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The Effect of R382W Mutation on Citrus paradisi Flavonol Specific 3-O-Glucosyltransferase

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The Effect of R382W Mutation on *Citrus paradisi* Flavonol Specific 3-O-Glucosyltransferase

By

Kathleen Alma King

An Undergraduate Thesis Submitted in Partial Fulfillment of the Requirements for the University Honors Scholars Program
Honors College
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Date
ABSTRACT

Flavonoids are a class of plant metabolites with C6-C3-C6 structure responsible for many biological functions, including coloration and defense. *Citrus paradisi*, grapefruit, contains a wide variety of flavonoids which are grouped by the extent of modification, examples of which are flavonols, flavones, and flavanones. A major modification is the addition of glucose by glucosyltransferases (GTs) to stabilize the structure and provide ease of transport. Glucosyltransferases can be highly substrate and regiospecific. With Cp3OGT, glucose is added at the 3-hydroxy position. This 3GT only accepts flavonols as its substrate; however, a *Vitis vinifera* (grape) 3-GT can accept both flavonols and anthocyanidins. Homology modeling using the crystallized structure of the *V. vinifera* GT predicted sites of amino acids that could influence substrate binding. The 382 position was of particular interest with arginine in *C. paradisi* and tryptophan in *V. vinifera*. This research was designed to test the hypothesis that a R382W mutation would result in altered substrate specificity. Site-directed mutagenesis was performed to form the R382W mutant Cp3OGT and the gene was transformed into yeast for protein expression. Western blot determined the optimal protein induction period for the cells, after which the cells were broken to extract the recombinant mutant protein. Purification of the R382W 3GT allowed for enzyme analysis to be performed by measuring the incorporation of radioactive glucose into the reaction product. Docking analysis was performed using AutoDock software with both the wild type and R382W mutant protein with the substrates that showed interesting activity. The results of this study indicate that the point mutation of arginine to tryptophan at position 382 broadened substrate and regiospecificity to include flavanones.
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CHAPTER 1

INTRODUCTION

I. Flavonoids

Flavonoids are one of the three major classes of secondary metabolites, the others being terpenoids and alkaloids. Flavonoids are derived from phenylalanine which is produced via the Shikamate pathway (Figure 1). Phenylalanine is then modified through the phenylpropanoid pathway to produce the characteristic 15 carbon structure of the flavonoids. This structure is arranged in a C6-C3-C6 configuration, with two benzene rings connected by a three carbon chain. The basic structure is further modified to yield the subclasses of flavonoids, consisting of chalcones, aurones, flavanones, flavones, isoflavones, dihydroflavonols, flavonols, leucoanthocyanidins, and anthocyanidins (Figure 2). The oxidation and substitution of the heterocycle of the basic flavonoid structure serves to subdivide the flavonoids into classes and each class has a different function within the plant, such as UV protection, antifungal, antimicrobial, and anti-inflammatory properties (Harborne and Williams 2000), among many others.
Figure 1: Biosynthesis of flavonoids directly following the formation of Shikimate through the Shikimate pathway (Adapted from Crozier et al 2000).
II. Glycosylation of Flavonoids

Glycosylation is a common modification of flavonoids, resulting in over 6000 flavonoid compounds found in nature (Ferrer et al 2008). Glycosylation refers to the addition of any sugar molecule, while glucosylation refers to the addition of a glucose molecule. The production and accumulation of flavonoid di-glycosides are characteristic of citrus fruits and vegetative tissues (Owens and McIntosh 2009, and references therein). Glucosylation serves to stabilize the structure of the flavonoid, affect solubility which has implications in transportation, and regulates bioavailability (Vogt and Jones 2000). The type, extent, and site of glucosylation can vary widely between glucosyltransferases from different species and tissues within the same plant (Schwinn 1997). In *Vitis vinifera* alone, 200 putative flavonoid glucosides have been identified (Bonisch 2014).

The enzymes responsible for glucosylation reactions are glucosyltransferases (GTs) that transfer a glucose molecule from a UDP-glucose molecule to a sugar acceptor. Some UGTs are highly substrate and regiospecific (McIntosh and Mansell 1990, Owens and McIntosh 2009) while others can accept a broad
range of substrates (Osmani et al 2009); the variation in activity could contribute to the large amount of structural variation observed in GTs (Schwab 2003). All putative glucosyltransferases contain a conserved Plant Secondary Product Glucosytransferase (PSPG) box which is a UDP-glucose binding region (Owens and McIntosh 2016 and references therein). However, biochemical characterization has only been done to determine function of some of the putative enzymes in databases that have been assigned the identity of glucosyltransferase due to the presence of the PSPG box.

![Figure 3: The reaction catalyzed by Cp3GT (Owens and McIntosh 2009).](image)

III. Flavonoid effects on human and plant tissues

Some flavonoids appear to directly inhibit HIV activity, while others inhibit the enzymes required for viral replication (Harborne 2000 and references therein). Flavonoids can act as scavengers of oxidizing species, particularly 3’,4’-dihydroxyflavonoids, and have been found in vitro to inhibit oxidation of low-density lipoprotein (Harborne 2000). In mammals, some flavonoids can act as antispasmodylytic agents by relaxing smooth muscle (Middleton et al 2000). Isoflavones have been shown to possess oestrogenic activity (Harborne 2000). Flavonoids such as gossypin (Viswanathan et al 1984), epicatechin (Viswanathan et al 1984), morin and rutin (Thirugnanasamambantham et al 1985) show analgesic activity. Flavonoids have also been shown to have anti-proliferative, anti-tumor, and pro-apoptotic properties (Taylor and Grotewold 2005 and references therein).
Flavonoids play a direct role in the protection of plants from insect and mammalian herbivory (Harborne and Williams 2000, Ferreyra et al 2012). They function as photoalexins and have allelopathic properties (Taylor and Grotewold 2005). The majority of constitutive antifungal agents in plants are isoflavonoids, flavans, or flavanones (Harborne and Williams 2000). Flavonoids provide antimicrobial protection and many protect from harmful UV-B radiation (Harborne and Williams 2000, Ferreyra et al 2012).

IV. Grapefruit model system

The plant model for this research is Citrus paradisi, or the grapefruit. Citrus paradisi is known to accumulate a wide variety of flavonoids including flavonol, flavone, flavanone, dihydroflavonol, and chalcone glycosides (Owens and McIntosh 2011 and references therein, Daniel et al 2011 and references therein, Berhow 1998). The genus Citrus itself is known to accumulate flavanone glycosides (Jourdan et al 1985 and references therein). Grapefruit produce derivatives of several early flavonoid classes which are not found (flavonones and flavones) in high quantities in many other plants, as well as all the later flavonoid classes except anthocyanins. Varying amounts of flavonoids are found throughout all stages and tissues of the grapefruit (McIntosh and Mansell 1990 and references therein). Naringin is the primary flavonoid di-glycoside in grapefruit plants, making up approximately 40-70% of the dry weight of young leaves and fruit (McIntosh and Mansell 1990) and is rapidly synthesized during cell division (Jourdan et al 1985). Naringin accumulation is responsible for the bitter taste of grapefruit (McIntosh and Mansell 1990, Jourdan et al 1985).

V. Flavonoid GT Structure and Function

Glucosyltransferases are categorized based on a variety of properties including evolutionary relationships, sugar donors, and substrate preference. One current classification system is the CAZy
database which currently separates putative GTs from a variety of organisms into 97 families (Lombard et al. 2014) and each family has at least one biochemically characterized member. However, the sequence data in the database are not sufficient to determine the function of noncharacterized GTs. The sequence homology between GTs is very low, even between those that share substrate, regio- and/or stereospecificity (McIntosh and Owens 2016 and references therein). As such, biochemical analysis is still the most accurate method to determine function.

The tertiary structure of glucosyltransferases generally follows a β-α-β Rossmann-like domain pattern and fall into two major categories, GT-A and GT-B (McIntosh and Owens 2016 and references therein). Cp3GT is a GT-B enzyme, wherein the two Rossmann-like domains face each other and are connected by a linker region. The N-terminus of the enzyme is correlated to substrate specificity and the C-terminus, which is more conserved, is associated with sugar binding (Wang 2009 and references therein). The active site of the enzyme is located between the two domains (McIntosh and Owens 2016).

VI. Current study

Through this research, a potential change in substrate specificity of Cp3GT was analyzed using a site-directed mutation. This enzyme facilitates the transfer of a glucose molecule from UDP-glucose to the 3-OH position of some of the flavonol class of flavonoids. The mutation chosen is the change of arginine to tryptophan at the 382nd position, hereafter referred to as R382W. Arginine is a charged molecule, while tryptophan is bulky and aromatic. This mutation was chosen through sequence alignment and homology modeling with the flavonoid 3-GT found in V. vinifera. The 3-O-glucosyltransferase in V. vinifera can accept both anthocyanidins and flavonols as substrates; however, the 3-O-GT in C. paradisi will only accept flavonols as substrate (McIntosh and Owens 2016). Thus Cp3GT is considered highly substrate specific. This is hypothesized to be a result of the shape and orientation of
the binding pocket. The hypothesis of this research is that site-directed mutagenesis of R382W may alter the substrate and/or regiospecificity of 3-O-glucosyltransferase in *C. paradisi*. 
CHAPTER 2

MATERIALS AND METHODS

Materials

All reagents used for this study were analytical grade. Quercetin, kaempferol, fisetin, gossypetin, naringenin, hesperitin, eriodictyol, isosakuranetin, apigenin, luteolin, diosmetin, scutallarein, dihydroquercetin, 4′-acetoxy-7-hydroxy-6-methoxyisoflavone, and cyanidin were purchased from Indofine Chemical Inc. (Hillsborough, NJ). Uridine-5′-diphospho-[U-14C]- glucose (UDP-G; 50 μCi/2.5 mL) was purchased from Perkin Elmer. UDP was obtained from Sigma (St. Louis, MO). The pPICZA vector and X-33 strain of Pichia pastoris were from Sonje Roje of Washington State University (Pullman, WA) and one shot Top 10 competent E. coli cells were purchased from Invitrogen (Carlsbad, CA).

Zeocin™ was purchased from Research Products International Corps (Illinois). Luria-Bertani broth powder, agar powder, Tris-base, β-mercaptoethanol (BME), tetramethylethylenediamine (TEMED, electrophoresis grade), ammonium persulfate (APS), Coomassie brilliant blue, nitroblue tetrazolium chloride (NBT), nitrocellulose membrane (0.4 μm pore size), ethyleneglycol monomethylether, Whatman chromatography paper, 3,5-bromo-4-chloro-3′-indolylphosphate p-toluidine salt (BCIP), sodium phosphate (NH₄PO₄·H₂O), phenol, chloroform, isoamyl alcohol, 100% ethanol, acrylamide 40% solution (acrylamide:bisacrylamide, 19:1), peptone powder, glacial acetic acid, yeast extract, and yeast nitrogen base powder were purchased from Fisher Scientific (Pittsburg, PA).

Mouse anti C-myc monoclonal antibody was purchased from Sigma Aldrich (St. Louis, MO) and goat anti-mouse IgG alkaline phosphatase (AP) conjugate was purchased from Novagen (Madison, WI). Phenylmethylsulfonyl fluoride (PMSF) was purchased from MP Biomedicals (Solon, Ohio). GoTaq® DNA polymerase and 5X green/colorless buffer for polymerase chain reaction (PCR), deoxyribonucleotide
triphosphate bases (dNTP), and restriction enzymes Sac1 and Dpn1 were purchased from Promega (Madison, WI). TALON Metal Affinity Resin was purchased from Clontech (Mountain View, CA). Amicon Centrifugal filters were purchased from Millipore (Billerica, MA). All primers were synthesized by Integrated DNA Technologies.

**Site-directed mutagenesis**

The mutation R382W was designed and site-directed mutagenesis performed by SK Devaiah. The primers were as follows:

Forward 5′-tgccgatgatcgggtggccattctttggg-3’

Reverse 5′-ccaaagaatgccccagcagcatacggca-3’

The primers were designed using Stratagene QuikChange® Site-Directed Mutagenesis program (Agilent Technologies). PCR was performed using the forward and reverse mutagenic primers purchased from Integrate DNA Technologies and the QuikChange® Lightning Site-Directed Mutagenesis Kit protocol (Agilent Technologies). The parent DNA strands of the PCR productes were subsequently digested with DPN1 restriction enzyme (Agilent Technologies) and the mutant plasmid transformed into Top 10 competent *E. coli* cells (Invitrogen) via heat shock. The plasmid DNA from transformed cells was extracted using Quantum Prep™ Plasmid Midiprep Kit (Bio-rad), linearized using Sac1 restriction enzyme (Promega) and transformed into competent *Pichia pastoris* (Mut+) cells for protein expression as described in Invitrogen User Manual catalog number K1740-01.

**Homology Modeling**

EasyModeller 4.0 was used to model the protein structure of both the WT and R382W proteins. The sequence to be modeled was copied into the text box and the spaces removed. The .pdb file for the *Vitis vinefera* protein to be modeled against (2C1Z_NCBI) was selected and the box was checked in the
“Serial Number” column. The “Align Templates” tab was opened and the “Align Templates” button was selected. Next, the “Align Query” tab was opened and “Align Query with Templates” button was selected. The “Build Model” tab was opened and “Generate Model” button was selected. Nine models to be generated was selected and NO for heteroatoms and automatically refine loops. Once the options were chosen, OK was selected and Modeller began generating models. In the command prompt window, the GA341 and DOPE (discrete optimized protein energy) scores were evaluated. A model with a GA341 score lower than 0.6 is not a viable model and a lower DOPE score indicates a better model. The model with the lowest DOPE score was copied to the desktop for ease of location and this was the model tested against. Swiss PDB Viewer was opened and the generated model was loaded. All residues were selected (under Select>All) and an energy minimization run (under Tools>Energy Minimization). The nine models were saved as PDB files.

**Docking Simulations**

PyRX was opened and the Vina Wizard tab in the bottom left of the screen selected, after which “Start” was clicked. The generated model was loaded by clicking the “Add Macromolecule” button. The folder of the model was clicked and the ligands were selected here for docking. Any ligands needed that were not already downloaded into PyRx were downloaded from the European Protein Data Bank. The plus button next to the protein name was clicked to display a list of amino acids in the model. The amino acids which are predicted to be important in the binding site were selected and the labels displayed by selecting Display, then Label, then Atoms. For UDP-Glucose, positions 20, 21, 290, 292, 293, 319, 345, 346, 348, 363, 365, 366, 367, 368, and 371 were displayed. For aglycones, positions 22, 87, 124, 150, 154, 191, 203, and 207 were displayed. The selection box in the protein view window was positioned so that all labeled amino acids were within the box. The “Exhaustiveness” number was set to 50 to increase accuracy and the “Run Vina” button was selected. Nothing else was running at the same
time as Vina uses the maximum available processing power in order to produce the model. The resulting file gave the nine most likely conformations for that ligand's docking configuration as well as the binding affinity in kcal/mol.

**Interpreting Docking Results**

AutoDock Tools was opened and the docking results were opened. The results were displayed as one molecule with multiple conformations. The model used for the docking simulation was opened (Analyze>Macromolecule>Open Macromolecule). The Show Interactions command was selected (Analyze>Docking>Show Interactions) to view only the amino acids interacting with the ligand. Msms was turned off as well as the spheres showing close contacts. The distances were measured between catalytic residues (His22, Asp122, and Ser20) and their corresponding interactions on the ligand.

**Expression of Mutant Recombinant Protein in *Pichia pastoris***

YPD media consisting of 1% yeast extract, 2% peptone, 2% dextrose, and 100 mg/L zeocin was prepared and yeast colonies of WT and R382W were inoculated to 3 mL each aliquots of the media. The cultures were incubated overnight at 28°C shaking. The OD at 600 nm was checked, with an optimum between 2 and 6. From the overnight cultures, 1 mL of each sample was inoculated into separate 100 mL BMGY media (1 M potassium phosphate buffer at pH 6.0, 10X YNB, 500X biotin, and 10X glycerol) in autoclaved 500 mL Erlenmeyer flasks. These cultures were incubated overnight, then centrifuged at 3000xg and washed once with water and once with BMMY (1 M potassium phosphate buffer at pH 6.0, 10X YNB, 500X biotin, and 10X methanol), and the pellets were re-suspended in three separate 100 mL BMMY media in 500 mL Baffled flasks. The cells were harvested 24 hours after induction with methanol for the wild type and 18 hours after induction with methanol for the R382W mutant and split into two 50 mL tubes per sample, then centrifuged at 3000xg at 4°C for five minutes. The supernatant was removed and the pellet stored at -80°C.
GT purification

The pellet stored at -80°C was resuspended in 5 mL of breaking buffer (50 mM sodium phosphate buffer at pH 7.5, 1 mM PMSF, 1 mM EDTA, 5% glycerol, 5 mM BME) and lysed using the French Press (Fisher Scientific) four times at 1250psi or until the sample appeared semitransparent, with samples collected on ice. The samples were centrifuged at 13,000xg and 4°C for 20 minutes. The supernatant was kept on ice. A PD-10 column and a cobalt IMAC column were equilibrated using equilibration buffer (50 mM sodium phosphate buffer at pH 7.5, 300 mM NaCl, 5 mM BME) at 4°C. Protein extract was added to the PD-10 column and the flow through discarded. Equilibration buffer was added to the column and the eluted protein solution collected. The desalted protein solution was added to the IMAC column and the flow through discarded. The column was washed with equilibration buffer to an OD < 0.09 at 280 nm. The bound protein was eluted using elution buffer (50 mM sodium phosphate buffer at pH 7.5, 300 mM NaCl, 5 mM BME, 150 mM imidazole) and collected in 2 mL fractions. Protein concentrations were measured with the nano drop at 280 nm. The fractions which showed the highest OD were concentrated using Amicon 30 MWCO filter, centrifuging at 3000xg and 4°C for 10-12 minutes. Two mL of final assay buffer (50 mM sodium phosphate buffer at pH 7.5, 14 mM BME) was added to the filter and centrifuged until the volume reached 500 µL. The protein solution was placed into a 2 mL tube and 500 µL of final assay buffer used to wash the Amicon filter then added to the sample and the concentration measured using the nano drop at 280 nm. The protein solution was held on ice and used immediately for assay, but could be stored in buffer containing 10% glycerol at -80°C for 24 hours if absolutely necessary.

Enzyme Assay

For the screening assay, the reaction mixture was made as follows: 50 nmol of aglycones in 5 µL ethylene glycol monoethyl ether, 50 µL of assay buffer (50 mM sodium phosphate buffer at pH 7.5, 14
mM BME), 100 nmol/10 µL UDP-\(^{14}\text{C}\) glucose (20,000 cpm) in 10 µL of 50 mM phosphate buffer (pH 7.5), and 2-5 µg of purified protein. Each reaction was done in duplicate and incubated five minutes at 37°C. The reaction was stopped by adding 15 µL of 6 M HCl and the mixture was vortexed. Next, 250 µL of ethyl acetate was added to each tube, vortexed, and centrifuged for 10 seconds. Approximately 150 µL of the upper phase was removed and added to 2 mL of scintillation liquid in a 7 mL scintillation vial for each sample. The incorporation was measured using a Beckman LS 6500 scintillation counter.

**SDS-PAGE Analysis**

Two 10% SDS-PAGE gels were prepared using the following recipe.

**Table 1: Recipe for 10% separating gel for SDS-Page.**

<table>
<thead>
<tr>
<th>10 % Separating Gel</th>
<th>1 gel</th>
<th>2 gel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deionized water</td>
<td>2.3 mL</td>
<td>4.8 mL</td>
</tr>
<tr>
<td>40% acrylamide</td>
<td>1.3 mL</td>
<td>2.5 mL</td>
</tr>
<tr>
<td>1.5 M Tris (pH 8.8)</td>
<td>1.3 mL</td>
<td>2.5 mL</td>
</tr>
<tr>
<td>10% SDS</td>
<td>0.05 mL</td>
<td>0.1 mL</td>
</tr>
<tr>
<td>10% Ammonium persulfate</td>
<td>0.05 mL</td>
<td>0.1 mL</td>
</tr>
<tr>
<td>Tetramethylethylenediamine</td>
<td>0.002 mL</td>
<td>0.004 mL</td>
</tr>
</tbody>
</table>
Table 2: Recipe for stacking gel for SDS-Page.

<table>
<thead>
<tr>
<th>Stacking gel</th>
<th>1 gel</th>
<th>2 gel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deionized water</td>
<td>0.73 mL</td>
<td>4.5 mL</td>
</tr>
<tr>
<td>40% acrylamide</td>
<td>0.13 mL</td>
<td>0.78 mL</td>
</tr>
<tr>
<td>1.5 M Tris (pH 6.8)</td>
<td>0.13 mL</td>
<td>0.75 mL</td>
</tr>
<tr>
<td>10% SDS</td>
<td>0.01 mL</td>
<td>0.06 mL</td>
</tr>
<tr>
<td>10% Ammonium persulfate</td>
<td>0.01 mL</td>
<td>0.06 mL</td>
</tr>
<tr>
<td>Tetramethylethylenediamine</td>
<td>0.001 mL</td>
<td>0.006 mL</td>
</tr>
</tbody>
</table>

The separating gel was allowed to solidify for approximately an hour. A 1:1 ratio of butanol-water mix was overlaid to make a straight separating gel after pouring. Once solidified, the butanol-water mix was poured off and the stacking gel was added and allowed to solidify. Samples to be loaded onto the gel were prepared in 1.5 mL autoclaved labelled Eppendorf tubes using 5 µL of 4X SDS-PAGE dye with BME and 10 µL of samples made to the same protein concentration for a total of 15 µL final volume. Duplicate samples were prepared, one for Western blot analysis and the other to be stained in Coomassie blue (0.006 g Coomassie Brilliant Blue, 10 mL acetic acid, 40 mL methanol, made up to 100 mL with deionized water). Reaction samples were boiled for 5 minutes to denature the protein. The SDS-PAGE apparatus was set up with the gel between two glass plates and placed in the tank. 1X Tris-glycine electrophoresis buffer was added to the tank and the samples were loaded, beginning with Low range prestained (Bio-rad) protein marker followed by a positive control and 15 µL of the sample reaction mixtures. Electrophoresis was performed at 100 V for 90 minutes or until the dye front reached the end of the gel. Subsequently, one gel was placed into Coomassie blue stain and allowed to shake on an ORBITRON shaker (BOEKEL), and the other was transferred to the Western blot apparatus. After 24
hours, the stained gel was placed into destaining solution (300 mL methanol, 100 mL acetic acid, made up to 1 L with deionized water).

**Western Blot Analysis**

Two porous pads (sponges) were soaked in cold 1X transfer buffer in a glass dish and a nitrocellulose membrane (Thermo Scientific) was cut to the size of the SDS-gels to transfer the protein from the gel to the nitrocellulose membrane. The electro blotting kit was set up by placing the negative terminal of the transfer cassette on the bench with a porous pad on top of it, followed by chromatography paper, the gel and nitrocellulose membrane, followed again by chromatography paper and a porous pad. The positive terminal of the transfer cartridge then closed the set and the transfer cassette was placed in cold 1X transfer buffer in the tank. An ice pack and stir bar were added to the tank. The SDS-gel was electroblotted for 75 minutes at 100 V.

After electroblotting, the membrane was carefully removed and placed in blocking solution (5% non-fat milk powder in 1X TBS and 0.02% sodium azide) using laboratory tongs, and allowed to shake gently for 2 hours or overnight at room temperature. It was then rinsed with 30 mL of 1X TBS containing 30 µL of 20% sodium azide three times for five minutes each. The membrane was incubated in a 1:2500 dilution with 1X TBS of monoclonal anti C-myc primary antibody solution from mouse for two hours. The primary antibody binds to the C-myc epitope (EQKLISEEDL) in the recombinant protein which was designed in the expressed protein for identification and purification of the enzyme. The membrane was next washed with 30 mL of 1X TBS containing no sodium azide three times for five minutes each and transferred to a fresh tray where it was incubated in a 1:10,000 dilution with 1X TBS of secondary antibody solution (Goat anti-mouse immunoglobin G-alkaline phosphatase conjugate) (Novagen) for two hours at room temperature, gently shaking.
The membrane was washed 3 times with 1X TBS solution as earlier and then transferred into 30 mL of CHES buffer solution (pH=9.5), 60 µL of BCIP, and 60 µL of NBT. The membrane was placed on the shaker until protein bands appeared, approximately 3-5 minutes. Once developed, the membrane was washed with deionized water to stop the developing reaction. Both the destained gel and the developed western blot were photographed for future reference (Appendix A).
CHAPTER 3

RESULTS AND DISCUSSION

Homology modeling was done for both the wild type Cp3GT and R382W mutant proteins based on the crystallized structure of the *Vitis vinifera* 3-O-GT (Offen et al 2006) and Clitoria ternata GT (Hiromoto et al 2013 and 2015). *Citrus paradisi* Cp3GT is flavonol specific, *Vitis vinifera* 3-O-GT is flavonol and anthocyanidin specific, and *Clitoria ternatea* 3-O-GT is anthocyanidin specific. There is a 56% sequence identity and an 87% homology between Cp3GT and Vv3GT. There is a 43% identity and a 74.4% homology between Cp3GT and Ct3GT. However, the substrate specificity of the three enzymes differs. Homology modeling was performed using the crystallized structures of *Vitis vinifera* 3-O-GT and *Clitoria ternatea* 3-O-GT to create a proposed model for Cp3GT that reflects its specificity towards flavonols. The mutant protein was modeled from the proposed Cp3GT model. The overall shape of the WT and mutant proteins is similar (Figure 4); however, when overlaid (Figure 5) clear structural differences emerge. The single point mutation of arginine to tryptophan at position 382, located in the C-terminal domain, causes changes in folding of the loop regions circled (Figure 5). The models predict that a single point mutation at this location could be significant enough to change the native conformation. These changes could alter substrate specificity due to differential ability of aglycones to fit in the binding pocket.
Figure 4: Quaternary structure of Cp3GT (left) and R382W (right).

Figure 5: Initial model of Cp3OGT overlaid with R382W mutant protein. The WT protein is shown in blue, the mutant in brown. Quercetin and UDP-Glucose are docked. The WT arginine at position 382 is marked in green and the mutant tryptophan is marked in red.
Next, multiple sequence alignment was performed with *Citrus paradisi*, *Vitis vinifera*, and *Clitoria ternatea* 3GTs (Figure 6). The mutation chosen is located within the conserved PSPG box; all three enzymes contain an arginine at the 382nd position. However, homology modeling showed that a point mutation in that position could cause substantial changes in folding which could result in altered substrate and regiospecificity.

![Sequence alignment](image)

Figure 6: Sequence alignment of wild type Cp3GT and mutant R382W using Bioedit software. The PSPG box is outlined in black. * indicate identical amino acids for all three enzymes.
Table 3: Initial screening of Cp3GT and R382W performed by Dr. Shivakumar Devaiah. Verification performed by KAK. Initial screening reported in % relative activity to Cp3GT with quercetin; verification reported in % relative activity to R382W with quercetin.

<table>
<thead>
<tr>
<th></th>
<th>Initial Screening</th>
<th>Verification</th>
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<tr>
<td></td>
<td>Cp3GT</td>
<td>R382W</td>
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<tr>
<td><strong>Flavonols</strong></td>
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<td>Kaempferol</td>
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<tr>
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<td>6</td>
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<td>------------------------</td>
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</tr>
<tr>
<td>Eriodictyol</td>
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<td>1</td>
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<tr>
<td>Isosakuranetin</td>
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It has been previously established that Cp3GT is specific for the flavonols quercetin, kaempferol, fisetin, and gossypetin (McIntosh and Owens 2016), which was also observed in the initial activity screening (Table 3). The initial screening assay is highly sensitive and can therefore produce false positives. However, this assay will not produce false negatives. These results are reported in percent relative activity of WT with quercetin as Cp3GT favors this flavonol over the others. The R382W mutant still shows a preference for the flavonols, with almost identical activity with the flavonols as the WT. However, the mutant protein also shows an increase in activity with naringenin and hesperitin which are both flavanones. This assay was repeated to confirm verify prior screening results of R382W (Table 3) and was reported in percent relative activity of R382W with quercetin. The verification assay shows the same pattern as the initial screening. This indicates that substrate specificity may have been broadened in the mutant protein. However, the assay also suggests an increase in activity with luteolin, dihydroquercetin, 4'-acetoxy-7-hydroxy-6-methoxy-isoflavone, and cyanidin. Dihydroquercetin and 4'-acetoxy-7-hydroxy-6-methoxy-isoflavone activities have been shown previously to be false positives. However, luteolin and cyanidin activity could be real and should be tested with a kinetic assay.
Docking analysis was used for in silico analysis to predict whether reactions could occur with the substrates that looked promising. Distances were measured from the three catalytic residues Histidine 22, Aspartic acid 122, and Serine 20 to their respective positions of interaction on the aglycone substrates (Offen et al 2006). The nitrogen on histidine 22 serves as a Bronstead base, removing electrons from a OH group to encourage addition of the glucose molecule at that position (Offen et al 2006). In the case of Cp3GT, this interaction is with the 3-OH group. Aspartic acid 122 stabilizes the charge on His22, interacting with the side chain nitrogen (Offen et al 2006). Serine 20 is responsible for stabilizing the aglycone molecule in the binding pocket by interacting with the 4-keto functional group (Offen et al 2006). Distances smaller than 5Å have been proposed to be favorable for reactions to occur (Offen et al 2006). However, preference will be given to activity screening results and docking simulations will be used only to confirm and visualize the results.

The first docking simulation performed was Cp3GT and R382W with quercetin (Figure 7). The activity screening showed identical activity between the WT and mutant proteins (Table 3) so the distances of the three catalytic residues to the substrate are expected to be similar as it is hypothesized that these distances impact enzymatic activity. This is indeed the case for the predicted model. The distance between His22 and the 3-OH position is 3.443Å for the WT and 3.017Å for the mutant protein. Both of these distances are well under 5Å and thus favorable for the reaction to occur. The distance between Ser20 and the 4-keto group is 3.443Å in WT and 3.017 in the mutant protein. These distances are both favorable for a reaction to occur. The distances between Asp122 and the side chain nitrogen of His22 in both the WT and mutant proteins are also favorable, at 4.591Å for WT and 3.017Å for R382W. This is validated by the activity screening which shows a preference of both proteins for quercetin. However, the WT model chosen was not the most energetically favorable model given by the software and was the only predicted model with distances under 5Å. Several of the predicted models for the
mutant protein were favorable for a reaction; thus, crystallization is needed to confirm the real protein structure. This is because there is not a crystal structure for a flavonol specific GT.

Figure 7: Models of A) Cp3GT and B) R382W docked with Quercetin.

The next docking simulation done was WT and R382W proteins with kaempferol. Both proteins had reduced activity as compared to quercetin but still favor the reaction (Table 3). This is seen as well in the predicted models. The distance from His22 to the 3-OH position is 3.235Å for the WT and 4.188Å for the mutant protein (Figure 8). Both of these are favorable distances for the reaction. The distances of Ser20 to the 4-keto position in both WT and R382W are favorable for the reaction, 3.235Å and 4.362Å respectively, but higher than those distances for quercetin (Figure 7) which could explain the reduction in activity. The distances between Asp122 and the side chain nitrogen of His22 are both favorable for a reaction to occur at 4.591Å for WT and 2.737Å for R382W. Again, the models chosen were not the most energetically favorable models given by the EasyModeller software. However, these models are validated by the activity screening. Several of the other predicted models also show favorable distances.
Therefore, both proteins would have to be crystallized to confirm the real protein structure and the interaction with kaempferol.

![Figure 8: Models of A) Cp3GT and B) R382W docked with Kaempferol.](image)

The activity screening showed activity with fisetin (Table 3). The distance between His22 and the 3-OH group of fisetin are 5.512Å for the WT and 5.404Å for the mutant protein (Figure 9). These distances are both over 5Å but because globular proteins are not static a reaction may still occur if at some times His22 is positioned closer to the substrate. The distances between Ser20 and the 4-keto group are well above the favorable 5Å. However, it is possible that binding of the substrate in the binding pocket could change the conformation enough to bring Ser20 within range for the reaction to occur. This could be responsible for the reduction of activity from quercetin. The distance between Asp122 and the side chain nitrogen of His22 in both Cp3GT and R382W are favorable as well. Again, both models were not the most energetically favorable models given by the modeling software. The most energetically favorable models gave distances too large for a reaction to occur; the activity screening shows activity so models were chosen to reflect the screening. However, this activity may be a
false positive which could be confirmed with a kinetic assay. Crystallization with and without fisetin for both Cp3GT and R382W would also confirm if the conformation of the binding pocket changes after the substrate binds to bring Ser20 into range for a reaction to occur.

Figure 9: Models of A) Cp3GT and B) R382W docked with Fisetin.

The last of the flavonols docked with Cp3GT and R382W was gossypetin. The activity screening showed a slight decrease in activity from fisetin (Table 3). The distance between His22 and the 3-OH group was 5.068Å for WT and 5.525Å for the mutant protein (Figure 10). These distances are both slightly above the favorable distance but because globular proteins are not static a reaction could occur if there are moments in which His22 is closer to the substrate. The distance between Ser20 and the 4-keto group was 5.984Å for WT and 3.715Å for R382W. There is an interesting balance that emerges between His22 and Ser20 where one distance is smaller in WT and larger in R382W and vice versa. This is likely to be the reason for a reduction in activity with gossypetin. The distances between Asp122 and the side chain nitrogen of His22 are both favorable for the reaction. The chosen models were again not the most energetically favorable conformations given by the modeling software. This is the first of the
substrates to have a completely different orientation within the binding pocket between WT and R382W. Crystallization of the two proteins with gossypetin would confirm these configurations.

![Models of A) Cp3GT and B) R382W docked with Gossypetin.](image)

Figure 10: Models of A) Cp3GT and B) R382W docked with Gossypetin.

Naringenin was docked with Cp3GT and R382W proteins because the activity screening showed a sharp increase in activity in the mutant protein (Table 3). This increase is readily apparent in the predicted models. Naringenin is a flavanone, which does not have a 3-OH group; however, HPLC analysis done previously in this lab with other mutants shows that the glucose molecule is added to the 7-OH group. It has not yet been confirmed that this is the position on R382W that receives the glucose molecule; however, for these models, it was assumed that the 7-OH position is the site of addition. The distance between His22 and the 7-OH group in the WT protein is 7.235 Å which is much further than is favorable for a reaction to occur. However, the distance in the mutant protein is 4.602 Å which is well within the 5 Å guideline (Figure 11). These models suggest that a reaction is possible for the mutant protein and a broadening in substrate specificity to include flavanones. This would also suggest a broadening of regiospecificity as the glucose molecule is added to the 7-OH group instead of a 3-OH group. The distance between Ser20 and the 4-keto group is similar in both the WT and mutant proteins,
indicating that His22 is the most important residue for a reaction to occur. The Ser20 distance is too far for a reaction to occur but it is possible that substrate binding changes the conformation to become favorable. However, this is not the determining factor as is seen in the WT. The WT model is the second most energetically favorable model; however, the R382W is the least energetically favorable model given by the EasyModeller software. This illustrates the limitations of the modeling software. Therefore, it is imperative to do experiments and not to rely solely on in silico analysis.

Figure 11: Models of A) Cp3GT and B) R382W docked with Naringenin.

The next substrate to show altered activity was luteolin which is a flavone. The activity screening showed a decrease in activity with the mutant protein compared to the WT in the initial screening but a huge increase in the verification (Table 3). This can be resolved using HPLC or by doing a kinetic assay. Docking analysis puts the distance between His22 and the 7-OH group at 4.078Å for the WT which is within the distance to favor a reaction (Figure 12). However, Cp3GT is not able to do a reaction using luteolin. Therefore, this model is likely wrong but still shows that a reaction could theoretically happen. The same distance for the mutant protein is 7.940Å which is too far away for a reaction to be possible.
The distance between Ser20 and the 4-keto group is 6.890Å in the WT which is not close enough to favor a reaction. Only a change in conformation after substrate binding would bring Ser20 into range for a reaction to occur. This distance in R382W is 7.652Å, which is too far to predict a change in conformation and the orientation of the substrate itself is not favorable. The distances between Asp122 and the side chain nitrogen of His22 are favorable for both proteins; however, because the other distances are not favorable in the mutant protein, the reaction probably does not occur.

Figure 12: Models of A) Cp3GT and B) R382W docked with Luteolin.

Cyanidin is an interesting substrate because there is not a 4-keto group for Ser20 to interact with. Therefore, distances were only measured between His22 and the 7-OH group, as well as Asp122 to the side chain nitrogen of His22. The distance between His22 and the 7-OH group in the WT is 5.825Å and 5.251Å in the mutant. Both of these distances are outside of the 5Å guideline which suggests that the reaction does not occur (Figure 13). It is possible that at certain times His22 is closer to the substrate than at other times but this is not likely. However, it has been shown that this reaction does not occur.
for WT in a kinetic assay (Owens and McIntosh 2009). This distance for R382W is closer than the WT distance, making the reaction slightly more favorable for the mutant but not enough to be considered expanded substrate specificity. The distance between Asp122 and the side chain nitrogen of His22 is 4.591 in WT and 2.737 in the mutant, both of which are favorable distances.

Figure 13: Models of A) Cp3GT and B) R382W docked with Cyanidin.
CHAPTER 4

CONCLUSIONS AND DIRECTIONS FOR FUTURE RESEARCH

Site directed mutagenesis was performed to create the mutation R382W which was an arginine changed to a tryptophan at position 382. This mutation was chosen based on homology modeling and multiple sequence alignment between Cp3GT and Vv3GT. Cp3GT is highly specific to some flavonols while Vv3GT is specific for anthocyanidins but can also glucosylate flavonols. The purpose of this research was to determine structure and function of key amino acid residues in Cp3GT. This mutation was hypothesized to alter the substrate and/or regiospecificity of Cp3GT to include anthocyanidins.

This mutation is within the conserved PSPG box so no variation was observed between the amino acids in Cp3GT and Vv3GT during sequence alignment; both enzymes contain an arginine residue at the 382nd position. However, modeling of the quaternary structure of both Cp3GT and R382W showed substantial structural differences in the mutant, primarily in the loop regions. Therefore, it was likely that this mutation would alter substrate specificity. This is indeed the case in the initial activity screening. R382W showed a sharp increase in activity with naringenin and hesperitin which are both flavanones. The mutant also retained activity with the flavonols and had a slight increase with anthocyanidin. However, this assay is highly sensitive and can produce false positives. As such, these results should be confirmed with a kinetic assay (Owens and McIntosh 2009).

Docking simulations were performed to visualize the predicted binding pocket conformation when bound with the substrates of interest. Distances between three catalytic residues (His22, Asp122, and Ser20) and the substrate were used to determine if reactions could be favorable. Any distance under 5Å was considered favorable for these docking models (Offen et al 2006). The docking simulations showed orientations that favor a reaction for only the flavonols with Cp3GT. However, with R382W there were also favorable models for naringenin. This indicates that the activity observed in the initial
screening with naringenin could be accurate and not a false positive. This could be should be confirmed using HPLC product identification and kinetic analysis. Crystallization of both proteins should be done to visualize the correct structure of the binding pocket as these models are only proposed and based off of Vv3GT.

In the future, the initial screening results should be confirmed. In addition, full biochemical characterization should be performed for the mutant protein. Kinetic assays should also be done to determine if a change occurred in catalytic efficiency due to the point mutation. HPLC analysis should be done to confirm the reaction products of these reactions. These results will be used to refine the in silico models to give a more accurate depiction of the binding site and substrate binding. Crystallization should also be done for both the WT and mutant proteins to elucidate the exact structure of the proteins. However, the results from this study indicate that the point mutation of R382W broadened substrate and regiospecificity of Cp3GT to include flavanones.
REFERENCES


APPENDIX A

Figure A1: Site-directed mutagenesis performed using QuikChange Lightning Site-Directed Mutagenesis Kit from Agilent Technologies.

Figure A2: Western Blot of Cp3GT and R382W to confirm size. The proteins with tags are approximately 65 kDa in size.
MAQTQSQPRP HIAVLNFPFS THASSVLSII KRLAVSAPTA LFTFFSTPQS NKALFSTGQQ RHLPSNVKPY
DVSDGVPEGH VFSGKRQEDI ELFMNAADAN FRKAVEAAVA ETGRPLTCLV TDAFIWFFAE MAREWNNVPW
VPCWPAGPNLSAHLTYTDII RDKIGTQSQN QDQQLIHFIP GMNKIRVADL PEGVVSGDLD SVFSVMLHQM
GRQLPKAAAV FINSFEELDP ELTNHLTKF NNKFLSVPNF KLLLASDQQP SSATDLDEY GCLAWLKDQK
KKPASVAYVS FGTVPATPSPN EIVAIAEAEL ANKVPIWSL RHRSQANLPN GFLERTRSDG IVVDWAPQVN
VLAHEAVGF VTHCGWGSIL ESIAAGVPMI GRPFFGDQRI NGRMMEQVWG VGVAVDGGGI CTKEGLSSL
DLILCQEKGI KIREKVTKLK QLQCNAIGPG GSSMQLDVL VDMISRSY

Figure A3: Cp3GT amino acid sequence.

MAQTQSQPRP HIAVLNFPFS THASSVLSII KRLAVSAPTA LFTFFSTPQS NKALFSTGQQ RHLPSNVKPY
DVSDGVPEGH VFSGKRQEDI ELFMNAADAN FRKAVEAAVA ETGRPLTCLV TDAFIWFFAE MAREWNNVPW
VPCWPAGPNLSAHLTYTDII RDKIGTQSQN QDQQLIHFIP GMNKIRVADL PEGVVSGDLD SVFSVMLHQM
GRQLPKAAAV FINSFEELDP ELTNHLTKF NNKFLSVPNF KLLLASDQQP SSATDLDEY GCLAWLKDQK
KKPASVAYVS FGTVPATPSPN EIVAIAEAEL ANKVPIWSL RHRSQANLPN GFLERTRSDG IVVDWAPQVN
VLAHEAVGF VTHCGWGSIL ESIAAGVPMI GRPFFGDQRI NGRMMEQVWG VGVAVDGGGI CTKEGLSSL
DLILCQEKGI KIREKVTKLK QLQCNAIGPG GSSMQLDVL VDMISRSY

Figure A4: R382W amino acid sequence.
Figure A5: Partial sequencing of Cp3GT and R382W to confirm that the mutation had been inserted correctly during site directed mutagenesis. The circled region is the location of the mutation from arginine to tryptophan.

Figure A6: Gel electrophoresis to confirm purification of the enzyme. Purification of WT and R382W.

From right to left: Crude WT(1), crude R382W (2), crude D344P(3), E7 R382W(4), E7 D344P(5), final WT(6), final R382W(7), final D344P(8).
Figure A7: Western blot time course of protein induction at 0hr (Lane 2), 6hr (Lane 3), 12hr (Lane 4), and 24hr (Lane 5) of the transformed *Pichia pastoris* to determine optimal growth time. Lane 1 is a positive control and Lane M is the low molecular weight marker.
APPENDIX B

VITA

KATHLEEN KING

Education:

East Tennessee State University

Class of 2016, University Honors Scholar

Bachelor of Science in Biology, Biochemistry concentration, degree earned August, 2016

Professional Memberships:

Beta Beta Beta, Biological Honors Society

Phytochemical Society of North America

Honors/Awards:

Summer 2015: Best Undergraduate Poster, Phytochemical Society of North America, Annual Conference, Urbana, Illinois, $250 award

Summer 2015: Travel grant from ETSU Honors College, $500 to attend PSNA annual conference

Fall 2014: Student faculty collaborative research grant, Effect of R382W on Substrate Specificity of Cp3GT, $1000 to purchase lab reagents and supplies

Presentations:


Fall 2015: Poster Presentation at Appalachian Student Research Forum, East Tennessee State University, Analysis of Impact of R382W Mutation on Substrate Specificity of Grapefruit Flavonol Specific 3-Glucosyltransferase. Co-authors: Shivakumar P. Devaiah, Cecilia A. McIntosh.