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#### Tandem Cancer Drug Delivery via Gold Nanoparticle Targeting Platforms

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### Introduction

According to the National Cancer Institute, there are several hundred drugs that have been developed with anti-cancer abilities. However, cancer is still the second leading cause of death in the United States according to the CDC. The effectiveness of most anticancer drugs is not limited by its potency but instead by its nontargeted delivery and unwanted side effects. Many cancer treatments such as chemotherapy can effectively kill cancer cells, but lack the selectivity and specificity to avoid wreaking havoc on the rest of the body. Thus, gold nanoparticles (AuNP) have been employed in an effort to overcome some of the downfalls of current chemotherapy options. AuNPs have the unique ability to heat upon exposure to near infrared irradiation and their nanometer sized dimensions aid in their selective delivery to solid tumors. Using AuNPs as a platform, the ability to selectively deliver two different anti-cancer therapeutics via heat sensitive conjugations was investigated. Two DNA strands of different melting temperatures were developed separately as potential temperaturesensitive tethers for drug attachment to AuNPs. Fluorophores were utilized to tag DNA strands during simulation of drug release and track attachment to the AuNP platform. Single-stranded DNA tethers were successfully conjugated to AuNPs by gold-thiol bond formation using a 5' thiol DNA modifier, but synthesizing the full AuNP platform proved to be more challenging. After attempting several attachment sequences and methods, there were no visible indications of successful attachment of dsDNA tethers to the AuNP platform. Future work will consist of identifying what is preventing the full dsDNA tethers from attaching or if its attachment is being masked. Finally, efforts will turn to synthesizing AuNPs with both DNA tethers attached which are able to denature independently at different temperatures to achieve a timed and temperaturedependent dual-drug release.

# General Structure of AuNP Targeting Platform



FAM fluorophore TYE<sup>665</sup> fluorophore Gold-Thiol bond



The AuNP tumor targeting platform consists of two dsDNA strands of different lengths and different melting temperatures (Tm) conjugated to a gold nanoparticle via gold-thiol bonds. Each non-thiolated complementary strand of DNA is tagged with either fluorescein (FAM) fluorophore or TYE665 fluorophore (a modified Cy5 substitute) for tracking temperature-sensitive release. The platform is also stabilized by Polyethylene Glycol (PEG) not shown in this diagram.

# **Tandem Cancer Drug Delivery via Gold Nanoparticle Targeting Platforms** Jeffrey Schulte, Sam Lohmar, Daniel Scott Department of Biochemistry, DePauw University, Greencastle, IN 46135, USA

# **DNA Tether Sequences**

Thiolseq44.2 : 5' - Thiol - ATA AGT CAT CGT ATT GTA TAG -3'

3'TYEcompseq44.2 : 5' - CTA TAC AAT ACG ATG ACT TAT - (TYE) - 3'

5'TYEcompseq44.2 : 5' - (TYE) - CTA TAC AAT ACG ATG ACT TAT - 3'

Two complementary sequences with TYE665 attached at opposite ends were developed in order to investigate the effect of fluorescent quenching

Thiolseq60.7 : 5' - Thiol - TAT GTG GCG TAA GTC CTA AGA GTT -3'

3'FAMcompseq60.7 : 5' - ATA CAC CGC ATT CAG GAT TCT CAA - (FAM) - 3'

5'FAMcompseq60.7 : 5' - (FAM) - ATA CAC CGC ATT CAG GAT TCT CAA - 3'

The FAM tagged dsDNA sequence was developed with a melting temperature significantly higher than the TYE strand (~15°C higher) in order to achieve timed release. This difference arises from its greater length and higher GC content.

# FAM and TYE665 Emission/Excitation Spectra



Wavelength (nm)

## **Conjugation and Hybridization of DNA-AuNPs**

Two approaches were implemented to form the DNA-AuNP platform: 1.) Complementary thiolated and fluorescent ssDNA strands were hybridized together and then conjugated to the AuNP by a gold-thiol bond; and 2.) The thiolated ssDNA strand was conjugated to the AuNP first, followed by hybridization of its complementary fluorescent ssDNA strand.

Unsuccessful hybridization of **Fluorescent DNA** 



Samples of the newly synthesized DNA-AuNP platform did not yield any fluorescence, indicating either unsuccessful hybridization or quenching of the fluorescent DNA by 'laying down' on the AuNP as shown above.

