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PRENYLTRANSFERASES

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## **Abstract**

Chemoenzymatic synthesis is a leading technique in the diversification and late stage modification of complex molecules, due to the high regio and stereo specificities of the enzymes. Over the last decade, the broad aromatic substrate scope for a class of prenyltransferases known as aromatic prenyltransferases has been extensively studied. Aromatic prenyltransferase catalyzed prenylation reactions are highly prevalent in nature and contribute to the amazing structural and functional diversity of natural products. Given their natural function and broad aromatic substrate promiscuity, aromatic prenyltransferases have been investigated as biocatalytic tool essential to facilitate chemoenzymatic natural product diversification. Although many studies have accomplished highlighting the acceptor substrate scope of this class of enzyme none have explored the enzyme's potential to transfer unnatural alkyl groups. Herein, we investigate the alkyl donor specificity of four aromatic prenyltransferases, SirD, FgaPT2, NphB and CdpNPT, in combination with our extensive library of synthetic alkyl pyrophosphates. To further highlight the feasibility of aromatic prenyltransferases as a powerful biocatalyst, we successfully diversified FDA approved drug compounds, subsequently resulting in the improved *in vitro* activity of the modified compound.

## Chapter 1: Introduction to Aromatic Prenyltransferases

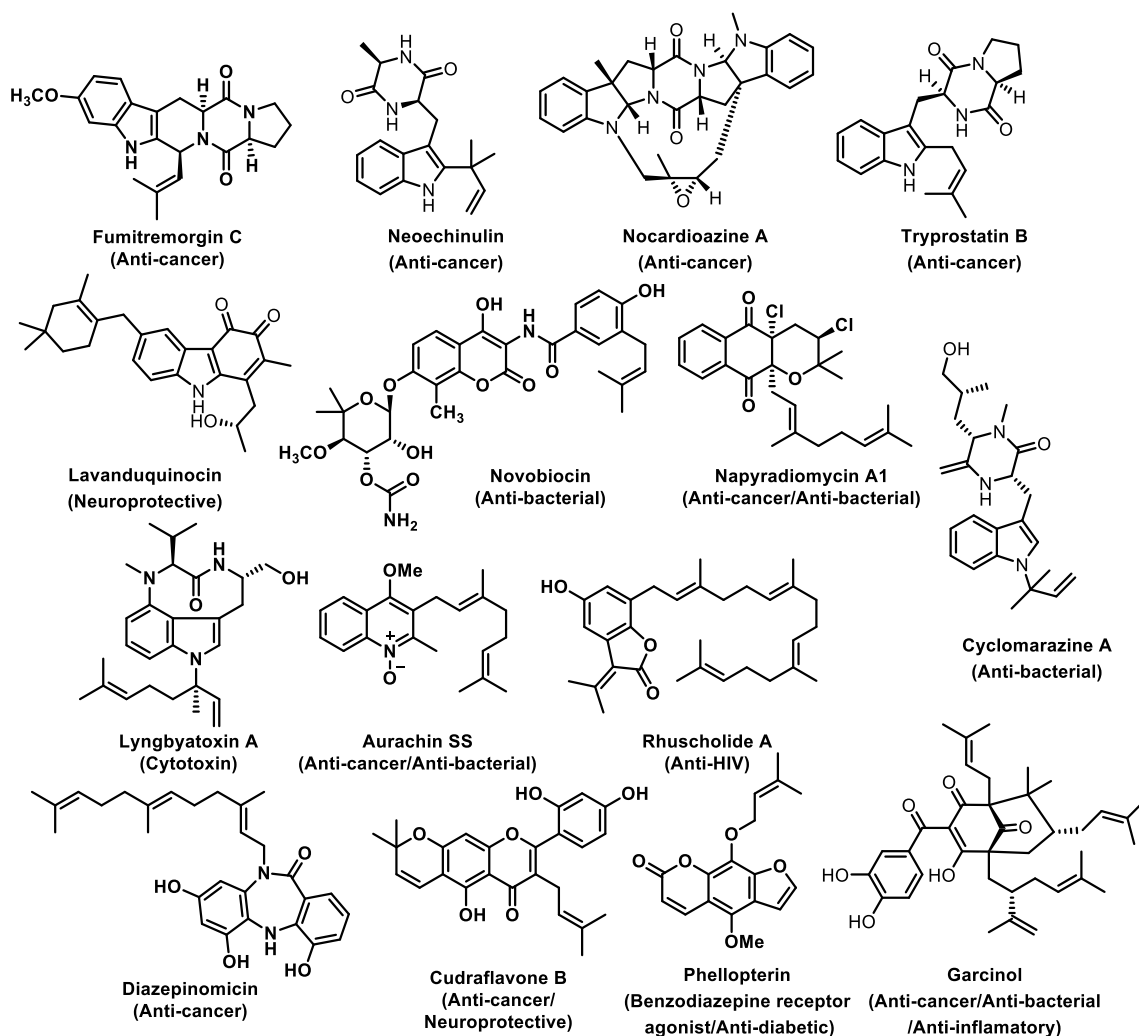
### Important Natural Products

Prenylation is ubiquitous across all biological systems and has been known to enhance lipophilicity of compounds giving the compounds a stronger affinity to biological membranes, as well as altering binding specificity and providing isoprenoid chains to be incorporated into the carbon scaffold. As a result, prenylated natural products display different pharmacological activities distinct from their non prenylated precursors attracting attention amongst the medical and scientific community (Figure 1)[1]. Indole diketopiperazines, derived from L-tryptophan and a second amino acid, typically and L-proline, L-tryptophan, L-histidine, L-phenylalanine or L-leucine, are a unique class of prenylated natural products with a wide spectrum of biological and pharmacological activities.[2] This class includes the tryprostatins A and B, nocardioazines, fumitremorgin C and Echinulin. Tryprostatin A and B, originally cultured from the fungus *Aspergillus fumigatus* and *A. tamari*, are found to inhibit cell cycle progression of various cancer cell lines.[3] It has been determined that tryprostatin A is active against tubulin polymerization, disrupting the interaction between microtubule associated proteins (MAP) and the C-terminal domain of tubulin. Although tryprostatin B also arrest cell cycle progression at lower concentrations than tryprostatin A it has a different mechanism of action as compared to tryprostatin A.[3] Echinulin and cyclomarazine A and B possess biological and pharmacological activities.. Specifically, echinulin and its analogues show antiviral, anticancer, and neuroprotective activity.[5][6] Cyclomarazine A and B, a reverse N-prenylated compound, has moderate antibacterial activity against

vancomycin-resistant *Enterococcus faecium* and methicillin-resistant *Staphylococcus aureus*. [7]

Aurachins, a unique family of natural products featuring a quinoline chromophore with a C-3 or C-4 farnesyl or geranyl substitution, display multiple biologically active properties. [8] Due to their structural similarities to ubiquinol and vitamin K, aurachins act as inhibitors of the electron transport chain, specifically by inhibiting various cytochrome complexes. They have been reported to possess strong antimicrobial, antifungal and antiplasmodial characteristics. Napyradiomycins, a group of natural products isolated from *Actinomycetes*, were previously identified as antibacterial antibiotics; however, they have recently been identified to exhibit cytotoxicity against human cancer cell lines. It has been speculated that the inhibitory effect against mitochondrial electron transport thus decreasing ATP production explains their mechanism of action against cancer cells. [9, 10] Diazepinomycin, an unusual metabolite found in bacterium *Micromonospora* sp. RV115, features a benzodiazepine derivative with a farnesyl side chain and underwent phase II clinical trials for the treatment of glioblastoma multiforme. Diazepinomycin inhibits the Ras-MAPK signaling pathway, crosses the blood-brain barrier and accumulates in gliomas possibly binding to the peripheral benzodiazepine receptor, a component of the mitochondrial permeability transition pore which is involved in the regulation and initialization of apoptosis. [11-13] Cudraflavone B, a prenylated flavone produced in *C. tricuspidata* with a long list of pharmacological activities such as anti-proliferative, monoamine oxidase (MAO) inhibitory effects, apoptotic activity in gastric carcinoma and melanoma cells as well as

hepatoprotective activity, neuroprotective effects and reactive oxygen species inhibition against glutamate-induced neurotoxicity.[14] Therefore, this natural product has the potential to treat neurodegenerative diseases such as Alzheimer's, Parkinson's disease, ischemia and may prove useful in treating variety of cancers.[15] These compounds illustrate the vast structural and pharmacological diversity of prenylated natural products.



**Figure 1. Prenylated compounds**

*Examples of prenylated compounds displaying biological and pharmacological activities.*

## **Prenyltransferases**

Prenyltransferases are a large and diverse family of enzymes responsible for catalyzing the transfer of prenyl groups onto an assortment of acceptor compounds.[16] Based on their biochemical and structural characteristics, as well as their primary amino acid sequences, this large family of enzymes is organized into smaller subcategories. Isopentenyl pyrophosphate (IPP) and dimethylallyl pyrophosphate (DMAPP) are products from the mevalonate pathway that can serve as the initial precursors for the prenyl pyrophosphate biosynthesis as well as prenyl donors for protein, peptide, tRNA and aromatic prenyltransferase enzymes.[17] In the biosynthesis of linear isoprenoids, prenyl diphosphate synthases catalyze the transfer of prenyl groups onto alkene carbons through sequential condensation reactions of IPP with other prenyl diphosphates[18]. Protein prenyltransferases are capable of transferring C<sub>15</sub> and C<sub>20</sub> carbon chains to nucleophilic cysteine residues near the C-terminus of the protein, while peptide prenyltransferases of the TruF family catalyze the posttranslational side chain prenylation of peptides.[19, 20] Prenyltransferases are also capable of modifying nucleic acids found in tRNA by attaching prenyl and geranyl groups onto the amino group of adenosine and sulfur atom in 5-methylaminomethyl-2-thiouridine nucleotides.[21, 22] Further, a large and well-studied family of aromatic prenyltransferase (PT) enzymes catalyze the transfer of prenyl moieties from prenyl donors such as dimethylallyl pyrophosphate (DMAPP, C<sub>5</sub>), geranyl pyrophosphate (GPP, C<sub>10</sub>), farnesyl pyrophosphate (FPP, C<sub>15</sub>) onto a large variety of aromatic acceptors.

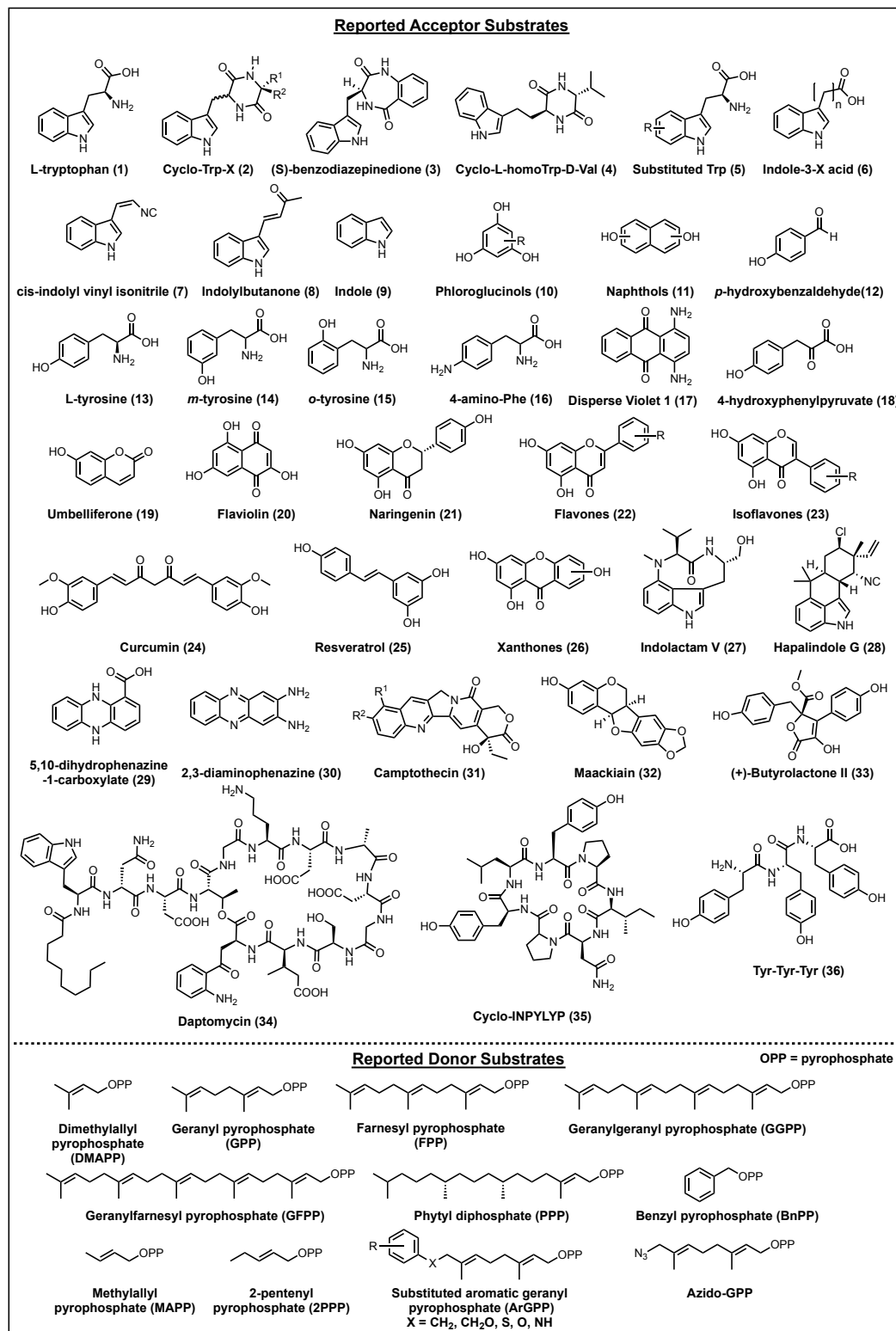


## **Aromatic Prenyltransferase**

In nature, aromatic PTs perform prenylation reactions on naphthalene, flavonoid, xanthone, coumarin, polyphenol and indole alkaloid aromatic compounds to generate multiple secondary metabolites in plants, fungi and bacteria.[23-25] Prenylation can occur late or early in the biosynthetic pathway, wherein the prenylated products can be further tailored by cyclization, hydroxylation, oxidation or further functionalization. Therefore, aromatic PTs contribute to the bioavailability and structural diversity of secondary metabolites by increasing the lipophilicity or contributing to the overall carbon scaffold. Prenylated secondary metabolites exhibit remarkable biological activities, e.g., anti-microbial activity, anti-cancer activity, anti-virus activity, and anti-oxidant activity[26-28]. Aromatic PTs are capable of performing C-C, C-O, C-N bond formation, and in certain conditions C-S prenylation.[12, 29-32] The reactions can take place in a forward or “normal” prenylation, where the C1 position of the prenyl donor connects directly to the aromatic moiety of the acceptor compound; alternately the reactions can take place in a “reverse” manner, where the C3 or tertiary carbon of the prenyl donor connects to the aromatic acceptor.[33] This class of enzymes all shares a common  $\alpha\alpha\beta\beta$  structural motif, known as the PT barrel.[34-41]

Within the large group of soluble aromatic PT enzymes, two main classes exist, the super family dimethylallyl tryptophan synthases (DMATS) and the CloQ/NphB group.[28, 36, 42-44] The DMATS superfamily is mostly responsible for prenylating nitrogen containing compounds such as indole alkaloids and indole alkaloid derivatives and are found primarily in fungi although some can be found in bacteria. Aromatic PTs belonging to the CloQ/NphB group utilize aromatic compounds such as phenols,

naphthalenes, phenazines, and flavonoids as aromatic substrates and are found in both bacteria and fungi.[45] Many of these soluble aromatic PTs exhibit considerable substrate promiscuity towards both donor and acceptor compounds accepting unnatural and synthetic analogues, a property critical to the development of synthetically useful biocatalysts (Figure 2). These enzymes are tools capable of regio-specifically functionalizing complex compounds in ways not currently achievable *via* synthetic means. Aromatic PTS are becoming a valuable tool for the late stage modification of complex natural and unnatural products, which is useful for the development of novel pharmaceutical candidates.



**Figure 2.** Aromatic prenyltransferase substrates. Known aromatic acceptor substrates and pyrophosphate donor substrates accepted by aromatic prenyltransferases.

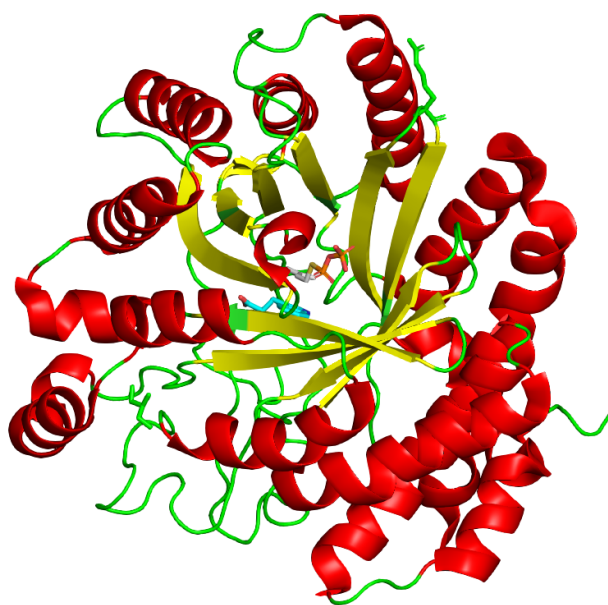
## **Global fold and Overall Structure**

### **Global fold**

To date, all enzymes belonging to the super family of soluble aromatic prenyltransferases adopt a similar  $\beta$ -barrel fold found exclusively within this class.[34, 36, 38, 40, 46-49] This fold was originally observed in the enzyme NphB, a prenyltransferase involved in the biosynthesis of the prenylated polyketide naphterpin in *Streptomyces* sp. Strain CL190.[36] This novel fold, which was termed a PT-barrel, is reminiscent of the better-known TIM barrel, which is a  $(\beta\alpha)_8$ -barrel fold, but differing in the connectivity of the secondary structure elements.[50] The PT-barrel is comprised of a cylindrical  $\beta$ -sheet of ten anti-parallel  $\beta$ -strands surrounded by a secondary ring of solvent exposed  $\alpha$ -helices. The secondary connectivity forms an AABB repeating motif; however, variability exists within known structures of soluble aromatic prenyltransferase due to helical kinks, additional helices and  $\beta$ -strands. The ten  $\beta$ -strands are arranged centrally, in a sequential antiparallel configuration generating a generous solvent accessible active site (Figure 3)[51].

As stated, most variations of the overall fold can be found within helices and loops surrounding the cylindrical  $\beta$ -sheet. For example, in the structure of EpzP, a phenazine prenyltransferase found in *Streptomyces cinnamonensis* DSM 1042, three additional  $3_{10}$ -helices were found, one of which is located on the resolved C-terminus, owing responsibility for sealing up one end of the barrel.[46] Similarly, C-terminal helices can be found on several other PT especially those belonging to the CloQ/NphB family (EpzP, FamD1/AmbP3, NphB, CloQ). In the case of CloQ, two  $\alpha$ -helices can be found following the final repeating unit; however, from the crystal structure only  $\alpha_{11}$  is

covering the opening of the barrel and participating in substrate interactions.[37] Moreover, much of the variance in the global fold can be attributed to the overall size differences between aromatic PTs. Aromatic PTs range in size from 302 to 464 residues. The additional residues of the larger enzymes are found in the extended loop regions located on either side of the barrel opening. These regions are typically seen in the DMATS superfamily of enzymes such as FgaPT2 and CdpNPT and have been shown to occlude the barrel opening and participate in substrate binding.[49] [35]



**Figure 3. Ribbon structure of FgaPT2 (PDB ID 314X)**

*Ribbon structure of FgaPT2 showing 10  $\beta$ -strands (yellow)  $\alpha$ -helices (red) of the PT-fold. The ligands tryptophan and DMSPP are seen bound in the central cavity of the barrel.*

### Structure

The overall interior architecture of the  $\beta$ -barrel's catalytic chamber is shared among the enzymes of this class.[35, 47, 52] The central cavity can be separated into three distinct

regions. Located at the top of the barrel is the prenyl donor binding pocket, lined with basic, positively charged residues, and non-charged polar residues which participate in ionic and hydrogen bonding interactions between the sidechains of the residues and the negatively charged phosphate moiety of the prenyl donor. The middle region of the cavity possess a conserved cluster of aromatic residues, mainly tyrosine. The hydroxyl group of the tyrosine form a “tyrosine shield”, protecting the allylic cation generated by the cleavage of the phosphate group from unwanted nucleophilic attack by solvent molecules. The third and bottom region of the cavity is composed mainly of hydrophobic residues responsible for the interaction and stability of the aromatic acceptor substrate and necessary intermediates. In many structures, the base of the barrel is occluded by structural elements such as rigid helices,  $\alpha$ -helix or  $3_{10}$ -helix, or extended loops.[36, 49, 53] The bottom region of the cavity offers the greatest amount variability, dictating substrate specificities and well as regio-specificities of the prenylated product.

### **Prenyl Binding Site**

Due to the ubiquitous nature of the pyrophosphate moiety of the donor substrates in prenyltransferase reactions, it is not surprising that the residues found in this region are highly conserved. The  $\alpha$ - and  $\beta$ -phosphates are anchored by hydrogen bonds with water molecules and polar residues as well as ionic interactions with positively charged basic residues lowering the ionization energy for the phosphate moiety.[19] Among the most structurally conserved residues surrounding the pyrophosphate binding site include arginine, lysine, asparagine and glutamine. The basic residues interact with the

pyrophosphate group and neutralize the negative charge while the polar residues and water mediated interactions fix the substrate into the active site via hydrogen bonding. The positively-charged pyrophosphate binding site does differ slightly between the magnesium-independent and magnesium-dependent PTs. It can be seen in the structure of NphB, a magnesium-dependent PTs, that two structurally conserved basic residues in the magnesium-dependent PT are missing, the loss of the positive charge is compensated by a magnesium ion that is coordinately via the aspartate side chain of Asp62.[36] Although the majority of the residues participating in the binding of the  $\alpha$ - and  $\beta$ -phosphate groups are conserved, some variability is found. Similarly, water mediated hydrogen bonding between polar residues and the electronegative oxygen atoms is seen in many structures of aromatic PTs but is not strictly conserved throughout. [36, 49] A highly structurally and functionally conserved cluster of tyrosine and other aromatic residues reside in this location of the active site surrounding the bridging oxygen to the prenyl group. Believed to be responsible for shielding the carbocation intermediate of the activated prenyl group, the ring of tyrosine and aromatic residues are a catalytic characteristic of all aromatic PTs. [23, 40, 47, 49, 54] Although the prenyl donors for aromatic PTs are variable in chain length, upon comparison of the structures it is clear that neither the pyrophosphate binding domain nor the “tyrosine shield” region is dependent on the chain length of the pyrophosphate donor nor aromatic acceptor.

## **Aromatic Binding Site**

Aromatic prenyltransferases are well known for their relaxed substrate specificities. [44, 54, 55] Tolerance is attributed to the spacious hydrophobic binding pocket, accommodating both the aromatic acceptor substrate and the carbon chain tail of the prenyl donor. The pocket is formed by nonpolar and hydrophobic residues of different sizes with varied numbers of interactions to the substrate. Structural similarities and differences of residues in contact with the substrate are of particular interest, as even small changes in the binding pocket have been known to alter the catalytic function. [56] Despite obvious differences in substrate specificity and regio-selectivity, residues found in the active site of aromatic PTs are relatively similar.

In many cases, the binding site residues remain functionally conserved but differ in the size of the amino acid side chain, changing the dimensions of the catalytic chamber. This is seen when comparing the acceptor binding site residues of aromatic PTs found in the DMATS family members, AtaPT, AnaPT, CdpNPT, FgaPT2 and FtmPT1. [47] The residues found in the aromatic acceptor binding pocket of AtaPT have relatively short side chains, lending the enzyme one of the largest solvent accessible surface area as well as greatest overall volume of the structurally characterized aromatic PTs . For example, residues Leu83, Gln172, Ser192 and G251 are found lining the active site of AtaPT, whereas the residues in equivalent positions of AnaPT are Phe103, Gln192, Phe212 and Phe265. Other spatial differences in the aromatic binding site can be seen in CdpNPT upon comparison to FgaPT2. Residue Thr102 and Arg244 in FgaPT2 are replaced with Ala271 and Ala313 in CdpNPT, enlarging the binding site relative to the



first. Additionally, positions Phe103, Gly126, Ile266, and Val340 on AnaPT are replaced with larger amino acids in CdpNPT corresponding to Leu109, Ala313, Phe272, and Met349. Variation in size of the active site is not limited to indole PT, size difference can be seen in the CloQ/NphB family of aromatic PTs as well. The binding site of EpzP was found to be larger than that of CloQ when comparing the larger amino acids Phe68, Phe161, Tyr233 and Trp295 to those smaller residues Thr64, Thr161, Ser236, Leu298 in EpzP. The various sizes of the aromatic binding site can explain the catalytic specificities of each PTs towards different aromatic substrates, prenyl donors as well as the orientation of the substrate.

Because the aromatic binding site accommodates both the prenyl chain of the donor and the aromatic substrate, and is the most variable amongst aromatic PTs, the majority of mutational studies have focused attention on this region. Mechanistic mutational studies have been conducted to better understand the essential residues required for catalysis. For example, a conserved glutamate residue, found within most indole prenyltransferases, was proven to be necessary for catalysis through selective mutations.[56] In the case of FtmPT1, single mutant E102Q was generated and incubated with its native substrates. This enzyme variant was unable to catalyze the conversion to tryprostatin B, to the limits of detection, in the presence of the precursor compound, brevianamide F. Interestingly, while both glutamate and glutamine are capable of forming hydrogen bonds with the N-1 atom of brevianamide F, only glutamate contains a negative charge at that position that can serve as a catalytic base, an apparent requirement for catalysis. As a follow up to these results, a similar more recent mutational study was performed in bacterial indole PT, AmbP1 which catalysis

the magnesium dependent selective C-3 geranylation of 3-((Z)-2'-isocyanoethenyl) indole. To study the importance of glutamate at position 209, mutations E209A and E209L were generated, thus replacing the carboxyl group with one that lacks hydrogen-bonding capabilities, and for Leu also or presents a bulkier isopropyl group. Both AmbP1 variants were inactive, further supporting the previous studies results, implicating the importance of the conserved glutamate in catalysis. In addition to a better understanding of the catalytic mechanism, the mutation of key residues has resulted in altered donor and acceptor selectivity, increased catalytic abilities, and new regio-specificities of enzyme.[38, 39] Recently, a structure-based engineering study was conducted on the endophenazine prenyltransferase, EpzP, wherein the active site was modeled after a more efficient endophenazine enzyme, PpzP.[47] The resulting single residue mutation, A285Q, yielded a 14-fold increase in catalytic activity when compared to the wild-type enzyme and a two-fold increase in catalytic activity compared to PpzP[46]. Similarly, indolactam prenyltransferases, TleC and MpnD which perform the reverse prenylation and geranylation of indolactam V respectively, a underwent single, double, and triple mutations switching the preference for either geranyl pyrophosphate (GPP) or DMAPP and changing the regio- and stereo-selectivity of the enzymes.[53] Mutant K174A of tryptophan prenyltransferase FgaPT2, generated through site-directed mutagenesis, was able to switch prenylation positions and prenyl attachment from its native C-4 normal prenylation capabilities to C-3 reverse reaction [57]. Aromatic PTs are becoming a well-studied biosynthetic enzyme family with strong potential for rational design and engineering to further advance the field of chemoenzymatic synthesis.

## Reactions mechanism

Despite a diverse substrate scope of aromatic PTs, the catalytic mechanism is expected to proceed in a similar manner. Specifically, the mechanism of the alkylation reaction has been characterized to proceed via a dissociative  $S_N1$ -like mechanism involving a carbocation-mediated electrophilic aromatic substitution [56, 58, 59]. To discern between the dissociative and associative mechanism, positional isotope exchange (PIX) experiments were performed using FgaPT2, a magnesium dependent prenyltransferase from *Aspergillus fumigatus*. [58] In this experiment, DMAPP was synthesized with an  $^{18}O$  -label on the bridging oxygen between the C1 dimethylallyl and pyrophosphate group [56]. The isotopically label DMAPP was enzymatically reacted with tryptophan and partially converted into product. The unreacted starting material was analyzed for isotopic scrambling via  $^{31}P$  NMR spectroscopy revealing that 15% of the isotopically labelled DMAPP contained the  $^{18}O$ -label in a non-bridging position. These results could possibly indicate a carbocation formation stable enough to allow for a P-O bond rotation in the active site by which the C-O bond cleavage is a reversible step. This would support a  $S_N1$ -like dissociative mechanism by which a partially stabilized carbocation could proceed forward to form the arenium ion with the aromatic substrate, or the reaction could proceed backwards, re-associating with the pyrophosphate group to re-form the starting material, DMAPP. To further test the mechanism, a known unreactive tryptophan analog, 6-fluorotryptophan, was used in the identical PIX experiment and revealed a similar ratio of non-bridging to bridging isotopic label in the

unreacted starting material. This is believed to further support the existence of a carbocation intermediate mediating the stepwise dissociative mechanism.

Solving the crystal structure of aromatic PTs provided further insight into the enzymatic mechanism. Structures were solved in the presence of the bound substrates providing a snap-shot of the active site immediately before catalysis. Generally, the pyrophosphate prenyl donors, DMAPP or GPP, bind to a highly conserved binding site, comprised primarily of positively charged and nonpolar residues, neutralizing the negative charge of the pyrophosphate moiety. Most soluble aromatic PTs are metal ion independent, excluding NphB whose prenyl binding site is dependent on the presence of a  $Mg^{2+}$  metal ion. Activation of the prenyl moiety involves the release of the pyrophosphate group, generating an allylic carbocation intermediate capable of multiple resonance forms leading to either a forward (C-2) or a reverse (C-3) addition of the alkyl group. Studies have indicated the double bond at the  $\beta$ -position of the prenyl donor is crucial for the stability of the carbocation, loss or change location did not yield any product[60]. As seen in the crystal structure of both NphB/CloQ and DMATS family of aromatic prenyltransferase, the carbon chain moiety of the prenyl donor is sandwiched between a conserved tyrosine residue and the aromatic substrate, facilitating the carbocation intermediate *via* stabilizing  $\pi$ -cation interactions and protecting its reaction with unwanted nucleophiles including solvents. Key charged residues, such as glutamate and lysine, are believed to assist during catalysis participating in deprotonation as the substrate undergoes an electrophilic aromatic substitution. Once the arenium ion is formed, the last deprotonation step is thought to be performed either directly by a key

residue or via water mediated interactions, restoring aromaticity and yielding the final product.

### **Conclusions and future directions.**

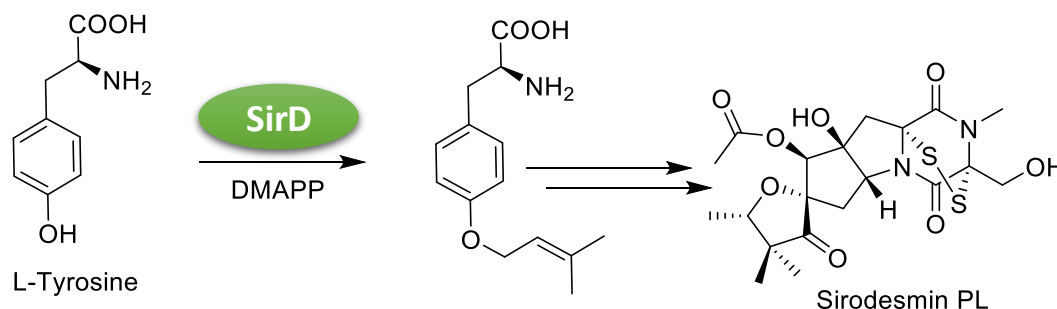
Over the last decade, many advances have been made towards the structural and biochemical characterization of aromatic PTs, resulting in over 50 biochemically characterized and 14 known structures for the enzymes of this class. This family of aromatic PTs has a diverse and vast acceptor substrate scope while also possessing the ability to accept multiple non-native substrates as well. Since aromatic PTs have the capability to functionally diversify biologically relevant scaffolds, thus producing natural and unnatural products they have the potential to chemoenzymatically synthesize a rich and untapped source of biologically and pharmacologically interesting compounds. High structural complexity and limited late stage functionalization methods prevent cost effective total synthesis of many natural products, limiting their usefulness as pharmaceuticals. Consequently, the high demand for chemo- and regio-selective catalysts is driving the development of biomimetic chemoenzymatic synthetic methodology. Although much of the proposed utility of aromatic PTs has been directed at their promiscuous aromatic substrate scope, we aim to showcase their ability to diversify natural products and pharmaceutical compounds through non-native alkylation. The successful utilization of unnatural acceptor and donor substrates with aromatic PTs will dramatically enhance the accessibility to a large unexplored chemical space.

## Chapter 2: Determination of Alkyl-Donor Specificity of Tyrosine-*O*

### Prenyltransferase SirD from *Leptosphaeria maculans*<sup>a</sup>

#### Introduction

Sirodesmin PL is a non-host specific phytotoxin produced by the pathogenic fungus *Leptosphaeria maculans*, known for causing the devastating blackleg disease in canola. Sirodesmin PL is a member of the epipolythiodioxopiperazine (ETP) class of secondary metabolites found in fungi. Like other ETPs, sirodesmin PL features a disulphide bridge across a diketopiperazine ring and exhibits antifungal, antibacterial, antiviral activities[61-63]. Interest in the properties of this secondary metabolite sparked a drive to investigate into the biosynthetic gene cluster and biosynthetic pathway for sirodesmin PL, leading to the discovery of a putative prenyltransferase gene *sirD*[64]. SirD was found to be responsible for the *O*-prenylation of L-Tyrosine in the first pathway specific step for the biosynthesis of sirodesminPL (Figure 4)[65]



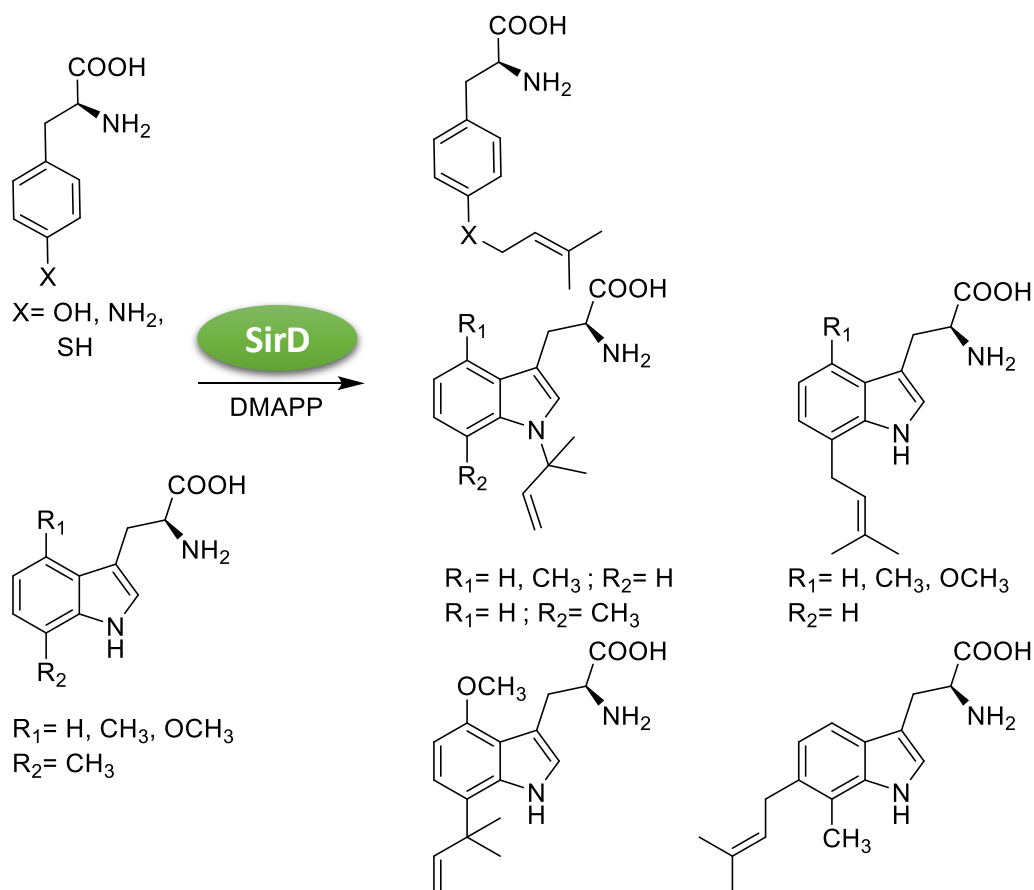
#### Figure 4. Sirodesmin biosynthetic pathway.

*Biosynthetic pathway for sirodesmin PL involving SirD in the first committed step of the pathway.*

<sup>a</sup> Reprinted with permission from: John Wiley and Sons and ChemBioChem, Authors Chandrasekhar Bandari, Erin M. Scull, Johanna M. Masterson, Rachel H. Q. Tran, Steven B. Foster, Kenneth M. Nicholas, and Shanteri Singh. “Determination of Alkyl-Donor Promiscuity of Tyrosine Prenyltransferase SirD from *Leptosphaeria maculans*” *ChemBioChem* **2017**, *18* (23), 2323-2327 DOI 10.1002/cbic.201700469. Copyright 2017 ChemBioChem. Copy-right permission has been included in the appendix.

Due to its high sequence similarity to known DMATS enzymes, prenyltransferase SirD is presumed to have the characteristic PT-fold, although the structure remains unknown. The functionality of SirD is independent of metal cations alike with other indole PTs. In addition to its native function in the synthesis of dimethylallyltyrosine, SirD has been shown to accept tryptophan, tryptophan derivatives, tyrosine and tyrosine derivatives such as 4-aminophenylalanine, 4-methoxytryptophan, 7-methyltryptophan, 4-mecraptophenylalanine, and L-3, 4- dihydroxyphenylalanine (Figure 5).[66, 67] Through the structural elucidation of the enzymatic products, SirD is capable of catalyzing O-, C-, N- and S-, prenylation[30, 66]. Specifically, SirD catalyzes normal S-prenylation of the sulfhydryl group of mecraoptophenylalanine as well as the normal N-prenylation of the amino group of 4-aminophenylalanine. SirD also catalyzes the N- and C- prenylations of tryptophan and tryptophan derivatives with the ability of transferring the prenyl group onto three different sites of the indole ring (N1, C6, C7), performing a normal prenylation at C6 and a reverse prenylation at C7 and N1. Through these studies, it was concluded that the electronic properties of the substituents on the indole ring played a role in determining the regio-specificity of the prenylation. Incubation with 4-methyltryptophan yielded a normal prenylation at C7 and a reverse prenylation at N1, while 4-methoxytrptophan gave a normal and reverse prenylation only a C7. Similar to other DMATs[60, 68, 69], SirD catalyzed reactions when incubated with unnatural pyrophosphate donors methylallyl pyrophosphate (MAPP), 2-pentenyl pyrophosphate (2-pen-PP), and benzyl pyrophosphate[32]. Further, it is believed that these studies provide evidence that SirD follows the dissociative electrophilic alkylation mechanism proposed for the prenylation by other DMATSs where the orientation of the

substrate in the enzymatic active site and electronic effects play a pivotal role in determining position and prenylation type.



**Figure 5. Products of SirD**

*SirD enzymatic products after incubation with tyrosine or tryptophan derivatives with DMAPP*

### Research Objectives

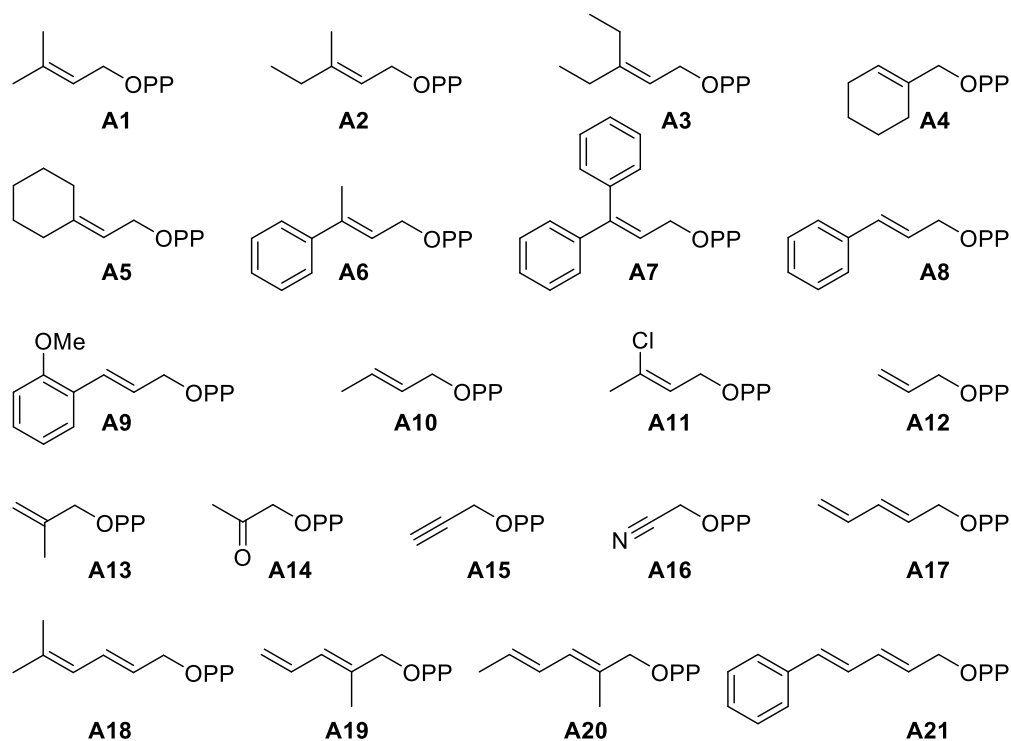
Studies thus far by different groups were focused on investigation of acceptor substrate specificity of SirD. These studies have demonstrated that SirD accepts derivatives of L-Tyr and L-Trp, catalyzing normal (attachment via C1' of DMAPP) or reverse (attachment via C3' of DMAPP) prenylation of *O*-, *N*-, *S*- and *C*- nucleophilic centers (Figure 5).[30, 66] However, apart from a couple of unnatural alkyl donor (2-pentenyl-



and benzyl- groups[70]) studies, not much is known about the unnatural alkyl-donor specificity of SirD. The goal of this study is to understand the alkyl donor specificity of SirD. To do this, we assessed the alkyl donor specificity of SirD using 21 synthetic alkyl pyrophosphate (alkyl-PP) analogues, performed the kinetic studies and identified the regio-specificity of the alkylated products using NMR spectroscopy.

## Contribution

In this study, the library of alkyl-donors (Figure 6) was synthesized by Dr. Chandrashekar Bandari, Johanna Masterson, and Rachel Tran. The large-scale tyrosine alkylation reaction, HRMS analysis and NMR characterization was done by Dr. Chandrashekar Bandari. My contributions involve expression and purification of SirD, reverse phases high-performance liquid chromatography (RP-HPLC) assessment of alkyl-donor specificity of SirD and determination of kinetic parameters for SirD catalyzed reaction with diverse alkyl-donors. Most of the contents including text, tables and figures for this chapter were directly taken from the publication .[71]



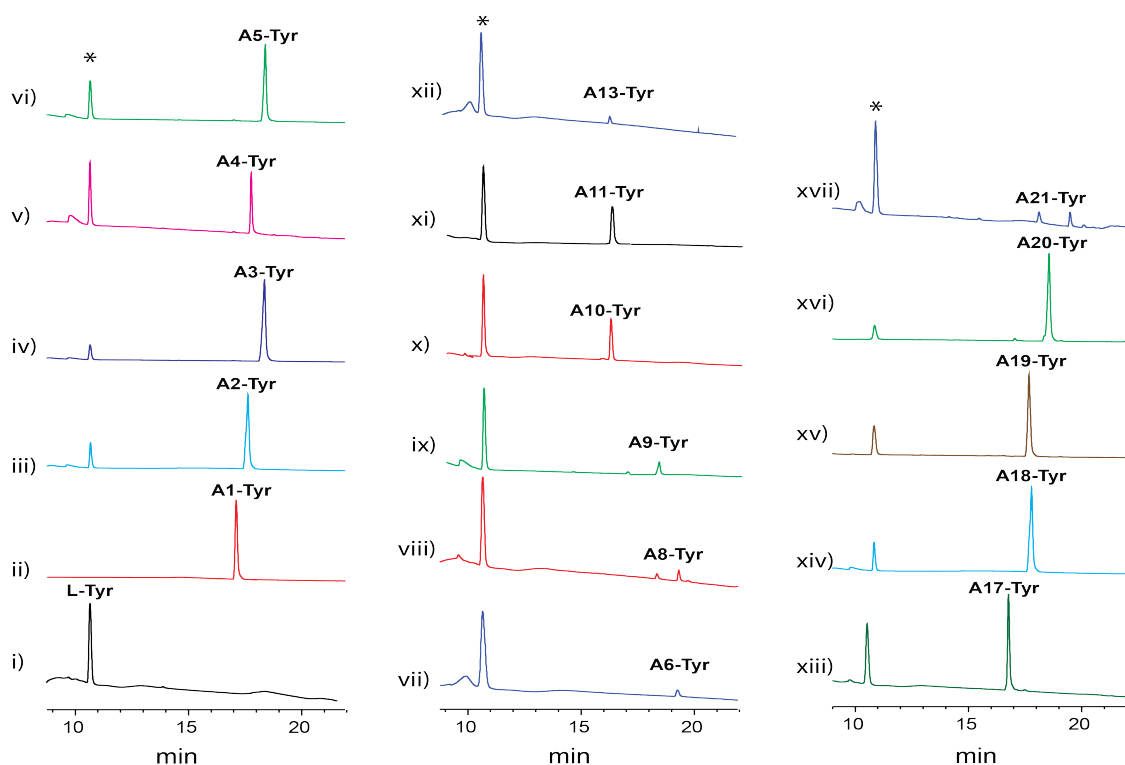
**Figure 6. Library of synthetic alkyl-PP**

*Pyrophosphate analogues assessed with SirD catalyzed reaction with L-Tyr.*

## Results and Discussion

The recombinant SirD was overexpressed in *Escherichia coli* BL21(DE3) cells transformed with a codon optimized synthetic *sirD* gene in pET28a vector. The

resulting SirD with *N*-terminal His<sub>6</sub>-fusion protein was purified to homogeneity via Ni-NTA affinity chromatography. Standard uniform assay conditions (1.2 mM alkyl-PP analog, 1 mM L-Tyr, 5 μM SirD, 25 mM Tris, 5 mM MgCl<sub>2</sub>, 50 mM KCl, pH 7.5, 16 h at 35 °C) were adopted to facilitate the assessment of donor substrate specificity of the SirD with synthetic alkyl-PP analogues (Figure 6). Production of alkyl-L-Tyr derivatives was determined by a RP-HPLC end point assay (Figure 7) with subsequent confirmation by HRMS coupled with liquid chromatography (LC) analysis for all positive reactions to afford complete analysis of the desired product (Table 1).

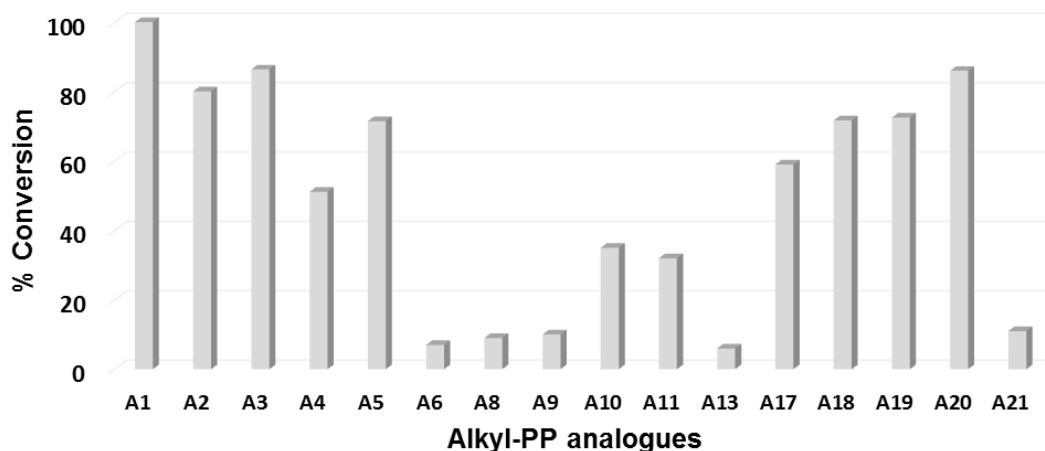


**Figure 7. RP-HPLC chromatograms of SirD catalyzed reactions.**

*Reactions were incubated with L-Tyr and alkyl-PP analogues that led to the formation of corresponding alkyl- L-Tyr analogues. The remaining L-Tyr and product of the reaction are labeled with \* symbol and compound number respectively.*

## RP-HPLC and HRMS Studies

An initial assessment of the SirD catalyzed alkylation of L-Tyr with 21 synthetic alkyl-PPs based upon RP-HPLC of analytical scale reaction is illustrated in Figure 7, wherein 16 of 21 synthetic analogues in SirD catalyzed reactions resulted in corresponding alkylated L-Tyr products. Fourteen of the 16 positive SirD reactions produced a single corresponding mono-alkylated L-Tyr products, whereas, two analogues (A8 and A21) resulted in two mono-alkylated L-Tyr products each as confirmed by HRMS analysis (Table 1). Of the 21 putative substrates tested with SirD, 9 led to appreciable (>50%) production of the corresponding *O*-alkyl- L-Tyr, an additional 2 led to moderate (25-50%) conversion, and 5 offered detectable product (<25%) under the conditions described (Figures 9 and 10). Except for A7, A12 and A14, all analogues with a double bond at the  $\beta$ -position served as substrates.



**Figure 8. Percent conversion**

*Percent conversion of alkyl-PP analogues to the corresponding alkyl-L-Tyr catalyzed by SirD based upon RP-HPLC. No product formation was observed in the absence of SirD or alkyl-PP analog.*

**Table 1.** Summary of HRMS data of alkyl-*O*-Tyr analogues from SirD catalyzed alkylation reaction with L-Tyr and synthetic alkyl-PP analogues

Enzyme Product	Chemical Formula	Calculated Mass	Observed Mass
<b>A1-Tyr</b>	C <sub>14</sub> H <sub>20</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	250.1443	250.1446
<b>A2-Tyr</b>	C <sub>15</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	264.1600	264.1599
<b>A3-Tyr</b>	C <sub>16</sub> H <sub>24</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	278.1756	278.1756
<b>A4-Tyr</b>	C <sub>16</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	276.1600	276.1585
<b>A5-Tyr</b>	C <sub>17</sub> H <sub>24</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	290.1756	290.1757
<b>A6-Tyr</b>	C <sub>19</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	312.1600	312.1599
<b>A8-Tyr</b>	C <sub>18</sub> H <sub>20</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	298.1443	298.1444
<b>A9-Tyr</b>	C <sub>19</sub> H <sub>22</sub> NO <sub>4</sub> [M+H] <sup>+</sup>	328.1549	328.1553
<b>A10-Tyr</b>	C <sub>13</sub> H <sub>18</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	236.1287	236.1274
<b>A11-Tyr</b>	C <sub>13</sub> H <sub>17</sub> ClNO <sub>3</sub> [M+H] <sup>+</sup>	270.0897	270.0895
<b>A13-Tyr</b>	C <sub>13</sub> H <sub>18</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	236.1282	236.1274
<b>A17-Tyr</b>	C <sub>14</sub> H <sub>18</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	248.1287	248.1283
<b>A18-Tyr</b>	C <sub>16</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	276.1600	276.1605
<b>A19-Tyr</b>	C <sub>15</sub> H <sub>20</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	262.1443	262.1443
<b>A20-Tyr</b>	C <sub>16</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	276.1600	276.1604
<b>A21-Tyr</b>	C <sub>20</sub> H <sub>22</sub> NO <sub>3</sub> [M+H] <sup>+</sup>	324.1600	324.1597

### Kinetic Studies

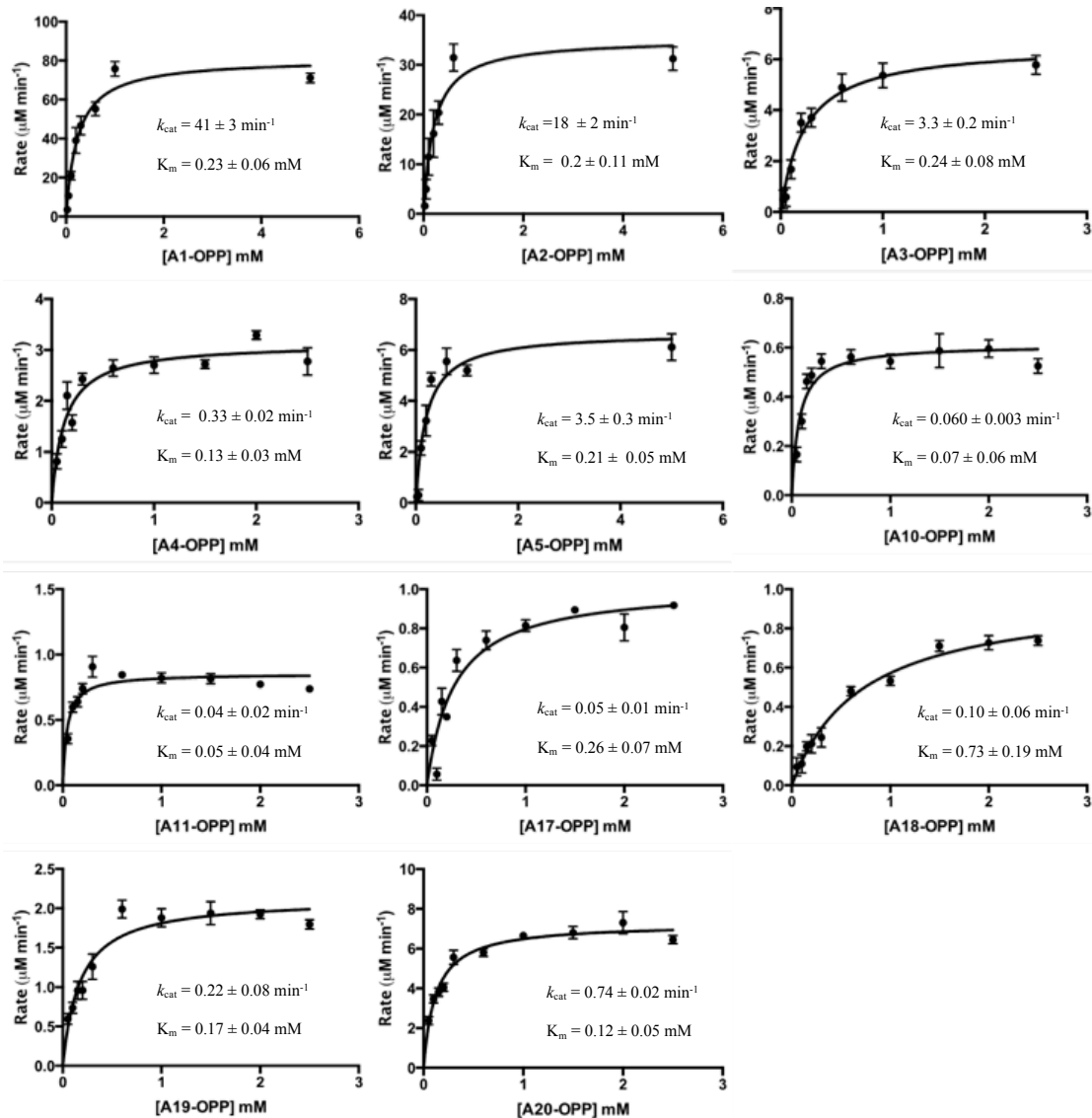
The SirD steady-state kinetic parameters for reactions with alkyl-L-Tyr yield > 25% were determined based on product formation monitored by HPLC (Table 2, Figure 9). Importantly, the  $K_M$  values for most of the analogues remained similar except for **A18**, where a 5 - 6-fold increase in the  $K_M$  compared to **A20** is observed, likely caused by the steric factors in the active site of the enzyme due to the presence of extra terminal methyl group. Surprisingly, **A11**, where an electron withdrawing chloro group replaced one of the methyl groups of DMAPP, and **A10**, which lacks a methyl group of DMAPP, were equally efficient (Table 2). However, both **A10** and **A11** were ~200 fold worse compared to the natural alkyl donor, **A1**, even though their  $K_M$  was 3 – 4 times lower than **A1**. Interestingly, where direct comparisons could be made, the analogues that contained groups to enhance the stability of carbocation[72] via increased conjugation

or resonance stabilization correlated with the increased catalytic efficiency. For example, the  $k_{cat}$  values of the reaction correlated with conjugation by methyl group in the order, **A1** > **A2** > **A3** > **A10** > **A17** (Table 2). Even among the analogues containing conjugated double bonds (**A17**- **A20**), the increase in the  $k_{cat}$  values of the reaction correlated with the presence of methyl group in the order, **A20** > **A19** > **A18** > **A17** (Table 2). While the literature precedence suggests cinnamyl carbocations to be highly stable,[73] cinnamyl analogues did not serve as good alkyl donors for the SirD reaction with L-Tyr. This is likely attributed to the steric hindrance in the active site of the enzyme due to the aromatic group in the analog (Table 2). Similar results were obtained for other analogues containing aromatic groups. However, the analogues that contained a triple bond at the  $\beta$ -carbon (**A15** and **A16**) did not serve as donors, likely due to the reduced stability of the carbocation[72] by the attached  $sp$ -hybridized alkynyl group in addition to the steric requirement of the enzyme. Similarly, the analogues with electron withdrawing groups at the  $\beta$ -carbon (**A14**) did not serve as donors.

**Table 2.** Kinetic parameters for SirD with 1 mM L-Tyr and 0.025 - 6 mM alkyl-PP analog in 25 mM Tris, 5 mM MgCl<sub>2</sub>, 50 mM KCl, pH 7.5 at 35 °C.

Alkyl-PP analog	$k_{cat}$ (min <sup>-1</sup> )	$K_M$ (mM)	$k_{cat} / K_M$ (mM <sup>-1</sup> min <sup>-1</sup> )
<b>A1</b>	41 ± 3	0.23 ± 0.06	178 ± 15
<b>A2</b>	18 ± 2	0.20 ± 0.11	90 ± 31
<b>A3</b>	3.3 ± 0.2	0.24 ± 0.08	14 ± 3
<b>A4</b>	0.33 ± 0.02	0.13 ± 0.03	2.5 ± 0.6
<b>A5</b>	3.5 ± 0.3	0.21 ± 0.05	16 ± 2
<b>A10</b>	0.060 ± 0.003	0.07 ± 0.06	0.9 ± 0.2

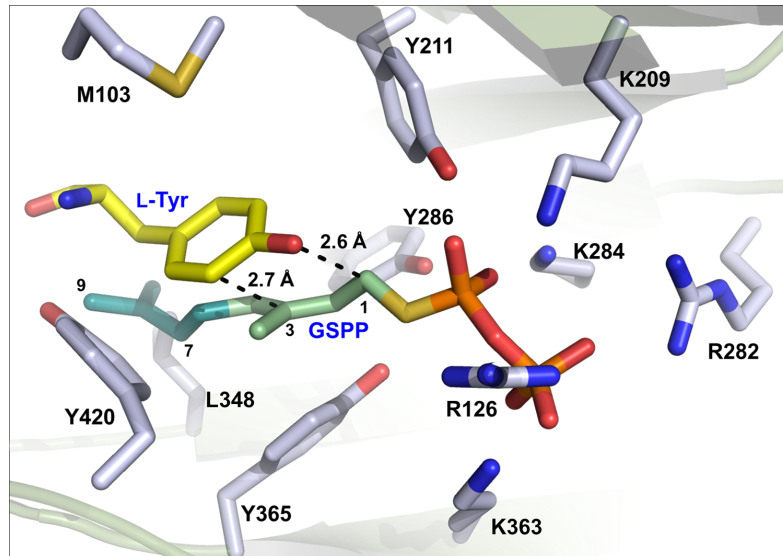
A11	$0.04 \pm 0.02$	$0.05 \pm 0.04$	$1.0 \pm 0.3$
A17	$0.05 \pm 0.01$	$0.26 \pm 0.07$	$0.20 \pm 0.06$
A18	$0.10 \pm 0.06$	$0.73 \pm 0.19$	$0.14 \pm 0.02$
A19	$0.22 \pm 0.08$	$0.17 \pm 0.04$	$1.3 \pm 0.3$
A20	$0.74 \pm 0.02$	$0.12 \pm 0.05$	$6.1 \pm 0.8$



**Figure 9. Michaelis-Menten kinetic plots for SirD reactions**  
*Kinetic parameters determined by Michaelis-Menten plots with 1mM Tyrosine and varying concentrations of pyrophosphate analogues*

## SirD Structural Model

In order to understand the alkyl-PP chain-length selectivity by SirD, we generated a homology model of SirD based on the template structure of FgaPT2 (PDB accession number 3I4X) that covered 92% sequence length of SirD (a.a. 27-440), and shared 30% sequence identity and 65% sequence similarity with SirD using SWISS-MODE.[74] In the SirD structural model, the L-Tyr and the non-hydrolyzable form of DMAPP (DMSPP) were docked posed for catalysis in a similar manner as in other ABBA PT structures (Figure 10).[24, 36, 40, 41, 47, 49]



**Figure 10: Structural model of SirD.**

*Structure is based on the template structure of FgaPT2 (PDB 3I4X) with docked L-Tyr (colored in yellow) and dimethylallyl thiopyrophosphate (DMSPP, colored in green). Colored in light-blue stick models are the active site residues of SirD interacting with DMSPP and L-Tyr.*

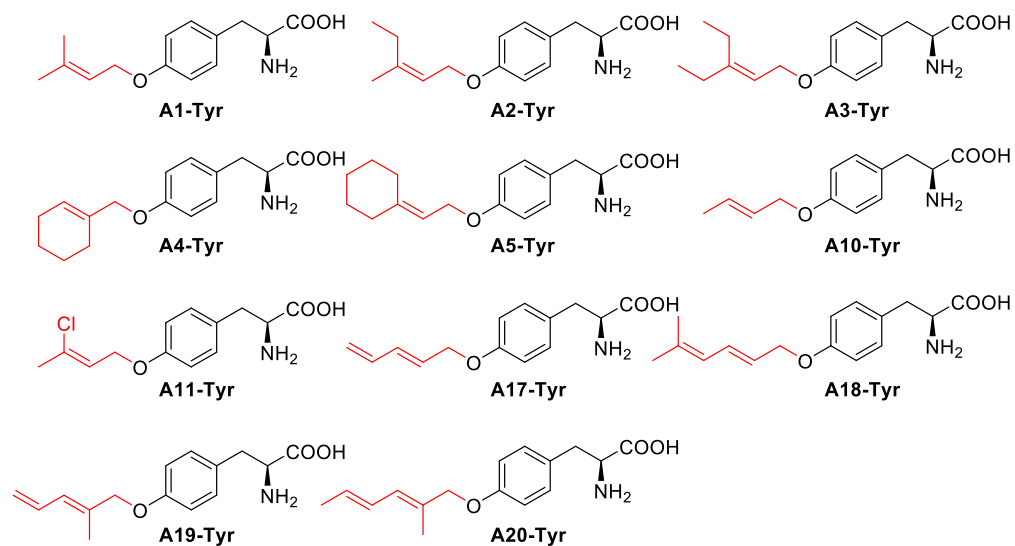
Briefly, the docking of the DMSPP and L-Tyr in the active site of SirD was performed to be consistent with the conserved interactions found among the class of ABBA PTs.[24, 36, 40, 41, 47, 49] These include hydrogen bonding interactions of the amine side chain of conserved basic residues (R126, K209, R282, K284, and K363; Figure 10)



and the conserved aromatic residues (Y211, Y286, Y365 and Y420) positioned to shield and stabilize the alkyl carbocation via  $\pi$ -cation interactions (Figure 10). Interestingly, according to the SirD model displayed in Figure 10, the distance between -OH of L-Tyr and -C1' of DMSPP; as well as C-3 of L-Tyr and -C3' of DMSPP are  $\sim 2.6$  Å. While the distance may facilitate either C1'- normal *O*-prenylation or C3'-reverse prenylation, the nucleophilicity of the -OH group seems to drive the observed normal *O*-alkylation of L-Tyr by SirD. In the SirD model, the alkyl chain of the natural donor, **A1** seem to stack on the aromatic ring of the Y365. The current SirD model suggests steric clash to be a potential contributing factor to the low or no turn-over of alkyl donors containing long carbon chain or bulky aromatic groups.

### **NMR Studies**

The SirD alkylation reactions with product yield > 25% were scaled up and purified via RP-HPLC to understand the regio-chemistry of alkyl group transfer. Based on the NMR characterization of the regioselectivity of SirD catalyzed alkyl-transfer performed by Dr. Bandari, all alkyl-L-Tyr (Figure 6, **A1**, **A2**, **A3**, **A4**, **A5**, **A10**, **A11**, **A17**, **A18**, **A19**, **A20**) were confirmed as *O*-alkylation of L-Tyr analogues (Figure 11). Additionally, SirD produced two mono-alkylated L-Tyr products in the presence of **A8** and **A21**. This suggests possibility of alkylation of L-Tyr at other nucleophilic center apart from -OH of L-Tyr. Based on the proximity of -C3 of L-Tyr to -C3' of DMSPP in the current SirD model (Figure 10), the reverse alkylation at -C3 of L-Tyr seems more likely, in addition to the normal *O*-alkylation. However, a better understanding of the relationship between substrate and SirD regioselectivity requires more studies including the crystal structures of SirD bound to different substrates.



**Figure 11. NMR confirmed structures.**

*Structures of alkyl-tyrosine products of SirD reaction with alkyl-PP analogues determined by NMR.*

## Conclusions

In summary, the presented work demonstrates the pliable nature of SirD towards a wide range of allylic alkyl donors, which is expected to be further expanded via enzyme engineering. In addition, our work also demonstrates SirD to catalyse mostly regiospecific, normal *O*-alkylation of L-Tyr. Thus, these studies highlight the potential utility of SirD for chemoenzymatic differential alkylation of natural products and open the door for interrogation of the alkyl donor specificity of other prenyltransferases.

## Experimental

Unless otherwise stated, all chemicals and reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA), Acros (New Jersey, USA), Alfa-Aesar (Ward Hill, MA, USA) or TCI (Portland, OR, USA) and were reagent grade or better. PD-10 column and Ni-NTA superflow columns were purchased from GE Healthcare (Piscataway, NJ).

## General methods

All reactions were conducted in oven- or flame-dried glassware under a nitrogen atmosphere with anhydrous solvents, unless otherwise noted. Reactions were monitored by thin-layer chromatography (TLC) (EMD Millipore Corp, Billerica, MA, USA) and visualization was accomplished with UV light (254 nm) followed by staining with phosphomolybdic acid solution or anisaldehyde solution and heating. Flash column chromatography was performed using ACS grade solvents and silica gel (SiliCycle Inc, P60, particle size 40-63  $\mu\text{m}$ ). HPLC was accomplished using Agilent 1220 system equipped with a DAD detector. **Method A (HPLC):** To monitor SirD reactions, analytical reverse-phase (RP) HPLC employed a Gemini-NX, C-18 (5  $\mu\text{m}$ , 4.6 mm  $\times$  250 mm) column (Phenomenex, Torrance, California, USA) [gradient of 1% B to 10% B over 10 min, 10% B to 50% B over 5 min, 50% B to 100% B for 12 min, 100% B to 1% B over 1 min, 1% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 1 mL min<sup>-1</sup>; A<sub>220</sub>]. The reaction was monitored by the retention time difference between starting material and product. **Method B (HPLC):** Semi-preparative RP HPLC was conducted on a Gemini-NX, C-18 (5  $\mu\text{m}$ , 10 mm  $\times$  250 mm) column (Phenomenex, Torrance, California, USA) to purify the L-Tyr analogues [gradient of 1% B to 10% B over 10 min, 10% B to 50% B over 5 min, 50% B to 100% B for 12 min, 100% B to 1% B over 1 min, 1% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 2 mL min<sup>-1</sup>; A<sub>220</sub>]. High-resolution mass spectrometric (HRMS) data and liquid chromatography mass spectrometric (LCMS) were obtained on Agilent 6545-QTOF W/1290 HPLC mass spectrometer at the University of Oklahoma, Department of Chemistry and Biochemistry.

### **Overexpression and purification of SirD**

The *Leptosphaeria maculans* SirD in pET28a vector bearing *N*-terminal-His<sub>6</sub> fusion was provided by Prof. Jon Thorson (University of Kentucky). Recombinant SirD with *N*-terminal-His<sub>6</sub> (herein referred to simply as SirD), was expressed in a *E. coli* BL21(DE3) host in the presence of 50 µg mL<sup>-1</sup> of kanamycin at 37 °C to an OD<sub>600</sub> of ~0.6, reduced the temperature to 20 °C, induced with 0.5 mM IPTG, and continued to grow at 20 °C for another 16 h. The cells were harvested by centrifugation (30 min, 5000 rpm), resuspended in buffer A (20 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 10 mM imidazole, pH 7.8). The cells were lysed for 30 min on ice by sonication (Fisher Scientific Model FB505; Thermo Fisher Scientific, Waltham, MA; 500 W, 4 x 30 s pulses, ~1 min between pulses) on ice. SirD was purified via affinity chromatography (5 mL HiTrap Ni-NTA chelating column, GE Healthcare, Piscataway, NJ) following standard linear gradient (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, pH 8.0 with a linear gradient of imidazole of 10-500 mM) using an NGC™ liquid chromatography systems (Bio-Rad, USA). Buffer exchange of pooled fractions containing the purified protein was accomplished using a PD-10 column (GE Healthcare) eluted with 25 mM Tris-HCl, 50 mM KCl, pH 8. Protein concentrations were determined by Bradford assay (Bio-Rad, Hercules, CA, USA) using BSA as a standard. All studies were conducted with SirD, that retained the *N*-terminal-His<sub>6</sub> affinity tag.

### **In-vitro SirD assay**

In vitro SirD reactions were conducted in a volume of 20 µl with 1.2 mM alkylpyrophosphate analog (compound **A1** – **A21**), 1 mM L-Tyr and 5 µM purified SirD in 25 mM Tris buffer pH 8.0, 5 mM MgCl<sub>2</sub>, 50 mM KCl, incubated at 35 °C for 16 h.

Reactions were quenched by adding an equal volume of methanol followed by centrifugation (10,000 x g for 15 min) to remove the precipitated protein and product formation for each reaction was subsequently analyzed by RP-HPLC using method A. For each reaction, percent yield was based upon the integration of species at 254 nm and calculated by dividing the integrated area of product by the sum of integrated area of product and/or and the remaining substrate. All putative products were subsequently confirmed by HRMS with positive (+) and/or negative (-) mode. Kinetic parameters were assessed using 1 mM L-Tyr and variable alkyl-PP analogues (0.1 - 5 mM). Assays were conducted in triplicate and all rates were confirmed to be linear. The kinetic curves were fit to Michaelis-Menton kinetics using Prism 5.04 (GraphPad Software, Inc. La Jolla, CA 92037 USA). The reactions with > 25% product yield were scaled (>500  $\mu$ M L-Tyr), purified by semi-prep HPLC using method B and the putative new products were confirmed by NMR.

### **SirD structural model**

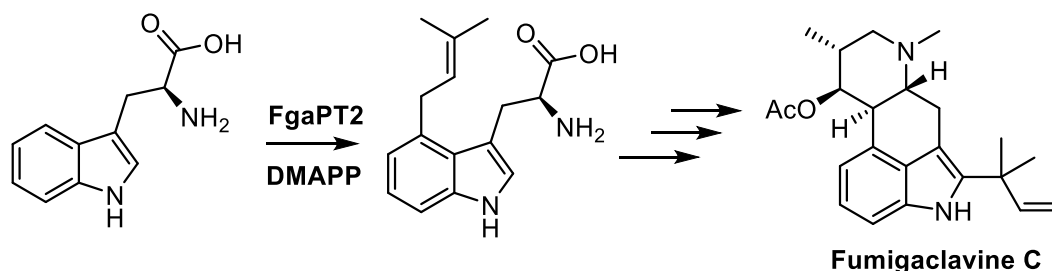
A SirD BLAST search against the structural database, PDB revealed SirD to have 30% sequence identity with Trp-C4-PT, FgaPT2 (PDB 3I4X). A SirD homology model was constructed based upon FgaPT2 using Swiss-Model.[74] Atomic coordinates of the ligands, L-Trp and DMSPP were obtained from FgaPT2 structure (PDB 3I4X). L-Tyr model was generated by manual editing of the L-Trp using the builder module of PyMol (<http://www.pymol.org>) with no additional energy minimization performed on the structures. Both L-Tyr and DMSPP were subsequently modeled in the active site of SirD in a manner consistent with the conserved interactions observed for other ABBA-PTs. [24, 36, 40, 41, 47, 49] The conserved interactions used for modeling-in the

ligands include the stacking interaction of L-Tyr with the prenyl moiety of DMSPP and the hydrogen bonding interactions of amine side chain of conserved basic residues (R128, K209, R282 and K284) with the pyrophosphate moiety. Figures were generated with PyMol.

## Chapter 3. Determination of Alkyl-Donor Specificity of a Tryptophan-C4- Prenyltransferase FgaPT2 from *Aspergillus fumigatus*.<sup>b</sup>

### Introduction

Filamentous fungi, such as *Aspergillus*, possess some of the most impressive chemical factories known in biology and are an incredible source of biologically active natural products.[27] Even more, their natural products are being used as scaffolds for more focused diversity oriented pharmacological and drug libraries.[75] In the biosynthesis of ergot alkaloids, DMATS has been identified to catalyze the first pathway-specific reaction, attaching a prenyl moiety onto L-tryptophan.[76] In 2005 a DMATS gene *fgaPT2*, was identified in the genome sequence for *A. fumigatus*. FgaPT2 is a soluble indole PT responsible for catalyzing the transfer of a prenyl group onto the C-4 position of L-tryptophan, which is the first step in the biosynthesis of many indole alkaloids such as the fumigaclavines (Figure 12).[77] This prenyl group can then be incorporated into a polycyclic carbon scaffold giving rise to diverse active natural products.



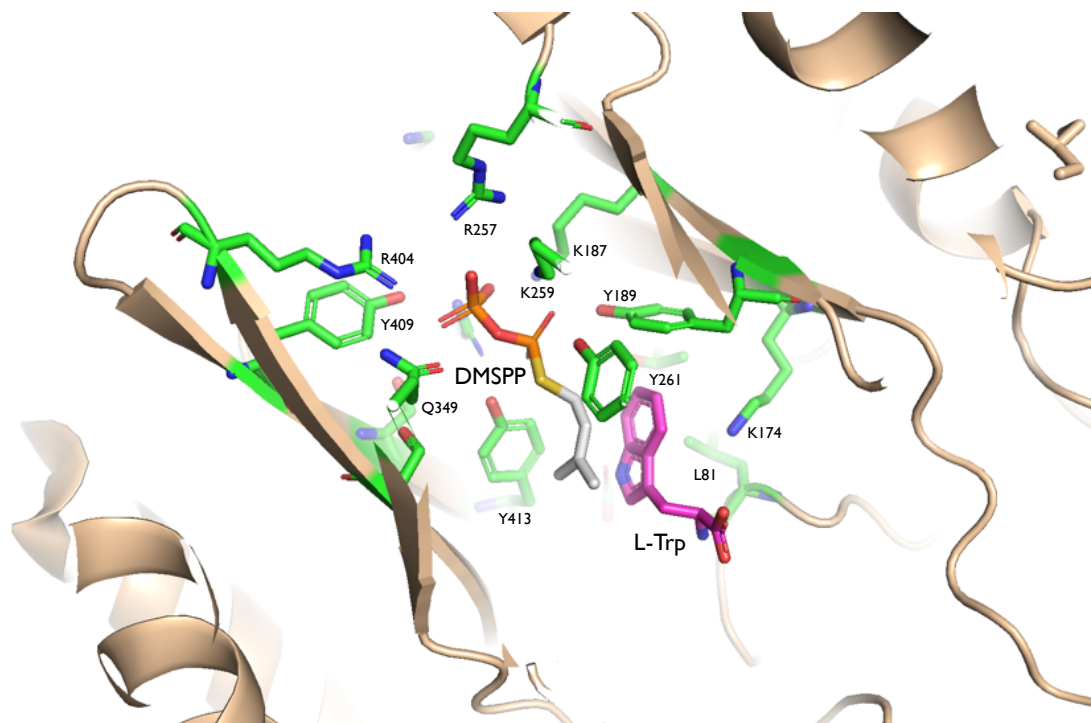
**Figure 12. The role of FgaPT2 in fumigaclavine C biosynthesis**

<sup>b</sup> Reprinted with permission from: Royal Society of Chemistry and MedChemComm, Authors Chandrasekhar Bandari, Erin M. Scull, Tejaswi Bavineni, Susan L. Nimmo, Eric D. Gardner, Ryan C. Bensen, Anthony W. Burgett and Shanteri Singh. "FgaPT2, a biocatalytic tool for alkyl-diversification of indole natural products" *MedChemComm* **2019**, MD-RES-03-2019-000177.R2. Copyright 2019 MedChemComm.

<https://pubs.rsc.org/en/journals/journalissues/md#!issueid=md010006&type=current&issnprint=2040-2503>

FgaPT2 is the first DMATS structure to be solved, shortly after the bacterial aromatic PT enzyme NphB structure was published.[36] FgaPT2 shares the PT barrel fold, common among all aromatic PTs, which contains 5 repeating  $\alpha\beta\beta$  units. The significant differences between NphB and FgaPT2 structures can be found in the connecting extended loop regions and extra  $\alpha$ -helices in FgaPT2. The structure of FgaPT2 complexed with the native substrate tryptophan and DMSPP revealed the architecture of the catalytic binding site and insight into understanding the mechanism of alkylation (Figure 13). L-tryptophan binds to a mostly hydrophobic pocket, forming a hydrogen bond between the indole NH and E89. [35] This likely enhances the reactivity toward the carbocation by stabilizing the positively charged intermediate. This hydrogen bond combined with several others is responsible for orienting the tryptophan within the catalytic chamber, ultimately determining its regioselectivity. Tryptophan forms hydrogen bonds with the side chains of Y191 and R244, and with main chain atoms of I80 and L81.[35] The basic side chain on K174 is located perfectly to help abstract the proton after C4 alkylation. Divalent cations  $Mg^{2+}$  and  $Ca^{2+}$  enhance the rate of FgaPT2 but are not required.[77] The negative charge of the pyrophosphate binds to the positively charged residues R100, K187, R257, K259, and R404. The oxygen atoms are able to hydrogen bond with Y189, Y261, Y345, Y413, and Q343 (Figure 14).[35] The carbocation that is formed is stabilized by the  $\pi$  electrons of the substrate indole ring and Y345. The carbocation is shielded from reacting with water by Y189, Y261, Y345, Y398, and Y413.[35]

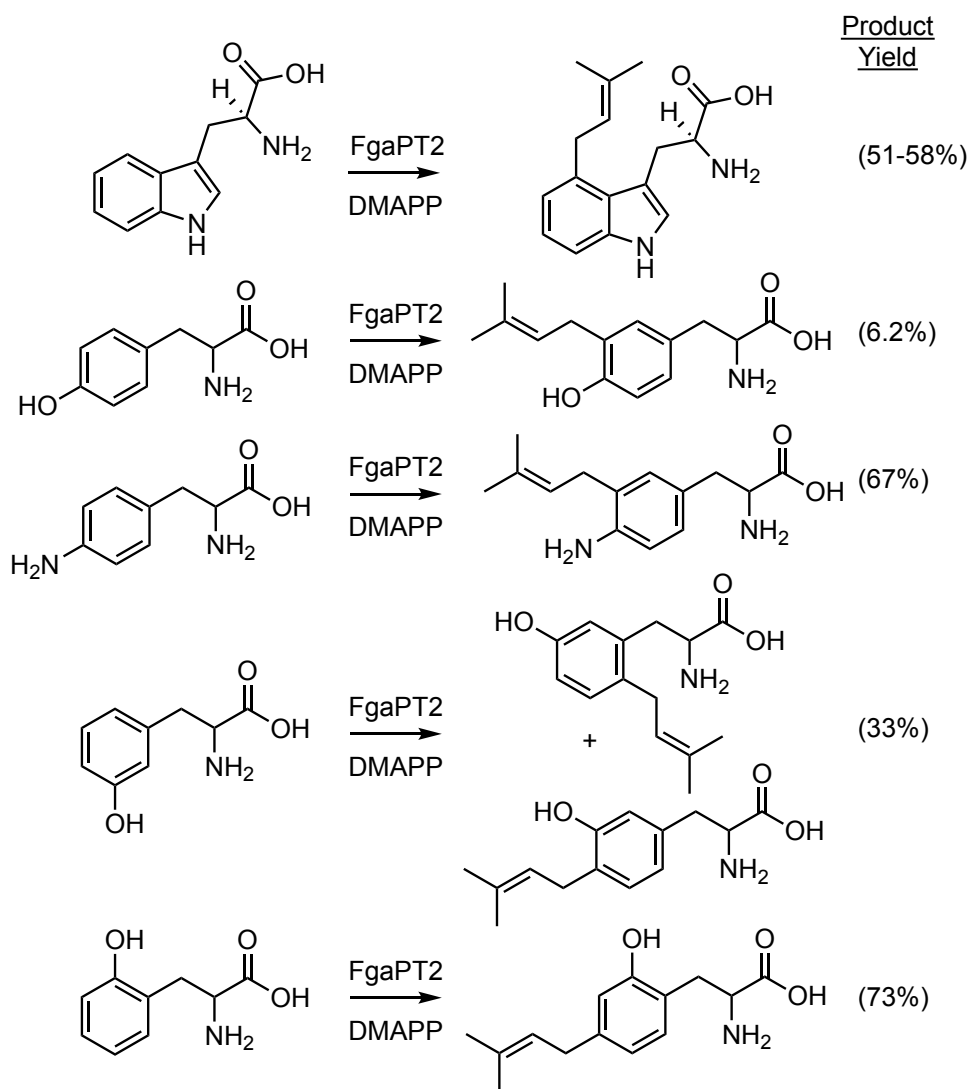




**Figure 13. Active site of FgaPT2**

*DMSPP in the active site side chains binding with pyrophosphate moiety. Residues in the active site interacting with L-tryptophan and carbon chain of DMSPP[35]*

In addition to its native catalytic function, FgaPT2 also accepts a variety of alternative substrates (Figure 16). For example, the enzyme can transfer a prenyl group to the 3 position of tyrosine, the 5 position of ortho-tyrosine, and the 4 and 6 position of meta-tyrosine.[78] Tyrosine and 4-amino-phenylalanine are accepted by FgaPT2 resulting in C3 prenylation, 6.2% and 67% yield respectively.[79] Indole-3-propionic acid and L- $\beta$ -homotryptophan were C4 prenylated by whole cells overexpressing FgaPT2.[80] Cyclo-L-homotryptophan-D-valine was well accepted as a substrate resulting in a 1:1 ratio of C4 and C5 prenylated products.[81] 1-naphthol has also been successfully prenylated with FgaPT2.[82] Daptomycin is also accepted by FgaPT2 to generate a indole N-1 prenylated product.[83]

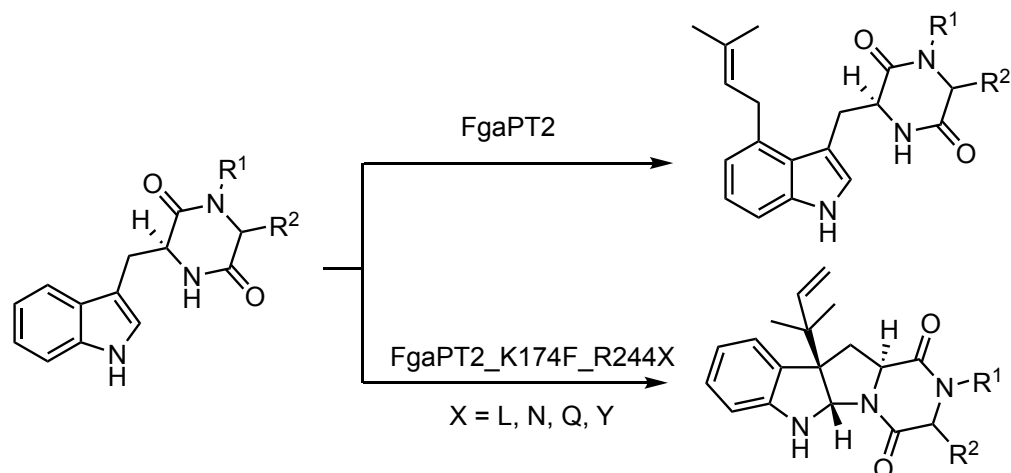


**Figure 14. Accepted aromatic substrates by FgaPT2.**

*Known reactions catalyzed by FgaPT2 and non-native substrates*

DMAPP is the natural donor substrate for FgaPT2, however a couple of unnatural pyrophosphates have been successfully transferred using this enzyme. Monomethylallyl pyrophosphate (MAPP) gave a 1:1 mixture of C4 and C5 alkylated tryptophans, while 2-pentenyl pyrophosphate alkylated the 5 position exclusively.[60] Benzyl pyrophosphate was also a substrate for FgaPT2 when incubated with tryptophan analogues, result in predominantly in C5 benzylation. The only 5 substituted indole accepted was 5-OH-Trp which was C6 benzylated by FgaPT2.[68] Native FgaPT2

cannot accept GPP or FPP as donors, however certain mutants have been engineered to accept longer chain donors. The FgaPT2 M328G/R244Q mutant was able to geranylate cyclo-L-Trp-L-Trp at the C4 position.[84] Met328 plays a major role in determining the maximum size of the donor molecule, with many of the smaller side chain mutants (C,A,T,S,G,V,N) having significant activity with GPP. FgaPT2 M328A was also able to accept FPP, although with very low turnover.[85] Site saturation mutagenesis was done on T102, K174, and R244 in attempt to increase the catalytic efficiency of FgaPT2 and alter substrate specificity.[86, 87] FgaPT2 K174F had almost no tryptophan activity but efficiently C3 prenylated L-tyrosine.[79] The related double mutants K174F/R244X (X= L,N,Q,Y) were able to reverse prenylate L-Trp cyclic dipeptides at the C3 position, resulting in cyclization (Figure 15).[87] FgaPT2 mutants R244N, R224L, R224Y and R244Q had substantially higher activity on a variety of L-Trp cyclic dipeptides while maintaining C4 prenylation selectivity.[86] The R244L mutant had the best overall substrate tolerance, converting 20-50% of most dipeptides screened. R244Y was the only mutant particularly good at prenylating cyclo-L-Trp-L-Phe.[86] FgaPT2 mutants K187E, K259E, and R257G all had >95% reduced activity because these residues are known to interact with the diphosphate group.[88]



**Figure 15. FgaPT2 mutants reactions.**

*FgaPT2 mutants have been engineered to alter the regioselectivity and substrate preference[87]*

### Research Objectives

Studies thus far have revealed that FgaPT2 has a broad and diverse acceptor substrate scope including, substituted tryptophans, L-Trp containing cyclic dipeptides, isomers of L-tyrosine, indolocarbazoles, as well as phenolic molecules.[78, 89-94] A few alternative alkyl donors have also been reported with FgaPT2 including the C<sub>10</sub> geranyl pyrophosphate (GPP), 2-butenyl pyrophosphate, 2-pentenyl pyrophosphate, and benzyl pyrophosphate.[60, 68] Based on our recent studies using tyrosine-*O*-prenyltransferase, SirD that demonstrated relaxed alkyl-donor substrate specificity, [95] in this chapter we describe the alkyl-donor substrate scope of FgaPT2 using 34 synthetic allylic, and benzylic alkyl donors with varying carbon chain length, and substituents. The alkyl donor analogues include 21 allylic analogues described in Chapter 2, and 13 additional benzylic analogues. For the sake of the flow of the manuscript (submitted to MedChemComm) all analogues have been renumbered as described in Figure 16. The study involves assessment of FgaPT2 catalyzed alkylation reaction using RP-HPLC and

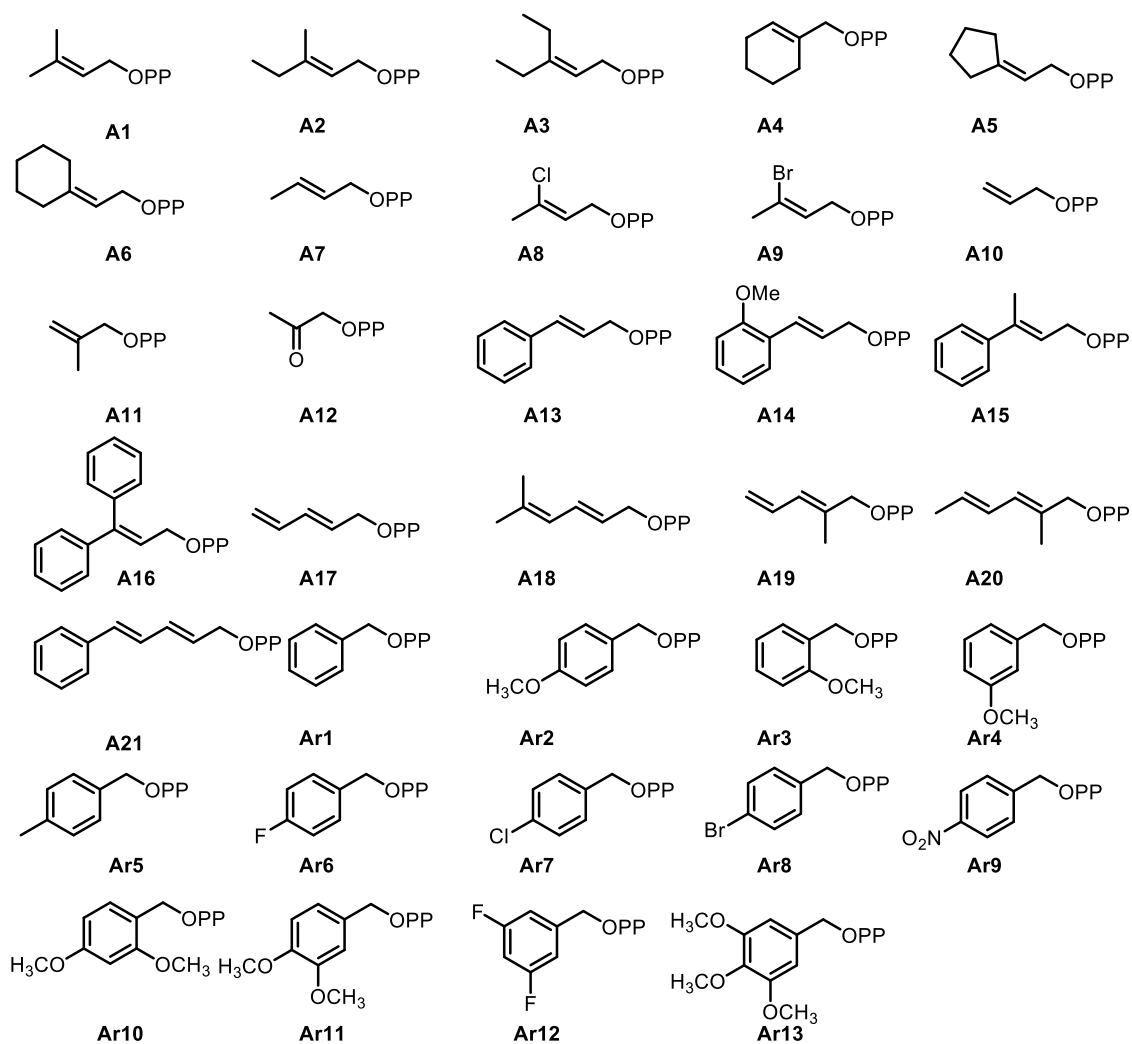
HRMS studies, systematic analysis of kinetic parameters and identification of the regio-specificity of the alkylated products using NMR spectroscopy.

### **Contribution**

The library of alkyl-donors (Figure 16) was synthesized by Dr. Chandrashekar Bandari, Johanna Masterson and Rachel Tran. The large-scale tryptophan alkylation reaction and HRMS analysis was carried out by Dr. Chandrashekar Bandari, NMR characterization of alkyl-tryptophan products was performed by Dr. Bandari and Dr. Susan Nimmo. My contributions involve expression and purification of FgaPT2, RP-HPLC based assessment of alkyl-donor specificity of FgaPT2, and determination of kinetic parameters for FgaPT2 catalyzed reaction with diverse alkyl-donors. Most of the contents including text, tables and figures for this chapter were directly taken from the manuscript submitted for publication to MedChemComm.

### **Results and Discussion**

In order to assess the ability of FgaPT2 to accept diverse unnatural alkyl-PP analogues, the recombinant FgaPT2 was overproduced in *Escherichia coli* BL21(DE3) cells transformed with codon-optimized synthetic *FgaPT2* gene in pET28a vector. The resulting FgaPT2 with *N*-terminal His<sub>6</sub>-fusion protein was purified to homogeneity via Ni-NTA affinity chromatography as described previously,[40] yielding ~40 mg L<sup>-1</sup> of pure *N*-terminal His<sub>6</sub>-fusion protein.

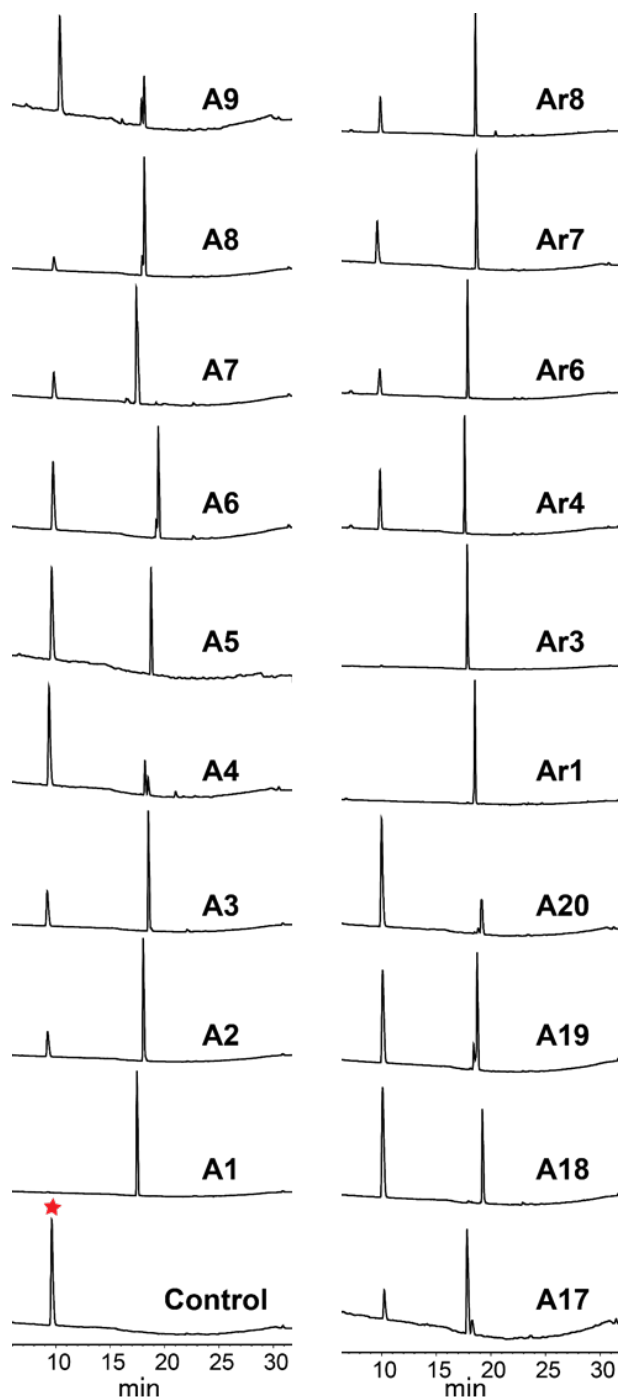


**Figure 16. Synthetic alkyl-PP analogues used in this study.**

### RP-HPLC and HRMS Studies

Standard uniform assay conditions (1.2 mM alkyl-PP analog, 1 mM L-Trp, 5  $\mu$ M FgaPT2, 25 mM Tris, 5 mM MgCl<sub>2</sub>, 50 mM KCl, pH 7.5, 16 h at 35 °C) were adopted to facilitate the initial assessment of donor substrate specificity of the FgaPT2 with synthetic alkyl-PP analogues. Production of alkyl-L-Trp derivatives was determined by the difference in the retention time of L-Trp and alkyl-L-Trp product using a reverse phase high-pressure liquid chromatography (RP-HPLC) endpoint assay (Figure 17) with

subsequent confirmation by HRMS coupled with liquid chromatography (LC) analysis for all positive reactions (Table 3) to afford complete analysis of the desired product.



**Figure 17. RP-HPLC chromatograms of FgaPT2 catalyzed reactions.**

*FgaPT2 was incubated with L-Trp and alkyl-PP analogues that led to the formation of corresponding alkyl-L-Trp analogues. The peak at retention time 10 min is L-Trp and the product peaks are after 15 min in the chromatograms respectively.*

**Table 3.** Summary of HRMS data of alkyl-Trp analogues.

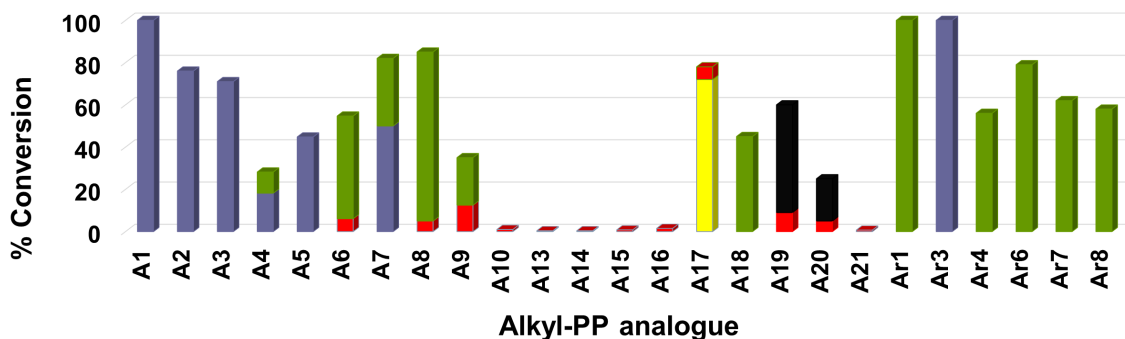
*FgaPT2 catalyzed alkylation reaction with L-Trp and synthetic alkyl-PP analogues. The subscript '2' represents the second product based on HPLC retention time.*

Enzyme Product	Chemical Formula	Calculated Mass (Da)	Observed Mass (Da)
A1-Trp	C <sub>16</sub> H <sub>21</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	273.1603	273.1602
A2-Trp	C <sub>17</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	287.1760	287.1754
A3-Trp	C <sub>18</sub> H <sub>25</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	301.1916	301.1916
A4-Trp	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1760	299.1756
A4-Trp <sub>2</sub>	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1760	299.1761
A5-Trp	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1760	299.1768
A6-Trp	C <sub>19</sub> H <sub>25</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	313.1916	313.1913
A6-Trp <sub>2</sub>	C <sub>19</sub> H <sub>25</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	313.1916	313.1911
A7-Trp	C <sub>15</sub> H <sub>19</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	259.1447	259.1449
A7-Trp <sub>2</sub>	C <sub>15</sub> H <sub>19</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	259.1447	259.1444
A8-Trp	C <sub>15</sub> H <sub>18</sub> ClN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	293.1057	293.1061
A8-Trp <sub>2</sub>	C <sub>15</sub> H <sub>18</sub> ClN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	293.1057	293.1062
A9-Trp	C <sub>15</sub> H <sub>18</sub> BrN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	337.0552	337.0552
A9-Trp <sub>2</sub>	C <sub>15</sub> H <sub>18</sub> BrN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	337.0552	337.0547
A10-Trp	C <sub>14</sub> H <sub>17</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	245.1290	245.1294
A13-Trp	C <sub>20</sub> H <sub>21</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	321.1603	321.1601
A14-Trp	C <sub>21</sub> H <sub>23</sub> N <sub>2</sub> O <sub>3</sub> [M+H] <sup>+</sup>	351.1709	351.1711
A15-Trp	C <sub>21</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	335.1760	335.1757
A16-Trp	C <sub>26</sub> H <sub>25</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	397.1916	397.1912
A17-Trp	C <sub>16</sub> H <sub>19</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	271.1447	271.1442
A17-Trp <sub>2</sub>	C <sub>16</sub> H <sub>19</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	271.1447	271.1444
A18-Trp	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1760	299.1768
A19-Trp	C <sub>17</sub> H <sub>21</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	285.1603	285.1602
A19-Trp <sub>2</sub>	C <sub>17</sub> H <sub>21</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	285.1603	285.1605
A20-Trp	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1715	299.1751
A20-Trp <sub>2</sub>	C <sub>18</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	299.1715	299.1749
A21-Trp	C <sub>22</sub> H <sub>23</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	347.1760	347.1767
Ar1-Trp	C <sub>18</sub> H <sub>19</sub> N <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	295.1447	295.1430
Ar3-Trp	C <sub>19</sub> H <sub>21</sub> N <sub>2</sub> O <sub>3</sub> [M+H] <sup>+</sup>	325.1552	325.1547
Ar4-Trp	C <sub>19</sub> H <sub>21</sub> N <sub>2</sub> O <sub>3</sub> [M+H] <sup>+</sup>	325.1552	325.1540
Ar6-Trp	C <sub>18</sub> H <sub>18</sub> FN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	313.1352	313.1338
Ar7-Trp	C <sub>18</sub> H <sub>18</sub> ClN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	329.1057	329.1039
Ar8-Trp	C <sub>18</sub> H <sub>18</sub> BrN <sub>2</sub> O <sub>2</sub> [M+H] <sup>+</sup>	373.0552	373.0531

The initial assessment of FgaPT2 catalyzed alkylation of L-Trp with 34 synthetic alkyl-PP analogues based upon RP-HPLC of analytical scale reaction is illustrated in Figure



17, wherein 25 of the 34 synthetic analogues (Figure 16) in FgaPT2 catalyzed reaction produced corresponding alkyl-L-Trp. All products were subsequently confirmed to be mono-alkylated products from HRMS coupled with liquid chromatography (LC) analysis (Table 3).



**Figure 18: The percent conversion of alkyl-PP analogues.**

*Percent conversion of alkyl-L-Trp catalyzed by FgaPT2 based upon RP-HPLC. Reactions were performed with 1.2 mM alkyl-PP analog, 1 mM L-Trp, 5  $\mu$ M FgaPT2, 25 mM Tris, 5 mM MgCl<sub>2</sub>, 50 mM KCl, pH 7.5, 16 h at 35 °C. No product formation was observed in the absence of FgaPT2 or alkyl-PP analog. The blue, green, yellow, and black colors in the bar diagram denote C4, C5, C3, and N1 alkyl-L-Trp regioisomers as confirmed by NMR, whereas the red color represents alkyl-L-Trp isomer with undetermined regio-chemistry.*

Of the 25 positive reactions, 12 led to appreciable (>50%) conversion, an additional seven led to moderate (20-50%) conversion, while six offered detectable product formation (<5%) under the conditions described (Figures 19 and 20). Analysis of HPLC and HRMS results suggested more than one product for eight (**Figure 18**, A4, A6–A9, A17, A19, A20) of the 19 analytical scale reactions with product yield > 20% (Figure 16, A1–A9, A17–A20, Ar1, Ar3, Ar4, Ar6–A8). Notably, the 3-carbon allyl pyrophosphate A10 was accepted as a substrate although with a very low conversion rate of 1%, resulting in the formation of allyl-L-Trp. However, A11 with a methyl

group substituted at the  $\beta$ -carbon of allyl pyrophosphate was not accepted by FgaPT2. No product formation was observed for **A12** that has a carbonyl group at  $\beta$ -carbon. The bulky cinnamyl analog (**A13**) and its derivatives (**A14**, **A15**, **A16**, and **A21**) produced only a trace amount of the alkylated product with low yield (< 2%). In addition, while *o*-methoxy (**Ar3**), *m*-methoxy (**Ar4**) and *p*-halogen (**Ar6-Ar8**) substituted benzyl analogues were accepted by FgaPT2, introduction of bulky groups at the para-position, including *p*-methoxy (**Ar2**), *p*-methyl (**Ar5**), *p*-nitro (**Ar9**) and di- and tri-substituted benzyl analogues (**Ar10 – Ar12**, and **A13**) did not serve as substrates for FgaPT2 catalyzed reaction with L-Trp due to the potential steric hindrance in the active site of the enzyme. These results suggest the FgaPT2 catalyzed reaction with L-Trp is sterically limited by bulky substitution on the alkyl-donor.

### **Kinetic studies**

Michaelis–Menten kinetic parameters for all 19 alkyl-PP analogues with product conversion > 20%, (Table 4) and for L-Trp with representative alkyl-PP analogues were carried out. The kinetic parameters, rates ( $k_{cat}$ ) and Michaelis constants ( $K_M$ ) were determined from a nonlinear regression fit of initial velocities versus concentration of alkyl-PP analog by GraphPad Prism 5.04 (Figure 19). Interestingly, the  $K_M$  of L-Trp for FgaPT2 with various alkyl-PP donors did not change drastically by the nature of the alkyl donors and remained within the range of  $0.20 \pm 0.08$  mM. Consistent with our previous observations with tyrosine-*O*-prenyltransferase SirD,[95] the alkyl-PP analogues that contained groups to enhance the stability of the carbocation via increased conjugation and resonance stabilization correlated with the increased turnover of the product. For example, the  $k_{cat}$  values of the reaction correlated with conjugation by a

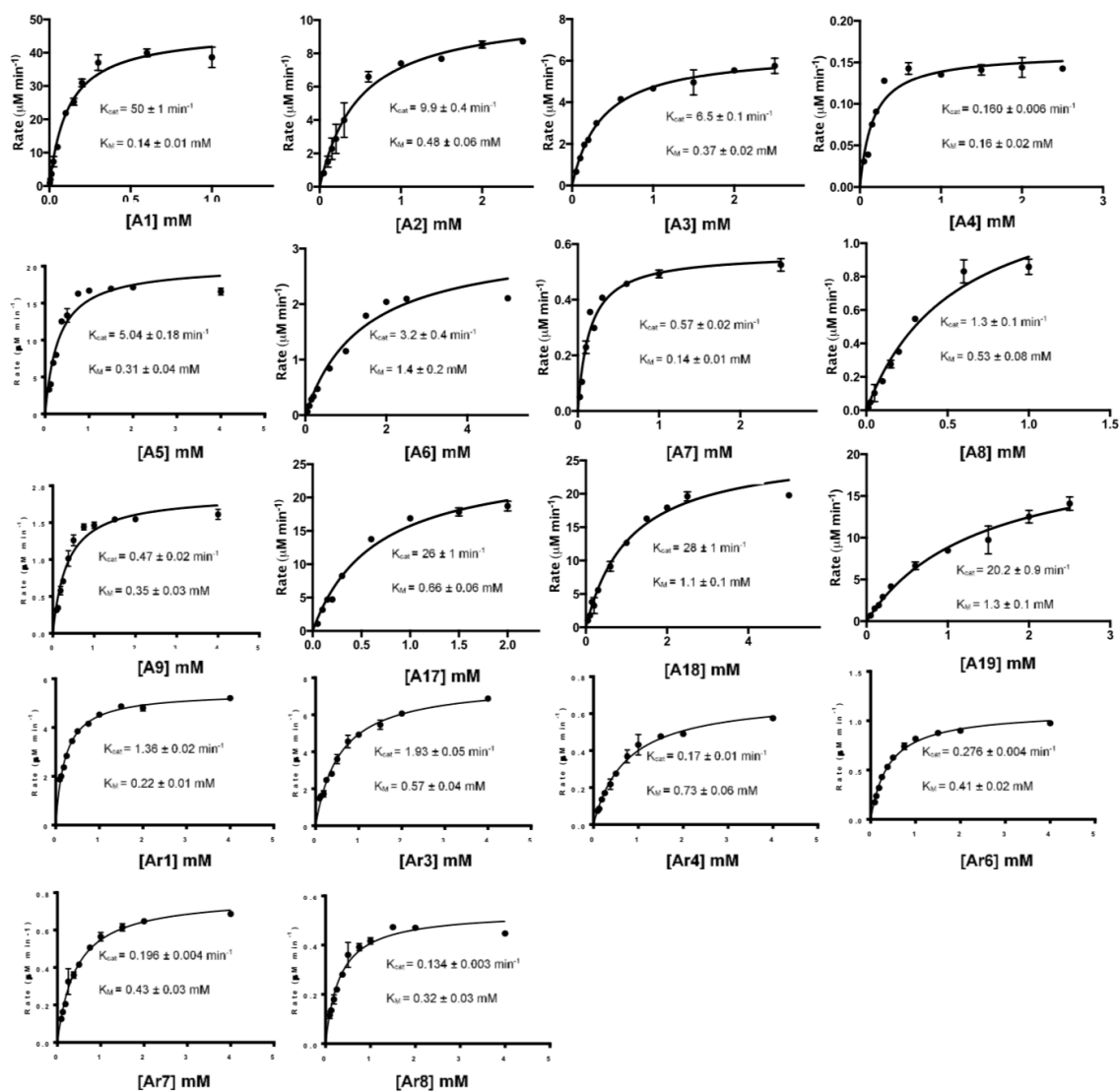
methyl group in the order, **A1** > **A2** > **A3** > **A7** (Table 4). In addition, the  $K_M$  values increased with the increased length of the carbon chain in the order **A1** < **A2**  $\cong$  **A3**  $\cong$  **A5** < **A6** and **A17** < **A18**  $\cong$  **A19**  $\cong$  **A20**. Removing one methyl group of **A1** decreased the efficiency ( $k_{cat}/K_M$ ) of **A7** by 88-fold compared to **A1**.

**Table 4. Kinetic parameters for FgaPT2 with L-Trp.**

*1.2 mM L-Trp, and varied concentration of alkyl-PP analogues in 25 mM Tris, 5 mM MgCl<sub>2</sub>, 50 mM KCl, pH 7.5, 35 °C. (\*Relative catalytic efficiency is defined as the percent  $k_{cat}/K_M$  when compared to **A1**)*

Alkyl-PP analog	$K_{cat}$ (min <sup>-1</sup> )	$K_M$ (mM)	$k_{cat} / K_M$ (mM <sup>-1</sup> min <sup>-1</sup> )	Relative catalytic efficiency*
<b>A1</b>	50 ± 1	0.14 ± 0.01	357	100
<b>A2</b>	9.9 ± 0.4	0.48 ± 0.06	21	5.88
<b>A3</b>	6.5 ± 0.1	0.37 ± 0.02	18	5.04
<b>A4</b>	0.160 ± 0.006	0.16 ± 0.02	1	0.28
<b>A5</b>	5.04 ± 0.18	0.31 ± 0.03	16	4.48
<b>A6</b>	3.2 ± 0.4	1.4 ± 0.2	2.28	0.64
<b>A7</b>	0.57 ± 0.02	0.14 ± 0.01	4.07	1.14
<b>A8</b>	1.3 ± 0.1	0.53 ± 0.08	2.5	0.70
<b>A9</b>	0.47 ± 0.01	0.35 ± 0.03	1.3	0.38
<b>A17</b>	26 ± 1	0.66 ± 0.06	39	10.92
<b>A18</b>	28 ± 1	1.1 ± 0.1	25.5	7.14
<b>A19</b>	20.2 ± 0.9	1.3 ± 0.1	15.5	4.34
<b>A20</b>	16.9 ± 2	1.39 ± 0.2	12.1	3.39
<b>Ar1</b>	1.36 ± 0.02	0.22 ± 0.01	6.2	1.74
<b>Ar3</b>	1.93 ± 0.05	0.57 ± 0.04	3.4	0.95
<b>Ar4</b>	0.170 ± 0.005	0.73 ± 0.06	0.23	0.064

Ar6	$0.276 \pm 0.004$	$0.4 \pm 0.02$	0.69	0.19
Ar7	$0.196 \pm 0.004$	$0.43 \pm 0.03$	0.46	0.13
Ar8	$0.134 \pm 0.003$	$0.32 \pm 0.03$	0.42	0.12



**Figure 19. Michaelis-Menten inetic plots for FgaPT2 reactions**

*Kinetic parameters detemered by Michaelis-Menten plots with 1mM tryptophan and varying concentrations of pyrophosphate analogues incubated at 35 °C*

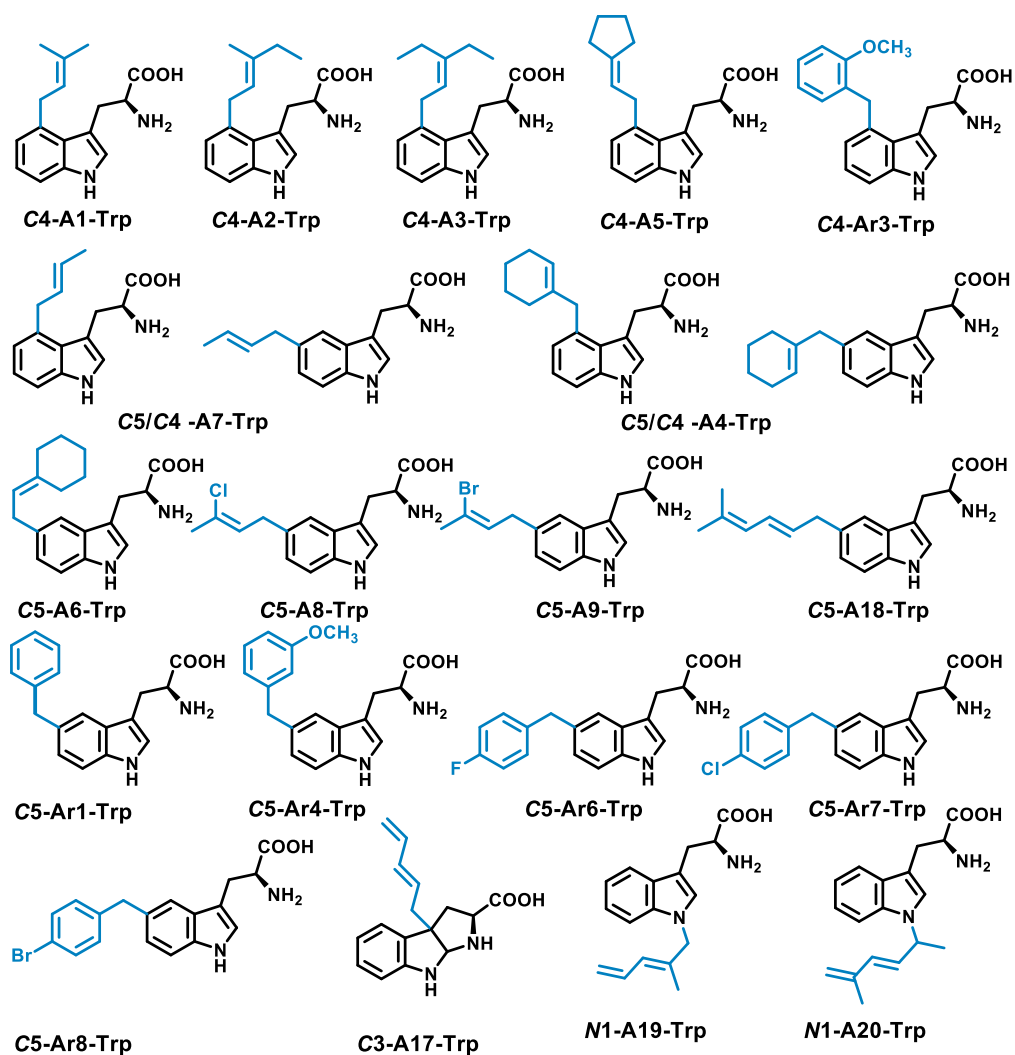
Further, changing one or both methyl groups of **A1** to an ethyl group (**A2** and **A3**) lowered the enzyme efficiency 17-20 times compared to **A1**. Similarly, the  $k_{cat}$  values of the reaction correlated with resonance stabilization in the order **A17** > **A7**; **Ar1** > **A4**; **Ar1** > **Ar6** > **Ar7** > **Ar8**. Furthermore, a 9-fold higher  $k_{cat}$  value of **Ar1** compared to **A4** suggests resonance stabilization of the carbocation by the benzylic group contributes to the rate of the reaction. However, the  $k_{cat}$  value of the cyclohexyl (**A4**) and benzyl analogues (**Ar1**) were nearly 310 and 36, fold lower than **A1**. When an electron withdrawing group such as chloro (**A8**) or bromo (**A9**) replaced one of the methyl groups of **A1**, the efficiency of the enzyme was lowered nearly 150–250 fold compared to the natural alkyl donor, **A1**, which explains the importance of methyl group in stabilizing the carbocation. Surprisingly, the dienyl analogues (**A17**, **A18**, **A19**, and **A20**) displayed relatively higher rates with just ~2 fold lower  $k_{cat}$  values compared to the natural alkyl donor, **A1**, implying extended resonance stabilization of the carbocation favors the FgaPT2 catalyzed reaction. However, the  $K_M$  values of the dienyl analogues were 5-10-fold higher than the natural alkyl donor, **A1** and the trend of  $K_M$  values correlated with the increasing length of the carbon chain (C5–C7), in the order, **A17** < **A19** < **A20**, again pointing at the steric factors and the orientation of the alkyl group in the active site of the enzyme to contribute to the enzyme efficiency. All mono-substituted benzylic analogues (**Ar3**, **Ar4**, **Ar6**, **Ar7**, **Ar8**) displayed a similar  $K_M$  value, which was 2–3 fold higher than the benzyl analog, **Ar1**. The 10-fold higher  $k_{cat}$  value of *o*-methoxy (**Ar3**) compared to the *m*-methoxy (**Ar4**), suggests benzylic cation stabilization by an *o*-methoxy group increases the efficiency of the FgaPT2 catalyzed alkylation reaction. Among the para-substituted halogens, the  $k_{cat}$  values followed the

order *p*-fluoro (**Ar6**) > *p*-chloro (**Ar7**)  $\cong$  *p*-bromo (**Ar8**), suggesting a combination of steric factors and possible stabilization of carbocation of the **Ar6** carbocation via back donation of the non-bonded electron pairs of the fluorine atom to the **Ar6** carbocation through effective 2p-2p orbital interaction. The observed  $k_{cat}$  values of para-substituted halogens were ~5-10 fold lower than **Ar1**, which is consistent with the inductive effect of the halogens. Surprisingly, while the para-substituted halogens were accepted by FgaPT2, the *p*-methoxy and *p*-methyl were not substrates for FgaPT2 catalyzed reaction even with their ability to contribute to the stabilization of the benzylic cation, likely due to the steric factors in the active site of the enzyme. These results emphasize the fact that the FgaPT2 catalyzed Friedel-Craft alkylation reaction proceeds via stabilized carbocation,[60] and the steric factor and the orientation of alkyl group in the active site has a major role in the ability of FgaPT2 to utilize alkyl donors.

### **NMR Studies**

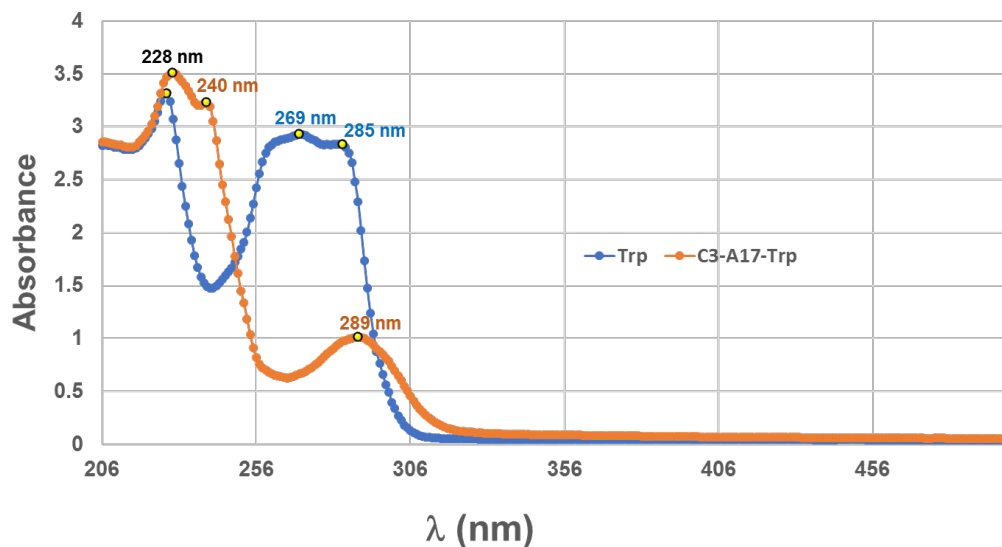
To assess the regio-specificity of FgaPT2 catalyzed alkyl-transfer, the 19 analytical scale reactions with an alkyl-tryptophan yield > 20% (reaction involving **A1-A9**, **A17-A20**, **Ar1**, **Ar3** **Ar4**, **Ar6-Ar8**, **Figure 16**) were scaled up, purified via RP-HPLC, and characterized via <sup>1</sup>H and 2D NMR experiments by Dr. Chandrashekar Bandari. While eight of the 19 alkyl-transfer reactions involving alkyl donors **A4**, **A6**, **A7**, **A8**, **A9**, **A17**, **A19** and **A20** yielded two or more mono-alkylated products based on RP-HPLC and HRMS analysis, regio-specific alkyl-group attachment of minor products of reactions involving **A6**, **A8**, **A9**, **A17**, **A19**, and **A20** could not be obtained due to insufficient intensity of NMR signals. The regiospecificity of 21 alkyl-tryptophan analogues was determined using detailed 2D NMR analysis (Figure 20). Among the 21

alkyl-tryptophan analogues characterized by NMR, the alkyl group of seven were at C4 (derived from **A1-A5**, **A7**, and **Ar3**), 11 at C5 (derived from **A4**, **A6**, **A7**, **A8**, **A9**, **A18**, **Ar1**, **Ar4**, **Ar6-Ar8**), one at C3 (derived from **A17**) forming hexahydro-pyrroloindoline product as verified by UV (Figure 21), and two were at the N1 (derived from **A19**, **A20**) position of tryptophan respectively (Figure 20).



**Figure 20. NMR characterized structures**

*Structure of isolated alkyl-tryptophan products obtained from FgaPT2 reaction with L-Trp and alkyl-PP analogues determined by NMR analysis.*

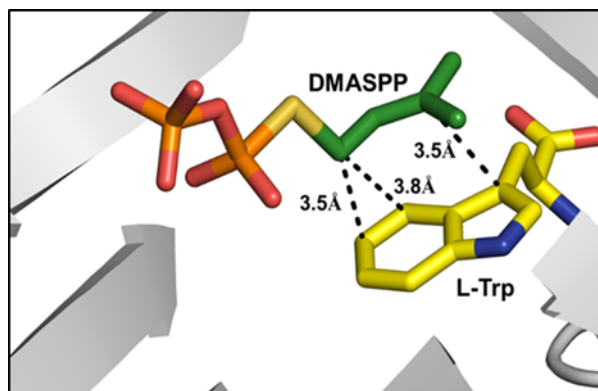


**Figure 21: UV absorption spectra of L-Trp and C3-A17-Trp.**  
*Showing signature absorption maxima for hexahydro-pyrroloindoline tryptophan product.*

Intriguingly, apart from the strict C4 alkylation of tryptophan by FgaPT2 using the natural alkyl donor **A1**, the unnatural alkyl donors, **A2**, **A3** and **A5**, which contained an ethyl group (**A2** and **A3**) in place of one or both methyl group/s of **A1**, or the similarly sized conformationally restricted cyclopentyl analog (**A5**) selectively added the alkyl group at the C4 position of tryptophan. Alkyl-PP analogues with smaller alkyl chain in the linear direction compared to **A1** (**A4** and **A7**) resulted in a mixture of C4 and C5 regio-isomers, and alkyl-PP analogues with a methyl group of **A1** replaced with a halogen (**A8** and **A9**) or larger alkyl chain (**A6** and **A18**) predominantly produced the C5 alkylated product. While most of the unnatural benzylic analogues (**Ar1**, **Ar4**, **Ar6**, **Ar7**, and **Ar8**) produced selectively C5 alkylation of tryptophan, the *o*-methoxy benzyl analog (**Ar3**) produced a single C4-alkylated tryptophan. Careful analysis of the C4 alkylating analogues (**A1**, **A2**, **A3**, **A5**, and **Ar3**) suggests that C4 substitution is



preferred when the carbon chain length is less than 6 while C3' is also disubstituted to facilitate stacking interaction with the indole ring. Therefore, the active site steric factors in combination with improper stacking interaction of the alkyl group with the indole ring conceivably displaces the carbocation resulting in a direct addition of alkyl group to C5 of tryptophan as seen for 11 of the C5-alkyl-Trp products. The addition of alkyl groups at the C5 position of tryptophan by most of the alkyl-PP analogues is consistent with the orientation of DMASPP with respect to the indole ring in the active site of the FgaPT2 crystal structure,[51] where C5 of tryptophan is just at 3.5 Å from the C1' of the alkyl group compared to a distance of 3.8 Å between C4 of tryptophan to the C1' of the alkyl group (Figure 22).



**Figure 22: The active site of FgaPT2 structure (PDB:3I4X)**

*FgaPT2 in complex with L-Trp and DMASPP displaying distances between C1' of DMASPP and C4 and C5 of L-Trp; and C3' of DMASPP and C3 of L-Trp.*

## Conclusions

In conclusion, this work has revealed that FgaPT2 has a remarkable ability to accept diverse allylic and benzylic unnatural alkyl-donor substrates. While FgaPT2 can transfer alkyl groups at C4, C5, N1 and C3 position of L-Trp, the C5 position is the

preferred alkylation position for FgaPT2 with unnatural alkyl-donors. Thus, these studies highlight the potential utility of FgaPT2 for broader applications such as alkyl-diversification of indole-containing natural products for drug discovery and other synthetic applications. Future enzyme mutational studies along with substrate engineering is expected to drive the development of more efficient and selective FgaPT2-based biocatalysts.

## **Experimental**

### **General materials**

Unless otherwise stated, all chemicals and reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA), Acros (New Jersey, USA), Alfa-Aesar (Ward Hill, MA, USA) or TCI (Portland, OR, USA) and were reagent grade or better. PD-10 column and Ni-NTA superflow columns were purchased from GE Healthcare (Piscataway, NJ).

### **General methods**

HPLC was accomplished using Agilent 1220 system equipped with a DAD detector. High-resolution mass spectrometric (HRMS) data and liquid chromatography mass spectrometric (LCMS) were obtained on Agilent 6545-QTOF W/1290 HPLC mass spectrometer at the University of Oklahoma, Department of Chemistry and Biochemistry. NMR spectra were obtained on Varian VNMRS 400 or 500 MHz instruments at the NMR facility of the Department of Chemistry and Biochemistry at the University of Oklahoma using 99.9% DMSO-d<sub>6</sub> with 0.05% v/v TMS, or 99.9% D<sub>2</sub>O or 99.8% MeOH-d<sub>4</sub>, 99.9% Acetone-d<sub>6</sub> (Cambridge Isotope Laboratories, MA, USA).

### ***In-vitro* FgaPT2 assay**

The recombinant FgaPT2 was overproduced in *Escherichia coli* BL21(DE3) cells transformed with codon-optimized synthetic *FgaPT2* gene in pET28a vector. The resulting FgaPT2 with *N*-terminal His<sub>6</sub>-fusion protein was purified to homogeneity via Ni-NTA affinity chromatography as described previously.[40] *In vitro* FgaPT2 reactions were conducted in a volume of 20 µl with 1.2 mM alkyl-PP analog (**A1 – A21**, **Ar1 – Ar13**), 1 mM L-Trp and 5 µM purified FgaPT2 in 25 mM Tris buffer pH 7.5, 5 mM MgCl<sub>2</sub>, 50 mM KCl, incubated at 35 °C for 16 h. Reactions were quenched by adding an equal volume of methanol followed by centrifugation (10,000 *xg* for 15 min) to remove the precipitated protein and product formation for each reaction was subsequently analyzed by RP-HPLC using **Method A** or **Method C**. For each reaction, percent yield was based upon the integration of species at 254 nm and calculated by dividing the integrated area of the product by the sum of the integrated area of product and/or and the remaining substrate. All putative products were subsequently confirmed by HRMS with positive (+) and/or negative (-) mode. Kinetic parameters were assessed using 1.2 mM L-Trp and variable alkyl-PP analogues (0.1 - 6 mM). Assays were conducted in triplicate and all rates were confirmed to be linear. The kinetic curves were fit to Michaelis-Menton kinetics using Prism 5.04 (GraphPad Software, Inc. La Jolla, CA 92037 USA). The reactions with > 20% product yield were performed in 2-5 mL volume under standard condition, purified by semi-prep HPLC using **Method B** and the putative new products were confirmed by NMR. **Method A:** To monitor FgaPT2 reactions of L-Tryptophan, analytical reverse-phase (RP) HPLC employed a Gemini-NX, C-18 (5 µm, 4.6 mm × 250 mm) column (Phenomenex, Torrance,

California, USA) [gradient of 10% B to 25% B over 10 min, 25% B to 100% B over 15 min, 100% B for 3 min, 100% B to 10% B over 0.1 min, 10% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 1 mL min<sup>-1</sup>; A<sub>254</sub>]. The reaction was monitored by the retention time difference between starting material and product.

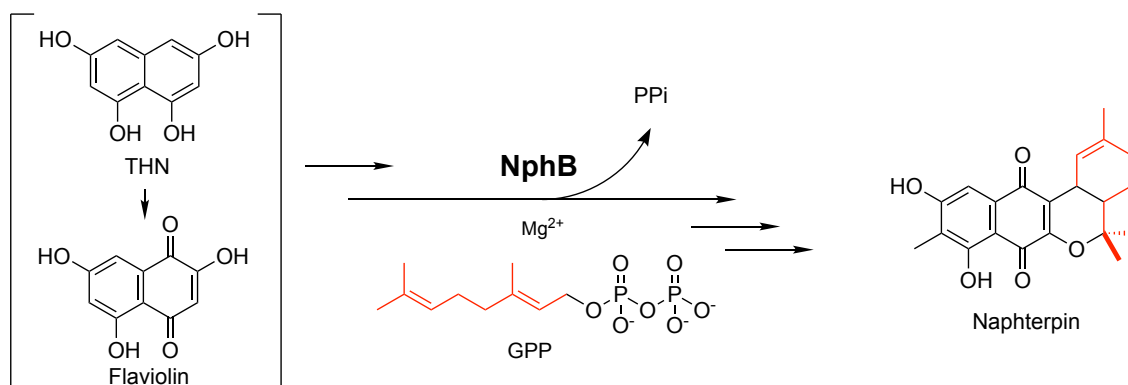
**Method B** :Semi-preparative RP HPLC was conducted on a Gemini-NX, C-18 (5 μm, 10 mm × 250 mm) column (Phenomenex, Torrance, California, USA) to purify the L-Trp analogues [gradient of 10% B to 25% B over 10 min, 25% B to 100% B over 15 min, 100% B for 3 min, 100% B to 10% B over 0.1 min, 10% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 2 mL min<sup>-1</sup>; A<sub>254</sub>].

## Chapter 4. Alkyl donor specificity of a Naphterpin prenyltransferase,

### NphB

#### Introduction

NphB, previously referred to as *orf2*, is a soluble aromatic PT with significant  $Mg^{2+}$  dependent activity found in the Naphterpin biosynthetic pathway.[96] Naphterpin is a powerful antioxidant produced in the microorganism *Streptomyces* sp CL190. The core is comprised of a naphthoquinone moiety with an attached 10-carbon geranyl group forming a cyclohexene ring with a geminal dimethyl group.[97] Briefly, the biosynthesis begins with the polyketide 1,3,6,8-tetrahydroxynaphthalene (THN), formed from the malonyl-CoA by a type III polyketide synthase, which undergoes an oxidative transformation to flaviolin. The resulting THN, flaviolin, or THN-derived metabolite are tailored by addition of a geranyl co-substrate and further modified by subsequent cyclization. The gene cluster responsible for naphterpin production revealed an open reading frame (*orf2*) with a high degree of homology to the characterized PTs CloQ and NovQ, leading to the proposal that *orf2* encodes for the PTase responsible for the transfer of the geranyl group on THN or THN derivative (Figure 23). [36] Although the precise native substrate of NphB currently remains ambiguous, in common practice 1,6 dihydroxynaphthalene is used as a surrogate acceptor substrate.



**Figure 23. NphB catalyzed step in the biosynthesis of naphtherpin[36]**

As mentioned, the true biological substrate has not been identified; however, various aromatic acceptors have been tested revealing a highly promiscuous nature towards aromatic substrates. Initially, enzyme activity was assessed by incubating with prenyl donors, DMAPP, GPP, or FPP, in the presence of aromatic acceptors; further, this was later tested with an azido-substituted GPP.[36, 54] For the isoprenoid pyrophosphate substrates, NphB did not display any enzymatic activity with DMAPP, as isoprenoid pyrophosphate substrate. There was detectable activity with FPP and azido-GPP and the highest activity with GPP.[36, 98]

NphB has been assayed for utilization of various aromatic substrates including flavonoids, isoflavonoids, plant polyketides and 1,3,6,8-tetrahydroxynaphthalene (THN) analogues. The enzyme is capable of GPP additions on several positions of the aromatic ring including C- and O- geranylations yielding several different alkylated products. A comparison of the reaction products showed that most contain a geranyl group in the ortho position relative to the hydroxyl group.[54, 99] With the flavonoids, NphB performed an O-prenylation of a hydroxyl group as well as the C6 position of

naringenin and apigenin and the C8 position of daidzein[36, 54]. This demonstrates that, consistent with other PTs, NphB catalyzes donor-dependent prenylation reactions. The structure of NphB consists of a single domain PT- fold with some variability in the overall  $\alpha\alpha\beta\beta$  repeating motif resulting in missing the  $\alpha 9$  helix between strands  $\beta 8$  and  $\beta 9$ . Additionally, NphB has three helices outside of the PT-fold. The first helix,  $3_{10}$ , is found between  $\beta 1$ -  $\beta 2$  strands and positioned above the metal binding site within the barrel. The second additional helix follows the end of strand  $\beta 4$  and partially overlaps with the opening at the C-terminal end of the  $\beta$ -barrel. NphB also has an extra C-terminal  $\alpha$ -helices, that fully extends across the aromatic binding domain opening of the barrel. The interior of the barrel contains a number of hydrophobic residues used to sequester the geranyl tail of GPP, with the head groups positioned towards the polar end of the barrel thus allowing contact with the coordinated  $Mg^{2+}$  cation. Notably, the sequence does not contain the (N/D)DXXD recognition motif found in most  $Mg^{2+}$  dependent isoprenoid diphosphate enzymes.

### **Research Objective**

The aim of this study is to define the alkyl pyrophosphate donor scope of the polyketide prenyltransferase NphB, using an extensive synthetic library of 65 pyrophosphate analogues, in the presence of the native substrate 1,6-DHN or of the drug compound sulfabenzamide. The library includes 44 new synthetic analogues added from the work reported in Chapter 3, and all analogues have been renumbered as described in Figure 24. In this chapter we report an extensive alkyl pyrophosphate donor scope of polyketide prenyltransferase NphB with 1,6-DHN and sulfabenzamide, and structure-

based engineering of NphB. Interestingly, a comparison of the *in vitro* activity assays with 1,6-DHN or with sulfabenzamide indicated a strong donor dependent alkylation reaction. In the presence of 1,6-DHN or sulfabenzamide, NphB was able to catalytically transfer 26 and 29 out of the 65 diverse pyrophosphate donors, respectively. Through the structural elucidation of the alkyl-diversified sulfabenzamide compounds, we observed a previously unreported functionality of NphB to perform regio specific N-alkylations. Moreover, we accomplished structure-based rational engineering of NphB to increase substrate acceptance capabilities, producing a gernalyated sulfabenzamide derivative.

### **Contributions**

All alkyl pyrophosphate analogues were synthesized by Dr. Chandrasekhar Bandari, Tejaswi Bavineni, Johanna Masterson, Rachel Tran, and Eric Gardener. Large-scale enzymatic reaction and subsequent purification was carried out in part by me and Andrea Batchev. Structural characterization of sulfabenzamide derivatives by NMR spectroscopy was performed in part by me, Dr. Susan Nimmo and Andrea Batchev. My contributions also involved expression and purification of NphB, RP-HPLC and HRMS based assessment of alkyl-donor specificity of NphB, and determination of kinetic parameters for NphB catalyzed reaction with diverse alkyl-donors. Some contents from this chapter including text, tables and figures will be directly used for a publication in a peer-reviewed journal.



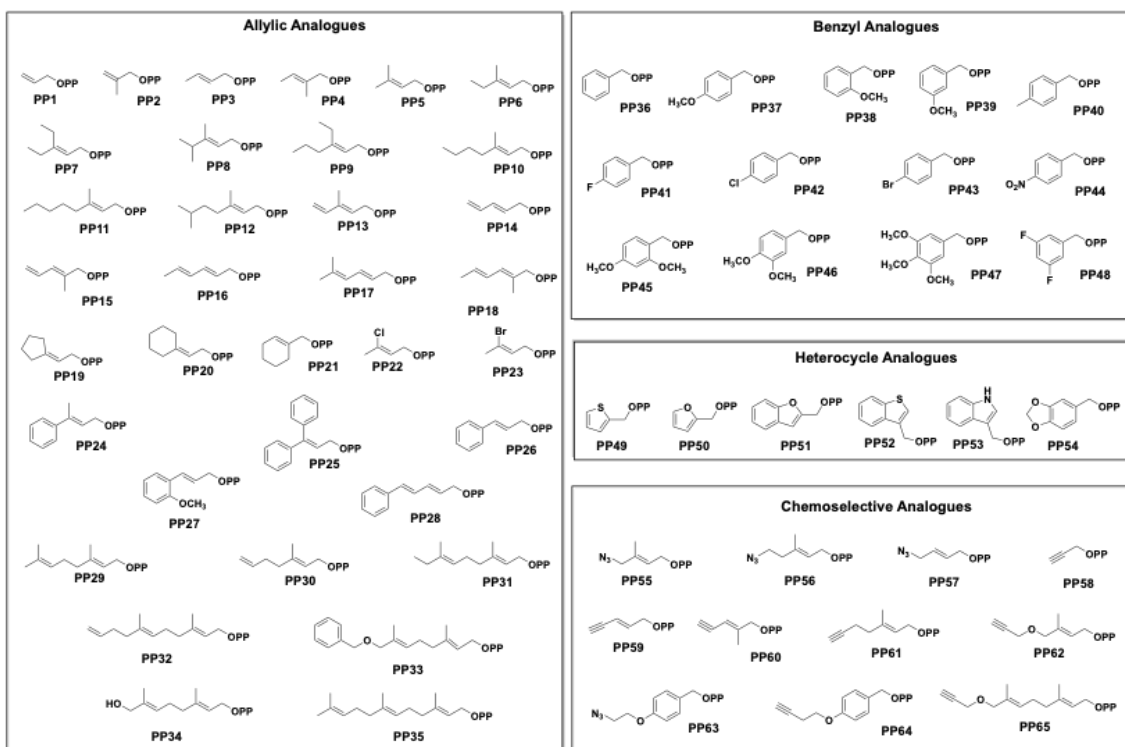
## Results

### Description of the alkyl pyrophosphate library.

In this study, we investigated the NphB catalyzed alkylation reaction of 1,6-DHN and sulfabenzamide, an antimicrobial compound with promising anti-tumor/ anti-cancer properties[96], against an internally-generated library of synthesized pyrophosphates. The library contains 65 electronically and structurally diverse compounds, designed to probe the donor permissiveness of the enzyme as well as potentially generating compounds with increased bioactivity (Figure 24). We have organized the compounds according to their chemical space and potential functionality, including; allylic-, benzylic- and heterocyclic analogues as well as analogues with chemo selective functionality.

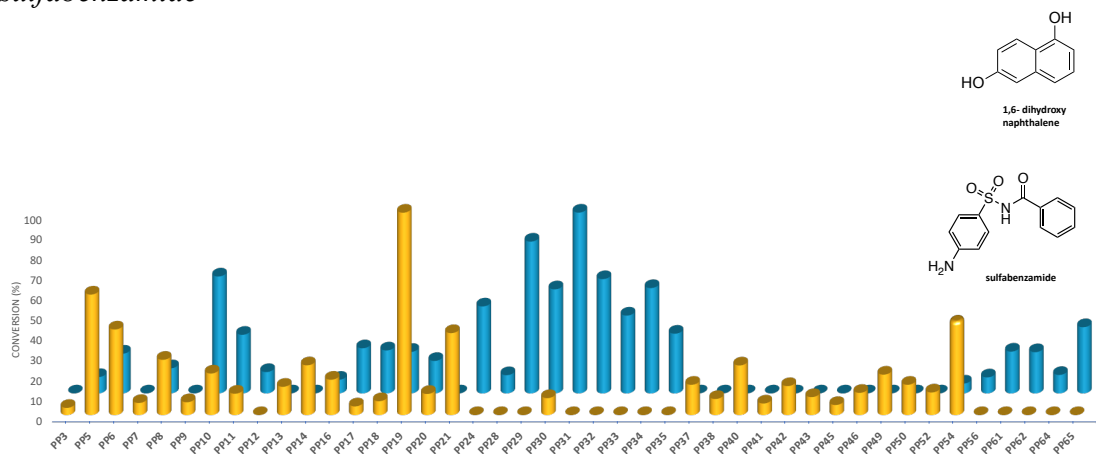
Compounds were designed with two different chemical handles; allylic analogues (**PP1-PP35**) and benzylic analogues (**PP36-PP48**). Amongst the allylic analogues, features including chain length, methylation positions, cyclization, degree of saturation, and functional groups were varied. In order to better understand the effect of chain length on substrate preference, compounds were synthesized with various lengths ranging from an allyl pyrophosphate (C<sub>3</sub>) (**PP1**) to farnesyl pyrophosphate (C<sub>15</sub>) (**PP35**). Derivatives of dimethyl allyl were created with assorted terminal saturated aliphatic chains (**PP6-PP12**), altering the overall length as well as bulk of the donor compound. Presumably, the saturated carbon chains could add more flexibility to the general structure when compared to polyunsaturated chains. Further, carbocation stability of the activated allylic group was investigated through conjugated system and the loss of the C3 methyl group, thought to stabilize the carbocation through hyper-conjugation. The necessity of

this methyl on the C3 carbon was investigated by either removal (**PP3**, **PP14**, **PP16**, **PP17**, and **PP26-PP28**), rearrangement on the C2 carbon position (**PP4**, **PP15**, **PP18**) or by replacement by a chloro- (**PP22**) or bromo- (**PP23**) halogen atom. Additionally, conjugated compounds were synthesized in either linear (**PP13-PP18**) or aromatic (**PP24-PP28**) form. Our library also includes several geranyl derivatives, formed by the addition of a methyl group (**PP31**), ester linked benzyl group (**PP33**), alcohol group (**PP34**) or alkene group (**PP32**) on the terminal methyl. Additionally, a subset of the library consists of benzylic analogues substituted with either electron-donating (methyl or methoxy) or electron-withdrawing (fluoro-, chloro-, bromo- or nitro-) substituents. To investigate the reactivity influenced by substituent orientation and degree of substitution, we synthesized ortho- and meta-methoxy (**PP38** and **PP39**), di-methoxy (**PP45** and **PP46**), di-fluoro (**PP48**) and tri methoxy (**PP48**) derivatives on the benzyl ring.



**Figure 24. Library of synthetic alkyl-PP analogues.**

Analogues were assessed with *NphB* catalyzed reaction with 1,6-DHN or sulfabenzamide



**Figure 25. Percent conversion.**

Percent conversion of library of alkyl pyrophosphate using 1,6-DHN (blue) or sulfabenzamide (orange). Assays were performed under standard condition, 1.2 mM alkyl pyrophosphate, 1 mM sulfabenzamide or 1,6-DHN, *NphB*, 25 mM Tris pH 7.8 5mM MgCl<sub>2</sub>, 50 mM KCl and incubated for 16hrs at 35 °C and evaluated by RP-HPLC.

**Table 5. Summary of HRMS data of 1,6-DHN analogues from NphB catalyzed alkylation reaction with 1,6-DHN and synthetic alkyl-PP analogues.**

Enzyme Product	Chemical Formula	Calculated Mass (Da)	Observed Mass (Da)
<i>1,6-DHN</i>	C <sub>10</sub> H <sub>8</sub> O <sub>2</sub> [M-H] <sup>-</sup>	159.0452	159.0457
<i>1,6-DHN-PP5</i>	C <sub>15</sub> H <sub>16</sub> O <sub>2</sub> [M-H] <sup>-</sup>	227.1078	227.1079
<i>1,6-DHN-PP6</i>	C <sub>16</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	241.1234	241.1237
<i>1,6-DHN-PP8</i>	C <sub>17</sub> H <sub>20</sub> O <sub>2</sub> [M-H] <sup>-</sup>	255.1390	255.1394
<i>1,6-DHN-PP10</i>	C <sub>18</sub> H <sub>22</sub> O <sub>2</sub> [M-H] <sup>-</sup>	269.1547	269.1548
<i>1,6-DHN-PP11</i>	C <sub>19</sub> H <sub>24</sub> O <sub>2</sub> [M-H] <sup>-</sup>	283.1703	283.2645
<i>1,6-DHN-PP16</i>	C <sub>16</sub> H <sub>16</sub> O <sub>2</sub> [M-H] <sup>-</sup>	239.1078	239.1081
<i>1,6-DHN-PP17</i>	C <sub>17</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	253.1234	253.1230
<i>1,6-DHN-PP18</i>	C <sub>17</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	253.1234	253.1238
<i>1,6-DHN-PP19</i>	C <sub>17</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	253.1234	253.1235
<i>1,6-DHN-PP20</i>	C <sub>18</sub> H <sub>20</sub> O <sub>2</sub> [M-H] <sup>-</sup>	267.1391	267.1379
<i>1,6-DHN-PP24</i>	C <sub>20</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	289.1234	289.1232
<i>1,6-DHN-PP28</i>	C <sub>21</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	301.1234	301.1235
<i>1,6-DHN-PP29</i>	C <sub>20</sub> H <sub>24</sub> O <sub>2</sub> [M-H] <sup>-</sup>	295.1703	295.1704
<i>1,6-DHN-PP30</i>	C <sub>18</sub> H <sub>20</sub> O <sub>2</sub> [M-H] <sup>-</sup>	267.1390	267.1391
<i>1,6-DHN-PP31</i>	C <sub>21</sub> H <sub>26</sub> O <sub>2</sub> [M-H] <sup>-</sup>	309.1860	309.1859
<i>1,6-DHN-PP32</i>	C <sub>23</sub> H <sub>28</sub> O <sub>2</sub> [M-H] <sup>-</sup>	355.2016	355.2098
<i>1,6-DHN-PP33</i>	C <sub>27</sub> H <sub>30</sub> O <sub>3</sub> [M-H] <sup>-</sup>	401.2122	401.2132
<i>1,6-DHN-PP34</i>	C <sub>20</sub> H <sub>24</sub> O <sub>3</sub> [M-H] <sup>-</sup>	311.1653	311.1660
<i>1,6-DHN-PP35</i>	C <sub>25</sub> H <sub>32</sub> O <sub>2</sub> [M-H] <sup>-</sup>	363.2329	363.2339
<i>1,6-DHN-PP37</i>	C <sub>18</sub> H <sub>16</sub> O <sub>3</sub> [M-H] <sup>-</sup>	279.1027	279.1026
<i>1,6-DHN-PP54</i>	C <sub>18</sub> H <sub>14</sub> O <sub>4</sub> [M-H] <sup>-</sup>	293.0819	293.0821
<i>1,6-DHN-PP56</i>	C <sub>16</sub> H <sub>17</sub> N <sub>3</sub> O <sub>2</sub> [M-H] <sup>-</sup>	282.1248	282.125
<i>1,6-DHN-PP61</i>	C <sub>18</sub> H <sub>18</sub> O <sub>2</sub> [M-H] <sup>-</sup>	265.1234	265.1244
<i>1,6-DHN-PP62</i>	C <sub>18</sub> H <sub>18</sub> O <sub>3</sub> [M-H] <sup>-</sup>	281.1183	281.1179
<i>1,6-DHN-PP64</i>	C <sub>20</sub> H <sub>16</sub> O <sub>3</sub> [[M-H] <sup>-</sup>	303.1027	303.1031
<i>1,6-DHN-PP65</i>	C <sub>23</sub> H <sub>26</sub> O <sub>3</sub> [M-H] <sup>-</sup>	349.1809	349.1801

### Chemoenzymatic syntheses of alkylated 1,6 dihydroxy naphthalene

NphB was heterologously expressed in *Escherichia coli* BL21 (DE3) cells as a N-terminal His-tagged fusion protein and purified to homogeneity by Ni-NTA affinity chromatography. Although the true native substrate for NphB is unknown, 1,6-DHN is used as a surrogate substrate to assess donor specificity. The purified recombinant

enzyme was incubated with 1,6-DHN and the synthesized alkyl pyrophosphate donors at standard assay conditions (1.2mM alkyl-PP, 1mM aromatic acceptor, 50uM NphB, 25mM Tris, 5mM MgCl<sub>2</sub>, 50mM KCl, pH 7.8, 16 h at 35 C). The formation of alkylated products was analyzed using reverse phase high-performance liquid chromatography (RP-HPLC) and confirmed by high resolution mass spectroscopy (HRMS) (Figure 25 and 28; Table 5).

Analysis of the analytical scale reactions demonstrated that 27 of the 65 pyrophosphate analogues were able to be catalytically transferred by NphB onto the aromatic substrate, 1,6-DHN (Figure 25). Although the HRMS data indicates only monopyrenylated products for all of the reactions, the HPLC chromatograms for 20 of the 27 catalyzed reactions indicated the presence of three different isomer products based off their retention times. Similarly, previous studies have determined the formation of three different geranylated products upon the incubation of NphB with 1,6-DHN and GPP[54].

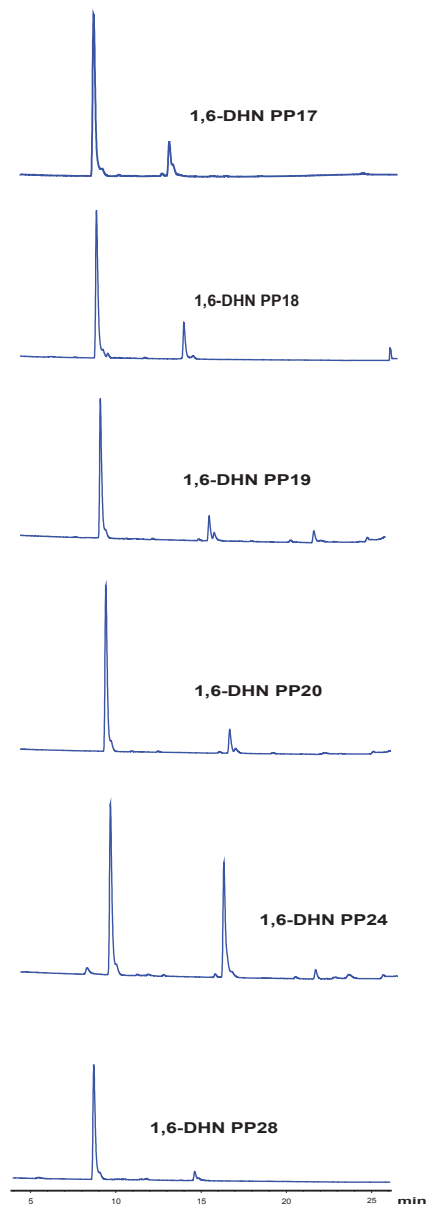
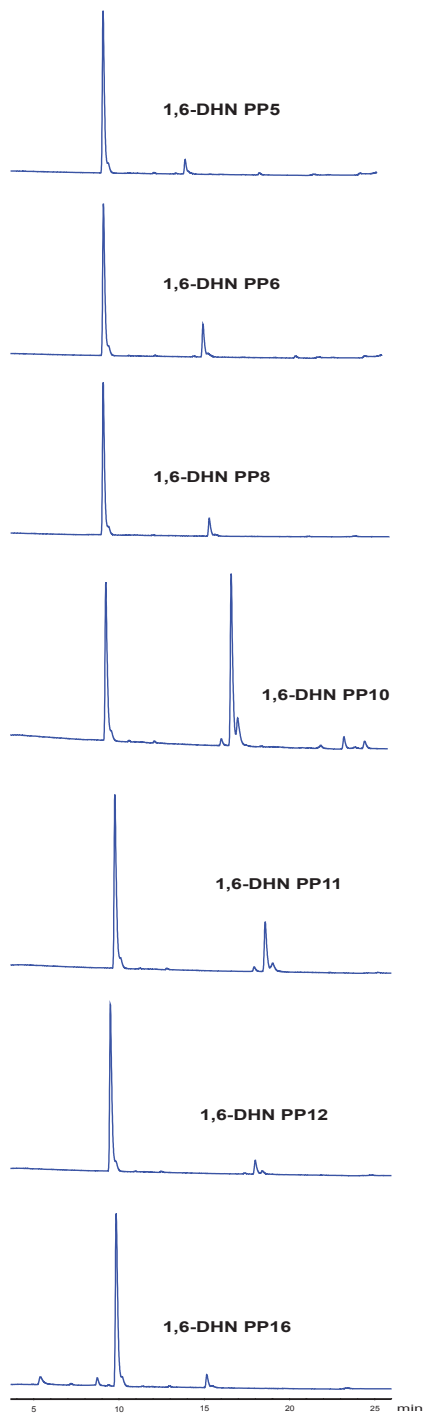
Of the 26-catalyzed reaction 16 displayed significant product formation (>20%) and six showed a >50% product formation. The six highest producing reactions included the native substrate GPP (**PP29**), **PP10**, **PP30-PP32** and **PP34**. As expected, the reaction containing GPP significant conversion of 75% when incubated for 16h. Both **PP10** (58% conversion) and **PP30** (51% conversion) are seven carbons in length and show a similar product conversion, **PP10** is a butane derivative of DMAPP and **PP30** is a butene derivative with a terminal alkene. **PP31** and **PP32** both have added aliphatic groups onto the terminal methyl of GPP. Interestingly, **PP31** with an additional methyl added and was able to show 90% product conversion, overall greater conversion

compared to the native substrate, GPP. However, **PP32** with an additional terminal allyl group, resulted in a decreased the conversion to only 57%. Analogue **PP34** features a terminal alcohol group on the geranyl tail and displays an impressive 52% product conversion.

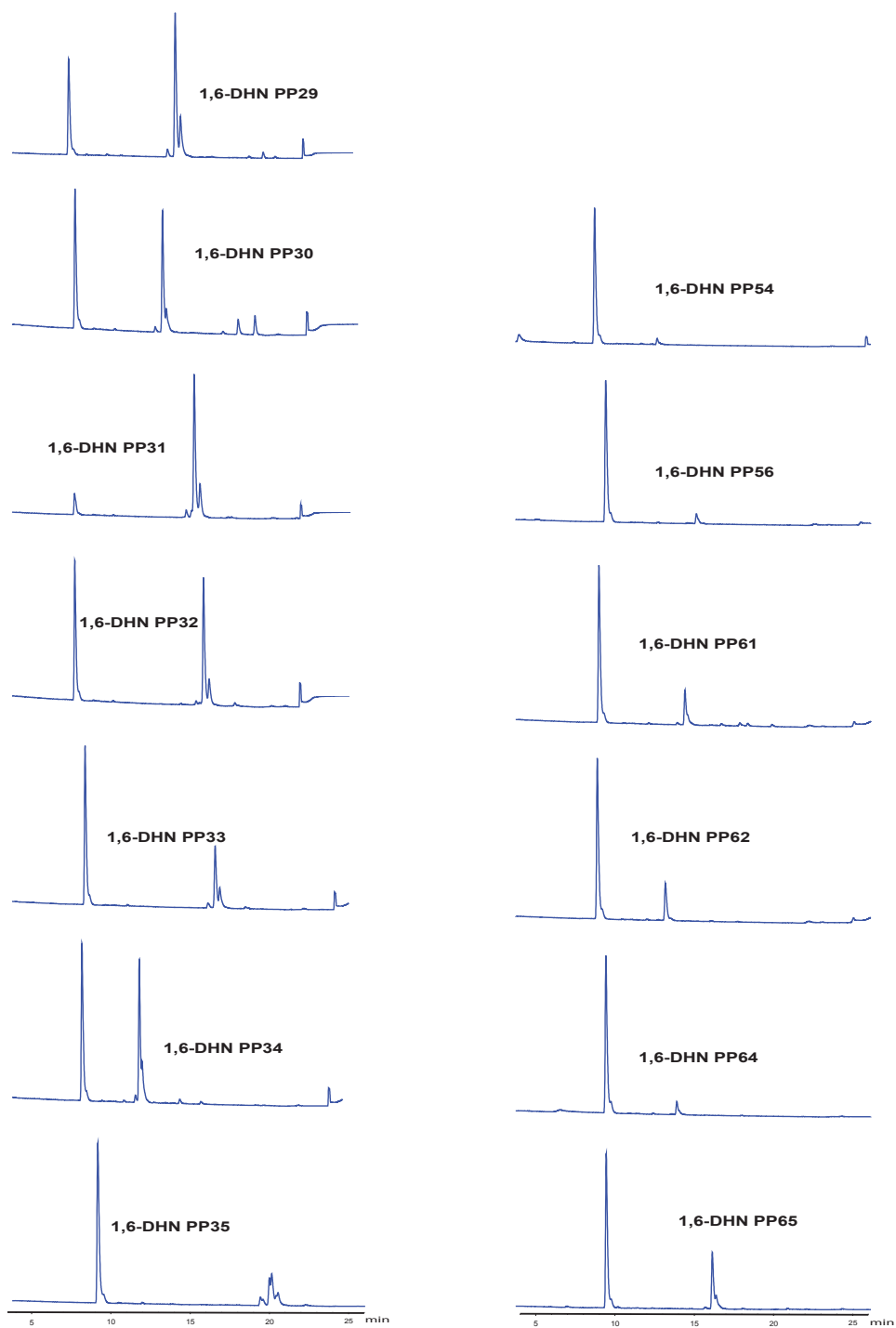
The remaining 10 compounds that showed an appreciable >20% conversion have a large distribution in the length of the alkyl moiety. The smallest compound that was catalytically transferred onto 1,6-DHN was an the allylic cyclopentane (**PP19** (21%)), while the longest were the ether linked benzyl GPP derivative (**PP33** (39%)) and FPP (**PP35** (30%)). FPP has previously been reported has a substrate for 1,6DHN[36] and showed a 30% conversion when incubated for 16h under the stated assay conditions. Additional allylic analogues, **PP6**, **PP11**, **PP18**, and **PP24** showed 20%, 29%, 21%, 43% conversion respectively. **PP6** is an ethyl DMAPP derivative, **P11** is a pentyl DMAPP derivative, while **PP18** is a C2 methylated 2,4-hexadiene. **PP24**, like **PP18**, has a conjugated system; however, it is in the form of a benzyl substituted DMAPP and mediated twice the product conversion compared to **PP18**. Furthermore, three propargyl containing analogues (**PP61**, **PP62** and **PP65**), ranging in linear length between 7 and 12 atoms, displayed appreciable conversion. **PP61** is a propargyl derivative of DMAPP with a terminal propargyl group showing and it underwent a 21% conversion. Both **PP62** and **PP65** are ether linked propargyl derivatives of DMAPP and GPP respectively, **PP62** underwent 20% conversion while **PP65** had a 33% conversion.

The remaining enzyme reactions produced <20% conversion. They included allylic analogues of varying length, methylation and conjugation; **PP5** (DMAPP) (8%), **PP8** (12%), **PP12**(11%), **PP16** (7%), and **PP17** (13%). Additionally, **PP20**, allylic

cyclohexane, and **PP28**, five-carbon chain aromatic displayed 16%, and 9% conversion. Of the 13 synthesized benzyl analogues only one was accepted as substrate; **PP37**, a para-methoxy substituted benzyl analogue, with a 18% conversion. Two additional chemoselective analogues, **PP64**, a para-propargyl ether substituted benzyl compound, and **PP56**, an azido DMAPP derivative showed slight product formation at 9% and 5% respectively. Finally, only one heterocycle analogue was accepted using 1,6-DHN as the substrate, **PP54**, a piperonyl pyrophosphate bearing a 3, 4 methylene dioxy moiety, which exhibited a 14% conversion.







**Figure 26 RP- HPLC chromatograms.**

*RP- HPLC chromatogram of incubation mixtures of 1,6-DHN with pyrophosphate analogues which lead to the NphB catalyzed formation of alkylated sulfabenzamide derivatives. The peak at approximately 9.5 min is 1,6-DHN and the product peaks are after 10 min in the chromatogram.*

### **Chemoenzymatic syntheses of alkylated sulfabenzamide**

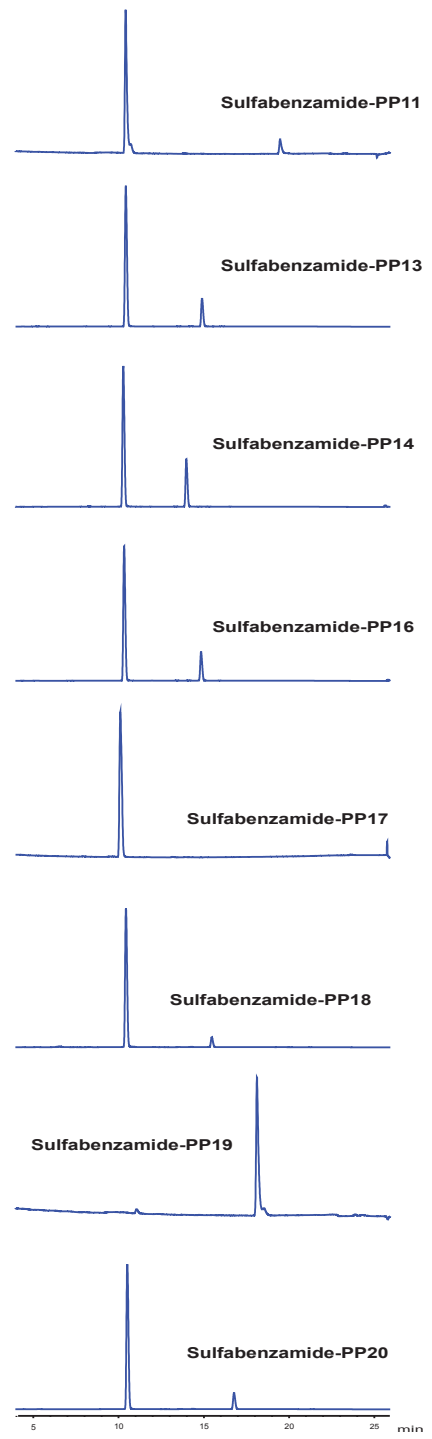
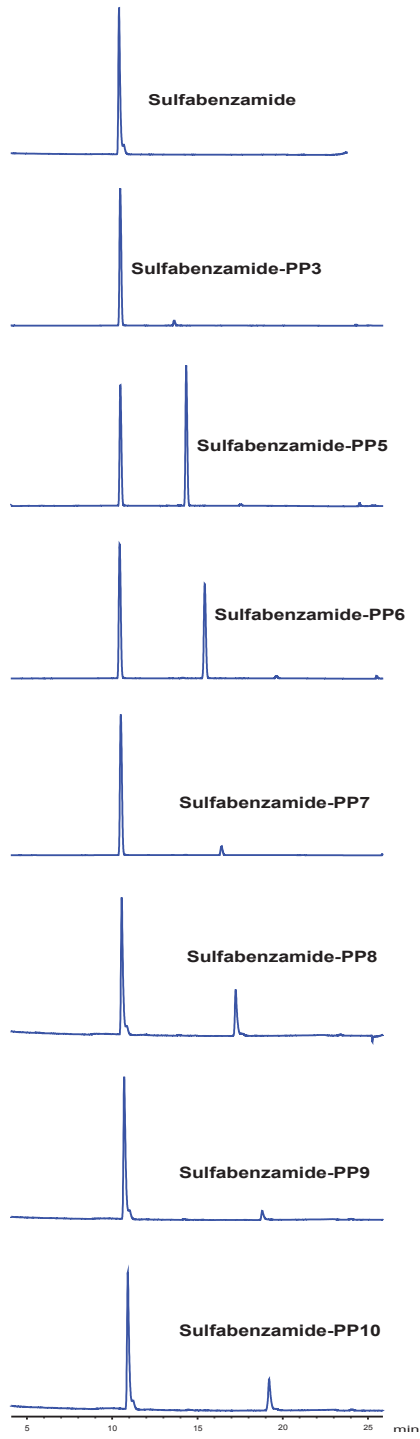
Next, to investigate the utility of NphB to perform catalytic modification of synthetic aromatic substrates, 11 known drug compounds (amoxicillin, methotrexate, tamoxifen, catharanthine, puromycin, camptothecin, yohimbine, sulfadoxin, chloramphenicol, ethynylestradiol and sulfabenzamide) containing aromatic moieties were incubated with prenyl pyrophosphates GPP and DMAPP. The HPLC and HRMS analysis of the enzyme reaction revealed that NphB accepted sulfabenzamide as a prenyl acceptor. In contrast to the native behavior of NphB, no product formation was detected with GPP as the donor substrate when sulfabenzamide is the aromatic acceptor. On the other hand, DMAPP was able to be catalytically transferred onto sulfabenzamide producing a prenylated derivative. The drastic shift in donor substrate preference prompted us to test the catalytic capability of NphB to transfer non-native alkyl pyrophosphates using our library of synthetic pyrophosphates, generating sulfabenzamide derivatives with potentially altered biological activity.

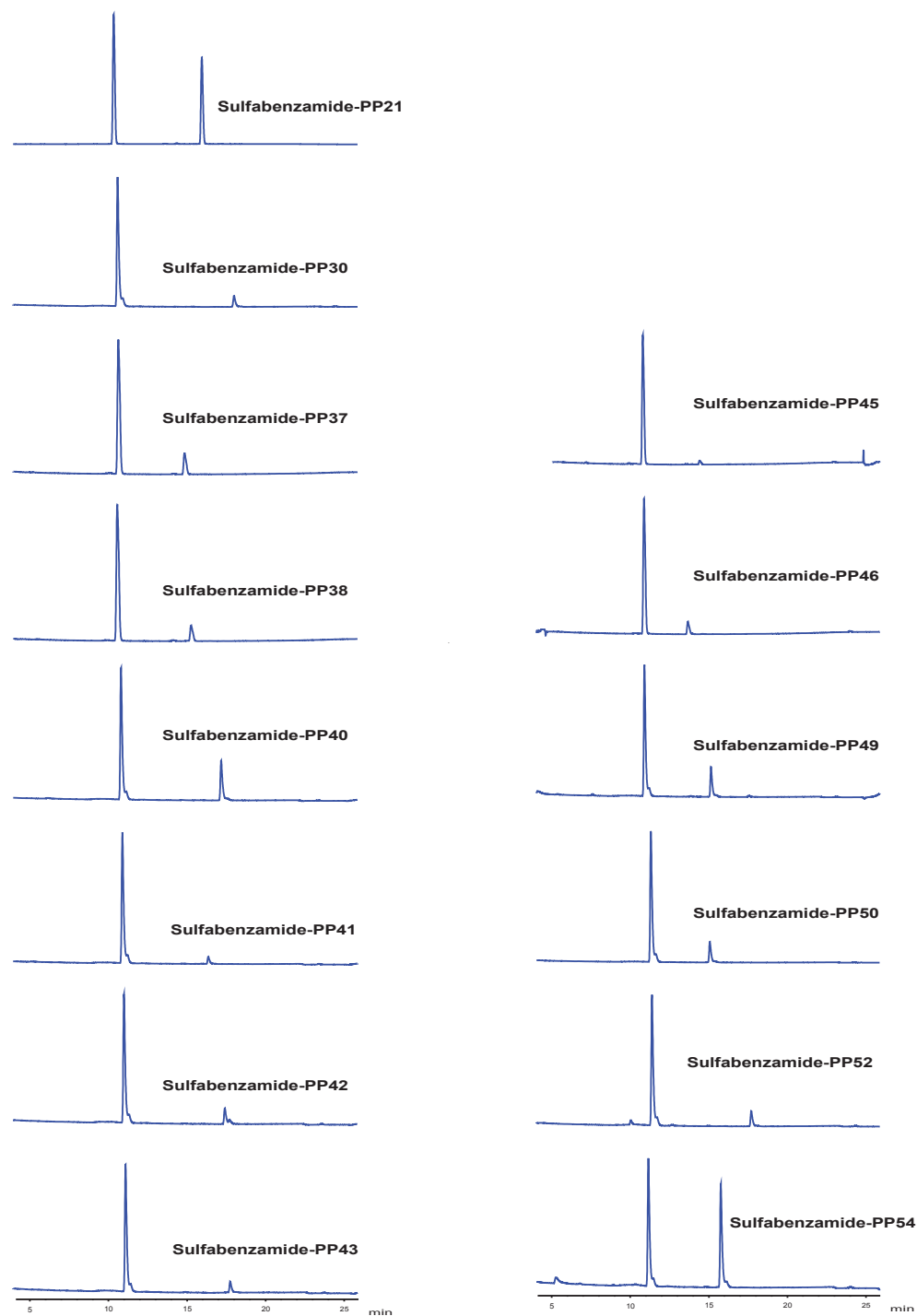
Similar to the analytical reactions involving 1,6-DHN aromatic acceptor, analysis of the product formation was performed using HPLC and HRMS. This revealed that 29 out of the 65 alkyl-PP enzymatic reactions resulted in alkylated sulfabenzamide products (Figure 25 and 29; Table 6). A variety of distinctive analogues were accepted as alkyl donors; markedly, of the 29 accepted donors, less than 50% were seen to be accepted using 1,6-DHN. Overall, the linear length of the accepted donors ranges from a four-carbon chain (**PP5 (DMAPP)** (60%), **PP19** (100%) and **PP3** (4%)) to a seven-carbon chain (**PP10**) (20%). Of the 29 positive reactions, ten pyrophosphate analogues (**PP5, PP6, PP19, PP21, PP14, PP10, PP8, PP40, PP49, PP54**) show an appreciable (>20%)

product conversion. Notably, of the analogues showing greater than 20% product conversion, half of these contain an aromatic (**PP40** (25%), **PP54** (46%), and **PP49** (20%)) or non-aromatic ring structures (**PP19** (100%) and **PP21** (40%)). The remaining five analogues contain various carbon chain length, methylation and conjugation. Diversity in the overall chain length can be seen in the acceptance of a four-carbon chain DMAPP (**PP5**) and the allylic cyclopentane (**PP19**), five-carbon chain ethyl DMAPP derivative (**PP6**), and isopropyl DMAPP derivative (**PP8** (27%)), and a seven-carbon chain, butyl DMAPP derivative (**PP10** (20%)) donor substrates. The C3 methyl group can be found on **PP5**, **PP6**, **PP10** and **PP8**. Compound **PP14** (25%) lacked methyl groups; however, it is the only analogue to show appreciable product formation with a linear conjugated system. Analysis of the subset of benzyl pyrophosphate analogues revealed that NphB is capable of transferring eight out of the 13 benzyl groups onto sulfabenzamide, including; a *para*- (**PP37** (13%)) and *ortho*- (**PP38** (8%)) methoxy substituted benzyl compounds, a *para*- methyl substituted benzyl compound (**PP40**(25%)), fluorine, chlorine and bromine halogenated benzyl compounds (**PP41** (6%), **PP42** (14%), **PP43**(9%)), *para*- and *ortho*- methoxy di-substituted benzyl (**PP45**(11%)), and a *para*- and *meta*-methoxy di-substituted benzyl (**PP46**(8%)). Additionally, four aromatic heterocycle analogues; thiopene (**PP49**(20%)), furane (**PP50**(14%)) a benzothiophen (**PP52**(11%)) and piperonyl pyrophosphate bearing a 3, 4 methylene dioxy moiety (**PP54**(46%)) pyrophosphate were also able to be transferred onto sulfabenzamide, displaying 20%, 15% 11% and 46% conversion respectively.

**Table 6. Summary of HRMS data of sulfabenzamide analogues from NphB catalyzed alkylation reaction with sulfabenzamide and synthetic alkyl-PP analogues.**

Enzyme Product	Chemical Formula	Calculated Mass (Da)	Observed Mass (Da)
<b>Sulfabenzamide</b>	C <sub>13</sub> H <sub>12</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	277.0641	277.0648
<b>Sulfabenzamide-PP3</b>	C <sub>17</sub> H <sub>18</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	331.1111	331.1120
<b>Sulfabenzamide-PP5</b>	C <sub>18</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	345.1267	345.1276
<b>Sulfabenzamide-PP6</b>	C <sub>19</sub> H <sub>22</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	359.1424	359.1442
<b>Sulfabenzamide-PP7</b>	C <sub>20</sub> H <sub>24</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	373.158	373.1592
<b>Sulfabenzamide-PP8</b>	C <sub>20</sub> H <sub>24</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	395.1406	395.1395
<b>Sulfabenzamide-PP9</b>	C <sub>21</sub> H <sub>26</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	409.1562	409.1576
<b>Sulfabenzamide-PP10</b>	C <sub>21</sub> H <sub>26</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	409.1562	409.1577
<b>Sulfabenzamide-PP11</b>	C <sub>22</sub> H <sub>28</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	423.1719	423.1733
<b>Sulfabenzamide-PP13</b>	C <sub>19</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	357.1267	357.1270
<b>Sulfabenzamide-PP14</b>	C <sub>18</sub> H <sub>18</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	343.1111	343.1118
<b>Sulfabenzamide-PP16</b>	C <sub>19</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	357.1267	357.1277
<b>Sulfabenzamide-PP17</b>	C <sub>22</sub> H <sub>22</sub> N <sub>2</sub> O <sub>5</sub> S [M+H] <sup>+</sup>	371.1424	371.1432
<b>Sulfabenzamide-PP18</b>	C <sub>20</sub> H <sub>22</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	371.1424	371.1435
<b>Sulfabenzamide-PP19</b>	C <sub>20</sub> H <sub>22</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	393.1249	393.1266
<b>Sulfabenzamide-PP20</b>	C <sub>21</sub> H <sub>24</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	385.158	385.1590
<b>Sulfabenzamide-PP21</b>	C <sub>20</sub> H <sub>22</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	371.1424	371.1435
<b>Sulfabenzamide-PP29</b>	C <sub>23</sub> H <sub>28</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	435.1719	435.1723
<b>Sulfabenzamide-PP30</b>	C <sub>21</sub> H <sub>24</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	407.1406	407.1420
<b>Sulfabenzamide-PP37</b>	C <sub>21</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	419.1036	419.1029
<b>Sulfabenzamide-PP38</b>	C <sub>22</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+H] <sup>+</sup>	393.1267	393.1275
<b>Sulfabenzamide-PP40</b>	C <sub>21</sub> H <sub>20</sub> N <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	403.1093	403.1106
<b>Sulfabenzamide-PP41</b>	C <sub>20</sub> H <sub>17</sub> FN <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	407.0842	407.0858
<b>Sulfabenzamide-PP42</b>	C <sub>20</sub> H <sub>17</sub> ClN <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	423.0546	423.0553
<b>Sulfabenzamide-PP43</b>	C <sub>20</sub> H <sub>17</sub> BrN <sub>2</sub> O <sub>3</sub> S [M+Na] <sup>+</sup>	467.0042	467.0060
<b>Sulfabenzamide-PP45</b>	C <sub>22</sub> H <sub>22</sub> N <sub>2</sub> O <sub>5</sub> S [M+Na] <sup>+</sup>	453.1647	453.1673
<b>Sulfabenzamide-PP46</b>	C <sub>22</sub> H <sub>22</sub> N <sub>2</sub> O <sub>5</sub> S [M+Na] <sup>+</sup>	449.1147	449.1166
<b>Sulfabenzamide-PP49</b>	C <sub>18</sub> H <sub>16</sub> N <sub>2</sub> O <sub>3</sub> S <sub>2</sub> [M+Na] <sup>+</sup>	395.0500	395.0499
<b>Sulfabenzamide-PP50</b>	C <sub>18</sub> H <sub>16</sub> N <sub>2</sub> O <sub>4</sub> S [M+Na] <sup>+</sup>	379.0729	379.0733
<b>Sulfabenzamide-PP52</b>	C <sub>22</sub> H <sub>18</sub> N <sub>2</sub> O <sub>3</sub> S <sub>2</sub> [M+Na] <sup>+</sup>	445.0657	455.0665
<b>Sulfabenzamide-PP54</b>	C <sub>21</sub> H <sub>18</sub> N <sub>2</sub> O <sub>5</sub> S [M+Na] <sup>+</sup>	433.0834	433.0838



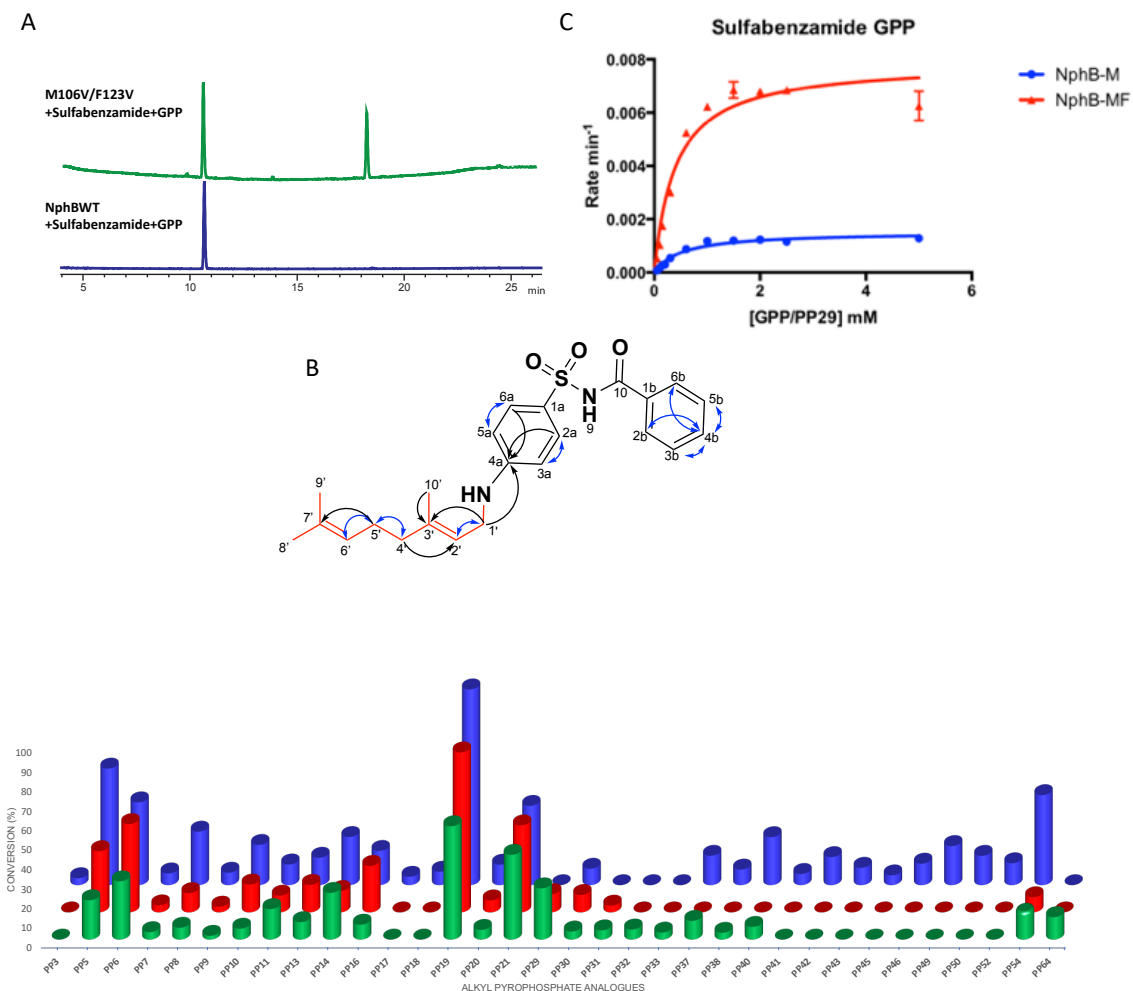


**Figure 27. RP- HPLC chromatograms.**

*Chromatograms of incubation mixtures of sulfabenzamide with pyrophosphate analogues which lead to the NphB catalyzed formation of alkylated sulfabenzamide derivatives.*

### **NphB mutational studies.**

Based on the analysis of the existing crystal structure of NphB, complexed with geranyl S-thiodiphosphate (GSPP) and 1,6-DHN (PDB ID, 1zb6), along with previously performed prenyltransferase mutational studies, we speculated altering residues found in the prenyl binding pocket would subsequently modify the substrate specificities. [36, 56] Guided by the mutational studies on TleC, MpnD, and PagF (TleC Phe170 and Trp97, MpnD Met159 and Tyr80, PagF Phe222), we therefore focused on key residues lining the distal end of the prenyl binding pocket: Met106 and Phe123. [53, 100] To evaluate the impact of the size of the side chain, we constructed the M106V variant. Further modifications to the prenyl binding pocket were made by construction of a double mutant, M106V/F123V. Variant enzymes were assayed against the alkyl-PP library with sulfabenzamide (Figure 28D). Both the single and double mutant, M106 and M106V/F123V, were able to accept GPP as a prenyl donor (M106V/F123V  $k_{cat}$   $7.8 \times 10^{-3} \text{ min}^{-1}$ ,  $K_M$  0.4 mM; M106V  $1.5 \times 10^{-3} \text{ min}^{-1}$ ,  $K_M$  0.6 mM), an activity not present in wild-type enzyme when incubated with sulfabenzamide as the acceptor (Figure 28 A-C). Furthermore, both mutants were able to transfer **PP31** (5% and 3.5%) and the double mutant accepted **PP32**, **PP33**, and **PP64** to a small degree (5%, 4%, and 11.5%). New activity shown in the NphB mutants was seen when incubated with longer pyrophosphate analogues and sulfabenzamide. Although the majority of wild-type activity was maintained in the mutants, a decrease in product formation was seen for all other reactions.



**Figure 28.** NphB mutation analysis.

**(A)** Reverse-phase HPLC analysis of NphB wild-type (bottom) and NphB M106V/F123V (top) enzyme assays using sulfabenzamide and GPP/PP29. **(B)** NMR correlations (blue arrow, COSY correlations, black arrow, HMBC correlations) of sulfabenzamide-GPP product generated through incubation NphB M106V/F123V. **(C)** Kinetic curves showing the enzyme activities of NphB M106 and M106V/F123V with sulfabenzamide and GPP/PP29. **(D)** Percent conversion of library of alkyl pyrophosphate incubated with sulfabenzamide using NphB-WT (blue) and NphB-M106V (red) and NphB-M106V/F123V (Green) Assays were performed under standard condition and incubated for 16hrs. Alkylated sulfabenzamide products marked with an asterisk were isolated using HPLC and characterized using NMR



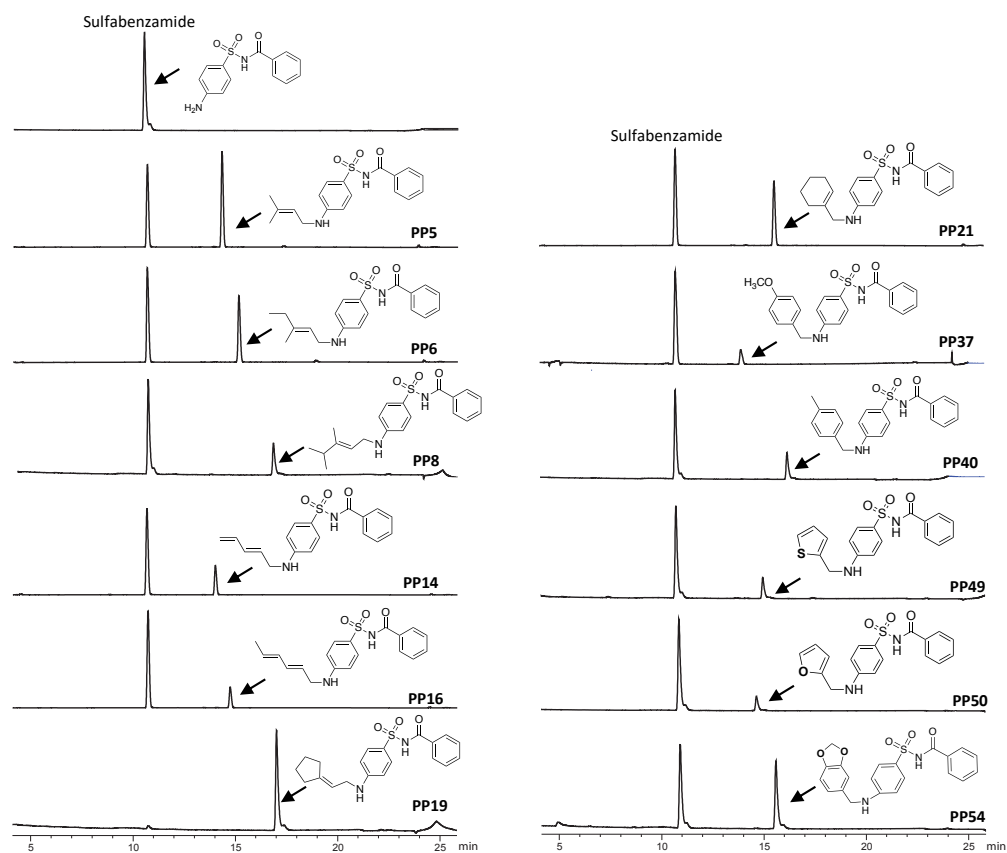
### Structure characterization of sulfabenzamide products.

To elucidate the structure of the alkylated sulfabenzamide products, which were generated for 12 out of 29 reactions, confirmed by HRMS and RP-HPLC to yield products, were scaled up and purified *via* a semi-preparative RP-HPLC protocol (See method B (HPLC) in Experimental Methods), and subjected to NMR analysis (Table 7). Sulfabenzamide products selected for characterization displayed >15% product conversion, as determined in the initial analytical reactions, and possess unique electronic and structural features, e.g. carbon chain length, conjugated systems, cyclization and aromaticity. Specifically, the characterized sulfabenzamide products were diversified through the attachment of allylic analogues, DMAPP (**PP5**), GPP (**PP29**), ethyl and isopropyl DMAPP derivative (**PP6** and **PP8**), as well as two diene analogues with C5 and C6 carbon chain length (**PP14** and **PP16**) and two cyclic pyrophosphates (**PP19** and **PP21**). Additionally, two benzylic derivatized sulfabenzamide compounds, (**PP37** and **PP40**), and three heterocyclic analogues (**PP49**, **PP50** and **PP54**) were also scaled up for characterization. The resulting purified sulfabenzamide products were subjected to <sup>1</sup>H and 2D NMR experiments to determine the exact position of derivatization on the parent compound. In addition to the <sup>1</sup>H experiments, all purified sulfabenzamide products have a COSY spectrum; however, due to limited available quantities, experiments such as HMBC and HSQC could only be performed on **PP29**, **PP8**, **PP19**, **PP49**, **PP50** and **PP54**. Previous studies have indicated that the position of the alkylation was influenced by the specific alkyl donor (Chapter 3, [56]); therefore, given the wide variety of structurally characterized

alkylated sulfabenzamide products, we expect insight into the regio selectivity and specificity of NphB catalyzed reaction.

We have concluded that all scaled up sulfabenzamide reactions resulted in the regiospecific N-alkylation (Figure 29, NMR Table 5) as characterization by NMR analysis, including chemical shift values, integration, splitting, and COSY and/or HMBC correlations. The  $^1\text{H}$ NMR of the sulfabenzamide derivatives, when compared to sulfabenzamide, revealed that all aromatic protons were accounted for on both rings of the parent compound found in the 6.0-8.0ppm range, strongly suggesting an N-substitution. Signals corresponding to the two aromatic rings on sulfabenzamide could be differentiated by their splitting pattern, as the benzamide ring displays three distinct peaks with multiplet splitting pattern integrating to a 2:2:1 ratio whereas the 4-amino benzenesulfonyl has two proton peaks with a doublet splitting pattern integrating to a 1:1 ratio. Assignments of the aromatic proton signals were further supported by the COSY spectra showing correlation peaks between the H2a-H3a and H6a-H5a on the 4-amino benzenesulfonyl ring as well as correlation peaks between H2b-H3b, H3b-H4b, H4b-H5b and H5b-H6b on the benzamide ring. In some cases, the COSY spectra also showed correlation between the proton on the amino NH and the H1' of the substituent alkyl or benzyl group. For the samples with sufficient quantities of purified compound, we were able to obtain additional 2D experiments, the  $^1\text{H} - ^{13}\text{C}$  HMBC and  $^1\text{H} - ^{13}\text{C}$  HSQC spectra were collected, displaying connectivity between H1' alkyl or benzyl group to the C4a of the 4-amino benzenesulfonyl ring, unequivocally confirming the N-alkyl addition. Each scale up reaction was additionally confirmed by HRMS data, supporting on the presence of only a mono-alkylated product. These findings verify the

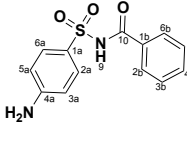
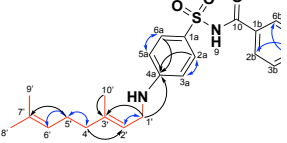
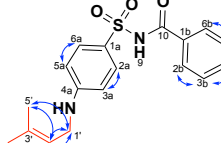
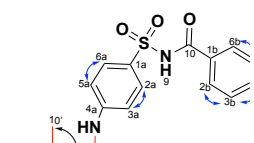
novel capabilities of NphB to catalyze the N-alkylation of sulfabenzamide using natural donors DMAPP and GPP, and non-native donors including dienes, para-substituted benzyl and heterocyclic pyrophosphate donors

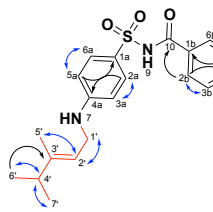
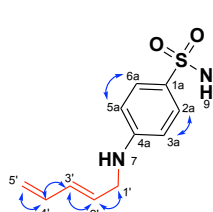
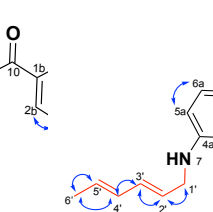
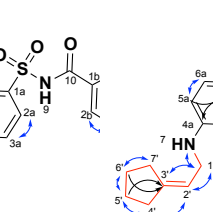


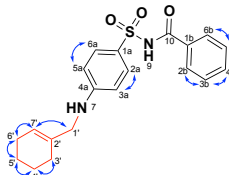
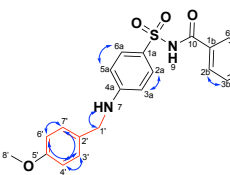
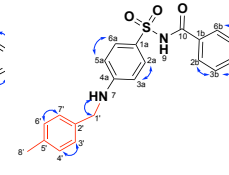
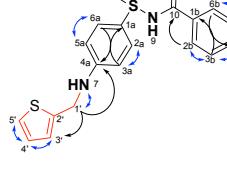
**Figure 29. Sulfabenzamide derivatives.**

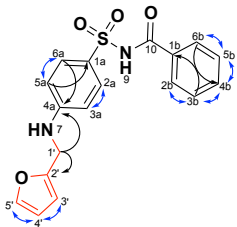
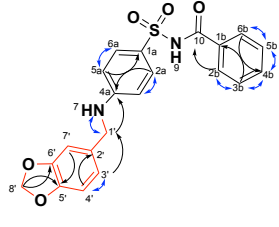
*Reverse-phase HPLC analysis of NphB wild-type catalyzed reactions that resulted in structure elucidated via NMR and RP-HRMS.*

**Table 7. NMR data of alkylated sulfabenzamide**

								
	<b>Sulfabenzamide</b>		<b>Sulfabenzamide-GPP_PP29</b>		<b>Sulfabenzamide-PP5</b>		<b>Sulfabenzamide-PP6</b>	
<b>Position</b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>
NH	12.07, s				8.13, s		8.13, s	
NH	6.15, s (NH <sub>2</sub> )				6.56, s		6.52, s	
1a								
2a	7.63, d (8.8)	130.6	7.78, d (8.9)	129.4	7.59, d (8.4)		7.62, d (8.7)	
3a	6.61, d (8.8)	112.7	6.63 d (8.9)	110.1	6.52, d (8.4)		6.57, d (8.7)	
4a				152.7				
5a	6.61, d (8.8)	112.7	6.63 d (8.9)	110.1	6.52, d (8.4)		6.57, d (8.7)	
6a	7.63, d (8.8)	130.6	7.78, d (8.9)	129.4	7.59, d (8.4)		7.62, d (8.7)	
1b								
2b	7.83, dd (8.4, 1.3)	128.7	7.79, dd (7.9, 1.0)	127.3			7.84, d (7.1)	
3b	7.47, dd (8.4, 7.4)	128.9	7.60, t (7.9)	127.8	7.36, m		7.40, t (7.1)	
4b	7.60, tt (7.4, 1.3)	133.5	7.47, tt (7.4, 1.0)	132.1	7.44, m		7.50, m	
5b	7.47, dd (8.4, 7.4)	128.9	7.6	127.8			7.40, t (7.1)	
6b	7.83, dd (8.4, 1.3)	128.7	7.79	127.3			7.84, d (7.1)	
1'			3.73, d (6.5)	39.9	3.64, dd (6.5, 5.9)		3.67, t (5.7)	
2'			5.26, t (6.5)	120.5	5.22, t (6.5)		5.22, t (5.7)	
3'				137.7				
4'			2.06, m	38.7	1.68, s		1.99, q (7.8)	
5'			2.11, m	25.5	1.69, s		0.96, t (7.8)	
6'			5.07, t (7.1)	123.1			1.68, s	
7'				130.3				
8'			1.57, s	15.8				
9'			1.57, s	15.8				
10'			1.73, s	14.5				
Solvent	DMSO-d <sub>6</sub>		CD <sub>3</sub> OD		DMSO-d <sub>6</sub>		DMSO-d <sub>6</sub>	

								
	<b>Sulfabenzamide-PP8</b>		<b>Sulfabenzamide-PP14</b>		<b>Sulfabenzamide-PP16</b>		<b>Sulfabenzamide-PP19</b>	
<b>Positi on</b>	<b><math>\delta_{\text{H}}</math>, multi (J)</b>	<b><math>\delta_{\text{C}}</math></b>	<b><math>\delta_{\text{H}}</math>, multi (J)</b>	<b><math>\delta_{\text{C}}</math></b>	<b><math>\delta_{\text{H}}</math>, multi (J)</b>	<b><math>\delta_{\text{C}}</math></b>	<b><math>\delta_{\text{H}}</math>, multi (J)</b>	<b><math>\delta_{\text{C}}</math></b>
NH			8.14, s		12.12, s		8.14, brs	
NH			6.56, s		6.54, s		6.37, brs	
1a		124.09						129.42
2a	7.93, d (8.9)	127.71	7.54, d (8.4)		7.63, d (8.7)		7.59, d (8.8)	129.45
3a	6.60, d (8.9)	111.44	6.49, d (8.4)		6.59, d (8.7)		6.52, d (8.8)	110.8
4a		152.8						151.8
5a	6.60, d (8.9)	111.44	6.49, d (8.4)		6.59, d (8.7)		6.52, d (8.8)	110.8
6a	7.94, d (8.9)	127.71	7.54, d (8.4)		7.63, d(8.7)		7.59, d (8.8)	129.49
1b		124.09						136.76
2b	7.79, dd (8.4,1.3)	130.97	7.84, dd (8.3, 1.6)		7.84, dd (8.3, 1.4)		7.86, d (8.0)	128.24
3b	7.44, t, (8.4, 7.4)	128.9	7.3		7.42, dd (8.3, 7.5)		7.36 t, (7.6)	128.24
4b	7.55, tt (7.5, 1.3)	133.22	7.34, m		7.53, tt (7.5, 1.4)		7.44, t (7.3)	131.49
5b	7.44, t, (8.4, 7.4)	128.9	7.30, m		7.42, dd (8.3, 7.5)		7.36 t, (7.6)	128.21
6b	7.79, dd (8.4,1.3)	130.97	7.84		7.84, dd (8.3, 1.4)		7.86, d (8.0)	128.7
1'	3.75, d (6.6)	41.24	3.75, t (5.6)		3.75, td(5.7, 1.5)		3.61, d (6.0)	42.35
2'	5.30, t (6.7)	117.84	5.79, dt (15.3, 5.6)		5.58, dt (15.1, 5.7)		5.34, p (4.1, 2.1)	
3'		146.11	6.22, dd (15.3, 10.5)		6.17, ddt (15.1, 10.5, 1.5)			145.5
4'	2.28, hept (7.3,7.0)	36.71	6.35, dt (16.9, 10.5)		6.05, ddd(15.1, 10.5, 1.5)		2.24, m	28.89

	 Sulfabenzamide-PP21		 Sulfabenzamide-PP37		 Sulfabenzamide-PP40		 Sulfabenzamide-PP49	
Position	$\delta_H$ , multi (J)	$\delta_C$	$\delta_H$ , multi (J)	$\delta_C$	$\delta_H$ , multi (J)	$\delta_C$	$\delta_H$ , multi (J)	$\delta_C$
NH	12.12, s		6.55, br s		8.41, s		8.3, brs	
NH	6.54, s				6.53, t (5.8)		6.62, t (6.0)	
1a								134.1
2a	7.60, d (8.4)		7.50, d (8.4)		7.49, d (8.5)		7.53, d (8.7)	128.6 8
3a	6.56, d (8.4)		6.47, d (8.4)		6.46, d (8.5)		6.55, d (8.9)	111.0 2
4a								150.0 3
5a	6.56, d (8.4)		6.47, d (8.4)		6.46, d (8.5)		6.55, d (8.7)	111.0 2
6a	7.60, d (8.4)		7.50, d (8.4)				7.53, d (8.7)	128.6 8
1b								139.6 4
2b	7.85, d (7.7)		7.85, d (7.1)		7.85, d (7.1)		7.87, dd (8.4,1.4)	128.7 4
3b	7.39, t (7.6)		7.27, t (7.1)		7.26, dd (7.1, 6.6)		7.27, tt (7.2, 1.4)	127.7 2
4b	7.49, m		7.32, t (7.1)		7.31, t (6.6)		7.33, tt (7.4, 1.4)	129.9 7
5b	7.39, t (7.6)		7.27, t (7.1)		7.26, dd (7.1, 6.6)		7.27, tt (7.2, 1.4)	127.7 2
6b	7.85, d (7.7)		7.85, d (7.1)		7.85, d (7.1)		7.87, dd (8.4,1.4)	128.7 4
1'	3.57, d (5.7)		4.20, d (5.9)		4.22, d (5.8)		4.45, d (4.45)	42.04
2'								
3'	1.94, m		7.25, d (8.1)		7.21, d (7.7)		7.03, dd (3.4, 1.1)	125.2 9
4'	1.57, m		6.87, d (8.4)		7.1, d (7.7)		6.94, dd (5.0,3.4)	127.1 9
5'	1.50, m						7.34, dd (5.1,1.3)	125
6'	1.95, m		6.87, d (8.4)		7.1, d (7.7)			

				
	<b>Sulfabenzamide-PP50</b>		<b>Sulfabenzamide-PP54</b>	
<b>Position</b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>	<b><math>\delta_H</math>, multi (J)</b>	<b><math>\delta_C</math></b>
NH	8.29, brs		8.16, brs	
NH	6.45, t (6.0)		6.64, brs	
1a		133.9		132.16
2a	7.53, d (8.7)	128.65	7.53, dd (8.8)	128.96
3a	6.56, d (8.8)	110.94	6.51, d (8.9)	110.93
4a		149.97		150.58
5a	6.56, d (8.8)	110.94	6.51, d (8.9)	110.93
6a	7.53, d (8.7)	128.65	7.53, d (8.8)	128.96
1b		140.79		138.85
2b	7.87, dd (7.9, 1.4)	128.69	7.86, dd (8.4,1.3)	128.71
3b	7.27, dd (8.2, 6.5)	127.73	7.30 t, (7.6,7.4)	127.9
4b	7.33, dd (8.2, 6.5)	129.94	7.36, t (7.5, 7.5)	130.47
5b	7.27, dd (8.2, 6.5)	127.73	7.30 t, (7.6,7.4)	127.9
6b	7.87, dd (7.9, 1.4)	128.69	7.86, dd (8.4,1.3)	128.71
1'	4.25, d (5.8)	40.07	4.18, d (5.1)	46.26
2'		153.52		134.13
3'	6.29, d (3.2)	107.36	6.89, s	108.08
4'	6.36, dd (3.2, 1.8)	110.76	6.82, m	108.5
5'				146.43
6'				147.68
7'			6.83, m	120.63
8'				5.95, s
10				169.14
Solvent	DMSO-d6		DMSO-d6	

## Discussion

The analysis of our *in vitro* activity assays revealed NphB capable of the successful catalytic transfer of several chemically and structurally distinct alkyl-PP onto two different aromatic substrates, 1,6-DHN and sulfabenzamide. Remarkably, these results indicate a clear donor-dependent alkylation reaction and a novel N-prenylation enzymatic functionality resulting in the diversification of the drug compound,

sulfabenzamide. Recently, a donor-dependent acceptor binding mechanism was reported for AtaPT from *A. terreus* NIH2624, wherein a sequential donor and acceptor binding order was determined through isothermal titration calorimetry.[53] The alkyl donor then acceptor binding order was also proposed for NphB, based on the crystal structure complexed with GSPP alone.[54] In the present study, we observed many examples of alkyl donors accepted as substrates by NphB when incubated with 1,6-DHN; however, the same alkyl donors would not result in a product formation using sulfabenzamide. Additionally, by employing structural engineering rationale we generated the single and double mutants M106V and M106V/F123V, and were able to revive the native geranylation capabilities of NphB when incubated with sulfabenzamide.

Alkyl-PP donor specificities were revealed by the enzymatic reaction with 1,6-DHN or sulfabenzamide. As seen in the results, a distinction in chain length preference is made according to the aromatic acceptor substrate. For reactions with 1,6-DHN, a minimal conversion of 8% was seen with DMAPP, which has a linear length of four carbons; whereas, when reacted with sulfabenzamide, a more substantial conversion of 60% was observed. It is our understanding that this is the first occurrence of NphB appreciably accepting DMAPP as a prenyl donor. It became apparent that the enzymatic reactions with sulfabenzamide loss the ability to transfer longer chain and bulkier analogues. GPP and derivatives (**PP29/GPP**, **PP31**, **PP32**, **PP65**, **PP33** and **PP34**) were solely transferred onto 1,6-DHN to varying product yields.



To explain these results, we offer the following explanations. NphB has been shown to geranylitate 1,6-DHN in three different positions on the naphthalene ring, signifying that 1,6-DHN must adopt three different orientations in the binding pocket that result in product formation.[54] On the basis of complexed crystal structure, 1,6-DHN is stabilized within the active site by a significant contribution from the geranyl tail of GSPP, and without these interactions, such as the case with DMAPP, 1,6-DHN could bind too loosely and therefore be unable to produce any appreciable product. In the case of NphB incubation with sulfabenzamide, the prenylation site resides in the same orientation as it would with 1,6-DHN within the active site, however additional hydrogen bonding potential and the larger size of the substrate is likely to contribute to its increased stability with respect to orientation in the active site in the absence of the geranyl tail of GPP. A fixed conformation could also explain the observation of a single N- alkylation mode for sulfabenzamide derivatives, a similar relationship between larger substrates and a single product have previously been reported for NphB reactions.[54] Because the larger prenyl donors could occupy the same space as the benzamide moiety of sulfabenzamide in the active site, the steric bulk is believed to prevent the binding of sulfabenzamide and therefore the formation of products, as seen in our results. The NphB mutants M106V and M106V/F123V relieved steric restrictions up the prenyl binding pocket, potentially decreasing the steric hindrance between the terminal end of the alkyl donor and benzamide moiety, as the mutants showed higher product conversion with longer chained alkyl-PP (Figure 28).

In conclusion, NphB demonstrated generous capabilities to transfer diverse unnatural and synthetic alkyl donors onto 1,6-DHN as well as the synthetic pharmaceutical drug, sulfabenzamide, ultimately generating novel N-alkylated sulfabenzamide derivatives. Furthermore, through structure-based enzyme engineering NphB was able to broaden its substrate acceptance and generate a geranylated sulfabenzamide derivative. These results implicate the plausibility of prenyltransferases in biotechnologies and expand on their catalytic abilities to feasibly generate novel drug derivatives.

## **Experimental**

### **Protein Expression and Purification**

The recombinant plasmid, pET-28a containing synthetic genes for NphB bearing N-terminal His<sub>6</sub>, was transformed into *E. coli* Rosetta cells. After overnight growth, 3 mL of bacterial culture was transferred to 4 L flasks containing 1.2 L Luria-Berani medium and 50 µg kanamycin ml<sup>-1</sup> and were grown at 37°C for approximately 4 h at 220 r.p.m. (O.D<sub>600</sub> =0.6-0.8). Protein expression was induced by the addition of IPTG (0.5 mM final) and continually shaken at 220 r.p.m. for an additional 16-19 h at 20°C. Bacterial cultures were harvested by centrifugation and the resulting cell pellets were resuspended in a lysis buffer (50mM NaH<sub>2</sub>PO<sub>4</sub>, 200mM NaCl, pH 7.8). Cells were lysed by way of sonication on ice 60 times for 10 s (Fisher Scientific Model FB505; Thermo Fisher Scientific, Waltham, MA). In order to remove unwanted cellular debris and insoluble protein, the lysed cells were spun down at 16000 r.p.m. for 1 h at 10°C. The supernatant, containing the expressed recombinant enzyme with a N-terminal His<sub>6</sub> tag, was purified using Ni Sepharose resin affinity chromatography column (GE

Healthcare, USA). Protein was eluted using 50 mM NaH<sub>2</sub>PO<sub>4</sub>, 200 mM NaCl, pH 7.8 and 500 mM Imidazole. The purified protein was concentrated and introduced to a new buffer condition (20 mM Tris-HCl, pH 8, 30 mM NaCl) through centrifugal filtration. Purity was evaluated by SDS-PAGE and the resulting protein bands indicate both a high grade of purity and the appropriate molecular weight for NphB. A Bradford assay was conducted to conclude a final enzyme concentration using BSA as a standard.

*Site-directed mutagenesis.* The plasmids containing the mutated NphB (M106V, M106V/F123V) were constructed by Genescript in a pET-28a vector. Each mutant protein contained an N-terminal His<sub>6</sub> and were expressed and purified identically to the wild type NphB.

### ***In vitro* Activity Assays**

*Analytical Assays.* NphB analytical reactions were performed in a final volume of 20ul, consisting of 1.2mM pyrophosphate analogues, 1mM sulfabenzamide (Sigma-Aldrich) or 1,6 Dihydroxy Naphthalene and 6uM purified NphB in a reaction buffer (25 mM Tris buffer pH 8.0, 5 mM MgCl<sub>2</sub>, 50 mM KCl) and incubated at 35 °C for 16 h. Reactions were terminated through cold methanol quenching followed by centrifugation (10,000 g for 30min) to remove precipitated protein. Reaction analysis was completed on RP-HPLC using method A (general method). NphB large scale reactions were conducted in a volume of 1.2ml or 10ml consisting of 4.16mM pyrophosphate analogues, 6mM sulfabenzamide and 0.3mM purified NphB in reaction buffer. Putative products were subsequently isolated by semi-prep HPLC through method B and confirmed by NMR and HRMS with positive (+) and/or negative (-) mode.

*HPLC and NMR methods.* HPLC was accomplished using Agilent 1220 system equipped with a DAD detector. **Method A (HPLC):** To monitor NphB reactions, analytical reverse-phase (RP) HPLC employed a Gemini-NX, C-18 (5  $\mu$ m, 4.6 mm  $\times$  250 mm) column (Phenomenex, Torrance, California, USA) [gradient of 1% B to 10% B over 10 min, 10% B to 50% B over 5 min, 50% B to 100% B for 12 min, 100% B to 1% B over 1 min, 1% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 1 mL min<sup>-1</sup>; A<sub>254/235</sub>]. The reaction was monitored by the retention time difference between starting material and product. **Method B (HPLC):** Semi-preparative RP HPLC was conducted on a Gemini-NX, C-18 (5  $\mu$ m, 10  $\times$  250 mm) column (Phenomenex, Torrance, California, USA) to purify the Sulfabenzamide analogues [gradient of 1% B to 10% B over 10 min, 10% B to 50% B over 5 min, 50% B to 100% B for 12 min, 100% B to 1% B over 1 min, 1% B for 7 min (A = ddH<sub>2</sub>O with 0.1% TFA; B = acetonitrile) flow rate = 2 mL min<sup>-1</sup>; A<sub>254</sub>]. High-resolution mass spectrometric (HRMS) data and liquid chromatography mass spectrometric (LCMS) were obtained on Agilent 6545-QTOF W/1290 HPLC mass spectrometer at the University of Oklahoma, Department of Chemistry and Biochemistry. NMR spectra were obtained on Varian VNMRS 500 MHz instruments at the NMR facility of the Department of Chemistry and Biochemistry of University at the Oklahoma using 99.9% DMSO-d<sub>6</sub> with 0.05% v/v TMS. <sup>1</sup>H, <sup>13</sup>C and <sup>31</sup>P chemical shifts were referenced to internal solvent resonances. Multiplicities are indicated by s (singlet), d (doublet), t (triplet), q (quartet), quin (quintet), m (multiplet), and br (broad). Chemical shifts are reported in parts per million (ppm) and coupling constants *J* are given in Hz. All NMR spectra were recorded at ambient temperature and processed using MestReNova software.

*Kinetic characterization sulfabenzamide derivatives for NphB and NphB mutants.*

Specific kinetic measurements involving NphB mutants, sulfabenzamide and GPP were accomplished by monitoring product formation on HPLC (254nm). Reactions took place in 20ul total volume, including 2mM sulfabenzamide, appropriate amount of NphB and various concentrations of GPP (0.05 – 5mM) in a reaction buffer (25 mM Tris buffer pH 8.0, 5 mM MgCl<sub>2</sub>, 50 mM KCl), The reactions were performed at 35°C and were stopped by the addition of 30ul cold methanol. Each reaction was conducted in triplicate and optimized through enzyme concentration and reaction time to ensure analysis in the linear range. Specific kinetic constants,  $K_m$  and  $k_{cat}$  were determined by fitting the data to the Michaelis-Menten equations using non-linear regression analysis provided by Prism 5.04 (GraphPad Software, Inc. La Jolla, CA 92037 USA)

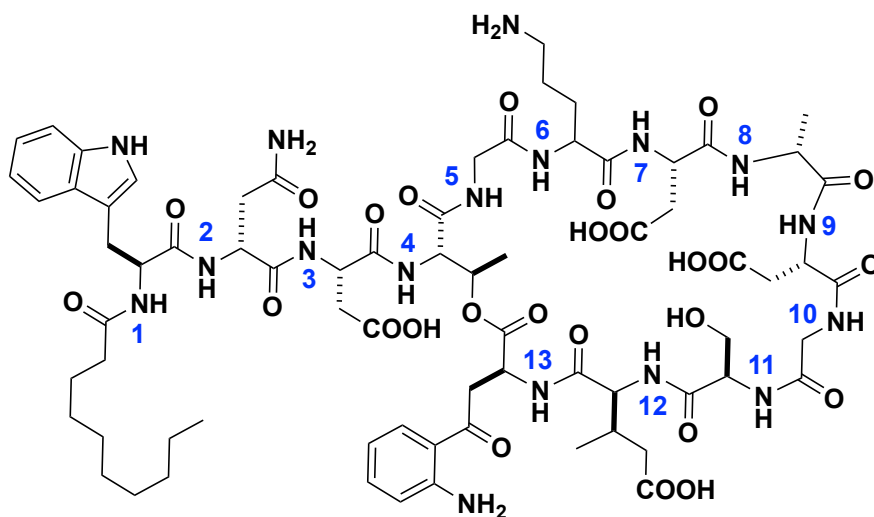
## **Chapter 5. Biocatalytic applications of prenyltransferases in drug diversification: A proof-of-concept study using FDA approved antibiotic, daptomycin**

### **Introduction**

Daptomycin is produced in *Streptomyces roseosporus* as a nonribosomally assembled lipopeptide and belonging to the group of calcium-dependent antibiotics. [101-103] Daptomycin was first discovered in 1980 and has been an FDA approved pharmaceutical since 2003 to treat skin infections caused by gram-positive bacteria *S. aureus*, *Streptococcus pyogenes*, *Streptococcus agalactiae*, *Streptococcus dysgalactiae* as well as *E. faecalis* and in 2006 it was approved for non-topical treatment of bacteraemia and endocarditis caused by MSSA and MRSA. [104-110] Additionally, daptomycin has rapid bactericidal activity against many resistant gram-positive organisms including penicillin-, vancomycin-, and methicillin- resistant bacteria making daptomycin a first-line drug of last resort.[111, 112] Notably, daptomycin cannot be used to treat pneumonia and respiratory infections due to inactivation by the pulmonary surfactant. [113] The structure of daptomycin is made up of 13 amino acids, six of which are non-standard including B-Asn, ornithine (Orn), D-Ala, D-Ser, (2S, 3R)-methylglutamate (MeGlu ) and kynurenine (Kyn). The macrocyclic core contains 10 amino acids and a ring-closing ester bond between the C-terminal end of Kyn and the side chain of Thr. The exocyclic portion of daptomycin protrudes from the ring and features a distinct acyl chain attached to Trp. (Figure 30). [103] The exact daptomycin-mediated bactericidal mechanism remains poorly understood and highly speculative. To date it has been suggested that the mode of action involves the interaction between

the gram-positive cell membrane and daptomycin, potentially involving insertion into the cell membrane in a  $\text{Ca}^{2+}$  - dependent action, generating pores and leading to overall membrane depolarization and cell death. It is believed that the pore formation is dependent upon the development of daptomycin aggregates facilitated by free  $\text{Ca}^{2+}$ . [114-117] However, the studies that reported these conclusions were all performed at concentrations much greater than the reported MIC values. It can therefore be speculated that these events could not perceivably take place *in vivo*. Additionally, it has been proposed that daptomycin inhibits peptidoglycan and/or lipoteichoic acid synthesis within the cell although the mechanism is also not understood. Although bacterial resistance remains uncommon, resistance in *S. aureus*, *E. faecium* and *E. faecalis* are appearing during treatment with increasing frequency. [118-120] The exact mechanism of resistance remains unresolved; however, it has been proposed that daptomycin resistance in *S. aureus* could be mediated by increasing the overall positive charge of the surface of the cell instigating an electrostatic repulsion of the daptomycin- $\text{Ca}^{2+}$  complex. The rise in cases of resistant strains prompts interest in developing new daptomycin derivatives to further investigate the mechanism of action as well developing new drug analogues with increased antibiotic activity. Many strategies have been implemented to develop daptomycin analogues including total chemical synthesis, chemoenzymatic synthesis and solid phase approaches, focusing on the alteration of amino acids, acyl chain length and ornithine derivatives. [101, 121-132] In order to generate daptomycin derivatives and maintain the overall potent scaffold of the molecule, we propose to employ a decorator enzyme to perform late-stage modification of the natural product. In this study we will use the aromatic prenyltransferase CdpNPT

from *Aspergill fumigatus* to generate alkyl diversified Trp of daptomycin and evaluate the antibacterial properties.



**Figure 30. Daptomycin structure**

*The structure of daptomycin comprised of 13 (numbered in blue) amino acids.*

### Research Objectives

In this study we sought to test the capability of producing daptomycin derivatives using chemoenzymatic methodology. Specifically, we utilized indole prenyltransferase CdpNPT, known to accept daptomycin as a substrate, to generate daptomycin analogues substituted with diverse alkyl moieties[40]. CdpNPT was found to catalytically transfer alkyl groups onto the N1, C2, C5 or C6 position of the tryptophan residue of daptomycin. The structural analysis and subsequent *in vitro* biological activity assays revealed a SAR in daptomycin susceptible and resistant strains yielding high improvements in the MIC values for some of daptomycin derivatives.

### Contributions

Syntheses of alkyl pyrophosphates were carried out by Dr. Chandrasekhar Bandari, Tejaswi Bavineni, Johanna Masterson, Rachel Tran, and Eric Gardner. Large-scale



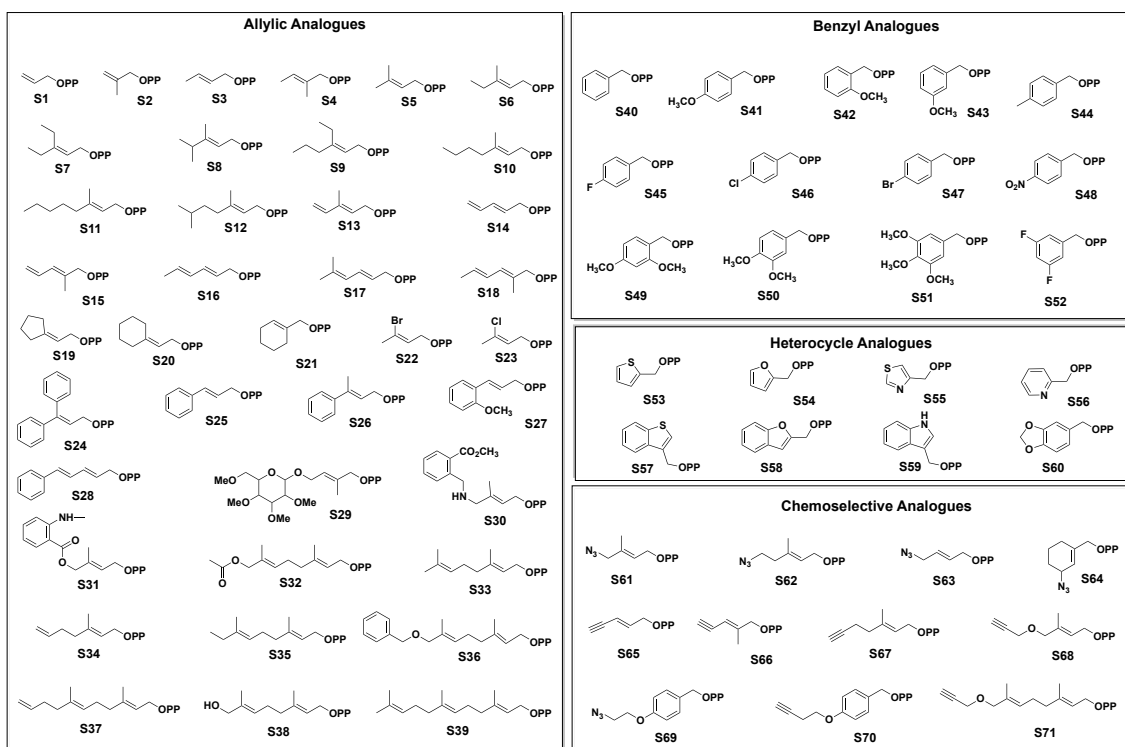
enzymatic reaction and purification of daptomycin analogs were carried out in part by Dr. Singh, Dr. Bandari and me. NMR data collection of daptomycin analogs was done at NMRFAM (Madison, WI) by Dr. Marco Tonelli, NMR data analysis and structural characterization was performed by Dr. Singh, and reconfirmation of NMR assignments was performed by Eric Gardner. Cytotoxicity assay was carried out by Dr. JianLan You. I carried out the expression, purification of CdpNPT, RP-HPLC based assessment, antibacterial activity assays, and calculation of LogP values. Most of the contents in this chapter including text, tables and figures will be directly used for the publication in a peer-reviewed journal.

## **Results and discussion**

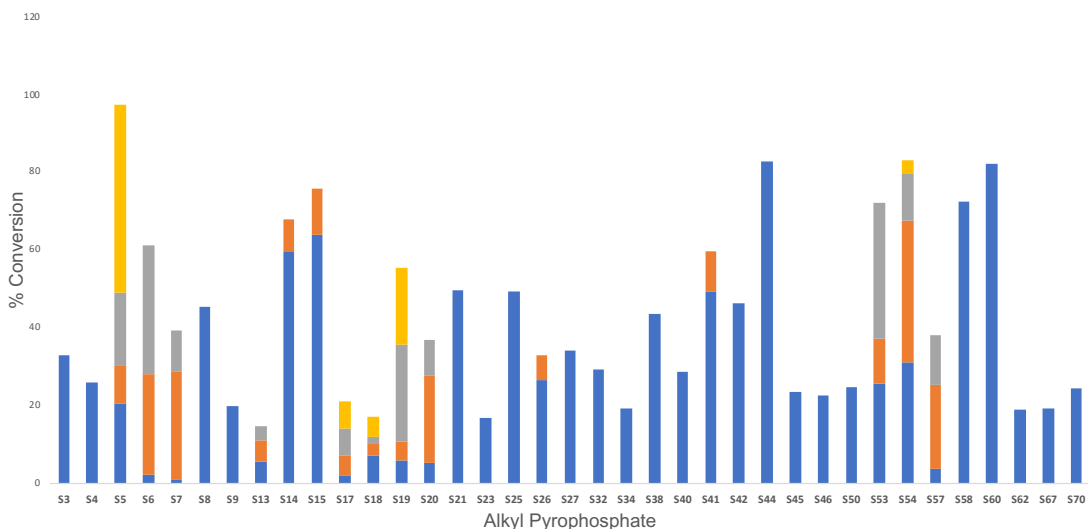
### **Assaying the alkyl pyrophosphate substrate tolerance of CdpNPT to construct daptomycin analogues.**

To explore the ability of CdpNPT to catalyze the differential alkylation of daptomycin, we synthesized a 71-member alkyl pyrophosphate donor library (Figure 31). Specifically, the alkyl pyrophosphate library includes various carbon chain length (C<sub>3</sub>-C<sub>15</sub>), benzylic analogues, heterocyclic analogues, as well as pyrophosphate analogues with chemoselective components such as azides and alkynes. For the assay, recombinant CdpNPT was expressed and soluble in *E. coli* containing a N-terminal His<sub>6</sub> tag and cultured in a total of 4L and purified by Ni-NTA affinity chromatography resulting in a yield of 48.5 mg mL<sup>-1</sup>. Assaying CdpNPT mediated daptomycin alkylation revealed the successful transfer of 37 pyrophosphate analogues, including the previously determined DMAPP, onto daptomycin (Figure 29).[40] Remarkably, the conversion to alkylated daptomycin was >20% for 33 alkyl analogues and >50% for 13

alkyl analogues. Overall, we were able to enzymatically produce daptomycin derivatives featuring DMAPP analogues (S3-S9, S13-S15, S17-21), cinnamyl analogues (S25-S27), GPP analogues (S32, S34, S38), benzyl analogues (S40-S42, S44-S46, S50), heterocycle analogue (S53, S54, S57, S58, S60), and chemoselective analogues with azido (S62) and alkyne groups (S67, S70). Several reactions indicated the formation of multiple product isomers, corresponding to different retention times upon HPLC analysis, with some reactions displaying as many as 4 distinct product peaks (Figure 32). In all cases, product formation of daptomycin derivatives was verified through HPLC and high-resolution LC-MS (Table 8).

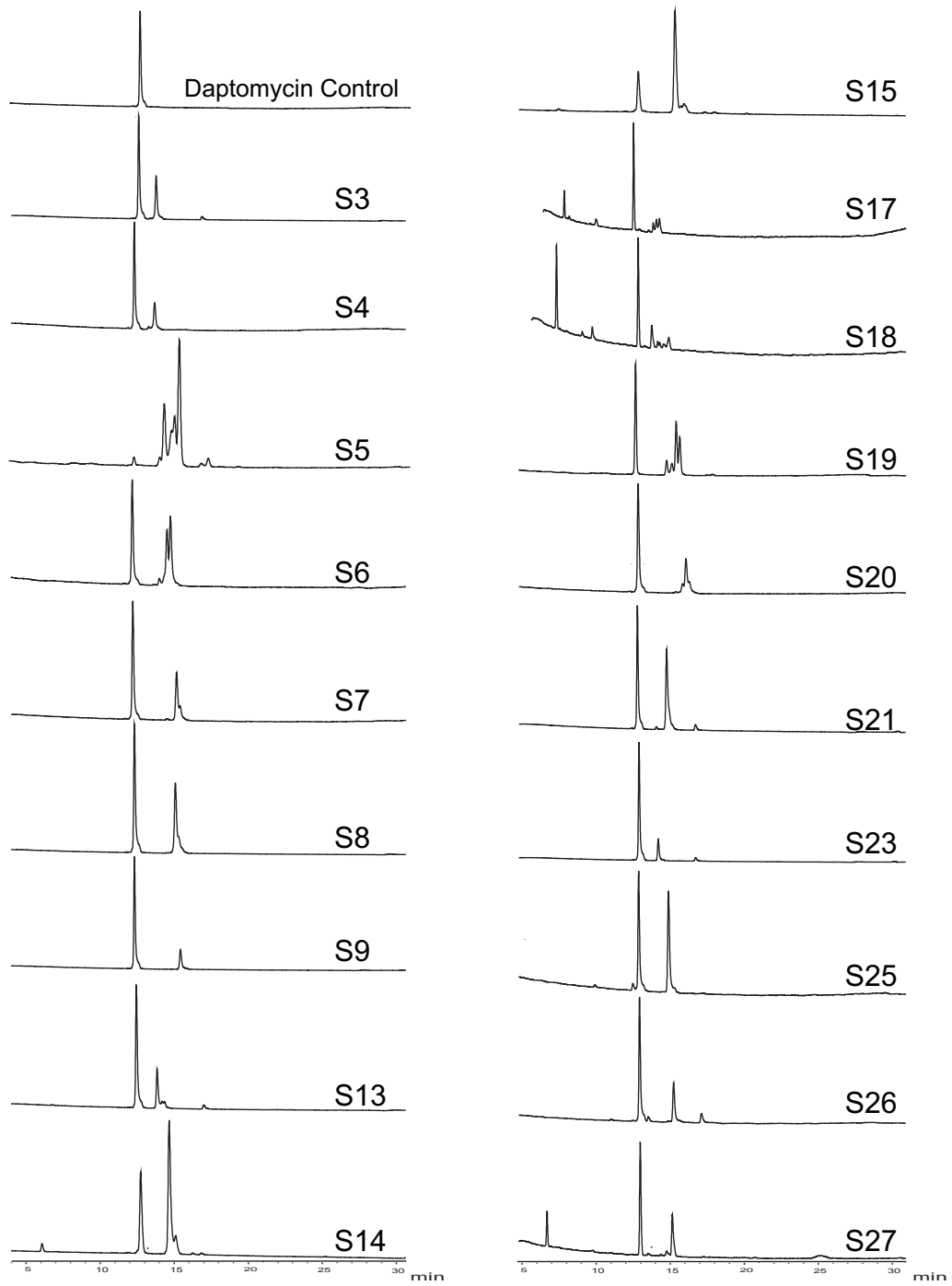


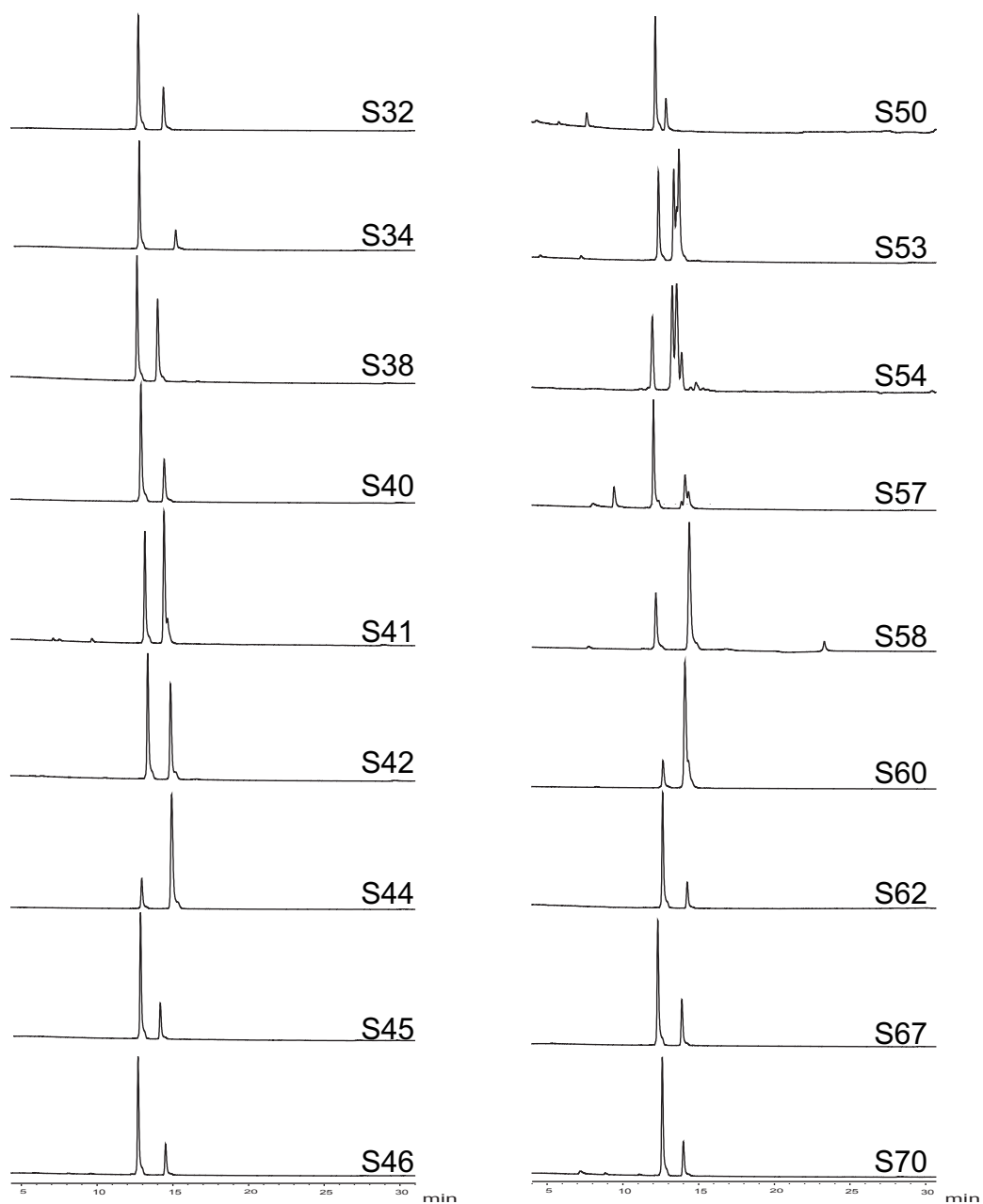
**Figure 31.** Synthetic alkyl pyrophosphate library



**Figure 32. Percent Conversion Daptomycin analogues using CdpNPT**

*The percent conversion of alkyl-pyrophosphate analogues with daptomycin catalyzed by CdpNPT under standard conditions based upon RP-HPLC. The blue, orange, grey and yellow colors correspond to different product peaks as seen on the RP-HPLC chromatogram, Reactions were performed with 1.2 mM alkyl-pyrophosphate analogue, 1 mM daptomycin, CdpNPT, 25 mM Tris, 5 mM CaCl<sub>2</sub>, 50 mM KCl, pH 7.5, 16 h at 35 °C. No product formation was observed in the absence of CdpNPT or alkyl-pyrophosphate analogue.*





**Figure 33. HPLC chromatograms**

*HPLC chromatograms of the CdpNPT catalyzed reaction using synthetic alkyl pyrophosphate library with daptomycin. Reactions were analyzed after incubation under standard conditions for 16hrs at 35 °C*

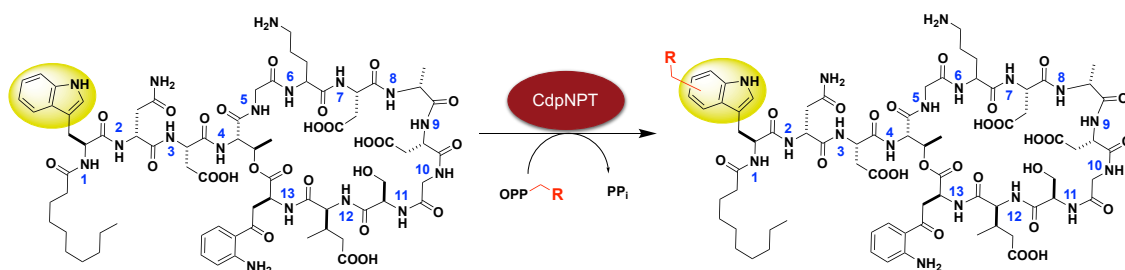
**Table 8 HRMS analytical scale daptomycin analogues**

Enzyme Product	Chemical Formula [M+H] <sup>+</sup>	Calculated Mass	Observed Mass	Retention Time (mins)
<b>S3-DAP</b>	C <sub>76</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub>	1674.7646	1674.7670 <sup>*</sup>	4.499
<b>S4-DAP</b>	C <sub>77</sub> H <sub>109</sub> N <sub>17</sub> O <sub>26</sub>	1687.7724	1688.7753 <sup>*</sup>	4.610
<b>S5-DAP</b>	C <sub>77</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1688.7802	1688.7813 <sup>a*</sup>	4.424
	C <sub>77</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1688.7802	1688.7824 <sup>b*</sup>	4.49
	C <sub>77</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1688.7802	1688.7818 <sup>c*</sup>	4.722
	C <sub>82</sub> H <sub>118</sub> N <sub>17</sub> O <sub>26</sub>	1756.8428	1756.8416 <sup>d#</sup>	4.722
<b>S6-DAP</b>	C <sub>78</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1702.7959	1702.7978 <sup>a*</sup>	4.641
	C <sub>78</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1702.7959	1702.7973 <sup>b*</sup>	4.79
<b>S7-DAP</b>	C <sub>79</sub> H <sub>114</sub> N <sub>17</sub> O <sub>26</sub>	1716.8115	1716.8121 <sup>*</sup>	4.852
<b>S8-DAP</b>	C <sub>79</sub> H <sub>113</sub> N <sub>17</sub> O <sub>26</sub>	1716.8115	1716.8091 <sup>*</sup>	4.783
<b>S9-DAP</b>	C <sub>80</sub> H <sub>115</sub> N <sub>17</sub> O <sub>26</sub>	1730.8272	1730.8261 <sup>*</sup>	4.780
<b>S13-DAP</b>	C <sub>78</sub> H <sub>109</sub> N <sub>17</sub> O <sub>26</sub>	1699.7724	1700.7802	4.508
<b>S14-DAP</b>	C <sub>77</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub>	1686.7646	1686.7638 <sup>a*</sup>	4.564
	C <sub>82</sub> H <sub>114</sub> N <sub>17</sub> O <sub>26</sub>	1752.8115	1752.8094 <sup>b#</sup>	4.713
<b>S15-DAP</b>	C <sub>78</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1700.7802	1700.7830 <sup>a*</sup>	4.431
	C <sub>78</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1700.7802	1700.7823 <sup>b*</sup>	4.762
<b>S17-DAP</b>	C <sub>79</sub> H <sub>113</sub> N <sub>17</sub> O <sub>26</sub> [M+2H] <sup>2+</sup>	857.9016	857.9004 <sup>*</sup>	4.688
<b>S18-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7982 <sup>a*</sup>	4.578
	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7984 <sup>b*</sup>	4.752
<b>S19-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7935	4.750
<b>S20-DAP</b>	C <sub>80</sub> H <sub>114</sub> N <sub>17</sub> O <sub>26</sub>	1728.8115	1728.8111 <sup>*</sup>	4.835
<b>S21-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7970 <sup>a*</sup>	4.538
	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7995 <sup>b*</sup>	4.669
	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1714.7959	1714.7986 <sup>c*</sup>	4.794
<b>S25-DAP</b>	C <sub>81</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1736.7802	1736.7806 <sup>*</sup>	4.706
<b>S26-DAP</b>	C <sub>82</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub>	1750.7959	1750.7978 <sup>*</sup>	4.734
<b>S27-DAP</b>	C <sub>82</sub> H <sub>112</sub> N <sub>17</sub> O <sub>27</sub>	1766.7908	1766.7914 <sup>*</sup>	4.366
<b>S34-DAP</b>	C <sub>80</sub> H <sub>113</sub> N <sub>17</sub> O <sub>26</sub>	1728.8115	1728.8067 <sup>*</sup>	4.756
<b>S38-DAP</b>	C <sub>82</sub> H <sub>118</sub> N <sub>17</sub> O <sub>27</sub>	1773.8456	1773.8328	4.521
<b>S40-DAP</b>	C <sub>79</sub> H <sub>107</sub> N <sub>17</sub> O <sub>26</sub>	1710.7646	1710.7569 <sup>*</sup>	4.536
<b>S41-DAP</b>	C <sub>80</sub> H <sub>110</sub> N <sub>17</sub> O <sub>27</sub>	1740.7752	1740.7767 <sup>a*</sup>	4.41
	C <sub>80</sub> H <sub>110</sub> N <sub>17</sub> O <sub>27</sub>	1740.7752	1740.7749 <sup>b*</sup>	4.617
<b>S42-DAP</b>	C <sub>80</sub> H <sub>110</sub> N <sub>17</sub> O <sub>27</sub>	1740.7752	1740.7669 <sup>*</sup>	4.565
<b>S44-DAP</b>	C <sub>80</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub>	1724.7802	1724.7731 <sup>*</sup>	4.687
<b>S45-DAP</b>	C <sub>79</sub> H <sub>106</sub> FN <sub>17</sub> O <sub>26</sub>	1728.7552	1728.7498 <sup>*</sup>	4.517
<b>S50-DAP</b>	C <sub>81</sub> H <sub>111</sub> N <sub>17</sub> O <sub>28</sub>	1770.7857	1770.7799 <sup>*</sup>	4.230
<b>S54-DAP</b>	C <sub>77</sub> H <sub>105</sub> N <sub>17</sub> O <sub>27</sub>	1700.7439	1700.7363	4.414
<b>S57-DAP</b>	C <sub>81</sub> H <sub>107</sub> N <sub>17</sub> O <sub>26</sub> S	1766.7367	1766.7290 <sup>*</sup>	4.730

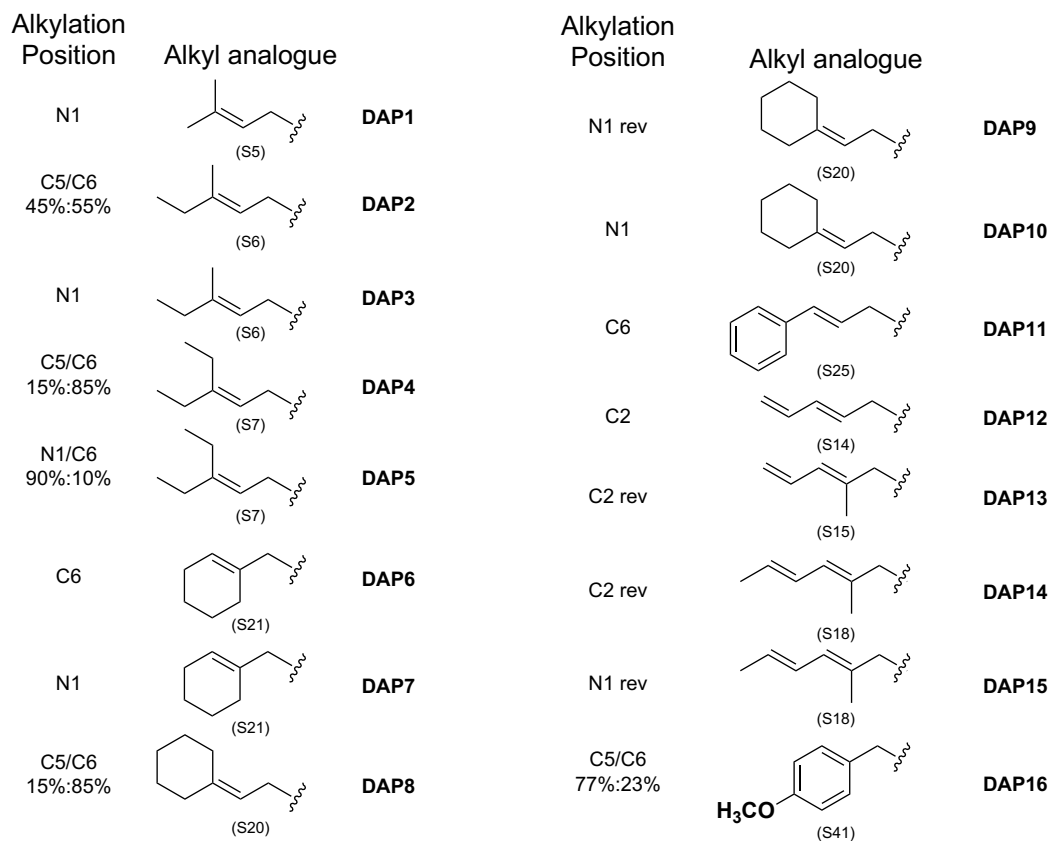
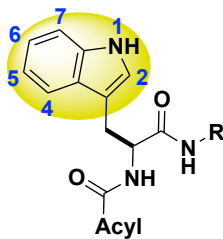
<b>S58-DAP</b>	C <sub>81</sub> H <sub>107</sub> N <sub>17</sub> O <sub>27</sub>	1750.7595	1750.7522*	4.708
<b>S60-DAP</b>	C <sub>80</sub> H <sub>107</sub> N <sub>17</sub> O <sub>28</sub>	1754.7544	1754.7481*	4.459
<b>S62-DAP</b>	C <sub>78</sub> H <sub>111</sub> N <sub>20</sub> O <sub>26</sub>	1743.7973	1743.7892*	4.621

### Derivatization and determination of bioactivity of daptomycin.

The analytical reactions described above suggest CdpNPT is a permissive and effective alkylation catalyst towards daptomycin, which lead us into the investigation of antibacterial potency of the daptomycin analogues. Therefore, to explore the significance of the derivatization of daptomycin we carried out semi preparative-scale reactions involving 10 alkyl pyrophosphates (**S5**, **S6**, **S7**, **S14**, **S15**, **S18**, **S20**, **S21**, **S25**, **S41**). The reactions were purified using reverse phase HPLC where each product peak was separated based on retention times. The resulting purified products were subjected to ESI-HMRS and NMR analysis for structure elucidation (Table 8; Figure 35). NMR analysis confirmed N and C alkylation was performed solely on the tryptophan residue of daptomycin, as previously reported.[40] Additionally, several regio- and structural isomers were identified amongst the purified compounds, some of which were inseparable isomeric mixtures. Ultimately, the semi-preparative purification resulted in 16 distinct daptomycin analogues (**DAP1-DAP16**) involving N1, C2, C5, C6 alkylation of the indole ring (Figure 34 and 37).



**Figure 34. CdpNPT catalyzed reaction producing daptomycin derivatives**  
*CdpNPT catalyzes the alkylation of daptomycin by diversifying the tryptophan (colored yellow) found on the exocycle portion of the compound*



**Figure 35.** Daptomycin derivatives and placement on the indole ring. Shown are the 16 different structurally elucidated daptomycin derivatives. Placement on the tryptophan was determined through  $^1\text{H}$  and 2D NMR analysis. In some cases, an inseparable mixture of isomers was found.



**Table 9. HRMS data of purified alkyl daptomycin analogues**

Enzyme Product	Chemical Formula	Calculated Mass	Observed Mass
<b>N1-S5-DAP</b>	C <sub>77</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1686.7651	1686.7659
<b>N1-S6-DAP</b>	C <sub>78</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1700.7808	1700.7818
<b>C5-S6-DAP: C6-S6-DAP</b>	C <sub>78</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1700.7808	1700.7816*
<b>N1-S7-DAP: C6-S7-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1714.7964	1714.7978*
<b>C5-S7-DAP: C6-S7-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1714.7964	1714.7979*
<b>N1-S21-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M+H] <sup>+</sup>	1714.7964	1714.7970
<b>C6-S21-DAP</b>	C <sub>79</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M+H] <sup>+</sup>	1714.7964	1714.7995
<b>N1<sup>R</sup>-S20-DAP</b>	C <sub>80</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1726.7964	1726.7957
<b>N1-S20-DAP</b>	C <sub>80</sub> H <sub>112</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1726.7964	1726.7956
<b>C5-S20-DAP: C6-S20-DAP</b>	C <sub>80</sub> H <sub>114</sub> N <sub>17</sub> O <sub>26</sub> [M+H] <sup>+</sup>	1728.8121	1728.8111*
<b>C6-S25-DAP</b>	C <sub>81</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1734.7651	1734.7659
<b>C2-S14-DAP</b>	C <sub>77</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub> [M+H] <sup>+</sup>	1686.7651	1686.7638
<b>C2<sup>R</sup>-S15-DAP</b>	C <sub>78</sub> H <sub>108</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1698.7651	1698.7605
<b>N1<sup>R</sup>-S18-DAP</b>	C <sub>79</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1712.7808	1712.7814
<b>C2<sup>R</sup>-S18-DAP</b>	C <sub>79</sub> H <sub>110</sub> N <sub>17</sub> O <sub>26</sub> [M-H] <sup>-</sup>	1712.7808	1712.7800
<b>C5-S41-DAP: C6-S41-DAP</b>	C <sub>80</sub> H <sub>108</sub> N <sub>17</sub> O <sub>27</sub> [M-H] <sup>-</sup>	1738.7601	1738.7578*
<b>DAP</b>	C <sub>72</sub> H <sub>102</sub> N <sub>17</sub> O <sub>26</sub> [M+H] <sup>+</sup>	1620.7182	1620.7177

\*Inseparable mixture of isomers.

**DAP1-DA16** were evaluated for antibacterial activity against 4 bacterial species, *Staphylococcus aureus*, *Bacillus subtilis*, *Enterococcus faecalis* and *Staphylococcus epidermidis*, including 12 clinical isolates using the micro dilution method (see experimental methods) (Table 10). To assess the possibility of improved antibiotic susceptibility using daptomycin derivatives, three daptomycin resistant strains were also tested, two transposon mutants, *S. aureus* (SAUSA300 1713 NE573, SAUSA300 1715 NE1656) as well as hospital acquired *E. faecalis* S712. *S. aureus* ATCC 25923 was used as a quality control strain and tested with daptomycin to ensure proper techniques and analysis. The MIC value for the parent compound was 0.6 µg ml<sup>-1</sup> as recommended by the CLSI and FDA (0.12-1.0 µg mL<sup>-1</sup>). [133, 134] In order to gauge the relative

antibiotic activity, daptomycin was also evaluated against all strains used in this study and compared against MICs for derivatives (Figure 36). Daptomycin showed an MIC range of 0.3-16.2 $\mu\text{g mL}^{-1}$  against the clinical isolates, not including daptomycin resistant strains. For the daptomycin resistant strains, the parent compound displayed an MIC range 16.2->32.4 $\mu\text{g mL}^{-1}$ , consistent with the noted high resistance. C-2 substituted compounds, such as **DAP12**, **DAP13** and **DAP14** display the lowest antibacterial activity with MICs ranging from 4.2 - >33.7 $\mu\text{g mL}^{-1}$ , 4.2 - >34.0 $\mu\text{g mL}^{-1}$  and 2.1 - > 34.3  $\mu\text{g mL}^{-1}$ , respectively. Amongst the compounds with highest antibacterial activity are C5 and C6 substituted, including **DAP2**, **DAP4**, **DAP5**, **DAP6**, and **DAP8** with MICs ranging from 0.1-8.5 $\mu\text{g mL}^{-1}$ , 0.5-2.7 $\mu\text{g mL}^{-1}$ , 0.3-4.3 $\mu\text{g mL}^{-1}$ , 0.1- 6.2 $\mu\text{g mL}^{-1}$ , and 0.2-2.2 $\mu\text{g mL}^{-1}$ , respectively (Table 10). Interestingly, **DAP6** and **DAP7** are both substituted with alkyl analogue S21 at different positions on the indole ring. **DAP6**, a C6 substituted compound, displayed up to an 80-fold MIC improvement compared to N1 substituted **DAP7**. To assess the effect of the lipophilicity of each derivative towards the antibacterial activity, logP values of daptomycin and **DAP1**-**DAP16** were plotted against their corresponding MIC relative to daptomycin MIC[135] (Figure 37). As anticipated, the addition of the alkyl group increases the logP value from -0.47 (daptomycin) to 0.25 (**DAP11**). A correlation between increasing logP values of the daptomycin derivatives and overall MIC improvement is most strongly seen with *E. faecalis* S613 and daptomycin resistant strains. **DAP2**, **DAP4**, **DAP6** and **DAP8** display an impressive 15 – 80-fold improvement over the parent compound. As seen in the LogP correlation plots, LogP values are a characteristic of both the alkyl group as well as the position on the indole ring (Figure 37). C-2 substituted analogues

exhibit the lowest logP values (-0.2 – -0.1) while C-5 and C-6 gave rise to the highest logP values (-0.17 – 0.25). A comparison of MIC and logP for compounds with high logP values (0.0 – 0.25), such as **DAP11**, **DAP8**, **DAP16**, and **DAP4**, did not show any direct correlation. However, when tested with the resistant strains **DAP16** and **DAP11** had lower improvement despite being C-5/C-6 substituted with logP values similar to the most effective analogues. Overall, the properties of the alkyl group and the alkylation position on the indole ring influences the antibacterial activity of the daptomycin derivatives. These results imply the aromatic moiety could potentially be contributing to the bioactivity of these compounds. Notably, the bacterial strains respond differently to the DAP analogues either indicating a different mechanism of action or a different cellular environment.

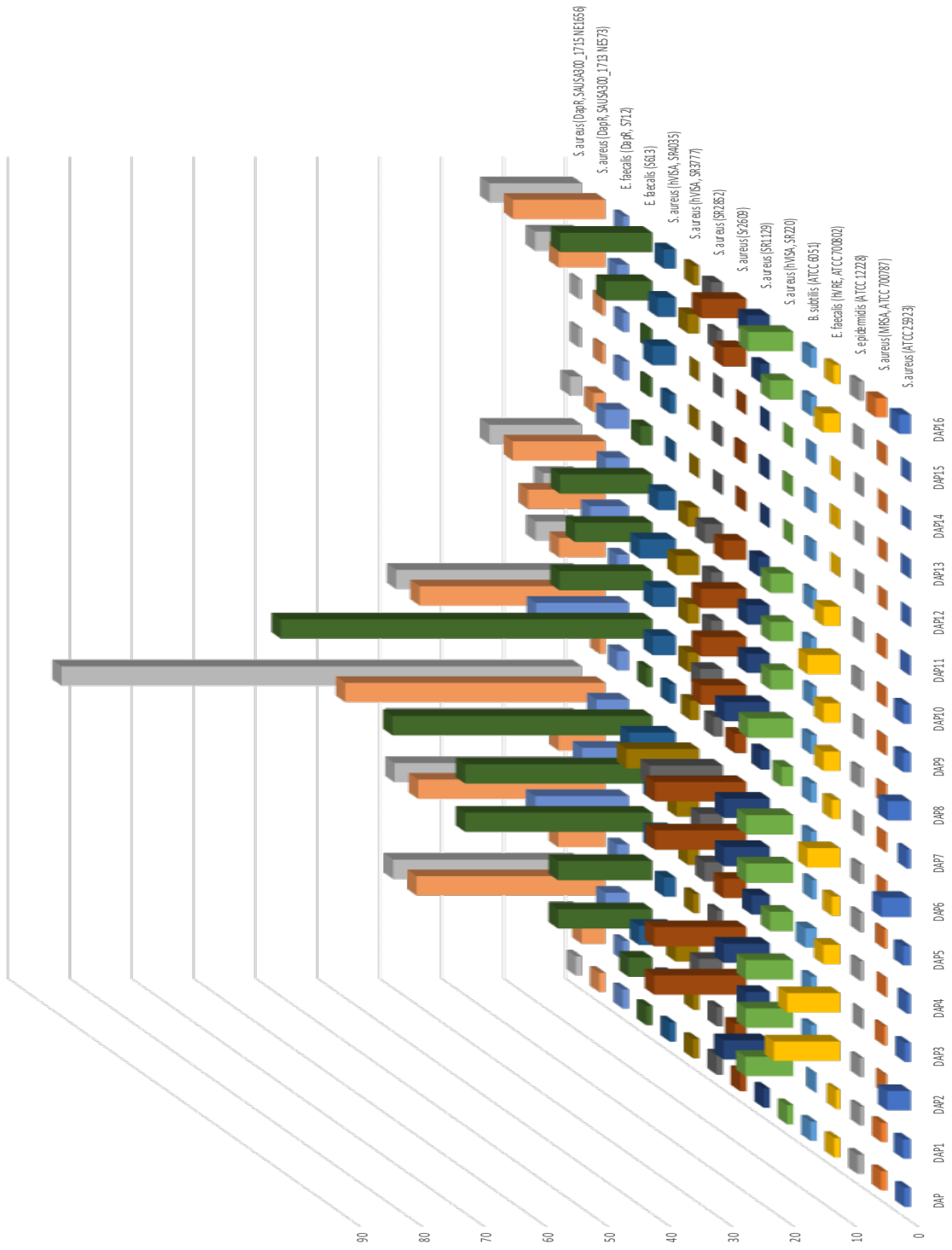
**Table 10.** In-vitro MIC ( $\mu\text{g mL}^{-1}$ ) values for Daptomycin and daptomycin derivatives.

MIC ( $\mu\text{g mL}^{-1}$ ) values for Daptomycin and Daptomycin derivatives	DAP	DAP1 Trp-N1-S5	DAP2 Trp-C5/C6-S6 (45%:55%)	DAP3 Trp-N1-S6	DAP4 Trp-C5/C6-S7 (15%:85%)	DAP5 Trp-N1/C6-S7 (90%:10%)	DAP6 Trp-C6-S21	DAP7 Trp-N1-S21	DAP8 Trp-C5/C6 (15%:85%)
<i>S. aureus</i> (ATCC 25923)	0.6	0.5	0.2	0.7	1.1	0.5	0.1	1.1	0.2
<i>S. aureus</i> (MRSA, ATCC 700787)	0.6	0.7	2.1	1.2	2.7	1.4	2.1	2.7	2.2
<i>S. epidermidis</i> (ATCC 12228)	0.3	0.5	0.5	1.2	0.5	0.5	0.5	1.1	0.5
<i>E. faecalis</i> (hVRE, ATCC 700802)	1.5	2.1	0.1	0.2	0.5	1.1	0.3	1.1	0.5
<i>B. subtilis</i> (ATCC 6051)	2.0	8.4	2.1	2.1	1.1	2.7	2.1	2.1	2.2
<i>S. aureus</i> (hVISA, SR220)	4.0	0.5	0.5	0.5	1.1	0.5	0.5	2.1	0.6
<i>S. aureus</i> (SR1129)	2.0	0.3	0.5	0.3	0.7	0.3	0.3	1.4	0.3
<i>S. aureus</i> (SR2609)	4.0	2.1	0.3	0.3	1.1	0.3	0.3	2.1	0.6
<i>S. aureus</i> (SR2852)	2.0	2.1	0.5	2.1	0.7	0.5	0.2	1.4	0.6
<i>S. aureus</i> (hVISA, SR3777)	2.0	2.1	0.5	2.1	1.1	0.5	0.2	1.4	1.1
<i>S. aureus</i> (hVISA, SR4035)	8.1	4.2	1.4	4.3	2.1	2.1	1.1	8.6	2.2
<i>E. faecalis</i> (S613)	16.2	4.2	1.1	1.1	0.5	0.5	0.4	17.1	0.3
<i>E. faecalis</i> (DapR, S712)	>32.4	33.7	8.5	17.0	2.1	4.3	6.2	>17.1	2.2
<i>S. aureus</i> (DapR, SAUSA300_1713 NE573)	16.2	4.2	0.5	2.1	0.5	2.1	0.4	17.1	0.5
<i>S. aureus</i> (DapR, SAUSA300_1715 NE1656)	16.2	4.2	0.5	2.1	0.5	0.5	0.2	17.1	0.5

MIC ( $\mu\text{g mL}^{-1}$ ) values for Daptomycin and Daptomycin derivatives	DAP9 Trp-N1rev-S20	DAP10 Trp-N1-S20	DAP11 Trp-C6-S25	DAP12 Trp-C2-S14	DAP13 Trp-C2rev-S15	DAP14 Trp-C2rev-S18	DAP15 Trp-N1rev-S18	DAP16 Trp-C5/C6-S41 (77%:23%)
<i>S. aureus</i> (ATCC 25923)	0.5	0.5	2.8	16.9	8.5	>34.3	17.1	0.3
<i>S. aureus</i> (MRSA, ATCC 700787)	2.8	2.8	2.4	>33.7	>34.0	>34.3	4.3	0.3
<i>S. epidermidis</i> (ATCC 12228)	1.1	0.7	1.1	16.9	8.5	34.3	1.1	0.4
<i>E. faecalis</i> (hVRE, ATCC 700802)	0.5	0.3	0.5	33.7	8.5	34.3	0.5	1.4
<i>B. subtilis</i> (ATCC 6051)	2.8	2.2	2.8	4.2	4.2	8.6	2.1	2.2
<i>S. aureus</i> (hVISA, SR220)	1.1	1.1	1.1	33.7	17.0	>34.3	1.1	0.6
<i>S. aureus</i> (SR1129)	0.6	0.6	1.1	16.9	8.5	34.3	1.4	0.6
<i>S. aureus</i> (SR2609)	0.6	0.6	1.1	16.9	8.5	34.3	1.1	0.6
<i>S. aureus</i> (SR2852)	1.1	1.1	0.7	16.9	8.5	34.3	2.1	1.1
<i>S. aureus</i> (hVISA, SR3777)	1.1	0.6	1.1	16.9	8.5	34.3	1.1	2.2
<i>S. aureus</i> (hVISA, SR4035)	2.2	1.4	2.8	>33.7	8.5	2.1	2.7	4.2
<i>E. faecalis</i> (S613)	1.1	1.3	1.1	>8.4	>34.0	>34.3	2.1	1.1
<i>E. faecalis</i> (DapR, S712)	17.3	5.2	8.7	>8.4	>34.0	>34.3	17.1	>34.3
<i>S. aureus</i> (DapR, SAUSA300_1713 NE573)	2.2	1.3	1.1	>8.4	>34.0	>34.3	2.1	1.1
<i>S. aureus</i> (DapR, SAUSA300_1715 NE1656)	2.2	2.6	1.1	>8.4	>34.0	>34.3	2.1	1.1

\* All strains were acquired from BPEI Resources

Daptomycin analogue improvement

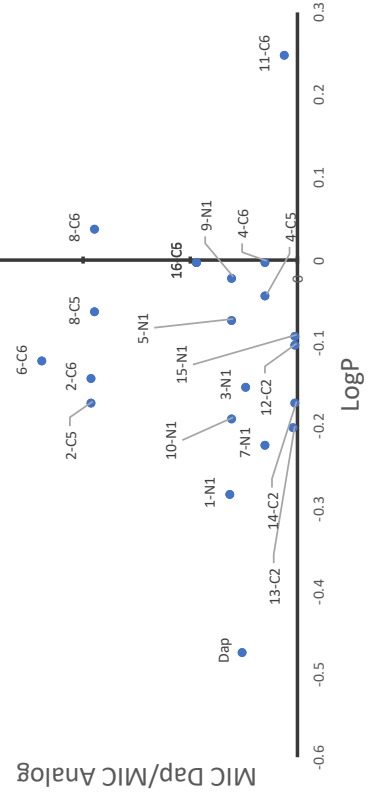


**Figure 36. MIC improvement of Daptomycin analogues compared to daptomycin.**

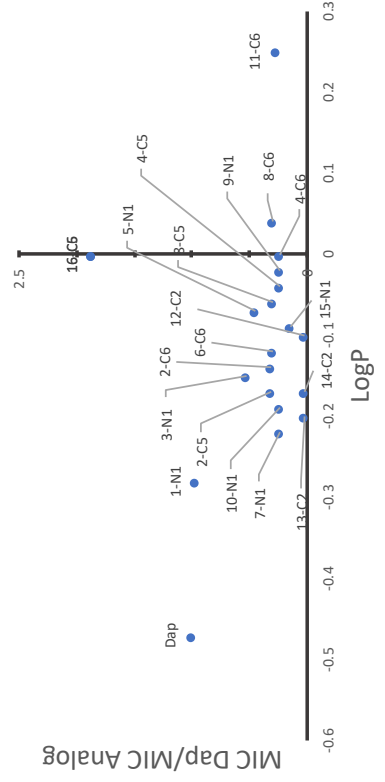
*MIC values from daptomycin derivatives were compared to daptomycin for each*

*bacterial strain. Fold improvement was found by the ratio of MIC daptomycin/ MIC daptomycin derivatives; MIC values found in **Table 10**.*

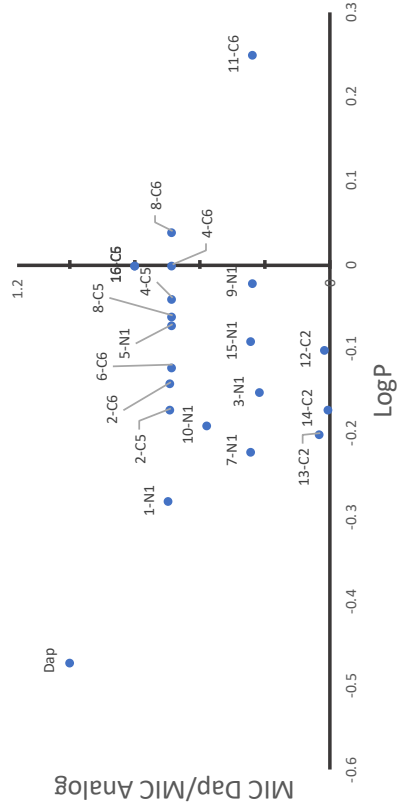
*S. aureus* (ATCC 25923)



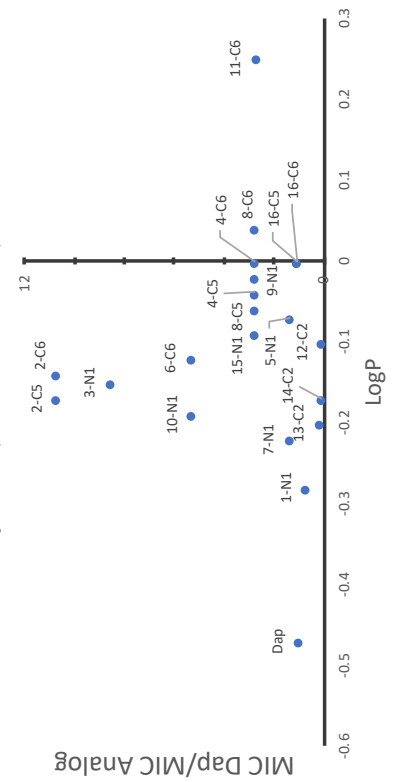
*S. aureus* (MRSA, ATCC 700787)



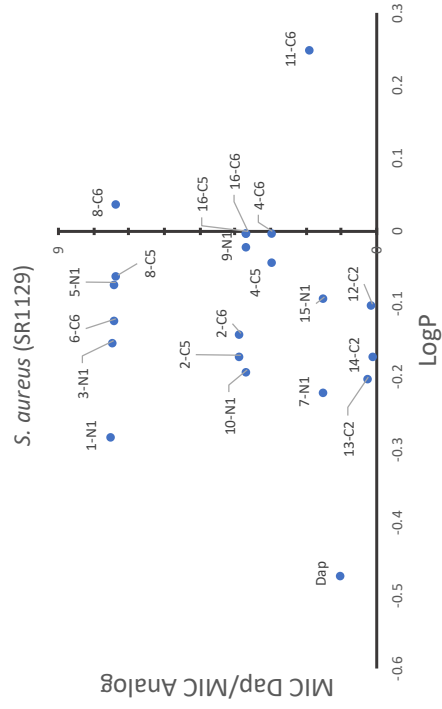
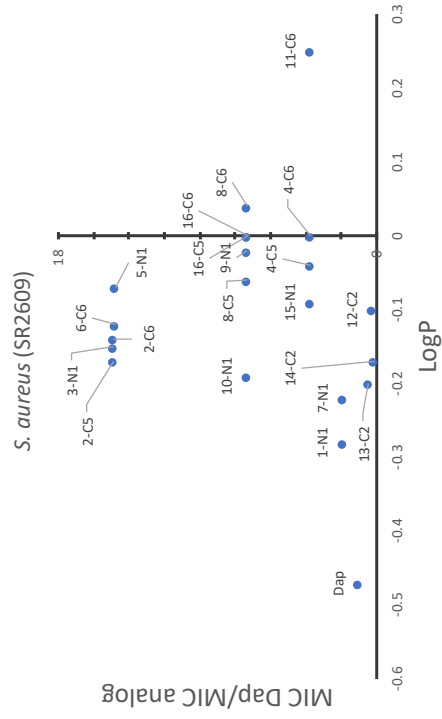
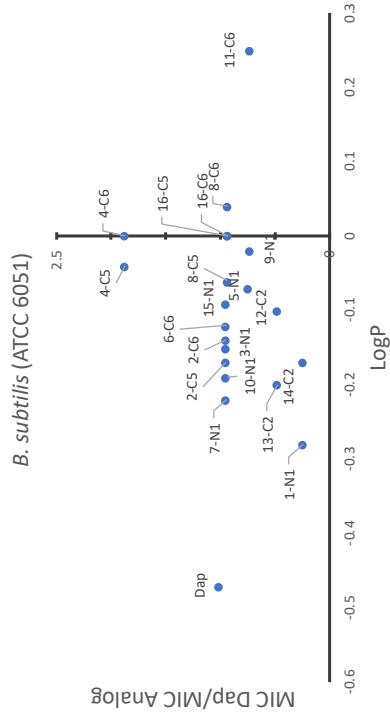
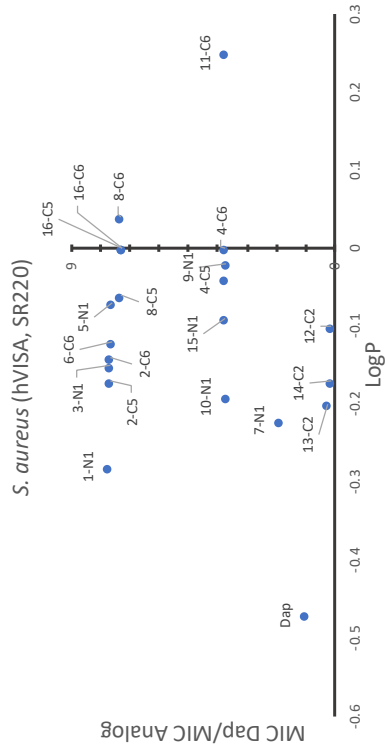
*S. epidermidis* (ATCC 12228)

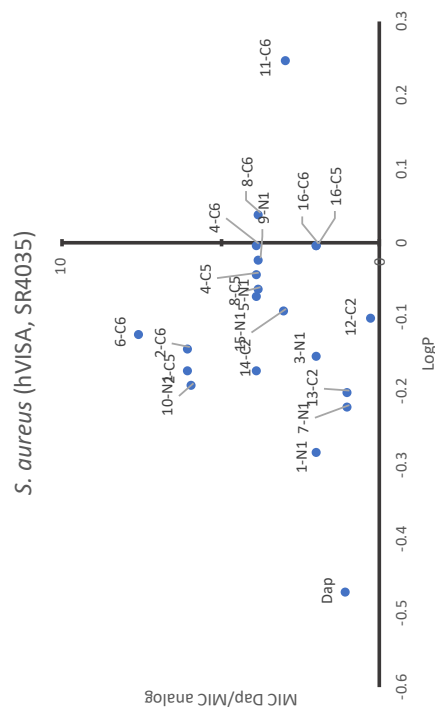
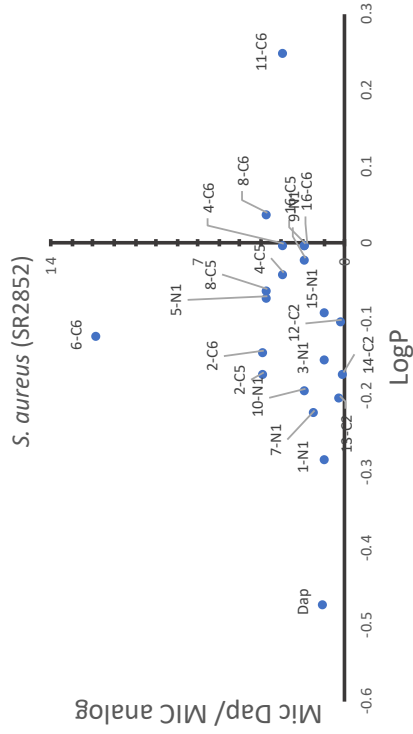
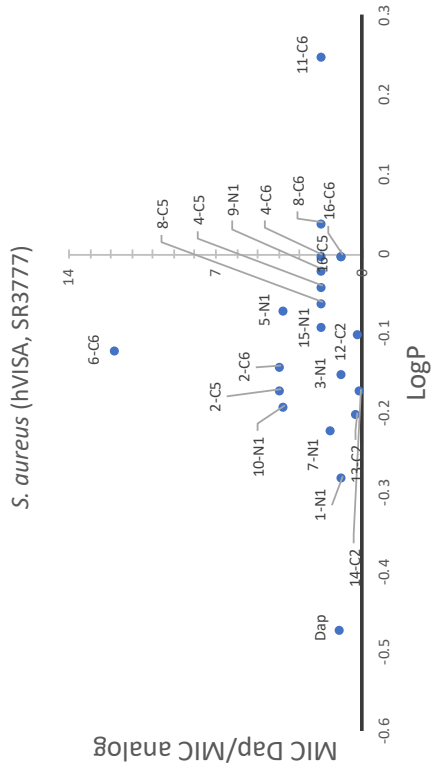


*E. faecalis* (hvRE, ATCC 700802)









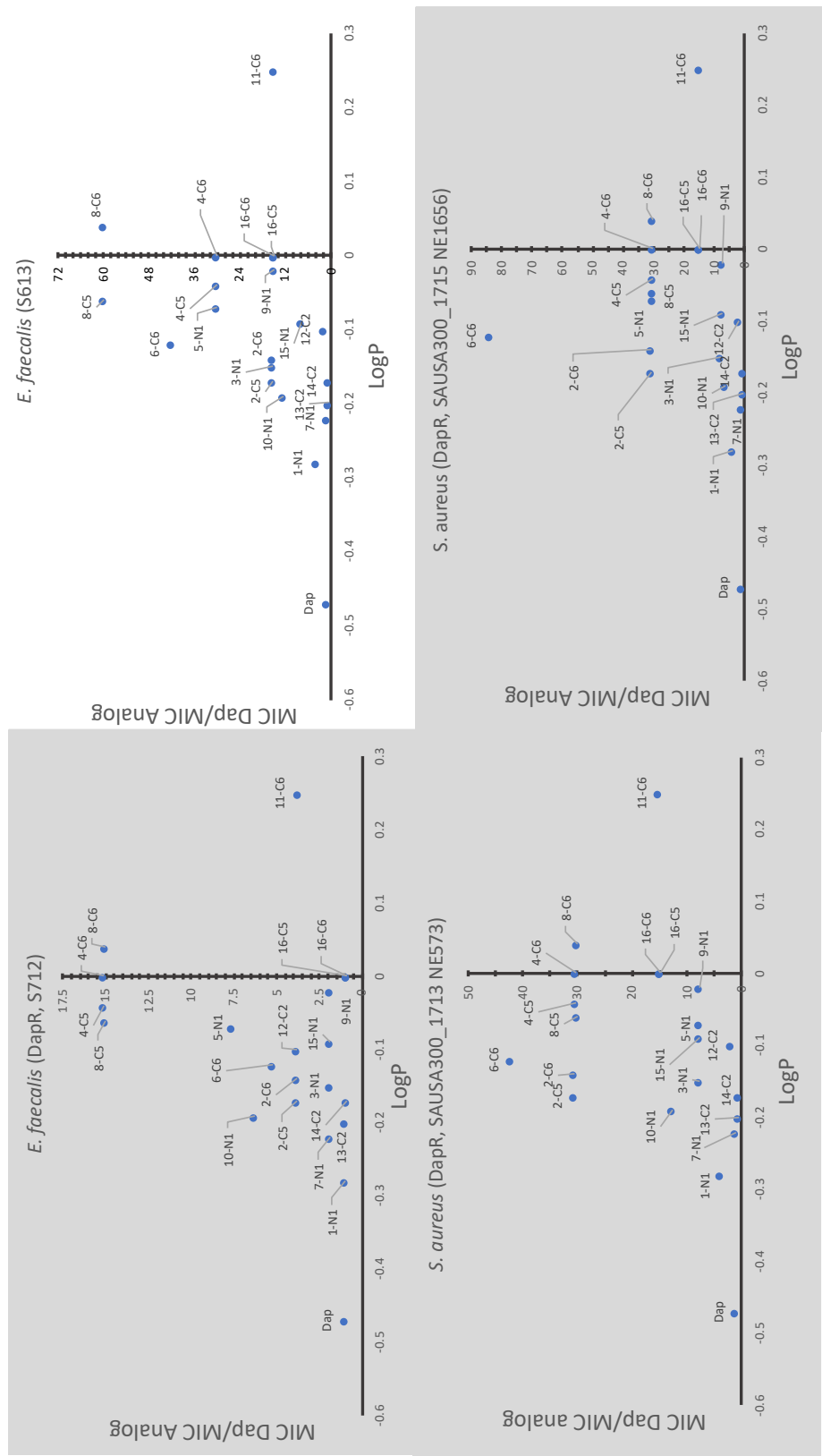


Figure 37. LogP vs MIC improvement plots for daptomycin and daptomycin analogues.

*LogP values for daptomycin and daptomycin derivatives were correlated to their MIC improvement as compared to daptomycin for each bacterial strain. Gray plots represent the daptomycin resistant strains. LogP values were calculated using ALOGPS.[135]*

Nonribosomal peptides represent a highly promising chemical space with great potential for pharmaceutical relevance and medical advancement. Among these compounds, daptomycin, a clinically approved antibiotic, represents a unique class of macrolactone antibiotics. Due to the fact that daptomycin is a highly complex molecule to approach synthetically, the chemoenzymatic strategy that we describe is a successful methodology to generate tryptophan substituted daptomycin derivatives.

Our chemoenzymatic strategy involves the utilization of the highly permissive prenyltransferase CdpNPT which uniformly transfers an alkyl group onto the indole moiety of daptomycin. Through this strategy we have revealed the transfer of a large variety of alkyl groups onto the C5, C6, C2 and N1 position of the indole ring. As reported in Table 8, the position of the substitution has a crucial influence on the antibacterial behavior of the derivatized compounds. The C-2 substituted variants display a significant increase of MIC values (up to 80-fold) compared to the N1, C-5 and C-6 substituted compounds. Based on the results of the logP values, the alkylation position as well as the substituent affects the overall lipophilicity of the compound.

In conclusion, through our chemoenzymatic strategy we were able to specifically modify the tryptophan of the highly chemically complex natural product daptomycin. These daptomycin derivatives show great improvement of MIC values over the parent

compound towards clinically isolated strains including highly relevant daptomycin resistant strains. Many issues that have arisen with daptomycin could potentially be addressed by further synthesis and evaluation of daptomycin analogues such as the ones generated here.

## **Experimental**

**General methods.** HPLC analysis was accomplished using Agilent Technologies 1220 Infinity LC system equipped with a diode array detector. *Method A (HPLC):* Monitoring CdpNPT catalyzed reaction was completed using an analytical reverse-phase (RP) HPLC Gemini-NX, C-18 (5  $\mu\text{m}$ , 4.6 mm  $\times$  250 mm) column (Phenomenex, Torrance, California, USA) employing a gradient of 10% B to 55% B over 10 min, 55% B to 100%B for 15 min, 10%B for 5 min (A=H<sub>2</sub>O with 0.1% TFA; B= acetonitrile), flow rate = 1mL min<sup>-1</sup> ; A<sub>280</sub>. Reaction progression was monitored by the different retention times between product and starting material. *Method B (HPLC):* Semi-preparative RP HPLC was conducted using a Gemini-NX, C-18 (5  $\mu\text{m}$ , 10  $\times$  250 mm) column (Phenomenex, Torrance, California, USA), peak separation and product purification was completed using a shallow gradient of 35% B to 55% B in 20 min, 55% B to 100% B in 3 min, 35%B for 7 min. (A=H<sub>2</sub>O with 0.05% formic acid; B= acetonitrile) flow rate = 2 mL min<sup>-1</sup>; A<sub>280</sub>. High-resolution mass spectrometric (HRMS) data and liquid chromatography mass spectrometric (LCMS) were obtained on Agilent 6545-QTOF W/1290 HPLC mass spectrometer at the University of Oklahoma, Department of Chemistry and Biochemistry. All logP values were calculated using ALOGPS software found in virtual computational chemistry laboratory[135]

**Prenyltransferase CdpNPT expression and purification.** The synthetic gene for CpdNPT, bearing an N-terminal His<sub>6</sub> tag, was inserted into vector pET-28a and transformed into *E. coli* Rosetta cells. After overnight growth, 3 mL of bacterial culture was used to inoculate 1.2 L Luria-Berani medium containing 50 µg kanamycin mL<sup>-1</sup>. Cultures were grown at 37°C for approximately 4 h at 220 r.p.m. (O. D<sub>600</sub> = 0.6-0.8). Protein expression was induced by the addition of isopropyl β-D-1-thiogalactopyranoside (IPTG) (0.5 mM final) and continually shaken at 220 r.p.m. for an additional 16-19 h at 20°C. Bacterial cultures were harvested by centrifugation and the resulting cell pellets were resuspended in a lysis buffer (50mM NaH<sub>2</sub>PO<sub>4</sub>, 200mM NaCl, pH 7.8) and stored at -80°C until use. Resuspended cells were thawed then lysed by way of sonication on ice 60 times for 10 s. In order to separate insoluble and undesirable cellular components, lysed cells were centrifuged for 1 h at 10°C at 16000 r.p.m. The supernatant, containing the expressed recombinant CdpNPT with an N-terminal His<sub>6</sub> tag, was purified using nickel sepharose resin affinity chromatography column (GE Healthcare, USA). Protein was eluted using an imidazole gradient from 10mM to 500mM in elution buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 200 mM NaCl, pH 7.8). The purified protein was concentrated and introduced to a new buffer condition (25 mM Tris-HCl, pH 8, 50 mM KCl) using a desalting column (PD-10 desalting column, GE healthcare). Eluents were evaluated by SDS-PAGE for purity, concentration was determined using NanoDrop One (Thermo Scientific, USA) and the purified protein was stored flash frozen in 20µL aliquots at -80°C.

***In-vitro* CpdNPT assay with daptomycin.** *In vitro* CdpNPT analytical reactions with daptomycin were performed in a final volume of 20µL, consisting of 1.2mM

pyrophosphate analogues, 1mM daptomycin (TCI, TCI America) and 8 $\mu$ M purified CdpNPT in reaction buffer (Tris-HCl pH 7.8, 5 mM CaCl<sub>2</sub>, 50mM KCl) and incubated at 35 °C for 16 h. Reactions were terminated by addition of 60 $\mu$ L cold methanol followed by centrifugation (10,000 g for 30 min) in order to remove precipitated enzyme. Analysis of the reactions was completed on RP-HPLC using method A (general method). For each reaction the percent conversion was calculated by the integration of the product peak at 280nm divided by the overall sum of the integrated product peak and starting material at 280nm. All enzymatic products were confirmed by using ESI-HRMS. Large scale reactions of daptomycin analogues were performed on a 10 mL scale consisting of 1mM daptomycin, 1.25mM pyrophosphate analogue, and 100 $\mu$ M purified CdpNPT in reaction buffer. Putative products were subsequently isolated by semi-prep HPLC using method B (general method, above) and confirmed by NMR and ESI-HRMS.

**Antibacterial assays.** All daptomycin analogue stocks were calibrated by absorbance (Calibration curve at  $\lambda$ = 280nm,  $\epsilon_{366}$  = 4,000 L mol<sup>-1</sup> cm<sup>-1</sup>)[101] and all bioactivity assays were conducted in triplicate. The strains for which MICs were determined included, thirteen daptomycin sensitive (**DapS**) and three daptomycin resistant (**DapR**) strains. The **DapS** strains are: *S. aureus* (ATCC 25923), methicillin-resistant *S. aureus* ATCC 700787, *S. epidermidis* ATCC 12228, vancomycin-resistant *Enterococcus faecalis* ATCC 700802, *Bacillus subtilis* (ATCC 6051), hetero-vancomycin intermediate *S. aureus* SR220 (BEI product NR-50512), *S. aureus*, Strain SR1129 (NR-50506), *S. aureus*, Strain SR2609 (NR-50507), *S. aureus*, Strain SR2852 (NR-50508),

hetero-vancomycin intermediate *S. aureus* SR3777 (BEI product NR-50509), hetero-vancomycin intermediate *S. aureus*, SR4035 (BEI product NR-50510), and *Enterococcus faecalis* S613 (HM-334). The **DapR** strains are *S. aureus*, Strain JE2, Transposon Mutant NE573 (NR-47116), *S. aureus* JE2, Transposon Mutant NE1656 (BEI product NR-48198), and *Enterococcus faecalis* R712 (HM-335). Enterococcus Faecalis Strains S613 and R712 were isolated from the blood of a 64- year old hemodialysis patient with fatal bacteremia. Strain S613 was isolated from the patient before treatment with daptomycin and is reported to be susceptible to daptomycin. Strain R712 was isolated from blood drawn after treatment of daptomycin. (Minimum inhibitory concentration (MIC) testing against all strains was performed in Mueller-Hinton Broth (MHB) or Brain Heart Infusion (BHI) broth, for DapS and DapR strains respectively, supplemented with 50 mg L<sup>-1</sup> of calcium [136, 137] following the CLSI guidelines for broth microdilution methods using inoculum of  $1 \times 10^5 - 5 \times 10^5$  CFU/mL.[138] All Dap analogue stocks were made in DMSO. Serial two-fold dilutions of the compounds ranged from 20 to 0.16  $\mu$ M, 5 to 0.039  $\mu$ M and 1.6 to 0.0125  $\mu$ M. Briefly, the overnight cultures were grown at 35 °C in MHB. Daptomycin and daptomycin analogues were serially diluted in MHB followed by inoculation of serially diluted media with 10,000 counts of bacteria from the overnight culture. The culture plates were incubated with shaking at 35 °C for 20–24 h. Growth was evaluated using the absorbance at 600 nm. The lowest concentration causing 80% inhibition microbial growth was defined as the MIC. Growth and sterility controls were included in each experiment. In addition, two representative gram-negative bacteria *E. coli* (ATCC 11775) and *P. aeruginosa* (ATCC 10145) were included in the activity assays. The



cancer cell-line cytotoxicity assays as a measure of general eukaryotic cell toxicity employed the human hepatocyte carcinoma cell line HepG2. The cytotoxicity of daptomycin and its analogues were tested against HepG2 cell line at 20  $\mu$ M by using MTT assay.[139]

# Appendix

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