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A Drug Discovery Partnership for Personalized Breast Cancer Therapy

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INTRODUCTION:

This project brings together the skills and resources of Xavier University and Tulane University researchers to form a collaborative team in the areas of drug design and validation. This inter-university collaboration whereby a number of in vitro and in vivo approaches are applied for validation of lead compounds designed and synthesized at Xavier University involves training of Xavier researchers and students in drug target validation, biological assays of drug efficacy, evaluation of resistance pathways, and identification of synergistic drug combinations. At Xavier University, Dr. Thomas Wiese is an expert in computer-aided drug design methods, Dr. Maryam Foroozesh is an expert in the design and synthesis of small biologically active organic compounds, and Dr. Jayalakshmi Sridhar is an expert in both fields. On the Tulane Cancer Center side, Dr. Frank Jones is an expert in HER2 positive breast cancers and has an active drug validation program, and Drs. Barbara Beckman and Matthew Burow are experts in the areas of endocrine- and chemo-resistant breast cancers. The distinct contributions of each institution places this collaboration in a unique position to successfully design and validate drugs to address the most pressing challenges in breast cancer therapy. The specific aims of this collaborative project are to develop, promote, and sustain independent, competitive breast cancer research at Xavier University of Louisiana while developing a true partnership between the two institutions.

BODY:

Foroozesh/Beckman/Burow Subproject (Novel Ceramide Analogs as Anti-Cancer Agents)

The research accomplishments of this subproject include the following syntheses and bioassays:

Task 1- Hire research associate to assist in project. (Month 1)

Dr. Jiawang Liu has been working on the DoD project since the beginning of this grant project. He is an expert in the design, synthesis, and bioassays of biologically active molecules.

Task 2- Identify student(s) to assist in project. (Months 1-3)

Over the past three years, a number of Xavier Undergraduate students (13) have been involved in this project.

Task 3- Design and synthesize new ceramide mimicking agents in order to perform structure-activity studies of these novel compounds in hopes of determining important and essential structural features leading to enhanced apoptosis induction. (Months 1-48)

Year 1: The synthesis of ceramide analogs (401, 402, 403, 404 and 406) with different backbones were achieved based on the previously published methods and reported in the Year 1 Progress Report. These specific analogs were designed to help us identify the contributions of the ceramide backbone in the anti-cancer activities. The synthetic scheme for this group of analogs is shown in Figure 1.
Figure 1. Synthesis scheme for analogs 401-406.

Ten ceramide analogs containing the imine group (409, 410, 412, 413, 415, 416, 417, 3T1, 3T2, and 3T3) were also prepared through a facile reaction. Since one of our previously synthesized ceramide analogs containing an imine group has shown the most potent anti-cancer activities, the modifications here are expected to increase the anti-cancer activity and/or provide us with important structure-activity relationship information. The structures of these compounds are shown in Figure 2.

Figure 2. Structures of ceramide analogs 409, 410, 412, 413, 415, 416, 417, 3T1, 3T2, and 3T3.
A ceramide analog with the 1-hydroxy group blocked (503) was prepared using relatively inexpensive staring materials and simple synthetic route as shown in Figure 3. Compared with the previous synthetic methods, our new synthetic route avoids the usage of expensive starting material methyl (S)-(−)-3-Boc-2,2-dimethyl-4-oxazolidinecarboxylate and excessive protecting and de-protecting steps. Using the starting material O-benzyl-serine, the 1-position blocked ceramide analog (503) was synthesized through a facile four-step route shown below.

**Figure 3.** The four-step synthetic route used for the synthesis of analog 503.

**Year 2:** In Year 2, we focused on the synthesis of fluorescent building blocks for incorporation into the new ceramide analogs. Evidence shows that rigid modification of ceramide backbone enhances pro-apoptotic efficacy. Employing flavone and coumarin cores as rigid moieties, the long conjugated system-modified ceramide analogs were designed. The synthetic route used is presented in Figure 4. Incorporation of the long conjugated system adds molecular rigidity and fluorescence to the ceramide, facilitating the determination and tracking of the ceramide analogs in bioassays.

**Figure 4.** Synthetic route for flavonyl-modified ceramide analogs. Step A) propargyl...
bromide, potassium carbonate/acetone; Step B1) 200°C in N,N-diethylaniline; Step B2) 200°C cesium fluoride in N,N-diethylaniline; Step C) liquid bromine, aluminum chloride/dichloromethane; Step D) methyl phenyl sulfoxide, lithium diisopropylamine/tetrahydrofuran; Step E) potassium carbonate/N,N-dimethylformamide; Step F) trifluoroacetic acid/dichloromethane; and Step G) octanoic acid, DCC, HOBT/tetrahydrofuran.

In Year 2, we successfully synthesized 11 pyranoflavone and 4 furano flavone/coumarin building blocks for the preparation of fluorescent ceramide analogs (Figure 5). Through altering the reaction condition of Claisen rearrangement, pyrano- and furano- derivatives were obtained in a facile method and good yields.

Figure 5. The molecular structures of pyranoflavones, furanoflavones, and furanocoumarins synthesized.
In Year 2, we also focused on the determination of molecular conformation of oxazolidine ceramide analogs. Oxazolidine ceramide analogs have shown high anticancer activity in previous studies. Mechanism investigations show that they could induce apoptosis as well as inhibit sphingosine kinase, a ceramide-metabolizing enzyme. Because of the cyclization of 1-position OH group and 3-position amino group, the oxazolidine ceramide analogs possess considerable rigidity and fixed conformations, which are useful for investigating their interactions with molecular targets. Ceramide analogs, 401 and 402 are oxazolidine ceramide analogs. Thus, a conformational investigation of these compounds was carried out in Year 2. 1D NMR spectra show the two sets of signals for each compound. 2D NMR spectra clarify the stereochemistry of compounds. Through molecular simulation and conformational analysis, dual-conformation model of oxazolidine ceramide analogs is generated as seen in Figure 6.

![Figure 6](image)

**Figure 6.** Hypothesized conformations for the conformational isomers of analogs 401. The bulky group –COCH=CHCH=CHC\(_{10}\)H\(_{21}\) is located above the amide bond plane in the half-chair form, while below the amide plane in the envelope form. In both conformations α-H is in an axial position.

The sp\(^2\) hybridization of N in the amide group results in a planar amide functional group (pink residue). Due to the presence of a bulky group (–COCH=CHCH=CHC\(_{10}\)H\(_{21}\)) on the carbon next to the N in the five-membered oxazoline ring, steric effects lead to the –COCH=CHCH=CHC\(_{10}\)H\(_{21}\) group locating in the space above or below of the amide bond plane. Since these two poses cannot freely interchange, two relatively stable conformations are formed which are reflected by the two sets of signals in the NMR spectra. Figure 6 shows the two possible conformations of 401 in half-chair and envelope forms, respectively. In both conformations the α-H (in blue) is axial, which is
consistent with the observations in $^1$H NMR spectra. This new finding, disclosing the conformational isomerism phenomenon of oxazolidine.

**Year 3:** In Year 3, we successfully synthesized 11 pyrano-, furano-, dioxolo-, and pyridino-flavone/coumarin building blocks (Figure 7). We have established an effective method to synthesize flavone and coumarin derivatives, which could be used as fluorescent blocks of ceramide analogs.

![Figure 7. The molecular structures of 11 pyrano-, furano-, dioxolo-, and pyridino-flavones or coumarins synthesized.](image)

In Year 3, we also successfully synthesized 6 ethynylflavones (Figure 8) as cytochrome P450 inhibitors. Enzymatic inhibition assays show that 4-ethynylflavone and 6-ethynylflavone are highly potent and selective inhibitors of P450 1A1 ($K_i$ values of 0.035 and 0.039 µM). Since P450 1A1 is responsible for the conversion of environment pollutants to carcinogens in human body, 4-ethynylflavone and 6-ethynylflavone are potential cancer prevention agents.

![Figure 8. The molecular structures of ethynylflavones.](image)

**Task 4** - Determination of Anti-Cancer Activities of the Ceramide Analogs. (Months 3-60)
The anti-cancer activities of ceramide analogs 401, 402, 403, 404, and 406 were tested using a cellular viability assay and a clonogenic survival assay in MCF-7, MDA-MB-231, and MCF-7TN-R cells. Compounds 401 and 406 were the most effective compounds across all cell lines with IC$_{50}$ values of 4.05 ± 1.3 µM (p<0.001) and 4.26 ± 1.48 µM (p<0.001) respectively, in the chemo-resistant MCF-7TN-R cell line. Interestingly, IC$_{50}$ values for all analogs except analog 401 were lower in the chemo-resistant MCF-7TN-R and hormone therapy-resistant MDA-MB-231 cell lines, indicating that these compounds exhibit increased therapeutic potential in drug-resistant cancers (Table 1). The fact that two compounds with the 3-ketone-4,6-diene backbone (406 and 401) have shown the most potent anti-cancer activities in this group suggests that the 3-hydroxy-4-ene backbone is not necessary for bioactivity of ceramides as previously believed. The raw results (Figures 9 and 10) are shown below.

Our results in apoptosis assays show that analog 406 induces a 4.3 ± 1.1 fold (p<0.05) increase over control in the induction of apoptosis, compared to C8-Cer with a 2.34 ± 0.79 fold increase. Analog 406-induced cell death is mediated through the intrinsic apoptotic pathways, with 3.59 ± 0.45 (p<0.05) fold increase in caspase-9 activity following treatment with the analog. In conjunction with our previous studies, these results suggest that development of ceramide analogs with a diene component in the sphingosine backbone may be well suited for the treatment of chemo-resistant breast cancer.
Table 1. IC_{50} values of ceramide analogs in the MTT viability assay and the clonogenic survival assay (µM). The values are the means of three independent experiments.

<table>
<thead>
<tr>
<th>IC_{50} values in viability assay (µM)</th>
<th>IC_{50} values in survival assay (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCF-7</td>
<td>MCF-7-NTR</td>
</tr>
<tr>
<td>MDA-MB-231</td>
<td>MCF-7-NTR</td>
</tr>
<tr>
<td>MDA-MB-231</td>
<td>MCF-7-NTR</td>
</tr>
<tr>
<td>401 3.906</td>
<td>4.047 26.76</td>
</tr>
<tr>
<td>26.44 9.908</td>
<td>45.54 5.692</td>
</tr>
<tr>
<td>37.28 4.742</td>
<td>35.79 4.175</td>
</tr>
<tr>
<td>233.6 28.96</td>
<td>NE 10.16</td>
</tr>
<tr>
<td>22.03 4.263</td>
<td>81.94 3.403</td>
</tr>
</tbody>
</table>

Figure 9. Effect of ceramide analogs on breast cancer viability. MCF-7, MDA-MB-231, and MCF-7TN-R cells were treated with increasing concentrations of analogs for 24h. The values are the mean ±SE of three independent experiments.
Figure 10. Effect of ceramide analogs on breast cancer clonogenic survival. MCF-7, MDA-MB-231, and MCF-7TN-R cells were treated with increasing concentrations of analogs and allowed to grow until colony formation was noted (generally 10-12 days). The values are the mean ± SE of three independent experiments.

A longitudinal activity comparison of analogs 315, 406, 415 and 503 was performed. These cell viability assays were performed on NCI/ADR-RES, NCI/ADR, OVCAR8, MCF-7, and MCF-7/Dox cells. The results showed that among these analogs, compounds 406 and 415 show the most potent activities. The raw data is provided below in Figure 11.

Glucosylceramide synthase (GCS) inhibition assays showed that analog 406 has a mild or no GCS inhibition activity in OVCAR8, NCI/ADR-RES, and NCI/ADR cells. This observation suggests that cytotoxicity of analog 406 is not a result of the inhibition of GCS enzyme. On the other hand, analog 503 showed a significant GCS inhibition activity in all of the tested cell lines. This observation confirms our hypothesis that GCS activity can be inhibited through modification of ceramide's 1-position. These results provide us with a great perspective for designing novel inhibitors of GCS, an enzyme considered to be critical in cancer drug-resistance. The results of the GCS activity assays are shown below in Figure 12.
Figure 11. Effect of ceramide analogs on cancer viability. NCI/ADR-RES, NCI/ADR, MCF-7, and MCF-7/Dox cells were treated with increasing concentrations of analogs for 72 h. Analyzed by CellTiter-Glo luminescent cell viability assay from the Promega.

Figure 12. Results of GCS inhibition activity assays in OVCAR8, NCI/ADR-RES, and NCI/ADR cells demonstrated by thin layer chromatograph (TLC). Treatments for 48 hours in 5% FBS RPMI-1640 medium; analyzed by fluorescence enzymatic assay (Gupta V et al J Lipid Res 2010, 51:866-74), three times.
**Wiese/Burow Subproject (Identification of novel estrogens and antiestrogens in the USDA Phytochemical and FDA Marketed Drugs databases)**

Research accomplishments of this subproject include the following tasks in Specific Aim 1: **Develop structure-based pharmacophore models and ligand-receptor (docking) models for estrogens based on the crystal structures of ER alpha and beta (with bound agonists or antagonists) and then virtually screen the USDA Phytochemical, Chinese Herbal Medicine, and the FDA Marketed Drug Databases for new estrogens.**

**Task 1- Identify student to assist in project. (Month 1)**

**Year 1:** Pharmacy students Chioma Obih and Felicia Gibson who have worked in the Wiese lab for the previous 2 years were assigned to this project in Fall 2011. Both students were supported by the College of Pharmacy Center of Excellence Grant. Dr. Wiese trained Ms. Obih on structure-based modeling methods using the MOE software and she worked with Dr. Wiese on Task 2. Ms. Gibson focused on working with Ms. Candace Hopgood on *in vitro* bioassays.

**Year 2:** Ms. Obih graduated in Spring 2013 and left the group. Starting January 2013, Ms. Gabriela Barbarini, a pharmacy exchange student from Brazil started working in the Wiese lab learning bioassay techniques used in Task 6. Ms. Barbarini was a third year pharmacy student doing one year in the US at Xavier’s college of pharmacy and selected to work in the Wiese lab. During summer of 2013, Ms. Barbarini worked in the laboratory of Dr. Burow at Tulane (DOD Project Collaborator of Dr. Wiese) and learned additional bioassay methods. In Fall 2013, she was back in the Wiese lab, applying her skills to evaluate the estrogen activity of compounds identified in this project in a 3 credit research experience course.

**Year 3:** Ms. Barbarini completed her fall semester research experience where she performed reporter gene assays on the 8 stilbenes identified in the initial screen as having some estrogen activity. This produced dose response data indicating the relative potency of these compounds. She then finished her exchange program and returned to Brazil in December 2013. A new P1 pharmacy student, Ms. Tamara Mitchell, joined the lab in Fall 2013 and quickly learned cell culture techniques and then applied these skills to testing the 8 stilbenes shown to have estrogen activity in the MCF-7 breast cancer cell proliferation assay. This produced dose response data for these compounds. Tamara then presented this data in a poster at the Spring 2014 ACS meeting. In spring 2014, Tamara started to learn molecular modeling methods relevant to the project and will apply these skills to modeling analysis of the stilbene data in Y4.

**Task 2- Develop structure-based pharmacophore models for estrogens. (Months 1-4)**
**2a - Obtain all crystal structures of ER LBDs (Month 1)**

**Year 1:** A search of the Protein Database in Fall 2011 resulted in the identification of 62 crystal structures of the human Estrogen Receptor (ER) ligand-binding domain (LBD), all of which contained one bound ligand. These LBD structures were processed and aligned relative to each other so that similarities and differences in ligand-binding pockets could be identified.

**2b - Sort LBD structures by cavity shape and helix-12 position (Months 1-3)**

In preliminary studies prior to this project, we have shown that ligand receptor docking (or virtual screening using docking) can produce very different results between ER LBD structures containing steroid or stilbene ligands, even though both ligands are agonists and the LBD cavity sizes are very similar. The Xavier Molecular Structure and Modeling Core was utilized to compare the ligand-binding cavity sizes of the 62 processed structures. At the same time, a manual sorting was undertaken to group ER LBD crystal structures by bound ligand type, cavity size, and position of helix 12. This process resulted in the identification of 26 structures in the antagonist configuration (helix 12 open) and 36 structures in the agonist configuration with helix 12 closed. While cavity volume did not clearly group these structures, a combination of cavity size and bound ligand type was used to select representative agonist and antagonist crystal structures of the ER LBD. These 5 agonist structures (1ERE, 2G50, 2P15, 2QH6, and 3ERD) and 3 antagonist structures (1ERR, 3DT3, and 3ERT) will be used in the structure-based database screening.

**Year 2:** A computational mythology was identified that can utilize all of the ER LBD structures for the virtual screening rather than a subset of 5. The new strategy was to use as many of the 62 ER LBD structures as needed for docking the target databases based on the similarity of each database compound to the bound ligands of each available ER LBD. The critical part of this method involves sorting the databases by similarity to the bound ligands of all 62 ER LBD structures. Then, database members most similar to any bound ligand are docked into the corresponding ER LBD crystal structure. This approach should reduce false positives and negatives in the virtual screening since the ER LBD is known to take on slightly different ligand binding cavity shapes as it binds to different ligands. In order to apply this methodology to this project, Dr. Wiese attended a one-day training workshop held by Chemical Computing Group (Makers of MOE software) in May 2013. From this training, a process was developed to presort the databases before docking and the skills were obtained to carry out the process. This new workflow for the screening and validation process has been refined to the following:
1. Wash and Filter Databases to molecules that may bind to ER LBD (same as original plan).

2. Sort databases into subsets by similarity to the ligands bound to the 62 ER LBD structures.

3. Dock sorted database subsets into corresponding ER LBD that have been aligned by binding cavity.

4. Repeat steps 2 and 3 after Meteor-derived Phase I metabolites are generated.

5. Select compounds for validation with bioassays based on docking scores.

**Year 3:** This process is continued in Y3.

2c - Develop pharmacophore models from representative LBD structures (Months 1-4)

Since all of the ligands in the selected ER LBD structures bind to the ER using similar interactions, the development of classical pharmacophore models for the ER LBD models was determined to be unnecessary. Structure-based screening for this project will utilize docking to the selected crystal structures where typical pharmacophore interactions are part of the ligand pose generation and score process.

**Task 3-** Mine phytochemical and marketed drug databases with pharmacophore models. (Months 3-5)

3a - Evaluate Docking methods for virtual screening of estrogens (Months 4-8)

**Year 1:** The Xavier Molecular Structure and Modeling Core was utilized to quickly evaluate the potential for the MOE, Gold, Glide, and Surflex Dock to be used for docking into the ER LBD. FlexX is no longer used in this lab, and the Glide method is an additional method used by the Molecular Structure and Modeling Core. This fast study using default setting of the software to replace the bound ligand into the binding cavity did not identify significant differences in the performance of these methods. Considering that the Wiese lab has the most experience with the MOE software and the fact that the MOE software is the only docking package available to us that can include a force field optimization, MOE was selected for further optimization studies.

**Year 2:** Each ER LBD structure was evaluated and processed for docking using the MOE Structure Preparation Module.

3b - Ligand replacement optimization docking of representative LBD structures using MOE, Gold, FlexX, Surflex Dock (Months 4-7)
Year 1: A systematic ligand replacement study was performed using the MOE software, and an optimal configuration of the MOE docking method was identified that produced ligand replacement very close to the crystal structure (low RMSD). The MOE docking software, with its multiple ligand placement and scoring methods, was systematically evaluated for ability to replace the bound ligand in representative examples of the ER agonist and antagonist LBD structures.

Year 2: A method using the Triangle Matches Placement Method and the London dG scoring followed by a ligand minimization and final rescoring with method GBVI/WSA dG was found to produce ligand replacements very close to the crystal structure.

Task 4 - Refine pharmacophore selection of estrogens using docking. (Months 6-8)

Year 1: Not initiated.

Year 2: The phytochemical (76,451 compounds) and marketed drug databases (16,096 compounds) were obtained from the Xavier Molecular Structure and Modeling Core in SDF format.

Year 3: The processing these databases for virtual screening continued in Y3. This includes creating all tautomers, isomers, enantiomers, and filtering out compounds too large to bind ER using MOE. In addition, the software Meteor will be used to create potential Phase I metabolites of each structure. The process of sorting these databases by similarity to the bound ligands of the 62 ER LBDs continues (see Task 2b above) and a new set of test compounds should be identified in first half of Y4.

The ER docking results for the 8 stilbenes found to have ER agonist or antagonist activity are shown below. Note that the stilbenes with agonist activity share the binding mode of the potent agonist diethylstilbestrol (DES) shown in dark magenta while the stilbenes with antagonist activity have only a partial overlap with the binding mode of 4OH-Tamoxifen (dark magenta). Thia binding mode allows the helix 12 of ER to close and initiate transcription. The antagonist stilbenes 4 and 9 are cis (bent) and occupy both the estradiol binding pocket and the channel created by the aryl amine of tamoxifen in the docking simulation. This is the binding mode of typical ER antagonists that directly push away the helix 12 of the ER by creating a new channel. Stilbene 28 is trans (not bent) and docking places the ligand in the estradiol binding pocket where it may act as a indirect ER antagonist and indirectly acting on helix 12 of ER.

ER Docking of Agonist Stilbenes 10 and 11 in 3ERD
ER Docking of Antagonist Stilbenes 4 and 28 in 3ERT

Task 5- Hire research associate to assist with *in vitro* assays. (Month 7)

**Year 1:** Ms. Candace Hopgood had worked in the Wiese lab for 2 years and was transferred to this project briefly in spring and summer of 2012. She spent the summer testing some of the bioassays to be used in the validation phase of this project including the Lantha Screen ER binding assay and the labs MVLN and T47D reporter gene assays. Ms. Hopgood left the Wiese lab in Fall 2012 to focus on her application to Xavier Pharmacy School where she is now a student.

**Year 2:** In January 2013, Ms. Peng Ma was partially reassigned to the bioassay component of the project (50% effort). Ms. Ma also continues to work with Dr. Wiese in the RCMI (NIH-funded Research Centers in Minority Institutions) Cell and Molecular Biology Core (at 50% effort) that he directs. Ms. Ma is very skilled with most of the *in vitro* methods used to validate the virtual screening in this project. Ms. Ma has also been involved in training and working with Ms. Barbarini (a new student) in the lab.
Year 3: Ms Ma continues to dedicate 50% of her time to this project maintaining cells, performing bioassays, training and working with students doing bio assays and learning the ER binding assay method.

Task 6 - *In vitro* validation of estrogen activity. (Months 8-30)

6a - Obtain samples of 10-20 test compounds selected in virtual screen (Months 8-9)

Year 1: Not initiated.

Year 2: A preliminary database of 29 stilbene analogs has been obtained from the USDA Natural Products Utilization Research Unit in Mississippi. The stilbene structure core has been used as the basis for potent ER agonists and antagonists that are in the registered pharmaceuticals and herbal medicine databases. The 29 analogs obtained have been characterized for anticancer effects, but not evaluated for estrogen activity.
If these Stilbenes, 19 have a trans (extended) configuration like the well studied and potent estrogen agonist diethylstilbestrol (DES), one is a stilbene with no potential for E, Z isomers and nine are in the cis configuration where the overall structure is bent. We expect that some of the extended stilbenes will have estrogen agonist activity and that some of the bent (cis) stilbenes might have some antiestrogen activity.

6b - Perform FP ER-alpha/beta binding determinations of test compounds (Months 9-15)

**Year 1:** Not initiated.

**Year 2:** Dr. Wiese has been training Ms. Ma to carry out the ER binding assays and she has developed her proficiency to a level that she will soon run ER alpha and beta binding curves for the 29 stilbenes.

**Year 3:** ER binding was not completed for the 8 active stilbenes in Y3 due to problems with the FP instrument and data collection from the instrument in the Wiese Lab. A new computer workstation was obtained and binding determinations will be done in Y4. In addition, a method development process has started where the FP binding assays can be read using the plate reader in the Xavier RCMI CMB core lab.

6c - Perform MVLN reporter gene agonist/antagonist determinations of test compounds (Months 12-18)

**Year 1:** Not initiated.

**Year 2:** The 29 stilbene compounds have been used as a test set to train and standardize the ER responsive reporter gene assays in the Wiese Lab. All compounds were tested in three experiments at 10 uM in the T47Dkb-Luc cells to check for agonist and/or antagonist activity. Rather than using the MVLN reporter gene cells, the T47Dkb-Luc cells have been used since they provide a stronger estrogen response and since they are 10X more sensitive to estrogens. Representative data are shown below. Five stilbenes were found to be ER agonists, 3 exhibit antagonist activity and 7 stilbenes potentiate the activity of estradiol.
**Year 3:** Ms. Barbarini performed reporter gene assays for the 8 active stilbenes producing dose response data over the range of 10 nM to 10μM characterizing stilbenes 10, 11, 15, 33 and 34 as weak agonists active only at high concentrations. While weakly potent, stilbenes 10 and 33 where shown to induce exaggerated efficacy of 300% compared to the estradiol control. One the other hand, stilbenes 9 and 28 were shown to inhibit the estrogen receptor induced reporter gene more than 90% at high concentrations. Summary data is shown below.
To further characterize the potential for the 8 “active” stilbenes to induce agonist or antagonist estrogen activity, the MCF-7 proliferation assay was used. These determinations attempt to show efficacy in terms of a natural cellular response: proliferation. Comparison of the reporter gene and proliferation results has shown that while the reporter gene assay is a very sensitive screen for ER activity, the impact on cell proliferation may be more muted. Samples data is shown below.
From these *in vitro* characterizations of the 28 stilbenes, 8 have shown to have significant activity in the reporter gene and cell proliferation systems. In year 4, we will focus on these 8 compounds and determine ER alpha and beta binding activity (Task 6B) and include only these compounds in Tasks 6d-h. We will then focus on the most active of the 8 compounds in Task 7 (Stilbenes 4, 9, 11).

**6d** - Perform ER alpha/beta selective reporter gene assays of test compounds (Months 12-20)

**6e** - Evaluate test compounds in estrogen/breast cancer PCR array (Months 16-30)

**6f** - Data analysis of PCR array data (Months 28-33)

**6g** - Evaluate coactivator ER interactions with test compounds bound using LanthaScreen™ TR-FRET ER alpha/beta Coactivator Assay (Months 16-35)

**6h** - Perform genome wide shRNA library screen coupled with gene expression arrays of sensitive cells to identify drug targets, drug sensitizers, and drug-resistance pathways (Months 18-30)
Tasks 6d-6h have not been initiated.

**Task 7** - *In vivo* validation of estrogen activity. (Months 33-48)

7a - Test compounds for uterotrophic activity in mice (months 33-37)

7b - Examine antiestrogenic capacity of test compounds in uterotrophic assays (months 34-38).

7c - Evaluate activity of test compounds on breast cancer xenografts (months 38-48).

Task 7 has not been initiated.

**Sridhar/Jones/Stevens Subproject (Identification of a New Class of Tyrosine Kinase Inhibitors)**

The research accomplishments of this subproject include the following:

**Task 1**- Hire research associate to assist in project. (Month 1)

Dr. Jayalakshmi was hired as a research associate. Her expertise in organic chemistry and skills in molecular modeling made her an ideal fit for this project. In August of 2012, she joined the Xavier University Chemistry Department in a new position as a tenure-track Assistant Professor. In her new capacity, she co-directs this subproject with Dr. Cheryl Stevens who has left Xavier for a position as the Dean of the College of Science and Engineering at Western Kentucky University. Dr. Stevens and Dr. Sridhar have agreed to continue collaborating on this subproject with the goal of developing Dr. Sridhar into a prolific and well-trained cancer researcher.

**Task 2**- Identify student to assist in project. (Month 3)

Year 1: Due to Dr. Stevens leaving Xavier University in January 2012, no students were hired on this project in Year 1.

Year 2: Two students worked on the project in Year 2 (Thuy-Linh Nguyen, and Jasmine Thompson).

Year 3: Four students have been working on the project in Year 3 (Nancy Pham, Phan Tram, Jasmine Geathers, and Don Q. Nguyen).

**Task 3**- Identify novel small molecules related to quinazoline, tyrphostin, emodin, and dasatinib that inhibit HER2 activity. (Months 1-24)

3a - Identify detailed pharmacophore and determine geometric, electronic, and lipophilic characteristics required for tyrosine kinase inhibition (Months 1-12)

HER2 is a growth factor receptor protein belonging to the tyrosine kinase receptor family. HER2 is overexpressed in 25-30% of breast cancer patients and its overexpression has been detected in several other cancers including prostate cancer, ovarian cancer, lung cancer, mammary carcinoma, liver tumors, and colorectal cancers.
Trastuzumab is a humanized antibody targeting the extracellular domain of HER2 that is currently being used clinically. Among the many tyrosine kinase inhibitors developed so far, only Lapatinib is in clinical use. Several other HER2 kinase inhibitors are in various stages of clinical trials.

The splice variant HER2Δ16 isoform lacking exon 16 preceding the transmembrane domain shows low sensitivity to Trastuzumab. This makes the development of a HER2 kinase inhibitor a more reasonable approach. Castiglioni, et al.\textsuperscript{1} have shown that emodin (1,3,8-trihydroxy-6-methyl-anthraquinone) was the only drug that inhibited the therapeutically resistant oncogenic HER2 isoform, HER2Δ16. Based on these reports, emodin was chosen as the lead structure for development of HER2Δ16 inhibitors. Emodin and Iressa were first docked onto the HER2 homology model to study their binding modes with the help of MOE docking tools. Iressa did not bind to the hinge region residues of the protein. However, emodin did bind to the hinge region of the protein and three binding modes were identified (Figure 1). Based on the orientation of emodin in the binding pocket of the protein, residues that could be targeted for developing a good inhibitor were identified. These were Thr95, Gln96, Met98, Asp160, Lys50, Glu67, Thr159 (Figure 1).

\textbf{Figure 1}. Binding modes of emodin onto HER2 protein homology model and a picture of the binding pocket with the potential target residues depicted in stick mode.

3b - Identify new compounds to be tested for tyrosine kinase inhibition with conformationally flexible searches of compound databases using detailed pharmacophore and CoMFA QSAR results. (Months 9-24).

Year 1: 1,3,8-trihydroxyanthraquinone was taken as the pharmacophore for a UNITY 2D-search of all the databases available to us. Hits were obtained from ACD (10 hits), TSCA (3 hits), and NCI databases (39 hits). NCI database hits overlapped significantly with the compounds contained in ACD hits.
**Table 1:** High-throughput screening of database hits against MCF7-HER2Δ16 cell line.

<table>
<thead>
<tr>
<th>Compound</th>
<th>% of E2 at 10^{-5}M</th>
</tr>
</thead>
<tbody>
<tr>
<td>AG-650/41069241</td>
<td>91.27</td>
</tr>
<tr>
<td>AG-650/41069319</td>
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<td>AG-650/41069355</td>
<td>98.16</td>
</tr>
<tr>
<td>AP-782/41885488</td>
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</tr>
<tr>
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</tr>
<tr>
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<tr>
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</tr>
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<td>NSC93419</td>
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</table>
Year 2: Based on the initial high-throughput assay against MCF7-HER2Δ16 cell lines (Table 1), 13 compounds were chosen for further analysis (Table 2). Structures of these compounds are shown in figure 2. Additionally 3D and 2D-databases are being compiled using the commercially available compounds from TIMTEC, LC laboratories, Maybridge, and Pubchem. Based on the docking studies (explained in Task 4a) of the active compounds 4, 5 and 13 from Table 2; the essential features required for the inhibitor pharmacophore has been reduced to 5-hydroxy-2-(hydroxymethyl)naphthalene-1,4-dione. Database searches are currently underway to find new compounds that satisfy this pharmacophore. The hits obtained from the database searches will be docked and common core structures will be identified as new lead molecules.

Table 2: The 13 compounds that were chosen for further analysis

<table>
<thead>
<tr>
<th>No.</th>
<th>Compound</th>
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</tr>
<tr>
<td>13</td>
<td>AP-782/41885488</td>
</tr>
</tbody>
</table>
Year 3: During our database search using the pharmacophore designed earlier for novel core structures, 1H-indazol-3-ol (Indazolone) was found to dock well in the hinge region of the HER2 kinase. Two hydrogen bonds were formed between the indazolone and the HER2 residues Gln102 & Met104. Functionalization of the phenyl ring with suitable substituents can be achieved using well-known reactions (Figure 3). Hence this scaffold was chosen for further exploration. Three compounds were synthesized as initial lead structures (Figure 4).

![Figure 3: Binding mode of 1H-indazol-3-ol (indazolone) to the ATP-binding region of HER2 kinase. The ligand is shown as ball and stick model.](image)

**Figure 4:** Structures of the three 1H-indazol-3-ol synthesized.

Task 4- Explore the mechanism of HER2 tyrosine kinase inhibition. (Months 12-48)

4a - Dock proven and proposed TKIs into the tyrosine kinase ATP binding site using multiple poses, and score results. (Months 12-24)

Year 1: All of the hits described in 3b were then docked onto the homology model of HER2 using MOE dock tools. The docking results were then studied manually. Binding of the molecule to one of the hinge region residues THR95, GLN96, MET98 was taken as a prerequisite. The number of ligand-protein hydrogen bond interactions, the extent of penetration of ligand into the pocket and the nature of ligand solvent exposure (hydrophobic/hydrophilic) were also considered.

Year 2: The three compounds that showed significant potency against the MCF-7 pcDNA, MCF7-HER2 and MCF-7 HER2Δ16 were subjected to docking studies onto the 3D-structure of HER2 kinase region (PDB ID: 3CRD.pdb) using MOE docking module and Surflex in SYBYL-X1.3. The consensus binding modes for these three compounds are shown in Figure 5. The docking modes of the compounds show that the phenolic group forms a hydrogen bond with the hinge region residue Gln799 and an additional hydrogen bond by the side chain hydroxyl group either with the invariable Lys753 for compound 4 or with Asp863 for compound 5. Compound 13 made only one hydrogen
bond with the protein which could account for its lower potency (refer to bioassay results in Task 4d). This lead us to the optimal core structure of 5-hydroxy-2-(hydroxymethyl)naphthalene-1,4-dione as our lead structure.

![Figure 5](image)

**Figure 5**: (A) 4, (B) 5 and (C) 13 in the active site of HER2 kinase. The protein (PDB ID: 3CRD.pdb) residues are shown as stick models and the compounds are shown as ball & stick model.

4b - Optimize molecular structures to maximize ability of compounds to inhibit HER2. (Months 15-30)

**Year 1**: Not initiated.

**Year 2**: From the bioassay results (Task 4d) and the docking studies it was clear that two core structures that can maintain the two hydrogen bonds shown by compound 5 can serve as the lead molecules LM1 and LM2. The synthetic protocol for the lead molecule LM1 has been designed as given in the Scheme 1 involving many known reactions (Tanoue and Terada, *Bull. Chem. Soc. Jpn.*, (1988) 61, 2039). The reactions until the formation of compound 17 have been standardized. Different Grignard reagents will be used to obtain variation in the R-group on compound 17. The purification of compound 17 and its conversion to LM1 by oxidation with ceric ammonium nitrate is currently being pursued.

Scheme 2 provides an alternate route for the synthesis of the lead compound LM1 by using the Diels-Alder reaction (Paull et al., *J. Med. Chem.*, (1976), 19, 337). The reactions in this scheme for the formation of the dihydroxynaphthoquinone 18 have been standardized. Further reactions are presently being carried out. Once the final diol is obtained from compound 18, further reactions will be performed to obtain derivatives with different functional groups on the side chains and with differing chain lengths.
The synthesis of the lead molecule LM2 will be pursued using Scheme 3 which is similar to Scheme 2 in that both the schemes involve the Diels-Alder reaction as the first step. The first reaction to compound 18 in Scheme 3 is presently being standardized. Upon standardization of the reactions, different derivatives of LM2 will be obtained by using suitably substituted starting materials and/or by derivatization of the hydroxynaphthoquinone LM2.
Year 3: Toxicology studies on HeLa cells (given in section 4d) revealed that among the three compounds that showed good inhibition of the MCF7/HER2 and MCF7/HER2Δ16 cell lines, two of them drugs 4 and 5 were highly toxic. Toxicology studies for the quinones that showed inhibition activity were also subjected to toxicology studies on DU-145 cells. All of these studies revealed that compounds that contain the oxygen atom marked by the red circle (Figure 13) were highly toxic. Our strategy is to avoid toxicity and improve the potency of the drug molecules.

Hence our lead compound structure has been modified to target the residues Gln799 and Asp863. We have synthesized several derivatives of quinones LM1 (R = H-, CH3-, CH3CH2-) using the Scheme 1 outlined earlier. The following schemes were employed for the syntheses of 5-hydroxy-7-methylnaphthalene-1,4-dione and 5,8-dihydroxy-2-methylnaphthalene-1,4-dione which were then converted to their bromomethyl derivatives and subsequently the alcohol derivatives (Scheme 4). These compounds will be subjected to in-vitro inhibition assays against MCF7 cell lines the within the next couple of months to check on their activity and toxicity. These derivatives will be used for the synthesis of further analogs.
The synthesis of the indazolone core structures was initiated. Three starting molecules are being synthesized (Scheme 6). The dinitroindazolone can be used to prepare many different analogues as the nitro groups can be reduced and functionalized further using suitable substituents on them to give us diamines or diamides. The reduction of the nitro groups using H₂/Pd will be attempted next followed by acylation and alkylation to get the final compounds. The dibromoindazolone can be used to prepare many different derivatives using coupling reactions such as Suzuki coupling to provide us with alkyl and aryl substituents on the core indazolone moiety. The Suzuki coupling reaction is presently being standardized. Indazolone, dinitroindazolone and dibromoindazolone have been prepared and are presently being subjected to bioassay for inhibition of MCF7 cell lines.

Scheme 5: Synthesis of 1H-indazol-3-ol/indazolone series of compounds
4c - Attempt to identify alternate binding sites. (Months 18-30)

No progress yet.

4d - Perform \textit{in vitro} kinase inhibition and binding assays. (Months 18-48)

\textbf{Year 1:} A total of 28 compounds were procured from the Developmental Therapeutics Program NCI/NIH and Specs chemicals (Table 1). An initial high-throughput assay was performed to determine the inhibition of proliferation of MCF-7 cell line. The compounds that showed good inhibition activity were then subjected to \textit{in-vitro} assay against HER2Δ16 cell line activity. Two compounds that showed low inhibition activity were included to confirm the activity profile of this set of compounds. Two of the tested compounds (NSC322354 and AG-650-41069319) showed low micro molar activity against the HER2Δ16 cell line (< 10 mM) (Table 1).

\textbf{Year 2:} We tested the ability of 13 small molecules (selected based on the high-throughput screening results given in Table 1) that are structurally similar to emodin to inhibit cell viability in MCF-7 breast cancer cell lines that express HER2, HER2Δ16 or empty vector. To do this we treated the cells for 48 hours with a concentration of 10µM for each set of drugs. After treatment, cell viability was tested using the CellTiter-Glo Assay (Promega) (Figure 6A). Of the 13 compounds, we found three that suppressed cell viability potently in all three cell lines (NSC322354 (Drug 4), NSC227279 (Drug 5), and AP-782/41885488 (Drug 13)). After 48 hours, lapatinib decreased cell viability by <50% while the three test drugs decreased cell viability by >90% at the same concentration (Figure 6B).

To determine the ability of the drug treatments to inhibit the activation of receptor tyrosine kinase HER2, Western blots were performed to detect total phosphorylated protein after treating each cell line with the three most potent drugs or lapatinib at 10µM for 2 hours. As expected, lapatinib dramatically decreases HER2 activating phosphorylation at auto-phosphorylation site Y1248 respectively in each of the cell lines. These sites were suppressed to the same extent in the cell lines after exposure to each of the test drugs, indicating that, like lapatinib, activation of both receptors was repressed by the test compounds effectively and quickly. We also tested phosphorylation of HER2 after exposure to one of the compounds (AP-650/41069356 (Drug 10)) that we found to have no effect on cell viability. This drug had no effect on the phosphorylation status of HER2, suggesting that the ability of the drugs to inhibit these receptor tyrosine kinases is critical to their ability to induce cell death.
Figure 6: Inhibition of MCF7-pcDNA, MCF7-HER2 and MCF7-HER2Δ16 cell lines (A) High-throughput assay of emodin and compounds 1 – 13, (B) Inhibition assay of compounds 4, 5 and 13 at different concentrations.

Figure 7: Western blot analysis of autophosphorylation at the HER2 residue Y1248. Compounds 4, 5 and 13 inhibit the phosphorylation at Y1248 in MCF7-pcDNA, MCF/HER2 and MCF7/HER216. Compound 10 which did not show notable inhibition in the high-throughput assay shows significant phosphorylation at Y1248 in all of the three cell lines.
Finally, we measured the IC$_{50}$ of each drug by treating each of the cell lines with different drug concentrations for 48 hours then using the CellTiter-Glo Assay to detect cell viability (Figure 8). Cells overexpressing HER2$\Delta$16 showed resistance to lapatinib (IC$_{50}$ = 19.22µM) compared to wildtype HER2 overexpression (IC$_{50}$ = 15.79µM). All three of the test drugs had a low µM IC$_{50}$ for each of the cell lines. We found that the two most effective drugs at inhibiting cell viability (NSC322354 (Drug 4), NSC227279 (Drug 5)) also had similar IC$_{50}$ concentrations for both the HER2 (0.87µM, 0.82µM) and HER2$\Delta$16 (0.91µM, 0.93µM) overexpression cell lines. With IC$_{50}$ values of <1µM for HER2$\Delta$16 cells, these two drugs are also more effective at inducing cell death compared to lapatinib with an IC$_{50}$ >10µM. These results indicate the potential of either of these drugs to effectively inhibit HER2$\Delta$16 action and thus combat the drug resistance seen in HER2$\Delta$16 expressing tumors.

**Year 3:** Both EGFR and HER2 are members of the EGFR-family of receptor tyrosine kinases. All ligand activated or constitutively active members of this family have the ability to dimerize with and transphosphorylate any other member of the family. In these specific cell lines, overexpression of HER2 or HER2$\Delta$16 results in constitutive activation of the receptor. The constitutively active receptors heterodimerize with and transphosphorylate coexpressed EGFR. To determine the ability of the drug treatments
to inhibit the activation of receptor tyrosine kinase EGFR, Western blots were performed to detect total phosphorylated protein after treating each cell line with the three most potent drugs or lapatinib at 10µM for 2 hours (Figure 9). The parental MCF-7 cell line lacks significant levels of HER2 phosphorylated activation; therefore HER2 fails to dimerize with and transphosphorylate EGFR. This explains why EGFR also remains unphosphorylated in parental MCF-7 cell line. As expected, lapatinib dramatically decreases EGFR activating phosphorylation at auto-phosphorylation site Y1068 respectively in each of the cell lines. These sites were suppressed to the same extent in the cell lines after exposure to each of the test drugs, indicating that, like lapatinib, activation of both receptors was repressed by the test compounds effectively and quickly. We also tested phosphorylation of EGFR after exposure to one of the compounds (AP-650/41069356 (Drug 10)) that we found to have no effect on cell viability. This drug had no effect on the phosphorylation status of EGFR, suggesting that the ability of the drugs to inhibit these receptor tyrosine kinases is critical to their ability to induce cell death.

![Western blot analysis of autophosphorylation at the EGFR residue Y1068.](image)

**Figure 9:** Western blot analysis of autophosphorylation at the EGFR residue Y1068. Compounds 4, 5 and 13 inhibit the phosphorylation at Y1068 in MCF7/pcDNA, MCF/HER2 and MCF7/HER216. Compound 10 which did not show notable inhibition in the high-throughput assay shows significant phosphorylation at Y1068 in all of the three cell lines.

We tested the ability of the compounds to inhibit HER2 (Figure 10) and EGFR (Figure 11) kinase activity directly by using the ADP-Glo in vitro kinase assay kit from Promega, according to the manufacturer’s instructions. Briefly, 11ng of purified HER2/EGFR kinase was incubated with varying concentrations of each inhibitor in the presence of 10µM ATP to establish an inhibition curve for each. The data were fit to a logistic sigmoid function using Origin software, which calculated IC50 values for each drug. The well-known HER2 and EGFR dual inhibitor lapatinib had an IC50 of 0.05µM and 0.027µM respectively. The other three drugs had IC50’s for HER2 and EGFR kinase inhibition as follows: compound 4 (10.8µM and 13µM), compound 5 (11.4µM and 15µM), compound 13 (30.8µM and 2µM), respectively.
Figure 10: Dose response curves for inhibition of HER2 kinase by compounds 4, 5, 13 and lapatinib.

Figure 11: Dose response curves for inhibition of EGFR kinase by compounds 4, 5, 13 and lapatinib.
Toxicology Studies:

HeLa cells were seeded in a 96-well plate at 0.9x10^5 cell/ml. The next day the media is replaced with 0.2% FBS media for 72 hours. At this point, an Alamar Blue assay is performed to calculate cell number per well. Drug is then added and the cells are counted again after an additional 72 hours. The ratio between the number of cells before drug and after incubation with drug is then calculated. The results indicated that compounds 4 and 13 were toxic at a concentration of 10^{-6} \mu M. Only compound 5 did not show any toxicity (Figure 12). Considering the toxicology assay results on HeLa cells and on the DU-145 cells (Task 5, Year 3), we could see compounds containing the oxygen atom marked with a 'red circle' were toxic (Figure 13). The hydroxyl group marked with the 'blue circle' seemed sufficient to maintain the same inhibition potency without toxicity. Hence we decided to pursue the synthesis of compounds containing only the 5,8-dihydroxy-2-

![Figure 12](chart.png)

**Figure 12:** HeLa Alamar Blue cell viability assay for the compounds 4, 5 and 13.

![Figure 13](structure.png)

**Figure 13:** Essential structural features for the lead structure based on bioactivity and toxicity.
(hydroxymethyl)naphthalene-1,4-dione moiety in them.

**Task 5**- Determine preclinical activity and specificity of novel HER2-targeting molecules: determine influence of targeting molecules on HER2 oncogenic signaling and cellular responses using multiple validated preclinical models of breast tumorigenesis and metastasis. (Months 12-24)

**Year 1**: The compound NSC-322354 which showed the best inhibition activity against HER2Δ16 cell line was taken for an analysis of its cross kinase activity. The compound was sent to KinomeScan ([www.kinomescan.com](http://www.kinomescan.com)) for KINOMEscan's *in vitro* competition binding assay against a panel of 96 representative kinases. KINOMEscan™ is based on a competition binding assay that quantitatively measures the ability of a compound to compete with an immobilized, active site-directed ligand. The assay is performed by combining three components: DNA-tagged kinase; immobilized ligand; and a test compound. The ability of the test compound to compete with the immobilized ligand is measured via quantitative PCR of the DNA tag (description of method taken from [www.kinomescan.com](http://www.kinomescan.com)). The compound was tested at a concentration of 10 mM. The results are given in Table 3. The S-score, selectivity score (the number of kinases that bind to the compound divided by the total number of distinct kinases tested), which is a quantitative measure of the compound selectivity, was 0.022. The compound showed good selectivity for two of the kinases, Casein Kinase 1 D and PIM kinases (more selective for PIM1 and PIM3 kinases). Both of these kinases are serine/threonine kinases. PIM1 is an oncogene. The PIM1 gene was initially identified as a proviral integration site in Moloney Murine leukemia virus-induced mouse T-cell lymphomas²,³. Pim kinases are implicated in the development of solid tumors. DNA microarray analyses showed the overexpression of PIM1 in human prostate cancer in relation to the grade of the prostate cancer. CK1d is a member of the ubiquitous casein kinase-1 family, and alterations in the expression and/or activity of CK1 have been observed in breast cancer⁴. CK1d, has been identified as a novel kinase implicated in the modulation of physiological aspects of both ERα (estrogen receptor alpha) and AIB1 (amplified in breast cancer-1 protein). The compound in fact did not show good inhibition of HER2 (ERBB2) kinase. The *in vitro* high-throughput assay of the compounds is currently being performed for the proteins PIM1 kinase, and casein kinase 1 D. This will be followed by a dose response curve assay to determine the IC₅₀ value for these kinases. These compounds will also be tested for inhibition of other breast cancer cell lines.
Year 2: A total of 40 compounds were subjected to high-throughput screening for inhibition of Pim1 kinase though an \textit{in vitro} kinase assay. Out of these 9 compounds (Figure 14) showed notable inhibition potency ranging from 1.3 $\mu$M to 57.1 $\mu$M (Table 4).

### Table 3. Matrix of Compound NSC322354 Screen

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<th>Kinase target</th>
<th>%Control @ 10$\mu$M</th>
<th>Kinase target</th>
<th>%Control @ 10$\mu$M</th>
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<tr>
<td>GSK3B</td>
<td>85</td>
<td>SNARK</td>
<td>90</td>
</tr>
<tr>
<td>IGF1R</td>
<td>100</td>
<td>SRC</td>
<td>91</td>
</tr>
<tr>
<td>IKKa-alpha</td>
<td>87</td>
<td>SRPK3</td>
<td>93</td>
</tr>
<tr>
<td>IKK-beta</td>
<td>91</td>
<td>TGFBR1</td>
<td>100</td>
</tr>
<tr>
<td>INSR</td>
<td>97</td>
<td>TIE2</td>
<td>89</td>
</tr>
<tr>
<td>JAK2(JH1domain-catalytic)</td>
<td>94</td>
<td>TRKA</td>
<td>78</td>
</tr>
<tr>
<td>JAK3(JH1domain-catalytic)</td>
<td>61</td>
<td>TSSK1B</td>
<td>89</td>
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<tr>
<td>JNK1</td>
<td>96</td>
<td>TYK2(JH1domain-catalytic)</td>
<td>57</td>
</tr>
<tr>
<td>JNK2</td>
<td>84</td>
<td>ULK2</td>
<td>88</td>
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<td>VEGFR2</td>
<td>100</td>
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<tr>
<td>KIT</td>
<td>76</td>
<td>YANK3</td>
<td>89</td>
</tr>
<tr>
<td>KIT(D816V)</td>
<td>100</td>
<td>ZAP70</td>
<td>100</td>
</tr>
</tbody>
</table>
Four of these compounds were shown to inhibit the prostate cancer cell line DU-145 cells with potencies similar to that of emodin (Figure 14). Table 4: Inhibition of Pim1 kinase by emodin and compounds 1 to 9.

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC₅₀ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emodin</td>
<td>2.5</td>
</tr>
<tr>
<td>1</td>
<td>21.4</td>
</tr>
<tr>
<td>2</td>
<td>11.6</td>
</tr>
<tr>
<td>3</td>
<td>7.4</td>
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<tr>
<td>4</td>
<td>19.2</td>
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<tr>
<td>5</td>
<td>57.1</td>
</tr>
<tr>
<td>6</td>
<td>1.3</td>
</tr>
<tr>
<td>7</td>
<td>3.6</td>
</tr>
<tr>
<td>8</td>
<td>23.8</td>
</tr>
<tr>
<td>9</td>
<td>3.5</td>
</tr>
</tbody>
</table>

Figure 14: Structure of emodin and compounds 1 to 9 investigated for inhibition of Pim1 kinase.

Figure 15: Growth inhibition curve for DU-145 cells treated with compounds 1, 4, 7, 8 and Emodin.
Year 3: Toxicology assay on DU-145 cells for the compounds in Table 4

DU-145 cells were seeded in a 96-well plate at 0.9x10^5 cell/ml. The next day the media is replaced with 0.2% FBS media for 72 hours. At this point, an Alamar Blue assay is performed to calculate cell number per well. Drug is then added and the cells are counted again after an additional 72 hours. The ratio between the number of cells before drug and after incubation with drug is then calculated. Emodin, compounds 7 and 9 were found to exhibit toxicity on these cells (Figure 16).

![Alamar Blue Viability Assay](image)

**Figure 16:** Toxicology assay on DU-145 cells for emodin and the compounds 1-9 in Table 4.

5a - Perform genome-wide shRNA library screen coupled with gene expression arrays of sensitive cells to identify Drug Targets, Drug Sensitizers, and Drug Resistance Pathways. (Months 18-30)

Not initiated.

5b - Identify and validate drug combinations to improve efficacy and overcome resistance in preclinical models. (Months 24-36)

Not initiated.
5c - Confirm efficacy of drug combinations in preclinical in vivo xenograft and transgenic mouse models of breast cancer. (Months 30-48)

Not initiated.

Programmatic Activities

The breast cancer research group members have met multiple times to discuss and plan the various aspects of the project. A number of seminars/workshops were also held for the group:

1- Monday September 16, 2013, Dr. Thomas Huckaba, Biology Department, “Parsing the catalytic core of kinesin molecules for the forward design of anti-tumor drugs”
2- Monday September 23, 2013, Dr. Melyssa Bratton, Lab Manger, Louisiana Cancer Research Consortium Core Lab, “Cell and Molecular Biology Core: New Core Services Update”
3- Monday September 30, 2013, Dr. Ren Wei Chen, Department of Chemical Engineering, University of California, Santa Barbara, “Development of Proapoptotic Peptide to Intratumorally Spreading Cancer Therapy”
4- Monday October 21, 2013, Dr. Jiawang Liu, Foroozesh Lab, Chemistry Department, “Developing new ceramides for the treatment of drug-resistant breast cancer”
5- Monday October 28, 2013, Dr. Astrid Engel, Tulane University School of Public Health and Tropical Medicine, “LINEs and SINEs differ in their retrotransposition requirements and cellular interactions”
6- Monday November 18, 2013, Dr. Anup Kundu, Biology Department, “Formulation of a targeted nanoparticle system for the treatment of breast cancer”
7- Monday November 25, 2013, Dr. Partha Bhattacharjee, Biology Department, “Peptidomimetic therapy of angiogenesis”
8- Monday December 2, 2013, Dr. Hector Biliran, Biology Department, “TLE1 as a novel regulator of EMT in lung cancer”
9- Monday December 9, 2013, Dr. Jayalakshmi Sridhar, Chemistry Department, "Quinones as Kinase inhibitors"
10- Monday December 16, 2013, Dr. Robert Stratford, College of Pharmacy
11- Monday January 27, 2014, Dr. Daniel Sarpong, Xavier Health Disparity Center
12- Monday February 3, 2014, Dr. Florastina Payton-Stewart, Chemistry Department
13- Monday February 10, 2014, Dr. Syreeta Tilghman, College of Pharmacy
14- Monday February 17, 2014, Dr. Tarun Mandal, College of Pharmacy, “The impact of nanomedicine on translational research”
15- Monday February 24, 2014, Dr. Torrie Harris, Director, Tobacco Free Living
16- Monday March 17, 2014, Dr. Guangdi Wang, Chemistry Department
17- Monday March 24, 2014, Dr. KiTani Parker-Johnson, College of Pharmacy
18- Monday March 31, 2014, Dr. Cecily Bennett, Biology Department
In addition, the researchers on each subproject (faculty, staff, and students) have met multiple times during the past grant year to discuss project direction and experimental designs and results for their projects.

KEY RESEARCH ACCOMPLISHMENTS:

**Foroozesh/Beckman/Burow Subproject**

- Successfully developed a facile synthetic route to prepared 3-ketone-4,6-diene and 1-position-modified ceramide analogs, and obtained 16 novel ceramide analogs.
- Discovered a highly potent ceramide analog (406). The mechanism investigation showed that analog 406 leads to cell apoptosis through intrinsic apoptotic pathway and does not interrupt the function of GCS.
- Designed, synthesized, and determined a novel GCS inhibitor (503, a 1-position-modified ceramide analog), which is extremely useful for the development of highly potent GCS inhibitors.
- Successfully developed a facile synthetic route to prepare fluorescent building blocks for ceramide analogs. Eleven pyranoflavones, three furanoflavones, one pyridinoflavone, one dioxoloflavone, four pyranocoumarin, four furanocoumarin, one pyridinocoumarin, and one dioxolocoumarin were synthesized.
- Discovered the conformational isomerism of oxazolidine ceramide analogs. According to the evidences from NMR spectra, successfully constructed the molecular models of two conformational isomers.
- Discovered highly potent and selective cytochrome P450 1A1 inhibitors, 4-ethynylflavone and 6-ethynylflavone.

**Wiese/Burow Subproject**

- Identified representative ER LBD structures to be used for virtual screening.
- Identified optimal ligand receptor (Docking) method for virtual screening.
- Obtained phytochemical and marketed drug databases for processing.
- Trained three pharmacy students, one in molecular modeling, and both in bioassays.
- Developed new, comprehensive method for virtual screening with ER LBDs.
- Trained one foreign exchange pharmacy student in bioassays.
- Obtained *in vitro* ER agonist and antagonist activity data for 29 naturally occurring stilbenes.
- 5 stilbenes have been characterized as ER agonists, 3 as ER antagonists.
- Modeling of potential binding modes suggest that antagonist stilbenes may act as direct and indirect antagonists of ER.

**Sridhar/Jones/Stevens Subproject**
Identified three compounds that inhibit MCF7-pcDNA, MCF7-HER2Δ16 and MCF7-HER2 overexpressing breast cancer cell lines with sub-micromolar potency.

Performed Western blots to detect total phosphorylated protein after treating each cell line with the three most potent drugs. IC_{50} values for the three compounds were measured for MCF7-pcDNA, MCF7-HER2Δ16 and MCF7-HER2 cell lines.

Performed docking studies on the identified compounds revealing the two hydrogen bonds made by the quinones with the HER2 kinase. The first hydrogen bond made by all three compounds was to the hinge region residue Glu799. The second hydrogen bond was formed by the two compounds showing higher potency to one of the two residues- invariable Lys753 or Asp863.

Discovered two new lead compounds for derivatization LM1 and LM2.

Established synthetic protocols for the synthesis of different derivatives of LM1. The synthetic scheme for LM2 synthesis is presently being standardized.

Performed Western blot analysis of autophosphorylation at the EGFR residue Y1068 and HER2 residue Y1248.

Determined IC_{50} values of inhibition for HER2 and EGFR kinases.

Performed Toxicology assays on HeLa cells and DU-145 cells.

Determined the lead structure for achieving inhibition of MCF7-HER2Δ16 and MCF7-HER2 overexpressing breast cancer cell lines while limiting toxicity.

Found a new core structure 1H-indazol-3-ol/indazolone as a potential inhibitor through computational database searches.

Synthesized several quinone derivatives and three indazolone derivatives that will be subjected to in-vitro inhibition assays against MCF7 cell line.

Program Accomplishments

Organized meetings in order to introduce the students, staff, and faculty members working on the different subprojects to each other and the various projects.

Organized meetings for each subproject’s researchers to meet to discuss project direction and experimental designs and results.

Organized training workshops/seminars.

REPORTABLE OUTCOMES:

Publications

Foroozesh/Beckman/Burow Subproject


“Pyranoflavones: a group of small-molecule probes for exploring the active site cavities of cytochrome P450 enzymes 1A1, 1A2, and 1B1”, J Liu, S. Taylor, P. Dupart, C. Arnold,


**Wiese/Burow Subproject**
None yet.

**Sridhar/Jones/Stevens Subproject**


“Functionalization and Modification of Naphthoquinone Analogs as HER2 Kinase Inhibitors” Divya Jyothi Lella, M.S. Thesis, Western Kentucky University, May 2014.

- Presentations

**Foroozesh/Beckman/Burow Subproject**

“The Design and Synthesis of Benzoate Esters as Potential Anti-proliferation Agents and Inhibitors of Cytochrome P450 Enzymes”, C. Arnould, P. Dupart, J. Liu, and M. Foroozesh, the Annual LaSPACE Council Meeting, and the American Chemical Society Local Section Student Poster Presentation, New Orleans, LA, October 2012.

“Propargyl Flavones as Inhibitors of Human Cytochrome P450s 1A1, and 1A2”, S. Taylor, J. Liu, P. Dupart, and M. Foroozesh, the American Chemical Society Local Section Student Poster Presentation, New Orleans, LA, October 2012.

“Quest for New Mechanism-Based Inhibitors of Cytochrome P450 Enzymes 1A1 and
“Pyranoflavones and 5-Hydroxy-pyranoflavones as Small-molecule Probes into the Active Site Cavities of P450s 1A1 and 1A2”, J. Liu, S. Taylor, P. Dupart, C. Arnold, and M. Foroozesh, the Louisiana Cancer Research Consortium Annual Retreat, New Orleans, LA, March 2013.


“Quest for New Mechanism-Based Inhibitors of Cytochrome P450 Enzymes 1A1 and 1A2”, J. Sridhar, J. Liu, M. Foroozesh, C.L. Stevens, the American Chemical Society National Meeting, New Orleans, LA, April 2013.


“Ethynyl Flavones as Inhibitors of Cytochrome P450 Enzymes”, S. Taylor, J. Liu, P. Dupart, and M. Foroozesh, the American Chemical Society National Meeting, New Orleans, LA, April 2013.


“The Design and Synthesis of Resorufin Propargyl Ethers as Potential Cytochrome
P450 Inhibitors”, S. Bellow, L. Lovings, J. Liu, and M. Foroozesh, the American Chemical Society National Meeting, Dallas, TX, April 2014.

“The Design and Synthesis of Benzoate Esters as Potential Anti-proliferation Agents and Inhibitors of Cytochrome P450 Enzyme”, E. McClain, L. Lovings, J. Liu, and M. Foroozesh, the American Chemical Society National Meeting, Dallas, TX, April 2014.

“Synthesis of Resorufin Derivatives as Inhibitor Indicators of Cytochrome P450 Enzymes”, L. Lovings, J. Liu, and M. Foroozesh, the American Chemical Society National Meeting, Dallas, TX, April 2014.

**Wiese/Burow Subproject**


**Sridhar/Jones/Stevens Subproject**


“Functionalization and modification of Shikonin compounds as HER2 inhibitors”, D.J. Lella, J. Sridhar, C.L. Stevens, and B. Yan. Western Kentucky University – Reach Week poster, April, 2013.


- Employment or Research Opportunities

Individuals trained in the first and second years of this DoD breast cancer project:

Jiawang Liu, Postdoctoral Fellow at Xavier University (Foroozesh Lab)

Jayalakshmi Sridhar, Postdoctoral Fellow at Xavier University (Stevens Lab, currently a new tenure-track faculty member at the Xavier University Department of Chemistry)

James Antoon, Medical Student at Tulane University (Beckman Lab, received his M.D. in May 2012 after receiving his Ph.D. in 2010 at Tulane University. He is currently doing his residency in pediatrics at the University of North Carolina.)

Barbara Drew, Medical Student at Tulane University (Beckman Lab, is currently in her residency in obstetrics and gynecology in Connecticut.)

Tony Wang, Medical Student at Tulane University (Beckman Lab, working on this DoD subproject)

Thong T. Nguyen, Undergraduate Student at Xavier University (Foroozesh Lab, graduated in May 2012 and is now pursuing a Ph.D. in Chemistry at Tulane University)

Adharsh P. Ponnapakkam, Undergraduate Student at Tulane University (Beckman Lab, graduated in May 2012 and is currently continuing his work on this DoD project as a Masters student at Tulane University)

Patrick Dupart, Technician at Xavier University (Foroozesh Lab, Xavier graduate, joined Virginia Commonwealth University as a graduate student in July of 2013)

Shannon Taylor, Technician at Xavier University (Foroozesh Lab, Xavier graduate)

Corey Arnold, Undergraduate Student at Xavier University (Foroozesh Lab)

Erika McClain, Undergraduate Students at Xavier University (Foroozesh Lab)

Brandon Dotson, Undergraduate Students at Xavier University (Foroozesh Lab)

Charne’sa Tutwiler, Undergraduate Students at Xavier University (Foroozesh Lab)

La’Nese Lovings, Technician at Xavier University (Foroozesh Lab, Xavier graduate)
Megan McKay, Undergraduate Students at Xavier University (Foroozesh Lab)
Sydni Bellow, Undergraduate Students at Xavier University (Foroozesh Lab)
Lydia Mensah, Undergraduate Students at Xavier University (Foroozesh Lab)
Amari Chatters, Undergraduate Students at Xavier University (Foroozesh Lab)
Peter Pham, Undergraduate Students at Xavier University (Foroozesh Lab)
Candace Hopgood, Technician at Xavier University (Wiese Lab, Xavier graduate, started Xavier Pharmacy School in August 2013)
Peng Ma, Technician at Xavier University (Wiese Lab)
Chioma Obih, Pharmacy Student at Xavier University (Wiese Lab, graduated in May 2013)
Gabriela Barbarini, Pharmacy Exchange Student at Xavier University (Wiese and Burow Labs)
Felicia Gibson, Pharmacy Student at Xavier University (Wiese Lab)
Elizabeth Martin, Graduate Student at Tulane University (Burow Lab)
Felicia Huynh, Graduate Student at Tulane University (Jones Lab)
Hope Burks, Graduate Student at Tulane University (Burow Lab)
Lucas Chan, Masters Student at Tulane University (Beckman Lab)
Lyndsay Rhodes, Postdoctoral Fellow at Tulane University (Burow Lab)
Melyssa Bratton, Instructor at Tulane University (Burow Lab, currently Research Associate at Xavier University Pharmacy)
Steven Elliott, Lab Supervisor at Tulane University (Burow Lab)
Van Hoang, Graduate Student at Tulane University (Burow Lab)
Han Wen, Graduate Student at Tulane University (Jones Lab)
Mary Sfondouris, Postdoctoral Fellow at Tulane University (Jones Lab)
Thuy-Linh Nguyen, Undergraduate Student at Xavier University (Sridhar lab)
Jasmine Thompson, Undergraduate Student at Xavier University (Sridhar lab)
CONCLUSION:

Foroozesh/Beckman/Burow Subproject

In Year 1, our results have shown that extending the conjugated system in the backbone of ceramide analogs can lead to an increase in the anti-cancer activity. This observation is expected to assist us in designing more potent anti-cancer ceramide analogs.

We have also discovered the conformational isomers of pro-apoptotic ceramide analogs, 401 and 402. This isomerism leads to the possibility that oxazolidine ceramide analogs could act on their molecular targets with two conformations. This finding provides us with a new perspective for the investigation of ceramide-receptor interactions.

In Year 2, we found that the modification of the 1-position of ceramides can lead to novel glucosylceramide synthase (GCS) inhibitors. This finding provides us with a new perspective for the design of effective GCS inhibitors. However, inhibition of GCS is not directly correlated with the ability of ceramide analogs to selectively kill chemotherapy-resistant cancer cell. This indicates that ceramide analogs could inhibit cancer multidrug resistance through two pathways.

In Years 2 and 3, we synthesized 26 novel fluorescent flavone and coumarin derivatives, which are important building blocks of fluorescently visible ceramide analogs. These various fluorescent building blocks include pyranoflavones, furanoflavones, dioxoloflavones, pyridinoﬂavones, pyranocoumarins, furanocoumarins, dioxolocoumarins, and pyridinocoumarins.

Wiese/Burow Subproject
In year one, we developed the methods we use to perform virtual screening of the phytochemical and marketed drug databases. This involved obtaining all crystal structures of the ERalpha ligand binding domain, sorting these structures by ligand type and structure characteristics, and then comparing and optimizing ligand receptor docking protocols. At the same time, we obtained the phytochemical and marketed drug databases, and started the process of filtering for compounds with potential to bind ER that will go into the virtual screening process. Two pharmacy students were trained and then involved in the molecular modeling as well as trained for the in vitro validation phase of the project.

In Year 2, we developed a better way to sort and screen the two molecule databases based on the similarity of the database members to the bound ligands in all available ER LBD crystal structures. To carry out this method, training was obtained and a resorting of the databases is underway. Ligand-receptor docking methods have been evaluated and a standard protocol showing high performance in ER LBDs has been established. One pharmacy student and a new research assistant were trained and are started generating bioassay data. A test set of 29 natural product stilbenes has been obtained and characterized for ER agonist and antagonist activity in a sensitive reporter gene system.

In Year 3 we have completed the dose response characterization of the 8 stilbenes shown to be active in Year 2. This data was presented in a poster at the ACS meeting in Spring 2014. Modeling analysis of these compounds presents ER binding modes consistent with the activity shown. We have identified 5 stilbenes with estrogen agonist activity and 3 stilbenes with antiestrogen activity. Of these antiestrogens 2 may bind ER in the typical direct antagonist mode while one may bind as in indirect antagonist.

**Sridhar/Jones/Stevens Subproject**

Over the years 1 and 2, we were able to identify molecules that target two kinases, namely, PIM1 and CK1d, which play important roles in prostate cancer and breast cancer. Several compounds were found that inhibited MCF7 breast cancer cell line and HER2Δ16. Development of these lead compounds using molecular modeling and organic synthesis will give us potential drug candidates for breast cancer and prostate cancer. The dose response curve studies are ongoing for these two kinases. Based on the results further modification of the lead molecules will be attempted towards the goal of achieving better potency and selectivity for these two kinases. In the meantime, new database searches will be initiated based on the docking studies of known kinase inhibitors on HER2 to identify new core structures as lead molecules with the final goal of finding a new drug candidate for breast cancer.

In year 3 we have performed kinase inhibition assays and toxicology assays for the compounds that showed good inhibition of MCF7/HER2 and MCF7/HER2Δ16 cell lines. Based on the bioactivity data and toxicity data, we have narrowed down the
minimum structural features of the lead quinone moieties. We have synthesized several new derivatives of the lead quinone molecules as potential drug candidates. These will be subjected to inhibition assays and toxicology studies. Additionally, we have found a new lead molecule 1H-indazol-3-ol (indazolone) through computational studies. Two derivatives of the 1H-indazol-3-ol molecules have been synthesized. Further synthetic work is in progress.

**Program**

In addition to the significant amount of scientific research performed and data collected during the first, second, and third years of the Project, it is important to note the valuable partnership developed between the two institutions involved. The productive collaboration formed between the Xavier University and Tulane Cancer Center researchers participating in this program, once again proves the value and importance of inter-institutional research/training projects. The different training activities and the number of trainees involved in the various aspects of the subprojects also positively impact the future cancer research environment in the area. This breast cancer research project is still in its early stages and is expected to develop significantly over the next years.

**REFERENCES:**

**Foroozesh/Beckman/Burow Subproject**


Wiese/Burow Subproject

NA

Sridhar/Jones/Stevens Subproject


