. Moreover, the \(\Delta G_{\text{binding}}\) of GTFase with RA, cynarine, and CGA was predicted to be more negative than that of WP1066, suggesting that these three compounds can attach to the GTFase catalytic site more tightly than the WP1066 (Figure 2).

The estimated \(\Delta G_{\text{binding}}\) and Ki values for all tested compounds in this study are presented in Table 2. The details of energies among top-ranked cinnamic acid derivatives and GTFase catalytic site are illustrated in Table 3. The interaction modes between top-ranked cinnamic derivatives

![Figure 1. Chemical Structure of cinnamic acid.](image-url)
### Table 1. Two-dimensional Structures of the Tested Ligands in This Study for the Identification of Potential GTFase Inhibitors

| Compound Name       | Sources                                                                 | Two-Dimensional Structure | Reference |
|---------------------|-------------------------------------------------------------------------|----------------------------|-----------|
| RA                  | Rosemary, *Perilla frutescens*, and *Salvia miltiorrhiza*               | ![RA Structure](image1)   | (56)      |
| Cynarine            | *Vernonia anthelmintica*                                                | ![Cynarine Structure](image2) | (57)      |
| CGA                 | Apples, artichoke, betel, burdock, carrots, coffee beans, eggplants, *Eucommia*, and grapes | ![CGA Structure](image3)  | (58)      |
| Caffeic Acid 3-glucoside | American cranberry                                                         | ![Caffeic Acid 3-glucoside Structure](image4) | (59)      |
| N-p-Coumaroyltyramine | *Crinum biflorum Rotb*                                                  | ![N-p-Coumaroyltyramine Structure](image5) | (60)      |
| Compound Name | Sources                                                                 | Two-Dimensional Structure | Reference |
|---------------|-------------------------------------------------------------------------|---------------------------|-----------|
| CAPE          | Propolis and grains                                                     |                           | (61, 62)  |
| O-Coumaric Acid | Barley, rye, corn, berries, grapes, apples, beans, peas, hazelnut, pecan, celery, tomato, garlic, flax, mustard, and tea |                           | (63-66)   |
| CA            | Blueberries, kiwis, plums, cherries, and apples                         |                           | (67)      |
| Ferulic acid  | Grains, spinach, parsley, grapes, rhubarb, and cereal seeds             |                           | (68)      |
| Sinapinic Acid | Rhizome of *Hydnophytum formicarum*                                     |                           | (69)      |
| P-Coumaric acid | Barley, rye, corn, berries, grapes, apples, beans, peas, hazelnut, pecan, celery, tomato, garlic, flax, mustard, and tea |                           | (63-66)   |
Possible GTFase Inhibition by Cinnamic Acid Derivatives

| Compound Name | Sources | Two-Dimensional Structure |
|---------------|---------|--------------------------|
| Cinnamic acid | Cinnamomum cassia, Panax ginseng, grains, and honey | ![Two-Dimensional Structure](image) |

Table 1.

| PubChem ID | Ligand Name     | ΔG_{binding} | K_i   |
|------------|-----------------|--------------|-------|
| 5281792    | RA              | -9.10        | 212.34 nM |
| 6124212    | Cynarine        | -8.97        | 265.18 nM |
| 1794427    | CGA             | -8.70        | 419.70 nM |
| 5281759    | Caffeic acid 3-glucoside | -8.42          | 669.37 nM |
| 5372945    | N-p-Coumaroyltyramine | -8.27          | 864.04 nM |
| 5281787    | CAPE            | -7.92        | 1.56 uM |
| 637540     | O-Coumaric acid | -5.01        | 212.28 uM |
| 689043     | CA              | -4.32        | 687.05 uM |
| 445858     | Ferulic acid    | -4.01        | 1.16 mM |
| 637775     | Sinapinic acid  | -3.99        | 1.18 mM |
| 637542     | p-Coumaric acid | -3.56        | 2.47 mM |
| 444539     | Cinnamic acid   | -3.17        | 4.74 mM |
| 41774      | Acarbose (Ctrl)| -11.45       | 138.56 pM |
| 6255       | Maltose (Ctrl)  | -10.94       | 9.53 mM |
| 11210478   | WP1066 (Ctrl)   | -8.58        | 511.17 nM |

Table 2.

Table 2. Estimated Binding Energy and K_i Value of all Compounds Tested in This Study After Molecular Docking With GTFase

**Discussion**

Tooth decay is one of the most common chronic diseases worldwide (32). It is a multifactorial disorder in which matrix metalloproteinases and *S. mutans* are most responsible for degrading the organic and mineral texture of the teeth, respectively (33,34). GTFase of *S. mutans* plays an essential role in biofilm formation, leading to more bacterial cohesion, acid production, and dental caries (11-13,35). To discover potential GTFase inhibitors, the binding affinity of several plant-based compounds including cinnamic acid and its 11 derivatives with GTFase catalytic sites were estimated using a molecular docking approach. The obtained results predicted that RA, cynarine, CGA, caffeic acid 3-glucoside, and N-p-Coumaroyltyramine could potentially inhibit the GTFase active site at the nanomolar scale. In addition, it was found that three of these compounds (i.e., RA, cynarine, and CGA) were more tightly bonded to the enzyme compared with WP1066 as one of the standard inhibitors of the enzyme.

CA is a water-soluble metabolite that can be synthesized in herbs with several beneficial properties such as antioxidant, antiviral, antibacterial, antitumorigenic, as well as liver and cardiovascular protective effects (36). Sorgi et al (37) conducted a study to examine CA’s antibacterial and anti-inflammatory properties in macrophage response against *S. mutans*. The authors demonstrated that *S. mutans* displayed an antibacterial effect at the half-maximal inhibitory concentration (IC_{50}) = 2.938 mM without illustrating cytotoxicity. Moreover, CA led to downregulation of nitrite, tumor necrosis factor alpha, and prostaglandin E_2 through the nuclear factor kappa B dependent pathway, demonstrating its anti-inflammatory effects within the macrophages. Furthermore, Nakamura et al (38) reported that CA solution significantly increased the antibacterial effect of 385 nM LED irradiation against cariogenic *S. mutans* biofilms. According to the present study, CA is a promising candidate for the treatment of dental caries.

**Figure 2.** Comparing the Binding Affinity Between the GTFase Catalytic Site and its Top-ranked Inhibitors From Cinnamic Acid Derivatives. Note. FTFase: Glucosyltransferase. Acarbose, Maltose, and WP1066 were considered as the standard GTFase inhibitors. The x-axis corresponds to the ligand name. The y-axis represents the score of ΔG_{binding} in terms of kcal/mol.
results, caffeic acid 3-glucoside was estimated to bond to the GTFase active site at the nanomolar scale (Ki = 669.37 nM) with a considerable ΔGbinding of −8.42 kcal/mol, while the estimated binding energy between CA and GTFase catalytic site was −4.32 kcal/mol, suggesting that binding of a sugar moiety to CA has enhanced the binding affinity of the compound to GTFase active site. Caffeic acid 3-glucoside demonstrated three hydrogens one hydrophobic and one electrostatic interaction with the Asp477, Glu515, and Asp588 inside the GTFase active site. A pi-charge was detected between caffeic acid 3-glucoside and Asp477, Glu515, and Asp588 inside the GTFase active site. The authors demonstrated that PVE and RA could significantly reduce lipopolysaccharide-induced damages in gingival fibroblasts due to their anti-inflammatory purposes in periodontal diseases. Further, previous studies have reported a link between inflammatory periodontal diseases and dental plaque (46).

According to the present results, RA demonstrated the highest binding affinity to GTFase active site among 12 cinnamic acid derivatives. It was predicted that RA could attach to the GTFase catalytic site at the nanomolar concentration (Ki = 212.34 nM) with a salient ΔGbinding of −9.10 kcal/mol. It revealed three hydrogen, two electrostatic, and one miscellaneous interactions with the enzyme.

### Table 3. Details of Energies Between Top-ranked Cinnamic Acid Derivatives, Control Inhibitors, and GTFase Catalytic Site Achieved From Molecular Docking Analysis

| Ligand Name | Final Intermolecular Energy (kcal/mol) | Final Total Internal Energy (kcal/mol) | Torsional Free Energy (kcal/mol) | Unbound System’s Energy (kcal/mol) | Estimated Free Energy of Binding (kcal/mol) |
|-------------|---------------------------------------|---------------------------------------|---------------------------------|-----------------------------------|--------------------------------------------|
| RA          | -8.1                                  | -5.72                                 | 3.88                            | -0.84                             | -9.10                                      |
| Cynarine    | -10.97                                | -4.67                                 | 5.37                            | -1.3                              | -8.97                                      |
| CGA         | -6.53                                 | -6.97                                 | 3.58                            | -1.22                             | -8.70                                      |
| N-p-coumaroyltetramine | -8.32 | -2.64                                 | 2.39                            | -0.3                              | -8.42                                      |
| Caffeic acid 3-glucoside | -8.2 | -4.75                                 | 3.58                            | -0.94                             | -8.27                                      |
| Acarbose (Ctrl) | -11.33 | -11.72                                | 6.56                            | -3.04                             | -13.45                                     |
| Maltose (Ctrl) | -8.95 | -7.01                                 | 3.58                            | -1.44                             | -10.94                                     |
| WP1066 (Ctrl) | -8.47 | -2.6                                 | 1.79                            | -0.7                              | -8.58                                      |

Note: GTFase: Glucosyltransferase; RA: Rosmarinic acid; CGA: Chlorogenic acid.

### Table 4. Interaction Modes Between Top-ranked Cinnamic Acid Derivatives and Residues Inside the GTFase Active Site

| Ligand Name | Hydrogen Bond (Distance Å) | Hydrophobic Interaction (Distance Å) | Electrostatic: Pi-charge (Distance Å) | Miscellaneous (Distance Å) | Halogen (Distance Å) |
|-------------|----------------------------|-------------------------------------|--------------------------------------|---------------------------|----------------------|
| RA          | Asp477 (4.11, 4.37, 4.39); | NA                                  | Glu515 (7.65); Asp909 (6.40)         | Tyr916 (3.60)             | NA                   |
| Cynarine    | Asp909 (4.23); Asn481 (3.99); Glu509 (1.83); Ser589 (3.21); Asp593 (3.88) | Tyr916 (6.97); Trp517 (4.91) | NA                                  | NA                      | NA                   |
| CGA         | Asn481 (3.98, 4.73) | Trp517 (6.01) | Asp909 (7.51) | NA | NA |
| N-p-Coumaroyltetramine | Asn481 (4.86); Ala478 (3.53); Glu592 (4.73, 4.78) | Phe907 (7.06); Tyr916 (5.55); His587 (7.17) | NA | NA | NA |
| Caffeic acid 3-glucoside | Asp477 (3.53, 4.31); Glu515 (4.83) | Trp517 (6.06) | Asp588 (5.86) | NA | NA |
| Acarbose (Ctrl) | Gly429 (3.60); Asp477 (4.35); Asp909 (4.61); Glu515 (4.98); Asn481 (4.81); Trp517 (4.68); Gly428 (3.51); Ser518 (3.89) | NA | NA | NA | NA |
| Maltose (Ctrl) | Asn481 (4.04, 4.98); Glu592 (4.62, 4.91) | NA | NA | NA | NA |
| WP1066 (Ctrl) | Asp588 (4.29) | Trp517 (5.99); His587 (5.90); Leu433 (7.12); Ala478 (6.31) | Glu515 (7.37); Asp909 (6.20) | NA | His587 (5.72) |

Note: GTFase: Glucosyltransferase; RA: Rosmarinic acid; CGA: Chlorogenic acid. Acarbose, Maltose, and WP1066 were considered control inhibitors of the enzyme.

### Table 5. Analysis of Energy Calculation from Molecular Docking Analysis

| Ligand Name | Hydrogen Bond (Distance Å) | Hydrophobic Interaction (Distance Å) | Electrostatic: Pi-charge (Distance Å) | Miscellaneous (Distance Å) | Halogen (Distance Å) |
|-------------|----------------------------|-------------------------------------|--------------------------------------|---------------------------|----------------------|
| RA          | Asp477 (4.11, 4.37, 4.39); | NA                                  | Glu515 (7.65); Asp909 (6.40)         | Tyr916 (3.60)             | NA                   |
| Cynarine    | Asp909 (4.23); Asn481 (3.99); Glu509 (1.83); Ser589 (3.21); Asp593 (3.88) | Tyr916 (6.97); Trp517 (4.91) | NA                                  | NA                      | NA                   |
| CGA         | Asn481 (3.98, 4.73) | Trp517 (6.01) | Asp909 (7.51) | NA | NA |
| N-p-Coumaroyltetramine | Asn481 (4.86); Ala478 (3.53); Glu592 (4.73, 4.78) | Phe907 (7.06); Tyr916 (5.55); His587 (7.17) | NA | NA | NA |
| Caffeic acid 3-glucoside | Asp477 (3.53, 4.31); Glu515 (4.83) | Trp517 (6.06) | Asp588 (5.86) | NA | NA |
| Acarbose (Ctrl) | Gly429 (3.60); Asp477 (4.35); Asp909 (4.61); Glu515 (4.98); Asn481 (4.81); Trp517 (4.68); Gly428 (3.51); Ser518 (3.89) | NA | NA | NA | NA |
| Maltose (Ctrl) | Asn481 (4.04, 4.98); Glu592 (4.62, 4.91) | NA | NA | NA | NA |
| WP1066 (Ctrl) | Asp588 (4.29) | Trp517 (5.99); His587 (5.90); Leu433 (7.12); Ala478 (6.31) | Glu515 (7.37); Asp909 (6.20) | NA | His587 (5.72) |

Note: GTFase: Glucosyltransferase; RA: Rosmarinic acid; CGA: Chlorogenic acid. Acarbose, Maltose, and WP1066 were considered control inhibitors of the enzyme.
Possible GTFase Inhibition by Cinnamic Acid Derivatives

Figure 3. Left: Two-dimensional Images of Interaction Modes Between (a) Rosmarinic Acid, (b) Cynarine, (c) Chlorogenic acid, (d) Caffeic acid 3-glucoside, (e) N-p-Coumaroyltyramine, (f) Acarbose, (g) Maltose, (h) WP1066, and Residues Within the GTFase Catalytic Site. Right: Three-dimensional Docked Pose of the Corresponding Ligands. Note: GTFase: Glucosyltransferase.
Asp477, Glu515, Tyr916, and Asp909 within the GTFase active site. Further, the interactions among Glu515, Asp909, and SA were of pi-charge type.

CGA is an ester of CA and quinic acid with antioxidant activity (47,48) which is mainly found in coffee, apples, berries, pears, and aubergines (49). Previous studies have shown that coffee has exhibited anti GTFase activity in S. mutans, leading to dental caries prevention (11,50). Moreover, Lin et al (51) triggered a study to examine the effect of CGA on tooth decay in rats. The authors investigated the antibacterial properties of CGA on S. mutans ATCC 10449 and S. sobrinus OMZ65. The obtained results revealed that the MIC and MBC of S. mutans were 2.5 and 7.5 mg/mL, respectively. Therefore, the authors suggested that CGA may be considered as a potential anti tooth caries compound by inhibiting the growth of S. mutans. Hu et al (52) found that CGA elevated the osteogenic differentiation of human dental pulp stem cells through the Wnt signaling pathway. The authors suggested that CGA may be useful for alveolar bone damage repairmen in patients with periodontal disease. According to the results of the present study, CGA formed two hydrogen, one hydrophobic, and one pi-charged interactions with Asn481, Trp517, and Asp909 within the GTFase active site. It was also estimated that CGA can attach to the GTFase catalytic domain at the nanomolar concentration (Ki = 419.70 nM) with a ∆G binding of -8.70 kcal/mol.

N-p-coumaroyltyramine is a phenolic compound primarily found in Tribulus terrestris that has demonstrated several pharmaceutical properties such as anti-cariogenic effect against S. mutans. According to previous studies, T. terrestris have significantly reduced the growth, adhesion, acid production, as well as synthesis of glucan within the S. mutans (53,54). According to the results of the present study, N-p-Coumaroyltyramine could block the GTFase activity at the nanomolar scale (Ki = 864.04 nM) with a ∆G binding = -8.27 kcal/mol, suggesting the potential anti-tooth decay property of the compound. N-p-Coumaroyltyramine demonstrated four hydrogen and three hydrophobic interactions with Ala478, Asn481, His587, Gln592, Phe907, and Tyr916 inside the GTFase catalytic site.

Cynarine is a polar component mainly found in the roots of Echinacea angustifolia (55). It revealed a considerable binding affinity to the GTFase catalytic site (ΔG binding = -8.97 kcal/mol) and was found to inhibit the enzyme activity at the nanomolar scale (Ki = 265.18 nM). Cynarine displayed five hydrogen and two hydrophobic interactions with the Asn481, Glu509, Trp517, Ser589, Asp593, Asp909, and Tyr916 within the GTFase active site. It should be noted that the electrostatic between Trp517 and cynarine is of pi-pi stack pairing type.

Previous studies have demonstrated that acarbose and maltose are potent GTFase inhibitors (22). Moreover, Tsurumaki et al (23) reported that WP1066 (PubChem ID: 11210478), a well-known JAK/STAT3 signaling pathway inhibitor, revealed inhibitory effects on ceramide GTFase. Therefore, these three compounds were considered as control inhibitors of GTFase in this study. Acarbose showed a salient binding affinity to the GTFase active site with ΔG binding and Ki values of -13.45 kcal/mol and 138.56 picomolar (pM), respectively. It demonstrated eight hydrogen bonds with the Gly428, Gly429, Asp477, Glu515, Trp517, Ser518, Asn481, and Asp909 residues within the GTFase catalytic site. In addition, maltose revealed a high binding affinity to the GTFase active site. The ΔG binding and Ki values for this compound were calculated to be -10.94 kcal/mol and 9.53 nM, respectively. It illustrated four hydrogen interactions with the Asn481 and Gln592 residues within the GTFase catalytic site. In addition, maltose revealed a high binding affinity to the GTFase active site. The ΔG binding and Ki values for this compound were calculated to be -10.94 kcal/mol and 9.53 nM, respectively. It illustrated four hydrogen interactions with the Asn481 and Gln592 residues within the GTFase catalytic site. Further, WP1066 formed one hydrogen, four hydrophobic, and two pi-charge interactions with the Leu433, Ala478, Glu515, Trp517, His587, Asp588, and Asp509 within the GTFase active site. The ΔG binding and Ki values for this compound were calculated to be -10.94 kcal/mol and 9.53 nM, respectively. It illustrated four hydrogen interactions with the Asn481 and Gln592 residues within the GTFase catalytic site. Further, WP1066 formed one hydrogen, four hydrophobic, and two pi-charge interactions with the Leu433, Ala478, Glu515, Trp517, His587, Asp588, and Asp509 within the GTFase active site.

Figure 4. Possible Connections Between Top-ranked Cinnamic Acid Derivatives and Amino Acids Incorporated Within the GTFase Catalytic Site. Note: GTFase: Glucosyltransferase.
values of WP1066 regarding the enzyme were calculated to be $-8.58$ kcal/mol and $511.17$ nM, respectively.

**Conclusions**

The present study suggests that RA, cyanine, CGA, caffeic acid 3-glucoside, and N-p-coumaroyltyramine potentially have inhibitory effects on GTFase of *S. mutans* at nanomolar concentration. Therefore, these compounds may be helpful for preventing dental caries; however, these findings should be confirmed by wet-lab techniques.

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**Authors’ Contributions**

AT and ZK designed the study. FG and AT performed Docking operations. AT processed all images were. AT, ZK, and FG analyzed and discussed the results. AT wrote the manuscript, and finally, all authors read and approved the final version of the manuscript.

**Availability of Data and Materials**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Conflict of Interests**

The authors declare no conflict of interests.

**Ethical Approval**

Not applicable.

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